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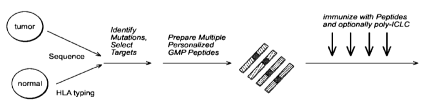


FIG. 1

(57) Abstract: The invention provides a method of making a personalized neoplasia vaccine for a subject diagnosed as having a neoplasia, which includes identifying a plurality of mutations in the neoplasia, analyzing the plurality of mutations to identify a subset of at least five neo-antigenic mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof, and producing, based on the identified subset, a personalized neoplasia vaccine.





COMPOSITIONS AND METHODS FOR PERSONALIZED NEOPLASIA VACCINES

STATEMENT OF RIGHTS TO INVENTIONS MADE UNDER FEDERALLY SPONSORED RESEARCH

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RELATED APPLICATIONS

This application claims the benefit of and priority to U.S. Provisional Patent Application No. 61/809,406, filed April 7, 2013 and U.S. Provisional Patent Application No. 61/869,721, filed August 25, 2013, the contents of which are incorporated herein by reference.

15 FIELD OF THE INVENTION

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The present invention relates to personalized strategies for the treatment of neoplasia. More particularly, the present invention relates to the identification and use of a patient specific pool of tumor specific neo-antigens in a personalized tumor vaccine for treatment of the subject.

20 BACKGROUND

Approximately 1.6 million Americans are diagnosed with neoplasia every year, and approximately 580,000 people in the United States are expected to die of the disease in 2013. Over the past few decades there been significant improvements in the detection, diagnosis, and treatment of neoplasia, which have significantly increased the survival rate for many types of neoplasia. However, only about 60% of people diagnosed with neoplasia are still alive 5 years after the onset of treatment, which makes neoplasia the second leading cause of death in the United States.

Currently, there are a number of different existing cancer therapies, including ablation techniques (e.g., surgical procedures, cryogenic/heat treatment, ultrasound, radiofrequency, and radiation) and chemical techniques (e.g., pharmaceutical agents, cytotoxic/chemotherapeutic agents, monoclonal antibodies, and various combinations thereof). Unfortunately, such therapies are frequently associated with serious risk, toxic side effects, and extremely high costs, as well as

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uncertain efficacy.

There is a growing interest in cancer therapies that seek to target cancerous cells with a patient's own immune system (e.g., cancer vaccines) because such therapies may mitigate/eliminate some of the above-described disadvantages. Cancer vaccines are typically composed of tumor antigens and immunostimulatory molecules (e.g., cytokines or TLR ligands) that work together to induce antigen-specific cytotoxic T cells that target and destroy tumor cells. Current cancer vaccines typically contain shared tumor antigens, which are native proteins (i.e. – proteins encoded by the DNA of all the normal cells in the individual) that are selectively expressed or over-expressed in tumors found in many individuals. While such shared tumor antigens are useful in identifying particular types of tumors, they are not ideal as immunogens for targeting a T-cell response to a particular tumor type because they are subject to the immune dampening effects of self-tolerance. Accordingly, there is a need for methods of identifying more effective tumor antigens that may be used for neoplasia vaccines.

SUMMARY OF THE INVENTION

The present invention relates to a strategy for the personalized treatment of neoplasia, and more particularly to the identification and use of a personalized cancer vaccine consisting essentially of a pool of tumor-specific and patient-specific neo-antigens for the treatment of tumors in a subject. As described below, the present invention is based, at least in part, on the discovery that whole genome/exome sequencing may be used to identify all, or nearly all, mutated neo-antigens that are uniquely present in a neoplasia/tumor of an individual patient, and that this collection of mutated neo-antigens may be analyzed to identify a specific, optimized subset of neo-antigens for use as a personalized neoplasia vaccine for treatment of the patient's neoplasia/tumor.

In one aspect, the invention provides a method of making a personalized neoplasia vaccine for a subject diagnosed as having a neoplasia, which includes identifying a plurality of sequences comprising mutations in the neoplasia, analyzing the plurality of sequences comprising mutations to identify a subset of at least five neo-antigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof, said analyzing comprising

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identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising: (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation; (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≥ 1000 nM; (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM; (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≥ 500 nM encoded by a sequence comprising a neoORF mutation; (v) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a neoORF mutation; (v) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM; and producing, based on the identified subset, a personalized neoplasia vaccine.

In a further aspect, the present invention provides a personalized neoplasia vaccine when used in a method for treating a subject diagnosed as having a neoplasia, said method comprising: identifying a plurality of sequences comprising mutations in the neoplasia, analyzing the plurality of sequences comprising mutations to identify a subset of at least five neo-antigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising: (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation; (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≥ 1000 nM; (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM; (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation; (v) a polypeptide that binds to

the HLA of the subject with a Kd of 150- ≤ 500 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of 150- ≤ 500 nM; producing, based on the identified subset, a personalized neoplasia vaccine; and administering the personalized neoplasia vaccine to the subject, thereby treating the neoplasia.

In an embodiment, the invention provides that the identifying step further includes sequencing the genome, transcriptome, or proteome of the neoplasia.

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In another embodiment, the analyzing step may further include determining one or more characteristics associated with the subset of at least five neo-antigenic mutations predicted to encode neo-antigenic peptides, the characteristics selected from the group consisting of molecular weight, cysteine content, hydrophilicity, hydrophobicity, charge, and binding affinity; and ranking, based on the determined characteristics, each of the neo-antigenic mutations within the identified subset of at least five neo-antigenic mutations. In an embodiment, the top 5-30 ranked neo-antigenic mutations are included in the personalized neoplasia vaccine. In another embodiment, the neo-antigenic mutations are ranked according to the order shown in FIG. 8.

In one embodiment, the personalized neoplasia vaccine comprises at least about 20 neoantigenic peptides corresponding to the neo-antigenic mutations.

In another embodiment, the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least about 20 neo-antigenic peptides corresponding to the neo-antigenic mutations. In another embodiment, the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 20 neo-antigenic peptides corresponding to the neo-antigenic mutations.

In embodiments, the personalized neoplasia vaccine comprises neoORF mutations predicted to encode a neoORF polypeptide having a Kd of ≤ 500 nM.

In another embodiment, the personalized neoplasia vaccine comprises missense mutations predicted to encode a polypeptide having a Kd of \leq 150 nM, wherein the native cognate protein has a Kd of \geq 1000 nM or \leq 150 nM.

In another embodiment, the at least about 20 neo-antigenic peptides range from about 5 to about 50 amino acids in length. In another embodiment, the at least about 20 neo-antigenic peptides range from about 15 to about 35 amino acids in length. In another embodiment, the at least about 20 neo-antigenic peptides range from about 18 to about 30 amino acids in length. In another embodiment, the at least about 20 neo-antigenic peptides range from about 6 to about 15 amino acids in length. In yet another embodiment, the at least about 20 neo-antigenic peptides are 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 amino acids in length.

In one embodiment, the personalized neoplasia vaccine further includes an adjuvant. In other embodiments, the adjuvant is selected from the group consisting of poly-ICLC, 1018 ISS,

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aluminum salts, Amplivax, AS15, BCG, CP-870,893, CpG7909, CyaA, dSLIM, GM-CSF, IC30, IC31, Imiquimod, ImuFact IMP321, IS Patch, ISS, ISCOMATRIX, Juvlmmune, LipoVac, MF59, monophosphoryl lipid A, Montanide IMS 1312, Montanide ISA 206, Montanide ISA 50V, Montanide ISA-51, OK-432, OM-174, OM-197-MP-EC, ONTAK, PepTel.RTM, vector system, PLGA microparticles, resiquimod, SRL172, Virosomes and other Virus-like particles, YF-17D, VEGF trap, R848, beta-glucan, Pam3Cys, Aquila's QS21 stimulon, vadimezan, and/or AsA404 (DMXAA). In a preferred embodiment, the adjuvant is poly-ICLC.

In another aspect, the invention includes a method of treating a subject diagnosed as having a neoplasia with a personalized neoplasia vaccine, comprising: identifying a plurality of sequences comprising mutations in the neoplasia; analyzing the plurality of sequences comprising mutations to identify a subset of at least five neo-antigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and ranking the neoantigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising: (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation; (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≥ 1000 nM; (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM; (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation, (v) a polypeptide that binds to the HLA of the subject with a Kd of 150- \leq 500 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of 150- \leq 500 nM; producing, based on the identified subset, a personalized neoplasia vaccine; and administering the personalized neoplasia vaccine to the subject, thereby treating the neoplasia.

In another embodiment, the identifying step may further include sequencing the genome, transcriptome, or proteome of the neoplasia.

In yet another embodiment, the analyzing step may further include determining one or

more characteristics associated with the subset of at least five neo-antigenic mutations predicted to encode expressed neo-antigenic peptides, the characteristics selected from the group consisting of molecular weight, cysteine content, hydrophilicity, hydrophobicity charge, and binding affinity; and ranking, based on the determined characteristics, each of the neo-antigenic mutations within the identified subset of at least five neo-antigenic mutations.

In one embodiment, the top 5-30 ranked neo-antigenic mutations are included in the personalized neoplasia vaccine. In another embodiment, the neo-antigenic mutations are ranked according to the order shown in FIG. 8.

In one embodiment, the personalized neoplasia vaccine comprises at least 20 neoantigenic peptides corresponding to the neo-antigenic mutations.

In another embodiment, the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least 20 neo-antigenic peptides corresponding to the neoantigenic mutations.

In one embodiment, the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 20 neo-antigenic peptides corresponding to the neo-antigenic mutations.

In one embodiment, the personalized neoplasia vaccine comprises neoORF mutations predicted to encode a neoORF polypeptide having a Kd of ≤ 500 nM.

In another embodiment, the personalized neoplasia vaccine comprises missense mutations predicted to encode a polypeptide having a Kd of \leq 150 nM, wherein the native cognate protein has a Kd of \geq 1000 nM or \leq 150 nM.

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In one embodiment, the at least 20 neo-antigenic peptides range from about 5 to about 50 amino acids in length. In one embodiment, the at least 20 neo-antigenic peptides range from about 15 to about 35 amino acids in length. In one embodiment, the at least 20 neo-antigenic peptides range from about 18 to about 30 amino acids in length. In one embodiment, the at least 20 neo-antigenic peptides range from about 6 to about 15 amino acids in length. In one embodiment, the at least 20 neo-antigenic peptides are 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 amino acids in length.

In one embodiment, the administering further includes dividing the produced vaccine into two or more sub-pools; and injecting each of the sub-pools into a different location of the patient. In one embodiment, each of the sub-pools injected into a different location comprises neo-antigenic peptides such that the number of individual peptides in the sub-pool targeting any single patient HLA is one, or as few above one as possible.

In one embodiment, the administering step further includes dividing the produced vaccine into two or more sub-pools, wherein each sub-pool comprises at least five neo-antigenic peptides selected to optimize intra-pool interactions.

In one embodiment, optimizing comprises reducing negative interaction among the neoantigenic peptides in the same pool.

In another aspect, the invention includes a personalize neoplasia vaccine prepared according to the above-described methods.

Definitions

To facilitate an understanding of the present invention, a number of terms and phrases are defined below:

Unless specifically stated or obvious from context, as used herein, the term "about" is understood as within a range of normal tolerance in the art, for example within 2 standard deviations of the mean. About can be understood as within 50%, 45%, 40%, 35%, 30%, 25%, 20%, 15%, 10%, 9%, 8%, 7%, 6%, 5%, 4%, 3%, 2%, 1%, 0.5%, 0.1%, 0.05%, or 0.01% of the stated value. Unless otherwise clear from context, all numerical values provided herein are modified by the term about.

By "agent" is meant any small molecule chemical compound, antibody, nucleic acid molecule, or polypeptide, or fragments thereof.

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By "ameliorate" is meant decrease, suppress, attenuate, diminish, arrest, or stabilize the development or progression of a disease (e.g., a neoplasia, tumor, etc.).

By "alteration" is meant a change (increase or decrease) in the expression levels or activity of a gene or polypeptide as detected by standard art known methods such as those described herein. As used herein, an alteration includes a 10% change in expression levels, preferably a 25% change, more preferably a 40% change, and most preferably a 50% or greater change in expression levels.

By "analog" is meant a molecule that is not identical, but has analogous functional or structural features. For example, a tumor specific neo-antigen polypeptide analog retains the biological activity of a corresponding naturally-occurring tumor specific neo-antigen polypeptide, while having certain biochemical modifications that enhance the analog's function relative to a naturally-occurring polypeptide. Such biochemical modifications could increase the analog's protease resistance, membrane permeability, or half-life, without altering, for example, ligand binding. An analog may include an unnatural amino acid.

The phrase "combination therapy" embraces the administration of a pooled sample of neoplasia/tumor specific neo-antigens and one or more additional therapeutic agents as part of a specific treatment regimen intended to provide a beneficial (additive or synergistic) effect from the co-action of these therapeutic agents. The beneficial effect of the combination includes, but is not limited to, pharmacokinetic or pharmacodynamic co-action resulting from the combination of therapeutic agents. Administration of these therapeutic agents in combination typically is carried out over a defined time period (usually minutes, hours, days, or weeks depending upon the combination selected). "Combination therapy" is intended to embrace administration of these therapeutic agents in a sequential manner, that is, wherein each therapeutic agent is

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administered at a different time, as well as administration of these therapeutic agents, or at least two of the therapeutic agents, in a substantially simultaneous manner. Substantially simultaneous administration can be accomplished, for example, by administering to the subject a single capsule having a fixed ratio of each therapeutic agent or in multiple, single capsules for each of the therapeutic agents. For example, one combination of the present invention may comprise a pooled sample of tumor specific neo-antigens and at least one additional therapeutic agent (e.g., a chemotherapeutic agent, an anti-angiogenesis agent, an immunosuppressive agent, an anti-inflammatory agent, and the like) at the same or different times or they can be formulated as a single, co-formulated pharmaceutical composition comprising the two compounds. As another example, a combination of the present invention (e.g., a pooled sample of tumor specific neo-antigens and at least one additional therapeutic agent) may be formulated as separate pharmaceutical compositions that can be administered at the same or different time. Sequential or substantially simultaneous administration of each therapeutic agent can be effected by any appropriate route including, but not limited to, oral routes, intravenous routes, sub-cutaneous routes, intramuscular routes, direct absorption through mucous membrane tissues (e.g., nasal, mouth, vaginal, and rectal), and ocular routes (e.g., intravitreal, intraocular, etc.). The therapeutic agents can be administered by the same route or by different routes. For example, one component of a particular combination may be administered by intravenous injection while the other component(s) of the combination may be administered orally. The components may be administered in any therapeutically effective sequence.

The phrase "combination" embraces groups of compounds or non-drug therapies useful as part of a combination therapy.

Where any or all of the terms "comprise", "comprises", "comprised" or "comprising" are used in this specification (including the claims) they are to be interpreted as specifying the presence of the stated features, integers, steps or components, but not precluding the presence of one or more other features, integers, steps or components.

The discussion of documents, acts, materials, devices, articles and the like is included in this specification solely for the purpose of providing a context for the present invention. It is not suggested or represented that any or all of these matters formed part of the prior art base or were common general knowledge in the field relevant to the present invention as it existed before the priority date of each claim of this application.

By "control" is meant a standard or reference condition.

By "disease" is meant any condition or disorder that damages or interferes with the normal function of a cell, tissue, or organ.

By "effective amount" is meant the amount required to ameliorate the symptoms of a disease (e.g., a neoplasia/tumor) relative to an untreated patient. The effective amount of active compound(s) used to practice the present invention for therapeutic treatment of a disease varies depending upon the manner of administration, the age, body weight, and general health of the subject. Ultimately, the attending physician or veterinarian will decide the appropriate amount and dosage regimen. Such amount is referred to as an "effective" amount.

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By "fragment" is meant a portion of a polypeptide or nucleic acid molecule. This portion contains, preferably, at least 5%, 10%, 20%, 30%, 40%, 50%, 60%, 70%, 80%, or 90% of the entire length of the reference nucleic acid molecule or polypeptide. A fragment may contain 5, 10, 20, 30, 40, 50, 60, 70, 80, 90, or 100, 200, 300, 400, 500, 600, 700, 800, 900, 1000 or more nucleotides or amino acids.

"Hybridization" means hydrogen bonding, which may be Watson-Crick, Hoogsteen or reversed Hoogsteen hydrogen bonding, between complementary nucleobases. For example, adenine and thymine are complementary nucleobases that pair through the formation of hydrogen bonds.

By "inhibitory nucleic acid" is meant a double-stranded RNA, siRNA, shRNA, or antisense RNA, or a portion thereof, or a mimetic thereof, that when administered to a mammalian cell results in a decrease (e.g., by 10%, 25%, 50%, 75%, or even 90-100%) in the expression of a target gene. Typically, a nucleic acid inhibitor comprises at least a portion of a target nucleic acid molecule, or an ortholog thereof, or comprises at least a portion of the complementary strand of a target nucleic acid molecule. For example, an inhibitory nucleic acid molecule comprises at least a portion of any or all of the nucleic acids delineated herein.

By "isolated polynucleotide" is meant a nucleic acid (*e.g.*, a DNA) that is free of the genes which, in the naturally-occurring genome of the organism—or in the genomic DNA of a neoplasia/tumor derived from the organism—the nucleic acid molecule of the invention is derived. The term therefore includes, for example, a recombinant DNA (*e.g.*, DNA coding for a neoORF, read-through, or InDel derived polypeptide identified in a patient's tumor) that is incorporated into a vector; into an autonomously replicating plasmid or virus; or into the genomic DNA of a prokaryote or eukaryote; or that exists as a separate molecule (for example, a cDNA or a genomic or cDNA fragment produced by PCR or restriction endonuclease digestion) independent of other sequences. In addition, the term includes an RNA molecule that is

transcribed from a DNA molecule, as well as a recombinant DNA that is part of a hybrid gene encoding additional polypeptide sequence.

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By an "isolated polypeptide" is meant a polypeptide of the invention that has been separated from components that naturally accompany it. Typically, the polypeptide is isolated when it is at least 60%, by weight, free from the proteins and naturally-occurring organic molecules with which it is naturally associated. Preferably, the preparation is at least 75%, more preferably at least 90%, and most preferably at least 99%, by weight, a polypeptide of the invention. An isolated polypeptide of the invention may be obtained, for example, by extraction from a natural source, by expression of a recombinant nucleic acid encoding such a polypeptide; or by chemically synthesizing the protein. Purity can be measured by any appropriate method, for example, column chromatography, polyacrylamide gel electrophoresis, or by HPLC analysis.

A "ligand" is to be understood as meaning a molecule which has a structure complementary to that of a receptor and is capable of forming a complex with the receptor. According to the invention, a ligand is to be understood as meaning a peptide or peptide fragment that has a suitable length and suitable binding motifs in its amino acid sequence, so that the peptide or peptide fragment is capable of forming a complex with proteins of MHC class I or MHC class II.

"Mutation" for the purposes of this document means a DNA sequence found in the tumor DNA sample of a patient that is not found in the corresponding normal DNA sample of that same patient. "Mutation" may also refer to patterns in the sequence of RNA from a patient that are not attributable to expected variations based on known information for an individual gene and are reasonably considered to be novel variations in, for example, the splicing pattern of one or more genes that has been specifically altered in the tumor cells of the patient.

"Neo-antigen" or "neo-antigenic" means a class of tumor antigens that arises from a tumor-specific mutation(s) which alters the amino acid sequence of genome encoded proteins.

By "neoplasia" is meant any disease that is caused by or results in inappropriately high levels of cell division, inappropriately low levels of apoptosis, or both. For example, cancer is an example of a neoplasia. Examples of cancers include, without limitation, leukemia (e.g., acute leukemia, acute lymphocytic leukemia, acute myelocytic leukemia, acute myeloblastic leukemia, acute promyelocytic leukemia, acute myelomonocytic leukemia, acute monocytic leukemia, acute erythroleukemia, chronic leukemia, chronic myelocytic leukemia, chronic lymphocytic

leukemia), polycythemia vera, lymphoma (e.g., Hodgkin's disease, non-Hodgkin's disease), Waldenstrom's macroglobulinemia, heavy chain disease, and solid tumors such as sarcomas and carcinomas (e.g., fibrosarcoma, myxosarcoma, liposarcoma, chondrosarcoma, osteogenic sarcoma, chordoma, angiosarcoma, endotheliosarcoma, lymphangiosarcoma,

lymphangioendotheliosarcoma, synovioma, mesothelioma, Ewing's tumor, leiomyosarcoma, rhabdomyosarcoma, colon carcinoma, pancreatic cancer, breast cancer, ovarian cancer, prostate cancer, squamous cell carcinoma, basal cell carcinoma, adenocarcinoma, sweat gland carcinoma, sebaceous gland carcinoma, papillary carcinoma, papillary adenocarcinomas, cystadenocarcinoma, medullary carcinoma, bronchogenic carcinoma, renal cell carcinoma, hepatoma, nile duct carcinoma, choriocarcinoma, seminoma, embryonal carcinoma, Wilm's tumor, cervical cancer, uterine cancer, testicular cancer, lung carcinoma, small cell lung carcinoma, bladder carcinoma, epithelial carcinoma, glioma, astrocytoma, medulloblastoma, craniopharyngioma, ependymoma, pinealoma, hemangioblastoma, acoustic neuroma,

oligodenroglioma, schwannoma, meningioma, melanoma, neuroblastoma, and retinoblastoma).

Lymphoproliferative disorders are also considered to be proliferative diseases.

Unless specifically stated or obvious from context, as used herein, the term "or" is

understood to be inclusive. Unless specifically stated or obvious from context, as used herein,

the terms "a," "an," and "the" are understood to be singular or plural.

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The term "patient" or "subject" refers to an animal which is the object of treatment, observation, or experiment. By way of example only, a subject includes, but is not limited to, a mammal, including, but not limited to, a human or a non-human mammal, such as a non-human primate, bovine, equine, canine, ovine, or feline.

"Pharmaceutically acceptable" refers to approved or approvable by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopeia or other generally recognized pharmacopeia for use in animals, including humans.

"Pharmaceutically acceptable excipient, carrier or diluent" refers to an excipient, carrier or diluent that can be administered to a subject, together with an agent, and which does not destroy the pharmacological activity thereof and is nontoxic when administered in doses sufficient to deliver a therapeutic amount of the agent.

A "pharmaceutically acceptable salt" of pooled tumor specific neo-antigens as recited herein may be an acid or base salt that is generally considered in the art to be suitable for use in contact with the tissues of human beings or animals without excessive toxicity, irritation, allergic response, or other problem or complication. Such salts include mineral and organic acid salts of basic residues such as amines, as well as alkali or organic salts of acidic residues such as carboxylic acids. Specific pharmaceutical salts include, but are not limited to, salts of acids such as hydrochloric, phosphoric, hydrobromic, malic, glycolic, fumaric, sulfuric, sulfamic, sulfamilic, formic, toluenesulfonic, methanesulfonic, benzene sulfonic, ethane disulfonic, 2hydroxyethylsulfonic, nitric, benzoic, 2-acetoxybenzoic, citric, tartaric, lactic, stearic, salicylic, glutamic, ascorbic, pamoic, succinic, fumaric, maleic, propionic, hydroxymaleic, hydroiodic, phenylacetic, alkanoic such as acetic, HOOC-(CH₂)_n-COOH where n is 0-4, and the like. Similarly, pharmaceutically acceptable cations include, but are not limited to sodium, potassium, calcium, aluminum, lithium and ammonium. Those of ordinary skill in the art will recognize further pharmaceutically acceptable salts for the pooled tumor specific neo-antigens provided herein, including those listed by Remington's Pharmaceutical Sciences, 17th ed., Mack Publishing Company, Easton, PA, p. 1418 (1985). In general, a pharmaceutically acceptable acid or base salt can be synthesized from a parent compound that contains a basic or acidic moiety by any conventional chemical method. Briefly, such salts can be prepared by reacting the free acid or base forms of these compounds with a stoichiometric amount of the appropriate base or acid in an appropriate solvent.

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As used herein, the terms "prevent," "preventing," "prevention," "prophylactic treatment," and the like, refer to reducing the probability of developing a disease or condition in a subject, who does not have, but is at risk of or susceptible to developing a disease or condition.

"Primer set" means a set of oligonucleotides that may be used, for example, for PCR. A primer set would consist of at least 2, 4, 6, 8, 10, 12, 14, 16, 18, 20, 30, 40, 50, 60, 80, 100, 200, 250, 300, 400, 500, 600, or more primers.

"Proteins or molecules of the major histocompatibility complex (MHC)," "MHC molecules," "MHC proteins" or "HLA proteins" are to be understood as meaning, in particular, proteins capable of binding peptides resulting from the proteolytic cleavage of protein antigens and representing potential T-cell epitopes, transporting them to the cell surface and presenting

them to specific cells there, in particular naïve T-cells, cytotoxic T-lymphocytes or T-helper cells. The major histocompatibility complex in the genome comprises the genetic region whose gene products are expressed on the cell surface and are important for binding and presenting endogenous and/or foreign antigens, and thus for regulating immunological processes. The major histocompatibility complex is classified into two gene groups coding for different proteins: molecules of MHC class I and MHC class II. The molecules of the two MHC classes are specialized for different antigen sources. The molecules of MHC class I typically present but are not restricted to endogenously synthesized antigens, for example viral proteins and tumor antigens. The molecules of MHC class II present protein antigens originating from exogenous sources, for example bacterial products. The cellular biology and the expression patterns of the two MHC classes are adapted to these different roles.

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MHC molecules of class I consist of a heavy chain and a light chain and are capable of binding a peptide of about 8 to 11 amino acids, but usually 9 or 10 amino acids, if this peptide has suitable binding motifs, and presenting it to naïve and cytotoxic T- lymphocytes. The peptide bound by the MHC molecules of class I typically but not exclusively originates from an endogenous protein antigen. The heavy chain of the MHC molecules of class I is preferably an HLA-A, HLA-B or HLA-C monomer, and the light chain is β-2-microglobulin.

MHC molecules of class II consist of an α -chain and a β -chain and are capable of binding a peptide of about 15 to 24 amino acids if this peptide has suitable binding motifs, and presenting it to T-helper cells. The peptide bound by the MHC molecules of class II usually originates from an extracellular or exogenous protein antigen. The α -chain and the β -chain are in particular HLA-DR, HLA-DQ and HLA-DP monomers.

Ranges provided herein are understood to be shorthand for all of the values within the range. For example, a range of 1 to 50 is understood to include any number, combination of numbers, or sub-range from the group consisting of 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32, 33, 34, 35, 36, 37, 38, 39, 40, 41, 42, 43, 44, 45, 46, 47, 48, 49, or 50, as well as all intervening decimal values between the aforementioned integers such as, for example, 1.1, 1.2, 1.3, 1.4, 1.5, 1.6, 1.7, 1.8, and 1.9. With respect to sub-ranges, "nested sub-ranges" that extend from either end point of the range are specifically contemplated. For example, a nested sub-range of an exemplary range of 1 to 50

may comprise 1 to 10, 1 to 20, 1 to 30, and 1 to 40 in one direction, or 50 to 40, 50 to 30, 50 to 20, and 50 to 10 in the other direction.

A "receptor" is to be understood as meaning a biological molecule or a molecule grouping capable of binding a ligand. A receptor may serve, to transmit information in a cell, a cell formation or an organism. The receptor comprises at least one receptor unit and frequently contains two or more receptor units, where each receptor unit may consist of a protein molecule, in particular a glycoprotein molecule. The receptor has a structure that complements the structure of a ligand and may complex the ligand as a binding partner. Signaling information may be transmitted by conformational changes of the receptor following binding with the ligand on the surface of a cell. According to the invention, a receptor may refer to particular proteins of MHC classes I and II capable of forming a receptor/ligand complex with a ligand, in particular a peptide or peptide fragment of suitable length.

A "receptor/ligand complex" is also to be understood as meaning a "receptor/peptide complex" or "receptor/peptide fragment complex," in particular a peptide- or peptide fragment-presenting MHC molecule of class I or of class II.

By "reduces" is meant a negative alteration of at least 10%, 25%, 50%, 75%, or 100%.

By "reference" is meant a standard or control condition.

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A "reference sequence" is a defined sequence used as a basis for sequence comparison. A reference sequence may be a subset of, or the entirety of, a specified sequence; for example, a segment of a full-length cDNA or genomic sequence, or the complete cDNA or genomic sequence. For polypeptides, the length of the reference polypeptide sequence will generally be at least about 10-2,000 amino acids, 10-1,500, 10-1,000, 10-500, or 10-100. Preferably, the length of the reference polypeptide sequence may be at least about 10-50 amino acids, more preferably at least about 10-40 amino acids, and even more preferably about 10-30 amino acids, about 10-20 amino acids, about 15-25 amino acids, or about 20 amino acids. For nucleic acids, the length of the reference nucleic acid sequence will generally be at least about 50 nucleotides, preferably at least about 60 nucleotides, more preferably at least about 75 nucleotides, and even more preferably about 100 nucleotides or about 300 nucleotides or any integer thereabout or there between.

By "specifically binds" is meant a compound or antibody that recognizes and binds a polypeptide of the invention, but which does not substantially recognize and bind other molecules in a sample, for example, a biological sample.

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Nucleic acid molecules useful in the methods of the invention include any nucleic acid molecule that encodes a polypeptide of the invention or a fragment thereof. Such nucleic acid molecules need not be 100% identical with an endogenous nucleic acid sequence, but will typically exhibit substantial identity. Polynucleotides having "substantial identity" to an endogenous sequence are typically capable of hybridizing with at least one strand of a double-stranded nucleic acid molecule. By "hybridize" is meant pair to form a double-stranded molecule between complementary polynucleotide sequences (e.g., a gene described herein), or portions thereof, under various conditions of stringency. (See, e.g., Wahl, G. M. and S. L. Berger (1987) Methods Enzymol. 152:399; Kimmel, A. R. (1987) Methods Enzymol. 152:507).

For example, stringent salt concentration will ordinarily be less than about 750 mM NaCl and 75 mM trisodium citrate, preferably less than about 500 mM NaCl and 50 mM trisodium citrate, and more preferably less than about 250 mM NaCl and 25 mM trisodium citrate. Low stringency hybridization can be obtained in the absence of organic solvent, e.g., formamide, while high stringency hybridization can be obtained in the presence of at least about 35% formamide, and more preferably at least about 50% formamide. Stringent temperature conditions will ordinarily include temperatures of at least about 30°C, more preferably of at least about 37°C, and most preferably of at least about 42°C. Varying additional parameters, such as hybridization time, the concentration of detergent, e.g., sodium dodecyl sulfate (SDS), and the inclusion or exclusion of carrier DNA, are well known to those skilled in the art. Various levels of stringency are accomplished by combining these various conditions as needed. In a preferred: embodiment, hybridization will occur at 30°C in 750 mM NaCl, 75 mM trisodium citrate, and 1% SDS. In a more preferred embodiment, hybridization will occur at 37° C in 500 mM NaCl, 50 mM trisodium citrate, 1% SDS, 35% formamide, and 100 µg/ml denatured salmon sperm DNA (ssDNA). In a most preferred embodiment, hybridization will occur at 42°C in 250 mM NaCl, 25 mM trisodium citrate, 1% SDS, 50% formamide, and 200 μg/ml ssDNA. Useful variations on these conditions will be readily apparent to those skilled in the art.

For most applications, washing steps that follow hybridization will also vary in stringency. Wash stringency conditions can be defined by salt concentration and by temperature.

As above, wash stringency can be increased by decreasing salt concentration or by increasing temperature. For example, stringent salt concentration for the wash steps will preferably be less than about 30 mM NaCl and 3 mM trisodium citrate, and most preferably less than about 15 mM NaCl and 1.5 mM trisodium citrate. Stringent temperature conditions for the wash steps will ordinarily include a temperature of at least about 25°C, more preferably of at least about 42°C, and even more preferably of at least about 68°C. In a preferred embodiment, wash steps will occur at 25°C in 30 mM NaCl, 3 mM trisodium citrate, and 0.1% SDS. In a more preferred embodiment, wash steps will occur at 42°C in 15 mM NaCl, 1.5 mM trisodium citrate, and 0.1% SDS. In a more preferred embodiment, wash steps will occur at 68°C in 15 mM NaCl, 1.5 mM trisodium citrate, and 0.1% SDS. Additional variations on these conditions will be readily apparent to those skilled in the art. Hybridization techniques are well known to those skilled in the art and are described, for example, in Benton and Davis (Science 196:180, 1977); Grunstein and Hogness (Proc. Natl. Acad. Sci., USA 72:3961, 1975); Ausubel et al. (Current Protocols in Molecular Biology, Wiley Interscience, New York, 2001); Berger and Kimmel (Guide to Molecular Cloning Techniques, 1987, Academic Press, New York); and Sambrook et al., Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory Press, New York.

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By "substantially identical" is meant a polypeptide or nucleic acid molecule exhibiting at least 50% identity to a reference amino acid sequence (for example, any one of the amino acid sequences described herein) or nucleic acid sequence (for example, any one of the nucleic acid sequences described herein). Preferably, such a sequence is at least 60%, more preferably 80% or 85%, and more preferably 90%, 95% or even 99% identical at the amino acid level or nucleic acid to the sequence used for comparison.

Sequence identity is typically measured using sequence analysis software (for example, Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin Biotechnology Center, 1710 University Avenue, Madison, Wis. 53705, BLAST, BESTFIT, GAP, or PILEUP/PRETTYBOX programs). Such software matches identical or similar sequences by assigning degrees of homology to various substitutions, deletions, and/or other modifications. Conservative substitutions typically include substitutions within the following groups: glycine, alanine; valine, isoleucine, leucine; aspartic acid, glutamic acid, asparagine, glutamine; serine, threonine; lysine, arginine; and phenylalanine, tyrosine. In an exemplary

approach to determining the degree of identity, a BLAST program may be used, with a probability score between e^{-3} and e^{-100} indicating a closely related sequence.

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A "T-cell epitope" is to be understood as meaning a peptide sequence that can be bound by MHC molecules of class I or II in the form of a peptide-presenting MHC molecule or MHC complex and then, in this form, be recognized and bound by naïve T-cells, cytotoxic T-lymphocytes or T-helper cells.

As used herein, the terms "treat," "treated," "treating," "treatment," and the like refer to reducing or ameliorating a disorder and/or symptoms associated therewith (e.g., a neoplasia or tumor). It will be appreciated that, although not precluded, treating a disorder or condition does not require that the disorder, condition, or symptoms associated therewith be completely eliminated.

The term "therapeutic effect" refers to some extent of relief of one or more of the symptoms of a disorder (e.g., a neoplasia or tumor) or its associated pathology. "Therapeutically effective amount" as used herein refers to an amount of an agent which is effective, upon single or multiple dose administration to the cell or subject, in prolonging the survivability of the patient with such a disorder, reducing one or more signs or symptoms of the disorder, preventing or delaying, and the like beyond that expected in the absence of such treatment. "Therapeutically effective amount" is intended to qualify the amount required to achieve a therapeutic effect. A physician or veterinarian having ordinary skill in the art can readily determine and prescribe the "therapeutically effective amount" (e.g., ED50) of the pharmaceutical composition required. For example, the physician or veterinarian could start doses of the compounds of the invention employed in a pharmaceutical composition at levels lower than that required in order to achieve the desired therapeutic effect and gradually increase the dosage until the desired effect is achieved.

The pharmaceutical compositions typically should provide a dosage of from about 0.0001 mg to about 200 mg of compound per kilogram of body weight per day. For example, dosages for systemic administration to a human patient can range from 0.01-10 μg/kg, 20-80 μg/kg, 5-50 μg/kg, 75-150 μg/kg, 100-500 μg/kg, 250-750 μg/kg, 500-1000 μg/kg, 1-10 mg/kg, 5-50 mg/kg, 25-75 mg/kg, 50-100 mg/kg, 100-250 mg/kg, 50-100 mg/kg, 250-500 mg/kg, 500-750 mg/kg, 750-1000 mg/kg, 1000-1500 mg/kg, 1500-2000 mg/kg, 5 mg/kg, 20 mg/kg, 50 mg/kg, 100 mg/kg, of 200 mg/kg. Pharmaceutical dosage unit forms are prepared to provide from about

0.001 mg to about 5000 mg, for example from about 100 to about 2500 mg of the compound or a combination of essential ingredients per dosage unit form.

A "vaccine" is to be understood as meaning a composition for generating immunity for the prophylaxis and/or treatment of diseases (e.g., neoplasia/tumor). Accordingly, vaccines are medicaments which comprise antigens and are intended to be used in humans or animals for generating specific defense and protective substance by vaccination.

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The recitation of a listing of chemical groups in any definition of a variable herein includes definitions of that variable as any single group or combination of listed groups. The recitation of an embodiment for a variable or aspect herein includes that embodiment as any single embodiment or in combination with any other embodiments or portions thereof.

Any compositions or methods provided herein can be combined with one or more of any of the other compositions and methods provided herein.

BRIEF DESCRIPTION OF THE DRAWINGS

The above-mentioned and other features and advantages of the present disclosure will be better understood when reading the following detailed description taken together with the following drawings in which:

Figure 1 depicts a flow process for making a personalized cancer vaccine according to an exemplary embodiment of the invention.

Figure 2 shows a flow process for pre-treatment steps for generating a cancer vaccine for a melanoma patient according to an exemplary embodiment of the invention.

Figure 3 is a flowchart depicting an approach for addressing an initial patient population study according to an exemplary embodiment of the invention. Five patients may be treated in the first cohort at an anticipated safe dose level. If fewer than two of these five patients develop a dose limiting toxicity at, or prior to, the primary safety endpoint, then 10 more patients may be recruited at that dose level to expand the analysis of the patient population (e.g., to assess efficacy, safety, etc.). If two or more dose limiting toxicities (DLTs) are observed, then the dose of poly-ICLC may be reduced by 50% and five additional patients may be treated. If fewer than two of these five patients develop a dose limiting toxicity, then 10 more patients may be recruited at that dose level. However, if two or more patients at the reduced poly-ICLC level develop a DLT, then the study will be stopped.

Figures 4A and 4B show examples of different types of discrete mutations and neoORFs, respectively.

Figure 5 illustrates an immunization schedule based on a prime boost strategy according to an exemplary embodiment of the present invention. Multiple immunizations may occur over the first ~3 weeks to maintain an early high antigen exposure during the priming phase of immune response. Patients may then be rested for eight weeks to allow memory T cells to develop and these T cells will then be boosted in order to maintain a strong ongoing response.

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Figure 6 shows a time line indicating the primary immunological endpoint according to an exemplary aspect of the invention.

Figure 7 illustrates a time line for administering a co-therapy with checkpoint blockade antibodies to evaluate the combination of relief of local immune suppression coupled with the stimulation of new immunity according to an exemplary embodiment of the invention. As shown in the scheme, patients who enter as appropriate candidates for checkpoint blockade therapy, e.g., anti-PDL1 as shown here, may be entered and immediately treated with antibody, while the vaccine is being prepared. Patients may then be vaccinated. Checkpoint blockade antibody dosing can be continued or possibly deferred while the priming phase of vaccination occurs.

Figure 8 is a table that shows the ranking assignments for different neo-antigenic mutations according to an exemplary embodiment of the invention.

Figure 9 shows a schematic depicting drug product processing of individual neoantigenic peptides into pools of 4 subgroups according to an exemplary embodiment of the invention.

Figure 10 shows a schematic representation of a strategy to systematically discover tumor neoantigens according to an exemplary embodiment of the invention. Tumor specific mutations in cancer samples may be detected using whole-exome (WES) or whole-genome sequencing (WGS) and identified through the application of mutation calling algorithms (e.g., Mutect). Subsequently, candidate neoepitopes may be predicted using well-validated algorithms (e.g., NetMHCpan) and their identification may be refined by experimental validation for peptide-HLA binding and by confirmation of gene expression at the RNA level. These candidate neoantigens may be subsequently tested for their ability to stimulate tumor-specific T cell responses.

Figures 11A-C show the frequency of classes of point mutations that have the potential to generate neoantigens in chronic lymphocytic leukemia (CLL). Analysis of WES and WGS data generated from 91 CLL cases reveals that (**A**) missense mutations are the most frequent class of the somatic alterations with the potential to generate neo-epitopes, while (**B**) frameshift insertions and deletions and (**C**) splice-site mutations constitute less common events.

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Figures 12A-D depict the application of the NetMHCpan prediction algorithm to functionally-defined neoepitopes and CLL cases. FIG. 12 A shows the predicted binding (IC50) to their known restricting HLA allele of 33 functionally identified cancer neoepitopes reported in literature tested by NetMHCpan, sorted on the basis of predicted binding affinity. FIG. 12B shows the distribution of the number of predicted peptides with HLA binding affinity < 150 nM (black) and 150-500 nM (grey) across 31 CLL patients with available HLA typing information. FIG. 12C shows a graph comparing the predicted binding (IC50 < 500 nM by NetMHCpan) of peptides from 4 patients with the experimentally determined binding affinity for HLA-A and -B allele binding using a competitive MHC I allele-binding assay with synthesized peptides. The percent of predicted peptides with evidence of experimental binding (IC50 < 500 nM) are indicated. FIG. 12D shows that from 26 CLL patients for which HLA typing and Affymetrix U133 2.0+ gene expression data were available, the distribution of gene expression was examined for all somatically mutated genes (n=347), and for the subset of gene mutations encoding neoepitopes with predicted HLA binding scores of IC50 < 500 nM (n=180). No-low: genes within the lowest quartile expression; medium: genes within the 2 middle quartiles of expression; and high: genes within the highest quartile of expression.

Figures 13A-B show the same data as in Figure 12D but separately for 9-mer (FIG. 13A) and 10-mer peptides (FIG. 13B). In each case, percentages of peptides with predicted IC50 < 150 nM and 150-500 nM, with evidence of experimental binding are indicated.

Figures 14A-C depict that mutations in *ALMS1* and *C6ORF89* in Pt 1 generate immunogenic peptides. FIG. 14A shows that 25 missense mutations were identified in Pt 1 CLL cells from which 30 peptides from 13 mutations were predicted to bind to Pt 1's MHC class I alleles. A total of 14 peptides from 9 mutations were experimentally confirmed as HLA-binding. Post-transplant T cells (7 yrs) from Pt 1 were stimulated weekly *ex vivo* for 4 weeks with 5 pools of 6 mutated peptides with similar predicted HLA binding, per pool, and subsequently tested by IFN-γ ELISPOT assay. FIG. 14B shows that increased IFN-γ secretion by T cells was detected

against Pool 2 peptides. Negative control - Irrelevant Tax peptide; positive control - PHA. FIG. 14C shows that of Pool 2 peptides, Pt 1 T cells were reactive to mutated *ALMS1* and *C6ORF89* peptides (right panel; averaged results from duplicate wells are displayed). Left panel-The predicted and experimental IC50 scores (nM) of mutated and wildtype *ALMS1* and *C6ORF89* peptides.

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Figure 15 illustrates that the sequence context around the sites of mutations in *FNDC3B*, *C6orf89* and *ALMS1* lack evolutionary conservation. The neoepitopes generated from each of the genes are boxed. Red- conserved amino acids (aa) in all 4 species; blue- conserved aa in at least 2 of 4 species; black –absent conservation across species.

Figure 16 shows localization of somatic mutations reported in *FNDC3B*, *C6orf89* and *ALMS1* genes. Missense mutations identified in *FNDC3B*, *C6orf89* and *ALMS1* in CLL Pts 1 and 2 compared to previously reported somatic mutations in these genes (COSMIC database) across cancers.

Figure 17 shows that mutated FNDC3B generates a naturally immunogenic neoepitope in Pt 2. FIG. 17A shows 26 missense mutations were identified in Pt 2 CLL cells from which 37 peptides from 16 mutations were predicted to bind to Pt 2's MHC class I alleles. A total of 18 peptides from 12 mutations were experimentally confirmed to bind. Post-transplant T cells (~3 yrs) from Pt 2 were stimulated with autologous DCs or B cells pulsed with 3 pools of experimentally validated binding mutated peptides (18 peptides total) for 2 weeks ex vivo (See table S6). FIG. 17B shows increased IFN-y secretion was detected by ELISPOT assay in T cells stimulated with Pool 1 peptides. FIG. 17C shows that of Pool 1 peptides, increased IFN-y secretion was detected against the mut-FNDC3B peptide (bottom panel; averaged results from duplicated wells are displayed). Top panel - Predicted and experimental IC50 scores of mut- and wt- FNDC3B peptides. FIG. 17D illustrates that T cells reactive to mut-FNDC3B demonstrate specificity to the mutated epitope but not the corresponding wildtype peptide (concentrations: 0.1-10 μg/ml), and are polyfunctional, secreting IFN-γ, GM-CSF and IL-2 (Tukey post-hoc tests from two-way ANOVA modeling for comparisons between T cell reactivity against mut vs wt peptide). FIG. 17E shows that Mut-FNDC3B-specific T cells are reactive in a class I-restricted manner (left), and recognize an endogenously processed and presented form of mutated FNDC3B, since they recognized HLA-A2 APCs transfected with a plasmid encoding a minigene of 300bp encompassing the FNDC3B mutation (right) (two-sided t test). Top right - Western blot

analysis-confirming expression of minigenes encoding mut- and wt- *FNDC3B*. FIG. 17F shows that T cells recognizing the mut-*FNDC3B* epitope as detected by HLA-A2⁺/mut *FNDC3B* tetramers are more frequently detected in T cells in Pt 2 compared to T cells from a normal donor. FIG. 17G shows expression of *FNDC3B* (based on Affymetrix U133Plus2 array data) in Pt 2 (triangle), CLL-B cells (n=182) and normal CD19+ B cells from healthy adult volunteers (n=24).

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Figure 18 illustrates kinetics of the mut-*FNDC3B* specific T cell response in relation to the transplant course. FIG. 18 shows molecular tumor burden was measured in Pt 2 using a patient tumor-specific Taqman PCR assay based on the clonotypic IgH sequence at serial time points before and after HSCT (top panel). Middle panel- Detection of mut-*FNDC3B* reactive T cells in comparison to wt-*FNDC3B* or irrelevant peptides from peripheral blood before and after allo-HSCT by IFN-γ ELISPOT following stimulation with peptide-pulsed autologous B cells. The number of IFN-γ-secreting spots per cells at each time point was measured in triplicate (Welch *t* test; mut vs. wt). Inset – IFN-γ secretion of T cells from 6 months post-HSCT (purple) compared to 32 months post-HSCT (red) following exposure to APCs pulsed with 0.1-10 μg/ml (log scale) mut-*FNDC3B* peptide. Bottom panel - Detection of mut-*FNDC3B*-specific TCR Vβ11 cells by nested clone-specific CDR3 PCR before and after HSCT in peripheral blood of Pt 2 (See supplementary methods). Triangles – time points at which a sample was tested; NA- no amplification; black: amplification detected, where '+' indicates detectable amplification up to 2-fold and '++' indicates more than 2-fold greater amplification than the median level of all samples with detectable expression of the clone-specific Vβ11 sequence.

Figures 19A-D show the design of mut-FNDC3B specific TCR V β specific primers in Pt 2. FIG. 19A shows mut-FNDC3B specific T cells detected and isolated from Pt 2 PBMCs 6 months following HSCT using an IFN- γ catch assay. FIG. 19B shows RNA from FNDC3B-reactive T cells expressed TCR V β 11, generating an amplicon of 350bp in length. FIG. 19C shows V β 11-specific real time primers were designed based on the sequence of the mut-FNDC3B clone-specific CDR3 rearrangement, such that the quantitative PCR probe was positioned in the region of junctional diversity (orange). FIG. 19D shows FNDC3B-reactive T cells were monoclonal for V β 11, as detected by spectratyping.

Figures 20A-G illustrate the application of the neoantigen discovery pipeline across cancers. FIG. 20A shows the comparison of overall somatic mutation rate detected across

cancers by massively parallel sequencing. Red-CLL; blue-clear cell renal carcinoma (RCC) and green- melanoma. LSCC: Lung squamous cell carcinoma, Lung AdCa: Lung adenocarcinoma, ESO AdCa: Esophageal adenocarcinoma, DLBCL: Diffused large B- cell lymphoma, GBM: Glioblastoma, Papillary RCC: Papillary renal cell carcinoma, Clear Cell RCC: Clear cell renal carcinoma, CLL: Chronic lymphocytic leukemia, AML: Acute myeloid leukemia. Distribution of FIG. 20B shows the number of missense, frameshift and splice-site mutations per case in melanoma, clear cell RCC and CLL, FIG. 20C shows the average neoORF length generated per sample and FIG. 20D shows predicted neopeptides with IC50 < 150 nM (dashed lines) and < 500 nM (solid lines) generated from missense and frameshift mutations. FIGS. 20E depicts the distributions (shown by box plot) of the number of missense, frameshift and splice-site mutations per case across 13 cancers. FIG. 20F shows the summed neoORF length generated per sample. 20G shows the predicted neopeptides with IC50 < 150 nM and with < 500 nM generated from missense and frameshift mutations,. For all box plots, the left and right ends of the boxes represent the 25th and 75th percentile values, respectively, while the segment in the middle is the median. The left and right extremes of the bars extend to the minimum and maximum values..

DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to personalized strategies for the treatment of neoplasia, and more particularly tumors, by administering a therapeutically effective amount of a pharmaceutical composition (e.g., a cancer vaccine) comprising a plurality of neoplasia/tumor specific neo-antigens to a subject (e.g., a mammal such as a human). As described in more detail below, the present invention is based, at least in part, on the discovery that whole genome/exome sequencing may be used to identify all, or nearly all, mutated neo-antigens that are uniquely present in a neoplasia/tumor of an individual patient, and that this collection of mutated neo-antigens may be analyzed to identify a specific, optimized subset of neo-antigens for use as a personalized cancer vaccine for treatment of the patient's neoplasia/tumor. For example, as shown in FIG. 1, a population of neoplasia/tumor specific neo-antigens may be identified by sequencing the neoplasia/tumor and normal DNA of each patient to identify tumor-specific mutations, and determining the patient's HLA allotype. The population of neoplasia/tumor specific neo-antigens and their cognate native antigens may then be subject to bioinformatic analysis using validated algorithms to predict which tumor-specific mutations create epitopes

that could bind to the patient's HLA allotype, and in particular which tumor-specific mutations create epitopes that could bind to the patient's HLA allotype more effectively than the cognate native antigen. Based on this analysis, a plurality of peptides corresponding to a subset of these mutations may be designed and synthesized for each patient, and pooled together for use as a cancer vaccine in immunizing the patient. The neo-antigens peptides may be combined with an adjuvant (e.g., poly-ICLC) or another anti-neoplastic agent. Without being bound by theory, these neo-antigens are expected to bypass central thymic tolerance (thus allowing stronger anti-tumor T cell response), while reducing the potential for autoimmunity (e.g., by avoiding targeting of normal self-antigens).

The immune system can be classified into two functional subsystems: the innate and the acquired immune system. The innate immune system is the first line of defense against infections, and most potential pathogens are rapidly neutralized by this system before they can cause, for example, a noticeable infection. The acquired immune system reacts to molecular structures, referred to as antigens, of the intruding organism. There are two types of acquired immune reactions, which include the humoral immune reaction and the cell-mediated immune reaction. In the humoral immune reaction, antibodies secreted by B cells into bodily fluids bind to pathogen-derived antigens, leading to the elimination of the pathogen through a variety of mechanisms, e.g. complement-mediated lysis. In the cell-mediated immune reaction, T-cells capable of destroying other cells are activated. For example, if proteins associated with a disease are present in a cell, they are fragmented proteolytically to peptides within the cell. Specific cell proteins then attach themselves to the antigen or peptide formed in this manner and transport them to the surface of the cell, where they are presented to the molecular defense mechanisms, in particular T-cells, of the body. Cytotoxic T cells recognize these antigens and kill the cells that harbor the antigens.

The molecules that transport and present peptides on the cell surface are referred to as proteins of the major histocompatibility complex (MHC). MHC proteins are classified into two types, referred to as MHC class I and MHC class II. The structures of the proteins of the two MHC classes are very similar; however, they have very different functions. Proteins of MHC class I are present on the surface of almost all cells of the body, including most tumor cells. MHC class I proteins are loaded with antigens that usually originate from endogenous proteins or from pathogens present inside cells, and are then presented to naïve or cytotoxic T-lymphocytes

(CTLs). MHC class II proteins are present on dendritic cells, B-lymphocytes, macrophages and other antigen-presenting cells. They mainly present peptides, which are processed from external antigen sources, i.e. outside of the cells, to T-helper (Th) cells. Most of the peptides bound by the MHC class I proteins originate from cytoplasmic proteins produced in the healthy host cells of an organism itself, and do not normally stimulate an immune reaction. Accordingly, cytotoxic T-lymphocytes that recognize such self-peptide-presenting MHC molecules of class I are deleted in the thymus (central tolerance) or, after their release from the thymus, are deleted or inactivated, i.e. tolerized (peripheral tolerance). MHC molecules are capable of stimulating an immune reaction when they present peptides to non-tolerized T-lymphocytes. Cytotoxic T-lymphocytes have both T-cell receptors (TCR) and CD8 molecules on their surface. T-Cell receptors are capable of recognizing and binding peptides complexed with the molecules of MHC class I. Each cytotoxic T-lymphocyte expresses a unique T-cell receptor which is capable of binding specific MHC/peptide complexes.

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The peptide antigens attach themselves to the molecules of MHC class I by competitive affinity binding within the endoplasmic reticulum, before they are presented on the cell surface. Here, the affinity of an individual peptide antigen is directly linked to its amino acid sequence and the presence of specific binding motifs in defined positions within the amino acid sequence. If the sequence of such a peptide is known, it is possible to manipulate the immune system against diseased cells using, for example, peptide vaccines.

One of the critical barriers to developing curative and tumor-specific immunotherapy is the identification and selection of highly specific and restricted tumor antigens to avoid autoimmunity. Tumor neo-antigens, which arise as a result of genetic change (e.g., inversions, translocations, deletions, missense mutations, splice site mutations, etc.) within malignant cells, represent the most tumor-specific class of antigens. Neo-antigens have rarely been used in cancer vaccines due to technical difficulties in identifying them, selecting optimized neo-antigens, and producing neo-antigens for use in a vaccine. According to the present invention, these problems may be addressed by:

• identifying all, or nearly all, mutations in the neoplasia/tumor at the DNA level using whole genome, whole exome (e.g., only captured exons), or RNA sequencing of tumor versus matched germline samples from each patient;

• analyzing the identified mutations with one or more peptide-MHC binding prediction algorithms to generate a plurality of candidate neo-antigen T cell epitopes that are expressed within the neoplasia/tumor and may bind patient HLA alleles; and

• synthesizing the plurality of candidate neo-antigen peptides selected from the sets of all neoORF peptides and predicted binding peptides for use in a cancer vaccine.

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For example, translating sequencing information into a therapeutic vaccine may include:

- (1) Prediction of personal mutated peptides that can bind to HLA molecules of the individual. Efficiently choosing which particular mutations to utilize as immunogen requires identification of the patient HLA type and the ability to predict which mutated peptides would efficiently bind to the patient's HLA alleles. Recently, neural network based learning approaches with validated binding and non-binding peptides have advanced the accuracy of prediction algorithms for the major HLA-A and -B alleles.
- (2) Formulating the drug as a multi-epitope vaccine of long peptides. Targeting as many mutated epitopes as practically possible takes advantage of the enormous capacity of the immune system, prevents the opportunity for immunological escape by down-modulation of a particular immune targeted gene product, and compensates for the known inaccuracy of epitope prediction approaches. Synthetic peptides provide a particularly useful means to prepare multiple immunogens efficiently and to rapidly translate identification of mutant epitopes to an effective vaccine. Peptides can be readily synthesized chemically and easily purified utilizing reagents free of contaminating bacteria or animal substances. The small size allows a clear focus on the mutated region of the protein and also reduces irrelevant antigenic competition from other components (unmutated protein or viral vector antigens).
- (3) Combination with a strong vaccine adjuvant. Effective vaccines require a strong adjuvant to initiate an immune response. As described below, poly-ICLC, an agonist of TLR3 and the RNA helicase –domains of MDA5 and RIG3, has shown several desirable properties for a vaccine adjuvant. These properties include the induction of local and systemic activation of immune cells *in vivo*, production of stimulatory chemokines and cytokines, and stimulation of antigen-presentation by DCs. Furthermore, poly-ICLC can induce durable CD4⁺ and CD8⁺ responses in humans. Importantly, striking similarities in the upregulation of transcriptional and signal transduction pathways were seen in subjects vaccinated with poly-ICLC and in volunteers who had received the highly effective, replication-competent yellow fever vaccine. Furthermore,

>90% of ovarian carcinoma patients immunized with poly-ICLC in combination with a NY-ESO-1 peptide vaccine (in addition to Montanide) showed induction of CD4⁺ and CD8⁺ T cell, as well as antibody responses to the peptide in a recent phase 1 study. At the same time, poly-ICLC has been extensively tested in more than 25 clinical trials to date and exhibited a relatively benign toxicity profile.

The above-described advantages of the invention are described in further detail below.

Identification of Tumor Specific Neo-antigen Mutations

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The present invention is based, at least in part, on the ability to identify all, or nearly all, of the mutations within a neoplasia/tumor (e.g., translocations, inversions, large and small deletions and insertions, missense mutations, splice site mutations, etc.). In particular, these mutations are present in the genome of neoplasia/tumor cells of a subject, but not in normal tissue from the subject. Such mutations are of particular interest if they lead to changes that result in a protein with an altered amino acid sequence that is unique to the patient's neoplasia/tumor (e.g., a neo-antigen). For example, useful mutations may include: (1) nonsynonymous mutations leading to different amino acids in the protein; (2) read-through mutations in which a stop codon is modified or deleted, leading to translation of a longer protein with a novel tumor-specific sequence at the C-terminus; (3) splice site mutations that lead to the inclusion of an intron in the mature mRNA and thus a unique tumor-specific protein sequence; (4) chromosomal rearrangements that give rise to a chimeric protein with tumor-specific sequences at the junction of 2 proteins (i.e., gene fusion); (5) frameshift mutations or deletions that lead to a new open reading frame with a novel tumor-specific protein sequence; and the like. Peptides with mutations or mutated polypeptides arising from, for example, splice-site, frameshift, read-through, or gene fusion mutations in tumor cells may be identified by sequencing DNA, RNA or protein in tumor versus normal cells.

Also within the scope of the inventions is personal neo-antigen peptides derived from common tumor driver genes and may further include previously identified tumor specific mutations. For example, known common tumor driver genes and tumor mutations in common tumor driver genes may be found on the world wide web at (www)sanger.ac.uk/cosmic.

A number of initiatives are currently underway to obtain sequence information directly from millions of individual molecules of DNA or RNA in parallel. Real-time single molecule

sequencing-by-synthesis technologies rely on the detection of fluorescent nucleotides as they are incorporated into a nascent strand of DNA that is complementary to the template being sequenced. In one method, oligonucleotides 30-50 bases in length are covalently anchored at the 5' end to glass cover slips. These anchored strands perform two functions. First, they act as capture sites for the target template strands if the templates are configured with capture tails complementary to the surface-bound oligonucleotides. They also act as primers for the template directed primer extension that forms the basis of the sequence reading. The capture primers function as a fixed position site for sequence determination using multiple cycles of synthesis, detection, and chemical cleavage of the dye-linker to remove the dye. Each cycle consists of adding the polymerase/labeled nucleotide mixture, rinsing, imaging and cleavage of dye. In an alternative method, polymerase is modified with a fluorescent donor molecule and immobilized on a glass slide, while each nucleotide is color-coded with an acceptor fluorescent moiety attached to a gamma-phosphate. The system detects the interaction between a fluorescently-tagged polymerase and a fluorescently modified nucleotide as the nucleotide becomes incorporated into the de novo chain. Other sequencing-by-synthesis technologies also exist.

Preferably, any suitable sequencing-by-synthesis platform can be used to identify mutations. Four major sequencing-by-synthesis platforms are currently available: the Genome Sequencers from Roche/454 Life Sciences, the HiSeq Analyzer from Illumina/Solexa, the SOLiD system from Applied BioSystems, and the Heliscope system from Helicos Biosciences. Sequencing-by-synthesis platforms have also been described by Pacific Biosciences and VisiGen Biotechnologies. Each of these platforms can be used in the methods of the invention. In some embodiments, a plurality of nucleic acid molecules being sequenced is bound to a support (e.g., solid support). To immobilize the nucleic acid on a support, a capture sequence/universal priming site can be added at the 3' and/or 5' end of the template. The nucleic acids may be bound to the support by hybridizing the capture sequence to a complementary sequence covalently attached to the support. The capture sequence (also referred to as a universal capture sequence) is a nucleic acid sequence complementary to a sequence attached to a support that may dually serve as a universal primer.

As an alternative to a capture sequence, a member of a coupling pair (such as, e.g., antibody/antigen, receptor/ligand, or the avidin-biotin pair as described in, e.g., U.S. Patent Application No. 2006/0252077) may be linked to each fragment to be captured on a surface

coated with a respective second member of that coupling pair. Subsequent to the capture, the sequence may be analyzed, for example, by single molecule detection/sequencing, e.g., as described in the Examples and in U.S. Patent No. 7,283,337, including template-dependent sequencing-by- synthesis. In sequencing-by-synthesis, the surface-bound molecule is exposed to a plurality of labeled nucleotide triphosphates in the presence of polymerase. The sequence of the template is determined by the order of labeled nucleotides incorporated into the 3' end of the growing chain. This can be done in real time or in a step-and-repeat mode. For real-time analysis, different optical labels to each nucleotide may be incorporated and multiple lasers may be utilized for stimulation of incorporated nucleotides.

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Any cell type or tissue may be utilized to obtain nucleic acid samples for use in the sequencing methods described herein. In a preferred embodiment, the DNA or RNA sample is obtained from a neoplasia/tumor or a bodily fluid, e.g., blood, obtained by known techniques (e.g. venipuncture) or saliva. Alternatively, nucleic acid tests can be performed on dry samples (e.g. hair or skin).

A variety of methods are available for detecting the presence of a particular mutation or allele in an individual's DNA or RNA. Advancements in this field have provided accurate, easy, and inexpensive large-scale SNP genotyping. Most recently, for example, several new techniques have been described including dynamic allele-specific hybridization (DASH), microplate array diagonal gel electrophoresis (MADGE), pyrosequencing, oligonucleotide-specific ligation, the TaqMan system as well as various DNA "chip" technologies such as the Affymetrix SNP chips. These methods require amplification of the target genetic region, typically by PCR. Still other newly developed methods, based on the generation of small signal molecules by invasive cleavage followed by mass spectrometry or immobilized padlock probes and rolling-circle amplification, might eventually eliminate the need for PCR. Several of the methods known in the art for detecting specific single nucleotide polymorphisms are summarized below. The method of the present invention is understood to include all available methods.

PCR based detection means may include multiplex amplification of a plurality of markers simultaneously. For example, it is well known in the art to select PCR primers to generate PCR products that do not overlap in size and can be analyzed simultaneously.

Alternatively, it is possible to amplify different markers with primers that are differentially labeled and thus can each be differentially detected. Of course, hybridization based

detection means allow the differential detection of multiple PCR products in a sample. Other techniques are known in the art to allow multiplex analyses of a plurality of markers.

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Several methods have been developed to facilitate analysis of single nucleotide polymorphisms in genomic DNA or cellular RNA. In one embodiment, the single base polymorphism can be detected by using a specialized exonuclease-resistant nucleotide, as disclosed, e.g., U.S. Patent No. 4,656,127. According to the method, a primer complementary to the allelic sequence immediately 3' to the polymorphic site is permitted to hybridize to a target molecule obtained from a particular animal or human. If the polymorphic site on the target molecule contains a nucleotide that is complementary to the particular exonuclease-resistant nucleotide derivative present, then that derivative will be incorporated onto the end of the hybridized primer. Such incorporation renders the primer resistant to exonuclease, and thereby permits its detection. Since the identity of the exonuclease-resistant derivative of the sample is known, a finding that the primer has become resistant to exonucleases reveals that the nucleotide present in the polymorphic site of the target molecule was complementary to that of the nucleotide derivative used in the reaction. This method has the advantage that it does not require the determination of large amounts of extraneous sequence data.

In another embodiment of the invention, a solution-based method is used for determining the identity of the nucleotide of a polymorphic site. Cohen et al. (French Patent No. 2,650,840; PCT Application No. WO1991/02087). As in the method of U.S. Patent No. 4,656,127, a primer may be employed that is complementary to allelic sequences immediately 3' to a polymorphic site. The method determines the identity of the nucleotide of that site using labeled dideoxynucleotide derivatives, which, if complementary to the nucleotide of the polymorphic site, will become incorporated onto the terminus of the primer.

An alternative method, known as Genetic Bit Analysis or GBA® is described in PCT Application No. WO1992/15712). GBA® uses mixtures of labeled terminators and a primer that is complementary to the sequence 3' to a polymorphic site. The labeled terminator that is incorporated is thus determined by, and complementary to, the nucleotide present in the polymorphic site of the target molecule being evaluated. In contrast to the method of Cohen et al. (French Patent 2,650,840; PCT Application No. W01991/02087) the GBA® method is preferably a heterogeneous phase assay, in which the primer or the target molecule is immobilized to a solid phase.

Recently, several primer-guided nucleotide incorporation procedures for assaying polymorphic sites in DNA have been described (Komher, J. S. et al., Nucl. Acids. Res. 17:7779-7784 (1989); Sokolov, B. P., Nucl. Acids Res. 18:3671 (1990); Syvanen, A.-C, et al., Genomics 8:684-692 (1990); Kuppuswamy, M. N. et al., Proc. Natl. Acad. Sci. (U.S.A.) 88: 1143-1147 (1991); Prezant, T. R. et al., Hum. Mutat. 1: 159-164 (1992); Ugozzoli, L. et al., GATA 9: 107-112 (1992); Nyren, P. et al., Anal. Biochem. 208: 171-175 (1993)). These methods differ from GBA® in that they all rely on the incorporation of labeled deoxynucleotides to discriminate between bases at a polymorphic site. In such a format, since the signal is proportional to the number of deoxynucleotides incorporated, polymorphisms that occur in runs of the same nucleotide can result in signals that are proportional to the length of the run (Syvanen, A.-C, et al., Amer. J. Hum. Genet. 52:46-59 (1993)).

An alternative method for identifying tumor specific neo-antigens is direct protein sequencing. Protein sequencing of enzymatic digests using multidimensional MS techniques (MSn) including tandem mass spectrometry (MS/MS)) can also be used to identify neo-antigens of the invention. Such proteomic approaches permit rapid, highly automated analysis (see, e.g., K. Gevaert and J. Vandekerckhove, Electrophoresis 21:1145-1154 (2000)). It is further contemplated within the scope of the invention that high-throughput methods for de novo sequencing of unknown proteins may be used to analyze the proteome of a patient's tumor to identify expressed neo-antigens. For example, meta shotgun protein sequencing may be used to identify expressed neo-antigens (see e.g., Guthals et al. (2012) Shotgun Protein Sequencing with Meta-contig Assembly, Molecular and Cellular Proteomics 11(10):1084-96).

Tumor specific neo-antigens may also be identified using MHC multimers to identify neo-antigen-specific T-cell responses. For example, highthroughput analysis of neo-antigen-specific T-cell responses in patient samples may be performed using MHC tetramer-based screening techniques (see e.g., Hombrink et al. (2011) High-Throughput Identification of Potential Minor Histocompatibility Antigens by MHC Tetramer-Based Screening: Feasibility and Limitations 6(8):1-11; Hadrup et al. (2009) Parallel detection of antigen-specific T-cell responses by multidimensional encoding of MHC multimers, Nature Methods, 6(7):520-26; van Rooij et al. (2013) Tumor exome analysis reveals neoantigen-specific T-cell reactivity in an Ipilimumab-responsive melanoma, Journal of Clinical Oncology, 31:1-4; and Heemskerk et al. (2013) The cancer antigenome, EMBO Journal, 32(2):194-203). It is contemplated within the

scope of the invention that such tetramer-based screening techniques may be used for the initial identification of tumor specific neo-antigens, or alternatively as a secondary screening protocol to assess what neo-antigens a patient may have already been exposed to, thereby facilitating the selection of candidate neo-antigens for the vaccines of the invention.

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Design of Tumor Specific Neo-Antigens

The invention further includes isolated peptides (e.g., neo-antigenic peptides containing the tumor specific mutations identified by the methods of the invention, peptides that comprise know tumor specific mutations, and mutant polypeptides or fragments thereof identified by the method of the invention). These peptides and polypeptides are referred to herein as "neo-antigenic peptides" or "neo-antigenic polypeptides." The term "peptide" is used interchangeably with "mutant peptide" and "neo-antigenic peptide" and "wildtype peptide" in the present specification to designate a series of residues, typically L-amino acids, connected one to the other, typically by peptide bonds between the alpha-amino and alpha-carboxyl groups of adjacent amino acids. The polypeptides or peptides can be of a variety of lengths and will minimally include the small region predicted to bind to the HLA molecule of the patient (the "epitope") as well as additional adjacent amino acids extending in both the N- and C-terminal directions. The polypeptides or peptides can be either in their neutral (uncharged) forms or in forms which are salts, and either free of modifications such as glycosylation, side chain oxidation, or phosphorylation or containing these modifications, subject to the condition that the modification not destroy the biological activity of the polypeptides as herein described.

In certain embodiments the size of the at least one neo-antigenic peptide molecule may comprise, but is not limited to, about 8, about 9, about 10, about 11, about 12, about 13, about 14, about 15, about 16, about 17, about 18, about 19, about 20, about 21, about 22, about 23, about 24, about 25, about 26, about 27, about 28, about 29, about 30, about 31, about 32, about 33, about 34, about 35, about 36, about 37, about 38, about 39, about 40, about 41, about 42, about 43, about 44, about 45, about 46, about 47, about 48, about 49, about 50, about 60, about 70, about 80, about 90, about 100, about 110, about 120 or greater amino molecule residues, and any range derivable therein. In specific embodiments the neo-antigenic peptide molecules are equal to or less than 50 amino acids. In a preferred embodiment, the neo-antigenic peptide molecules are equal to about 20 to about 30 amino acids.

A longer peptide may be designed in several ways. For example, when the HLA-binding regions (e.g., the "epitopes") are predicted or known, a longer peptide may consist of either: individual binding peptides with an extension of 0-10 amino acids toward the N- and C-terminus of each corresponding gene product. A longer peptide may also consist of a concatenation of some or all of the binding peptides with extended sequences for each. In another case, when sequencing reveals a long (>10 residues) neo-epitope sequence present in the tumor (e.g. due to a frameshift, read-through or intron inclusion that leads to a novel peptide sequence), a longer peptide may consist of the entire stretch of novel tumor-specific amino acids. In both cases, use of a longer peptide requires endogenous processing by professional antigen presenting cells such as dendritic cells and may lead to more effective antigen presentation and induction of T cell responses. In some cases, it is desirable or preferable to alter the extended sequence to improve the biochemical properties of the polypeptide (properties such as solubility or stability) or to improve the likelihood for efficient proteasomal processing of the peptide (Zhang et al (2012) Aminopeptidase substrate preference affects HIV epitope presentation and predicts immune escape patterns in HIV-infected individuals. J. Immunol 188:5924-34; Hearn et al (2010) Characterizing the specificity and co-operation of aminopeptidases in the cytosol and ER during MHC Class I antigen presentation. J. Immunol 184(9):4725-32; Wiemerhaus et al (2012) Peptidases trimming MHC Class I ligands. Curr Opin Immunol 25:1-7).

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The neo-antigenic peptides and polypeptides may bind an HLA protein. In preferred aspects, the neo-antigenic peptides and polypeptides may bind an HLA protein with greater affinity than the corresponding native / wild-type peptide. The neo-antigenic peptide or polypeptide may have an IC50 of about less than 1000 nM, about less than 500 nM, about less than 200 nM, about less than 100 nM, about less than 100 nM, or about less than 50 nM.

In a preferred embodiment, the neo-antigenic peptides and polypeptides of the invention do not induce an autoimmune response and/or invoke immunological tolerance when administered to a subject.

The invention also provides compositions comprising a plurality of neo-antigenic peptides. In some embodiments, the composition comprises at least 5 or more neo-antigenic peptides. In some embodiments the composition contains at least about 6, about 8, about 10, about 12, about 14, about 16, about 18, or about 20 distinct peptides. In some embodiments the

composition contains at least 20 distinct peptides. According to the invention, 2 or more of the distinct peptides may be derived from the same polypeptide. For example, if a preferred neoantigenic mutation encodes a neoORF polypeptide, two or more of the neo-antigenic peptides may be derived from the neoORF polypeptide. In one embodiment, the two or more neoantigenic peptides derived from the neoORF polypeptide may comprise a tiled array that spans the polypeptide (e.g., the neo-antigenic peptides may comprise a series of overlapping neoantigenic peptides that spans a portion, or all, of the neoORF polypeptide). Without being bound by theory, each peptide is believed to have its own epitope; accordingly, a tiling array that spans one neoORF polypeptide may give rise to polypeptides that are targeted to different HLA molecules. Neo-antigenic peptides can be derived from any protein coding gene. Exemplary polypeptides from which the neo-antigenic peptides may be derived can be found for example at the COSMIC database (on the worldwide web at (www)sanger.ac.uk/cosmic). COSMIC curates comprehensive information on somatic mutations in human cancer. The peptide may contain the tumor specific mutation. In some aspects the tumor specific mutation is in a common driver gene or is a common driver mutation for a particular cancer type. For example, common driver mutation peptides may include, but are not limited to, the following: a SF3B1 polypeptide, a MYD88 polypeptide, a TP53 polypeptide, an ATM polypeptide, an Abl polypeptide, A FBXW7 polypeptide, a DDX3X polypeptide, a MAPK1 polypeptide, or a GNB1 polypeptide.

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The neo-antigenic peptides, polypeptides, and analogs can be further modified to contain additional chemical moieties not normally part of the protein. Those derivatized moieties can improve the solubility, the biological half-life, absorption of the protein, or binding affinity. The moieties can also reduce or eliminate any desirable side effects of the proteins and the like. An overview for those moieties can be found in Remington's Pharmaceutical Sciences, 20th ed., Mack Publishing Co., Easton, PA (2000).

For example, neo-antigenic peptides and polypeptides having the desired activity may be modified as necessary to provide certain desired attributes, e.g. improved pharmacological characteristics, while increasing or at least retaining substantially all of the biological activity of the unmodified peptide to bind the desired MHC molecule and activate the appropriate T cell. For instance, the neo-antigenic peptide and polypeptides may be subject to various changes, such as substitutions, either conservative or non-conservative, where such changes might provide for certain advantages in their use, such as improved MHC binding. Such conservative substitutions

may encompass replacing an amino acid residue with another amino acid residue that is biologically and/or chemically similar, e.g., one hydrophobic residue for another, or one polar residue for another. The effect of single amino acid substitutions may also be probed using D-amino acids. Such modifications may be made using well known peptide synthesis procedures, as described in e.g., Merrifield, Science 232:341-347 (1986), Barany & Merrifield, The Peptides, Gross & Meienhofer, eds. (N.Y., Academic Press), pp. 1-284 (1979); and Stewart & Young, Solid Phase Peptide Synthesis, (Rockford, III., Pierce), 2d Ed. (1984).

The neo-antigenic peptide and polypeptides may also be modified by extending or decreasing the compound's amino acid sequence, e.g., by the addition or deletion of amino acids. The neo-antigenic peptides, polypeptides, or analogs can also be modified by altering the order or composition of certain residues. It will be appreciated by the skilled artisan that certain amino acid residues essential for biological activity, e.g., those at critical contact sites or conserved residues, may generally not be altered without an adverse effect on biological activity. The non-critical amino acids need not be limited to those naturally occurring in proteins, such as L-a-amino acids, or their D-isomers, but may include non-natural amino acids as well, such as β - γ - δ -amino acids, as well as many derivatives of L-a-amino acids.

Typically, a neo-antigen polypeptide or peptide may be optimized by using a series of peptides with single amino acid substitutions to determine the effect of electrostatic charge, hydrophobicity, etc. on MHC binding. For instance, a series of positively charged (e.g., Lys or Arg) or negatively charged (e.g., Glu) amino acid substitutions may be made along the length of the peptide revealing different patterns of sensitivity towards various MHC molecules and T cell receptors. In addition, multiple substitutions using small, relatively neutral moieties such as Ala, Gly, Pro, or similar residues may be employed. The substitutions may be homo-oligomers or hetero-oligomers. The number and types of residues which are substituted or added depend on the spacing necessary between essential contact points and certain functional attributes which are sought (e.g., hydrophobicity versus hydrophilicity). Increased binding affinity for an MHC molecule or T cell receptor may also be achieved by such substitutions, compared to the affinity of the parent peptide. In any event, such substitutions should employ amino acid residues or other molecular fragments chosen to avoid, for example, steric and charge interference which might disrupt binding.

Amino acid substitutions are typically of single residues. Substitutions, deletions, insertions or any combination thereof may be combined to arrive at a final peptide. Substitutional variants are those in which at least one residue of a peptide has been removed and a different residue inserted in its place.

The neo-antigenic peptides and polypeptides may be modified to provide desired attributes. For instance, the ability of the peptides to induce CTL activity can be enhanced by linkage to a sequence which contains at least one epitope that is capable of inducing a T helper cell response. Particularly preferred immunogenic peptides/T helper conjugates are linked by a spacer molecule. The spacer is typically comprised of relatively small, neutral molecules, such as amino acids or amino acid mimetics, which are substantially uncharged under physiological conditions. The spacers are typically selected from, e.g., Ala, Gly, or other neutral spacers of nonpolar amino acids or neutral polar amino acids. It will be understood that the optionally present spacer need not be comprised of the same residues and thus may be a hetero- or homologomer. When present, the spacer will usually be at least one or two residues, more usually three to six residues. Alternatively, the peptide may be linked to the T helper peptide without a spacer.

The neo-antigenic peptide may be linked to the T helper peptide either directly or via a spacer either at the amino or carboxy terminus of the peptide. The amino terminus of either the neo-antigenic peptide or the T helper peptide may be acylated. Exemplary T helper peptides include tetanus toxoid 830-843, influenza 307-319, malaria circumsporozoite 382-398 and 378-389.

Production of Tumor Specific Neo-antigens

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The present invention is based, at least in part, on the ability to present the immune system of the patient with a pool of tumor specific neo-antigens. One of skill in the art will appreciate that there are a variety of ways in which to produce such tumor specific neo-antigens. In general, such tumor specific neo-antigens may be produced either in vitro or in vivo. Tumor specific neo-antigens may be produced in vitro as peptides or polypeptides, which may then be formulated into a personalized neoplasia vaccine and administered to a subject. As described in further detail below, such in vitro production may occur by a variety of methods known to one of skill in the art such as, for example, peptide synthesis or expression of a peptide/polypeptide

from a DNA or RNA molecule in any of a variety of bacterial, eukaryotic, or viral recombinant expression systems, followed by purification of the expressed peptide/polypeptide.

Alternatively, tumor specific neo-antigens may be produced in vivo by introducing molecules (e.g., DNA, RNA, viral expression systems, and the like) that encode tumor specific neo-antigens into a subject, whereupon the encoded tumor specific neo-antigens are expressed.

In Vitro Peptide/Polypeptide Synthesis

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Proteins or peptides may be made by any technique known to those of skill in the art, including the expression of proteins, polypeptides or peptides through standard molecular biological techniques, the isolation of proteins or peptides from natural sources, or the chemical synthesis of proteins or peptides. The nucleotide and protein, polypeptide and peptide sequences corresponding to various genes have been previously disclosed, and may be found at computerized databases known to those of ordinary skill in the art. One such database is the National Center for Biotechnology Information's Genbank and GenPept databases located at the National Institutes of Health website. The coding regions for known genes may be amplified and/or expressed using the techniques disclosed herein or as would be known to those of ordinary skill in the art. Alternatively, various commercial preparations of proteins, polypeptides and peptides are known to those of skill in the art.

Peptides can be readily synthesized chemically utilizing reagents that are free of contaminating bacterial or animal substances (Merrifield RB: Solid phase peptide synthesis. I. The synthesis of a tetrapeptide. J. Am. Chem. Soc. 85:2149-54, 1963).

A further aspect of the invention provides a nucleic acid (e.g., a polynucleotide) encoding a neo-antigenic peptide of the invention, which may be used to produce the neo-antigenic peptide in vitro. The polynucleotide may be, e.g., DNA, cDNA, PNA, CNA, RNA, either single- and/or double-stranded, or native or stabilized forms of polynucleotides, such as e.g. polynucleotides with a phosphorothiate backbone, or combinations thereof and it may or may not contain introns so long as it codes for the peptide. A still further aspect of the invention provides an expression vector capable of expressing a polypeptide according to the invention. Expression vectors for different cell types are well known in the art and can be selected without undue experimentation. Generally, the DNA is inserted into an expression vector, such as a plasmid, in proper orientation and correct reading frame for expression. If necessary, the DNA may be linked to the

appropriate transcriptional and translational regulatory control nucleotide sequences recognized by the desired host (e.g., bacteria), although such controls are generally available in the expression vector. The vector is then introduced into the host bacteria for cloning using standard techniques (see, e.g., Sambrook et al. (1989) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.).

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The invention further embraces variants and equivalents which are substantially homologous to the identified tumor specific neo-antigens described herein. These can contain, for example, conservative substitution mutations, i.e., the substitution of one or more amino acids by similar amino acids. For example, conservative substitution refers to the substitution of an amino acid with another within the same general class such as, for example, one acidic amino acid with another acidic amino acid, one basic amino acid with another basic amino acid, or one neutral amino acid by another neutral amino acid. What is intended by a conservative amino acid substitution is well known in the art.

The invention also includes expression vectors comprising the isolated polynucleotides, as well as host cells containing the expression vectors. It is also contemplated within the scope of the invention that the neo-antigenic peptides may be provided in the form of RNA or cDNA molecules encoding the desired neo-antigenic peptides. The invention also provides that the one or more neo-antigenic peptides of the invention may be encoded by a single expression vector. The invention also provides that the one or more neo-antigenic peptides of the invention may be encoded and expressed in vivo using a viral based system (e.g., an adenovirus system).

The term "polynucleotide encoding a polypeptide" encompasses a polynucleotide which includes only coding sequences for the polypeptide as well as a polynucleotide which includes additional coding and/or non-coding sequences. The polynucleotides of the invention can be in the form of RNA or in the form of DNA. DNA includes cDNA, genomic DNA, and synthetic DNA; and can be double-stranded or single-stranded, and if single stranded can be the coding strand or non-coding (anti-sense) strand.

In embodiments, the polynucleotides may comprise the coding sequence for the tumor specific neo-antigenic peptide fused in the same reading frame to a polynucleotide which aids, for example, in expression and/or secretion of a polypeptide from a host cell (e.g., a leader sequence which functions as a secretory sequence for controlling transport of a polypeptide from

the cell). The polypeptide having a leader sequence is a preprotein and can have the leader sequence cleaved by the host cell to form the mature form of the polypeptide.

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In embodiments, the polynucleotides can comprise the coding sequence for the tumor specific neo-antigenic peptide fused in the same reading frame to a marker sequence that allows, for example, for purification of the encoded polypeptide, which may then be incorporated into the personalized neoplasia vaccine. For example, the marker sequence can be a hexa-histidine tag supplied by a pQE-9 vector to provide for purification of the mature polypeptide fused to the marker in the case of a bacterial host, or the marker sequence can be a hemagglutinin (HA) tag derived from the influenza hemagglutinin protein when a mammalian host (e.g., COS-7 cells) is used. Additional tags include, but are not limited to, Calmodulin tags, FLAG tags, Myc tags, S tags, SBP tags, Softag 1, Softag 3, V5 tag, Xpress tag, Isopeptag, SpyTag, Biotin Carboxyl Carrier Protein (BCCP) tags, GST tags, fluorescent protein tags (e.g., green fluorescent protein tags), maltose binding protein tags, Nus tags, Strep-tag, thioredoxin tag, TC tag, Ty tag, and the like.

In embodiments, the polynucleotides may comprise the coding sequence for one or more of the tumor specific neo-antigenic peptides fused in the same reading frame to create a single concatamerized neo-antigenic peptide construct capable of producing multiple neo-antigenic peptides.

In embodiments, the present invention provides isolated nucleic acid molecules having a nucleotide sequence at least 60% identical, at least 65% identical, at least 70% identical, at least 75% identical, at least 80% identical, at least 85% identical, at least 90% identical, at least 95% identical, or at least 96%, 97%, 98% or 99% identical to a polynucleotide encoding a tumor specific neo-antigenic peptide of the present invention.

By a polynucleotide having a nucleotide sequence at least, for example, 95% "identical" to a reference nucleotide sequence is intended that the nucleotide sequence of the polynucleotide is identical to the reference sequence except that the polynucleotide sequence can include up to five point mutations per each 100 nucleotides of the reference nucleotide sequence. In other words, to obtain a polynucleotide having a nucleotide sequence at least 95% identical to a reference nucleotide sequence, up to 5% of the nucleotides in the reference sequence can be deleted or substituted with another nucleotide, or a number of nucleotides up to 5% of the total

nucleotides in the reference sequence can be inserted into the reference sequence. These mutations of the reference sequence can occur at the amino- or carboxy-terminal positions of the reference nucleotide sequence or anywhere between those terminal positions, interspersed either individually among nucleotides in the reference sequence or in one or more contiguous groups within the reference sequence.

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As a practical matter, whether any particular nucleic acid molecule is at least 80% identical, at least 85% identical, at least 90% identical, and in some embodiments, at least 95%, 96%, 97%, 98%, or 99% identical to a reference sequence can be determined conventionally using known computer programs such as the Bestfit program (Wisconsin Sequence Analysis Package, Version 8 for Unix, Genetics Computer Group, University Research Park, 575 Science Drive, Madison, WI 53711). Bestfit uses the local homology algorithm of Smith and Waterman, Advances in Applied Mathematics 2:482-489 (1981), to find the best segment of homology between two sequences. When using Bestfit or any other sequence alignment program to determine whether a particular sequence is, for instance, 95% identical to a reference sequence according to the present invention, the parameters are set such that the percentage of identity is calculated over the full length of the reference nucleotide sequence and that gaps in homology of up to 5% of the total number of nucleotides in the reference sequence are allowed.

The isolated tumor specific neo-antigenic peptides described herein can be produced in vitro (e.g., in the laboratory) by any suitable method known in the art. Such methods range from direct protein synthetic methods to constructing a DNA sequence encoding isolated polypeptide sequences and expressing those sequences in a suitable transformed host. In some embodiments, a DNA sequence is constructed using recombinant technology by isolating or synthesizing a DNA sequence encoding a wild-type protein of interest. Optionally, the sequence can be mutagenized by site-specific mutagenesis to provide functional analogs thereof. *See, e.g.* Zoeller et al., *Proc. Nat'l. Acad. Sci.* USA 81:5662-5066 (1984) and U.S. Pat. No. 4,588,585.

In embodiments, a DNA sequence encoding a polypeptide of interest would be constructed by chemical synthesis using an oligonucleotide synthesizer. Such oligonucleotides can be designed based on the amino acid sequence of the desired polypeptide and selecting those codons that are favored in the host cell in which the recombinant polypeptide of interest will be produced. Standard methods can be applied to synthesize an isolated polynucleotide sequence

encoding an isolated polypeptide of interest. For example, a complete amino acid sequence can be used to construct a back-translated gene. Further, a DNA oligomer containing a nucleotide sequence coding for the particular isolated polypeptide can be synthesized. For example, several small oligonucleotides coding for portions of the desired polypeptide can be synthesized and then ligated. The individual oligonucleotides typically contain 5' or 3' overhangs for complementary assembly.

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Once assembled (e.g., by synthesis, site-directed mutagenesis, or another method), the polynucleotide sequences encoding a particular isolated polypeptide of interest will be inserted into an expression vector and optionally operatively linked to an expression control sequence appropriate for expression of the protein in a desired host. Proper assembly can be confirmed by nucleotide sequencing, restriction mapping, and expression of a biologically active polypeptide in a suitable host. As well known in the art, in order to obtain high expression levels of a transfected gene in a host, the gene can be operatively linked to transcriptional and translational expression control sequences that are functional in the chosen expression host.

Recombinant expression vectors may be used to amplify and express DNA encoding the tumor specific neo-antigenic peptides. Recombinant expression vectors are replicable DNA constructs which have synthetic or cDNA-derived DNA fragments encoding a tumor specific neo-antigenic peptide or a bioequivalent analog operatively linked to suitable transcriptional or translational regulatory elements derived from mammalian, microbial, viral or insect genes. A transcriptional unit generally comprises an assembly of (1) a genetic element or elements having a regulatory role in gene expression, for example, transcriptional promoters or enhancers, (2) a structural or coding sequence which is transcribed into mRNA and translated into protein, and (3) appropriate transcription and translation initiation and termination sequences, as described in detail below. Such regulatory elements can include an operator sequence to control transcription. The ability to replicate in a host, usually conferred by an origin of replication, and a selection gene to facilitate recognition of transformants can additionally be incorporated. DNA regions are operatively linked when they are functionally related to each other. For example, DNA for a signal peptide (secretory leader) is operatively linked to DNA for a polypeptide if it is expressed as a precursor which participates in the secretion of the polypeptide; a promoter is operatively linked to a coding sequence if it controls the transcription of the sequence; or a ribosome binding site is operatively linked to a coding sequence if it is positioned so as to permit

translation. Generally, operatively linked means contiguous, and in the case of secretory leaders, means contiguous and in reading frame. Structural elements intended for use in yeast expression systems include a leader sequence enabling extracellular secretion of translated protein by a host cell. Alternatively, where recombinant protein is expressed without a leader or transport sequence, it can include an N-terminal methionine residue. This residue can optionally be subsequently cleaved from the expressed recombinant protein to provide a final product.

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The choice of expression control sequence and expression vector will depend upon the choice of host. A wide variety of expression host/vector combinations can be employed. Useful expression vectors for eukaryotic hosts, include, for example, vectors comprising expression control sequences from SV40, bovine papilloma virus, adenovirus and cytomegalovirus. Useful expression vectors for bacterial hosts include known bacterial plasmids, such as plasmids from *Escherichia coli*, including pCR 1, pBR322, pMB9 and their derivatives, wider host range plasmids, such as M13 and filamentous single-stranded DNA phages.

Suitable host cells for expression of a polypeptide include prokaryotes, yeast, insect or higher eukaryotic cells under the control of appropriate promoters. Prokaryotes include gram negative or gram positive organisms, for example *E. coli* or *bacilli*. Higher eukaryotic cells include established cell lines of mammalian origin. Cell-free translation systems could also be employed. Appropriate cloning and expression vectors for use with bacterial, fungal, yeast, and mammalian cellular hosts are well known in the art (see Pouwels *et al.*, *Cloning Vectors: A Laboratory Manual*, Elsevier, N.Y., 1985).

Various mammalian or insect cell culture systems are also advantageously employed to express recombinant protein. Expression of recombinant proteins in mammalian cells can be performed because such proteins are generally correctly folded, appropriately modified and completely functional. Examples of suitable mammalian host cell lines include the COS-7 lines of monkey kidney cells, described by Gluzman (*Cell* 23:175, 1981), and other cell lines capable of expressing an appropriate vector including, for example, L cells, C127, 3T3, Chinese hamster ovary (CHO), HeLa and BHK cell lines. Mammalian expression vectors can comprise nontranscribed elements such as an origin of replication, a suitable promoter and enhancer linked to the gene to be expressed, and other 5' or 3' flanking nontranscribed sequences, and 5' or 3' nontranslated sequences, such as necessary ribosome binding sites, a polyadenylation site, splice

donor and acceptor sites, and transcriptional termination sequences. Baculovirus systems for production of heterologous proteins in insect cells are reviewed by Luckow and Summers, *Bio/Technology* 6:47 (1988).

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The proteins produced by a transformed host can be purified according to any suitable method. Such standard methods include chromatography (e.g., ion exchange, affinity and sizing column chromatography, and the like), centrifugation, differential solubility, or by any other standard technique for protein purification. Affinity tags such as hexahistidine, maltose binding domain, influenza coat sequence, glutathione-S-transferase, and the like can be attached to the protein to allow easy purification by passage over an appropriate affinity column. Isolated proteins can also be physically characterized using such techniques as proteolysis, nuclear magnetic resonance and x-ray crystallography.

For example, supernatants from systems which secrete recombinant protein into culture media can be first concentrated using a commercially available protein concentration filter, for example, an Amicon or Millipore Pellicon ultrafiltration unit. Following the concentration step, the concentrate can be applied to a suitable purification matrix. Alternatively, an anion exchange resin can be employed, for example, a matrix or substrate having pendant diethylaminoethyl (DEAE) groups. The matrices can be acrylamide, agarose, dextran, cellulose or other types commonly employed in protein purification. Alternatively, a cation exchange step can be employed. Suitable cation exchangers include various insoluble matrices comprising sulfopropyl or carboxymethyl groups. Finally, one or more reversed-phase high performance liquid chromatography (RP-HPLC) steps employing hydrophobic RP-HPLC media, e.g., silica gel having pendant methyl or other aliphatic groups, can be employed to further purify a cancer stem cell protein-Fc composition. Some or all of the foregoing purification steps, in various combinations, can also be employed to provide a homogeneous recombinant protein.

Recombinant protein produced in bacterial culture can be isolated, for example, by initial extraction from cell pellets, followed by one or more concentration, salting-out, aqueous ion exchange or size exclusion chromatography steps. High performance liquid chromatography (HPLC) can be employed for final purification steps. Microbial cells employed in expression of a recombinant protein can be disrupted by any convenient method, including freeze-thaw cycling, sonication, mechanical disruption, or use of cell lysing agents.

In Vivo Peptide/Polypeptide Synthesis

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The present invention also contemplates the use of nucleic acid molecules as vehicles for delivering neo-antigenic peptides/polypeptides to the subject in vivo in the form of, e.g., DNA/RNA vaccines (see, e.g., WO2012/159643, and WO2012/159754, hereby incorporated by reference in their entirety).

In one embodiment, the personalized neoplasia vaccine may include separate DNA plasmids encoding, for example, one or more neo-antigenic peptides/polypeptides as identified in according to the invention. As discussed above, the exact choice of expression vectors will depend upon the peptide/polypeptides to be expressed, and is well within the skill of the ordinary artisan. The expected persistence of the DNA constructs (e.g., in an episomal, non-replicating, non-integrated form in the muscle cells) is expected to provide an increased duration of protection.

In another embodiment, the personalized neoplasia vaccine may include separate RNA or cDNA molecules encoding neo-antigenic peptides/polypeptides of the invention.

In another embodiment the personalized neoplasia vaccine may include a viral based vector for use in a human patient such as, for example, and adenovirus system (see, e.g., Baden et al. First-in-human evaluation of the safety and immunogenicity of a recombinant adenovirus serotype 26 HIV-1 Env vaccine (IPCAVD 001). J Infect Dis. 2013 Jan 15;207(2):240-7, hereby incorporated by reference in its entirety).

Pharmaceutical Compositions/Methods of Delivery

The present invention is also directed to pharmaceutical compositions comprising an effective amount of one or more compounds according to the present invention (including a pharmaceutically acceptable salt, thereof), optionally in combination with a pharmaceutically acceptable carrier, excipient or additive.

A "pharmaceutically acceptable derivative or prodrug" means any pharmaceutically acceptable salt, ester, salt of an ester, or other derivative of a compound of this invention which,

upon administration to a recipient, is capable of providing (directly or indirectly) a compound of this invention. Particularly favored derivatives and prodrugs are those that increase the bioavailability of the compounds of this invention when such compounds are administered to a mammal (e.g., by allowing an orally or ocularly administered compound to be more readily absorbed into the blood) or which enhance delivery of the parent compound to a biological compartment (e.g., the retina) relative to the parent species.

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While the tumor specific neo-antigenic peptides of the invention can be administered as the sole active pharmaceutical agent, they can also be used in combination with one or more other agents and/or adjuvants. When administered as a combination, the therapeutic agents can be formulated as separate compositions that are given at the same time or different times, or the therapeutic agents can be given as a single composition.

The tumor specific neo-antigenic peptides of the present invention may be administered by injection, orally, parenterally, by inhalation spray, rectally, vaginally, or topically in dosage unit formulations containing conventional pharmaceutically acceptable carriers, adjuvants, and vehicles. The term parenteral as used herein includes, into a lymph node or nodes, subcutaneous, intravenous, intramuscular, intrasternal, infusion techniques, intraperitoneally, eye or ocular, intravitreal, intrabuccal, transdermal, intranasal, into the brain, including intracranial and intradural, into the joints, including ankles, knees, hips, shoulders, elbows, wrists, directly into tumors, and the like, and in suppository form.

The pharmaceutically active compounds of this invention can be processed in accordance with conventional methods of pharmacy to produce medicinal agents for administration to patients, including humans and other mammals.

Modifications of the active compound can affect the solubility, bioavailability and rate of metabolism of the active species, thus providing control over the delivery of the active species. This can easily be assessed by preparing the derivative and testing its activity according to known methods well within the routine practitioner's skill in the art.

Pharmaceutical compositions based upon these chemical compounds comprise the abovedescribed tumor specific neo-antigenic peptides in a therapeutically effective amount for treating

diseases and conditions (e.g., a neoplasia/tumor), which have been described herein, optionally in combination with a pharmaceutically acceptable additive, carrier and/or excipient. One of ordinary skill in the art will recognize that a therapeutically effective amount of one of more compounds according to the present invention will vary with the infection or condition to be treated, its severity, the treatment regimen to be employed, the pharmacokinetics of the agent used, as well as the patient (animal or human) treated.

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To prepare the pharmaceutical compositions according to the present invention, a therapeutically effective amount of one or more of the compounds according to the present invention is preferably intimately admixed with a pharmaceutically acceptable carrier according to conventional pharmaceutical compounding techniques to produce a dose. A carrier may take a wide variety of forms depending on the form of preparation desired for administration, e.g., ocular, oral, topical or parenteral, including gels, creams ointments, lotions and time released implantable preparations, among numerous others. In preparing pharmaceutical compositions in oral dosage form, any of the usual pharmaceutical media may be used. Thus, for liquid oral preparations such as suspensions, elixirs and solutions, suitable carriers and additives including water, glycols, oils, alcohols, flavoring agents, preservatives, coloring agents and the like may be used. For solid oral preparations such as powders, tablets, capsules, and for solid preparations such as suppositories, suitable carriers and additives including starches, sugar carriers, such as dextrose, mannitol, lactose and related carriers, diluents, granulating agents, lubricants, binders, disintegrating agents and the like may be used. If desired, the tablets or capsules may be entericcoated or sustained release by standard techniques.

The active compound is included in the pharmaceutically acceptable carrier or diluent in an amount sufficient to deliver to a patient a therapeutically effective amount for the desired indication, without causing serious toxic effects in the patient treated.

Oral compositions will generally include an inert diluent or an edible carrier. They may be enclosed in gelatin capsules or compressed into tablets. For the purpose of oral therapeutic administration, the active compound or its prodrug derivative can be incorporated with excipients and used in the form of tablets, troches, or capsules. Pharmaceutically compatible binding agents, and/or adjuvant materials can be included as part of the composition.

The tablets, pills, capsules, troches and the like can contain any of the following ingredients, or compounds of a similar nature: a binder such as microcrystalline cellulose, gum tragacanth or gelatin; an excipient such as starch or lactose, a dispersing agent such as alginic acid or corn starch; a lubricant such as magnesium stearate; a glidant such as colloidal silicon dioxide; a sweetening agent such as sucrose or saccharin; or a flavoring agent such as peppermint, methyl salicylate, or orange flavoring. When the dosage unit form is a capsule, it can contain, in addition to material-of the above type, a liquid carrier such as a fatty oil. In addition, dosage unit forms can contain various other materials which modify the physical form of the dosage unit, for example, coatings of sugar, shellac, or enteric agents.

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Formulations of the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets or tablets each containing a predetermined amount of the active ingredient; as a powder or granules; as a solution or a suspension in an aqueous liquid or a non-aqueous liquid; or as an oil-in-water liquid emulsion or a water-in-oil emulsion and as a bolus, etc.

A tablet may be made by compression or molding, optionally with one or more accessory ingredients. Compressed tablets may be prepared by compressing in a suitable machine the active ingredient in a free-flowing form such as a powder or granules, optionally mixed with a binder, lubricant, inert diluent, preservative, surface-active or dispersing agent. Molded tablets may be made by molding in a suitable machine a mixture of the powdered compound moistened with an inert liquid diluent. The tablets optionally may be coated or scored and may be formulated so as to provide slow or controlled release of the active ingredient therein.

Methods of formulating such slow or controlled release compositions of pharmaceutically active ingredients, are known in the art and described in several issued US Patents, some of which include, but are not limited to, US Patent Nos. 3,870,790; 4,226,859; 4,369,172; 4,842,866 and 5,705,190, the disclosures of which are incorporated herein by reference in their entireties. Coatings can be used for delivery of compounds to the intestine (see, e.g., U.S. Patent Nos. 6,638,534, 5,541,171, 5,217,720, and 6,569,457, and references cited therein).

The active compound or pharmaceutically acceptable salt thereof may also be administered as a component of an elixir, suspension, syrup, wafer, chewing gum or the like. A

syrup may contain, in addition to the active compounds, sucrose or fructose as a sweetening agent and certain preservatives, dyes and colorings and flavors.

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Solutions or suspensions used for ocular, parenteral, intradermal, subcutaneous, or topical application can include the following components: a sterile diluent such as water for injection, saline solution, fixed oils, polyethylene glycols, glycerine, propylene glycol or other synthetic solvents; antibacterial agents such as benzyl alcohol or methyl parabens; antioxidants such as ascorbic acid or sodium bisulfite; chelating agents such as ethylenediaminetetraacetic acid; buffers such as acetates, citrates or phosphates and agents for the adjustment of tonicity such as sodium chloride or dextrose.

In one embodiment, the active compounds are prepared with carriers that will protect the compound against rapid elimination from the body, such as a controlled release formulation, including implants and microencapsulated delivery systems. Biodegradable, biocompatible polymers can be used, such as ethylene vinyl acetate, polyanhydrides, polyglycolic acid, collagen, polyorthoesters, polylactic acid, and polylactic-co-glycolic acid (PLGA). Methods for preparation of such formulations will be apparent to those skilled in the art.

A skilled artisan will recognize that in addition to tablets, other dosage forms can be formulated to provide slow or controlled release of the active ingredient. Such dosage forms include, but are not limited to, capsules, granulations and gel-caps.

Liposomal suspensions may also be pharmaceutically acceptable carriers. These may be prepared according to methods known to those skilled in the art. For example, liposomal formulations may be prepared by dissolving appropriate lipid(s) in an inorganic solvent that is then evaporated, leaving behind a thin film of dried lipid on the surface of the container. An aqueous solution of the active compound are then introduced into the container. The container is then swirled by hand to free lipid material from the sides of the container and to disperse lipid aggregates, thereby forming the liposomal suspension. Other methods of preparation well known by those of ordinary skill may also be used in this aspect of the present invention.

The formulations may conveniently be presented in unit dosage form and may be prepared by conventional pharmaceutical techniques. Such techniques include the step of

bringing into association the active ingredient and the pharmaceutical carrier(s) or excipient(s). In general, the formulations are prepared by uniformly and intimately bringing into association the active ingredient with liquid carriers or finely divided solid carriers or both, and then, if necessary, shaping the product.

Formulations and compositions suitable for topical administration in the mouth include lozenges comprising the ingredients in a flavored basis, usually sucrose and acacia or tragacanth; pastilles comprising the active ingredient in an inert basis such as gelatin and glycerin, or sucrose and acacia; and mouthwashes comprising the ingredient to be administered in a suitable liquid carrier.

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Formulations suitable for topical administration to the skin may be presented as ointments, creams, gels and pastes comprising the ingredient to be administered in a pharmaceutical acceptable carrier. A preferred topical delivery system is a transdermal patch containing the ingredient to be administered.

Formulations for rectal administration may be presented as a suppository with a suitable base comprising, for example, cocoa butter or a salicylate.

Formulations suitable for nasal administration, wherein the carrier is a solid, include a coarse powder having a particle size, for example, in the range of 20 to 500 microns which is administered in the manner in which snuff is administered, i.e., by rapid inhalation through the nasal passage from a container of the powder held close up to the nose. Suitable formulations, wherein the carrier is a liquid, for administration, as for example, a nasal spray or as nasal drops, include aqueous or oily solutions of the active ingredient.

Formulations suitable for vaginal administration may be presented as pessaries, tampons, creams, gels, pastes, foams or spray formulations containing in addition to the active ingredient such carriers as are known in the art to be appropriate.

The parenteral preparation can be enclosed in ampoules, disposable syringes or multiple dose vials made of glass or plastic. If administered intravenously, preferred carriers include, for example, physiological saline or phosphate buffered saline (PBS).

For parenteral formulations, the carrier will usually comprise sterile water or aqueous sodium chloride solution, though other ingredients including those which aid dispersion may be included. Of course, where sterile water is to be used and maintained as sterile, the compositions and carriers will also be sterilized. Injectable suspensions may also be prepared, in which case appropriate liquid carriers, suspending agents and the like may be employed.

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Formulations suitable for parenteral administration include aqueous and non-aqueous sterile injection solutions which may contain antioxidants, buffers, bacteriostats and solutes which render the formulation isotonic with the blood of the intended recipient; and aqueous and non-aqueous sterile suspensions which may include suspending agents and thickening agents. The formulations may be presented in unit-dose or multi-dose containers, for example, sealed ampules and vials, and may be stored in a freeze-dried (lyophilized) condition requiring only the addition of the sterile liquid carrier, for example, water for injections, immediately prior to use. Extemporaneous injection solutions and suspensions may be prepared from sterile powders, granules and tablets of the kind previously described.

Administration of the active compound may range from continuous (intravenous drip) to several oral administrations per day (for example, Q.I.D.) and may include oral, topical, eye or ocular, parenteral, intramuscular, intravenous, sub-cutaneous, transdermal (which may include a penetration enhancement agent), buccal and suppository administration, among other routes of administration, including through an eye or ocular route.

Application of the subject therapeutics may be local, so as to be administered at the site of interest. Various techniques can be used for providing the subject compositions at the site of interest, such as injection, use of catheters, trocars, projectiles, pluronic gel, stents, sustained drug release polymers or other device which provides for internal access. Where an organ or tissue is accessible because of removal from the patient, such organ or tissue may be bathed in a medium containing the subject compositions, the subject compositions may be painted onto the organ, or may be applied in any convenient way.

The tumor specific neo-antigenic peptides may be administered through a device suitable for the controlled and sustained release of a composition effective in obtaining a desired local or systemic physiological or pharmacological effect. The method includes positioning the sustained

released drug delivery system at an area wherein release of the agent is desired and allowing the agent to pass through the device to the desired area of treatment.

The tumor specific neo-antigenic peptides may be utilized in combination with at least one known other therapeutic agent, or a pharmaceutically acceptable salt of said agent. Examples of known therapeutic agents which can be used for combination therapy include, but are not limited to, corticosteroids (e.g., cortisone, prednisone, dexamethasone), non-steroidal anti-inflammatory drugs (NSAIDS) (e.g., ibuprofen, celecoxib, aspirin, indomethicin, naproxen), alkylating agents such as busulfan, cis-platin, mitomycin C, and carboplatin; antimitotic agents such as colchicine, vinblastine, paclitaxel, and docetaxel; topo I inhibitors such as camptothecin and topotecan; topo II inhibitors such as doxorubicin and etoposide; and/or RNA/DNA antimetabolites such as 5-azacytidine, 5-fluorouracil and methotrexate; DNA antimetabolites such as 5-fluoro-2'-deoxy-uridine, ara-C, hydroxyurea and thioguanine; antibodies such as Herceptin® and Rituxan®.

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It should be understood that in addition to the ingredients particularly mentioned above, the formulations of the present invention may include other agents conventional in the art having regard to the type of formulation in question, for example, those suitable for oral administration may include flavoring agents.

In certain pharmaceutical dosage forms, the pro-drug form of the compounds may be preferred. One of ordinary skill in the art will recognize how to readily modify the present compounds to pro-drug forms to facilitate delivery of active compounds to a targeted site within the host organism or patient. The routine practitioner also will take advantage of favorable pharmacokinetic parameters of the pro-drug forms, where applicable, in delivering the present compounds to a targeted site within the host organism or patient to maximize the intended effect of the compound.

Preferred prodrugs include derivatives where a group which enhances aqueous solubility or active transport through the gut membrane is appended to the structure of formulae described herein. See, e.g., Alexander, J. et al. *Journal of Medicinal Chemistry* **1988**, *31*, 318-322; Bundgaard, H. Design of Prodrugs; Elsevier: Amsterdam, 1985; pp 1-92; Bundgaard, H.; Nielsen, N. M. Journal of Medicinal Chemistry 1987, 30, 451-454; Bundgaard, H. A Textbook

of Drug Design and Development; Harwood Academic Publ.: Switzerland, 1991; pp 113-191; Digenis, G. A. et al. Handbook of Experimental Pharmacology 1975, 28, 86-112; Friis, G. J.; Bundgaard, H. A Textbook of Drug Design and Development; 2 ed.; Overseas Publ.: Amsterdam, 1996; pp 351-385; Pitman, I. H. Medicinal Research Reviews 1981, 1, 189-214. The prodrug forms may be active themselves, or may be those such that when metabolized after administration provide the active therapeutic agent in vivo.

Pharmaceutically acceptable salt forms may be the preferred chemical form of compounds according to the present invention for inclusion in pharmaceutical compositions according to the present invention.

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The present compounds or their derivatives, including prodrug forms of these agents, can be provided in the form of pharmaceutically acceptable salts. As used herein, the term pharmaceutically acceptable salts or complexes refers to appropriate salts or complexes of the active compounds according to the present invention which retain the desired biological activity of the parent compound and exhibit limited toxicological effects to normal cells. Nonlimiting examples of such salts are (a) acid addition salts formed with inorganic acids (for example, hydrochloric acid, hydrobromic acid, sulfuric acid, phosphoric acid, nitric acid, and the like), and salts formed with organic acids such as acetic acid, oxalic acid, tartaric acid, succinic acid, malic acid, ascorbic acid, benzoic acid, tannic acid, pamoic acid, alginic acid, and polyglutamic acid, among others; (b) base addition salts formed with metal cations such as zinc, calcium, sodium, potassium, and the like, among numerous others.

The compounds herein are commercially available or can be synthesized. As can be appreciated by the skilled artisan, further methods of synthesizing the compounds of the formulae herein will be evident to those of ordinary skill in the art. Additionally, the various synthetic steps may be performed in an alternate sequence or order to give the desired compounds. Synthetic chemistry transformations and protecting group methodologies (protection and deprotection) useful in synthesizing the compounds described herein are known in the art and include, for example, those such as described in R. Larock, *Comprehensive Organic Transformations*, 2nd. Ed., Wiley-VCH Publishers (1999); T.W. Greene and P.G.M. Wuts, *Protective Groups in Organic Synthesis*, 3rd. Ed., John Wiley and Sons (1999); L. Fieser

and M. Fieser, *Fieser and Fieser's Reagents for Organic Synthesis*, John Wiley and Sons (1999); and L. Paquette, ed., *Encyclopedia of Reagents for Organic Synthesis*, John Wiley and Sons (1995), and subsequent editions thereof.

The additional agents that may be included with the tumor specific neo-antigenic peptides of this invention may contain one or more asymmetric centers and thus occur as racemates and racemic mixtures, single enantiomers, individual diastereomers and diastereomeric mixtures. All such isomeric forms of these compounds are expressly included in the present invention. The compounds of this invention may also be represented in multiple tautomeric forms, in such instances, the invention expressly includes all tautomeric forms of the compounds described herein (e.g., alkylation of a ring system may result in alkylation at multiple sites, the invention expressly includes all such reaction products). All such isomeric forms of such compounds are expressly included in the present invention. All crystal forms of the compounds described herein are expressly included in the present invention.

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Preferred unit dosage formulations are those containing a daily dose or unit, daily subdose, as hereinabove recited, or an appropriate fraction thereof, of the administered ingredient.

The dosage regimen for treating a disorder or a disease with the tumor specific neoantigenic peptides of this invention and/or compositions of this invention is based on a variety of factors, including the type of disease, the age, weight, sex, medical condition of the patient, the severity of the condition, the route of administration, and the particular compound employed. Thus, the dosage regimen may vary widely, but can be determined routinely using standard methods.

The amounts and dosage regimens administered to a subject will depend on a number of factors, such as the mode of administration, the nature of the condition being treated, the body weight of the subject being treated and the judgment of the prescribing physician.

The amount of compound included within therapeutically active formulations according to the present invention is an effective amount for treating the disease or condition. In general, a therapeutically effective amount of the present preferred compound in dosage form usually ranges from slightly less than about 0.025 mg/kg/day to about 2.5 g/kg/day, preferably about 0.1

mg/kg/day to about 100 mg/kg/day of the patient or considerably more, depending upon the compound used, the condition or infection treated and the route of administration, although exceptions to this dosage range may be contemplated by the present invention. In its most preferred form, compounds according to the present invention are administered in amounts ranging from about 1 mg/kg/day to about 100 mg/kg/day. The dosage of the compound will depend on the condition being treated, the particular compound, and other clinical factors such as weight and condition of the patient and the route of administration of the compound. It is to be understood that the present invention has application for both human and veterinary use.

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For oral administration to humans, a dosage of between approximately 0.1 to 100 mg/kg/day, preferably between approximately 1 and 100 mg/kg/day, is generally sufficient.

Where drug delivery is systemic rather than topical, this dosage range generally produces effective blood level concentrations of active compound ranging from less than about 0.04 to about 400 micrograms/cc or more of blood in the patient.

The compound is conveniently administered in any suitable unit dosage form, including but not limited to one containing 0.001 to 3000 mg, preferably 0.05 to 500 mg of active ingredient per unit dosage form. An oral dosage of 10-250 mg is usually convenient.

The concentration of active compound in the drug composition will depend on absorption, distribution, inactivation, and excretion rates of the drug as well as other factors known to those of skill in the art. It is to be noted that dosage values will also vary with the severity of the condition to be alleviated. It is to be further understood that for any particular subject, specific dosage regimens should be adjusted over time according to the individual need and the professional judgment of the person administering or supervising the administration of the compositions, and that the concentration ranges set forth herein are exemplary only and are not intended to limit the scope or practice of the claimed composition. The active ingredient may be administered at once, or may be divided into a number of smaller doses to be administered at varying intervals of time.

In certain embodiments, the compound is administered once daily; in other embodiments, the compound is administered twice daily; in yet other embodiments, the compound is

administered once every two days, once every three days, once every four days, once every five days, once every six days, once every seven days, once every two weeks, once every three weeks, once every four weeks, once every two months, once every six months, or once per year. The dosing interval can be adjusted according to the needs of individual patients. For longer intervals of administration, extended release or depot formulations can be used.

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The compounds of the invention can be used to treat diseases and disease conditions that are acute, and may also be used for treatment of chronic conditions. In certain embodiments, the compounds of the invention are administered for time periods exceeding two weeks, three weeks, one month, two months, three months, four months, five months, six months, one year, two years, three years, four years, or five years, ten years, or fifteen years; or for example, any time period range in days, months or years in which the low end of the range is any time period between 14 days and 15 years and the upper end of the range is between 15 days and 20 years (e.g., 4 weeks and 15 years, 6 months and 20 years). In some cases, it may be advantageous for the compounds of the invention to be administered for the remainder of the patient's life. In preferred embodiments, the patient is monitored to check the progression of the disease or disorder, and the dose is adjusted accordingly. In preferred embodiments, treatment according to the invention is effective for at least two weeks, three weeks, one month, two months, three months, four months, five months, six months, one year, two years, three years, four years, or five years, ten years, fifteen years, twenty years, or for the remainder of the subject's life.

The invention provides for pharmaceutical compositions containing at least one tumor specific neo-antigen described herein. In embodiments, the pharmaceutical compositions contain a pharmaceutically acceptable carrier, excipient, or diluent, which includes any pharmaceutical agent that does not itself induce the production of an immune response harmful to a subject receiving the composition, and which may be administered without undue toxicity. As used herein, the term "pharmaceutically acceptable" means being approved by a regulatory agency of the Federal or a state government or listed in the U.S. Pharmacopia, European Pharmacopia or other generally recognized pharmacopia for use in mammals, and more particularly in humans. These compositions can be useful for treating and/or preventing viral infection and/or autoimmune disease.

A thorough discussion of pharmaceutically acceptable carriers, diluents, and other excipients is presented in *Remington's Pharmaceutical Sciences* (17th ed., Mack Publishing Company) and *Remington: The Science and Practice of Pharmacy* (21st ed., Lippincott Williams & Wilkins), which are hereby incorporated by reference. The formulation of the pharmaceutical composition should suit the mode of administration. In embodiments, the pharmaceutical composition is suitable for administration to humans, and can be sterile, non-particulate and/or non-pyrogenic.

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Pharmaceutically acceptable carriers, excipients, or diluents include, but are not limited, to saline, buffered saline, dextrose, water, glycerol, ethanol, sterile isotonic aqueous buffer, and combinations thereof.

Wetting agents, emulsifiers and lubricants, such as sodium lauryl sulfate and magnesium stearate, as well as coloring agents, release agents, coating agents, sweetening, flavoring and perfuming agents, preservatives, and antioxidants can also be present in the compositions.

Examples of pharmaceutically-acceptable antioxidants include, but are not limited to: (1) water soluble antioxidants, such as ascorbic acid, cysteine hydrochloride, sodium bisulfate, sodium metabisulfite, sodium sulfite and the like; (2) oil-soluble antioxidants, such as ascorbyl palmitate, butylated hydroxyanisole (BHA), butylated hydroxytoluene (BHT), lecithin, propyl gallate, alpha-tocopherol, and the like; and (3) metal chelating agents, such as citric acid, ethylenediamine tetraacetic acid (EDTA), sorbitol, tartaric acid, phosphoric acid, and the like.

In embodiments, the pharmaceutical composition is provided in a solid form, such as a lyophilized powder suitable for reconstitution, a liquid solution, suspension, emulsion, tablet, pill, capsule, sustained release formulation, or powder.

In embodiments, the pharmaceutical composition is supplied in liquid form, for example, in a sealed container indicating the quantity and concentration of the active ingredient in the pharmaceutical composition. In related embodiments, the liquid form of the pharmaceutical composition is supplied in a hermetically sealed container.

Methods for formulating the pharmaceutical compositions of the present invention are conventional and well known in the art (see Remington and Remington's). One of skill in the art

can readily formulate a pharmaceutical composition having the desired characteristics (e.g., route of administration, biosafety, and release profile).

Methods for preparing the pharmaceutical compositions include the step of bringing into association the active ingredient with a pharmaceutically acceptable carrier and, optionally, one or more accessory ingredients. The pharmaceutical compositions can be prepared by uniformly and intimately bringing into association the active ingredient with liquid carriers, or finely divided solid carriers, or both, and then, if necessary, shaping the product. Additional methodology for preparing the pharmaceutical compositions, including the preparation of multilayer dosage forms, are described in *Ansel's Pharmaceutical Dosage Forms and Drug Delivery Systems* (9th ed., Lippincott Williams & Wilkins), which is hereby incorporated by reference.

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Pharmaceutical compositions suitable for oral administration can be in the form of capsules, cachets, pills, tablets, lozenges (using a flavored basis, usually sucrose and acacia or tragacanth), powders, granules, or as a solution or a suspension in an aqueous or non-aqueous liquid, or as an oil-in-water or water-in-oil liquid emulsion, or as an elixir or syrup, or as pastilles (using an inert base, such as gelatin and glycerin, or sucrose and acacia) and/or as mouth washes and the like, each containing a predetermined amount of a compound(s) described herein, a derivative thereof, or a pharmaceutically acceptable salt or prodrug thereof as the active ingredient(s). The active ingredient can also be administered as a bolus, electuary, or paste.

In solid dosage forms for oral administration (e.g., capsules, tablets, pills, dragees, powders, granules and the like), the active ingredient is mixed with one or more pharmaceutically acceptable carriers, excipients, or diluents, such as sodium citrate or dicalcium phosphate, and/or any of the following: (1) fillers or extenders, such as starches, lactose, sucrose, glucose, mannitol, and/or silicic acid; (2) binders, such as, for example, carboxymethylcellulose, alginates, gelatin, polyvinyl pyrrolidone, sucrose and/or acacia; (3) humectants, such as glycerol; (4) disintegrating agents, such as agar-agar, calcium carbonate, potato or tapioca starch, alginic acid, certain silicates, and sodium carbonate; (5) solution retarding agents, such as paraffin; (6) absorption accelerators, such as quaternary ammonium compounds; (7) wetting agents, such as, for example, acetyl alcohol and glycerol monostearate; (8) absorbents, such as kaolin and

bentonite clay; (9) lubricants, such a talc, calcium stearate, magnesium stearate, solid polyethylene glycols, sodium lauryl sulfate, and mixtures thereof; and (10) coloring agents. In the case of capsules, tablets, and pills, the pharmaceutical compositions can also comprise buffering agents. Solid compositions of a similar type can also be prepared using fillers in soft and hard-filled gelatin capsules, and excipients such as lactose or milk sugars, as well as high molecular weight polyethylene glycols and the like.

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A tablet can be made by compression or molding, optionally with one or more accessory ingredients. Compressed tablets can be prepared using binders (for example, gelatin or hydroxypropylmethyl cellulose), lubricants, inert diluents, preservatives, disintegrants (for example, sodium starch glycolate or cross-linked sodium carboxymethyl cellulose), surface-actives, and/ or dispersing agents. Molded tablets can be made by molding in a suitable machine a mixture of the powdered active ingredient moistened with an inert liquid diluent.

The tablets and other solid dosage forms, such as dragees, capsules, pills, and granules, can optionally be scored or prepared with coatings and shells, such as enteric coatings and other coatings well known in the art.

In some embodiments, in order to prolong the effect of an active ingredient, it is desirable to slow the absorption of the compound from subcutaneous or intramuscular injection. This can be accomplished by the use of a liquid suspension of crystalline or amorphous material having poor water solubility. The rate of absorption of the active ingredient then depends upon its rate of dissolution which, in turn, can depend upon crystal size and crystalline form. Alternatively, delayed absorption of a parenterally-administered active ingredient is accomplished by dissolving or suspending the compound in an oil vehicle. In addition, prolonged absorption of the injectable pharmaceutical form can be brought about by the inclusion of agents that delay absorption such as aluminum monostearate and gelatin.

Controlled release parenteral compositions can be in form of aqueous suspensions, microspheres, microcapsules, magnetic microspheres, oil solutions, oil suspensions, emulsions, or the active ingredient can be incorporated in biocompatible carrier(s), liposomes, nanoparticles, implants or infusion devices.

Materials for use in the preparation of microspheres and/or microcapsules include biodegradable/bioerodible polymers such as polyglactin, poly-(isobutyl cyanoacrylate), poly(2-hydroxyethyl-L-glutamine) and poly(lactic acid).

Biocompatible carriers which can be used when formulating a controlled release parenteral formulation include carbohydrates such as dextrans, proteins such as albumin, lipoproteins or antibodies.

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Materials for use in implants can be non-biodegradable, e.g., polydimethylsiloxane, or biodegradable such as, e.g., poly(caprolactone), poly(lactic acid), poly(glycolic acid) or poly(ortho esters).

In embodiments, the active ingredient(s) are administered by aerosol. This is accomplished by preparing an aqueous aerosol, liposomal preparation, or solid particles containing the compound. A nonaqueous (e.g., fluorocarbon propellant) suspension can be used. The pharmaceutical composition can also be administered using a sonic nebulizer, which would minimize exposing the agent to shear, which can result in degradation of the compound.

Ordinarily, an aqueous aerosol is made by formulating an aqueous solution or suspension of the active ingredient(s) together with conventional pharmaceutically-acceptable carriers and stabilizers. The carriers and stabilizers vary with the requirements of the particular compound, but typically include nonionic surfactants (Tweens, Pluronics, or polyethylene glycol), innocuous proteins like serum albumin, sorbitan esters, oleic acid, lecithin, amino acids such as glycine, buffers, salts, sugars or sugar alcohols. Aerosols generally are prepared from isotonic solutions.

Dosage forms for topical or transdermal administration of an active ingredient(s) includes powders, sprays, ointments, pastes, creams, lotions, gels, solutions, patches and inhalants. The active ingredient(s) can be mixed under sterile conditions with a pharmaceutically acceptable carrier, and with any preservatives, buffers, or propellants as appropriate.

Transdermal patches suitable for use in the present invention are disclosed in *Transdermal Drug Delivery: Developmental Issues and Research Initiatives* (Marcel Dekker Inc., 1989) and U.S. Pat. Nos. 4,743,249, 4,906,169, 5,198,223, 4,816,540, 5,422,119, 5,023,084, which are hereby incorporated by reference. The transdermal patch can also be any transdermal

patch well known in the art, including transscrotal patches. Pharmaceutical compositions in such transdermal patches can contain one or more absorption enhancers or skin permeation enhancers well known in the art (*see*, *e.g.*, U.S. Pat. Nos. 4,379,454 and 4,973,468, which are hereby incorporated by reference). Transdermal therapeutic systems for use in the present invention can be based on iontophoresis, diffusion, or a combination of these two effects.

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Transdermal patches have the added advantage of providing controlled delivery of active ingredient(s) to the body. Such dosage forms can be made by dissolving or dispersing the active ingredient(s) in a proper medium. Absorption enhancers can also be used to increase the flux of the active ingredient across the skin. The rate of such flux can be controlled by either providing a rate controlling membrane or dispersing the active ingredient(s) in a polymer matrix or gel.

Such pharmaceutical compositions can be in the form of creams, ointments, lotions, liniments, gels, hydrogels, solutions, suspensions, sticks, sprays, pastes, plasters and other kinds of transdermal drug delivery systems. The compositions can also include pharmaceutically acceptable carriers or excipients such as emulsifying agents, antioxidants, buffering agents, preservatives, humectants, penetration enhancers, chelating agents, gel-forming agents, ointment bases, perfumes, and skin protective agents.

Examples of emulsifying agents include, but are not limited to, naturally occurring gums, e.g. gum acacia or gum tragacanth, naturally occurring phosphatides, e.g. soybean lecithin and sorbitan monooleate derivatives.

Examples of antioxidants include, but are not limited to, butylated hydroxy anisole (BHA), ascorbic acid and derivatives thereof, tocopherol and derivatives thereof, and cysteine.

Examples of preservatives include, but are not limited to, parabens, such as methyl or propyl p-hydroxybenzoate and benzalkonium chloride.

Examples of humectants include, but are not limited to, glycerin, propylene glycol, sorbitol and urea.

Examples of penetration enhancers include, but are not limited to, propylene glycol, DMSO, triethanolamine, N,N-dimethylacetamide, N,N-dimethylformamide, 2-pyrrolidone and

derivatives thereof, tetrahydrofurfuryl alcohol, propylene glycol, diethylene glycol monoethyl or monomethyl ether with propylene glycol monolaurate or methyl laurate, eucalyptol, lecithin, Transcutol[®], and Azone[®].

Examples of chelating agents include, but are not limited to, sodium EDTA, citric acid and phosphoric acid.

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Examples of gel forming agents include, but are not limited to, Carbopol, cellulose derivatives, bentonite, alginates, gelatin and polyvinylpyrrolidone.

In addition to the active ingredient(s), the ointments, pastes, creams, and gels of the present invention can contain excipients, such as animal and vegetable fats, oils, waxes, paraffins, starch, tragacanth, cellulose derivatives, polyethylene glycols, silicones, bentonites, silicic acid, talc and zinc oxide, or mixtures thereof.

Powders and sprays can contain excipients such as lactose, talc, silicic acid, aluminum hydroxide, calcium silicates and polyamide powder, or mixtures of these substances. Sprays can additionally contain customary propellants, such as chlorofluorohydrocarbons, and volatile unsubstituted hydrocarbons, such as butane and propane.

Injectable depot forms are made by forming microencapsule matrices of compound(s) of the invention in biodegradable polymers such as polylactide-polyglycolide. Depending on the ratio of compound to polymer, and the nature of the particular polymer employed, the rate of compound release can be controlled. Examples of other biodegradable polymers include poly(orthoesters) and poly(anhydrides). Depot injectable formulations are also prepared by entrapping the drug in liposomes or microemulsions which are compatible with body tissue.

Subcutaneous implants are well known in the art and are suitable for use in the present invention. Subcutaneous implantation methods are preferably non-irritating and mechanically resilient. The implants can be of matrix type, of reservoir type, or hybrids thereof. In matrix type devices, the carrier material can be porous or non-porous, solid or semi-solid, and permeable or impermeable to the active compound or compounds. The carrier material can be biodegradable or may slowly erode after administration. In some instances, the matrix is non-degradable but instead relies on the diffusion of the active compound through the matrix for the

carrier material to degrade. Alternative subcutaneous implant methods utilize reservoir devices where the active compound or compounds are surrounded by a rate controlling membrane, e.g., a membrane independent of component concentration (possessing zero-order kinetics). Devices consisting of a matrix surrounded by a rate controlling membrane also suitable for use.

Both reservoir and matrix type devices can contain materials such as polydimethylsiloxane, such as SilasticTM, or other silicone rubbers. Matrix materials can be insoluble polypropylene, polyethylene, polyvinyl chloride, ethylvinyl acetate, polystyrene and polymethacrylate, as well as glycerol esters of the glycerol palmitostearate, glycerol stearate, and glycerol behenate type. Materials can be hydrophobic or hydrophilic polymers and optionally contain solubilizing agents.

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Subcutaneous implant devices can be slow-release capsules made with any suitable polymer, e.g., as described in U.S. Pat. Nos. 5,035,891 and 4,210,644, which are hereby incorporated by reference.

In general, at least four different approaches are applicable in order to provide rate control over the release and transdermal permeation of a drug compound. These approaches are: membrane-moderated systems, adhesive diffusion-controlled systems, matrix dispersion-type systems and microreservoir systems. It is appreciated that a controlled release percutaneous and/or topical composition can be obtained by using a suitable mixture of these approaches.

In a membrane-moderated system, the active ingredient is present in a reservoir which is totally encapsulated in a shallow compartment molded from a drug-impermeable laminate, such as a metallic plastic laminate, and a rate-controlling polymeric membrane such as a microporous or a non-porous polymeric membrane, e.g., ethylene-vinyl acetate copolymer. The active ingredient is released through the rate controlling polymeric membrane. In the drug reservoir, the active ingredient can either be dispersed in a solid polymer matrix or suspended in an unleachable, viscous liquid medium such as silicone fluid. On the external surface of the polymeric membrane, a thin layer of an adhesive polymer is applied to achieve an intimate contact of the transdermal system with the skin surface. The adhesive polymer is preferably a polymer which is hypoallergenic and compatible with the active drug substance.

In an adhesive diffusion-controlled system, a reservoir of the active ingredient is formed by directly dispersing the active ingredient in an adhesive polymer and then by, e.g., solvent casting, spreading the adhesive containing the active ingredient onto a flat sheet of substantially drug-impermeable metallic plastic backing to form a thin drug reservoir layer.

A matrix dispersion-type system is characterized in that a reservoir of the active ingredient is formed by substantially homogeneously dispersing the active ingredient in a hydrophilic or lipophilic polymer matrix. The drug-containing polymer is then molded into disc with a substantially well-defined surface area and controlled thickness. The adhesive polymer is spread along the circumference to form a strip of adhesive around the disc.

A microreservoir system can be considered as a combination of the reservoir and matrix dispersion type systems. In this case, the reservoir of the active substance is formed by first suspending the drug solids in an aqueous solution of water-soluble polymer and then dispersing the drug suspension in a lipophilic polymer to form a multiplicity of unleachable, microscopic spheres of drug reservoirs.

Any of the above-described controlled release, extended release, and sustained release compositions can be formulated to release the active ingredient in about 30 minutes to about 1 week, in about 30 minutes to about 72 hours, in about 30 minutes to 24 hours, in about 30 minutes to 12 hours, in about 30 minutes to 6 hours, in about 30 minutes to 4 hours, and in about 3 hours to 10 hours. In embodiments, an effective concentration of the active ingredient(s) is sustained in a subject for 4 hours, 6 hours, 8 hours, 10 hours, 12 hours, 16 hours, 24 hours, 48 hours, 72 hours, or more after administration of the pharmaceutical compositions to the subject.

Dosages

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When the agents described herein are administered as pharmaceuticals to humans or animals, they can be given per se or as a pharmaceutical composition containing active ingredient in combination with a pharmaceutically acceptable carrier, excipient, or diluent.

Actual dosage levels and time course of administration of the active ingredients in the pharmaceutical compositions of the invention can be varied so as to obtain an amount of the active ingredient which is effective to achieve the desired therapeutic response for a particular patient, composition, and mode of administration, without being toxic to the patient. Generally, agents or pharmaceutical compositions of the invention are administered in an amount sufficient to reduce or eliminate symptoms associated with viral infection and/or autoimmune disease.

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Exemplary dose ranges include 0.01 mg to 250 mg per day, 0.01 mg to 100 mg per day, 1 mg to 100 mg per day, 10 mg to 100 mg per day, 1 mg to 10 mg per day, and 0.01 mg to 10 mg per day. A preferred dose of an agent is the maximum that a patient can tolerate and not develop serious or unacceptable side effects. In embodiments, the agent is administered at a concentration of about 10 micrograms to about 100 mg per kilogram of body weight per day, about 0.1 to about 10 mg/kg per day, or about 1.0 mg to about 10 mg/kg of body weight per day.

In embodiments, the pharmaceutical composition comprises an agent in an amount ranging between 1 and 10 mg, such as 1, 2, 3, 4, 5, 6, 7, 8, 9, or 10 mg.

In embodiments, the therapeutically effective dosage produces a serum concentration of an agent of from about 0.1 ng/ml to about 50-100 μg/ml. The pharmaceutical compositions typically should provide a dosage of from about 0.001 mg to about 2000 mg of compound per kilogram of body weight per day. For example, dosages for systemic administration to a human patient can range from 1-10 μg/kg, 20-80 μg/kg, 5-50 μg/kg, 75-150 μg/kg, 100-500 μg/kg, 250-750 μg/kg, 500-1000 μg/kg, 1-10 mg/kg, 5-50 mg/kg, 25-75 mg/kg, 50-100 mg/kg, 100-250 mg/kg, 50-100 mg/kg, 250-500 mg/kg, 500-750 mg/kg, 750-1000 mg/kg, 1000-1500 mg/kg, 1500-2000 mg/kg, 5 mg/kg, 20 mg/kg, 50 mg/kg, 100 mg/kg, 500 mg/kg, 1000 mg/kg, 1500 mg/kg, or 2000 mg/kg. Pharmaceutical dosage unit forms are prepared to provide from about 1 mg to about 5000 mg, for example from about 100 to about 2500 mg of the compound or a combination of essential ingredients per dosage unit form.

In embodiments, about 50 nM to about 1 μ M of an agent is administered to a subject. In related embodiments, about 50-100 nM, 50-250 nM, 100-500 nM, 250-500 nM, 250-750 nM, 500-750 nM, or 750 nM to 1 μ M of an agent is administered to a subject.

Determination of an effective amount is well within the capability of those skilled in the art, especially in light of the detailed disclosure provided herein. Generally, an efficacious or effective amount of an agent is determined by first administering a low dose of the agent(s) and then incrementally increasing the administered dose or dosages until a desired effect (e.g., reduce or eliminate symptoms associated with viral infection or autoimmune disease) is observed in the treated subject, with minimal or acceptable toxic side effects. Applicable methods for determining an appropriate dose and dosing schedule for administration of a pharmaceutical composition of the present invention are described, for example, in *Goodman and Gilman's The Pharmacological Basis of Therapeutics*, Goodman *et al.*, eds., 11th Edition, McGraw-Hill 2005, and *Remington: The Science and Practice of Pharmacy*, 20th and 21st Editions, Gennaro and University of the Sciences in Philadelphia, Eds., Lippencott Williams & Wilkins (2003 and 2005), each of which is hereby incorporated by reference.

Combination Therapies

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The tumor specific neo-antigen peptides and pharmaceutical compositions described herein can also be administered in combination with another therapeutic molecule. The therapeutic molecule can be any compound used to mitigate neoplasia, or symptoms thereof. Examples of such compounds include, but are not limited to, chemotherapeutic agents, anti—angiogenesis agents, checkpoint blockade antibodies or other molecules that reduce immune-suppression, and the like.

The tumor specific neo-antigen peptides can be administered before, during, or after administration of the additional therapeutic agent. In embodiments, the tumor specific neo-antigen peptides are administered before the first administration of the additional therapeutic agent. In embodiments, the tumor specific neo-antigen peptides are administered after the first administration of the additional therapeutic agent (e.g., 1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14 days or more). In embodiments, the tumor specific neo-antigen peptides are administered simultaneously with the first administration of the additional therapeutic agent.

Vaccines

In an exemplary embodiment, the present invention is directed to an immunogenic composition, e.g., a vaccine composition capable of raising a specific T-cell response. The

vaccine composition comprises mutant neo-antigenic peptides and mutant neo-antigenic polypeptides corresponding to tumor specific neo-antigens identified by the methods described herein.

A suitable vaccine will preferably contain a plurality of tumor specific neo-antigenic peptides. In an embodiment, the vaccine will include between 1 and 100 sets peptides, more preferably between 1 and 50 such peptides, even more preferably between 10 and 30 sets peptides, even more preferably between 15 and 25 peptides. According to another preferred embodiment, the vaccine will include approximately 20 peptides, more preferably 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, or 30 different peptides, further preferred 6, 7, 8, 9, 10 11, 12, 13, 14, 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 different peptides.

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In one embodiment of the present invention the different tumor specific neo-antigenic peptides and/or polypeptides are selected for use in the neoplasia vaccine so as to maximize the likelihood of generating an immune attack against the neoplasia/tumor of the patient. Without being bound by theory, it is believed that the inclusion of a diversity of tumor specific neo-antigenic peptides will generate a broad scale immune attack against a neoplasia/tumor. In one embodiment, the selected tumor specific neo-antigenic peptides/polypeptides are encoded by missense mutations. In a second embodiment, the selected tumor specific neo-antigenic peptides/polypeptides are encoded by a combination of missense mutations and neoORF mutations. In a third embodiment, the selected tumor specific neo-antigenic peptides/polypeptides are encoded by neoORF mutations.

In one embodiment in which the selected tumor specific neo-antigenic peptides/polypeptides are encoded by missense mutations, the peptides and/or polypeptides are chosen based on their capability to associate with the particular MHC molecules of the patient. Peptides/polypeptides derived from neoORF mutations can also be selected on the basis of their capability to associate with the particular MHC molecules of the patient, but can also be selected even if not predicted to associate with the particular MHC molecules of the patient.

The vaccine composition is capable of raising a specific cytotoxic T-cells response and/or a specific helper T-cell response.

The vaccine composition can further comprise an adjuvant and/or a carrier. Examples of useful adjuvants and carriers are given herein below. The peptides and/or polypeptides in the

composition can be associated with a carrier such as, e.g., a protein or an antigen-presenting cell such as e.g. a dendritic cell (DC) capable of presenting the peptide to a T-cell.

Adjuvants are any substance whose admixture into the vaccine composition increases or otherwise modifies the immune response to the mutant peptide. Carriers are scaffold structures, for example a polypeptide or a polysaccharide, to which the neo-antigenic peptides, is capable of being associated. Optionally, adjuvants are conjugated covalently or non-covalently to the peptides or polypeptides of the invention.

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The ability of an adjuvant to increase the immune response to an antigen is typically manifested by a significant increase in immune-mediated reaction, or reduction in disease symptoms. For example, an increase in humoral immunity is typically manifested by a significant increase in the titer of antibodies raised to the antigen, and an increase in T-cell activity is typically manifested in increased cell proliferation, or cellular cytotoxicity, or cytokine secretion. An adjuvant may also alter an immune response, for example, by changing a primarily humoral or Th2 response into a primarily cellular, or Th1 response.

Suitable adjuvants include, but are not limited to 1018 ISS, aluminum salts, Amplivax, AS15, BCG, CP-870,893, CpG7909, CyaA, dSLIM, GM-CSF, IC30, IC31, Imiquimod, ImuFact IMP321, IS Patch, ISS, ISCOMATRIX, Juvlmmune, LipoVac, MF59, monophosphoryl lipid A, Montanide IMS 1312, Montanide ISA 206, Montanide ISA 50V, Montanide ISA-51, OK-432, OM-174, OM-197-MP-EC, ONTAK, PepTel.RTM. vector system, PLG microparticles, resiguimod, SRL172, Virosomes and other Virus-like particles, YF-17D, VEGF trap, R848, beta-glucan, Pam3Cys, Aquila's QS21 stimulon (Aquila Biotech, Worcester, Mass., USA) which is derived from saponin, mycobacterial extracts and synthetic bacterial cell wall mimics, and other proprietary adjuvants such as Ribi's Detox. Quil or Superfos. Several immunological adjuvants (e.g., MF59) specific for dendritic cells and their preparation have been described previously (Dupuis M, et al., Cell Immunol. 1998; 186(1): 18-27; Allison A C; Dev Biol Stand. 1998; 92:3-11). Also cytokines may be used. Several cytokines have been directly linked to influencing dendritic cell migration to lymphoid tissues (e.g., TNF-alpha), accelerating the maturation of dendritic cells into efficient antigen-presenting cells for T-lymphocytes (e.g., GM-CSF, IL-1 and IL-4) (U.S. Pat. No. 5,849,589, specifically incorporated herein by reference in its entirety) and acting as immunoadjuvants (e.g., IL-12) (Gabrilovich D I, et al., J Immunother Emphasis Tumor Immunol. 1996 (6):414-418).

Toll like receptors (TLRs) may also be used as adjuvants, and are important members of the family of pattern recognition receptors (PRRs) which recognize conserved motifs shared by many micro-organisms, termed "pathogen-associated molecular patterns" (PAMPS). Recognition of these "danger signals" activates multiple elements of the innate and adaptive immune system. TLRs are expressed by cells of the innate and adaptive immune systems such as dendritic cells (DCs), macrophages, T and B cells, mast cells, and granulocytes and are localized in different cellular compartments, such as the plasma membrane, lysosomes, endosomes, and endolysosomes. Different TLRs recognize distinct PAMPS. For example, TLR4 is activated by LPS contained in bacterial cell walls, TLR9 is activated by unmethylated bacterial or viral CpG DNA, and TLR3 is activated by double stranded RNA. TLR ligand binding leads to the activation of one or more intracellular signaling pathways, ultimately resulting in the production of many key molecules associated with inflammation and immunity (particularly the transcription factor NF-κB and the Type-I interferons). TLR mediated DC activation leads to enhanced DC activation, phagocytosis, upregulation of activation and co-stimulation markers such as CD80, CD83, and CD86, expression of CCR7 allowing migration of DC to draining lymph nodes and facilitating antigen presentation to T cells, as well as increased secretion of cytokines such as type I interferons, IL-12, and IL-6. All of these downstream events are critical for the induction of an adaptive immune response.

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Among the most promising cancer vaccine adjuvants currently in clinical development are the TLR9 agonist CpG and the synthetic double-stranded RNA (dsRNA) TLR3 ligand poly-ICLC. In preclinical studies poly-ICLC appears to be the most potent TLR adjuvant when compared to LPS and CpG due to its induction of pro-inflammatory cytokines and lack of stimulation of IL-10, as well as maintenance of high levels of co-stimulatory molecules in DCs. Furthermore, poly-ICLC was recently directly compared to CpG in non-human primates (rhesus macaques) as adjuvant for a protein vaccine consisting of human papillomavirus (HPV)16 capsomers (Stahl-Hennig C, Eisenblatter M, Jasny E, et al. Synthetic double-stranded RNAs are adjuvants for the induction of T helper 1 and humoral immune responses to human papillomavirus in rhesus macaques. PLoS pathogens. Apr 2009;5(4)).

CpG immuno stimulatory oligonucleotides have also been reported to enhance the effects of adjuvants in a vaccine setting. Without being bound by theory, CpG oligonucleotides act by activating the innate (non- adaptive) immune system via Toll-like receptors (TLR), mainly

TLR9. CpG triggered TLR9 activation enhances antigen-specific humoral and cellular responses to a wide variety of antigens, including peptide or protein antigens, live or killed viruses, dendritic cell vaccines, autologous cellular vaccines and polysaccharide conjugates in both prophylactic and therapeutic vaccines. More importantly, it enhances dendritic cell maturation and differentiation, resulting in enhanced activation of Thl cells and strong cytotoxic T- lymphocyte (CTL) generation, even in the absence of CD4 T-cell help. The Thl bias induced by TLR9 stimulation is maintained even in the presence of vaccine adjuvants such as alum or incomplete Freund's adjuvant (IFA) that normally promote a Th2 bias. CpG oligonucleotides show even greater adjuvant activity when formulated or co-administered with other adjuvants or in formulations such as microparticles, nano particles, lipid emulsions or similar formulations, which are especially necessary for inducing a strong response when the antigen is relatively weak. They also accelerate the immune response and enabled the antigen doses to be reduced by approximately two orders of magnitude, with comparable antibody responses to the full-dose vaccine without CpG in some experiments (Arthur M. Krieg, Nature Reviews, Drug Discovery, 5, Jun. 2006, 471-484). U.S. Pat. No. 6,406,705 Bl describes the combined use of CpG oligonucleotides, non-nucleic acid adjuvants and an antigen to induce an antigen-specific immune response. A commercially available CpG TLR9 antagonist is dSLIM (double Stem Loop Immunomodulator) by Mologen (Berlin, GERMANY), which is a preferred component of the pharmaceutical composition of the present invention. Other TLR binding molecules such as RNA binding TLR 7, TLR 8 and/or TLR 9 may also be used.

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Xanthenone derivatives such as, for example, Vadimezan or AsA404 (also known as 5,6-dimethylaxanthenone-4-acetic acid (DMXAA)), may also be used as adjuvants according to embodiments of the invention. Alternatively, such derivatives may also be administered in parallel to the vaccine of the invention, for example via systemic or intratumoral delivery, to stimulate immunity at the tumor site. Without being bound by theory, it is believed that such xanthenone derivatives act by stimulating interferon (IFN) production via the stimulator of IFN gene ISTING) receptor (see e.g., Conlon et al. (2013) Mouse, but not Human STING, Binds and Signals in Response to the Vascular Disrupting Agent 5,6-Dimethylxanthenone-4-Acetic Acid, Journal of Immunology, 190:5216-25 and Kim et al. (2013) Anticancer Flavonoids are Mouse-Selective STING Agonists, 8:1396-1401).

Other examples of useful adjuvants include, but are not limited to, chemically modified CpGs (e.g. CpR, Idera), Poly(I:C)(e.g. polyi:CI2U), non-CpG bacterial DNA or RNA as well as immunoactive small molecules and antibodies such as cyclophosphamide, sunitinib, bevacizumab, celebrex, NCX-4016, sildenafil, tadalafil, vardenafil, sorafinib, XL-999, CP-547632, pazopanib, ZD2171, AZD2171, ipilimumab, tremelimumab, and SC58175, which may act therapeutically and/or as an adjuvant. The amounts and concentrations of adjuvants and additives useful in the context of the present invention can readily be determined by the skilled artisan without undue experimentation. Additional adjuvants include colony- stimulating factors, such as Granulocyte Macrophage Colony Stimulating Factor (GM-CSF, sargramostim).

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Poly-ICLC is a synthetically prepared double-stranded RNA consisting of polyI and polyC strands of average length of about 5000 nucleotides, which has been stabilized to thermal denaturation and hydrolysis by serum nucleases by the addition of polylysine and carboxymethylcellulose. The compound activates TLR3 and the RNA helicase-domain of MDA5, both members of the PAMP family, leading to DC and natural killer (NK) cell activation and production of a "natural mix" of type I interferons, cytokines, and chemokines. Furthermore, poly-ICLC exerts a more direct, broad host-targeted anti-infectious and possibly antitumor effect mediated by the two IFN-inducible nuclear enzyme systems, the 2'5'-OAS and the P1/eIF2a kinase, also known as the PKR (4-6), as well as RIG-I helicase and MDA5.

In rodents and non-human primates, poly-ICLC was shown to enhance T cell responses to viral antigens, cross-priming, and the induction of tumor-, virus-, and autoantigen-specific CD8⁺T-cells. In a recent study in non-human primates, poly-ICLC was found to be essential for the generation of antibody responses and T-cell immunity to DC targeted or non-targeted HIV Gag p24 protein, emphasizing its effectiveness as a vaccine adjuvant.

In human subjects, transcriptional analysis of serial whole blood samples revealed similar gene expression profiles among the 8 healthy human volunteers receiving one single s.c. administration of poly-ICLC and differential expression of up to 212 genes between these 8 subjects versus 4 subjects receiving placebo. Remarkably, comparison of the poly-ICLC gene expression data to previous data from volunteers immunized with the highly effective yellow fever vaccine YF17D showed that a large number of transcriptional and signal transduction canonical pathways, including those of the innate immune system, were similarly upregulated at peak time points.

More recently, an immunologic analysis was reported on patients with ovarian, fallopian tube, and primary peritoneal cancer in second or third complete clinical remission who were treated on a phase 1 study of subcutaneous vaccination with synthetic overlapping long peptides (OLP) from the cancer testis antigen NY-ESO-1 alone or with Montanide-ISA-51, or with 1.4 mg poly-ICLC and Montanide. The generation of NY-ESO-1-specific CD4+ and CD8⁺ T-cell and antibody responses were markedly enhanced with the addition of poly-ICLC and Montanide compared to OLP alone or OLP and Montanide.

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A vaccine composition according to the present invention may comprise more than one different adjuvant. Furthermore, the invention encompasses a therapeutic composition comprising any adjuvant substance including any of the above or combinations thereof. It is also contemplated that the peptide or polypeptide, and the adjuvant can be administered separately in any appropriate sequence.

A carrier may be present independently of an adjuvant. The function of a carrier can for example be to confer stability, to increase the biological activity, or to increase serum half-life. Furthermore, a carrier may aid presenting peptides to T-cells. The carrier may be any suitable carrier known to the person skilled in the art, for example a protein or an antigen presenting cell. A carrier protein could be but is not limited to keyhole limpet hemocyanin, serum proteins such as transferrin, bovine serum albumin, human serum albumin, thyroglobulin or ovalbumin, immunoglobulins, or hormones, such as insulin or palmitic acid. For immunization of humans, the carrier may be a physiologically acceptable carrier acceptable to humans and safe. However, tetanus toxoid and/or diptheria toxoid are suitable carriers in one embodiment of the invention. Alternatively, the carrier may be dextrans for example sepharose.

Cytotoxic T-cells (CTLs) recognize an antigen in the form of a peptide bound to an MHC molecule rather than the intact foreign antigen itself. The MHC molecule itself is located at the cell surface of an antigen presenting cell. Thus, an activation of CTLs is only possible if a trimeric complex of peptide antigen, MHC molecule, and APC is present. Correspondingly, it may enhance the immune response if not only the peptide is used for activation of CTLs, but if additionally APCs with the respective MHC molecule are added. Therefore, in some embodiments the vaccine composition according to the present invention additionally contains at least one antigen presenting cell.

The antigen-presenting cell (or stimulator cell) typically has an MHC class I or II molecule on its surface, and in one embodiment is substantially incapable of itself loading the MHC class I or II molecule with the selected antigen. As is described in more detail below, the MHC class I or II molecule may readily be loaded with the selected antigen in vitro.

Preferably, the antigen presenting cells are dendritic cells. Suitably, the dendritic cells are autologous dendritic cells that are pulsed with the neo-antigenic peptide. The peptide may be any suitable peptide that gives rise to an appropriate T-cell response. T-cell therapy using autologous dendritic cells pulsed with peptides from a tumor associated antigen is disclosed in Murphy et al. (1996) The Prostate 29, 371-380 and Tjua et al. (1997) The Prostate 32, 272-278.

Thus, in one embodiment of the present invention the vaccine composition containing at least one antigen presenting cell is pulsed or loaded with one or more peptides of the present invention. Alternatively, peripheral blood mononuclear cells (PBMCs) isolated from a patient may be loaded with peptides ex vivo and injected back into the patient. As an alternative the antigen presenting cell comprises an expression construct encoding a peptide of the present invention. The polynucleotide may be any suitable polynucleotide and it is preferred that it is capable of transducing the dendritic cell, thus resulting in the presentation of a peptide and induction of immunity.

Therapeutic Methods

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The invention further provides a method of inducing a neoplasia/tumor specific immune response in a subject, vaccinating against a neoplasia/tumor, treating and or alleviating a symptom of cancer in a subject by administering the subject a neo-antigenic peptide or vaccine composition of the invention.

According to the invention, the above-described cancer vaccine may be used for a patient that has been diagnosed as having cancer, or at risk of developing cancer. In one embodiment, the patient may have a solid tumor such as breast, ovarian, prostate, lung, kidney, gastric, colon, testicular, head and neck, pancreas, brain, melanoma, and other tumors of tissue organs and hematological tumors, such as lymphomas and leukemias, including acute myelogenous leukemia, chronic myelogenous leukemia, chronic lymphocytic leukemia, T cell lymphocytic leukemia, and B cell lymphomas.

The peptide or composition of the invention is administered in an amount sufficient to induce a CTL response.

The neo-antigenic peptide, polypeptide or vaccine composition of the invention can be administered alone or in combination with other therapeutic agents. The therapeutic agent is for example, a chemotherapeutic or biotherapeutic agent, radiation, or immunotherapy. Any suitable therapeutic treatment for a particular cancer may be administered. Examples of chemotherapeutic and biotherapeutic agents include, but are not limited to, aldesleukin, altretamine, amifostine, asparaginase, bleomycin, capecitabine, carboplatin, carmustine, cladribine, cisapride, cisplatin, cyclophosphamide, cytarabine, dacarbazine (DTIC), dactinomycin, docetaxel, doxorubicin, dronabinol, epoetin alpha, etoposide, filgrastim, fludarabine, fluorouracil, gemcitabine, granisetron, hydroxyurea, idarubicin, ifosfamide, interferon alpha, irinotecan, lansoprazole, levamisole, leucovorin, megestrol, mesna, methotrexate, metoclopramide, mitomycin, mitotane, mitoxantrone, omeprazole, ondansetron, paclitaxel (Taxol®), pilocarpine, prochloroperazine, rituximab, tamoxifen, taxol, topotecan hydrochloride, trastuzumab, vinblastine, vincristine and vinorelbine tartrate. For prostate cancer treatment, a preferred chemotherapeutic agent with which anti- CTLA-4 can be combined is paclitaxel (Taxol®).

In addition, the subject may be further administered an anti- immunosuppressive or immunostimulatory agent. For example, the subject is further administered an anti-CTLA antibody or anti-PD-1 or anti-PD-Ll. Blockade of CTLA-4 or PD-1/PD-L1 by antibodies can enhance the immune response to cancerous cells in the patient. In particular, CTLA-4 blockade has been shown effective when following a vaccination protocol (Hodi et al 2005).

The optimum amount of each peptide to be included in the vaccine composition and the optimum dosing regimen can be determined by one skilled in the art without undue experimentation. For example, the peptide or its variant may be prepared for intravenous (i.v.) injection, sub-cutaneous (s.c.) injection, intradermal (i.d.) injection, intraperitoneal (i.p.) injection, intramuscular (i.m.) injection. Preferred methods of peptide injection include s.c, i.d., i.p., i.m., and i.v. Preferred methods of DNA injection include i.d., i.m., s.c, i.p. and i.v. For example, doses of between 1 and 500 mg 50 µg and 1.5 mg, preferably 10 µg to 500 µg, of peptide or DNA may be given and will depend from the respective peptide or DNA. Doses of this range were successfully used in previous trials (Brunsvig P F, et al., Cancer Immunol Immunother. 2006; 55(12): 1553- 1564; M. Staehler, et al., ASCO meeting 2007; Abstract No

3017). Other methods of administration of the vaccine composition are known to those skilled in the art.

The inventive pharmaceutical composition may be compiled so that the selection, number and/or amount of peptides present in the composition is/are tissue, cancer, and/or patient-specific. For instance, the exact selection of peptides can be guided by expression patterns of the parent proteins in a given tissue to avoid side effects. The selection may be dependent on the specific type of cancer, the status of the disease, earlier treatment regimens, the immune status of the patient, and, of course, the HLA-haplotype of the patient. Furthermore, the vaccine according to the invention can contain individualized components, according to personal needs of the particular patient. Examples include varying the amounts of peptides according to the expression of the related neoantigen in the particular patient, unwanted side-effects due to personal allergies or other treatments, and adjustments for secondary treatments following a first round or scheme of treatment.

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Pharmaceutical compositions comprising the peptide of the invention may be administered to an individual already suffering from cancer. In therapeutic applications, compositions are administered to a patient in an amount sufficient to elicit an effective CTL response to the tumor antigen and to cure or at least partially arrest symptoms and/or complications. An amount adequate to accomplish this is defined as "therapeutically effective dose." Amounts effective for this use will depend on, e.g., the peptide composition, the manner of administration, the stage and severity of the disease being treated, the weight and general state of health of the patient, and the judgment of the prescribing physician, but generally range for the initial immunization (that is for therapeutic or prophylactic administration) from about 1.0 µg to about 50,000 µg of peptide for a 70 kg patient, followed by boosting dosages or from about 1.0 μg to about 10,000 μg of peptide pursuant to a boosting regimen over weeks to months depending upon the patient's response and condition and possibly by measuring specific CTL activity in the patient's blood. It should be kept in mind that the peptide and compositions of the present invention may generally be employed in serious disease states, that is, life-threatening or potentially life threatening situations, especially when the cancer has metastasized. For therapeutic use, administration should begin as soon as possible after the detection or surgical removal of tumors. This is followed by boosting doses until at least symptoms are substantially abated and for a period thereafter.

The pharmaceutical compositions (e.g., vaccine compositions) for therapeutic treatment are intended for parenteral, topical, nasal, oral or local administration. Preferably, the pharmaceutical compositions are administered parenterally, e.g., intravenously, subcutaneously, intradermally, or intramuscularly. The compositions may be administered at the site of surgical excision to induce a local immune response to the tumor. The invention provides compositions for parenteral administration which comprise a solution of the peptides and vaccine compositions are dissolved or suspended in an acceptable carrier, preferably an aqueous carrier. A variety of aqueous carriers may be used, e.g., water, buffered water, 0.9% saline, 0.3% glycine, hyaluronic acid and the like. These compositions may be sterilized by conventional, well known sterilization techniques, or may be sterile filtered. The resulting aqueous solutions may be packaged for use as is, or lyophilized, the lyophilized preparation being combined with a sterile solution prior to administration. The compositions may contain pharmaceutically acceptable auxiliary substances as required to approximate physiological conditions, such as pH adjusting and buffering agents, tonicity adjusting agents, wetting agents and the like, for example, sodium acetate, sodium lactate, sodium chloride, potassium chloride, calcium chloride, sorbitan monolaurate, triethanolamine oleate, etc.

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The concentration of peptides of the invention in the pharmaceutical formulations can vary widely, i.e., from usually less than about 0.1%, to at least about 2% to as much as 20% to 50% or more by weight, and will be selected primarily by fluid volumes, viscosities, etc., in accordance with the particular mode of administration selected.

A liposome suspension containing a peptide may be administered intravenously, locally, topically, etc. in a dose which varies according to, inter alia, the manner of administration, the peptide being delivered, and the stage of the disease being treated. For targeting to the immune cells, a ligand, such as, e.g., antibodies or fragments thereof specific for cell surface determinants of the desired immune system cells, can be incorporated into the liposome.

For solid compositions, conventional or nanoparticle nontoxic solid carriers may be used which include, for example, pharmaceutical grades of mannitol, lactose, starch, magnesium stearate, sodium saccharin, talcum, cellulose, glucose, sucrose, magnesium carbonate, and the like. For oral administration, a pharmaceutically acceptable nontoxic composition is formed by incorporating any of the normally employed excipients, such as those carriers previously listed,

and generally 10-95% of active ingredient, that is, one or more peptides of the invention, and more preferably at a concentration of 25%-75%.

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For aerosol administration, the immunogenic peptides are preferably supplied in finely divided form along with a surfactant and propellant. Typical percentages of peptides are 0.01 %-20% by weight, preferably 1%-10%. The surfactant will, of course, be nontoxic, and preferably soluble in the propellant. Representative of such agents are the esters or partial esters of fatty acids containing from 6 to 22 carbon atoms, such as caproic, octanoic, lauric, palmitic, stearic, linoleic, linolenic, olesteric and oleic acids with an aliphatic polyhydric alcohol or its cyclic anhydride. Mixed esters, such as mixed or natural glycerides may be employed. The surfactant may constitute 0.1%-20% by weight of the composition, preferably 0.25-5%. The balance of the composition is ordinarily propellant. A carrier can also be included as desired, as with, e.g., lecithin for intranasal delivery.

The peptides and polypeptides of the invention can be readily synthesized chemically utilizing reagents that are free of contaminating bacterial or animal substances (Merrifield RB: Solid phase peptide synthesis. I. The synthesis of a tetrapeptide. J. Am. Chem. Soc. 85:2149-54, 1963).

For therapeutic or immunization purposes, nucleic acids encoding the peptide of the invention and optionally one or more of the peptides described herein can also be administered to the patient. A number of methods are conveniently used to deliver the nucleic acids to the patient. For instance, the nucleic acid can be delivered directly, as "naked DNA". This approach is described, for instance, in Wolff et al., Science 247: 1465-1468 (1990) as well as U.S. Patent Nos. 5,580,859 and 5,589,466. The nucleic acids can also be administered using ballistic delivery as described, for instance, in U.S. Patent No. 5,204,253. Particles comprised solely of DNA can be administered. Alternatively, DNA can be adhered to particles, such as gold particles.

The nucleic acids can also be delivered complexed to cationic compounds, such as cationic lipids. Lipid-mediated gene delivery methods are described, for instance, in WO1996/18372; WO 1993/24640; Mannino & Gould-Fogerite, BioTechniques 6(7): 682-691 (1988); U.S. Patent No. 5,279,833; WO 1991/06309; and Feigner et al., Proc. Natl. Acad. Sci. USA 84: 7413-7414 (1987).

RNA encoding the peptide of interest can also be used for delivery (see, e.g., Kiken et al, 2011; Su et al, 2011).

The peptides and polypeptides of the invention can also be expressed by attenuated viral hosts, such as vaccinia or fowlpox. This approach involves the use of vaccinia virus as a vector to express nucleotide sequences that encode the peptide of the invention. Upon introduction into an acutely or chronically infected host or into a noninfected host, the recombinant vaccinia virus expresses the immunogenic peptide, and thereby elicits a host CTL response. Vaccinia vectors and methods useful in immunization protocols are described in, e.g., U.S. Patent No. 4,722,848,. Another vector is BCG (Bacille Calmette Guerin). BCG vectors are described in Stover et al. (Nature 351:456-460 (1991)). A wide variety of other vectors useful for therapeutic administration or immunization of the peptides of the invention, e.g., Salmonella typhi vectors and the like, will be apparent to those skilled in the art from the description herein.

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A preferred means of administering nucleic acids encoding the peptide of the invention uses minigene constructs encoding multiple epitopes. To create a DNA sequence encoding the selected CTL epitopes (minigene) for expression in human cells, the amino acid sequences of the epitopes are reverse translated. A human codon usage table is used to guide the codon choice for each amino acid. These epitope-encoding DNA sequences are directly adjoined, creating a continuous polypeptide sequence. To optimize expression and/or immunogenicity, additional elements can be incorporated into the minigene design. Examples of amino acid sequence that could be reverse translated and included in the minigene sequence include: helper T lymphocyte, epitopes, a leader (signal) sequence, and an endoplasmic reticulum retention signal. In addition, MHC presentation of CTL epitopes may be improved by including synthetic (e.g. poly-alanine) or naturally- occurring flanking sequences adjacent to the CTL epitopes.

The minigene sequence is converted to DNA by assembling oligonucleotides that encode the plus and minus strands of the minigene. Overlapping oligonucleotides (30-100 bases long) are synthesized, phosphorylated, purified and annealed under appropriate conditions using well known techniques. The ends of the oligonucleotides are joined using T4 DNA ligase. This synthetic minigene, encoding the CTL epitope polypeptide, can then cloned into a desired expression vector.

Standard regulatory sequences well known to those of skill in the art are included in the vector to ensure expression in the target cells. Several vector elements are required: a promoter

with a down-stream cloning site for minigene insertion; a polyadenylation signal for efficient transcription termination; an E. coli origin of replication; and an E. coli selectable marker (e.g. ampicillin or kanamycin resistance). Numerous promoters can be used for this purpose, e.g., the human cytomegalovirus (hCMV) promoter. See, U.S. Patent Nos. 5,580,859 and 5,589,466 for other suitable promoter sequences.

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Additional vector modifications may be desired to optimize minigene expression and immunogenicity. In some cases, introns are required for efficient gene expression, and one or more synthetic or naturally-occurring introns could be incorporated into the transcribed region of the minigene. The inclusion of mRNA stabilization sequences can also be considered for increasing minigene expression. It has recently been proposed that immuno stimulatory sequences (ISSs or CpGs) play a role in the immunogenicity of DNA' vaccines. These sequences could be included in the vector, outside the minigene coding sequence, if found to enhance immunogenicity.

In some embodiments, a bicistronic expression vector, to allow production of the minigene-encoded epitopes and a second protein included to enhance or decrease immunogenicity can be used. Examples of proteins or polypeptides that could beneficially enhance the immune response if co-expressed include cytokines (e.g., IL2, IL12, GM-CSF), cytokine-inducing molecules (e.g. LeIF) or costimulatory molecules. Helper (HTL) epitopes could be joined to intracellular targeting signals and expressed separately from the CTL epitopes. This would allow direction of the HTL epitopes to a cell compartment different than the CTL epitopes. If required, this could facilitate more efficient entry of HTL epitopes into the MHC class II pathway, thereby improving CTL induction. In contrast to CTL induction, specifically decreasing the immune response by co-expression of immunosuppressive molecules (e.g. TGF-β) may be beneficial in certain diseases.

Once an expression vector is selected, the minigene is cloned into the polylinker region downstream of the promoter. This plasmid is transformed into an appropriate E. coli strain, and DNA is prepared using standard techniques. The orientation and DNA sequence of the minigene, as well as all other elements included in the vector, are confirmed using restriction mapping and DNA sequence analysis. Bacterial cells harboring the correct plasmid can be stored as a master cell bank and a working cell bank.

Purified plasmid DNA can be prepared for injection using a variety of formulations. The simplest of these is reconstitution of lyophilized DNA in sterile phosphate-buffer saline (PBS). A variety of methods have been described, and new techniques may become available. As noted above, nucleic acids are conveniently formulated with cationic lipids. In addition, glycolipids, fusogenic liposomes, peptides and compounds referred to collectively as protective, interactive, non-condensing (PINC) could also be complexed to purified plasmid DNA to influence variables such as stability, intramuscular dispersion, or trafficking to specific organs or cell types.

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Target cell sensitization can be used as a functional assay for expression and MHC class I presentation of minigene-encoded CTL epitopes. The plasmid DNA is introduced into a mammalian cell line that is suitable as a target for standard CTL chromium release assays. The transfection method used will be dependent on the final formulation. Electroporation can be used for "naked" DNA, whereas cationic lipids allow direct in vitro transfection. A plasmid expressing green fluorescent protein (GFP) can be co-transfected to allow enrichment of transfected cells using fluorescence activated cell sorting (FACS). These cells are then chromium-51 labeled and used as target cells for epitope- specific CTL lines. Cytolysis, detected by 51 Cr release, indicates production of MHC presentation of mini gene-encoded CTL epitopes.

In vivo immunogenicity is a second approach for functional testing of minigene DNA formulations. Transgenic mice expressing appropriate human MHC molecules are immunized with the DNA product. The dose and route of administration are formulation dependent (e.g. IM for DNA in PBS, IP for lipid-complexed DNA). Twenty-one days after immunization, splenocytes are harvested and restimulated for 1 week in the presence of peptides encoding each epitope being tested. These effector cells (CTLs) are assayed for cytolysis of peptide-loaded, chromium-51 labeled target cells using standard techniques. Lysis of target cells sensitized by MHC loading of peptides corresponding to minigene-encoded epitopes demonstrates DNA vaccine function for in vivo induction of CTLs.

Peptides may be used to elicit CTL ex vivo, as well. The resulting CTL, can be used to treat chronic tumors in patients that do not respond to other conventional forms of therapy, or will not respond to a peptide vaccine approach of therapy. Ex vivo CTL responses to a particular tumor antigen are induced by incubating in tissue culture the patient's CTL precursor cells (CTLp) together with a source of antigen-presenting cells (APC) and the appropriate peptide. After an appropriate incubation time (typically 1-4 weeks), in which the CTLp are activated and

mature and expand into effector CTL, the cells are infused back into the patient, where they will destroy their specific target cell (i.e., a tumor cell). In order to optimize the in vitro conditions for the generation of specific cytotoxic T cells, the culture of stimulator cells is maintained in an appropriate serum-free medium.

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Prior to incubation of the stimulator cells with the cells to be activated, e.g., precursor CD8+ cells, an amount of antigenic peptide is added to the stimulator cell culture, of sufficient quantity to become loaded onto the human Class I molecules to be expressed on the surface of the stimulator cells. In the present invention, a sufficient amount of peptide is an amount that will allow about 200, and preferably 200 or more, human Class I MHC molecules loaded with peptide to be expressed on the surface of each stimulator cell. Preferably, the stimulator cells are incubated with $>2\mu g/ml$ peptide. For example, the stimulator cells are incubates with >3, 4, 5, 10, 15, or more $\mu g/ml$ peptide.

Resting or precursor CD8+ cells are then incubated in culture with the appropriate stimulator cells for a time period sufficient to activate the CD8+ cells. Preferably, the CD8+ cells are activated in an antigen- specific manner. The ratio of resting or precursor CD8+ (effector) cells to stimulator cells may vary from individual to individual and may further depend upon variables such as the amenability of an individual's lymphocytes to culturing conditions and the nature and severity of the disease condition or other condition for which the within-described treatment modality is used. Preferably, however, the lymphocyte: stimulator cell ratio is in the range of about 30: 1 to 300: 1. The effector/stimulator culture may be maintained for as long a time as is necessary to stimulate a therapeutically useable or effective number of CD8+ cells.

The induction of CTL in vitro requires the specific recognition of peptides that are bound to allele specific MHC class I molecules on APC. The number of specific MHC/peptide complexes per APC is crucial for the stimulation of CTL, particularly in primary immune responses. While small amounts of peptide/MHC complexes per cell are sufficient to render a cell susceptible to lysis by CTL, or to stimulate a secondary CTL response, the successful activation of a CTL precursor (pCTL) during primary response requires a significantly higher number of MHC/peptide complexes. Peptide loading of empty major histocompatability complex molecules on cells allows the induction of primary cytotoxic T lymphocyte responses.

Since mutant cell lines do not exist for every human MHC allele, it is advantageous to use a technique to remove endogenous MHC- associated peptides from the surface of APC,

followed by loading the resulting empty MHC molecules with the immunogenic peptides of interest. The use of non-transformed (non-tumorigenic), noninfected cells, and preferably, autologous cells of patients as APC is desirable for the design of CTL induction protocols directed towards development of ex vivo CTL therapies. This application discloses methods for stripping the endogenous MHC-associated peptides from the surface of APC followed by the loading of desired peptides.

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A stable MHC class I molecule is a trimeric complex formed of the following elements:

1) a peptide usually of 8 - 10 residues, 2) a transmembrane heavy polymorphic protein chain which bears the peptide-binding site in its al and a2 domains, and 3) a non-covalently associated non-polymorphic light chain, p2microglobulin. Removing the bound peptides and/or dissociating the p2microglobulin from the complex renders the MHC class I molecules nonfunctional and unstable, resulting in rapid degradation. All MHC class I molecules isolated from PBMCs have endogenous peptides bound to them. Therefore, the first step is to remove all endogenous peptides bound to MHC class I molecules on the APC without causing their degradation before exogenous peptides can be added to them.

Two possible ways to free up MHC class I molecules of bound peptides include lowering the culture temperature from 37°C to 26°C overnight to destabilize p2microglobulin and stripping the endogenous peptides from the cell using a mild acid treatment. The methods release previously bound peptides into the extracellular environment allowing new exogenous peptides to bind to the empty class I molecules. The cold-temperature incubation method enables exogenous peptides to bind efficiently to the MHC complex, but requires an overnight incubation at 26°C which may slow the cell's metabolic rate. It is also likely that cells not actively synthesizing MHC molecules (e.g., resting PBMC) would not produce high amounts of empty surface MHC molecules by the cold temperature procedure.

Harsh acid stripping involves extraction of the peptides with trifluoroacetic acid, pH 2, or acid denaturation of the immunoaffinity purified class I-peptide complexes. These methods are not feasible for CTL induction, since it is important to remove the endogenous peptides while preserving APC viability and an optimal metabolic state which is critical for antigen presentation. Mild acid solutions of pH 3 such as glycine or citrate -phosphate buffers have been used to identify endogenous peptides and to identify tumor associated T cell epitopes. The treatment is especially effective, in that only the MHC class I molecules are destabilized (and

associated peptides released), while other surface antigens remain intact, including MHC class II molecules. Most importantly, treatment of cells with the mild acid solutions do not affect the cell's viability or metabolic state. The mild acid treatment is rapid since the stripping of the endogenous peptides occurs in two minutes at 4°C and the APC is ready to perform its function after the appropriate peptides are loaded. The technique is utilized herein to make peptidespecific APCs for the generation of primary antigen-specific CTL. The resulting APC are efficient in inducing peptide-specific CD8+ CTL.

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Activated CD8+ cells may be effectively separated from the stimulator cells using one of a variety of known methods. For example, monoclonal antibodies specific for the stimulator cells, for the peptides loaded onto the stimulator cells, or for the CD8+ cells (or a segment thereof) may be utilized to bind their appropriate complementary ligand. Antibody- tagged molecules may then be extracted from the stimulator-effector cell admixture via appropriate means, e.g., via well-known immunoprecipitation or immunoassay methods.

Effective, cytotoxic amounts of the activated CD8+ cells can vary between in vitro and in vivo uses, as well as with the amount and type of cells that are the ultimate target of these killer cells. The amount will also vary depending on the condition of the patient and should be determined via consideration of all appropriate factors by the practitioner. Preferably, however, about 1×10^6 to about 1×10^{12} , more preferably about 1×10^8 to about 1×10^{11} , and even more preferably, about 1×10^9 to about 1×10^{10} activated CD8+ cells are utilized for adult humans, compared to about 5×10^6 - 5×10^7 cells used in mice.

Preferably, as discussed above, the activated CD8+ cells are harvested from the cell culture prior to administration of the CD8+ cells to the individual being treated. It is important to note, however, that unlike other present and proposed treatment modalities, the present method uses a cell culture system that is not tumorigenic. Therefore, if complete separation of stimulator cells and activated CD8+ cells is not achieved, there is no inherent danger known to be associated with the administration of a small number of stimulator cells, whereas administration of mammalian tumor-promoting cells may be extremely hazardous.

Methods of re-introducing cellular components are known in the art and include procedures such as those exemplified in U.S. Patent No. 4,844,893 to Honsik, et al. and U.S. Patent No. 4,690,915 to Rosenberg. For example, administration of activated CD8+ cells via intravenous infusion is appropriate.

CD8+ cell activity may be augmented through the use of CD4+ cells. The identification of CD4 T+ cell epitopes for tumor antigens has attracted interest because many immune based therapies against cancer may be more effective if both CD8+ and CD4+ T lymphocytes are used to target a patient's tumor. CD4+ cells are capable of enhancing CD8 T cell responses. Many studies in animal models have clearly demonstrated better results when both CD4+ and CD8+ T cells participate in anti-tumor responses (see e.g., Nishimura et al. (1999) Distinct role of antigen-specific T helper type 1 (TH1) and Th2 cells in tumor eradication in vivo. J Ex Med 190:617-27). Universal CD4+ T cell epitopes have been identified that are applicable to developing therapies against different types of cancer (see e.g., Kobayashi et al. (2008) Current Opinion in Immunology 20:221-27). For example, an HLA-DR restricted helper peptide from tetanus toxoid was used in melanoma vaccines to activate CD4+ T cells non-specifically (see e.g., Slingluff et al. (2007) Immunologic and Clinical Outcomes of a Randomized Phase II Trial of Two Multipeptide Vaccines for Melanoma in the Adjuvant Setting, Clinical Cancer Research 13(21):6386-95). It is contemplated within the scope of the invention that such CD4+ cells may be applicable at three levels that vary in their tumor specificity: 1) a broad level in which universal CD4+ epitopes (e.g., tetanus toxoid) may be used to augment CD8+ cells; 2) an intermediate level in which native, tumor-associated CD4+ epitopes may be used to augment CD8+ cells; and 3) a patient specific level in which neoantigen CD4+ epitopes may be used to augment CD8+ cells in a patient specific manner.

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CD8+ cell immunity may also be generated with neo-antigen loaded dendritic cell (DC) vaccine. DCs are potent antigen-presenting cells that initiate T cell immunity and can be used as cancer vaccines when loaded with one or more peptides of interest, for example, by direct peptide injection. For example, patients that were newly diagnosed with metastatic melanoma were shown to be immunized against 3 HLA-A*0201-restricted gp100 melanoma antigenderived peptides with autologous peptide pulsed CD40L/IFN-g-activated mature DCs via an IL-12p70-producing patient DC vaccine (see e.g., Carreno et al (2013) L-12p70-producing patient DC vaccine elicits Tc1-polarized immunity, Journal of Clinical Investigation, 123(8):3383-94 and Ali et al. (2009) In situ regulation of DC subsets and T cells mediates tumor regression in mice, Cancer Immunotherapy, 1(8):1-10). It is contemplated within the scope of the invention that neo-antigen loaded DCs may be prepared using the synthetic TLR 3 agonist Polyinosinic-Polycytidylic Acid-poly-L-lysine Carboxymethylcellulose (Poly-ICLC) to stimulate the DCs.

Poly-ICLC is a potent individual maturation stimulus for human DCs as assessed by an upregulation of CD83 and CD86, induction of interleukin-12 (IL-12), tumor necrosis factor (TNF), interferon gamma-induced protein 10 (IP-10), interleukin 1 (IL-1), and type I interferons (IFN), and minimal interleukin 10 (IL-10) production. DCs may be differentiated from frozen peripheral blood mononuclear cells (PBMCs) obtained by leukapheresis, while PBMCs may be isolated by Ficoll gradient centrifugation and frozen in aliquots.

Illustratively, the following 7 day activation protocol may be used. Day 1—PBMCs are thawed and plated onto tissue culture flasks to select for monocytes which adhere to the plastic surface after 1-2 hr incubation at 37°C in the tissue culture incubator. After incubation, the lymphocytes are washed off and the adherent monocytes are cultured for 5 days in the presence of interleukin-4 (IL-4) and granulocyte macrophage-colony stimulating factor (GM-CSF) to differentiate to immature DCs. On Day 6, immature DCs are pulsed with the keyhole limpet hemocyanin (KLH) protein which serves as a control for the quality of the vaccine and may boost the immunogenicity of the vaccine. The DCs are stimulated to mature, loaded with peptide antigens, and incubated overnight. On Day 7, the cells are washed, and frozen in 1 ml aliquots containing 4-20 x 10(6) cells using a controlled-rate freezer. Lot release testing for the batches of DCs may be performed to meet minimum specifications before the DCs are injected into patients (see e.g., Sabado et al. (2013) Preparation of tumor antigen-loaded mature dendritic cells for immunotherapy, J. Vis Exp. Aug 1;(78). doi: 10.3791/50085).

A DC vaccine may be incorporated into a scaffold system to facilitate delivery to a patient. Therapeutic treatment of a patients neoplasia with a DC vaccine may utilize a biomaterial system that releases factors that recruit host dendritic cells into the device, differentiates the resident, immature DCs by locally presenting adjuvants (e.g., danger signals) while releasing antigen, and promotes the release of activated, antigen loaded DCs to the lymph nodes (or desired site of action) where the DCs may interact with T cells to generate a potent cytotoxic T lymphocyte response to the cancer neo-antigens. Implantable biomaterials may be used to generate a potent cytotoxic T lymphocyte response against a neoplasia in a patient specific manner. The biomaterial-resident dendritic cells may then be activated by exposing them to danger signals mimicking infection, in concert with release of antigen from the biomaterial. The activated dendritic cells then migrate from the biomaterials to lymph nodes to induce a cytotoxic T effector response. This approach has previously been demonstrated to lead

to regression of established melanoma in preclinical studies using a lysate prepared from tumor biopsies (see e.g., Ali et al. (2209) In situ regulation of DC subsets and T cells mediates tumor regression in mice, Cancer Immunotherapy 1(8):1-10; Ali et al. (2009) Infection-mimicking materials to program dendritic cells in situ. Nat Mater 8:151-8), and such a vaccine is currently being tested in a Phase I clinical trial recently initiated at the Dana-Farber Cancer Institute. This approach has also been shown to lead to regression of glioblastoma, as well as the induction of a potent memory response to prevent relapse, using the C6 rat glioma model.24 In the current proposal. The ability of such an implantable, biomatrix vaccine delivery scaffold to amplify and sustain tumor specific dendritic cell activation may lead to more robust anti-tumor immunosensitization than can be achieved by traditional subcutaneous or intra-nodal vaccine administrations.

The practice of the present invention employs, unless otherwise indicated, conventional techniques of molecular biology (including recombinant techniques), microbiology, cell biology, biochemistry and immunology, which are well within the purview of the skilled artisan. Such techniques are explained fully in the literature, such as, "Molecular Cloning: A Laboratory Manual", second edition (Sambrook, 1989); "Oligonucleotide Synthesis" (Gait, 1984); "Animal Cell Culture" (Freshney, 1987); "Methods in Enzymology" "Handbook of Experimental Immunology" (Wei, 1996); "Gene Transfer Vectors for Mammalian Cells" (Miller and Calos, 1987); "Current Protocols in Molecular Biology" (Ausubel, 1987); "PCR: The Polymerase Chain Reaction", (Mullis, 1994); "Current Protocols in Immunology" (Coligan, 1991). These techniques are applicable to the production of the polynucleotides and polypeptides of the invention, and, as such, may be considered in making and practicing the invention. Particularly useful techniques for particular embodiments will be discussed in the sections that follow.

25 EXAMPLES

The following examples are put forth so as to provide those of ordinary skill in the art with a complete disclosure and description of how to make and use the assay, screening, and therapeutic methods of the invention, and are not intended to limit the scope of what the inventors regard as their invention.

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Example 1: Cancer Vaccine Testing Protocol

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The above-described compositions and methods may be tested on 15 patients with high-risk melanoma (fully resected stages IIIB, IIIC and IVM1a,b) according to the general flow process shown in FIG. 2. Patients may receive a series of priming vaccinations with a mixture of personalized tumor-specific peptides and poly-ICLC over a 4 week period followed by two boosts during a maintenance phase. All vaccinations will be subcutaneously delivered. The vaccine will be evaluated for safety, tolerability, immune response and clinical effect in patients and for feasibility of producing vaccine and successfully initiating vaccination within an appropriate time frame. The first cohort will consist of 5 patients, and after safety is adequately demonstrated, an additional cohort of 10 patients may be enrolled (see, e.g., FIG. 3 depicting an approach for an initial population study). Peripheral blood will be extensively monitored for peptide-specific T-cell responses and patients will be followed for up to two years to assess disease recurrence.

As described above, there is a large body of evidence in both animals and humans that mutated epitopes are effective in inducing an immune response and that cases of spontaneous tumor regression or long term survival correlate with CD8⁺ T-cell responses to mutated epitopes (Buckwalter and Srivastava PK. "It is the antigen(s), stupid" and other lessons from over a decade of vaccitherapy of human cancer. Seminars in immunology 20:296-300 (2008); Karanikas et al, High frequency of cytolytic T lymphocytes directed against a tumor-specific mutated antigen detectable with HLA tetramers in the blood of a lung carcinoma patient with long survival. Cancer Res. 61:3718-3724 (2001); Lennerz et al, The response of autologous T cells to a human melanoma is dominated by mutated neo-antigens. Proc Natl Acad Sci U S A.102:16013 (2005)) and that "immunoediting" can be tracked to alterations in expression of dominant mutated antigens in mice and man (Matsushita et al, Cancer exome analysis reveals a T-cell-dependent mechanism of cancer immunoediting Nature 482:400 (2012); DuPage et al, Expression of tumor-specific antigens underlies cancer immunoediting Nature 482:405 (2012); and Sampson et al, Immunologic escape after prolonged progression-free survival with epidermal growth factor receptor variant III peptide vaccination in patients with newly diagnosed glioblastoma J Clin Oncol. 28:4722-4729 (2010)).

Next-generation sequencing can now rapidly reveal the presence of discrete mutations such as coding mutations in individual tumors, most commonly single amino acid changes (e.g.,

missense mutations; FIG. 4A) and less frequently novel stretches of amino acids generated by frame-shift insertions/deletions/gene fusions, read-through mutations in stop codons, and translation of improperly spliced introns (e.g., neoORFs; FIG. 4B). NeoORFs are particularly valuable as immunogens because the entirety of their sequence is completely novel to the immune system and so are analogous to a viral or bacterial foreign antigen. Thus, neoORFs: (1) are highly specific to the tumor (i.e. there is no expression in any normal cells); (2) can bypass central tolerance, thereby increasing the precursor frequency of neoantigen-specific CTLs. For example, the power of utilizing analogous foreign sequences in a therapeutic anti-cancer vaccine was recently demonstrated with peptides derived from human papilloma virus (HPV). ~50% of the 19 patients with pre-neoplastic, viral-induced disease who received 3 - 4 vaccinations of a mix of HPV peptides derived from the viral oncogenes E6 and E7 maintained a complete response for ≥24 months (Kenter et a, Vaccination against HPV-16 Oncoproteins for Vulvar Intraepithelial Neoplasia NEJM 361:1838 (2009)).

Sequencing technology has revealed that each tumor contains multiple, patient-specific mutations that alter the protein coding content of a gene. Such mutations create altered proteins, ranging from single amino acid changes (caused by missense mutations) to addition of long regions of novel amino acid sequence due to frame shifts, read-through of termination codons or translation of intron regions (novel open reading frame mutations; neoORFs). These mutated proteins are valuable targets for the host's immune response to the tumor as, unlike native proteins, they are not subject to the immune-dampening effects of self-tolerance. Therefore, mutated proteins are more likely to be immunogenic and are also more specific for the tumor cells compared to normal cells of the patient.

Utilizing recently improved algorithms for predicting which missense mutations create strong binding peptides to the patient's cognate MHC molecules, a set of peptides representative of optimal mutated epitopes (both neoORF and missense) for each patient will be identified and prioritized and up to 20 or more peptides will be prepared for immunization (Zhang et al, Machine learning competition in immunology – Prediction of HLA class I binding peptides J Immunol Methods 374:1 (2011); Lundegaard et al Prediction of epitopes using neural network based methods J Immunol Methods 374:26 (2011)). Peptides ~20-35 amino acids in length will be synthesized because such "long" peptides undergo efficient internalization, processing and cross-presentation in professional antigen-presenting cells such as dendritic cells, and have been

shown to induce CTLs in humans (Melief and van der Burg, Immunotherapy of established (pre) malignant disease by synthetic long peptide vaccines Nature Rev Cancer 8:351 (2008)).

In addition to a powerful and specific immunogen, an effective immune response requires a strong adjuvant to activate the immune system (Speiser and Romero, Molecularly defined vaccines for cancer immunotherapy, and protective T cell immunity Seminars in Immunol 22:144 (2010)). For example, Toll-like receptors (TLRs) have emerged as powerful sensors of microbial and viral pathogen "danger signals", effectively inducing the innate immune system, and in turn, the adaptive immune system (Bhardwaj and Gnjatic, TLR AGONISTS: Are They Good Adjuvants? Cancer J. 16:382-391 (2010)). Among the TLR agonists, poly-ICLC (a synthetic double-stranded RNA mimic) is one of the most potent activators of myeloid-derived dendritic cells. In a human volunteer study, poly-ICLC has been shown to be safe and to induce a gene expression profile in peripheral blood cells comparable to that induced by one of the most potent live attenuated viral vaccines, the yellow fever vaccine YF-17D (Caskey et al, Synthetic double-stranded RNA induces innate immune responses similar to a live viral vaccine in humans J Exp Med 208:2357 (2011)). Hiltonol®, a GMP preparation of poly-ICLC prepared by Oncovir, Inc, will be utilized as the adjuvant.

Example 2: Target Patient Population

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Patients with stage IIIB, IIIC and IVM1a,b, melanoma have a significant risk of disease recurrence and death, even with complete surgical resection of disease (Balch et al, Final Version of 2009 AJCC Melanoma Staging and Classification J Clin Oncol 27:6199 – 6206 (2009)). An available systemic adjuvant therapy for this patient population is interferon- α (IFN α) which provides a measurable but marginal benefit and is associated with significant, frequently dose-limiting toxicity (Kirkwood et al, Interferon alfa-2b Adjuvant Therapy of High-Risk Resected Cutaneous Melanoma: The Eastern Cooperative Oncology Group Trial EST 1684 J Clin Oncol 14:7-17 (1996); Kirkwood et al , High- and Low-dose Interferon Alpha-2b in High-Risk Melanoma: First Analysis of Intergroup Trial E1690/S9111/C9190 J Clin Oncol 18:2444 – 2458 (2000)). These patients are not immuno-compromised by previous cancer-directed therapy or by active cancer and thus represent an excellent patient population in which to assess the safety and immunological impact of the vaccine. Finally, current standard of care for these patients does

not mandate any treatment following surgery, thus allowing for the 8-10 week window for vaccine preparation.

The target population will be cutaneous melanoma patients with clinically detectable, histologically confirmed nodal (local or distant) or in transit metastasis, who have been fully resected and are free of disease (most of stage IIIB (because of the need to have adequate tumor tissue for sequencing and cell line development, patients with ulcerated primary tumor but micrometastatic lymph nodes (T1-4b, N1a or N2a) will be excluded.), all of stage IIIC, and stage IVM1a, b). These may be patients at first diagnosis or at disease recurrence after previous diagnosis of an earlier stage melanoma.

Tumor harvest: Patients will undergo complete resection of their primary melanoma (if not already removed) and all regional metastatic disease with the intent of rendering them free of melanoma. After adequate tumor for pathological assessment has been harvested, remaining tumor tissue will be placed in sterile media in a sterile container and prepared for disaggregation. Portions of the tumor tissue will be used for whole-exome and transcriptome sequencing and cell line generation and any remaining tumor will be frozen.

Normal tissue harvest: A normal tissue sample (blood or sputum sample) will be taken for whole exome sequencing.

Patients with clinically evident locoregional metastatic disease or fully resectable distant nodal, cutaneous or lung metastatic disease (but absence of unresectable distant or visceral metastatic disease) will be identified and enrolled on the study. Entry of patients prior to surgery is necessary in order to acquire fresh tumor tissue for melanoma cell line development (to generate target cells for in vitro cytotoxicity assays as part of the immune monitoring plan).

Example 3: Dose and Schedule

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For patients who have met all pre-treatment criteria, vaccine administration will commence as soon as possible after the study drug has arrived and has met incoming specifications. For each patient, there will be four separate study drugs, each containing 5 of 20 patient-specific peptides. Immunizations may generally proceed according to the schedule shown in FIG. 5.

Patients will be treated in an outpatient clinic. Immunization on each treatment day will consist of four 1 ml subcutaneous injections, each into a separate extremity in order to target

different regions of the lymphatic system to reduce antigenic competition. If the patient has undergone complete axillary or inguinal lymph node dissection, vaccines will be administered into the right or left midriff as an alternative. Each injection will consist of 1 of the 4 study drugs for that patient and the same study drug will be injected into the same extremity for each cycle. The composition of each 1 ml injection is:

0.75 ml study drug containing 300 µg each of 5 patient-specific peptides

0.25 ml (0.5 mg) of 2 mg/ml poly-ICLC (Hiltonol®)

During the induction/priming phase, patients will be immunized on days 1, 4, 8, 15 and 22. In the maintenance phase, patients will receive booster doses at weeks 12 and 24.

Blood samples may be obtained at multiple time points: pre- (baseline; two samples on different days); day 15 during priming vaccination; four weeks after the induction/priming vaccination (week 8); pre- (week 12) and post- (week 16) first boost; pre- (week 24) and post- (week 28) second boost 50 – 150 ml blood will be collected for each sample (except week 16). The primary immunological endpoint will be at week 16, and hence patients will undergo leukapheresis (unless otherwise indicated based on patient and physician assessment).

Example 4: Immune Monitoring

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The immunization strategy is a "prime-boost" approach, involving an initial series of closely spaced immunizations to induce an immune response followed by a period of rest to allow memory T-cells to be established. This will be followed by a booster immunization, and the T-cell response 4 weeks after this boost is expected to generate the strongest response and will be the primary immunological endpoint. Global immunological response will be initially monitored using peripheral blood mononuclear cells from this time point in an 18 hr *ex vivo* ELISPOT assay, stimulating with a pool of overlapping 15mer peptides (11 aa overlap) comprising all the immunizing epitopes. Pre-vaccination samples will be evaluated to establish the baseline response to this peptide pool. As warranted, additional PBMC samples will be evaluated to examine the kinetics of the immune response to the total peptide mix. For patients demonstrating responses significantly above baseline, the pool of all 15mers will be deconvoluted to determine which particular immunizing peptide(s) were immunogenic. In addition, a number of additional assays will be conducted on a case-by-case basis for appropriate samples:

• The entire 15mer pool or sub-pools will be used as stimulating peptides for intracellular cytokine staining assays to identify and quantify antigen-specific CD4+, CD8+, central memory and effector memory populations

- Similarly, these pools will be used to evaluate the pattern of cytokines secreted by these cells to determine the T_H1 vs T_H2 phenotype
- Extracellular cytokine staining and flow cytometry of unstimulated cells will be used to quantify Treg and myeloid-derived suppressor cells (MDSC).
- If a melanoma cell line is successfully established from a responding patient and the activating epitope can be identified, T-cell cytotoxicity assays will be conducted using the mutant and corresponding wild type peptide
- PBMC from the primary immunological endpoint will be evaluated for "epitope spreading" by using known melanoma tumor associated antigens as stimulants and by using several additional identified mutated epitopes that were not selected to be among the immunogens, as shown in FIG. 6.
- Immuno-histochemistry of the tumor sample will be conducted to quantify CD4+, CD8+, MDSC, and Treg infiltrating populations.

Example 5: Clinical Efficacy in Patients with Metastatic Disease

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Vaccine treatment of patients with metastatic disease is complicated by their need for an effective therapy for the active cancer and the consequent absence of an off treatment time window for vaccine preparation. Furthermore, these cancer treatments may compromise the patient's immune system, possibly impeding the induction of an immune response. With these considerations in mind, settings may be chosen where timing of vaccine preparation fits temporally with other standard care approaches for the particular patient population and/or where such standard care is demonstrably compatible with an immunotherapeutic approach. There are two types of settings that may be pursued:

1. Combination with checkpoint blockade: Checkpoint blockade antibodies have emerged as an effective immunotherapy for metastatic melanoma (Hodi et al, Improved Survival with Ipilimumab in Patients with Metastatic Melanoma NEJM 363:711 – 723 (2010)) and are being actively pursued in other disease settings including non-small cell lung cancer (NSCLC) and renal cell carcinoma (Topalian et al, Safety, Activity, and Immune Correlates of Anti-PD-1

Antibody in Cancer NEJM 366:2443-2454 (2012); Brahmer et al, Safety and Activity of Anti-PD-L1 Antibody in Patients with Advanced Cancer NEJM 366:2455-2465(2012)). Although the mechanism of action is not proven, both reversal of relief from local immunosuppression and enhancement of an immune response are possible explanations. Integrating a powerful vaccine 5 to initiate an immune response with checkpoint blockade antibodies may provide synergies, as observed in multiple animal studies (van Elsas et al Combination immunotherapy of B16 melanoma using anti-cytotoxic T lymphocyte-associated antigen 4 (CTLA-4)and granulocyte/macrophage colony-stimulating factor (GM-CSF)-producing vaccines induces rejection of subcutaneous and metastatic tumors accompanied by autoimmune depigmentation J 10 Exp Med 190:35-366 (1999); Li et al, Anti-programmed death-1 synergizes with granulocyte macrophage colony-stimulating factor -secreting tumor cell immunotherapy providing therapeutic benefit to mice with established tumors Clin Cancer Res 15:1623 – 1634 (2009); Pardoll, D. M. The blockade of immune checkpoints in cancer immunotherapy Nature Reviews Cancer 12:252 – 264 (2012); Curran et al. PD-1 and CTLA-4 combination blockade expands infiltrating T cells and reduces regulatory T and myeloid cells within B16 melanoma tumors. 15 Proc Natl Acad Sci U S A. 2010 Mar 2;107(9):4275-80; Curran et al. Tumor vaccines expressing flt3 ligand synergize with ctla-4 blockade to reject preimplanted tumors. Cancer Res. 2009 Oct 1;69(19):7747-55). Patients can be immediately started on checkpoint blockade therapy while vaccine is being prepared and once prepared, the vaccine dosing can be integrated with antibody 20 therapy, as illustrated in FIG. 7; and

2. Combination with standard treatment regimens exhibiting beneficial immune properties.

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a) Renal cell carcinoma (RCC) patients who present with metastatic disease typically undergo surgical de-bulking followed by systemic treatment, which is commonly with one of the approved tyrosine kinase inhibitors (TKI) such as sunitinib, pazopanib and sorafenib. Of the approved TKIs, sunitinib has been shown to increase T_H1 responsiveness and decrease Treg and myeloid-derived suppressor cells (Finke et al, Sunitinib reverses Type-1 immune suppression and decreases T-regulatory cells in renal cell carcinoma patients Clin Can Res 14:6674 - 6682 (2008); Terme et al, VEGFA-VEGFR pathway blockade inhibits tumor-induced regulatory T cell proliferation in colorectal cancer (Cancer Research Author Manuscript published Online (2102)). The ability to immediately treat patients with an approved therapy that does not

compromise the immune system provides the needed window to prepare the vaccine and could provide synergy with a vaccine therapy. In addition, cyclophosphamide (CTX) has been implicated in multiple animal and human studies to have an inhibitory effect on Treg cells and a single dose of CTX prior to a vaccine has been recently shown to improve survival in RCC patients who responded to the vaccine (Walter et al, Multipeptide immune response to a cancer vaccine IMA901 after single-dose cyclophosphamide associates with longer patient survival Nature Medicine 18:1254- 1260 (2012)). Both of these immune-synergistic approaches have been utilized in a recently completed phase 3 study of a native peptide vaccine in RCC (ClinicalTrials.gov, NCT01265901 IMA901 in Patients Receiving Sunitinib for Advanced/Metastatic Renal Cell Carcinoma):

b) Alternatively, standard treatment of glioblastoma (GBM) involves surgery, recovery and follow-up radiation and low dose temozolomide (TMZ) followed by a four week rest period before initiating standard dose TMZ. This standard treatment provides a window for vaccine preparation followed by initiation of vaccination prior to starting standard dose TMZ. Interestingly, in a study in metastatic melanoma, peptide vaccination during standard dose TMZ treatment increased the measured immune responsiveness compared to vaccination alone, suggesting additional synergistic benefit (Kyte et al, Telomerase peptide vaccination combined with temozolomide: a clinical trial in stage IV melanoma patients Clin Cancer Res 17:4568 (2011)).

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Example 6: Vaccine Preparation

Patient tumor tissue will be surgically resected, and tumor tissue will be disaggregated and separate portions used for DNA and RNA extraction and for patient-specific melanoma cell line development. DNA and/or RNA extracted from the tumor tissue will be used for whole-exome sequencing (e.g., by using the Illumina HiSeq platform) and to determine HLA typing information. It is contemplated within the scope of the invention that missense or neoORF neo-antigenic peptides may be directly identified by protein-based techniques (e.g., mass spectrometry).

Bioinformatics analysis will be conducted as follows. Sequence analysis of the Exome and RNA – SEQ fast Q files will leverage existing bioinformatic pipelines that have been used and validated extensively in large-scale projects such as the TCGA for many patient samples

(e.g., Chapman et al, 2011, Stransky et al, 2011, Berger et al, 2012). There are two sequential categories of analyses: data processing and cancer genome analysis.

Data processing pipeline: The Picard data processing pipeline (picard.sourceforge.net/) was developed by the Sequencing Platform. Raw data extracted from (e.g., Illumina) sequencers for each tumor and normal sample is subjected to the following processes using various modules in the Picard pipeline:

- (i). Quality recalibration: Original base quality scores reported by the Illumina pipeline will be recalibrated based on the read-cycle, the lane, the flow cell tile, the base in question, and the preceding base.
- (ii). Alignment: BWA (Li and Durbin, 2009) will be used to align read pairs to the human genome (hg19).
- (iii). Mark duplicates: PCR and optical duplicates will be identified based on read pair mapping positions and marked in the final bam file.

The output of Picard is a bam file (Li et al, 2009) (samtools.sourceforge.net/SAM1.pdf) that stores the base sequences, quality scores, and alignment details for all reads for the given sample.

Cancer Mutation Detection Pipeline: Tumor and matched normal bam files from the Picard pipeline will be analyzed as described below:

1. Quality Control

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- (i). Sample mix-up during sequencing will be done by comparing initial SNP fingerprinting done on a sample at a few dozen sites with exome sequencing pileups at those sites.
- (ii). Intra-sample tumor/normal mixup will be checked by first comparing the insert size distribution of lanes that correspond to the same library for both tumor and normal samples, and discarding those lanes that have a different distribution. Bioinformatic analysis will be applied to tumor and matched normal exome samples to get the DNA copy number profiles. Tumor samples should also have more copy number variation than the corresponding normals. Lanes corresponding to normal

samples that do not have flat profiles will be discarded, as will tumor lanes that don't have profiles consistent with other lanes from the same tumor sample will be discarded.

(iii). Tumor purity and ploidy will be estimated based on the bioinformatic-generated copy number profiles.

(iv). ContEst (Cibulskis et al, 2011) will be used to determine the level of cross-sample contamination in samples.

2. Local realignment around putative indels

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True somatic and germline small indels with respect to the reference genome often result in misalignment and miscalls of missense mutations and indels. This will be corrected for by doing a local realignment using the GATK IndelRealigner module (on the worldwide web at (www)broadinstitute.org/gatk) (McKenna et al, 2010, Depristo et al, 2011) of all reads that map in the vicinity of putative indels and evaluating them comprehensively to ensure consistency and correctness of indel calls.

3. Identification of somatic single nucleotide variations (SSNVs)

Somatic base pair substitutions will be identified by analyzing tumor and matched normal samples from a patient using a Bayesian statistical framework called muTect (Cibulskis et al, 2013). In the preprocessing step, reads with a preponderance of low quality bases or mismatches to the genome are filtered out. Mutect then computes two log-odds (LOD) scores which encapsulate confidence in presence and absence of the variant in the tumor and normal samples respectively. In the post-processing stage candidate mutations are empirically filtered by various criteria to account for artifacts of capture, sequencing and alignment. One such filter, for example, tests for consistency between distributions of orientations of reads that harbor the mutation and the overall orientation distribution of reads that map to the locus to ensure that there is no strand bias. The final set of mutations will then be annotated with the

Oncotator tool by several fields including genomic region, codon, cDNA and protein changes.

4. Identification of somatic small insertions and deletions

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The local realignment output from section 2.2 will be used to predict candidate somatic and germline indels based on assessment of reads supporting the variant exclusively in tumor or both in tumor and normal bams respectively. Further filtering based on number and distribution of mismatches and base quality scores will be done (McKenna et al, 2010, DePristo et al, 2011). All indels will be manually inspected using the Integrated Genomics Viewer (Robinson et al, 2011) (on the worldwide web at (www)broadinstitute.org/igv) to ensure high-fidelity calls.

5. Gene fusion detection

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The first step in the gene fusion detection pipeline is alignment of tumor RNA-Seq reads to a library of known gene sequences following by mapping of this alignment to genomic coordinates. The genomic mapping helps collapse multiple read pairs that map to different transcript variants that share exons to common genomic locations. The DNA aligned bam file will be queried for read pairs where the two mates map to two different coding regions that are either on different chromosomes or at least 1 MB apart if on the same chromosome. It will also be required that the pair ends aligned in their respective genes be in the direction consistent with coding-->coding 5'-> 3' direction of the (putative) fusion mRNA transcript. A list of gene pairs where there are at least two such 'chimeric' read pairs will be enumerated as the initial putative event list subject to further refinement. Next, all unaligned reads will be extracted from the original bam file, with the additional constraint that their mates were originally aligned and map into one of the genes in the gene pairs obtained as described above. An attempt will then be made to align all such originally unaligned reads to the custom "reference" built of all possible exon-exon junctions (full length,

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boundary-to-boundary, in coding 5'-> 3' direction) between the discovered gene pairs. If one such originally unaligned read maps (uniquely) onto a junction between an exon of gene X and an exon of gene Y, and its mate was indeed mapped to one of the genes X or Y, then such a read will be marked as a "fusion" read. Gene fusion events will be called in cases where there is at least one fusion read in correct relative orientation to its mate, without excessive number of mismatches around the exon:exon junction and with a coverage of at least 10 bp in either gene. Gene fusions between highly homologous genes (ex. HLA family) are likely spurious and will be filtered out.

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6. Estimation of clonality

Bioinformatic analysis may be used to estimate clonality of mutations. For example, the ABSOLUTE algorithm (Carter et al, 2012, Landau et al, 2013) may be used to estimate tumor purity, ploidy, absolute copy numbers and clonality of mutations. Probability density distributions of allelic fractions of each mutation will be generated followed by conversion to cancer cell fractions (CCFs) of the mutations. Mutations will be classified as clonal or subclonal based on whether the posterior probability of their CCF exceeds 0.95 is greater or lesser than 0.5 respectively.

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7. Quantification of expression

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The TopHat suite (Langmead et al, 2009) will be used to align RNA-Seq reads for the tumor and matched normal bams to the hg19 genome. The quality of RNA-Seq data will be assessed by the RNA-SeQC (DeLuca et al, 2012) package. The RSEM tool (Li et al, 2011) will then be used to estimate gene and isoform expression levels. The generated reads per kilobase per million and tau estimates will be used to prioritize neo-antigens identified in each patient as described elsewhere.

8. Validation of mutations in RNA-Seq

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Mutations that will be identified by analysis of whole exome data (section 2.3) will be assessed for presence in the corresponding RNA-Seq tumor bam file of the patient. For each variant locus, a power calculation based on the beta-binomial distribution will be performed to ensure that there is at least 80% power to detect it in the RNA-Seq data. A capture identified mutation will be considered validated if there are at least 2 reads harboring the mutation for adequately powered sites.

10 Selection of Tumor-Specific Mutation-Containing Epitopes: All missense mutations and neoORFs will be analyzed for the presence of mutation-containing epitopes using the neural-network based algorithm netMHC, provided and maintained by the Center for Biological Sequence Analysis, Technical University of Denmark, Netherlands. This family of algorithms were rated the top epitope prediction algorithms based on a competition recently completed among a series of related approaches (ref). The algorithms were trained using an artificial neural network based approach on multiple different human HLA A and B alleles utilizing over 100,000 measured binding and non-binding interactions.

The accuracy of the algorithms were evaluated by conducting predictions from mutations found in CLL patients for whom the HLA allotypes were known. The included allotypes were A0101, A0201, A0310, A1101, A2402, A6801, B0702, B0801, B1501. Predictions were made for all 9mer and 10 mer peptides spanning each mutation using netMHCpan in mid-2011. Based on these predictions, seventy-four (74) 9mer peptides and sixty-three (63) 10mer peptides, most with predicted affinities below 500 nM, were synthesized and the binding affinity was measured using a competitive binding assay (Sette).

The predictions for these peptides were repeated in March 2013 using each of the most up to date versions of the netMHC servers (netMHCpan, netMHC and netMHCcons). These three algorithms were the top rated algorithms among a group of 20 used in a competition in 2012 (Zhang et al). The observed binding affinities were then evaluated with respect to each of the new predictions. For each set of predicted and observed values, the % of correct predictions for each range is given, as well as the number of samples. The definition for each range is as follows:

0-150 Predicted to have an affinity equal to or lower than 150 nM and measured to have an affinity equal to or lower than 150 nM.

0-150*: Predicted to have an affinity equal to or lower than 150 nM and measured to have an affinity equal to or lower than 500 nM.

151 - 500 nM: Predicted to have an affinity greater than 150 nM but equal to or lower than 500 nM and measured to have an affinity equal to or below 500 nM.

FN (> 500 nM): False Negatives – Predicted to have an affinity greater than 500 nM but measured to have an affinity equal to or below 500 nM.

For 9mer peptides (Table 1), there was little difference between the algorithms, with the slightly higher value for the 151-500 nM range for netMHC cons not judged to be significant because of the low number of samples.

Table 1

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Range (nM)	9mer PAN	9mer netMHC	9mer CONS
0.450	76%	78%	76%
0-150	(33)	(37)	(34)
0.450*	91%	89%	88%
0-150*	(33)	(37)	(34)
151 500	50%	50%	62%
151-500	(28)	(14)	(13)
EN (> E00)	38%	39%	41%
FN (>500)	(13)	(23)	(27)

15 For 10mer peptides (Table 2), again there was little difference between the algorithms except that netMHC produced significantly more false positives than netMHCpan or netMMHCcons. However, the precision of the 10mer predictions is slightly lower in the 0 – 150 nM and 0 – 150* nM ranges and significantly lower in the 151-500 nM range, compared to the 9mers.

Table 2.

Range (nM)	10mer PAN	10mer netMHC	10mer CONS
0.150	53%	50%	59%
0-150	(19)	(16)	(17)
0-150*	68%	69%	76%
	(19)	(16)	(17)
151 500	35%	42%	35%
151-500	(26)	(12)	(23)
[N] (> E00)	11%	23%	13%
FN (>500)	(18)	(35)	(23)

For 10mers, only predictions in the 0-150 nM range will be utilized due to the lower than 50% precision for binders in the 151-500 nM range.

The number of samples for any individual HLA allele was too small to draw any conclusions regarding accuracy of the prediction algorithm for different alleles. Data from the largest available subset (0 - 150* nM; 9mer) is shown in Table 3 as an example.

Table 3

Allele	Fraction		
	correct		
A0101	2/2		
A0201	9/11		
A0301	5/5		
A1101	4/4		
A2402	0/0		
A6801	3/4		
B0702	4/4		
B0801	1/2		
B1501	2/2		

Only predictions for HLA A and B alleles will be utilized as there is little available data on which to judge accuracy of predictions for HLA C alleles (Zhang et al).

An evaluation of melanoma sequence information and peptide binding predictions was conducted using information from the TCGA database. Information from 220 melanomas from different patients revealed that on average there were approximately 450 missense and 5 neoORFs per patient. 20 patients were selected at random and the predicted binding affinities were calculated for all the missense mutations using netMHC (Lundegaard et al Prediction of epitopes using neural network based methods J Immunol Methods 374:26 (2011)). As the HLA allotypes were unknown for these patients, the number of predicted binding peptides per allotype was adjusted based on the frequency of that allotype (Bone Marrow Registry dataset for the expected affected dominant population in the geographic area [Caucasian for melanoma]) to generate a predicted number of actionable mutant epitopes per patient. For each of these mutant epitopes (MUT), the corresponding native (WT) epitope binding was also predicted. Utilizing a single peptide for predicted missense binders with Kd ≤ 500 nM and a WT/MUT Kd ratio of >5X and over-lapping peptides spanning the full length of each neoORF, 80% (16 of 20) of patients were predicted to have at least 20 peptides appropriate for vaccination. For a quarter of the patients, neoORF peptides could constitute nearly half to all of the 20 peptides. Thus, there is an adequate mutational load in melanoma to expect a high proportion of patients to generate an adequate number of immunogenic peptides.

Example 7: Prioritization of Immunizing Peptides

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Peptides for immunization may be prioritized based on a number of criteria: neoORF vs. missense, predicted Kd for the mutated peptide, the comparability of predicted affinity for the native peptide compared to the mutated peptide, whether the mutation occurs in an oncogenic driver gene or related pathway, and # of RNA-Seq reads (see e.g., FIG. 8).

As shown in FIG. 8, peptides derived from segments of neoORF mutations that are predicted to bind (Kd < 500 nM) may be given the highest priority based on the absence of tolerance for these entirely novel sequences and their exquisite tumor-specificity.

The similar class of missense mutations in which the native peptide is not predicted to bind (Kd > 1000 nM) and the mutated peptide is predicted to bind with strong/moderate affinity (Kd < 150 nM) may be given the next highest priority. This class (Group I discussed above) represents approximately 20% of naturally observed T-cell responses.

The third highest priority may be given to the more tightly binding (< 150 nM) subset of the Group II class discussed above. This class is responsible for approximately almost 2/3 of naturally observed T-cell responses.

All the remaining peptides derived from the neoORF mutations may be given the fourth priority. Despite not being predicted to bind, these are included based on the known false negative rate, potential binding to HLA-C, potential for presence of Class II epitopes and the high value of utilizing totally foreign antigens.

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The fifth priority may be given to the subset of Group II with lower predicted binding affinities (150 - 500 nM). This class is responsible for approximately 10% of the naturally observed T-cell responses.

As the predicted affinity decreases, higher stringency may be applied to expression levels. Within each grouping, peptides may be ranked based on binding affinity (e.g., the lowest Kd may have the highest priority). Within a given grouping of missense mutations, oncogenic driver mutations may be given higher priority. A normal human peptidome library of ~12.6 million unique 9 and 10 mers curated from all known human protein sequences (HG19) has been created. Prior to final selection, any potential predicted epitopes derived from a missense mutation and all neoORF regions may be screened against this library, and perfect matches may be excluded. As discussed below, particular peptides predicted to have deleterious biochemical properties may be eliminated or modified.

According to the techniques herein, RNA levels may be analyzed to assess neoantigen expression. For example, RNA-Seq read-count may be used as a proxy to estimate neoantigen expression. However, there is no currently available information to assess the minimum RNA expression level required in a tumor cell needed to initiate cytolysis. Even the level of expression from "pioneer" translation of messages destined for nonsense mediated decay may be sufficient for target generation. Accordingly, the techniques herein initially set broad acceptance limits for RNA levels that may vary inversely with the priority group. As the predicted affinity decreases, higher stringency may be applied to expression levels. One of skill in the art will appreciate that as additional information becomes available, such limits may be adjusted.

Because of the high value of neoORFs as targets due to their novelty and exquisite tumor specificity, neoORFs with predicted binding epitopes ($Kd \le 500 \text{ nM}$) may be utilized even if there are no detectable mRNA molecules by RNA-Seq (Rank 1). Regions of neoORFs without

predicted binding epitopes (> 500 nM), may generally be utilized only if some level of RNA expression is detected (Rank 4). All missense mutations with strong to intermediate predicted MHC binding affinity (\leq 150 nM) may generally be utilized unless there were no RNA-Seq reads (Ranks 2 and 3). For missense mutations with lower predicted binding affinity (150 - \leq 500 nM), these will likely be utilized only if a slightly higher level of RNA expression is detected (Rank 5).

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Oncogenic drivers may represent a high priority group. For example, within a given grouping of missense mutations, oncogenic driver mutations may be of higher priority. This approach is based on the observed down-regulation of genes that are targeted by immune pressure (e.g., immunoediting). In contrast to other immune targets where down-regulation may not have a deleterious effect of cancer cell growth, continued expression of oncogenic driver genes may be crucial to cancer cell survival, thus shutting off a pathway of immune escape. Exemplary oncogenic drivers are listed in Table 3-1 (see e.g., Vogelstein et al; GOTERM_BP Assignment of genes to Gene Ontology Term - Biological Function on the worldwide web at (www)geneontology.org; BIOCARTA Assignment of genes to signaling pathways, on the worldwide web at (www)genome.jp/krgg/pathway.html; REACTOME Assignment of genes to pathways according to REACTOME pathways and gene interactions, on the worldwide web at (www)reactome.org).

Table 3-1 Exemplary Oncogenic Driver Genes

<u>Gene</u> Symbol	Gene Name	# Mutated Tumor Samples**	Onco- gene score*	Tumor Suppressor Gene score*	Classification*	<u>Core</u> pathway	Process
ABL1	c-abl oncogene 1, receptor tyrosine kinase	851	93%	0%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
AKT1	v-akt murine thymoma viral oncogene homolog 1	155	93%	1%	Oncogene	PI3K	Cell Survival
ALK	anaplastic lymphoma receptor tyrosine kinase	189	72%	1%	Oncogene	PI3K; RAS	Cell Survival
AR	androgen receptor	23	54%	0%	Oncogene	Transcriptional Regulation	Cell Fate
BCL2	B-cell CLL/lymphoma 2	45	27%	1%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
BRAF	v-raf murine sarcoma viral oncogene homolog B1	24288	100%	0%	Oncogene	RAS	Cell Survival
CARD11	caspase recruitment domain family, member 11	74	30%	1%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
CBL	Cas-Br-M (murine) ecotropic retroviral transforming sequence	168	57%	9%	Oncogene	PI3K; RAS	Cell Survival
CRLF2	cytokine receptor-like factor 2	10	100%	0%	Oncogene	STAT	Cell Survival
CSF1R	colony stimulating factor 1 receptor	48	50%	15%	Oncogene	PI3K; RAS	Cell Survival
CTNNB1	catenin (cadherin- associated protein), beta 1, 88kDa	3262	92%	1%	Oncogene	APC	Cell Fate
DNMT1	DNA (cytosine-5-)- methyltransferase 1	22	36%	5%	Oncogene	Chromatin Modification	Cell Fate
DNMT3A	DNA (cytosine-5-)- methyltransferase 3 alpha	788	74%	12%	Oncogene	Chromatin Modification	Cell Fate
EGFR	epidermal growth factor receptor (erythroblastic leukemia viral (v-erb- b) oncogene homolog, avian)	10628	97%	0%	Oncogene	PI3K; RAS	Cell Survival
ERBB2	v-erb-b2 erythroblastic leukemia viral oncogene homolog 2, neuro/glioblastoma derived oncogene homolog (avian)	164	67%	3%	Oncogene	PI3K; RAS	Cell Survival
EZH2	enhancer of zeste homolog 2 (Drosophila)	276	67%	12%	Oncogene	Chromatin Modification	Cell Fate
FGFR2	fibroblast growth factor receptor 2	121	49%	6%	Oncogene	PIЗК; RAS ; STAT	Cell Survival
FGFR3	fibroblast growth factor receptor 3	2948	99%	0%	Oncogene	PI3K; RAS ; STAT	Cell Survival

<u>Gene</u> Symbol	<u>Gene Name</u>	# Mutated Tumor Samples**	Onco- gene score*	Tumor Suppressor Gene score*	Classification*	<u>Core</u> pathway	Process
FLT3	fms-related tyrosine kinase 3	11520	98%	0%	Oncogene	RAS; PI3K; STAT	Cell Survival
FOXL2	forkhead box L2	330	100%	0%	Oncogene	TGF-β	Cell Fate
GATA2	GATA binding protein 2	45	53%	4%	Oncogene	NOTCH, TGF-β	Cell Fate
GNA11	guanine nucleotide binding protein (G protein), alpha 11 (Gq class)	110	92%	1%	Oncogene	PI3K; RAS; MAPK	Cell Survival
GNAQ	guanine nucleotide binding protein (G protein), q polypeptide	245	95%	1%	Oncogene	PI3K;RAS; MAPK	Cell Survival
GNAS	GNAS complex locus	422	93%	2%	Oncogene	APC; PI3K; TGF- β, RAS	Cell Survival/C ell Fate
НЗГЗА	H3 histone, family 3B (H3.3B); H3 histone, family 3A pseudogene; H3 histone, family 3A; similar to H3 histone, family 3B; similar to histone H3.3B	122	93%	0%	Oncogene	Chromatin Modification	Cell Fate
HIST1H3B	histone cluster 1, H3j; histone cluster 1, H3i; histone cluster 1, H3h; histone cluster 1, H3g; histone cluster 1, H3f; histone cluster 1, H3e; histone cluster 1, H3d; histone cluster 1, H3c; histone cluster 1, H3b; histone cluster 1, H3a; histone cluster 1, H2ad; histone cluster 2, H3a; histone cluster 2, H3c; histone cluster 2, H3c; histone cluster 2,	25	60%	0%	Oncogene	Chromatin Modification	Cell Fate
HRAS	v-Ha-ras Harvey rat sarcoma viral oncogene homolog	812	96%	0%	Oncogene	RAS	Cell Survival
IDH1	isocitrate dehydrogenase 1 (NADP+), soluble	4509	100%	0%	Oncogene	Chromatin Modification	Cell Fate
IDH2	isocitrate dehydrogenase 2 (NADP+), mitochondrial	1029	99%	0%	Oncogene	Chromatin Modification	Cell Fate
JAK1	Janus kinase 1	61	26%	18%	Oncogene	STAT	Cell Survival
JAK2	Janus kinase 2	32692	100%	0%	Oncogene	STAT	Cell Survival
JAK3	Janus kinase 3	89	60%	6%	Oncogene	STAT	Cell Survival

<u>Gene</u> Symbol	<u>Gene Name</u>	# Mutated Tumor Samples**	Onco- gene score*	Tumor Suppressor Gene score*	<u>Classification*</u>	<u>Core</u> pathway	<u>Process</u>
КІТ	similar to Mast/stem cell growth factor receptor precursor (SCFR) (Proto- oncogene tyrosine- protein kinase Kit) (c- kit) (CD117 antigen); v-kit Hardy- Zuckerman 4 feline sarcoma viral oncogene homolog	4720	90%	0%	Oncogene	PI3K; RAS; STAT	Cell Survival
KLF4	Kruppel-like factor 4	61	80%	4%	Oncogene	Transcriptional Regulation; WNT	Cell Fate
KRAS	v-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog	23261	100%	0%	Oncogene	RAS	Cell Survival
MAP2K1	mitogen-activated protein kinase kinase 1	13	67%	0%	Oncogene	RAS	Cell Survival
MED12	mediator complex subunit 12	337	84%	0%	Oncogene	Cell Cycle/Apoptosis; TGF-β	Cell Survival
MET	met proto-oncogene (hepatocyte growth factor receptor)	159	61%	4%	Oncogene	PI3K; RAS	Cell Survival
MPL	myeloproliferative leukemia virus oncogene	531	96%	0%	Oncogene	STAT	Cell Survival
MYD88	myeloid differentiation primary response gene (88)	134	92%	1%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
NFE2L2	nuclear factor (erythroid-derived 2)- like 2	102	74%	1%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
NRAS	neuroblastoma RAS viral (v-ras) oncogene homolog	2738	99%	0%	Oncogene	RAS	Cell Survival
PDGFRA	platelet-derived growth factor receptor, alpha polypeptide	653	84%	1%	Oncogene	PI3K; RAS	Cell Survival
PIK3CA	phosphoinositide-3- kinase, catalytic, alpha polypeptide	4560	95%	1%	Oncogene	РІЗК	Cell Survival
PPP2R1A	protein phosphatase 2 (formerly 2A), regulatory subunit A, alpha isoform	86	85%	2%	Oncogene	Cell Cycle/Apoptosis	Cell Survival
PTPN11	protein tyrosine phosphatase, non- receptor type 11; similar to protein tyrosine phosphatase, non-receptor type 11	410	90%	0%	Oncogene	RAS	Cell Survival
RET	ret proto-oncogene	500	86%	1%	Oncogene	RAS; PI3K	Cell Survival

<u>Gene</u> Symbol	Gene Name	# Mutated Tumor Samples**	Onco- gene score*	Tumor Suppressor Gene score*	Classification*	<u>Core</u> pathway	<u>Process</u>
SETBP1	SET binding protein 1	95	25%	4%	Oncogene	Chromatin Modification; Replication	Cell Fate
SF3B1	splicing factor 3b, subunit 1, 155kDa	516	91%	0%	Oncogene	Transcriptional Regulation	Cell Fate
SMO	smoothened homolog (Drosophila)	34	51%	3%	Oncogene	нн	Cell Fate
SPOP	speckle-type POZ protein	35	66%	3%	Oncogene	Chromatin Modification; HH	Cell Fate
SRSF2	SRSF2 serine/arginine-rich splicing factor 2	273	95%	2%	Oncogene	Transcriptional Regulation	Cell Fate
TSHR	thyroid stimulating hormone receptor	301	86%	0%	Oncogene	PI3K; MAPK	Cell Survival
U2AF1	U2 small nuclear RNA auxiliary factor 1	96	92%	1%	Oncogene	Transcriptional Regulation	Cell Fate

Example 8: Peptide Production and Formulation

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GMP neo-antigenic peptides for immunization will be prepared by chemical synthesis Merrifield RB: Solid phase peptide synthesis. I. The synthesis of a tetrapeptide. J. Am. Chem. Soc. 85:2149-54, 1963) in accordance with FDA regulations. Three development runs have been conducted of 20 ~20-30mer peptides each. Each run was conducted in the same facility and utilized the same equipment as will be used for the GMP runs, utilizing draft GMP batch records. Each run successfully produced > 50 mg of each peptide, which were tested by all currently planned release tests (e.g., Appearance, Identify by MS, Purity by RP-HPLC, Content by Elemental Nitrogen, and TFA content by RP-HPLC) and met the targeted specification where appropriate. The products were also produced within the timeframe anticipated for this part of the process (approximately 4 weeks). The lyophilized bulk peptides were placed on a long term stability study and will be evaluated at various time points up to 12 months.

Material from these runs has been used to test the planned dissolution and mixing approach. Briefly, each peptide will be dissolved at high concentration (50 mg/ml) in 100% DMSO and diluted to 2 mg/ml in an aqueous solvent. Initially, it was anticipated that PBS would be used as a diluent, however, a salting out of a small number of peptides caused a visible cloudiness. D5W (5% dextrose in water) was shown to be much more effective; 37 of 40 peptides were successfully diluted to a clear solution. The only problematic peptides are very hydrophobic peptides. The predicted biochemical properties of planned immunizing peptides

will be evaluated and synthesis plans may be altered accordingly (using a shorter peptide, shifting the region to be synthesized in the N- or C-terminal direction around the predicted epitope, or potentially utilizing an alternate peptide). Ten separate peptides in DMSO/D5W were subjected to two freeze/thaw cycles and showed full recovery. Two individual peptides were dissolved in DMSO/D5W and placed on stability at two temperatures (-20°C and -80°C). These peptides will be evaluated (RP-HPLC, MS and pH) for up to 6 months. To date, both peptides are stable at the 12 week time point with additional time points at 24 weeks to be evaluated.

As shown in FIG. 9, the design of the dosage form process is to prepare 4 pools of patient-specific peptides consisting of 5 peptides each. A RP-HPLC assay has been prepared and qualified to evaluate these peptide mixes. This assay achieves good resolution of multiple peptides within a single mix and can also be used to quantitate individual peptides.

Membrane filtration (0.2 μm pore size) will be used to reduce bioburden and conduct final filter sterilization. Four different appropriately sized filter types were initially evaluated and the Pall, PES filter (# 4612) was selected. To date, 4 different mixtures of 5 different peptides each have been prepared and individually filtered sequentially through two PES filters. Recovery of each individual peptide was evaluated utilizing the RP-HPLC assay. For 18 of the 20 peptides, the recovery after two filtrations was >90%. For two highly hydrophobic peptides, the recovery was below 60% when evaluated at small scale but were nearly fully recovered (87 and 97%) at scale. As stated above, approaches will be undertaken to limit the hydrophobic nature of the sequences selected.

GMP neo-antigenic peptides for immunization will be prepared by chemical synthesis Merrifield RB: Solid phase peptide synthesis. I. The synthesis of a tetrapeptide. J. Am. Chem. Soc. 85:2149-54, 1963) in accordance with FDA regulations.

25 Example 9: Endpoint Assessment

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The primary immunological endpoint of this study will be the assessment of T cell response measured by ex vivo IFN-γ ELISPOT. IFN-γ secretion occurs as a result of the recognition of cognate peptides or mitogenic stimuli by CD4⁺ and/or CD8⁺ T –cells. A multitude of different CD4⁺ and CD8⁺ determinants will likely be presented to T cells in vivo since the 20-30mer peptides used for vaccination should undergo processing into smaller peptides by antigen presenting cells. Without being bound by theory, it is believed that the combination of

personalized neo-antigen peptides, which are novel to the immune system and thus not subject to the immune-dampening effects of self-tolerance, and the powerful immune adjuvant poly-ICLC will induce strong CD4⁺ and/or CD8⁺ responses. The expectation is therefore that T cell responses are detectable ex vivo i.e. without the need for in vitro expansion of epitope specific T cells through short-term culture. Patients will initially be evaluated using the total pool of peptide immunogens as stimulant in the ELISPOT assay. For patients demonstrating a robust positive response, the precise immunogenic peptide(s) will be determined in follow-up analysis. The IFN-y ELISPOT is generally accepted as a robust and reproducible assay to detect ex vivo T cell activity and determine specificity. In addition to the analysis of the magnitude and determinant mapping of the T cell response in peripheral blood monocytes, other aspects of the immune response induced by the vaccine are critical and will be assessed. These evaluations will be performed in patients who exhibit an ex vivo IFN-γ ELISPOT response in the screening assay. They include the evaluation of T cell subsets (Th1 versus Th2, T effector versus memory cells), analysis of the presence and abundance of regulatory cells such as T regulatory cells or myeloid derived suppressor cells, and cytotoxicity assays if patient-specific melanoma cells lines are successfully established.

Example 10: Peptide synthesis

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GMP peptides will be synthesized by standard solid phase synthetic peptide chemistry and purified by RP-HPLC. Each individual peptide will be analyzed by a variety of qualified assays to assess appearance (visual), purity (RP-HPLC), identity (by mass spectrometry), quantity (elemental nitrogen), and trifluoroacetate counterion (RP-HPLC) and released.

The personalized neoantigen peptides may be comprised of up to 20 distinct peptides unique to each patient. Each peptide may be a linear polymer of ~20 - ~30 L-amino acids joined by standard peptide bonds. The amino terminus may be a primary amine (NH2-) and the carboxy terminus is a carbonyl group (-COOH). The standard 20 amino acids commonly found in mammalian cells are utilized (alanine, arginine, asparagine, aspartic acid, cysteine, glutamine, glutamic acid, glycine, histidine, isoleucine, leucine lysine, methionine, phenylalanine, proline, serine, threonine, tryptophan, tyrosine, valine). The molecular weight of each peptide varies based on its length and sequence and is calculated for each peptide.

Personalized neoantigen peptides may be supplied as a box containing 2 ml Nunc Cryo vials with color-coded caps, each vial containing approximately 1.5 ml of a frozen DMSO/D5W

solution containing up to 5 peptides at a concentration of 400 ug/ml. There may be 10 - 15 vials for each of the four groups of peptides. The vials are to be stored at -80oC until use. Ongoing stability studies support the storage temperature and time.

Storage and Stability: The personalized neoantigen peptides are stored frozen at -80oC. The thawed, sterile filtered, in process intermediates and the final mixture of personalized neoantigen peptides and poly-ICLC can be kept at room temperature but should be used within 4 hours.

Compatibility: The personalized neoantigen peptides will be mixed with 1/3 volume poly-ICLC just prior to use.

Example 11: Administration

Following mixing with the personalized neo-antigenic peptides/polypeptides, the vaccine (e.g., peptides + poly-ICLC) is to be administered subcutaneously.

Preparation of personalized neo-antigenic peptides/polypeptides pools: peptides will be mixed together in 4 pools of up to 5 peptides each. The selection criteria for each pool will be based on the particular MHC allele to which the peptide is predicted to bind.

Pool Composition: The composition of the pools will be selected on the basis of the particular HLA allele to which each peptide is predicted to bind. The four pools will be injected into anatomic sites that drain to separate lymph node basins. This approach was chosen in order to potentially reduce antigenic competition between peptides binding to the same HLA allele as much as possible and involve a wide subset of the patient's immune system in developing an immune response. For each patient, peptides predicted to bind up to four different HLA A and B alleles will be identified. Some neoORF derived peptides will not be associated with any particular HLA allele. The approach to distributing peptides to different pools will be to spread each set of peptides associated with a particular HLA allele over as many of the four pools as possible. It is highly likely there will be situations where there will be more than 4 predicted peptides for a given allele, and in these cases it will be necessary to allocate more than one peptide associated with a particular allele to the same pool. Those neoORF peptides not associated with any particular allele will be randomly assigned to the remaining slots. An example is shown below:

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A1 A2 B1 B2 X	HLA A	A0101 A1101 B0702 B6801 E (neoORF)	5 pep 2 pep 7 pep	etides etides etides etides etides
Pool#	1	2	3	4
	B2	B2	B2	B2
	B2	B2	B2	A2
	A2	A2	A2	A2
	A1	A1	A1	B1
	B1	X	X	X

Peptides predicted to bind to the same MHC allele will be placed into separate pools whenever possible. Some of the neoORF peptides may not be predicted to bind to any MHC allele of the patient. These peptides will still be utilized however, primarily because they are completely novel and therefore not subject to the immune-dampening effects of central tolerance and therefore have a high probability of being immunogenic. NeoORF peptides also carry a dramatically reduced potential for autoimmunity as there is no equivalent molecule in any normal cell. In addition, there can be false negatives arising from the prediction algorithm and it is possible that the peptide will contain a HLA class II epitope (HLA class II epitopes are not reliably predicted based on current algorithms). All peptides not identified with a particular HLA allele will be randomly assigned to the individual pools. The amounts of each peptide are predicated on a final dose of 300 µg of each peptide per injection.

For each patient, four distinct pools (labeled "A", "B", "C" and "D") of 5 synthetic peptides each will have been prepared manufacturer and stored at -80°C. On the day of immunization, the complete vaccine consisting of the peptide component(s) and poly-ICLC will be prepared in a laminar flow biosafety cabinet in the research pharmacy. One vial each (A, B, C and D) will be thawed at room temperature and moved into a biosafety cabinet for the remaining steps. 0.75 ml of each peptide pool will be withdrawn from the vial into separate syringes. Separately, four 0.25 ml (0.5 mg) aliquots of poly-ICLC will be withdrawn into separate syringes. The contents of each peptide-pool containing syringe will then be gently

mixed with a 0.25 ml aliquot of poly-ICLC by syringe-to-syringe transfer. The entire one ml of the mixture will be used for injection. These 4 preparations will be labeled "study drug A", "study drug B", "study drug C", and "study drug D".

Injections: At each immunization, each of the 4 study drugs will be injected subcutaneously into one extremity. Each individual study drug will be administered to the same extremity at each immunization for the entire duration of the treatment (i.e. study drug A will be injected into left arm on day 1, 4, 8 etc., study drug B will be injected into right arm on days 1, 4, 8 etc.). Alternative anatomical locations for patients who are status post complete axillary or inguinal lymph node dissection are the left and right midriff, respectively.

Vaccine will be administered following a prime/boost schedule. Priming doses of vaccine will be administered on days 1, 4, 8, 15, and 22 as shown above. In the boost phase, vaccine will be administered on days 85 (week 13) and 169 (week 25).

All patients receiving at least one dose of vaccine will be evaluable for toxicity. Patients will be evaluable for immunologic activity if they have received all vaccinations during the induction phase and the first vaccination (boost) during the maintenance phase.

Example 12: Pharmacodynamic Studies

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The immunization strategy is a "prime-boost" approach, involving an initial series of closely spaced immunizations to induce an immune response followed by a period of rest to allow memory T-cells to be established. This will be followed by a booster immunization, and the T-cell response 4 weeks after this boost (16 weeks after the first vaccination) is expected to generate the strongest response and will be the primary immunological endpoint. Immune monitoring will be performed in a step-wise fashion as outlined below to characterize the intensity and quality of the elicited immune responses. Peripheral blood will be collected and PBMC will be frozen at two separate time points prior to the first vaccination (baseline) and at different time points thereafter as illustrated in Schema B and specified in the study calendar. Immune monitoring in a given patient will be performed after the entire set of samples from the induction phase and the maintenance phase, respectively, have been collected. If sufficient

tumor tissue is available, a portion of the tumor will be used to develop autologous melanoma cell lines for use in cytotoxic T-cell assays.

Example 13: Screening ex vivo IFN-γ ELISPOT

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For each patient, a set of screening peptides will be synthesized. The screening peptides will be 15 amino acids in length (occasionally a 16mer or 17mer will be used), overlapping by 11 amino acids and covering the entire length of each peptide or the entire length of the neoORF for neoORF-derived peptides. The entire set of patient-specific screening peptides will be pooled together at approximately equal concentration and a portion of each peptide will also be stored individually. Purity of the peptide pool will be ascertained by testing PBMC from 5 healthy donors with established low background in ex vivo IFN-γ ELISPOTs. Initially, PBMC obtained at baseline and at week 16 (the primary immunological endpoint) will be stimulated for 18 hours with the complete pool of overlapping 15-mer peptides (11 amino acids overlap) to examine the global response to the peptide vaccine. Subsequent assays may utilize PBMC collected at other time points as indicated. If no response is identified at the primary immunological endpoint using the ex vivo IFN-γ ELISPOT assay, PBMC will be stimulated with the peptide pool for a longer time period (up to 10 days) and re-analyzed.

Example 14: Deconvolution of epitopes in follow-up ex vivo IFN-γ ELISPOT assays.

Once an ex vivo IFN- γ ELISPOT response elicited by an overlapping peptide pool is observed (defined as at least 55 spot forming units / 10^6 PBMC or increased at least 3 times over baseline), the particular immunogenic peptide eliciting this response will be identified by deconvoluting the peptide pool based into sub-pools based on the immunizing peptides and repeating the ex vivo IFN- γ ELISPOT assays. For some responses, an attempt will be made to precisely characterize the stimulating epitope by utilizing overlapping 8-10 mer peptides derived from confirmed, stimulating peptides in IFN- γ ELISPOT assays. Additional assays may be conducted on a case-by case basis for appropriate samples. For example,

 The entire 15mer pool or sub-pools will be used as stimulating peptides for intracellular cytokine staining assays to identify and quantify antigen-specific CD4+, CD8+, central memory and effector memory populations

- Similarly, these pools will be used to evaluate the pattern of cytokines secreted by these cells to determine the T_H1 vs T_H2 phenotype
- Extracellular cytokine staining and flow cytometry of unstimulated cells will be used to quantify Treg and myeloid-derived suppressor cells (MDSC).
- If a melanoma cell line is successfully established from a responding patient and the activating epitope can be identified, T-cell cytotoxicity assays will be conducted using the mutant and corresponding wild type peptide
- PBMC from the primary immunological endpoint will be evaluated for "epitope spreading" by using known melanoma tumor associated antigens as stimulants and by using several additional identified mutated epitopes that were not selected to be among the immunogens

Immuno-histochemistry of tumor samples will be conducted to quantify CD4+, CD8+, MDSC, and Treg infiltrating populations.

Example 15: Pipeline for the systematic identification of tumor neoantigens

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Recent advances in sequencing technologies and peptide epitope predictions were leveraged to generate a two-step pipeline to systematically discover candidate tumor-specific HLA-bound neoantigens. As depicted in FIG. 10, this approach starts with DNA sequencing of tumors (e.g., by either whole-exome (WES) or whole-genome sequencing (WGS)) in parallel with matched normal DNA to comprehensively identify non-synonymous somatic mutations (see e.g., Lawrence et al. 2013; Cibulski et al. 2012). Next, candidate tumor specific mutated peptides generated by tumor mutations with the potential to bind personal class I HLA proteins, and hence be presented to CD8⁺ T cells, may be predicted using prediction algorithms such as, for example, NetMHCpan (see e.g., Lin 2008; Zhang 2011). Candidate peptide antigens were further evaluated based on experimental validation of their binding to HLA and expression cognate mRNAs in autologous leukemia cells.

This pipeline was applied to a large dataset of sequenced CLL samples (see e.g., Wang et

al. 2011). From 91 cases that were sequenced by either WES or WGS, a total of 1838 nonsynonymous mutations were discovered in protein-coding regions, corresponding to a mean somatic mutation rate of 0.72 (±0.36 s.d.) per megabase (range, 0.08 to 2.70), and a mean of 20 non-synonymous mutations per patient (range, 2 to 76) (see e.g., Wang et al. 2011). Three general classes of mutations were identified that would be expected to generate regions of amino acid changes and hence potentially be recognized immunologically. The most abundant class included missense mutation that cause single amino acid (aa) changes, representing 90% of somatic mutations per CLL. Of 91 samples, 99% harbored missense mutations and 69% had between 10-25 missense mutations (see e.g., FIG. 2A). The other two classes of mutations, frameshifts and splice-site mutations (mutations at exon-intron junctions) have the potential to generate longer stretches of novel amino acid sequences entirely specific to the tumor (neo-open reading frames, or neoORFs), with a higher number of neoantigen peptides per given alteration (compared to missense mutations). However, consistent with data from other cancer types, neoORF-generating mutations were approximately 10 fold less abundant than missense mutations in CLL (see e.g., FIGS. 2B-C). Given the prevalence of missense mutations, subsequent experimental studies was focused on the analysis of neoepitopes generated by missense mutations.

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Example 16: Somatic missense mutations generate neopeptides predicted to bind to personal HLA class I alleles

T cell recognition of peptide epitopes by the T cell receptor (TCR) requires the display of peptides bound within the binding groove of HLA molecules on the surface of antigen-presenting cells. Recent comparative studies across the >30 available class I prediction algorithms have shown NetMHCpan to consistently perform with high sensitivity: and specificity across HLA alleles (see e.g., Zhang et al. 2011).

The NetMHCpan algorithm was tested against a set of 33 known mutated epitopes that were originally identified in the literature on the basis of their functional activity (i.e., ability to stimulate antitumor cytolytic T cell responses) or were characterized as immunogenic minor histocompatibility antigens to determine whether the algorithm would correctly predict binding for the 33 known mutated epitopes (see e.g., Tables 4 and 5). Tables 4 and 5 show HLA-peptide

binding affinities of known functionally derived immunogenic mutated epitopes across human cancers using NetMHCpan. Table 4 shows epitopes from missense mutations (NSCLC: non-small cell lung cancer; MEL: melanoma; CLL: chronic lymphocytic leukemia; RCC: clear cell renal carcinoma; BLD: bladder cancer; NR: not reported;). Yellow: IC50 < 150 nM, green: IC50 150-500 nM and grey: IC50 > 500 nM.

Table 4

Clinical response HLA Mutated
Response Allele MUT>>>WT
Yes Yes A*02:01
Yes Yes A*01:01
NR Yes A*02:01
Yes Yes A*11:01
Yes Yes A*68:01
Yes Yes B*07:02
Yes Yes A*02:01
Yes Yes A*03:01
NR Yes A*68:02
Yes Yes A*02:01
NR Yes A*02:01
Yes Yes A*02:01
Yes Yes A*11:01
Yes Yes A*01:01
Yes Yes A*02:01
Yes Yes A*01:01
Yes Yes B*44:02
NR Yes B*44:03

(61) (13) (13) (98) (28) (69)(09) (62) (64) (e9)219 312 236 9.0 457 35 19 4 N 14976 18746 88555 312 7566 1161 4960 4504 5701 62 ARDPHSGHFV TLGWLLQTPK KILDAVVAQE GSFGDIYLAI EINKNPKYK FLGGNEVGKT SYLDSGIHS KELEGILLP AQQITQTEV 41192 7314 ± 35 141 S3853 48 ဖ 32 41 9 FLEGNEVGKTY TLDWLLQTPK KILDAVVAQK ACDPHSGHFV GLFGDIYLAI KINKNPKYK SYLDSGIHF KELEGILLL AQQITKTEV ILDKVLVHL A*24:02 B*44:03 A*03:01 A*02:01 A*02:01 A*02:01 A*03:01 A*01:01 B*52:01 A*03:01 Yes NR Yes Yes ΝR NR NSCCC MEL MEL MEL MEL MEL MEL MEL MEL MEL CSNK1A1 CTNNB1 GPNMB MYH2 SNRP CDK4 CLPPNFYC6SO4 ო

Table 5 shows epitopes from minor histocompatibility antigens (MM: multiple myeloma; HM:

hematological malignancy; B-ALL: B cell acute lymphocytic leukemia).

Table 5

				T cell							
Group	Gene	Disease	Clinical Disease Response	response	Allele	Mutated	70			WT/MUT	Reference
				MUT>>>WT		Observed	Predicted IC50 (nM)	Observed	Predicted IC50 (nM)	IC50	
F	ECGF-1	MM	Yes	Yes	B*07:02	RPHAIRRPLAL	, E	ЯР	, a	2.0	(99)
-	KIAA022 3 (HA-1)	Σ I	Yes	R	A*02:01	VLHDDLLEA	17	VLRDDLLEA	140	80	(99)
-	BCL2A1	MΗ	S R	Yes	A*24:02	DYLQYVLQI	22	DYLQCVLQI	34	2	(29)
-	BCL2A1	ΣH	N R	Yes	A*24:02	KEFEDDIINW	36	KEFEDGIINW	27	8.0	(29)
-	HB-1	B-ALL	N	Yes	B*44:03	EEKRGSLHVW	81	EEKRGSLYVW	29	-	(89)

Among all tiled 9-mer and 10-mer possibilities, NetMHCpan identified all 33 functionally validated mutated epitopes as the best binding peptide among the possible choices for the given mutation. The median predicted binding affinity (IC50) to the known reported HLA restricting elements of each of the 33 mutated epitopes was 32 nM (range, 3-11, 192 nM). By setting the predicted IC50 cut-offs to 150 and 500 nM, 82 and 91% of the functionally validated peptides, respectively, were captured (see e.g., Tables 4 and 5 and FIG. 12A).

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On the basis of its high degree of sensitivity and specificity, NetMHCpan was then applied to the 31 of 91 CLL cases for which HLA typing information was available. By convention, peptides with IC50 < 150 nM were considered as strong to intermediate binders, IC50 150-500 nM as weak binders, and IC50 > 500 nM as non-binders, respectively (see e.g., Cai et al. 2012). For all 91 CLL cases, a median of 10 strong binding peptides (range, 2-40) and 12 intermediate to weak binding peptides (range, 2-41) was found. In total, a median of 22 (range, 6-81) peptides per case was predicted with IC50 < 500 nM (see e.g., FIG. 12B and Table 6). In particular, Table 6 shows that the numbers and affinity distributions of peptides predicted from 31 CLL cases with available HLA typing. Patients expressing the 8 most common HLA - A, -B alleles in the Caucasian population are marked in grey.

Table 6.

Pts		HLA -A	alleles			HLA -E	i alleles		Uncommon alleles		redicted reptides
(1)										•	`
	*01:01	*02:01	*24:02	*03:01	*07:02	*08:01	*15:01	*38:01		<150 (n%)	159- 5 99 (nM)
P46										10	32
P9*										47	20
932									8*44:02	13	8
940									A*68:01	14	19
SAS:									A*32:01; B*44:02	€	15
889									A*2%:02;B*44:03	17	26
25.3	ì								8*35:01; *57:01	8	3.2
ଷ୍ଟ	1								A*31:01; B*14:02	8	ä
944									8*51.01; *62:01	Š	.\$
P28									A*26:01; B*48:01	10	6
1234									8*18 :91; * 39:06	<u> </u>	7
354									8 * 51:01	11	13
P37									A*03:02; B*44:03	2	্ধ
P43									A*11:01; B*35:03	28	29
942		***************************************							A*26:01; B *35:02	10	18
P66			*****************						A*23:01; B*49:01	4	쓢
P82						000000000000000000000000000000000000000			A*26:01; B*36:06	13	29
983									B*08:01; *04:01	£	2
963		***************************************							A*26:01; *31:01	24	29
\$47									A*11:01; B*51:01	10	9
948				***************************************	*************************	-			A*11:01; B*51:01	13	37
P18									A*11:01 ; B*44:02; *61:01	13	12
1736									A*88:01; B*14:02	3	ő
P39									A *32:01; B*36:01; *44:01	\$ 0	43
884									A*26:01;3 *35:01	13	\$ \$
873									A*31.01; *68.01; 8*49.01	18	13
P\$(0)									A*29/32; B*45/31; *55/31	33	7
P45									A*66:02; B*44:02; *15:03	<u> </u>	ő
21									A*11:01; *23:01; B*35:01; *51:01	*	*7
P40									A*11:01; *32:01; B*40:01; *44:03	14	% 9
P41									A*29/03; *32:01; B*44:03	10	æ

Example 17: More than half of predicted HLA-binding neopeptides showed direct binding to HLA proteins in vitro

As shown in Table 7, IC50 nM scores generated by HLA-peptide binding predictions were validated using a competitive MHC I allele binding assay and focused on class I-A and –B alleles. To this end, 112 mutated peptides (9 or 10-mer mutated peptides) with predicted IC50 scores of less than 500 nM that were identified from 4 CLL cases (Pt 1-4) were synthesized. The experimental results correlated with the binding predictions. Experimental binding (defined as IC 50 < 500 NM) was confirmed in 76.5% and 36% of peptides predicted with IC50 of < 150 nM or 150-500 nM, respectively (see e.g., FIG. 12C). In total, ~54.5% (61/112) of predicted peptides were experimentally validated as binders to personal HLA alleles. Overall, the predictions for 9-mer peptides were more sensitive than for 10-mer peptides, as 60% vs 44.5% of predicted peptides (IC50 < 500 nM) could be experimentally validated, respectively, as shown in (FIG. 13).

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Table 7. Predicted and experimental HLA-binding results of candidate neoepitopes generated from 4 CLL cases.

1. THOC6 ELWCRQPPYR 10 A*33:01 Fredicted Experimental 1. THOC6 ELWCRQPPYR 10 A*33:01 10 18 1. CDC25A QSYCEPSSYR 10 A*68:12 25 11 1. CDC25A QSYCEPSSYR 10 A*68:12 25 11 1. WHSC11 EVQASKHTK 9 A*68:12 25 11 1. WHSC11 EVQASKHTK 9 A*68:12 25 11 1. WHSC11 EVQASKHTK 9 A*68:12 33 58 1. CAPYBA1 WVCYQYSGYR 10 A*68:12 70 14 1. CAPYBA1 WVCYQYSGYR 10 A*68:12 71 42 1. TH/NSL2 ATIESVQGAK 10 A*68:12 75 91 1. THOS6 ELWCRQPRY 9 A*68:12 75 91 1. THOS6 LUSCKDGKR 9 A*68:12 153 134 1. THOC6 LWCRQDAK 10 A*68:12 253	표	Gene	Sednence	Length	HLA allele	Candidate r	Candidate neoepitopes
THOC6 ELWCRQPPYR 10 A*33:01 10 THOC6 ELWCRQPPYR 10 A*68:12 59 CDC25A GSYCEPSSYR 10 A*68:12 23 ALMS1 TVPSSSFSHR 10 A*68:12 23 WHSC1L1 EVQASKHTK 9 A*68:12 25 WHSC1L1 EVQASKHTK 9 A*68:12 25 THNSL2 SYCEPSSYR 10 A*68:12 33 CDC25A SYCEPSSYR 9 A*68:12 71 ALMS1 TPTVPSSSF 9 A*68:12 75 RALGAPB WIMVLVLPK 10 A*68:12 95 THOC6 ELWCRQPRY 9 B*35:01 117 C60rf89 MPIEPGDIGC 10 A*68:12 157 WHSC1L1 LLNEVQASK 9 A*68:12 157 WHSC1L1 LLNEVQASK 9 A*68:12 224 XPO1 KTVVNIKLFK 9 A*68:12 253						IC50 Predicted	(nM) Experimental
THOC6 ELWCROPPYR 10 A'68:12 59 CDC25A QSYCEPSSYR 10 A'68:12 23 ALMS1 TVPSSSFSHR 10 A'68:12 25 WHSC1L1 EVQASKHTK 9 A'68:12 33 CRYBA1 WVCYQYSGYR 10 A'88:12 35 CRZ5A SYCEPSSYR 9 A'88:12 71 THNSL2 ATIESVQGAK 10 A'88:12 71 ALMS1 TPTVPSSSF 9 B'35:01 75 THOC6 ELWCRQPPY 9 A'88:12 75 THOC6 ELWCRQPPY 9 A'88:12 163 CG0r/89 WINNVLVLPK 10 A'88:12 153 STRAP LISACKDGKR 9 A'88:12 224 AFG18 DWINNVLVLPK 10 A'88:12 258 AFG4APB DWINNVLVLFK 9 A'88:12 258 AFO1 KTVVNIKLK 9 A'88:12 258	_	920Н1	ELWCRQPPYR	10	A*33:01	10	18
CDC25A QSYCEPSSYR 10 A*68:12 23 ALMS1 TVPSSSFSHR 10 A*68:12 25 WHSC1L1 EVQASKHTK 9 A*68:12 33 CRYBA1 WVCYQYSGYR 10 A*68:12 33 CDC25A SYCEPSSYR 9 A*33:01 70 THNSL2 ATIESVQGAK 10 A*68:12 71 ALMS1 TPTVPSSSF 9 B*35:01 75 PALGAPB WIMVLVLPK 9 B*35:01 112 RALGAPB DWIMVLVLPK 10 A*68:12 95 STRAP LISACKDGKR 9 B*35:01 170 WHSC1L1 LLNEVQASK 9 A*68:12 253 STRAP ISACKDGKR 9 A*68:12 253 AMGAZ KTVVNKLFK 9 A*68:12 253 THOC6 LWCRQAPB 9 A*68:12 253 POLR2A VQKIFHINPR 10 A*68:12 253	_	ЭЭОН1	ELWCRQPPYR	10	A*68:12	59	5.1
ALMS1 TVPSSSFSHR 10 A'68:12 25 WHSC1L1 EVQASKHTK 9 A'68:12 33 CRYBA1 WVCYQYSGYR 10 A'68:12 33 CDC25A SYCEPSSYR 10 A'73:01 70 THNSL2 ATIEVVBSSF 9 A'73:01 71 ALMS1 TPTVPSSF 9 A'68:12 71 ALGAPB WIMVLVLPK 9 A'68:12 95 RALGAPB DWIMVLVLPK 10 A'68:12 153 C60rt89 MPIEFGDIGC 10 A'68:12 163 STRAP LISACKDGKR 9 A'68:12 224 RALGAPB DWIMVLVLPK 10 A'68:12 224 FALGAPB DWIMVLVLPK 9 A'68:12 253 FALGAPB DWIMVLVLPK 9 A'68:12 253 FALGAPB DWIMVLLFK 9 A'68:12 253 FALGAP INSAENGDAK 9 A'68:12 253 <tr< td=""><td>-</td><td>CDC25A</td><td>QSYCEPSSYR</td><td>10</td><td>A*68:12</td><td>23</td><td>1.5</td></tr<>	-	CDC25A	QSYCEPSSYR	10	A*68:12	23	1.5
WHSC1L1 EVQASKHTK 9 A*68:12 33 CRYBA1 WVCYQYSGYR 10 A*33:01 44 CDC25A SYCEPSSYR 9 A*33:01 70 THNSL2 ATIESVQGAK 10 A*68:12 71 ALMS1 TPTVPSSSF 9 B*35:01 75 RALGAPB WIMVLVLPK 9 B*35:01 112 RALGAPB WIMVLVLPK 10 A*68:12 95 CGori89 MPIEPGDIGC 10 B*35:01 117 CGORI89 MPIEPGDIGC 10 A*68:12 163 STRAP LISACKDGKR 9 A*68:12 224 WHSC1L1 LLNEVQASK 9 A*68:12 254 RALGAPB DWIMVLVLFK 9 A*68:12 254 STRAP ISACKDGKR 9 A*68:12 254 HMGNZ KTVVNKLFK 9 A*68:12 258 HMGNZ NSAENGGDAK 9 A*68:12 258	_	ALMS1	TVPSSSFSHR	10	A*68:12	25	11
CRYBA1 WVCYQYSGYR 10 A*33:01 44 CDC25A SYCEPSSYR 9 A*33:01 70 THNSL2 ATIESVQGAK 10 A*68:12 71 ALMS1 TPTVPSSSF 9 B*35:01 75 ALMS1 TPTVPSSSF 9 B*35:01 75 RALGAPB WIMVLVLPK 9 B*35:01 112 RALGAPB DWIMVLVLPK 10 A*68:12 152 C6or/89 MPIEPGDIGC 10 B*35:01 170 WHSC1L1 LLNEVQASK 9 A*68:12 153 WHSC1L2 LLNEVQASK 9 A*68:12 224 RALGAPB DWIMVLVLFK 9 A*68:12 254 RALGAPB DWIMVLVLFK 9 A*68:12 254 RALGAPB DWIMVLVLFK 9 A*68:12 254 RALGAPB NSAENGGAK 9 A*68:12 254 HMGNZ KTVVNKLFK 9 A*68:12 258 <tr< td=""><td>-</td><td>WHSC1L1</td><td>EVQASKHTK</td><td>6</td><td>A*68:12</td><td>33</td><td>28</td></tr<>	-	WHSC1L1	EVQASKHTK	6	A*68:12	33	28
CDC25A SYCEPSSYR 9 A*33:01 70 THNSL2 ATIESVOGAK 10 A*68:12 71 ALMS1 TPTVPSSSF 9 B*35:01 75 RALGAPB WIMNULNLPK 9 A*68:12 95 RALGAPB WIMNULNLPK 10 A*33:01 112 CGO189 MPIEPGDIGC 10 A*68:12 152 CGO189 MPIEPGDIGC 10 A*68:12 153 CRYBA1 YQYSGYRGY 9 A*68:12 153 WHSC1L1 LLNEVQASK 9 A*68:12 157 WHSC1L1 LLNEVQASK 9 A*68:12 224 XPO1 KTVVNKLFK 9 A*68:12 253 YRO1 KTVNNKLFK 9 A*68:12 253 HMGN2 NSAENGDPK 9 A*68:12 258 POLR2A VQKIFHINPR 10 A*33:01 309 ALMS1 SSSFSHREK 9 A*68:12 314	-	CRYBA1	WVCYQYSGYR	10	A*33:01	44	972
THNSL2 ATIESVQGAK 10 A*68:12 71 ALMS1 TPTVPSSSF 9 B*35:01 75 RALGAPB WIMVLVLPK 9 A*68:12 75 THOC6 ELWCRQPPY 9 B*35:01 112 C60rt89 MPIEPGDIGC 10 A*68:12 152 STRAP LISACKDGKR 10 A*68:12 153 CRYBA1 YQYSGYRGY 9 A*68:12 153 WHSC1L1 LLNEVQASK 9 A*68:12 222 STRAP ISACKDGKR 9 A*68:12 224 XPO1 KTVVNKLFK 9 A*68:12 253 YMO1 KTVVNKLFK 9 A*68:12 253 HMGN2 NSAENGDAK 9 A*68:12 253 POLR24 VQKIFHINPR 10 A*33:01 308 POLR24 QSYCEPSSYR 9 A*33:01 309 ALMS1 SSSFSHREK 9 A*68:12 314	1	CDC25A	SYCEPSSYR	6	A*33:01	70	14
ALMS1 TPTVPSSSF 9 B°35.01 75 RALGAPB WIMVLVLPK 9 A°68:12 95 THOC6 ELWCRQPPY 9 B°35.01 112 RALGAPB DWIMVLVLPK 10 B°35.01 117 CG0489 MPIEPGDIGC 10 B°35.01 132 STRAP LISACKDGKR 9 A°68:12 153 WHSC1L1 LLNEVQASK 9 A°68:12 222 WHSC1L1 LLNEVQASK 9 A°68:12 224 STRAP ISACKDGKR 9 A°68:12 253 HMGN2 KTVVNKLFK 9 A°68:12 253 HMGN2 UWCRQPPYR 9 A°68:12 258 POLR2A VQKIFHINPR 10 A°33:01 308 POLR2A VQKIFHINPR 10 A°33:01 309 ALMS1 SSSFSHREK 9 A°68:12 314	_	ZTSNH1	ATIESVQGAK	10	A*68:12	71	42
RALGAPB WIMVLVLPK 9 A*68:12 95 THOC6 ELWCRQPPY 9 B*35:01 112 RALGAPB DWIMVLVLPK 10 A*33:01 117 CGOrf89 MPIEPGDIGC 10 B*35:01 132 STRAP LISACKDGKR 10 A*68:12 163 CRYBA1 YQYSGYRGY 9 A*68:12 197 WHSC1L1 LLNEVQASK 9 A*68:12 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 STRAP ISACKDGKR 9 A*68:12 253 HMGN2 NSAENGDAK 9 A*68:12 253 THOC6 LWCRQPPYR 9 A*68:12 253 POLR2A VQKIFHINPR 10 A*33:01 308 CDC25A QSYCEPSSYR 10 A*33:01 309 ALMS1 SSSFSHREK 9 A*68:12 314	-	ALMS1	TPTVPSSSF	6	B*35:01	75	91
THOC6 ELWCRQPPY 9 B*35:01 112 RALGAPB DWIMVLVLPK 10 A*33:01 117 117 C60r/89 MPIEPGDIGC 10 B*35:01 132 132 STRAP LISACKDGKR 9 A*68:12 163 170 WHSC1L1 LLNEVQASK 9 A*68:12 222 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 224 RALGAPB ISACKDGKR 9 A*68:12 224 253 HMGNZ NSAENGDAK 9 A*68:12 258 258 HMGNZ NSAENGDAK 9 A*68:12 258 258 POLRZA VQKIFHINPR 10 A*33:01 308 258 POLRZA QSYCEPSSYR 10 A*33:01 309 314	-	RALGAPB	WIMVLVLPK	6	A*68:12	95	218
RALGAPB DWIMVLVLPK 10 A*33:01 117 C6orf89 MPIEPGDIGC 10 B*35:01 132 STRAP LISACKDGKR 10 A*68:12 163 WHSC1L1 LLNEVQASK 9 B*35:01 170 WHSC1L1 LLNEVQASK 9 A*68:12 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 STRAP ISACKDGKR 9 A*68:12 224 XPO1 KTVVNKLFK 9 A*68:12 253 HMGN2 NSAENGDAK 9 A*68:12 253 POLR24 VQKIFHINPR 10 A*33:01 309 POLR24 VQKIFHINPR 10 A*33:01 309 ALMS1 SSSFSHREK 9 A*68:12 314	_	9)ОН1	ELWCRQPPY	6	B*35:01	112	13776
C601789 MPIEPGDIGC 10 B*35.01 132 STRAP LISACKDGKR 10 A*68:12 163 WHSC1L1 LLNEVQASK 9 B*35.01 170 WHSC1L1 LLNEVQASK 9 A*68:12 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 STRAP ISACKDGKR 9 A*68:12 224 YPO1 KTVVNKLFK 9 A*68:12 253 HMGN2 NSAENGDAK 9 A*68:12 258 POLR2A VQKIFHINPR 10 A*33:01 308 POLR2A VQKIFHINPR 10 A*33:01 308 CDC25A QSYCEPSSYR 9 A*68:12 314	_	RALGAPB	DWIMVLVLPK	10	A*33:01	117	37826
STRAP LISACKDGKR 10 A*68:12 163 163 WHSC1L1 LLNEVQASK 9 B*35:01 170 170 RALGAPB DWIMVLVLPK 10 A*68:12 222 197 STRAP ISACKDGKR 9 A*68:12 224 10 XPO1 KTVVNKLFK 9 A*68:12 253 10 HMGN2 NSAENGDAK 9 A*68:12 258 10 POLR2A VQKIFHINPR 10 A*33:01 308 10 CDC25A QSYCEPSSYR 10 A*33:01 309 10 ALMS1 SSSFSHREK 9 A*68:12 314 11	-	C6orf89	MPIEPGDIGC	10	B*35:01	132	131
CRYBA1 YQYSGYRGY 9 B*35:01 170 770 WHSC1L1 LLNEVQASK 9 A*68:12 197 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 224 XPO1 KTVVNKLFK 9 A*68:12 253 1 HMGN2 NSAENGDAK 9 A*68:12 258 1 POLR2A UQKIFHINPR 10 A*33:01 308 1 CDC25A QSYCEPSSYR 10 A*33:01 309 1 ALMS1 SSSFSHREK 9 A*68:12 314 1	1	STRAP	LISACKDGKR	10	A*68:12	163	15845
WHSC1L1 LLNEVQASK 9 A*68:12 197 197 RALGAPB DWIMVLVLPK 10 A*68:12 222 224<	-	CRYBA1	YQYSGYRGY	6	B*35:01	170	9851
RALGAPB DWIMVLVLPK 10 A*68:12 222 7 STRAP ISACKDGKR 9 A*68:12 224 7 XPO1 KTVVNKLFK 9 A*68:12 253 7 HMGN2 NSAENGDAK 9 A*68:12 258 7 POLR2A LWCRQPPYR 9 A*33:01 308 7 POLR2A VQKIFHINPR 10 A*33:01 308 7 ALMS1 SSSFSHREK 9 A*68:12 314 314	1	NHSC1L1	LLNEVQASK	6	A*68:12	197	7440
XPO1 KTVVNKLFK 9 A*68:12 224 7 HMGN2 NSAENGDAK 9 A*68:12 253 7 THOC6 LWCRQPPYR 9 A*33:01 297 7 POLR2A VQKIFHINPR 10 A*33:01 308 7 CDC25A QSYCEPSSYR 10 A*33:01 309 7 ALMS1 SSSFSHREK 9 A*68:12 314 11	-	RALGAPB	DWIMVLVLPK	10	A*68:12	222	2956
XPO1 KTVVNKLFK 9 A*68:12 253 P. HMGN2 NSAENGDAK 9 A*68:12 258 58 THOC6 LWCRQPPYR 9 A*33:01 297 308 POLR2A VQKIFHINPR 10 A*33:01 308 5 CDC25A QSYCEPSSYR 10 A*33:01 309 5 ALMS1 SSSFSHREK 9 A*68:12 314 5	1	STRAP	ISACKDGKR	6	A*68:12	224	6671
HMGNZ NSAENGDAK 9 A*68:12 258 7 THOC6 LWCRQPPYR 9 A*33:01 297 5 POLR2A VQKIFHINPR 10 A*33:01 308 5 CDC25A QSYCEPSSYR 10 A*33:01 309 5 ALMS1 SSSFSHREK 9 A*68:12 314 5	1	XPO1	KTVVNKLFK	6	A*68:12	253	25393
LWCRQPPYR 9 A*33:01 297 VQKIFHINPR 10 A*33:01 308 QSYCEPSSYR 10 A*33:01 309 SSSFSHREK 9 A*68:12 314	1	HMGN2	NSAENGDAK	6	A*68:12	258	141
VQKIFHINPR 10 A*33:01 308 70 QSYCEPSSYR 10 A*33:01 309 809 SSSFSHREK 9 A*68:12 314 9	1	<i>9</i> 00H1	LWCRQPPYR	6	A*33:01	297	915
55A QSYCEPSSYR 10 A*33:01 309 S1 SSSFSHREK 9 A*68:12 314	1	POLR2A	NQKIFHINPR	10	A*33:01	308	17699
S1 SSSFSHREK 9 A*68:12 314	1	CDC25A	QSYCEPSSYR	10	A*33:01	309	53
	1	ALMS1	SSSFSHREK	6	A*68:12	314	1496

812	237	653	541	20000	9195	1.1	6.4	8.5	6.8	11	۲١	58	40	62	94	107	122	123	130	137	175	274	378	299	743	803	855
314	335	338	393	478	480	10.63	4.21	8.13	414.37	41.91	413.95	443.97	2.67	63.7	22.26	28.18	382.07	98.15	245.43	179.31	454.23	302.94	37.77	13.74	145.51	340.37	243.46
A*68:12	A*33:01	A*68:12	B*35:01	B*35:01	B*35:01	A*02:01	A*02:01	A*01:01	B*08:01	A*02:01	A*02:01	B*08:01	A*02:01	A*02:01	A*02:01	A*02:01	A*02:01	A*02:01	B*08:01	B*07:02	A*02:01	A*02:01	A*02:01	A*02:01	B*08:01	B*08:01	B*07:02
6	10	6	6	6	10	10	6	6	6	6	6	6	10	6	6	10	6	10	6	6	6	10	10	6	6	10	10
SYCEPSSYR	TVPSSSFSHR	TIESVQGAK	MIWNVQKIF	QSYCEPSSY	SSIRSFVLQY	FLQEETLTQM	VVMSWAPPV	CSDSKLIGY	EMLIKPKEL	ILLMTVTSI	SLMEHWALG	LLRVHTEHV	SLMEHWALGA	KMTFLFPNL	GLVDEQQEV	ALPDPILQSI	GVWALPDPI	AVVMSWAPPV	TSIDRFLAV	GPSWGLSLM	LLRVHTEHV	WVNCSSMTFL	VMSWAPPVGL	ILYKDDMGV	NIQARAVVM	HVRCKSGNKF	LPDPILQSIL
CDC25A	ALMS1	THNSL2	POLR2A	CDC25A	DSCAML1	NIN	<i>FNDC3B</i>	SLC46A1	SYT15	F2R	ACSM2A	C16orf57	ACSM2A	TBC1D9B	SF3B1	LRBC41	LRRC41	<i>FNDC3B</i>	F2R	KIAA0467	C160rf57	C22orf28	<i>FNDC3B</i>	GDF2	FNDC3B	C16orf57	LBRC41
-	1	1	-	1	1	2	2	2	2	5	2	5	5	2	2	5	2	2	5	2	2	5	5	5	5	2	2

929	896	1056	1252	1423	1442	1651	1687	1775	1789	2322	3416	5074	6511	8085	85	40	273	739	1897	10452	41	30	21	59	51	62	125
301.24	314.16	471.62	23.04	107.39	162.61	424.59	280.32	140.39	285.7	327.97	132.77	437.05	128.72	228.47	205.26	443.32	116.69	343.52	164.9	227	26	122.42	17	25	19	12	117
A*02:01	A*02:01	B*07:02	A*02:01	B*07:02	A*02:01	A*02:01	B*07:02	A*02:01	B*08:01	B*08:01	B*07:02	A*02:01	A*02:01	B*07:02	B*15:01	A*03:01	A*03:01	A*01:01	A*03:01	A*01:01	A*03:01	A*03:01	B*08:01	A*03:01	B*08:01	A*03:01	B*15:01
10	6	6	10	10	6	10	<u></u>	10	6	10	10	10	6	10	6	10	6	10	10	10	6	6	10	6	6	6	6
SILLMTVTSI	LMEHWALGA	LPDPILQSI	VLLRVHTEHV	FPNLKDRDFL	MLIKPKELV	ILCSLMEHWA	FPNLKDRDF	SILYKDDMGV	NTFRHRVVV	EVRTISALAI	VPTKLSPISI	LLLQDSECKA	SQPGPSWGL	KPAVSSDSDI	ITHTGEKPY	ITHTGEKPYK	KFSNSNIYK	LTRGTFANIK	LTDFGLSKIM	LTDFGLSKI	KLTDFGLSK	SLSLGAHQK	FLKHKQSCAV	GVMTSCFLK	FLKHKQSCA	GLLCAFTLK	HLCGLLCAF
F2R	ACSM2A	LRBC41	C16orf57	TBC1D9B	SYT15	ACSM2A	TBC1D9B	GDF2	<i>TP53</i>	SF3B1	GDF2	ЕГКЗ	KIAA0467	RNF150	ZNF182	ZNF182	ZNF253	IREB2	TLK2	TLK2	TLK2	<i>MYD88</i>	PATE2	PATE2	PATE2	JTB	JTB
5	2	2	2	5	2	2	5	2	2	5	5	5	5	5	3	3	၉	က	က	၉	က	၉	၉	၉		3	ဇ

FE2 VMTSCFLKHK 10 7E2 MTSCFLKHK 9 3C5 KISSLEGRSK 10 3C5 KISSLEGRSK 10 3C5 LSIFKISSL 9 3C5 LSIFKISSL 9 7L2 EVAGFVFDKK 10 7L2 EVAGFVFDKK 10 DZ FSIVGGYGR 9 L1 YMKKAEAPL 9 L25 ISYLGRDRLR 10 DZ APITTTTTV 9 L1 YMKKAEAPLL 10 DZ RLRQEVYLSL 10 L1 YMKKAEAPLL 10 L25 RLRGEVYLNTAR 9 L33 FTAGGEPCLY 10 M1 DPKLGTAQPL 9 L2 CLT	OR13C5	5 LSIFKISSL	6	B*08:01	151	158
PATE2 MTSCFLKHK 9 OR13C5 KISSLEGRSK 10 OR13C5 LSIFKISSL 9 MAPK14 RPTFYRQGL 9 SCYL2 EVAGFVFDKK 10 SCYL2 EVAGFVFDKK 10 SCYL2 EVAGFVFDKK 9 COL5A3 FTAGGEPCLY 10 MDZ FSINGGYGR 9 CUL1 YMKKAEAPL 9 KDM5D HSIPLRQSVK 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 NUP98 APGENTTPA 9 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEPYLSL 9 COL53 FTAGGEPCLY 10 COL53 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 COL53 FTAGGEPCY 9 <th>PATE2</th> <th></th> <th>10</th> <th>A*03:01</th> <th>140</th> <th>174</th>	PATE2		10	A*03:01	140	174
OR13C5 KISSLEGRSK 10 OR13C5 LSIFKISSL 9 MAPK14 RPTFYRQGL 9 SCYL2 EVAGFVFDK 10 SCYL2 EVAGFVFDK 10 COL5A3 FTAGGEPCLY 10 MDZ FSINGGYGR 9 CUL1 YMKKAEAPL 9 KDM5D HSIPLRQSVK 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MDDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL543 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 CUL1 EAPLLEEQR 9	PATE2		6	A*03:01	147	218
OB13C5 LSIFKISSL 9 MAPK14 RPTFYRQGL 9 SCYL2 EVAGFVFDK 10 SCYL2 EVAGFVFDKK 10 COL5A3 FTAGGEPCLY 10 MDZ FSIVGGYGR 9 CUL1 YMKKAEAPL 9 MUC2 APITTITTV 9 KDM5D HSIPLRQSVK 10 MUD2 RPHGDLPITR 10 MDZ RPHGDLPITR 10 TBC1D25 RLRQEVLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 9 PPP2R2C QTHEPEFDY 9	OR13C		10	A*03:01	185	257
MAPK14 RPTFYRQGL 9 SCYL2 EVAGFVFDK 10 COL5A3 FTAGGEPCLY 10 MPDZ FSIVGGYGR 9 MUC2 APITTTTTV 9 MUC2 APITTTTTV 9 MUC2 APITTTTTV 9 MUC2 APITTTTTV 9 NUP98 APGFNTTPA 10 NUP98 APGFNTTPA 10 NUP98 APGFNTTPA 10 NUP98 APGFNTTPA 9 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRGEPYLSL 10 CUL1 YMKKAEAPLL 10 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 9 PPP2R2C QTHEPEFDY 9 CUL1 EAPLLEEQR	OR13C		6	B*15:01	152	368
SCYL2 EVAGEVEDKK 9 SCYL2 EVAGEVEDKK 10 COL5A3 FTAGGEPCLY 10 MPDZ FSIVGGYGR 9 MUC2 APITTTTTV 9 KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRILR 10 NUP98 APGFNTTPA 9 XNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVLSL 10 TBC1D25 RLRGEVALSL 10 TBC1D25 RLRGEVALSL 10 TBC1D25 RLRGEVALSL 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 COL5A3 FTAGGEPCLY 9 COL5A3 FTAGGEPCLY 9 PPP2R2C QTHEPEFDY 9 RMUC2 APPLTTTTTYT <th>MAPK1</th> <th>L</th> <th>6</th> <th>B*07:02</th> <th>2.9</th> <th>9/</th>	MAPK1	L	6	B*07:02	2.9	9/
SCYL2 EVAGFVFDKK 10 COL5A3 FTAGGEPCLY 10 MPDZ FSIVGGYGR 9 CUL1 YMKKAEAPL 9 KDM5D HSIPLRQSVK 10 KDM5D HSIPLRQSVK 10 NUP98 APGFNTTPA 9 NUP98 APGFNTTPA 10 MPDZ RPHGDLPIYV 10 MPDZ RPHGDLPIYV 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 9 MUC2 APITTITITYT 10 CUL1 EAPLLEEQR 9	SCYL2		თ	A*68:01	7.3	14
COL5A3 FTAGGEPCLY 10 MPDZ FSIVGGYGR 9 CUL1 YMKKAEAPL 9 MUC2 APITTTTTV 9 KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVLNTSL 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTYT 10 CUL1 EAPLLEEQR 9	SCX12	L	10	A*68:01	7.4	8.8
MPDZ FSIVGGYGR 9 CUL1 YMKKAEAPL 9 MUC2 APITTTTTV 9 KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEPCLY 10 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	COL5A	L	10	A*01:01	14	153
CUL1 YMKKAEAPL 9 MUC2 APITTTTTV 9 KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	ZOAW		6	A*68:01	50	2.6
MUC2 APITTTTTV 9 KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRGEPCLY 10 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10	CUL1		თ	B*08:01	36	34841
KDM5D HSIPLRQSVK 10 TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 TANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	MUC2		6	B*07:02	53	13
TBC1D25 ISYLGRDRLR 10 NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	KDM5D		10	A*68:01	22	45
NUP98 APGFNTTPA 9 ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	TBC1D2		10	A*68:01	106	929
ZNF330 KAFFCDDHTR 10 MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 TBC1D25 RLRQEVYLSL 10 COL543 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	NUP98		6	B*07:02	107	13
MPDZ RPHGDLPIYV 10 TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPPZR2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	ZNF330		10	A*68:01	137	102
TBC1D25 RLRQEVYLSL 10 CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTYT 10 CUL1 EAPLLEEQR 9	ZOAW	_	10	B*07:02	155	1321
CUL1 YMKKAEAPLL 10 TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	TBC1D2		10	B*08:01	165	1084
TBC1D25 RLRQEVYLSL 10 LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	CNT 1		10	B*08:01	168	138
LANCL1 CLTKRSIAF 9 COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	TBC1D2		10	B*07:02	183	114
COL5A3 FTAGGEPCLY 10 SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTTVT 10 CUL1 EAPLLEEQR 9	TANCE		6	B*08:01	202	47
SF3B1 EYVLNTTAR 9 CNN1 DPKLGTAQPL 10 PPP2R2C QTHEPEFDY 9 MUC2 APITTTTVT 10 CUL1 EAPLLEEQR 9	COL5A		10	A*68:01	230	11
CNN1DPKLGTAQPL10PPP2R2CQTHEPEFDY9MUC2APITTTTTVT10CUL1EAPLLEEQR9	SF3B1		6	A*68:01	301	651
PPP2R2CQTHEPEFDY9MUC2APITTITIVT10CUL1EAPLLEEQR9	CNN1		10	B*07:02	698	3974
MUC2 APITTTTVT 10 CUL1 EAPLLEEQR 9	PPP2R2		6	A*01:01	435	26184
CUL1 EAPLLEEQR 9	MUC2		10	B*07:02	436	3731
	CNT 1	EAPLLEEQR	6	A*68:01	454	36
4 LANCL1 CLTKRSIAFL 10 B*08:01	TANCE		10	B*08:01	467	640

4	86ANN	APGFNTTPAT	10	B*07:02	475	5744
4	Z)NW	TTAPITTT	6	A*68:01	625	118
4	CNT 1	YMKKAEAPL	6	B*07:02	480	1361
4	77X07	IPGFKFDNL	6	B*07:02	487	608

** An experimental binding assay for A*68:12 was not available. Because A*68:12 and A*68:01 have identical primary structures in the B and F main peptide binding pockets and have been predicted to have similar binding specificity (Sidney and Sette, 2007), experimental binding for peptides predicted to bind A*68:12 were assayed against A*68:01.

Example 18: Neoantigens are expressed in CLL tumors

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CTL responses against an epitope would only be useful if the gene encoding the epitope is expressed in the target cells. Of the 31 patient samples sequenced and typed for HLA, 26 were subjected to genome-wide expression profiling (see e.g., Brown et al. 2012). The expression level of 347 genes with mutations in CLL samples was classified as having low/absent (lowest quartile), medium (middle two quartiles), or high (highest quartile) expression. As shown in FIG. 12D, 80% of the 347 mutated genes (or 79% of the 180 mutations with predicted HLA-binding) were expressed at medium or high expression levels. A similar high frequency of expression was observed among the subset of 221 mutated genes (88.6%) with predicted class I binding epitopes.

RNA levels may be determined based on the number of reads per gene product, and ranked by quartiles. "H" - Top quartile; "M" - Middle two quartiles; "L" - Lowest quartile (excluding genes with no reads; "-" - no reads detectable. As the predicted affinity decreases, higher stringency may be applied to expression levels. NeoORFs with predicted binders were utilized even if there was no detectable mRNA molecules by RNA-Seq. There is no data currently available to assess what, if any, the minimum expression level required in a tumor cell would be for a neoORF to be useful as a target for activated T-cells. Even the level of expression of "pioneer" translation of messages destined for nonsense mediated decay may be sufficient for target generation ((Chang YF, Imam JS, Wilkinson MF: The nonsense-mediated decay RNA surveillance pathway. Annu Rev Biochem 76:51-74, 2007). Therefore, because of the high value of neoORFs as targets due to their novelty and exquisite tumor specificity, neoORFs may be utilized as immunogens even if expression at the RNA level is low or undetectable.

Example 19: T cells targeting candidate neoepitopes were detected in CLL Patient 1 following HSCT

The post-allogeneic hematopoietic stem cell transplantation (HSCT) setting in CLL was analyzed to determine whether an immune response against the predicted mutated peptides could develop in patients. Reconstitution of T cells from a healthy donor following HSCT can overcome endogenous immune defects of the host, and also allow priming against leukemia cells

in the host in vivo. Analysis focused on two patients who had both undergone unrelated reduced intensity conditioning allo-HSCT for advanced CLL and had achieved continuous remission for greater than 4 years following HSCT (see e.g., Table 8). Post-transplant T cells were collected 7 years (Patient 1) and 4 years (Patient 2) from the time of transplant.

Table 8 shows the clinical characteristics of CLL Pts 1 and 2. Both patients have achieved ongoing continuous remission following HSCT of greater than 7 (Pt 1) and 4 years (Pt 2). M: male; HSCT: hematopoietic stem cell transplantation; RIC: reduced intensity conditioning; Flu/Bu: Fludarabine/Busulfan; GvHD: graft vs host disease; URD: unrelated donor; Mis: missense; FS: frameshift.

PCT/US2014/033185

Table 8.

				Allogeneic HSCT	e HSCT		N	Number of Mutations	f Muta	tions	Neo€	Neoepitopes (IC50 < 500 nM)
		βĞ										
		Sex	a difference	Stem	Days to							
	HLA		Conditioning	cell	сGvHD	GvHD				Putative		
Ŧ	typing			source	Onset	meds	Total	Mis	FS	drivers	Predicted	Experimental
	A*33:01/											
	*68:12					lmatinib/						
	B*35:01/		RIC	URD		Prednisone						
-	*14:01	51/M	Flu/Bu	PBSC	448		33	25	∞	XPO1	30	4
	A*01:01/											
	*02:01											
	B*07:02/		RIC	URD						TP53,		
2	*08:01	72/M	Fu/Bu	PBSC	208	Imatinib	27	26	1	SF3B1	37	18

For Patient (Pt 1), 25 missense mutations were identified by WES. In total, 30 peptides from 13 mutations were predicted to bind to personal HLA (13 peptides with IC50 < 150; 17 peptides with IC50 150-500 nM). As shown in FIG. 14A, experimental validation of peptide predictions confirmed HLA binding for 14 peptides derived from 9 mutations. All 30 predicted HLA binding peptides were selected for T cell priming studies, and were organized into 5 pools of 6 peptides/pool (see e.g., Table 9). Peptides with similar predicted binding scores were put together within the same pool.

Table 9 provides a summary of peptides from Pt 1 missense mutations that were included in peptide pools for T cell stimulation studies. In Pt 1, all predicted peptides with IC50 < 500 nM binding to HLA -A and -B alleles were used. 5 pools of mutated peptides with 6 peptides/pool listed in decreasing order of predicted binding affinities to MHC class I alleles. The corresponding experimental HLA-peptide binding affinities, wildtype peptides and their predicted IC50 scores are included in the far right columns.

Table 9.

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				MU	T peptide		WT pep	tide
			HLA		Predicted	Experimental		Predicted
Pool	Gene	Length	allele	Sequence	IC50 (nM)	IC50 (nM)	Sequence	IC50 (nM)
	THOC6	10	A*33:01	ELWCRQPPYR	10	18	ELWRRQPPYR	11
	THOC6	10	A*68:12	ELWCRQPPYR	59	5.1	ELWRRQPPYR	61
	CDC25A	10	A*68:12	QSYCEPSSYR	23	1.5	QSYCEPPSYR	37
1	ALMS1	10	A*68:12	TVPSSSFSHR	25	11	TVPSGSFSHR	35
	WHSC1L1	9	A*68:12	EVQASKHTK	33	58	EVQASEHTK	34
	CRYBA1	10	A*33:01	WVCYQYSGYR	44	972	WVCYQYPGYR	50
	CDC25A	9	A*33:01	SYCEPSSYR	70	14	SYCEPPSYR	61
	THNSL2	10	A*68:12	ATIESVQGAK	71	42	AAIESVQGAK	470
	ALMS1	9	B*35:01	TPTVPSSSF	75	91	TPTVPSGSF	89
2	RALGAPB	9	A*68:12	WIMVLVLPK	95	218	WIMALVLPK	46
	THOC6	9	B*35:01	ELWCRQPPY	112	13776	ELWRRQPPY	126
	RALGAPB	10	A*33:01	DWIMVLVLPK	117	37826	DWIMALVLPK	171

	C6orf89	10	B*35:01	MPIEPGDIGC	132	131	MPIEPGDIGY	3
	STRAP	10	A*68:12	LISACKDGKR	163	15845	LISACKDGKP	38499
	CRYBA1	9	B*35:01	YQYSGYRGY	170	9851	YQYPGYRGY	171
3	WHSC1L1	9	A*68:12	LLNEVQASK	197	7440	LLNEVQASE	21454
	RALGAPB	10	A*68:12	DWIMVLVLPK	222	2956	DWIMALVLPK	299
	STRAP	9	A*68:12	ISACKDGKR	224	6671	ISACKDGKP	39393
	XPO1	9	A*68:12	KTVVNKLFK	253	25393	KTVVNKLFE	18346
	HMGN2	9	A*68:12	NSAENGDAK	258	141	NPAENGDAK	3679
4	THOC6	9	A*33:01	LWCRQPPYR	297	915	LWRRQPPYR	222
4	POLR2A	10	A*33:01	VQKIFHINPR	308	17699	AQKIFHINPR	738
	CDC25A	10	A*33:01	QSYCEPSSYR	309	53	QSYCEPPSYR	398
	ALMS1	9	A*68:12	SSSFSHREK	314	1496	SGSFSHREK	3554
	CDC25A	9	A*68:12	SYCEPSSYR	314	812	SYCEPPSYR	597
	ALMS1	10	A*33:01	TVPSSSFSHR	335	237	TVPSGSFSHR	378
E	THNSL2	9	A*68:12	TIESVQGAK	338	953	AIESVQGAK	3861
5	POLR2A	9	B*35:01	MIWNVQKIF	393	541	MIWNAQKIF	294
	CDC25A	9	B*35:01	QSYCEPSSY	478	50000	QSYCEPPSY	472
	DSCAML1	10	B*35:01	SSIRSFVLQY	480	9195	SSIRGFVLQY	391

T cells were tested for neoantigen reactivity by expanding them using autologous antigen presenting cells (APCs) pulsed with candidate neoantigen peptide pools (once per week X 4 weeks). As shown in FIG. 14B, reactivity in a IFN-γ ELISPOT assay was detected against Pool 2, but not against an irrelevant peptide (Tax peptide). Deconvolution of the pool revealed that the mutated (mut) *ALMS1* and *C6orf89* peptides within Pool 2 were immunogenic. *ALMS1* plays a role in ciliary function, cellular quiescence and intracellular transport, and mutations in this gene have been implicated in type II diabetes. *C6orf89* encodes a protein that interacts with bombesin receptor subtype-3, which is involved in cell cycle progression and wound repair of bronchial epithelial cells. Both mutated sites were not in conserved regions of the gene, and were not within genes previously reported to be mutated in cancer. Both of the target peptides were among the subset of 14 predicted peptides that could be experimentally confirmed to bind

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Pt 1's HLA alleles. The experimental binding scores of mut and wildtype (wt) *ALMS1* were 91 and 666 nM, respectively; and of mut- and wt-*C60RF89*, 131 and 1.7 nM, respectively (see e.g., FIG. 14C and Table 9). Both mutated genes localized to poorly conserved regions and did not localize to previously reported mutation sites in cancers (see e.g., FIGS. 15-16).

5 Example 20: CLL Patient 2 exhibited immunity against a mutated *FNDC3B* peptide that is naturally processed

In Patient 2, the ability personal neoantigens to contribute to memory T responses in the setting of long-lived remission was tested. From this individual, 26 non-synonymous missense mutations were identified. In total, 37 peptides from 16 mutations were predicted to bind to personal HLA alleles, of which 18 peptides from 12 mutations could be experimentally validated (15 with IC50 < 150; 3 with IC50 150-500 nM) (see e.g., FIG. 17A). In Pt 2, all 18 experimentally validated HLA-binding peptides were studied. T cell stimulations were performed using 3 pools of 6 peptides/pool (see e.g., Table 10). Table 10 shows a summary of peptides from Pt 2 missense mutations that were included in peptide pools for T cell stimulation studies. In Pt 2, all peptides that were experimentally confirmed to bind to HLA -A and -B alleles were used. 3 pools of peptides with 6 peptides/pool listed in decreasing order of experimental binding affinity of mutated peptides. The corresponding wildtype peptides and their predicted IC50 scores are included in the far right columns.

Table 10.

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				MU	T peptide		WT per	otide
			HLA		Predicted	Experimental		Predicted
Pool	Gene	Length	allele	Sequence	IC50 (nM)	IC50 (nM)	Sequence	IC50 (nM)
	NIN	10	A*02:01	FLQEETLTQM	10.63	1.1	FLQEERLTQM	45
	FNDC3B	9	A*02:01	VVMSWAPPV	4.21	6.2	VVLSWAPPV	9
1	SLC46A1	9	A*01:01	CSDSKLIGY	8.13	8.5	CWDSKLIGY	1778
	SYT15	9	B*080:1	EMLIKPKEL	414.37	8.9	EMLSKPKEL	785
	F2R	9	A*02:01	ILLMTVTSI	41.91	11	ILLMTVISI	53
	ACSM2A	9	A*02:01	SLMEHWALG	413.95	17	SLMEPWALG	1313
2	C16orf57	9	B*080:1	LLRVHTEHV	443.97	28	LLRVHTEQV	498.35

	ACSM2A	10	A*02:01	SLMEHWALGA	5.67	40	SLMEPWALGA	9.8
	TBC1D9B	9	A*02:01	KMTFLFPNL	63.7	62	KMTFLFANL	93
	SF3B1	9	A*02:01	GLVDEQQEV	22.26	94	GLVDEQQKV	51
	LRRC41	10	A*02:01	ALPDPILQSI	28.18	107	ALPGPILQSI	99
	LRRC41	9	A*02:01	GVWALPDPI	382.07	122	GVWALPGPI	963
	FNDC3B	10	A*02:01	AVVMSWAPPV	98.15	123	AVVLSWAPPV	89
	F2R	9	B*080:1	TSIDRFLAV	245.43	130	ISIDRFLAV	252
3	KIAA0467	9	B*07:02	GPSWGLSLM	179.31	137	GPSRGLSLM	39
	C16orf57	9	A*02:01	LLRVHTEHV	454.23	175	LLRVHTEQV	433.02
	C22orf28	10	A*02:01	WVNCSSMTFL	302.94	274	WVNRSSMTFL	835
	FNDC3B	10	A*02:01	VMSWAPPVGL	37.77	378	VLSWAPPVGL	48

Peptides with similar experimental binding scores were combined within the same pool. Responses were assessed after 2 rounds of weekly stimulations of T cells against mutated peptide pool-pulsed autologous APCs, and T cells were found to be reactive against Pool 1, as shown in FIG. 17B. Deconvolution of the pool revealed mut-*FNDC3B* to be the dominant immunogenic peptide among others within this pool (experimental IC50 of mut- and wt-*FNDC3B* were 6.2 and 2.7 nM, respectively; see e.g., FIG. 17C). The function of *FNDC3B* in blood malignancies is unclear, although down-regulation of *FNDC3B* expression is known to upregulate *miR-143* expression, which has been shown to differentiate prostate cancer stem cells and promote prostate cancer metastasis. Similar to *ALMS1* and *C6orf89*, the mutation in *FNDC3B* neither localized to evolutionarily conserved regions nor was it previously reported in other cancers (see e.g., FIGS. 15 and 16).

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T cell reactivity against mut-FNDC3B was polyfunctional (secreting GM-CSF, IFN- γ and IL-2), and specific to the mut-FNDC3B peptide but not its wildtype counterpart. Testing T cell reactivity against different concentrations of mut- and wt-FNDC3B peptides revealed a high avidity and specificity of mut-FNDC3B reactive T cells. T cell reactivity was abrogated by the presence of class I blocking antibody (W6/32), indicating that T cell reactivity was class I restricted (see e.g., FIGS. 17D-E). Moreover, the mut-FNDC3B peptide appeared to be a

naturally processed and presented peptide since T cell reactivity was detected against HLA-A2-expressing APCs that were transfected with a 300 basepair minigene encompassing the region of gene mutation but not the wildtype minigene, as shown in FIG. 17E, right panel.

Using a mut-FNDC3B/A2⁺-specific tetramer, a discrete population of mut-FNDC3B-reactive CD8⁺ T cells was detected within Pool 1-stimulated T cells (2.42% of the population) compared to control PBMCs from a healthy adult HLA-A2+ volunteer (0.38%), as shown in FIG. 17F. Gene expression analysis of FNDC3B in a large dataset of 182 CLL cases (including Pt 2) and 24 CD19⁺ B cells collected from normal volunteers revealed this gene to be relatively overexpressed in Patient 2 compared to other CLLs and normal B cells, as shown in FIG. 17G. Accordingly, it is clear that long-lived neoantigen-specific T cells could be tracked in CLL Patient 2.

To define the kinetics of mut-FNDC3B specific T cells in relationship to post-HSCT course, Pt 2 T cells isolated from different time points before and after HSCT were stimulated for 2 weeks and then tested for IFN- γ reactivity on ELISPOT. The emergence of mut-FNDC3B-specific T cells coincided with molecular remission and was sustained over time with continuous remission. As shown in FIG. 18 (top and middle panel), mut-FNDC3B T cell responses were not detected before or up to 3 months following HSCT. Molecular remission was first achieved 4 months following HSCT, and mut-FNDC3B-specific T cells were then first detected 6 months following HSCT. Antigen-specific reactivity subsequently waned (between 12 and 20 months post-HSCT), but was again strongly detected at 32 months post-HSCT. Based on molecular analysis of the TCR of the mut-FNDC3B-specific T cells, V β 11 was identified as the predominant CDR3 V β subfamily used by the reactive T cells, as shown in FIG. 19 and Table 11). Table 11 shows primers used for amplification of the TCR V β subfamily.

Table 11.

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		Amplicon size
Name	Forward primer sequence (5'-3')	(bp)
Vβ1	GCACAACAGTTCCCTGACTTGCAC	346

Vβ2	TCATCAACCATGCAAGCCTGACCT	349
Vβ3	GTCTCTAGAGAGAAGAAGGAGCGC	346
Vβ4	ACATATGAGAGTGGATTTGTCATT	378
Vβ5.1	ATACTTCAGTGAGACACAGAGAAAC	396
Vβ5.2	TTCCCTAACTATAGCTCTGAGCTG	343
Vβ6	AGGCCTGAGGGATCCGTCTC	340
Vβ7	CCTGAATGCCCCAACAGCTCTC	347
Vβ8	ATTTACTTTAACAACAACGTTCCG	404
V β9	CCTAAATCTCCAGACAAAGCTCAC	348
Vβ10	CCACGGAGTCAGGGGACACAGCAC	313
Vβ11	TCCAACCTGCAAAGCTTGAGGACT	312
Vβ12	CATGGGCTGAGGCTGATC	417
Vβ13.1	CAAGGAGAAGTCCCCAAT	372
Vβ13.2	GGTGAGGGTACAACTGCC	390
Vβ14	GTCTCTCGAAAAGAGAAGAGGAAT	349
Vβ15	AGTGTCTCCGACAGGCACAGGCT	352
Vβ16	AAAGAGTCTAAACAGGATGAGTCC	395
Vβ17	GGAGATATAGCTGAAGGGTA	372
Vβ18	GATGAGTCAGGAATGCCAAAGGAA	380
Vβ19	TCCTCTCACTGTGACATCGGCCCA	322
Vβ20	AGCTCTGAGGTGCCCCAGAATCTC	370

Vβ22	AAGTGATCTTGCGCTGTGTCCCCA	490
Vβ23	AGGACCCCCAGTTCCTCATTTC	435
Vβ24	CCCAGTTTGGAAAGCCAGTGACCC	509
Vβ25	TCAACAGTCTCCAGAATAAGGACG	352
Name	Reverse primer sequence (5'-3')	
External Cβ	GACAGCGGAAGTGGTTGCGGGGT	
Internal Cβ	FAM-CGGGCTGCTCCTTGAGGGGCTGCG	

This molecular information was used to develop a clone-specific nested PCR assay. Applying this assay, it was observed that T cells with the same specificity for mut-*FNDC3B* were not detected in PBMCs (n=3) and CD8⁺ T cells of normal healthy volunteers (see e.g., Table 12), but could be detected with similar kinetics as detection of IFN-γ secretion following HSCT in the patient as shown in FIG. 18, bottom panel. Although relative numbers of clone-specific T cells declined over time, lower concentrations of peptide antigen could stimulate T cell reactivity at 32 months compared to 6 months post-HSCT, indicating the emergence of potentially more antigen-sensitive memory T cells over time (see e.g., FIG. 18, inset).

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Table 12 shows detection of mut-*FNDC3B* specific TCR Vβ11, using T cell receptor-specific primers in Pt 2. A real-time PCR assay was designed to detect the mut-*FNDC3B*-specific TCR Vβ11 clone. This clone was not detectable in healthy donor PBMCs (n=3) or CD8 T cells, but clearly detectable in cDNA from mut-*FNDC3B* reactive T cells from Pt 2 (at 6 months post-HSCT). The PCR products were normalized over 18S ribosomal RNA. -, negative: no amplification; +, positive: amplification detected; ++, double positive: amplification detected and amplification level is more than median level of all positive samples.

Table 12.

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	Vβ11 Clone specific	
cDNA	PCR	18s ribosomal RNA
T cell clone	++	+
Healthy donor PBMCs		
(n=3)	-	+
Healthy donor CD8 T		
cells	-	+

Example 21: Large numbers of candidate neoantigens were predicted across diverse cancers

The overall somatic mutation rate of CLL is similar to other blood malignancies, but low in comparison to solid tumor malignancies (see e.g., FIG. 20A). To examine how tumor type and mutation rate impacts the abundance and quality of candidate neoantigens, the pipeline was applied to publicly available WES data from 13 malignancies – including high (melanoma (MEL)), lung squamous (LUSC) and adeno (LUAD) carcinoma, head and neck cancer (HNC), bladder cancer, colon and rectum adenocarcinoma, medium (glioblastoma (GBM), ovarian, clear cell renal carcinoma (clear cell RCC), and breast cancer) and low (CLL and acute myeloid leukemia (AML) cancers. To perform this analysis, a recently described algorithm that enables inference of HLA typing from the WES data was also implemented (Liu et al. 2013).

The overall mutation rate in these solid malignancies was an order of magnitude higher than for CLL and was associated with an increased median number of missense mutations. For example, melanoma displayed a median of 300 (range, 34-4276) missense mutations per case, while RCC had 41 (range, 10-101), respectively. Frameshift and splice-site mutations in RCC and melanoma were increased by only 2-3 fold in frequency as compared to CLL and summed neoORF length per sample were increased only moderately (by 5-13 fold). Overall, the median

number of predicted neopeptides with IC50 < 500 nM generated from missense and frameshift events per sample was proportional to the mutation rate; this was approximately 20- and 4-fold higher for melanoma (488; range, 18-5811) and RCC (80; range, 6-407)), respectively, compared to CLL (24; range 2-124). With a more stringent threshold of IC50 < 150 nM, the corresponding numbers of predicted neopeptides were 212, 35 and 10 for melanoma, RCC and CLL, respectively, as shown in FIG. 20B and Table 13).

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Table 13 shows the distribution of mutation classes, summed neoORF sizes and number of predicted binding peptides across 13 cancers. MEL:melanoma, LUSC: lung squamous cell carcinoma, LUAD: lung adenocarcinoma, BLCA: bladder, HNSC: head and neck cancer, COAD: colon adenocarcinoma, READ: renal adenocarcinoma, GBM: glioblastoma, OV: ovarian, RCC: clear cell renal carcinoma, BRCA: breast, CLL: chronic lymphocytic leukemia, AML: acute myeloid leukemia. *-predicted number of peptides based on missense and frameshift mutations.

Table 13.

Cancer type	# of r	# of mutations/sample median (range)	nple)	Summed NeoORF	# of predicted peptides median (range)*	ptides median je)*
	Missense	Frame shift	Splice site	length/Sample	IC50 < 150 (nM)	IC50 150-500 (nM)
MEL	300 (34- 4276)	2 (0-16)	4 (0-101)	48 (0-425)	212 (10-2566)	488 (18-5811)
TUSC	212 (0-2397)	3 (0-28)	5 (0-37)	86.5 (0-975)	149.5 (0-1320)	351.5 (0-2946)
LUAD	172.5 (0- 8971)	7 (0-61)	5 (0-127)	173.5 (0-2137)	122 (0-6999)	269.5 (1- 16360)
BLCA	161.5 28- 1194)	6 (0-22)	4 (0-22)	152 (0-780)	97 19-1073)	232.5 (59- 2337)
HNSC	95 (2-1400)	5 (0-106)	2 (0-29)	124.5 (0-2585)	66.5 (2-1139)	159.5 (3-2916)
COAD	93 (32- 5902)	4 (1-182)	(96-0) 0	121 (9-4794)	68 (15-2155)	172 (40-5199)
READ	72.5 (37- 1837)	2 (0-31)	0 (0-2)	51(0-929)	52 (14-1215)	114 (38-2750)
GBM	47 (0- 169)	2 (0-16)	1 (0-5)	(0-239)	39 (0-166)	90 (0-332)
۸٥	42 (9-149)	1 (0-7)	1 (0-6)	7.5 (0-328)	30 (3-181)	70 (13-420)
RCC	41(10-101)	6 (0-22)	1(0-8)	143 (0-813)	35 (2-223)	80 (6-407)
BRCA	25 (1-300)	2 (0-54)	1 (0-8)	37 (0-1415)	21 (0-346)	47 (0-781)
CLL	16 (0-75)	1 (0-9)	1 (0-6)	11 (0-427)	10 (0-20)	24 (2-124)
AML	7 (0-20)	1 (0-2)	0 (0-3)	6 (0-160)	4 (0-19)	8 (0-41)

* Refers only to predicted epitopes arising from missense mutations.

Example 22: Clinical strategies for addressing clonal mutations

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"Clonal" mutations are those that are found in all cancer cells within a tumor, while "subclonal" mutations are those that statistically are not in all cancer cells and therefore are derived from a sub population within the tumor.

According to the techniques herein, bioinformatic analysis may be used to estimate clonality of mutations. For example, the ABSOLUTE algorithm (Carter et al, 2012, Landau et al, 2013) may be used to estimate tumor purity, ploidy, absolute copy numbers and clonality of mutations. Probability density distributions of allelic fractions of each mutation may be generated followed by conversion to cancer cell fractions (CCFs) of the mutations. Mutations may be classified as clonal or subclonal based on whether the posterior probability of their CCF exceeds 0.95 is greater or lesser than 0.5 respectively.

It is contemplated within the scope of the disclosure that a neoantigen vaccine may include peptides to clonal, sub-clonal or both types of mutations. The decision may depend on the disease stage of the patient and the tumor sample(s) sequenced. For an initial clinical study in the adjuvant setting, it may not be necessary to distinguish between the two mutations types during peptide selection, however, one of skill in the art will appreciate that such information may be useful in guiding future studies for a number of reasons.

First, subject tumor cells may be genetically heterogeneous. Multiple studies have been published in which tumors representing different stages of disease progression have been evaluated for heterogeneity. These include examining the evolution from a pre-malignant disease (Myelodysplastic syndrome) to leukemia (secondary acute myelogenous leukemia [AML]) (Walter et al 2012), relapse following therapy-induced remission of AML(Ding et al 2012), evolution from primary to metastatic breast cancer and medulloblastomas (Ding et al

2012; Wu et al Nature 2012), and evolution from primary to highly metastatic pancreatic and renal cancers (Yachida et al 2012; Gerlinger et al 2012). Most studies utilized genome or exome sequencing but one study also evaluated copy number variations and CpG methylation pattern variations. These studies have shown that genetic events are acquired during cancer cell growth which alter the profile of mutations. Many, and usually most (40 % - 90%), of the earliest detectable mutations ("founder mutations") persist in all evolved variants but new mutations unique to evolved clones do arise and these may be distinct between different evolved clones. These changes can be driven by host/cancer cell "environmental" pressures and/or therapeutic intervention and thus more highly metastatic disease or prior therapeutic intervention generally lead to more significant heterogeneity.

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Second, it is contemplated that a single tumor for each patient may be initially sequenced, which may provide a snapshot of the profile of genetic variation for that particular point in time. The sequenced tumor may be derived from a clinically evident lymph node, in transit/satellite metastasis, or resectable visceral metastasis. None of the initially tested patients will have disease that has clinically progressed to multiple sites; however, it is contemplated that the techniques described herein in will be broadly applicable to patients have cancer that has progressed to multiple sites. Within this tumor cell population, "clonal mutations" may be comprised of both founder mutations and any novel mutations present in the cell that seeded the resected tumor and sub-clonal mutations represent those that evolved during growth of the resected tumor.

Third, the clinically important tumor cells for the vaccine induced T-cells to target are frequently not the resected tumor cells but rather other currently undetectable tumor cells within a given patient. These cells may have spread directly from the primary tumor or from the

resected tumor, may have derived from a dominant or sub-dominant population within the seeding tumor and may have genetically evolved further at the surgically resected site. These events are currently unpredictable.

Thus, for the surgically resected adjuvant setting, there is no a priori way to decide whether mutations found in the resected tumor that are clonal or subclonal represent the optimal choice for targeting other non-resected cancer cells. For example, mutations that are subclonal within the resected tumor may be clonal at other sites if those other sites were seeded from a subpopulation of cells containing the sub-clonal mutation within the resected tumor.

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In other disease settings however, such as settings in which patients carry multiple and metastatic lesions, sequencing of more than one lesion (or parts of lesion) or lesions from different time points may provide more information relative to effective peptide selection.

Clonal mutations may typically be prioritized in the design of neo-antigen epitopes for the vaccine. In some instances, especially as the tumor evolves and sequencing details from metastatic lesions are evaluated for an individual patient, certain subclonal mutations may be prioritized for consideration as part of peptide selection.

Example 23: Personalized cancer vaccines stimulate immunity against tumor neoantigens

The above-described detailed integration of comprehensive bioinformatics with functional data in CLL and other cancers provides several novel biological insights. First, although CLL is a relatively low mutation rate cancer, it was nonetheless possible to identify epitopes generated by somatic mutations that elicited long-term T cell responses. Whole-exome sequencing data from 31 CLL samples revealed that per case, a median of 22 peptides (range, 6-81) were predicted to bind to personal HLA-A and -B alleles with IC50 < 500nM originating from a median of 16 (range, 2-75) missense mutations. Approximately 75% and half (54.5%) of

predicted peptides with IC50 < 150 nM and 500 nM, respectively, were experimentally validated to bind to the patient's HLA alleles. RNA expression analysis showed that nearly 90% of the cognate genes corresponding to the predicted mutated peptides were confirmed to be expressed in CLL cells and expression of a transcript from the mutated allele was detected in each of the three (data not shown) examples tested. Only a fraction of all neoepitopes had generated a spontaneous T-cell response although this response was still detectable years after transplant; ~6% (3/48) of all predicted and tested mutated peptides or 9% (3/32) of experimentally validated and tested mutated peptides stimulated IFN-γ secretion responses from patient T cells. This rate of neo-epitope discovery in CLL, a low mutation rate tumor, is remarkably similar to the rates recently reported in melanoma (4.5%, or 11/247 peptides; Robbins PF, Lu YC, El-Gamil M, et al: Mining exomic sequencing data to identify mutated antigens recognized by adoptively transferred tumor-reactive T cells. Nat Med, 2013), a high mutation rate cancer. Hence, functional neoepitopes can be systematically discovered across the broad range of cancers including low mutation rate tumors.

A second key finding is that T cell responses against CLL neoepitopes were long-lived (on the order of several years), associated with continuous disease remission and were generated during *in vitro* stimulation in a timeframe consistent with memory T cell responses. These studies add to the growing literature that responses against tumor neoantigens contribute to efficacious immune responses. Thus, although approximately 5% of predicted peptides generated from missense mutations yielded detectable T cell responses, the kinetics of the response suggest a possible role in ongoing anti-leukemia surveillance functions. The functional impact of neoantigen-directed T-cell responses is supported by a recent study from Castle et al. (Castle JC, Kreiter S, Diekmann J, et al: Exploiting the mutanome for tumor vaccination. Cancer Res

72:1081-1091, 2012) who identified candidate neoepitopes by WES of B16 murine melanoma and prediction of peptide-HLA allele binders. A subset of these predicted epitopes not only elicited immune responses that were specific to the mutated peptide and not the wildtype counterpart, but could also control the disease both therapeutically and prophylactically. While it was difficult to directly compare the relative contributions of tumor neoantigens versus other types of CLL antigens such as overexpressed or shared native antigens (in contrast to melanoma, CLL tumor antigens are not well characterized) or to the GvL response, prior characterization of antigen-specific T cell responses from a melanoma patient with prolonged survival suggest that anti-neoantigen immunity is more prolonged and sustained over time than that against shared overexpressed tumor antigens.

Third, these results highlight the concept that targeting tumor-specific "trunk" mutations can be impactful from the immunologic standpoint. All three of the immunogenic neoantigens (mutated *FND3CB*, *ALMS1*, *C6orf89*) in the two patients appeared to be passenger mutations, not directly contributory to the oncogenic process, and were clonal, affecting the bulk of the cancer mass. Several features of these immunogenic mutations suggest them to be passenger mutations: lack of sequence conservation around the mutation and lack of previously reported mutations in other cancers at the observed sites. Because clonal evolution is a fundamental feature of cancer, it has been posited that immunologic targeting of cancer drivers would have the advantage of minimal antigenic drift, given their essentiality in tumor function that would require them to be maintained in the face of selective pressure. Although such an advantage may be possible, it is apparently not a requirement. Additionally, driver mutations may not necessarily generate immunogenic peptides. For example, the *TP53-S83R* mutation in Patient 2 did not generate a predicted epitope of < 500 nM against any of its class I HLA-A or -B alleles.

Finally, analysis of the binding characteristics of the neoantigen data from the literature (Table 4) as well as the candidate neoepitopes from the data in CLL revealed conceptual insights into the types of point mutations most likely to effectively create a T cell response. It was found that a consistent feature of immunogenic neoepitopes was a predicted binding affinity < 500 nM (3 of 3 of immunogenic CLL peptides and 30 of 33 [91%] of the historical functional 5 neoepitopes) and the majority of these (92%) displayed predicted affinities < 150 nM. Unexpectedly however, in most cases (3 of 3 immunogenic CLL peptides and 27 of 33 [82%] historical functional epitopes), the corresponding wild type epitopes were also predicted to bind with comparable strong/intermediate (< 150 nM, Group 1 in Table 4) or weak (150 – 500 nM, Group 2 in Table 4) affinity. The data support the idea that two types of mutations are 10 commonly observed among naturally occurring T-cell responses to neoantigens: (1) mutations at positions that lead to substantially better binding to the MHC allele (mutated ALMS1 as well as 6 of 33 (18%) of the historical functionally-identified neoepitopes ['Group 3', Table 4]), presumably due to improved interaction with MHC, or (2) mutations at positions that do not significantly interact with MHC but instead presumably alter the T cell receptor binding ((2 of 3 CLL epitopes [FNDC3B and C6orf89] and 24 of 33 (73%) naturally immunogenic neoepitopes ['Group 1' and 'Group 2', Table 4]). The distinction between these two types of mutations fits with the concept that the peptide can be considered as a "key', which must fit both the MHC and the TCR "locks" in order to stimulate cytolysis, allowing mutations to independently vary MHC or TCR binding. Excepting the contribution of minor histocompatibility antigens to graft-vs-20 host disease, there are no reports of auto-immune sequelae linked to neoantigens in these patients, even in those patients where a reaction occurs to a mutated peptide and the cognate native peptide is predicted to be a tight binder. This result is consistent with the idea that MHC-

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binding native peptides are normally involved in the negative selection process in which T cells bearing TCRs reactive to these native peptides are thymically deleted or rendered anergic, and yet the T cell repertoire can accommodate the development of a specific immune response to a neoeptiope peptide due to an altered presentation of the mutated peptide to the T cell receptor. It is clear that each individual tumor in a patient may harbor a broad spectrum of both shared and personal genetic alterations that may continue to evolve in response to the environment, and that this progression may often lead to resistance to therapy. Given the uniqueness and plasticity of tumors, an optimal therapy may need to be customized based on the exact mutations present in each tumor, and may need to target multiple nodes to avoid resistance. The vast repertoire of human CTLs has the potential to create such a therapy that targets multiple, personalized tumor antigens. As discussed above, the present disclosure shows that it is possible to systematically identify CTL target antigens harboring tumor-specific mutations by using massively parallel sequencing in combination with algorithms that effectively predict HLA-binding peptides. Advantageously, the present disclosure allows tumor neoantigens in a variety of low and high mutation rate cancers to be predicted, and experimentally identifies long-lived CTLs that target leukemia neoantigens in CLL patients. The present disclosure supports the existence of protective immunity targeting tumor neoantigens, and provides a method for selecting neoantigens for personalized tumor vaccines.

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As discussed in detail above, the techniques described herein were applied to a unique group of CLL patients who developed clinically evident durable remission associated with antitumor immune responses following allogeneic-HSCT. These graft-versus-leukemia responses have typically been attributed to allo-reactive immune responses targeting hematopoietic cells. However, the above described results indicate that the GvL response is also associated with

CTLs that recognize personal leukemia neoantigens. These results are consistent with data indicating that the existence of GvL-associated CTLs with specificity for tumor, rather than alloantigens. It has been postulated that neoantigen-reactive CTLs are important in cancer surveillance because the study of a long-term melanoma survivor found that CTLs targeting neoantigens are significantly more abundant and sustained than those against non-mutated overexpressed tumor antigens (Lennerz V, Fatho M, Gentilini C, et al: The response of autologous T cells to a human melanoma is dominated by mutated neoantigens. Proc Natl Acad Sci U S A 102:16013-8, 2005). The data presented above is consistent with this melanoma study because neoantigen-specific T cell responses in CLL patients were found to be long-lived (on the order of several years) memory T cells (based on their rapid stimulation kinetics *in vitro*) and associated with continuous disease remission. Accordingly, neoantigen-reactive CTLs likely play an active role in controlling leukemia in transplanted CLL patients.

More generally, the abundance of neoantigens across many tumors was estimated and found to be ~1.5 HLA-binding peptides with IC50<500nM per point mutation and ~ 4 binding peptides per frameshift mutation. As expected, the rate of predicted HLA binding peptides mirrored the somatic mutation rate per tumor type (see e.g., FIG. 20). Two approaches were used to study the relationship between predicted binding affinity and immunogenic neoantigens that induce CTLs. The above-described techniques were applied to published immunogenic tumor neoantigens (i.e. in which reactive CTLs were observed in patients) demonstrated that the vast majority (91%) of functional neoantigens are predicted to bind HLA with IC50<500nM (with ~70% of wild type counterpart epitopes predicted to bind at a similar affinity) (see e.g., Table 4). This test used a gold standard set of neoantigens confirmed that the techniques described herein correctly classify true positives. A prospective prediction of neoepitopes followed by functional

validation showed that 6% (3/48) of predicted epitopes were associated with neoantigen-specific T cell responses in patients -- comparable to the rate of 4.8% found recently for melanoma. The low proportion does not necessarily imply low prediction accuracy for the algorithm. Rather, the number of true neoantigens is greatly underestimated because: (i) allo-HSCT is a general cellular therapy likely to induce only a small number of neoantigen-specific T cell memory clones; and (ii) standard T cell expansion methods are not sensitive enough to detect naïve T cells that represent a much larger part of the repertoire but with much lower precursor frequencies.

Although the frequency of CTLs that target neoORFs has yet to be measured, it is specifically contemplated within the scope of the invention that this class of neoantigens may be an excellent candidate neoepitope because it is likely to be more specific (for lack of a wild type counterpart) and immunogenic (as a result of bypassing thymic tolerance).

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With the ongoing development of highly powerful vaccination reagents, the present disclosure provides techniques that make it feasible to generate personalized cancer vaccines that effectively stimulate immunity against tumor neoantigens.

MATERIALS AND METHODS

Patient samples: Heparinized blood was obtained from patients enrolled on clinical research protocols at the Dana-Farber Cancer Institute (DFCI). All clinical protocols were approved by the DFCI Human Subjects Protection Committee. Peripheral blood mononuclear cells (PBMCs) from patient samples were isolated by Ficoll/Hypaque density-gradient centrifugation, cryopreserved with 10% DMSO, and stored in vapor-phase liquid nitrogen until the time of analysis. For a subset of patients, HLA typing was performed by either molecular or serological typing (Tissue Typing Laboratory, Brigham and Women's Hospital, Boston, MA).

Whole exome capture sequencing data for CLL and other cancers: The list for melanoma was obtained from dbGaP database (phs000452.v1.p1) and for the 11 other cancers. through TCGA (available through the Sage Bionetworks' Synapse resource (on the worldwide web at (www)synapse.org/#!Synapse:syn1729383)). The HLA-A, HLA-B and HLA-C loci in 2488 samples across these 13 tumor types were sequenced using a two-stage likelihood based approach, and this data is summarized in Table 14. Briefly, a dedicated sequence library consisting of all known HLA alleles (6597 unique entries), based on the IMGT database, was constructed. From this resource, a secondary library of 38-mers was generated, and putative reads emanating from the HLA locus were extracted from total sequence reads based on perfect matches against it. The extracted reads were then aligned to the IMGT-based HLA sequence library using the Novoalign software (on the worldwide web at (www)novocraft.com), and HLA alleles were inferred through a two-stage likelihood calculation. In the first stage, populationbased frequencies were used as priors for each allele and the posterior likelihoods were calculated based on quality and insert size distributions of aligned reads. Alleles with the highest likelihoods for each of HLA-A, B and C genes were identified as the first set of alleles. A heuristic weighting strategy of the computed likelihoods in conjunction with the first set of winners were then used to identify the second set of alleles.

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Table 14 shows TCGA patient IDs for neoantigen load estimates across cancers. LUSC (lung squamous carcinoma), LUAD (lung adeno carcinoma), BLCA (bladder), HNSC (head and neck), COAD (colon) and READ (rectum), GBM (glioblastoma), OV (ovarian), RCC (clear cell renal carcinoma), AML (acute myeloid leukemia) and BRCA (breast),

Table 14

TCGA Barcodes	Disease	UUID
TCGA-BL-A0C8-01A-11D-A10S-08	BLCA	134b0a5e-a0ba-444d-bc4b-bdceb02d5b04
TCGA-BL-A13I-01A-11D-A13W-08	BLCA	aa490522-7bb9-4f82-8f19-eaf63f719bfe
TCGA-BL-A13J-01A-11D-A10S-08	BLCA	0c7aca3f-e006-4de3-afc2-20b4f727d4fd
TCGA-BL-A3JM-01A-12D-A21A-08	BLCA	b181ba68-f50f-4faf-b7b5-356e119b5f04
TCGA-BT-A0S7-01A-11D-A10S-08	BLCA	b2e5d244-94c1-4dbf-8d33-34b595903310
TCGA-BT-A0YX-01A-11D-A10S-08	BLCA	d61ccd8c-b798-46e0-aeed-f95b4f3ba4ff
TCGA-BT-A20J-01A-11D-A14W-08	BLCA	1d3c0ff9-d149-4d21-8955-5fb849fc5462
TCGA-BT-A20N-01A-11D-A14W-08	BLCA	341bbffe-7587-4ad0-b3b4-68e64080e216
TCGA-BT-A20O-01A-21D-A14W-08	BLCA	7df63263-de4e-4ed8-804f-9e8fee3be2d5
TCGA-BT-A20P-01A-11D-A14W-08	BLCA	e6c78a98-f45b-482b-a551-4f11b8c1ff8b
TCGA-BT-A20Q-01A-11D-A14W-08	BLCA	8c619cbc-9e91-4716-9711-5236e55d8f46
TCGA-BT-A20R-01A-12D-A16O-08	BLCA	e9bbbfc3-0beb-4f91-92a1-081bff7c4a07
TCGA-BT-A20T-01A-11D-A14W-08	BLCA	301d6ce3-4099-4c1d-8e50-c04b7ce91450
TCGA-BT-A20U-01A-11D-A14W-08	BLCA	4576527b-b288-4f50-a9ea-5d5dede22561
TCGA-BT-A20V-01A-11D-A14W-08	BLCA	973d0577-8ca4-44a1-817f-1d3c1bada151
TCGA-BT-A20W-01A-21D-A14W-08	BLCA	85ccdf9b-f787-4701-822f-ae0fce5b4fc5
TCGA-BT-A20X-01A-11D-A16O-08	BLCA	9b4586ee-4091-484f-8be8-5a5196fe7b6f
TCGA-BT-A2LB-01A-11D-A18F-08	BLCA	e7aea186-f13b-43b1-8693-f90f51e005dd
TCGA-BT-A2LD-01A-12D-A20D-08	BLCA	cc95719c-7fcc-4ed7-837e-1840c0a6bc27
TCGA-BT-A3PH-01A-11D-A21Z-08	BLCA	cda1a403-16b6-487c-a82a-c377d1d0f89d
TCGA-BT-A3PJ-01A-21D-A21Z-08	BLCA	b73523d7-f5a5-4140-8537-4df4d1ecf465
TCGA-BT-A3PK-01A-21D-A21Z-08	BLCA	4ad38e8e-e63e-41d9-9216-617be7fa1d75
TCGA-C4-A0EZ-01A-21D-A10S-08	BLCA	b01a7081-8eb5-4728-a517-52156cdfe7ed
TCGA-C4-A0F0-01A-12D-A10S-08	BLCA	612fd956-9a41-4201-9d74-6ab50f6ae987
TCGA-C4-A0F1-01A-11D-A10S-08	BLCA	9377460a-8497-41b8-b2c2-5f50cfeda1fe
TCGA-C4-A0F6-01A-11D-A10S-08	BLCA	608f8c75-40e4-44f2-bdde-5f07aa6b4bee
TCGA-C4-A0F7-01A-11D-A10S-08	BLCA	f389176f-d8f3-45c2-aae4-7378a3d6fc7f
TCGA-CF-A1HR-01A-11D-A13W-08	BLCA	69acf4f1-063f-453d-b148-681518c0bc39
TCGA-CF-A1HS-01A-11D-A13W-08	BLCA	b36e672b-c5d8-4481-bbb3-7be805215212
TCGA-CF-A27C-01A-11D-A16O-08	BLCA	acc629cb-ad03-4cec-9b21-922e4932ef3e
TCGA-CF-A3MF-01A-12D-A21A-08	BLCA	c66c92d5-df65-46e6-861d-d8a98808e6a3
TCGA-CF-A3MG-01A-11D-A20D-08	BLCA	4c89ce08-ed24-4179-8884-4706660b7da8
TCGA-CF-A3MH-01A-11D-A20D-08	BLCA	8867b16f-cd05-41e9-b3ca-4c72a1ebeb70
TCGA-CF-A3MI-01A-11D-A20D-08	BLCA	0afabd62-8454-41b4-9b02-386681589688
TCGA-CU-A0YN-01A-21D-A10S-08	BLCA	803ab221-b813-4bcc-95a9-1f686d172d3c
TCGA-CU-A0YO-01A-11D-A10S-08	BLCA	e80278f9-2059-4e98-92b2-3e9868fc5818
TCGA-CU-A0YR-01A-12D-A10S-08	BLCA	31382822-3792-47bc-99e8-8a1ee1e4e58b

TCGA-CU-A3KJ-01A-11D-A21A-08 BLCA e22c6a44-4f8e-44eb-8ca8-dff0f2fc5575	
TCGA-DK-A1A3-01A-11D-A13W-08 BLCA 2322f7cd-7d55-4a9f-b7f3-da3068089383	
TCGA-DK-A1A5-01A-11D-A13W-08 BLCA 448fe471-3f4e-4dc8-a4e0-6f147dc93abe	
TCGA-DK-A1A6-01A-11D-A13W-08 BLCA df8a913c-5160-4fc5-950d-7c890e24e820	
TCGA-DK-A1A7-01A-11D-A13W-08 BLCA 91f458e6-64b7-454d-a542-b0aa23638fd8	
TCGA-DK-A1AA-01A-11D-A13W-08 BLCA 804ffa2e-158b-447d-945c-707684134c87	
TCGA-DK-A1AB-01A-11D-A13W-08 BLCA 5f0fb2ba-0351-4ce0-8b74-31aa3deecae1	
TCGA-DK-A1AC-01A-11D-A13W-08 BLCA a5dc17f5-abda-4534-b0f8-34b59ed4faa3	
TCGA-DK-A1AD-01A-11D-A13W-08 BLCA 32398d56-8668-41b1-9c0b-c6aea6e3e787	
TCGA-DK-A1AE-01A-11D-A13W-08 BLCA abd2d959-d5ed-4eb3-9759-67eb1aa23325	
TCGA-DK-A1AF-01A-11D-A13W-08 BLCA fbdcd7f9-1901-4e90-8e3c-71b05dc96da1	
TCGA-DK-A1AG-01A-11D-A13W-08 BLCA 7d2a22eb-7344-4cba-ad7d-94c3f9ef3d7c	
TCGA-DK-A2HX-01A-12D-A18F-08 BLCA a8f0d416-2102-43ea-9cf1-465c37f9642a	
TCGA-DK-A2I1-01A-11D-A17V-08 BLCA f350676a-e308-42fe-8297-9d18ba7027b1	
TCGA-DK-A2I2-01A-11D-A17V-08 BLCA 537e0d59-dd1c-479e-877f-eb9523c0967e	
TCGA-DK-A2I4-01A-11D-A21A-08 BLCA d68074b8-ce96-4dc5-b14c-3bbc7ba92ad9	
TCGA-DK-A2I6-01A-12D-A18F-08 BLCA 97a755af-ca00-4116-8a32-0984dbfb1585	
TCGA-DK-A3IK-01A-32D-A21A-08 BLCA f730e341-8102-4405-95e2-46a3455a35cc	
TCGA-DK-A3IL-01A-11D-A20D-08 BLCA 4838b5a9-968c-4178-bffb-3fafe1f6dc09	
TCGA-DK-A3IM-01A-11D-A20D-08 BLCA 780f4201-4e59-47b8-b3b7-d322a6162b2d	
TCGA-DK-A3IN-01A-11D-A20D-08 BLCA 173c1518-6bcb-4e25-a119-de32dab91286	
TCGA-DK-A3IQ-01A-31D-A20D-08 BLCA c3da3cc2-2299-4a3e-9de8-7a1d0a10345d	
TCGA-DK-A3IS-01A-21D-A21A-08 BLCA 92a59313-da12-4896-b164-fd2d50684638	
TCGA-DK-A3IT-01A-31D-A20D-08 BLCA 07db4596-cb49-4a32-bc99-3b202ffe61a2	
TCGA-DK-A3IU-01A-11D-A20D-08 BLCA 52de410f-3ce3-4ee6-87f3-8ec2e829962f	
TCGA-DK-A3IV-01A-22D-A21A-08 BLCA 7cecfbbc-5fe4-4413-95fd-07533aacbb73	
TCGA-E5-A2PC-01A-11D-A202-08 BLCA 62b9f71c-2dab-455a-a454-579e8843f712	
TCGA-FD-A3B3-01A-12D-A202-08 BLCA 8e9fb61d-c90d-440b-857a-12e1048435ea	
TCGA-FD-A3B4-01A-12D-A202-08 BLCA df922c85-5a10-487f-a9d5-220d5090e2e4	
TCGA-FD-A3B5-01A-11D-A20D-08 BLCA d05f9b81-7ba9-4231-aae6-1d2c14df22d7	
TCGA-FD-A3B6-01A-21D-A20D-08 BLCA 36524c53-ac54-4a42-a982-bed2e4354268	
TCGA-FD-A3B7-01A-31D-A20D-08 BLCA fc76c5bd-315d-4981-ae53-705f40d2c078	
TCGA-FD-A3B8-01A-31D-A20D-08 BLCA 7957bb77-8329-43a0-b1a8-140f2cb6b91b	
TCGA-FD-A3N5-01A-11D-A21A-08 BLCA 418a3dec-96ff-4719-becb-e1a8260cce2f	
TCGA-FD-A3N6-01A-11D-A21A-08 BLCA d4615ca0-b5c7-4a5c-8593-bd50034a78ae	
TCGA-FD-A3NA-01A-11D-A21A-08 BLCA d079a32c-270b-4c43-8372-884e8d0c48ed	
TCGA-G2-A2EC-01A-11D-A17V-08 BLCA 1376c881-cea5-4470-8dc1-63c69f201570	
TCGA-G2-A2EF-01A-12D-A18F-08 BLCA 4e5917bd-2cb1-438c-a46c-5d8ca5b2fd0e	
TCGA-G2-A2EJ-01A-11D-A17V-08 BLCA 82f98ff9-7161-45c3-8107-033b47e25f21	
TCGA-G2-A2EK-01A-22D-A18F-08 BLCA eb73bb35-af99-47b8-8bbb-33b5374e5c74	
TCGA-G2-A2EL-01A-12D-A18F-08 BLCA 56924619-0724-4b3e-9c53-27c27d3789d6	

TCGA-G2-A2EO-01A-11D-A17V-08	BLCA	ebb5cdb6-df4a-436d-b4a6-1655d263e3dd
TCGA-G2-A2ES-01A-11D-A17V-08	BLCA	5c628df6-a848-4177-87b8-714788118980
TCGA-G2-A3IE-01A-11D-A20D-08	BLCA	ebacd09f-c204-4cd2-a087-07bc4f2c5b74
TCGA-GC-A3I6-01A-11D-A20D-08	BLCA	372feefe-ee84-4833-8651-8f023f38a56a
TCGA-GC-A3RB-01A-12D-A21Z-08	BLCA	eaf54383-4286-4416-9b18-be1081797df2
TCGA-GD-A2C5-01A-12D-A17V-08	BLCA	2b142863-b963-4cc9-8f8f-c72503c93390
TCGA-GD-A3OP-01A-21D-A21Z-08	BLCA	3e02d723-691a-448c-85e2-4e39a3696ba5
TCGA-GD-A3OQ-01A-32D-A21Z-08	BLCA	fb985b3d-b0f7-42a0-bc3c-f71d9c5f78d8
TCGA-GD-A3OS-01A-12D-A21Z-08	BLCA	9b3e164d-aaa0-4bb5-b7b8-6264b2746a47
TCGA-GV-A3JV-01A-11D-A21Z-08	BLCA	5fed4b8a-4b59-4424-bbf1-bc73ce041361
TCGA-GV-A3JW-01A-11D-A20D-08	BLCA	4534413b-d0d0-4b34-a3d4-f821705485ae
TCGA-GV-A3JX-01A-11D-A20D-08	BLCA	21525d6f-4222-4e0a-9f07-8adbbd55c54f
TCGA-GV-A3JZ-01A-11D-A21A-08	BLCA	074fc904-0a0e-4114-b569-89d51e7a89db
TCGA-GV-A3QG-01A-11D-A21Z-08	BLCA	90534196-b1d8-4054-b4d5-1d29943b52bc
TCGA-GV-A3QI-01A-11D-A21Z-08	BLCA	33a9da52-5471-456f-84cb-13c5de5b0994
TCGA-H4-A2HO-01A-11D-A17V-08	BLCA	2e327841-eef0-42dd-883e-7d5b5a0d3a93
TCGA-H4-A2HQ-01A-11D-A17V-08	BLCA	94108975-b7a0-40ba-ad39-e44cc62e8cc0
TCGA-HQ-A2OE-01A-11D-A202-08	BLCA	61324839-e90a-49f2-a9c9-629d7b125fe9
TCGA-A1-A0SB-01A-11D-A142-09	BRCA	db9d40fb-bfce-4c3b-a6c2-41c5c88982f1
TCGA-A1-A0SD-01A-11D-A10Y-09	BRCA	1847727f-ea57-4e2e-84e5-a10e764c9096
TCGA-A1-A0SE-01A-11D-A099-09	BRCA	0539776c-3943-41d0-972c-8dc833a603e5
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TCGA-A1-A0SG-01A-11D-A142-09	BRCA	39642c6d-9191-4746-8a9d-62d437bfdce8
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TCGA-A1-A0SJ-01A-11D-A099-09	BRCA	a55c6a44-c0f5-4300-8df4-4a70befe2d3b
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TCGA-A1-A0SM-01A-11D-A099-09	BRCA	2057b341-ff5c-45ef-83bb-005e29b2e740
TCGA-A1-A0SN-01A-11D-A142-09	BRCA	1b8d93f4-acc2-48ee-9ca8-a327eb0463c2
TCGA-A1-A0SO-01A-22D-A099-09	BRCA	b3568259-c63c-4eb1-bbc7-af711ddd33db
TCGA-A1-A0SP-01A-11D-A099-09	BRCA	d3ae9617-b6cd-4d98-b631-39bd4afd3c4e
TCGA-A1-A0SQ-01A-21D-A142-09	BRCA	9055ddce-a0ff-4980-af86-c07f949acbc3
TCGA-A2-A04N-01A-11D-A10Y-09	BRCA	389dd52b-a7b7-46f0-83ae-308e485466a8
TCGA-A2-A04P-01A-31D-A128-09	BRCA	a85cf239-ff51-46e7-9b88-4c2cb49c66b9
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TCGA-A2-A04U-01A-11D-A10Y-09	BRCA	f819433a-44db-4022-abdb-d6123cfa30b2
TCGA-A2-A04V-01A-21W-A050-09	BRCA	89501861-2778-4b88-9a44-939fed99850d
TCGA-A2-A04W-01A-31D-A10Y-09	BRCA	7822a6b1-68c8-4675-993c-c4b54a510c09
TCGA-A2-A04X-01A-21W-A050-09	BRCA	66a73891-2fea-450c-8224-0865d98b4346
TCGA-A2-A04Y-01A-21W-A050-09	BRCA	3669bbbd-2e75-4b57-a5a8-8eebc25a97c2

TCGA-A2-A0CL-01A-11D-A10Y-09			
TCGA-A2-A0CP-01A-11W-A050-09 BRCA a776e274-fe9f-49a9-83ab-95ca6819-96b TCGA-A2-A0CQ-01A-21W-A050-09 BRCA fa0d7183-8757-495-87b2-2366a1dbd508 TCGA-A2-A0CC-01A-11D-A10Y-09 BRCA fe96b832-68b6.4499-98a-5124a43d5c95 TCGA-A2-A0CU-01A-12W-A050-09 BRCA 2b412ad8-abda-4cR8-868-590bee80031e TCGA-A2-A0CU-01A-12W-A050-09 BRCA 2b412ad8-abda-4cR8-868-590bee80031e TCGA-A2-A0CU-01A-12W-A050-09 BRCA 5d1dead5-d9a5-42d3-a703-4c38ad6e8f57 TCGA-A2-A0CW-01A-21D-A10Y-09 BRCA da4f085-b16f-40fa-95c6-524d70d7ac4d TCGA-A2-A0CW-01A-21W-A019-09 BRCA 95dxb606-367a-46b5-b66-254d70d7ac4d TCGA-A2-A0CW-01A-11W-A019-09 BRCA 95dxb606-367a-46b5-b66-2624d70d7ac4d TCGA-A2-A0CW-01A-11W-A019-09 BRCA 3752b0f6-3631-46b5-b66-242ad-263f2e212 TCGA-A2-A0D0-01A-11W-A050-09 BRCA a762809e-15c9-485e-ad7a-de38d-aca3f2e2a2 TCGA-A2-A0D1-01A-11W-A050-09 BRCA a8183420a-7744-4024-b3db-6553ad293988 TCGA-A2-A0D2-01A-21W-A050-09 BRCA 8183420a-7744-4024-b3db-6553ad293988 TCGA-A2-A0E0-01A-11W-A050-09 BRCA 123ac2ad7-6866-467-9ec-8a068dd88d896 TCGA-A2-A0EW-01A-12W-A050-09 B	TCGA-A2-A0CL-01A-11D-A10Y-09	BRCA	a630ed59-dd23-45e1-aa16-4f7a98e32728
TCGA-A2-A0CQ-01A-21W-A050-09 BRCA fa0d7183-8757-495-8762-2366a1dbd508 TCGA-A2-A0CX-01A-11D-A10Y-09 BRCA fe96b832-cb86-4499-948a-5124a43d5c95 TCGA-A2-A0CT-01A-31W-A071-09 BRCA 2b412ad3 abda-4c78-8f68-59dbcc80031e TCGA-A2-A0CT-01A-31W-A071-09 BRCA a9aa68af-15fe-4ac0-987f-8ai49b85c231 TCGA-A2-A0CV-01A-31D-A10Y-09 BRCA da4f0f85-b16f-40fa-95e-524d70d7ac4d TCGA-A2-A0CW-01A-21D-A10Y-09 BRCA da4f0f85-b16f-40fa-95e-524d70d7ac4d TCGA-A2-A0CW-01A-21W-A050-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-A0CW-01A-11W-A050-09 BRCA 3120d0fe-anal-40f1-b2c1-7707093ac15 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 3762809c-15c9-485e-ad7a-c128a27750e9 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 3762809c-15c9-485e-ad7a-c128a27750e9 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 36656575-69c7-4745-889d-ca0568cb5559 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 8183420c-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D1-01A-11W-A050-09 BRCA f3accede-1716-d444-bad4-5127a9ebd675 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 61866-b2d4-965-b2d4-7ee7c68124f48 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 6266-b7660-47ae-b86e-652e99fa69ca TCGA-A2-A0EW-01A-11W-A050-09 BRCA 8269eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EW-01A-11D-A10Y-09 BRCA 31c4187e-9bf6-c423-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-11D-A10Y-09 BRCA 31c4187e-9bf6-c433-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 31c4187e-9bf6-c433-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 31c4187e-9bf6-c433-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 31c4187e-9bf6-da3-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 31c4187e-9bf6-da3-8cbb-10x1c0184331 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 343bf4f-23ba-4fc7-9503-1ad243d74225 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 343bf4f-23ba-4fc7-9503-1ad243d74225 TCGA-A2-A0EW-01A-2W-A050-09 BRCA 343bf4f-23ba-4fc7-9503-1ad243d74225 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 364696-4674-676-4678-698-6984049 TCGA-A2-A0EW-01A-2D-A099-09 BRCA 364696-4674-6698-862-6572e988047	TCGA-A2-A0CM-01A-31W-A050-09	BRCA	fe8023d4-5476-4c58-bf70-cbf65cdd4327
TCGA-A2-AOCS-01A-11D-A10Y-09 BRCA fe96b832-cb86-4499-948a-5124a43d5e95 TCGA-A2-AOCT-01A-31W-A071-09 BRCA 2b412ad8-abda-4cf8-8f68-59dbce80031e TCGA-A2-AOCV-01A-12W-A050-09 BRCA 39aa68af-f5fe-4ac0-987f-8af49b85c231 TCGA-A2-AOCV-01A-31D-A10Y-09 BRCA Sd1dead5-d95-42d3-a703-4c38ad6e8157 TCGA-A2-AOCW-01A-21D-A10Y-09 BRCA Sd1dead5-d95-42d3-a703-4c38ad6e8157 TCGA-A2-AOCW-01A-21D-A10Y-09 BRCA da4f0f85-b16f-40fa-95ec-524d70d7ac4d TCGA-A2-AOCW-01A-21D-A10Y-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-AOCZ-01A-11W-A019-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-AOD-01A-11W-A050-09 BRCA 3f20d0fe-aaa1-40f1-b2c1-7070f93acf5 TCGA-A2-AODD-01A-11W-A050-09 BRCA 3f20d0fe-aaa1-40f1-b2c1-7070f93acf5 TCGA-A2-AOD1-01A-11W-A050-09 BRCA 3f22080fe-3eaa1-40f1-b2c1-7070f93acf5 TCGA-A2-AOD2-01A-21W-A050-09 BRCA 3f320a-f15e9-485s-ad7a-c128427750e9 TCGA-A2-AOD3-01A-11D-A10Y-09 BRCA 8183420a-7f44-4024-b3db-653ad293988 TCGA-A2-AOD4-01A-11W-A050-09 BRCA BRCA Brack Brac	TCGA-A2-A0CP-01A-11W-A050-09	BRCA	a776e274-fe9f-49a9-83ab-95ca6819c96b
TCGA-A2-A0CT-01A-31W-A071-09 BRCA 2b412ad8-abda-4cf8-8f68-59dbce80031e TCGA-A2-A0CU-01A-12W-A050-09 BRCA a9aa68af-f5fe-4ac0-987f-8af49b85c231 TCGA-A2-A0CU-01A-21D-A10Y-09 BRCA 5d1dead5-d9a5-42d3-a703-4c38ad6c8f57 TCGA-A2-A0CW-01A-21D-A10Y-09 BRCA 5d1dead5-d9a5-42d3-a703-4c38ad6c8f57 TCGA-A2-A0CW-01A-21W-A019-09 BRCA 4d4f0f85-b16f-d0fa-95e-652d470d7ac4d TCGA-A2-A0CZ-01A-11W-A019-09 BRCA 95d5c606-367a-4db5-b663-deca3f42c2a2 TCGA-A2-A0DZ-01A-11W-A019-09 BRCA 3f2cd6fe-aaa1-40f1-b62-1-7070f93acf5 TCGA-A2-A0DJ-01A-11W-A019-09 BRCA 3f2cd6fe-aaa1-40f1-b62-1-7070f93acf5 TCGA-A2-A0DJ-01A-11W-A050-09 BRCA 05656575-69e7-4745-a89d-ca0568eb5559 TCGA-A2-A0DJ-01A-11W-A050-09 BRCA 3faccede-1716-4d44-bad4-bad4-5427-99ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 12362ad7-866-4e7a-9ec-8a0368df8896 TCGA-A2-A0EW-01A-11D-A099-09 BRCA 8e2f9eb7-0660-47a-9ec-8a0368df8896 TCGA-A2-A0EW-01A-11D-A090-09 BRCA 31ed187e-9bfe-4ca3-8eb-1-0c1e0184331 TCGA-A2-A0EW-01A-11D-A090-09 BRCA 3led187e-9bfe-4ca3-8eb-1-0c1e0184331 TCGA-A2-A0EW-01A-11D-A090-09 B	TCGA-A2-A0CQ-01A-21W-A050-09	BRCA	fa0d7183-8757-4f95-87b2-2366a1dbd508
TCGA-A2-AOCU-01A-12W-A050-09 BRCA a9aa68af-15fe-4ac0-987f-8af49b85c231 TCGA-A2-AOCW-01A-21D-A10Y-09 BRCA 5d1dead5-d9a5-42d3-a703-4c38ad6e8f57 TCGA-A2-AOCW-01A-21D-A10Y-09 BRCA da4f0f85-b16f-40fa-95e6-524d70d7ac4d TCGA-A2-AOCX-01A-21W-A019-09 BRCA 975ad5r6-3561-41a0b-959a-68da470816c7 TCGA-A2-AOCX-01A-11W-A019-09 BRCA 975d5c606-36fa-adea3i42e2a2 TCGA-A2-AODD-01A-11W-A019-09 BRCA 3f20d0fa-aaa1-40f1-b2e1-f8070f93aef5 TCGA-A2-AODD-01A-11W-A050-09 BRCA a762809e-15c9-485e-ad7a-ef28427750e9 TCGA-A2-AODD-01A-11W-A050-09 BRCA a762809e-15c9-485e-ad7a-ef28427750e9 TCGA-A2-AODD-01A-11W-A050-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-AOD3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-AOD4-01A-11W-A050-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-AOD4-01A-11W-A050-09 BRCA 61866-89ed4-965-b247-ee768124f8 TCGA-A2-AOEM-01A-11W-A050-09 BRCA 82e29b7-6660-47ae-b66-652e99fa69ca TCGA-A2-AOEM-01A-11W-A050-09 BRCA 82e29b7-6660-47ae-b66-652e99fa69ca TCGA-A2-AOEW-01A-11W-A050-09 BRCA 82e29b7-6660-47ae-b66-652e99fa69ca TCGA-A2-AOEW-01A-11W-A050-09 BRCA 2c449ea9-e3ff-4726-8566-5933e2b7056d TCGA-A2-AOEW-01A-11D-A10Y-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10e1e0184331 TCGA-A2-AOEW-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-AOEW-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-AOEW-01A-11W-A050-09 BRCA 9433bf47-23ba-4fe7-9503-1ad243d74225 TCGA-A2-AOEW-01A-21W-A050-09 BRCA 9433bf47-23ba-4fe7-9503-1ad243d74225 TCGA-A2-AOEW-01A-11W-A050-09 BRCA 9433bf47-23ba-4fe7-9503-1ad243d74225 TCGA-A2-AOEW-01A-11W-A050-09 BRCA 9433bf47-23ba-4fe7-954a-e2f58e1d5bf TCGA-A2-AOEW-01A-11D-A099-09 BRCA 6469f44-f64-4afc-a5a-c5f7689d1d43 TCGA-A2-AOEW-01A-11D-A099-09 BRCA 6469f44-f64-4afc-a5a-c5f7689d1d43 TCGA-A2-AOEW-01A-11D-A099-09 BRCA 6469f44-f64-4afc-a5a-c5f7689d1d43 TCGA-A2-AOEW-01A-11D-A099-09 BRCA 6469f44-f64-4afc-a5a-c5f7689d1d43	TCGA-A2-A0CS-01A-11D-A10Y-09	BRCA	fe96b832-cb86-4499-948a-5124a43d5c95
TCGA-A2-A0CV-01A-31D-A10Y-09 BRCA 5d1dead5-d9a5-42d3-a703-4c38ad6e8f57 TCGA-A2-A0CW-01A-21D-A10Y-09 BRCA da4f0f85-b16f-40fa-95c6-524d70d7ac4d TCGA-A2-A0CX-01A-21W-A019-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-A0CX-01A-11W-A050-09 BRCA 95d5c606-367a-46b5-b663-deca3f42c2a2 TCGA-A2-A0D0-01A-11W-A050-09 BRCA 3f2c0dfe-aa1-40f1-b2c1-7f670f93acf5 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 3f2c0dfe-aa1-40f1-b2c1-7f670f93acf5 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 05656575-69c7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6653ad293988 TCGA-A2-A0D4-01A-11W-A050-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA d01c6b8-9edd-4965-b247-ee768124f48 TCGA-A2-A0EM-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 2e449ea9-c3ff-472-6e6-8a0a68df8896 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 31ed187e-9bf6-4ca3-8ebe-65c-59a3c2b7056d TCGA-A2-A0EM-01A-11W-A050-09 BRCA 31ed187e-9bf6-4ca3-8eb-10c1e0184331 TCGA-A2-A0EW-01A-11W-A050-09 BRCA </td <td>TCGA-A2-A0CT-01A-31W-A071-09</td> <td>BRCA</td> <td>2b412ad8-abda-4cf8-8f68-59dbce80031e</td>	TCGA-A2-A0CT-01A-31W-A071-09	BRCA	2b412ad8-abda-4cf8-8f68-59dbce80031e
TCGA-A2-A0CW-01A-21D-A10Y-09 BRCA da4f0f85-b16f-40fa-95c6-524d70d7ac4d TCGA-A2-A0CX-01A-21W-A019-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-A0D0-01A-11W-A050-09 BRCA 95d5c606-367a-46b5-b663-dcea3f42c2a2 TCGA-A2-A0D1-01A-11W-A050-09 BRCA 3f20d0fc-aaa1-40f1-b2c1-f707093acf5 TCGA-A2-A0D1-01A-11W-A050-09 BRCA a762809c-15c9-485e-ad7a-ef28427750e9 TCGA-A2-A0D2-01A-21W-A050-09 BRCA 05656575-69e7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 183a42ce-7444-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A019-09 BRCA 18accede-1716-d444-bad4-5427z9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 22449ea9-c3ff-4726-8566-5933c2b7056d TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EQ-01A-11D-A10Y-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EQ-01A-11D-A10Y-09 BRCA 4d42c62-5c2d-49f5-856e-793ace188044 TCGA-A2-A0EQ-01A-11W-A050-09 BRCA	TCGA-A2-A0CU-01A-12W-A050-09	BRCA	a9aa68af-f5fe-4ac0-987f-8af49b85c231
TCGA-A2-A0CX-01A-21W-A019-09 BRCA 975adb76-3561-41a0-959a-68da470816c7 TCGA-A2-A0CZ-01A-11W-A050-09 BRCA 95d5c606-367a-46b5-b663-dcea3f42c2a2 TCGA-A2-A0D0-01A-11W-A019-09 BRCA 3f20d0fc-aaa1-40f1-b2c1-7f070f93acf5 TCGA-A2-A0D12-01A-21W-A050-09 BRCA a762809c-15c9-485e-ad7a-ef28427750e9 TCGA-A2-A0D2-01A-21W-A050-09 BRCA 05656575-69c7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A019-09 BRCA 13accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 12362ad7-6866-4c7a-e9ce-8a0a68df8896 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 12362ad7-6866-4c7a-e9ce-8a0a68df8896 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 8e219eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EW-01A-11W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 4fb40023-4ad-4cf3-8c3-8cb-10c1e0184331 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 6d442c62-5c2d-49f5-8b6-72beef88044d TCGA-A2-A0EW-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fc7-3e3-5c10ddcbc0fb TCGA-A2-A0EW-01A-21W-A050-09 BR	TCGA-A2-A0CV-01A-31D-A10Y-09	BRCA	5d1dead5-d9a5-42d3-a703-4c38ad6e8f57
TCGA-A2-A0CZ-01A-11W-A050-09 BRCA 95d5c606-367a-46b5-b663-dcea3i42e2a2 TCGA-A2-A0D0-01A-11W-A019-09 BRCA 3f20d0fe-aaa1-40f1-b2c1-7f070f93aef5 TCGA-A2-A0D1-01A-11W-A050-09 BRCA a762809c-15c9-485e-ad7a-ef28427750e9 TCGA-A2-A0D2-01A-21W-A050-09 BRCA 05656575-69c7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420c-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0EM-01A-11W-A019-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee768124f48 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ac-986e-680268df8896 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 2e2449ea9-c3ff-4726-856e-52e99fa69ca TCGA-A2-A0EC-01A-11W-A050-09 BRCA 31ed187c-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EC-01A-11W-A050-09 BRCA 31ed187c-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EC-01A-21W-A050-09 BRCA 4fb40023-4adc-4c7d-ac73-5c10ddcbc0fb TCGA-A2-A0EC-01A-11W-A050-09 BRCA 4d30a8f-903f-428-a63d-59625fe858a9 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9433b4f-23ba-4fe70-950-1ad24374225 TCGA-A2-A0EW-01A-11W-A050-09 BRCA <td>TCGA-A2-A0CW-01A-21D-A10Y-09</td> <td>BRCA</td> <td>da4f0f85-b16f-40fa-95c6-524d70d7ac4d</td>	TCGA-A2-A0CW-01A-21D-A10Y-09	BRCA	da4f0f85-b16f-40fa-95c6-524d70d7ac4d
TCGA-A2-A0D0-01A-11W-A019-09 BRCA 3f20d0fe-aaa1-40f1-b2c1-7f070f93aef5 TCGA-A2-A0D1-01A-11W-A050-09 BRCA a762809c-15c9-485e-ad7a-ef28427750e9 TCGA-A2-A0D2-01A-21W-A050-09 BRCA 05656575-69c7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A019-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 12362ad7-6866-427a-9ec6-8a0a68df8896 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 3led187e-9bfe-4ca3-8bcb-10c1e0184331 TCGA-A2-A0ER-01A-21W-A050-09 BRCA 4d42c62-5c2d-49f5-856e-72beef8804dd TCGA-A2-A0EU-01A-22W-A071-09 BRCA d630488f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9405a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-11W-A050-09 BRCA	TCGA-A2-A0CX-01A-21W-A019-09	BRCA	975adb76-3561-41a0-959a-68da470816c7
TCGA-A2-A0D1-01A-11W-A0S0-09 BRCA a762809c-15c9-485e-ad7a-ef28427750e9 TCGA-A2-A0D2-01A-21W-A0S0-09 BRCA 05656575-69e7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A019-09 BRCA 63accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EN-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 3led187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ER-01A-21W-A050-09 BRCA 64442c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ER-01A-11D-A01C-09 BRCA 6442c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad24374225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad24374225 TCGA-A2-A0EW-01A-11W-A050-09 BRCA </td <td>TCGA-A2-A0CZ-01A-11W-A050-09</td> <td>BRCA</td> <td>95d5c606-367a-46b5-b663-dcea3f42e2a2</td>	TCGA-A2-A0CZ-01A-11W-A050-09	BRCA	95d5c606-367a-46b5-b663-dcea3f42e2a2
TCGA-A2-A0D2-01A-21W-A050-09 BRCA 05656575-69e7-4745-a89d-ca0568eb5559 TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A050-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EW-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0EQ-01A-11D-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0EE-01A-21D-A10Y-09 BRCA 64d42c2-5c2d-49f5-856e-72beef8804dd TCGA-A2-A0EU-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EY-01A-11W-A050-09 BRCA 3038f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA	TCGA-A2-A0D0-01A-11W-A019-09	BRCA	3f20d0fe-aaa1-40f1-b2c1-7f070f93aef5
TCGA-A2-A0D3-01A-11D-A10Y-09 BRCA 8183420e-7f44-4024-b3db-6b53ad293988 TCGA-A2-A0D4-01A-11W-A019-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EN-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ER-01A-21D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0EU-01A-22W-A071-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddebc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA dd669f44-f64d-afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA dd669f44-f64d-afc-a5ac-5f7769d1db43 TCGA-A2-A0SW-01A-11D-A099-09 BRCA<	TCGA-A2-A0D1-01A-11W-A050-09	BRCA	a762809c-15c9-485e-ad7a-ef28427750e9
TCGA-A2-A0D4-01A-11W-A019-09 BRCA f3accede-1716-4d44-bad4-5427a9ebd675 TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EN-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-11W-A050-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 38cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0EY-01A-11W-A050-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA d63206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRC	TCGA-A2-A0D2-01A-21W-A050-09	BRCA	05656575-69e7-4745-a89d-ca0568eb5559
TCGA-A2-A0EM-01A-11W-A050-09 BRCA 0e01c6b8-9edd-4965-b247-ee7e68124f48 TCGA-A2-A0EN-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0EU-01A-31D-A045-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-22W-A071-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EV-01A-21D-A10Y-09 BRCA 39308f50c-1320-4c45-acc7-38f43b6f9336 TCGA-A2-A0EY-01A-11W-A050-09 BRCA 38cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA 6d3206e6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SV-01A-11D-A099-09 BR	TCGA-A2-A0D3-01A-11D-A10Y-09	BRCA	8183420e-7f44-4024-b3db-6b53ad293988
TCGA-A2-A0EN-01A-13D-A099-09 BRCA 12362ad7-6866-4e7a-9ec6-8a0a68df8896 TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ES-01A-31D-A045-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ET-01A-31D-A045-09 BRCA de30da8f-903f-428e-a63d-59625fe858a9 TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fe858a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 3045a04e-4f7b-4f9a-3733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA 36d669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SY-01A-11D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6ca206f-1458-417p-94a-e2f58ed163bf TCGA-A2-A0SW-01A-11D-A099-09 BRCA<	TCGA-A2-A0D4-01A-11W-A019-09	BRCA	f3accede-1716-4d44-bad4-5427a9ebd675
TCGA-A2-A0EO-01A-11W-A050-09 BRCA 8e2f9eb7-0660-47ae-b86e-652e99fa69ca TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ET-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-ac73-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA dc30da8f-903f-428e-a63d-59625fc85a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EX-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9c7f-45d079893176 TCGA-A2-A0SY-01A-12D-A099-09 BRCA dc669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ca2f20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-AOSV-01A-11D-A099-09 BRCA 554bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-AOSY-01A-31D-A099-09 BRCA	TCGA-A2-A0EM-01A-11W-A050-09	BRCA	0e01c6b8-9edd-4965-b247-ee7e68124f48
TCGA-A2-A0EQ-01A-11W-A050-09 BRCA 2c449ea9-c3ff-4726-8566-5933e2b7056d TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ES-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddebc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EU-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA 3045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EX-01A-21W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0EY-01A-11D-A059-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SW-01A-11D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SW-01A-31D-A099-09 BRCA a6a9c0b-c14b-4141-b48e-cc2c6b89ab73 TCGA-A2-A0T4-01A-21D-A099-09 BRCA	TCGA-A2-A0EN-01A-13D-A099-09	BRCA	12362ad7-6866-4e7a-9ec6-8a0a68df8896
TCGA-A2-A0ER-01A-21W-A050-09 BRCA 31ed187e-9bfe-4ca3-8cbb-10c1e0184331 TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ET-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-aer3-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SX-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0T4-01A-31D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA	TCGA-A2-A0EO-01A-11W-A050-09	BRCA	8e2f9eb7-0660-47ae-b86e-652e99fa69ca
TCGA-A2-A0ES-01A-11D-A10Y-09 BRCA 64d42c62-5c2d-49f5-856e-72beef88044d TCGA-A2-A0ET-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9336 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SX-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A10Y-09 BRCA	TCGA-A2-A0EQ-01A-11W-A050-09	BRCA	2c449ea9-c3ff-4726-8566-5933e2b7056d
TCGA-A2-A0EU-01A-31D-A045-09 BRCA f7b40023-4adc-4c7d-ae73-5c10ddcbc0fb TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EW-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SW-01A-11D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc26b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 7918143-dbce-45b3-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T4-01A-31D-A099-09 BRCA<	TCGA-A2-A0ER-01A-21W-A050-09	BRCA	31ed187e-9bfe-4ca3-8cbb-10c1e0184331
TCGA-A2-A0EU-01A-22W-A071-09 BRCA de30da8f-903f-428e-a63d-59625fc858a9 TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EX-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0SY-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SX-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc2c6b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 9515373a-d982-45fa-b8f9-363f9ba8649f TCGA-A2-A0T2-01A-11W-A097-09 BRCA c7918143-dbce-45b3-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T6-01A-11D-A099-09 BRC	TCGA-A2-A0ES-01A-11D-A10Y-09	BRCA	64d42c62-5c2d-49f5-856e-72beef88044d
TCGA-A2-A0EV-01A-11W-A050-09 BRCA 9433bf4f-23ba-4fe7-9503-1ad243d74225 TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EX-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SW-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc2c6b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 7918143-dbce-45b3-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T4-01A-31D-A099-09 BRCA 0f1b1fda-4956-498a-b8ff-e98b5d64e509 TCGA-A2-A0T6-01A-11D-A099-09 BRCA	TCGA-A2-A0ET-01A-31D-A045-09	BRCA	f7b40023-4adc-4c7d-ae73-5c10ddcbc0fb
TCGA-A2-A0EW-01A-21D-A10Y-09 BRCA a045a04e-4f7b-4f9a-a733-47ad24475496 TCGA-A2-A0EX-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SW-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc2c6b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 9515373a-d982-45fa-b8f9-363f9ba8649f TCGA-A2-A0T2-01A-11W-A097-09 BRCA c7918143-dbce-45ba-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T6-01A-11D-A099-09 BRCA 0f1b1fda-4956-498a-b8ff-e98b5d64e509 TCGA-A2-A0T6-01A-11D-A099-09 BRC	TCGA-A2-A0EU-01A-22W-A071-09	BRCA	de30da8f-903f-428e-a63d-59625fc858a9
TCGA-A2-A0EX-01A-21W-A050-09 BRCA 9308f50c-1320-4c45-acc7-38f43b6f9a36 TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SX-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc2c6b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 9515373a-d982-45fa-b8f9-363f9ba8649f TCGA-A2-A0T2-01A-11W-A097-09 BRCA c7918143-dbcc-45b3-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T4-01A-31D-A099-09 BRCA 0f1b1fda-4956-498a-b8ff-e98b5d64e509 TCGA-A2-A0T6-01A-11D-A099-09 BRCA e4dcb280-c309-4ebb-a58d-e6389a0306ee	TCGA-A2-A0EV-01A-11W-A050-09	BRCA	9433bf4f-23ba-4fe7-9503-1ad243d74225
TCGA-A2-A0EY-01A-11W-A050-09 BRCA a8cde596-e3f5-4b20-9e7f-45d079893176 TCGA-A2-A0ST-01A-12D-A099-09 BRCA dd669f44-f64d-4afc-a5ac-5f7769d1db43 TCGA-A2-A0SU-01A-11D-A099-09 BRCA 6ceaf20f-1458-4f7f-954a-e2f58ed163bf TCGA-A2-A0SV-01A-11D-A099-09 BRCA 6d3206c6-0ca8-4b2b-a160-b1719217f9c7 TCGA-A2-A0SW-01A-11D-A099-09 BRCA 7fbd2807-a5bb-4030-a299-524ec3ab4543 TCGA-A2-A0SW-01A-12D-A099-09 BRCA b54bc31e-bdcc-4ad5-998e-5a9c542f83bb TCGA-A2-A0SY-01A-31D-A099-09 BRCA efaa9c0b-c14b-4141-b48c-cc2c6b89ab73 TCGA-A2-A0T0-01A-22D-A099-09 BRCA 3c107ce4-a6ac-469b-b1c0-cd86674b5766 TCGA-A2-A0T1-01A-21D-A099-09 BRCA 9515373a-d982-45fa-b8f9-363f9ba8649f TCGA-A2-A0T2-01A-11W-A097-09 BRCA c7918143-dbce-45b3-8d24-2993a9e2b7f4 TCGA-A2-A0T3-01A-21D-A10Y-09 BRCA 0ca029bb-3b3a-48ec-8ade-5591e8e8629f TCGA-A2-A0T4-01A-31D-A099-09 BRCA 0f1b1fda-4956-498a-b8ff-e98b5d64e509 TCGA-A2-A0T6-01A-11D-A099-09 BRCA e4dcb280-c309-4ebb-a58d-e6389a0306ee	TCGA-A2-A0EW-01A-21D-A10Y-09	BRCA	a045a04e-4f7b-4f9a-a733-47ad24475496
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TCGA-A2-A259-01A-11D-A16D-09	BRCA	93febb0a-587c-47f2-9a59-117f7aa475c5
TCGA-A2-A25A-01A-12D-A16D-09	BRCA	5739a7e1-7fa3-434c-b1c3-c0a9e570c858
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TCGA-A7-A0CG-01A-11W-A019-09	BRCA	351275c7-70ca-4ddc-be76-a6ff4dc7655e
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TCGA-A7-A13G-01A-11D-A13L-09	BRCA	ef847b83-eb88-435b-bcfd-4b51d4dfa5fe
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TCGA-A7-A26F-01A-21D-A167-09	BRCA	fc73db72-d0ac-48d0-b809-2f7540482ec5
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TCGA-A8-A06Q-01A-11W-A050-09	BRCA	473d5422-978a-48be-be32-2b7516d6d2d5
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TCGA-A8-A06X-01A-21W-A019-09	BRCA	dc306402-3a55-4996-b786-f3f738f13dd3
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TCGA-AO-A03N-01B-11D-A10M-09 BRCA ef5987f1-46ac-430a-b94a-49afa0e286d4
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TCGA-AO-A03R-01A-21W-A050-09 BRCA 6d2dc4e3-f1ed-4ef0-ae83-e09c87756d56
TCGA-AO-A03T-01A-21W-A050-09 BRCA cbea866d-da66-4f7c-994b-c1ec35aa2d4b

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	BRCA	d88c365f-366a-49d5-9860-b930aab3eb1b
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TCGA-AR-A24V-01A-21D-A167-09 BRCA bb77af66-bb8f-4590-9be8-5f729373c555 TCGA-AR-A24W-01A-11D-A17G-09 BRCA 454e7cd4-8424-4cad-8fbb-f69affa5d1bf TCGA-AR-A24X-01A-11D-A167-09 BRCA 53d55f5a-df86-44d7-a3a2-2decc2557b7b TCGA-AR-A24X-01A-11D-A167-09 BRCA c11f2060-d3fb-4c3d-8058-b8cce44af519 TCGA-AR-A250-01A-31D-A167-09 BRCA f7d9a372-fcd1-4462-9e0b-7eb46ddb68fd TCGA-AR-A250-10A-12D-A167-09 BRCA 68b4de6d-352d-44e8-911a-f454f128fc78 TCGA-AR-A252-01A-11D-A167-09 BRCA 6800d9b3-32a1-48eb-840b-9a3bec9d1f6e TCGA-AR-A252-01A-11D-A167-09 BRCA 6800d9b3-32a1-48eb-840b-9a3bec9d1f6e TCGA-AR-A255-01A-11D-A167-09 BRCA fc2bdac0-832e-4268-bd8f-5dcfffda1979 TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A255-01A-11D-A167-09 BRCA 6804d948-82ce-4dbc7f3776de TCGA-B6-A0I2-01A-11W-A050-09 BRCA a9cac7c8-a62b-46ad-98b-82ce0b5iddf00 TCGA-B6-A0I3-01A-11W-A100-09 BRCA a9cac7c8-a62b-46ad-98b-82ce0b5iddf00 TCGA-B6-A0I3-01A-11W-A100-09 BRCA a876398c-5bid-44df-a360-5fc2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA a876398c-5bid-44f1b-8b45-b617c2861ce7 TCGA-B6-A0I8-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-60737cf3acea TCGA-B6-A0I8-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-60737cf3acea TCGA-B6-A0I8-01A-11W-A050-09 BRCA f180d5cd-7acd-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA f180d5cd-7acd-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA f180d5cd-7acd-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA d23f4730-0a18-423b-a2cd-f1a4231c2b53 TCGA-B6-A0IB-01A-11W-A050-09 BRCA d23f4730-0a18-423b-a2cd-f1a4231c2b53 TCGA-B6-A0IB-01A-11W-A050-09 BRCA d23f4730-0a18-423b-a2cd-f1a4231c2b53 TCGA-B6-A0IB-01A-11W-A050-09 BRCA d23f4730-0a18-423b-a2cd-f1a4231c2b53 TCGA-B6-A0IB-01A-11W-A050-09 BRCA d25f16a-d50a-d60a-d50a-d50a-d50a-d50a-d50a-d50a-d50a-d5	TCGA-AR-A24T-01A-11D-A167-09	BRCA	09991de6-2e8e-476f-987b-98d9a85dac7d
TCGA-AR-A24W-01A-11D-A167-09 BRCA 45467cd4-8424-4cad-8fbb-f69affa5d1bf TCGA-AR-A24X-01A-11D-A167-09 BRCA 53d55f5a-df86-44d7-a3a2-2dccc2557b7b TCGA-AR-A24Z-01A-11D-A167-09 BRCA 61f2060-d3fb-4c3d-8058-b8ccc44af519 TCGA-AR-A250-01A-31D-A167-09 BRCA 68b4de6d-352d-44e8-911a-f454ff28fc78 TCGA-AR-A251-01A-12D-A167-09 BRCA 68b4de6d-352d-44e8-911a-f454ff28fc78 TCGA-AR-A252-01A-11D-A167-09 BRCA 68b4de6d-352d-44e8-911a-f454ff28fc78 TCGA-AR-A252-01A-11D-A167-09 BRCA 68c0d9b3-32a1-48eb-840b-9a3bec9d1f6e TCGA-AR-A255-01A-11D-A167-09 BRCA 62bdac0-832e-4268-bd8f-5dcfffdal979 TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0048-4f1c-a142-651605158bf3 TCGA-AR-A255-01A-11D-A167-09 BRCA 62bdac0-832e-4268-bd8f-5dcfffdal979 TCGA-B6-A013-01A-11W-A050-09 BRCA 62a43434b-197e-48ac-ac2e-46bc7f3776de 6443434b-197e-48ac-ac2e-46bc7f3776de 7CGA-B6-A0I3-01A-11W-A100-09 BRCA 61139266-fade-4d27-ac67-60870c666295 TCGA-B6-A0I3-01A-11W-A050-09 BRCA 61139266-fade-4d27-ac67-60870c666295 TCGA-B6-A0I8-01A-11W-A050-09 BRCA 62291482-9bb-4f8f-a56b-c0737cf3acea TCGA-B6-A0I3-01A-11W-A050-09 BRCA 62291482-9bb-4f8f-a56b-c0737cf3acea TCGA-B6-A0I3-01A-11W-A050-09 BRCA 62291482-9bb-4f8f-a56b-c0737cf3acea TCGA-B6-A0I3-01A-11W-A050-09 BRCA 623fd730-0a18-4c3b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 623fd730-0a18-4d2b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 623fd730-0a18-4c3b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 623fd730-0a18-4c3b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 623fd730-0a18-4c3b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 623fd730-0a18-4c3b-ac2d-f1a4231c2b53 TCGA-B6-A0II-01A-11W-A050-09 BRCA 6246488b-7404-4cb1-aref-ce94c32b942 TCGA-B6-A0II-01A-11W-A050-09 BRCA 639ddb-6301-400e-ac8e-197ea2efc75 TCGA-B6-A0II-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0II-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0II-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0II-01A-11	TCGA-AR-A24U-01A-11D-A167-09	BRCA	567cdc6c-df03-4642-8cbc-a269769ce1a1
TCGA-AR-A24X-01A-11D-A167-09 BRCA 53d55f5a-df86-44d7-a3a2-2dece2557b7b TCGA-AR-A24Z-01A-11D-A167-09 BRCA c11f2060-d3fb-4e3d-8058-b8cce44af519 TCGA-AR-A250-01A-31D-A167-09 BRCA f7d9a372-fcd1-4462-9e0b-7eb46ddb68fd TCGA-AR-A251-01A-12D-A167-09 BRCA 68b4de6d-352d-44e8-911a-fd541f28fc78 TCGA-AR-A252-01A-11D-A167-09 BRCA e800d9b3-32a1-48eb-840b-9a3bec9d1f6e TCGA-AR-A254-01A-21D-A167-09 BRCA fe2bdac0-832e-4268-bd8f-5dcfffda1979 TCGA-AR-A255-01A-11D-A167-09 BRCA s05f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A256-01A-11D-A167-09 BRCA a9cae7c8-a62b-46ad-398b-82e6b5fddf00 TCGA-BC-A012-01A-11W-A050-09 BRCA d9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-BC-A015-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-BC-A016-01A-11W-A050-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-BC-A016-01A-11W-A050-09 BRCA d2291482-9bbb-4f8-a65b-c0737cf3acea TCGA-BC-A010-01A-11W-A050-09 BRCA f7e5ada6-Kf3-4765-a874-5ee9d258ad6a TCGA-BC-A01A-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-BC-A01B-01A-11W-A050-09 BRCA<	TCGA-AR-A24V-01A-21D-A167-09	BRCA	bb77af66-bb8f-4590-9be8-5f729373c555
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TCGA-AR-A251-01A-12D-A167-09 BRCA 68b4de6d-352d-44e8-911a-f4541f28fc78 TCGA-AR-A252-01A-11D-A167-09 BRCA e800d9b3-32a1-48eb-840b-9a3bec9d1f6e TCGA-AR-A254-01A-21D-A167-09 BRCA fe2bdac0-832e-4268-bd8f-5defffda1979 TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A256-01A-11D-A167-09 BRCA ea43434b-197e-48ac-ae2e-46bc7f3776de TCGA-B6-A012-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A015-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A016-01A-11W-A050-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A018-01A-11W-A050-09 BRCA d280b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A018-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A018-01A-11W-A050-09 BRCA f7e5ada6-8f3-4765-a874-5ee9d258ad6a TCGA-B6-A01A-01A-11W-A050-09 BRCA ff80d5cd-7acd-499f-a472-153cc40f65de TCGA-B6-A01B-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IG-01A-11W-A050-09 BRCA d283f50-5031-4b08-ba3a-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA<	TCGA-AR-A24Z-01A-11D-A167-09	BRCA	c11f2060-d3fb-4e3d-8058-b8cce44af519
TCGA-AR-A252-01A-11D-A167-09 BRCA e800d9b3-32a1-48eb-840b-9a3bee9d1f6e TCGA-AR-A254-01A-21D-A167-09 BRCA fe2bdac0-832e-4268-bd8f-5dcfffdal979 TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A256-01A-11D-A167-09 BRCA ea43434b-197e-48ac-ae2e-46bc7f3776de TCGA-B6-A012-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A016-01A-11D-A128-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A016-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A016-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A018-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A01B-01A-11W-A050-09 BRCA f180d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A01B-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IG-01A-11W-A050-09 BRCA deb04519-d928-dfd3-bae2-84585aa4f022 TCGA-B6-A0IG-01A-11W-A050-09 BRCA e8046519-d928-dfd3-bae2-84585aa4f022 TCGA-B6-A0IG-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IM-01A-11W-A050-09 BRC	TCGA-AR-A250-01A-31D-A167-09	BRCA	f7d9a372-fcd1-4462-9e0b-7eb46ddb68fd
TCGA-AR-A254-01A-21D-A167-09 BRCA fe2bdac0-832e-4268-bd8f-5defffda1979 TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A256-01A-11D-A167-09 BRCA ea43434b-197e-48ac-ae2e-46bc7f3776de TCGA-B6-A0I2-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A0I6-01A-11D-A128-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A0I6-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fc2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IG-01A-11W-A050-09 BRCA 4cb39f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA 4a4488b9-74d9-4eb1-a7ef-c884c32db942 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IM-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IM-01A-11W-A050-09 BRC	TCGA-AR-A251-01A-12D-A167-09	BRCA	68b4de6d-352d-44e8-911a-f4541f28fc78
TCGA-AR-A255-01A-11D-A167-09 BRCA 505f1398-0bd8-4f1c-a142-651605158bf3 TCGA-AR-A256-01A-11D-A167-09 BRCA ea43434b-197e-48ac-ae2e-46bc7f3776de TCGA-B6-A0I2-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A0I5-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A0I6-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617ce861ce7 TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0IA-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IG-01A-11W-A050-09 BRCA e8046519-d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IG-01A-11W-A050-09 BRCA e8046519-d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efc75 TCGA-B6-A0IK-01A-12W-A071-09 BRCA c5b1f426-562e-44e4-bcce-ce2fff6d96c8 TCGA-B6-A0IN-01A-11W-A050-09 BRC	TCGA-AR-A252-01A-11D-A167-09	BRCA	e800d9b3-32a1-48eb-840b-9a3bec9d1f6e
TCGA-AR-A256-01A-11D-A167-09 BRCA ea43434b-197e-48ac-ae2e-46bc7f3776de TCGA-B6-A0I2-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A0I5-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A0I6-01A-11W-A050-09 BRCA a876398c-5b1d-444f-a360-5fc2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A0I8-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0I9-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IB-01A-11W-A050-09 BRCA f7e3da6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IC-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IC-01A-11W-A050-09 BRCA e8046519-0d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IG-01A-11W-A050-09 BRCA 4a4488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0IL-01A-11W-A050-09 BRCA c5b1f426-562e-44e4-bce-ce2fff6969c8 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0ID-01A-11W-A050-09 BRCA	TCGA-AR-A254-01A-21D-A167-09	BRCA	fe2bdac0-832e-4268-bd8f-5dcfffda1979
TCGA-B6-A0I2-01A-11W-A050-09 BRCA a9cae7c8-a62b-46ad-a98b-82e6b5fddf00 TCGA-B6-A0I5-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A0I6-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A0I8-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0I9-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IA-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA dcb39f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA de8046519-d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA da4488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0II-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IW-01A-11W-A050-09 BRCA c99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IW-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IP-01A-11W-A050-09 BR	TCGA-AR-A255-01A-11D-A167-09	BRCA	505f1398-0bd8-4f1c-a142-651605158bf3
TCGA-B6-A0I5-01A-11W-A100-09 BRCA f1139266-fade-4d27-ac67-60870e666295 TCGA-B6-A0I6-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861cc7 TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acca TCGA-B6-A0I9-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IC-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IC-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IC-01A-11W-A050-09 BRCA d239f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA a44488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA d23fd9db-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IJ-01A-11W-A050-09 BRCA c5b1f426-562e-44e4-bcce-ce2fff6d969c8 TCGA-B6-A0IM-01A-11W-A050-09 BRCA c99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IP-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0IP-01A-11W-A050-09 BRC	TCGA-AR-A256-01A-11D-A167-09	BRCA	ea43434b-197e-48ac-ae2e-46bc7f3776de
TCGA-B6-A0I6-01A-11D-A128-09 BRCA a876398c-5b1d-444f-a360-5fe2db697480 TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0IA-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IB-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IE-01A-11W-A050-09 BRCA dcb39f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA a8046519-d928-4fd3-ba22-84585aa4f022 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA da4488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0IJ-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IJ-01A-11W-A050-09 BRCA c5b1f426-562e-44e4-bcce-ce2ff6d969c8 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e2ce198-cea3-4a54-b96b-834a70c30d2f TCGA-B6-A0ID-01A-11W-A050-09 BRCA 583964cf-84ad-4ef1-90d1-2f6bfbeb245a TCGA-B6-A0IQ-01A-11W-A071-09 BRCA	TCGA-B6-A0I2-01A-11W-A050-09	BRCA	a9cae7c8-a62b-46ad-a98b-82e6b5fddf00
TCGA-B6-A0I8-01A-11W-A050-09 BRCA ba80b13a-e20a-441b-b845-b617cc861ce7 TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0IA-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IC-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IE-01A-11W-A050-09 BRCA 4cb39f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA 48046519-d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA 4a4488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0IH-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IK-01A-12W-A071-09 BRCA c5b1f426-562e-44e4-bcce-ce2ff6d969c8 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IN-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0IP-01A-11D-A045-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0IQ-01A-11W-A050-09 BRCA 583964cf-84ad-4ef1-90d1-2f6bfbeb245a TCGA-B6-A0RG-01A-11W-A071-09 BRC	TCGA-B6-A0I5-01A-11W-A100-09	BRCA	f1139266-fade-4d27-ac67-60870e666295
TCGA-B6-A0I9-01A-11W-A050-09 BRCA d2291482-9bbb-4f8f-a65b-c0737cf3acea TCGA-B6-A0IA-01A-11W-A050-09 BRCA f7e5ada6-8f53-4765-a874-5ee9d258ad6a TCGA-B6-A0IB-01A-11W-A050-09 BRCA ff80d5cd-7aed-499f-a472-153cc40f65de TCGA-B6-A0IC-01A-11W-A050-09 BRCA f23fd730-0a18-4e3b-a2ed-f1a4231c2b53 TCGA-B6-A0IE-01A-11W-A050-09 BRCA 4cb39f50-5031-4b08-baa3-1a366ada6514 TCGA-B6-A0IG-01A-11W-A050-09 BRCA e8046519-d928-4fd3-b3e2-84585aa4f022 TCGA-B6-A0IH-01A-11D-A10Y-09 BRCA 4a4488b9-74d9-4eb1-a7ef-c894c32db942 TCGA-B6-A0IH-01A-11W-A050-09 BRCA c63f9ddb-6301-400e-a0e8-197eea2efe75 TCGA-B6-A0IK-01A-12W-A071-09 BRCA c5b1f426-562e-44e4-bcce-ce2ff6d969c8 TCGA-B6-A0IM-01A-11W-A050-09 BRCA e99a4753-10db-4823-953d-e878a90e6b01 TCGA-B6-A0IN-01A-11W-A050-09 BRCA e2c9198-cea3-4a54-b96b-834a70c30d2f TCGA-B6-A0IO-01A-11W-A050-09 BRCA 648cee86-f2e7-45a0-abf2-0ab0037e2eee TCGA-B6-A0IQ-01A-11W-A050-09 BRCA 583964cf-84ad-4ef1-90d1-2f6bfbeb245a TCGA-B6-A0IQ-01A-11W-A071-09 BRCA 583964cf-84ad-4ef1-90d1-2f6bfbeb245a TCGA-B6-A0RE-01A-11W-A071-09 BRCA	TCGA-B6-A0I6-01A-11D-A128-09	BRCA	a876398c-5b1d-444f-a360-5fe2db697480
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TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA d3c1b30c-b768-4a98-b285-5284bffa66f9 TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA d3c1b30c-aag-245Rs-bc28-eccel192a0fab TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA f0ca4831-d56d-d4bac-b304-bb43c5d2f09b TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA f75de986-bc8a-4ffe-9b35-011eec3a1446 TCGA-BH-A18I-O1A-32D-A12B-09 BRCA f75de986-bc8a-4ffe-9b35-011eec3a1446 TCGA-BH-A18I-O1A-32D-A12B-09 BRCA 883ed3e9-2681-4822-8b22-29149a027514 TCGA-BH-A18I-O1A-32D-A12B-09 BRCA 883ed3e9-2681-4822-8b22-29149a027514 TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA 3c48ed-e-b67-4432-8b12-b262a1ef949 TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA 3d466680-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18I-O1A-1ID-A12B-09 BRCA 42facac2-8140-4a9f-b416-1de89a7662fc TCGA-BH-A18I-O1A-1ID-A135-09 BRCA 6150d25-a8f4-4d9f-9da0-956855ab67d TCGA-BH-A1EV-01A-1ID-A135-09 BRCA 6150d25-a8f4-4d9f-9da0-956855ab67d TCGA-BH-A1EF-01A-1ID-A135-09 BRCA 6150d25-a8f4-4d9f-abfa-e2d063e5cd6d TCGA-BH-A1EV-01A-1ID-A135-09 BRCA 62578675-e63c-4bdf-abfa-e2d063e5cd6d TCGA-BH-A1EV-01A-1ID-A135-09 BRCA 63646648-6947-4b47-38b66e7e427 TCGA-BH-A1EV-01A-1ID-A135-09 BRCA 63646648-6947-4b46-33ea-33ea-6055d1c8bal TCGA-BH-A1EV-01A-1ID-A131-09 BRCA 63666486-4348-409-3600-5144738560279 TCGA-BH-A1EV-01A-1ID-A131-09 BRCA 63666486-4348-409-6055-6366-4364-63664		1	
TCGA-BH-A18L-01A-11D-A12B-09 BRCA f0sa4831-d56d-4bae-b304-bb43c5d2f09b TCGA-BH-A18L-01A-11D-A12B-09 BRCA fd9923db-2a27-432e-a0c6-4c44e6ee1f53 TCGA-BH-A18L-01A-32D-A12B-09 BRCA f75de986-bc8a-dffe-9b35-011eec3a1446 TCGA-BH-A18L-01A-32D-A12B-09 BRCA f75de986-bc8a-dffe-9b35-011eec3a1446 TCGA-BH-A18L-01A-32D-A12B-09 BRCA 883cd3e2-2681-4822-8b22-29149a027514 TCGA-BH-A18M-01A-11D-A12B-09 BRCA d85de1e-cbb7-4432-8112-bb262a1er9d9 TCGA-BH-A18M-01A-11D-A12B-09 BRCA d36d24a3-57e9-46be-9bcc-3e53d7c2drb7 TCGA-BH-A18M-01A-11D-A12B-09 BRCA add624a3-57e9-46be-9bcc-3e53d7c2drb7 TCGA-BH-A18M-01A-11D-A12B-09 BRCA d4f66680-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18M-01A-11D-A12B-09 BRCA d4f66e80-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18M-01A-11D-A12B-09 BRCA d4f6ce-847-4132-bdcc-aace2c027b28 TCGA-BH-A18M-01A-11D-A12B-09 BRCA de0dfcb-847-4132-bdcc-aace2c027b28 TCGA-BH-A18M-01A-11D-A12B-09 BRCA de0dfcb-847-4132-bdcc-aace2c027b28 TCGA-BH-A18M-01A-11D-A12B-09 BRCA d6150d25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EM-01A-11D-A12B-09 BRCA d150d25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EM-01A-11D-A135-09 BRCA d131381-8a11-425d-8954-980c6ec7e427 TCGA-BH-A1EM-01A-11D-A135-09 BRCA d26d613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EM-01A-11D-A135-09 BRCA d26d613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EM-01A-11D-A135-09 BRCA d37bc2a9-078a-4bc2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA d37bc2a9-078a-4bc2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA d37bc2a9-078a-4bc2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA d37bc2a9-078a-4bc2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A131-09 BRCA d57bb-85-0882-48ec-a38a-a05b5d1e8ba1 TCGA-BH-A1EW-01A-11D-A131-09 BRCA d57bb-85-3444-bc9-c677-9855329ed TCGA-BH-A1F0-01A-11D-A131-09 BRCA d57bb-85-3444-bc9-c677-985524ed d67bb-85-444-bc9-c677-9856ed TCGA-BH-A1F0-01A-11D-A131-09 BRCA d57bc9-3444-bc9-bc6-51ba6b2-26b6 TCGA-BH-A1F0-01A-11D-A131-09 BRCA	TCGA-BH-A18F-01A-11D-A12B-09	BRCA	d414b3fe-b768-4a98-b285-5284bffa66f9
TCGA-BH-A18L-01A-11D-A12B-09 BRCA fd9923db-2a27-432e-a0c6-4c44e6ee1f53 TCGA-BH-A18K-01A-11D-A12B-09 BRCA f75de986-bc8a-4ffc-9b35-011eee3a1446 TCGA-BH-A18L-01A-32D-A12B-09 BRCA 883cd3s9-2681-4822-8b22-29149a027514 TCGA-BH-A18M-01A-11D-A12B-09 BRCA 0c548c1e-cbb7-4432-8112-bb262a1ef949 TCGA-BH-A18N-01A-11D-A12B-09 BRCA 13c38ac4-c410-4602-83e3-9b80b4693839 TCGA-BH-A18N-01A-11D-A12B-09 BRCA adde6689-33c3-4f6f-8696-353d7c2dfb7 TCGA-BH-A18Q-01A-11D-A12B-09 BRCA adde6689-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18R-01A-11D-A12B-09 BRCA adde6689-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18N-01A-11D-A12B-09 BRCA adde6689-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18N-01A-11D-A12B-09 BRCA addef689-36c3-466-8b69-ebe6d4f59c2b TCGA-BH-A18N-01A-11D-A12B-09 BRCA addef6-847-4132-bdee-aaee2e027b28 TCGA-BH-A18N-01A-11D-A12B-09 BRCA addef6-847-4132-bdee-aaee2e027b28 TCGA-BH-A18N-01A-11D-A12B-09 BRCA added66-847-4132-bdee-aaee2e027b28 TCGA-BH-A18N-01A-11D-A12B-09 BRCA added66-847-4152-6bc3-edce4d8f6d22 TCGA-BH-A18N-01A-11D-A135-09 BRCA ac100def0-bd5-415f-909d-717226fd0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-98066ec7e427 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-98066ec7e427 TCGA-BH-A1EN-01A-11D-A135-09 BRCA 20466613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 20466613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 20566613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 3666613-88ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 3666613-88ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A131-09 BRCA 366661-38ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A131-09 BRCA 366661-38ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A131-09 BRCA 366661-38ad-42b-a-500-b144738fe18f TCGA-BH-A1F0-01A-11D-A131-09 BRCA 366661-38ad-42b-a-500-b144738fe18f TCGA-BH-A1F0-01A-11D-A131-09 BRCA 366661-38ad-42b-a-500-b144	TCGA-BH-A18H-01A-11D-A12B-09	BRCA	d3c1b990-aae2-45f8-be28-8ccd192a0fab
TCGA-BH-A18K-01A-11D-A12B-09 BRCA	TCGA-BH-A18I-01A-11D-A12B-09	BRCA	f0ca4831-d56d-4bae-b304-bb43c5d2f09b
TCGA-BH-A18L-01A-32D-A12B-09 BRCA 883cd3e9-2681-4822-8b22-29149a027514 TCGA-BH-A18M-01A-11D-A12B-09 BRCA 0e548c1e-cbb7-4432-8112-bb262a1ef9d9 TCGA-BH-A18N-01A-11D-A12B-09 BRCA 13c38ac4-c410-4602-83e3-9b80b4f93839 TCGA-BH-A18P-01A-11D-A12B-09 BRCA addc643a-37e9-46be-9bec-3e53d7c2dfb7 TCGA-BH-A18Q-01A-12D-A12B-09 BRCA adde6680-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a7662fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA a01c12fc-a33e-4a66-8b69-ebc6d4f59c2b TCGA-BH-A18T-01A-11D-A12B-09 BRCA adddfcb-8e47-4132-bdce-aaec2e027b28 TCGA-BH-A18T-01A-11D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18T-01A-11D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18T-01A-11D-A12B-09 BRCA a6150dd25-a8f4-4d9f-9da0-P56855ab67d TCGA-BH-A1EN-01A-11D-A135-09 BRCA a100ef0-be45-415f-9094-7172261d0084 TCGA-BH-A1ED-01A-11D-A135-09 BRCA 20131381-8a11-4254-8954-980e6ec7e427 TCGA-BH-A1ED-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EW-01A-11D-A135-09 BRCA dc578e75-e63-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EW-01A-11D-A131-09 BRCA dc578e74-d884-4bfa-b9e-d077396ed2de TCGA-BH-A1EP-01A-11D-A131-09 BRCA dc578e74-d884-34bfa-b9e-d077396ed2de TCGA-BH-A1EP-01A-11D-A131-09 BRCA dc58e74-b98-de64-4b6-8be4-74bbe2-3229ed TCGA-BH-A1F6-01A-11D-A131-09 BRCA dc68e74-db8-3de4-4b6-8be4-74bbe2-3de779900 TCGA-BH-A1F6-01A-11D-A131-09 BRCA dc69e64-db6-8be4-74bbe2-d077396e	TCGA-BH-A18J-01A-11D-A12B-09	BRCA	fd9923db-2a27-432e-a0c6-4c44e6ee1f53
TCGA-BH-A18M-01A-11D-A12B-09 BRCA 0e548c1e-ebb7-4432-8112-bb262a1ef9d9 TCGA-BH-A18N-01A-11D-A12B-09 BRCA 13c38ac4-c410-4602-83e3-9b80b4f93839 TCGA-BH-A18P-01A-11D-A12B-09 BRCA add624a3-57e9-46be-9bce-3e53d7c2db7 TCGA-BH-A18Q-01A-1D-A12B-09 BRCA add6680-33c3-44f61-6896-4553d70a00bcb TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a76e2fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA add6680-33c3-44f61-6896-4c5edfdf5e2fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA add6dfeb-e8d7-44132-bdce-aaec2e027b28 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-edcc4d816d22 TCGA-BH-A18U-01A-11D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-edcc4d816d22 TCGA-BH-A18U-01A-11D-A12B-09 BRCA a6150d25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A18U-01A-11D-A135-09 BRCA ac100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EN-01A-11D-A135-09 BRCA ac100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 20131381-8a11-4256-8954-98066e67c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 45f8e26-36-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EW-01A-11D-A135-09 BRCA 45f8e26-36-4bd8-360-4bd8-360-360-360-360-360-360-360-360-360-360	TCGA-BH-A18K-01A-11D-A12B-09	BRCA	f75de986-bc8a-4ffe-9b35-011eee3a1446
TCGA-BH-A18N-01A-11D-A12B-09 BRCA 13c38ac4-c410-4602-83e3-9b80b4f93839 TCGA-BH-A18P-01A-11D-A12B-09 BRCA add624a3-57e9-46be-9bcc-3e53d7c2dfb7 TCGA-BH-A18Q-01A-12D-A12B-09 BRCA add6624a3-57e9-46be-9bcc-3e53d7c2dfb7 TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a76c2fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a76c2fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA 4e0ddfcb-e847-4132-bdce-aaeee2e027b28 TCGA-BH-A18V-01A-21D-A12B-09 BRCA 4e0ddfcb-e847-4132-bdce-aaeee2e027b28 TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EV-01A-11D-A13F-09 BRCA c1100cf0-be45-415f-909d-7172261d0084 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 7ccda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc-bc75-e855329091f0 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc-bc75-e855329091f0 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc-bc75-e855329091f0 TCGA-BH-A1EV-01A-11D-A13L-09 BRC	TCGA-BH-A18L-01A-32D-A12B-09	BRCA	883cd3c9-2681-4822-8b22-29149a027514
TCGA-BH-A18P-01A-11D-A12B-09 BRCA add624a3-57e9-46be-9bcc-3e53d7c2dfb7 TCGA-BH-A18Q-01A-12D-A12B-09 BRCA add6680-33c3-4f6f-8696-453470a00bcb TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a7662fc TCGA-BH-A18R-01A-11D-A12B-09 BRCA 4c0ddfcb-e8d7-4132-bdce-aace2c027b28 TCGA-BH-A18R-01A-11D-A12B-09 BRCA 4c0ddfcb-e8d7-4132-bdce-aace2c027b28 TCGA-BH-A18R-01A-11D-A12B-09 BRCA 4e0ddfcb-e8d7-4132-bdce-aace2c027b28 TCGA-BH-A18R-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9dao-956855ab67d TCGA-BH-A18R-01A-11D-A13B-09 BRCA c100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EN-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1ER-01A-11D-A135-09 BRCA 43be2a9-078a-4be2-b67c-88553329091 fo TCGA-BH-A1ER-01A-11D-A135-09 BRCA 43be2a9-078a-4be2-b67c-88553329091 fo TCGA-BH-A1EX-01A-11D-A135-09 BRCA 6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 37b1685-0882-48ee-a38a-a05b5d1e8ba1 TCGA-BH-A1EX-01A-11D-A13L-09 BRCA </td <td>TCGA-BH-A18M-01A-11D-A12B-09</td> <td>BRCA</td> <td>0e548c1e-cbb7-4432-8112-bb262a1ef9d9</td>	TCGA-BH-A18M-01A-11D-A12B-09	BRCA	0e548c1e-cbb7-4432-8112-bb262a1ef9d9
TCGA-BH-A18Q-01A-12D-A12B-09 BRCA adde680-33-3-4f6f-8696-453470a00bcb TCGA-BH-A18R-01A-11D-A12B-09 BRCA d2facac2-81d9-4a9f-b4f6-1de89a7662fc a01c12fc-a33e-4a06-8b69-ebe6d4f59c2b TCGA-BH-A18T-01A-11D-A12B-09 BRCA de0ddfcb-e847-4132-bdce-aaee2e027b28 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18U-01A-11D-A13E-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18U-01A-11D-A13G-09 BRCA cal00ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA cal00ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 7ccda44b-e942-4077-9418-2a844ec53c9d TCGA-BH-A1EU-01A-11D-A135-09 BRCA 7ccda44b-e942-4077-9418-2a844ec53c9d TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63e-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63e-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EU-01A-11D-A135-09 BRCA d3fbc2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EU-01A-11D-A135-09 BRCA d578e75-e63e-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EU-01A-11D-A135-09 BRCA d578e75-e63e-4bdf-abfa-e2d063e9cd6d TCGA-BH-A1EU-01A-11D-A131-09 BRCA d578b1-A1EU-01A-11D-A135-09 BRCA d578e75-e63e-4bdf-abfa-e3d063e9cd6d TCGA-BH-A1EU-01A-11D-A131-09 BRCA d578b1-A1EU-01A-11D-A131-09 BRCA d578b1-A1EU-01A-11D-A131-09 BRCA d578b1-A1EU-01A-11D-A131-09 BRCA d564b1-A1EU-01A-11D-A131-09 BRCA d564b1-A1EU-01A-11D-A131-09 BRCA d564b1-A1EU-01A-11D-A131-09 BRCA d56749-A4843-A4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-11D-A131-09 BRCA d5664b1-A1EU-01A-11D-A131-09 BRCA d5666-A1A-A1EU-01A-11D-A131-09 BRCA d56664b1-A1EU-01A-11D-A131-09 BRCA d56664b1-A1EU-01A-11D-A131-09 BRCA d56664b1-A1EU-01A-11D-A131-	TCGA-BH-A18N-01A-11D-A12B-09	BRCA	13c38ac4-c410-4602-83e3-9b80b4f93839
TCGA-BH-A18R-01A-11D-A12B-09 BRCA 42facac2-81d9-4a9f-b4f6-1de89a7662fc TCGA-BH-A18S-01A-11D-A12B-09 BRCA a01c12fc-a33e-4a06-8b69-ebe6d4f59c2b TCGA-BH-A18T-01A-11D-A12B-09 BRCA 4e0ddfcb-e847-4132-bdce-aaec2e027b28 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EN-01A-11D-A135-09 BRCA ca100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EV-01A-11D-A131-09 BRCA 537b1685-0882-48ec-a38a-a505b51c8ba1 TCGA-BH-A1EV-01A-11D-A131-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1FC-01A-11D-A131-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1FC-01A-11D-A131-09 BRC	TCGA-BH-A18P-01A-11D-A12B-09	BRCA	add624a3-57e9-46be-9bcc-3e53d7c2dfb7
TCGA-BH-A18S-01A-11D-A12B-09 BRCA a01c12fc-a3ac-4a06-8b69-ebe6d4f59c2b TCGA-BH-A18T-01A-11D-A12B-09 BRCA 4e0ddfcb-e847-4132-bdce-aace2e027b28 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EN-01A-11D-A135-09 BRCA ca100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1EU-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc2-bc7c-b855329091f0 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc2-bc7c-b855329091f0 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 43fbc2a9-078a-4bc2-bc7c-b855329091f0 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 537b1685-0882-48ec-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 32121518-98d6-44b6-8bc4-74bbc232a9ed TCGA-BH-A1F5-01A-11D-A13L-09 BRCA 34cb095d-3d44-4c59-9cf5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 34cb095d-3d44-4c59-9cf5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 34cb095d-3d44-4c59-9cf5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 37c56d-438-4cd3-95e0-8708cc5437e7 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FB-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173	TCGA-BH-A18Q-01A-12D-A12B-09	BRCA	a4de6680-33c3-4f6f-8696-453470a00bcb
TCGA-BH-A18T-01A-11D-A12B-09 BRCA 4e0ddfcb-e847-4132-bdce-aaee2e027b28 TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EN-01A-11D-A135-09 BRCA cal00ef0-bc45-415f-909d-717226fd0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ccda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1ET-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1FQ-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 3567494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 3121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 372b5cd-3424-4899-9cf5-94592ba97900 TCGA-BH-A1FG-01A-11D-A13L-09 BRCA </td <td>TCGA-BH-A18R-01A-11D-A12B-09</td> <td>BRCA</td> <td>42facac2-81d9-4a9f-b4f6-1de89a7662fc</td>	TCGA-BH-A18R-01A-11D-A12B-09	BRCA	42facac2-81d9-4a9f-b4f6-1de89a7662fc
TCGA-BH-A18U-01A-21D-A12B-09 BRCA a8400863-c145-4c6c-bcf3-e4cc4d816d22 TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EN-01A-11D-A17G-09 BRCA ca100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ccda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1EU-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F5-01A-11D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1FD-01A-11D-A13L-09 BRC	TCGA-BH-A18S-01A-11D-A12B-09	BRCA	a01c12fc-a33e-4a06-8b69-ebe6d4f59c2b
TCGA-BH-A18V-01A-11D-A12B-09 BRCA 6150dd25-a8f4-4d9f-9da0-f956855ab67d TCGA-BH-A1EN-01A-11D-A17G-09 BRCA ca100ef0-be45-415f-909d-7172261d0084 TCGA-BH-A1EN-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1EU-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbc2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA 6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 82121518-98d6-44b6-8be4-74bbe232a9ed TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97000 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97000 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA	TCGA-BH-A18T-01A-11D-A12B-09	BRCA	4e0ddfcb-e847-4132-bdce-aaee2e027b28
TCGA-BH-AIEN-01A-11D-A13C-09 BRCA ca100ef0-be45-415f-909d-7172261d0084 TCGA-BH-AIEO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7e427 TCGA-BH-AIES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-AIET-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-AIEU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-AIEU-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-AIEW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-AIEW-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-AIEW-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-AIFD-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-AIFD-01A-11D-A13L-09 BRCA 82121518-98d6-44b6-8be4-74bbe232a9ed TCGA-BH-AIFD-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-AIFD-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-AIFD-01A-11D-A13L-09 BRCA 37eb5cd-4c38-4c43-95e0-8708e5437e7 TCGA-BH-AIFD-01A-11D-A13L-09 BRCA<	TCGA-BH-A18U-01A-21D-A12B-09	BRCA	a8400863-c145-4c6c-bcf3-e4cc4d816d22
TCGA-BH-A1EO-01A-11D-A135-09 BRCA 20131381-8a11-425d-8954-980e6ec7c427 TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53c9d TCGA-BH-A1ET-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA 3567494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F2-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3112f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-11D-A13L-09 BRCA </td <td>TCGA-BH-A18V-01A-11D-A12B-09</td> <td>BRCA</td> <td>6150dd25-a8f4-4d9f-9da0-f956855ab67d</td>	TCGA-BH-A18V-01A-11D-A12B-09	BRCA	6150dd25-a8f4-4d9f-9da0-f956855ab67d
TCGA-BH-A1ES-01A-11D-A135-09 BRCA 7ecda44b-e942-4077-9d18-2a844ec53e9d TCGA-BH-A1ET-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EW-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fc18f TCGA-BH-A1EW-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA 32667494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1FD-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 571fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FD-01A-11D-A13L-09 BRCA	TCGA-BH-A1EN-01A-11D-A17G-09	BRCA	ca100ef0-be45-415f-909d-7172261d0084
TCGA-BH-A1ET-01A-11D-A135-09 BRCA 9bd66613-68ad-42c1-ab43-dac1386027f9 TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA 32121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 37eb5d-4c38-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA 571fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FD-01A-11D-A13L-09 BRCA 561fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FD-01A-11D-A13L-09 BRCA 56db4486-6371-4892-863e-64838fcea624 TCGA-BH-A1FD-01A-11D-A13L-09 BRCA <td>TCGA-BH-A1EO-01A-11D-A135-09</td> <td>BRCA</td> <td>20131381-8a11-425d-8954-980e6ec7c427</td>	TCGA-BH-A1EO-01A-11D-A135-09	BRCA	20131381-8a11-425d-8954-980e6ec7c427
TCGA-BH-A1EU-01A-11D-A135-09 BRCA dc578e75-e63c-4bdf-abfa-e2d063c9cd6d TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1FO-01A-11D-A13L-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 32121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6e TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA 5671fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FD-01A-11D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FL-01A-11D-A13L-09 BRC	TCGA-BH-A1ES-01A-11D-A135-09	BRCA	7ecda44b-e942-4077-9d18-2a844ec53c9d
TCGA-BH-A1EV-01A-11D-A135-09 BRCA 43fbe2a9-078a-4be2-b67c-b855329091f0 TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1FO-01A-11D-A135-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA 35c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA 55c71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FD-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FD-01A-11D-A13L-09 BRCA 31f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FL-01A-11D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FM-01A-11D-A13L-09 BRCA	TCGA-BH-A1ET-01A-11D-A135-09	BRCA	9bd66613-68ad-42c1-ab43-dac1386027f9
TCGA-BH-A1EW-01A-11D-A135-09 BRCA c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A135-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 5c71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FH-01A-12D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FJ-01A-11D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FL-01A-11D-A13L-09 BRCA bb84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FM-01A-11D-A13L-09 BRC	TCGA-BH-A1EU-01A-11D-A135-09	BRCA	dc578e75-e63c-4bdf-abfa-e2d063c9cd6d
TCGA-BH-A1EX-01A-11D-A13L-09 BRCA 537b1685-0882-48ee-a38a-a05b5d1c8ba1 TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A135-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-12D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FL-01A-11D-A13L-09 BRCA bb84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FN-01A-11D-A13L-09 BRCA bb84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FN-01A-11D-A13L-09 BRC	TCGA-BH-A1EV-01A-11D-A135-09	BRCA	43fbe2a9-078a-4be2-b67c-b855329091f0
TCGA-BH-A1EY-01A-11D-A13L-09 BRCA 7c035023-8ea9-4504-8f03-9573745cb6ef TCGA-BH-A1F0-01A-11D-A135-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 5e71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-12D-A13L-09 BRCA d66bd486-6371-4892-863e-64838fcea624 TCGA-BH-A1FJ-01A-11D-A13L-09 BRCA b8CA TCGA-BH-A1FM-01A-11D-A13L-09 BRCA b8Acbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FM-01A-11D-A13L-09 BRCA b8CA TCGA-BH-A1FM-01A-11D-A13L-09 BRCA b8CA TCGA-	TCGA-BH-A1EW-01A-11D-A135-09	BRCA	c6f4b1b6-a8dd-4a9a-a500-b14a738fe18f
TCGA-BH-A1F0-01A-11D-A135-09 BRCA 3903b485-366d-4318-b17d-a0194f032bd8 TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 5e71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-12D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FJ-01A-11D-A13L-09 BRCA b84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FM-01A-11D-A13L-09 BRCA 7cb17736-03da-4f77-8397-145585a25b1e TCGA-BH-A1FN-01A-11D-A13L-09 BRCA bf92d76e-31ff-4273-82ea-982c4c26394b	TCGA-BH-A1EX-01A-11D-A13L-09	BRCA	537b1685-0882-48ee-a38a-a05b5d1c8ba1
TCGA-BH-A1F2-01A-31D-A13L-09 BRCA a5c67494-d843-4b14-ba9c-d077396ed2dc TCGA-BH-A1F5-01A-12D-A13L-09 BRCA 82121518-98d6-4db6-8be4-74bbe232a9ed TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 5e71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-12D-A13L-09 BRCA fd6bd486-6371-4892-863e-64838fcea624 TCGA-BH-A1FJ-01A-11D-A13L-09 BRCA b8C4 TCGA-BH-A1FL-01A-11D-A13L-09 BRCA bb84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FM-01A-11D-A13L-09 BRCA 7cb17736-03da-4f77-8397-145585a25b1e TCGA-BH-A1FN-01A-11D-A13L-09 BRCA bf92d76e-31ff-4273-82ea-982c4c26394b	TCGA-BH-A1EY-01A-11D-A13L-09	BRCA	7c035023-8ea9-4504-8f03-9573745cb6ef
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TCGA-BH-A1F6-01A-11D-A13L-09 BRCA 34eb095d-3d44-4c59-9ef5-94592ba97900 TCGA-BH-A1F8-01A-11D-A13L-09 BRCA 030cfc8a-7b43-4d73-8bfa-b68a47749e49 TCGA-BH-A1FC-01A-11D-A13L-09 BRCA 84c77098-03d0-4b22-afb1-797703e85c6c TCGA-BH-A1FD-01A-11W-A14Q-09 BRCA b372b5cd-4c38-4cd3-95e0-8708ce5437e7 TCGA-BH-A1FE-01A-11D-A13L-09 BRCA 5e71fc3a-a2f4-4899-9c1f-8fee1ef29e2e TCGA-BH-A1FG-01A-11D-A13L-09 BRCA 311f2f1a-75c8-4fee-b31d-0815d71a3173 TCGA-BH-A1FH-01A-12D-A13L-09 BRCA fd6bd486-6371-4892-863e-64838fcea624 TCGA-BH-A1FJ-01A-11D-A13L-09 BRCA dc62eafd-b5ad-42b4-9665-11ba6b22cff5 TCGA-BH-A1FL-01A-11D-A13L-09 BRCA bb84cbb1-7244-4d92-8977-a37dbafc47b4 TCGA-BH-A1FM-01A-11D-A13L-09 BRCA 7cb17736-03da-4f77-8397-145585a25b1e TCGA-BH-A1FN-01A-11D-A13L-09 BRCA bf92d76e-31ff-4273-82ea-982c4c26394b	TCGA-BH-A1F2-01A-31D-A13L-09	BRCA	a5c67494-d843-4b14-ba9c-d077396ed2dc
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TCGA-D8-A1JP-01A-11D-A13L-09	BRCA	1e21a355-0cb6-4a43-b134-50ff88dacf92
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TOOL DO LIVE OLD ALD ALLY OR	DDC.	21 20725 4125 44 2 04 15 04 15 20005 5
TCGA-D8-A1JT-01A-31D-A13L-09	BRCA	3be3972f-4125-44c3-94d6-0ddba2008fcf
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TCGA-D8-A1X5-01A-11D-A14K-09	BRCA	db4526d4-e344-4b5a-bb66-fd43b41764ca
TCGA-D8-A1X7-01A-11D-A14K-09	BRCA	1951aa38-481b-464c-9a78-0819312a0a93
TCGA-D8-A1X7-01A-11D-A14K-09	BRCA	7acb4232-db95-4889-942e-f1be897b4f2a
TCGA-D8-A1X8-01A-11D-A14K-09	BRCA	78c3c787-5731-4c38-8d7a-e5b503b11c36
TCGA-D8-A1X9-01A-12D-A159-09	BRCA	b5f65c3a-b922-4a81-863d-59b72b08d1bf
TCGA-D8-A1XA-01A-11D-A14G-09	BRCA	a362780b-8917-4438-9693-ec9fa84c352a
TCGA-D8-A1XB-01A-11D-A14G-09	BRCA	e5ca0f82-6fa9-4d54-adc7-385721f351f3
TCGA-D8-A1XC-01A-11D-A14G-09	BRCA	68fd3045-073d-4242-8a41-41b707fca625

TCCA DO ALVE OLA LLD ALAC OO	DDCA	a1597622 260 4062 0744 h45h0f14ha46
TCGA-D8-A1XF-01A-11D-A14G-09	BRCA	e1587f32-2ff9-40f3-97dd-b45b0f14be46
TCGA-D8-A1XG-01A-11D-A14G-09	BRCA	800ff536-a1d2-4213-b85e-7780851c6378
TCGA D8 A1VL 01A 11D A14V 00	BRCA	a37b27a2-c3b0-4f62-82a2-94e9205b1d6e 28d44e6e-c73f-4788-8ad4-2bd6572f643d
TCGA-D8-A1XL-01A-11D-A14K-09	BRCA	
TCGA-D8-A1XM-01A-21D-A14K-09	BRCA	07418962-0a82-43a2-a66f-614903ea8380
TCGA-D8-A1XD-01A-11D-A14K-09	BRCA	b5ff68a2-da74-4608-941e-dbac40153077
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TCGA-D8-A1XS-01A-11D-A14K-09	BRCA	5d302c04-302e-4040-9429-37cd672e8d53
TCGA-D8-A1XT-01A-11D-A14K-09	BRCA	bc13601e-3e03-4d7d-8e6e-5b05ff500ea3
TCGA-D8-A1XV-01A-11D-A14K-09	BRCA	55c547ee-7cc9-4b7a-aaca-22f2a8c8c3a4
TCGA-D8-A1XV-01A-11D-A14K-09	BRCA	a76adfd1-8c89-4c13-b570-5ccc47043a70
TCGA-D8-A1XW-01A-11D-A14K-09	BRCA	f29405cc-d712-4562-ac02-ca3c89fb82af
TCGA-D8-A1XY-01A-11D-A14K-09	BRCA	edb6d161-8f50-4c11-8246-487c4ea9a55d
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TCGA-D8-A1Y2-01A-11D-A159-09	BRCA	9dbf62eb-0de7-4410-b44b-fdf59026d8e6
TCGA-D8-A1Y3-01A-11D-A159-09	BRCA	64fa29ff-534f-4b22-b0c4-513e8657edb1
TCGA-D8-A27E-01A-11D-A16D-09	BRCA	eab47cbb-eab0-4dd6-9cd0-f2700e5b6227
TCGA-D8-A27F-01A-11D-A16D-09	BRCA	fc6d77a9-121b-48ab-a899-713c3d1319a2
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TCGA-D8-A27M-01A-11D-A16D-09	BRCA	cb9257f9-ca3f-4c14-a680-6632175dd526
TCGA-D8-A27N-01A-11D-A16D-09	BRCA	6a411174-582a-4c68-bb04-5ea2e504bf7c
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TCGA-D8-A27W-01A-11D-A16D-09	BRCA	b045d675-286b-4cf8-aed4-c7ff81a78919
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TCGA-E2-A107-01A-11D-A10M-09	BRCA	5804fc1c-063b-429d-a652-22b0de416bd6
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TCGA-E2-A14O-01A-31D-A10Y-09	BRCA	d6ab6f8d-0e65-40a3-bf98-7249e4075395
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TCCA F2 A140 01A 11D A12B 00	DDCA	as51af64 251f 49f9 ab02 620a27a50a0f
TCGA-E2-A14Q-01A-11D-A12B-09	BRCA	ee51cf6d-351f-48f8-ab93-639c27c50e9f
TCGA-E2-A14R-01A-11D-A10Y-09	BRCA	c7212115-1007-40cf-b9b5-7b25e2f5f2a4
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TCGA-E2-A14V-01A-11D-A12B-09	BRCA	703314fe-bfd5-45d5-9ed5-fcdce8a19fd6
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TCGA-E2-A14X-01A-11D-A10Y-09	BRCA	74039acd-5aca-4c65-818c-3b577d295be0
TCGA-E2-A14Z-01A-11D-A10Y-09	BRCA	c83eaaca-ced5-4630-abb5-ef34db888753
TCGA-E2-A150-01A-11D-A12B-09	BRCA	446064de-ff64-4113-9080-360e5bf6d5e4
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TCGA-E2-A154-01A-11D-A10Y-09	BRCA	336e39fb-d407-4ced-b7bb-e8ff5329abdb
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TCGA-E2-A15A-01A-11D-A12B-09	BRCA	b7e3eff1-65d5-491f-a726-35dc6752b370
TCGA-E2-A15C-01A-31D-A12B-09	BRCA	10c594a1-0843-4740-9d96-00211a9509fb
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TCGA-E2-A15I-01A-21D-A135-09	BRCA	9bec02b4-7cf0-4797-b1ac-253ef78a34af
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TCGA-E2-A1IK-01A-11D-A17G-09	BRCA	8577ac01-1274-4bd5-ab04-380eaa78d95b

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TCGA-E9-A1N9-01A-11D-A14G-09	BRCA	2aa7a1db-40a5-421b-97ab-1031e6fa7f04
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TCGA-E9-A1NE-01A-21D-A14K-09	BRCA	dbd34322-ac40-41f0-acc7-7bfd06afdf67
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TCGA-E9-A1NG-01A-21D-A14K-09	BRCA	1cbf389d-1ec8-4543-880f-4ef64c55a44b
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TCGA-EW-A1IX-01A-12D-A142-09	BRCA	01ea194f-dc06-4e15-9b9e-1c73668040e0
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TCGA-EW-A1IZ-01A-11D-A188-09	BRCA	18db4143-48cc-424c-8d23-46cf23056528
TCGA-EW-A1J1-01A-11D-A188-09	BRCA	4b8d51b3-8393-45d4-a73d-3c22c561d6f3
TCGA-EW-A1J2-01A-21D-A13L-09	BRCA	c906931e-dc1a-434c-96cd-58088762f1e7
TCGA-EW-A1J3-01A-11D-A13L-09	BRCA	ac13b81a-ca05-432c-918a-0c9c8170bf46
TCGA-EW-A1J5-01A-11D-A13L-09	BRCA	98bb3025-0637-4106-8621-12df7b5d662f
TCGA-EW-A1J6-01A-11D-A188-09	BRCA	d95c5cb1-d081-47fa-8ac0-1ade7652a0af
TCGA-EW-A1OV-01A-11D-A142-09	BRCA	e27ca8f5-3f76-4531-87ea-ba3a44f6830d
TCGA-EW-A1OX-01A-11D-A142-09	BRCA	7828f9cf-aa93-44a0-8070-efdf90a677f0
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TCGA-EW-A1P4-01A-21D-A142-09	BRCA	204e4ef3-e6b8-469f-9024-56c6f6f07afd
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TCGA-EW-A1P7-01A-21D-A142-09	BRCA	402abf40-5a01-467d-a5be-b9101743f34b
TCGA-EW-A1P8-01A-11D-A142-09	BRCA	e55f338f-97e2-4394-ae23-c92606069485
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TCGA-EW-A1PH-01A-11D-A14K-09	BRCA	ce860c6f-c87a-4a45-92df-ca34bfb2e8b2
TCGA-GI-A2C8-01A-11D-A16D-09	BRCA	535a899d-67ca-4500-8dda-63a331a3611c
TCGA-AA-3664-01A-01W-0900-09	COAD	9cff122a-9960-4f2e-ba5b-94736bad7f2b
TCGA-AA-3666-01A-02W-0900-09	COAD	d7065ea5-88b0-4b56-a367-5defa0d9ed27
TCGA-AA-3667-01A-01W-0900-09	COAD	c2799cdc-c6f7-44ba-a72c-e1632b434575
TCGA-AA-3672-01A-01W-0900-09	COAD	04dc0b16-834c-4351-b3b9-58fe558c634d
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TCGA-AA-3678-01A-01W-0900-09	COAD	968fea30-df40-425f-87ba-935942dbd450
TCGA-AA-3679-01A-02W-0900-09	COAD	94cfbc05-df22-4db0-9aa0-808faab01c61
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TCGA-AA-3693-01A-01W-0900-09	COAD	45ea6cb9-8d5e-4470-bd07-a2c59ddc5cf0
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TCGA-AA-3715-01A-01W-0900-09	COAD	554258ce-99c3-49a3-bfbf-131ec867a0e9
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TCGA-AA-3814-01A-01W-0900-09	COAD	733e8b21-718b-405d-b860-ed36c70a8411
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TCGA-AA-3837-01A-01W-0900-09	COAD	888c1825-a44b-49cb-bed1-09db01e54b75
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TCGA-AA-3852-01A-01W-0900-09	COAD	lee1ab0a-cd8c-49d5-ab8c-0d2a2f94724f
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TCGA-AA-3856-01A-01W-0900-09	COAD	7a07d137-7936-486d-aeb5-6d9598fe4660
TCGA-AA-3858-01A-01W-0900-09	COAD	99e41f17-b760-4b34-8230-39aa42db46fd
TCGA-AA-3860-01A-02W-0900-09	COAD	57869735-96fd-4439-ba2d-583df6fc32a0
TCGA-AA-3875-01A-01W-0900-09	COAD	06e6b2e8-634e-4b03-989e-0d192b60b64a

TCCA AA 2066 01A 01W 1072 00	COAD	69061-40 4215 491- 9105 754900-171-44
TCGA AA 2004 01A 01W 1073 00	COAD	689f1a40-4315-48bc-8b05-75d800e17b44
TCGA-AA-3994-01A-01W-1073-09	COAD	4348f66a-e104-4fdd-bdee-2f346832835d
TCGA-AA-A004-01A-01W-A00E-09	COAD	0b856311-aa63-44b7-a191-9d6d8308c3d0
TCGA-AA-A00N-01A-02W-A00E-09	COAD	dfb1aec9-d196-49e6-bdb1-9318222b8121
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TCGA-AY-4070-01A-01W-1073-09	COAD	a7a74785-31cf-4527-bae2-991d7df97b5f
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TCGA-02-0003-01A-01D-1490-08	GBM	458f13e0-34f3-4a92-b3b3-9a3c2ee3ef23
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TCGA-02-0047-01A-01D-1490-08	GBM	ce03026e-b756-43a2-972d-b3a4dcda5491
TCGA-02-0055-01A-01D-1490-08	GBM	9cd89af4-5118-4adb-aa1d-fbd03bf42a33
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TCGA-06-0119-01A-08D-1490-08	GBM	0cda6181-c62b-4ced-a543-d6138fd2e94a
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TCGA-06-0130-01A-01D-1490-08	GBM	c09f0ebd-d604-49a3-9738-0c65fd47fbf9
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TCGA-06-0137-01A-01D-1490-08	GBM	37c11dfc-c37c-4cb6-bd81-9e0a7789b0f1
TCGA-06-0139-01A-01D-1490-08	GBM	c84ff17d-436d-49c1-aef2-b998ffe4a693
TCGA-06-0140-01A-01D-1490-08	GBM	18c94086-d2cc-45cd-9bad-f8968a042d5e
TCGA-06-0141-01A-01D-1490-08	GBM	5af251d5-e76b-480c-8142-6d6fbfce0b2a
TCGA-06-0142-01A-01D-1490-08	GBM	4bce79ce-c59c-4d86-b25f-28c8edda1651

TCGA-06-0145-01A-01W-0224-08 GBM 8f904068-2967-4b38-8813-3ad0a99e4af8 TCGA-06-0151-01A-01D-1491-08 GBM 5fea9ebc-8c1b-4078-af87-79c7f5b5470b TCGA-06-0152-01A-02W-0323-08 GBM 79062efd-2b09-4798-a504-0a18ca30ef2d TCGA-06-0154-01A-03D-1491-08 GBM 15045707-3ddd-4ade-959a-b368437752fb TCGA-06-0155-01B-01D-1492-08 GBM 2dc59e9b-3a60-4178-9fa0-81cf5171622d TCGA-06-0157-01A-01D-1491-08 GBM b1e62d8e-24d2-4118-8cd0-3142acebdd5b TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0166-01A-01D-1491-08 GBM d530c696-235d-4a41-944c-77ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-ddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0168-01A-01D-1490-08 GBM 39520be3-a2af-4189-acf9-9239363333a TCGA-06-0169-01A-01D-1491-08 GBM 39520be3-a2af-4189-acf9-9239363333a TCGA-06-0173-01A-01D-1491-08 GBM 017e9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM 0
TCGA-06-0152-01A-02W-0323-08 GBM 79062efd-2b09-4798-a504-0a18ca30ef2d TCGA-06-0154-01A-03D-1491-08 GBM f5045707-3ddd-4ade-959a-b368437752fb TCGA-06-0155-01B-01D-1492-08 GBM 2dc59e9b-3a60-4178-9fa0-81cf5171622d TCGA-06-0157-01A-01D-1491-08 GBM b1e62d8e-24d2-4118-8cd0-3142acebdd5b TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0166-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e7f7ac21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2e-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1491-08 GBM 2053a41-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0169-01A-01D-1491-08 GBM 3952bba3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0178-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0188-01A-01W-0254-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0188-01A-01W-0254-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0188-01A-01W-0254-08 GBM <
TCGA-06-0154-01A-03D-1491-08 GBM f5045707-3ddd-4ade-959a-b368437752fb TCGA-06-0155-01B-01D-1492-08 GBM 2dc5999b-3a60-4178-9fa0-81cf5171622d TCGA-06-0157-01A-01D-1491-08 GBM b1e62d8e-24d2-4118-8cd0-3142acebdd5b TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0166-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e717ac21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2e-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-017-02A-11D-2280-08 GBM 39520be3-a2a1-4189-acf4-9d239363333a TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38feb5668c TCGA-06-018-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38feb5668c TCGA-06-018-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-018-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-018-01A-01D-01-491-08 GBM <td< td=""></td<>
TCGA-06-0155-01B-01D-1491-08 GBM 2dc59e9b-3a60-4178-9fa0-81cf5171622d TCGA-06-0157-01A-01D-1491-08 GBM b1e62d8e-24d2-4118-8cd0-3142acebdd5b TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-94d-e7f7ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda6366f33 TCGA-06-0169-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-d30-fb38fcb5668c TCGA-06-018-01A-01D-1491-08 GBM a4fa779b-d116-469-b170-60f3e215e9fb TCGA-06-018-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-018-01A-01D-1491-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-018-01A-01D-1491-08 GBM 25c
TCGA-06-0157-01A-01D-1491-08 GBM b1e62d8e-24d2-4118-8cd0-3142acebdd5b TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-94dd-e7f7ac21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1491-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb568c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-018-01A-01D-1491-08 GBM a5a2e50f-dc7e-44ce-bffe-b675a707bf53 TCGA-06-018-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765f3d7439 TCGA-06-018-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 2
TCGA-06-0158-01A-01D-1491-08 GBM 14580533-4a0c-47ca-bb51-c233700de35c TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e7f7ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0179-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 26c4c57d-b75a-4da-4d96-b40d-4c2c57a701e TCGA-06-0190-01A-01D-1491-08 GBM
TCGA-06-0165-01A-01D-1491-08 GBM 1728988e-0877-4194-92c5-92c1ee6c5f5b TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e7f7ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 23c2fc52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0195-01B-01D-1491-08 GBM
TCGA-06-0166-01A-01D-1491-08 GBM 70157018-a3c5-4ef8-9314-f8715a3438a4 TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e7f7ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0189-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0199-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM c065761d-f775-457f-bda0-4c7c257a701e TCGA-06-0195-01B-01D-1491-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0209-01A-01D-1491-08 GBM
TCGA-06-0167-01A-01D-1491-08 GBM d530c696-235d-4a41-944d-e7f7ae21aa17 TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0189-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM
TCGA-06-0168-01A-01D-1491-08 GBM 2b3bab1e-dddd-4c2c-b5ec-7bb6e700e070 TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0189-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4e9c-b91b-ded40492e61c TCGA-06-0213-01A-01D-1491-08 GBM
TCGA-06-0169-01A-01D-1490-08 GBM 06053a14-2d9a-4df0-a79b-81bda36bf3c3 TCGA-06-0171-02A-11D-2280-08 GBM 39520be3-a2af-4189-acf4-9d239363333a TCGA-06-0173-01A-01D-1491-08 GBM 0908aac1-d3b7-4eec-96f2-a28c3738388c TCGA-06-0174-01A-01D-1491-08 GBM 017c9167-0354-41e4-ad50-fb38fcb5668c TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0188-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM c065761d-f775-457f-bda0-4c7c257a701e TCGA-06-0195-01B-01D-1491-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b40392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0213-01A-01D-1491-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM
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TCGA-06-0178-01A-01D-1491-08 GBM a4fa779b-d116-4696-b170-60f3e215e9fb TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0188-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM c065761d-f775-457f-bda0-4c7c257a701e TCGA-06-0192-01B-01W-0348-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0211-02A-02D-2280-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
TCGA-06-0184-01A-01D-1491-08 GBM a5a2e50f-dc7e-44cc-bffe-b675a707bf53 TCGA-06-0185-01A-01W-0254-08 GBM bc62d57d-b536-41ab-a344-e765fd3f7439 TCGA-06-0188-01A-01W-0254-08 GBM cc0c78e7-1d76-45e6-b043-dc209bb9a32a TCGA-06-0189-01A-01D-1491-08 GBM 25c64c53-746c-4e92-976a-8bd947fb9c7f TCGA-06-0190-02A-01D-2280-08 GBM c065761d-f775-457f-bda0-4c7c257a701e TCGA-06-0192-01B-01W-0348-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0211-02A-02D-2280-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
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TCGA-06-0192-01B-01W-0348-08 GBM 43d7bc6f-be9b-4d5e-bcec-4fb30b0d9b65 TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0211-02A-02D-2280-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
TCGA-06-0195-01B-01D-1491-08 GBM 2a2fac52-44aa-41f7-ae27-de6b7eba8ff1 TCGA-06-0209-01A-01D-1491-08 GBM b4a7de67-14b6-4b8c-abbe-9eaa990d905e TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0211-02A-02D-2280-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
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TCGA-06-0210-02A-01D-2280-08 GBM b60392fb-43d9-4c9c-b91b-ded40492e61c TCGA-06-0211-02A-02D-2280-08 GBM 3914c02e-44ad-4c96-8464-61aa95b42c49 TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
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TCGA-06-0213-01A-01D-1491-08 GBM 885f9df7-fc27-43c2-9acc-833c410b2db1 TCGA-06-0214-01A-02D-1491-08 GBM 08ac57ec-0036-4134-a9bb-f22eaa27ab0d
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TCGA-06-0878-01A-01W-0424-08 GBM 07869e29-9ced-4be5-9a6c-8fd3c29ae487 TCGA-06-0879-01A-01W-0424-08 GBM f96b8966-e0c2-4fb6-b3f6-e76d7953d537
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TCGA-06-5411-01A-01D-1696-08 GBM 2fdab641-d73b-4f9a-aa4c-c1944f131a69
TCGA-06-5412-01A-01D-1696-08 GBM b6be0866-b8ae-4767-8cdc-e1dd4f78f440
TCGA-06-5413-01A-01D-1696-08 GBM 72c13e51-0dd2-4e96-af37-aa471407436f
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TCGA-06-5415-01A-01D-1486-08 GBM fca08ee9-b480-4dc7-be56-f1eb03b56f7c
TCGA-06-5417-01A-01D-1486-08 GBM 66350d36-6662-4d4c-9cf8-e052a17cddba
TCGA-06-5418-01A-01D-1486-08 GBM ae28fd78-d254-46fa-aba1-1353931aa414
TCGA-06-5856-01A-01D-1696-08 GBM 0bd9b573-712b-4da1-9c33-7b7f43d4af31
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TCGA-06-5859-01A-01D-1696-08 GBM bb404507-ab63-4d82-99c6-f3297bffc46f
TCGA-06-6388-01A-12D-1845-08 GBM c9214f8b-6684-4e29-812c-2a44963e8914
TCGA-06-6389-01A-11D-1696-08 GBM 10911471-5404-42d5-817e-f9616e7dacfc

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TCGA-06-6391-01A-11D-1696-08	GBM	40fc77dc-46df-4487-925f-1d87c5326661
TCGA-06-6693-01A-11D-1845-08	GBM	45ca8f53-6d0e-4659-a81f-258184b7a70e
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TCGA-06-6695-01A-11D-1845-08	GBM	13817acd-8c1e-4154-8b88-7cdc5f2660a7
TCGA-06-6697-01A-11D-1845-08	GBM	7d947ed1-1315-459e-b973-f3dd624d9e39
TCGA-06-6698-01A-11D-1845-08	GBM	d605a279-c0ea-467c-a423-cdf21547f87e
TCGA-06-6699-01A-11D-1845-08	GBM	90ba858d-e3bb-40d8-98ee-eeb127c58409
TCGA-06-6700-01A-12D-1845-08	GBM	6da42a38-94dd-49b7-8a03-df0f7174ca6f
TCGA-06-6701-01A-11D-1845-08	GBM	fad178f1-385b-4f94-bd29-567c1aa0a8fc
TCGA-08-0386-01A-01D-1492-08	GBM	90bf7f8f-4b8c-410f-afa6-2b439ec82f97
TCGA-12-0615-01A-01D-1492-08	GBM	a6068793-51e4-4762-9150-cdfb030e8ade
TCGA-12-0616-01A-01D-1492-08	GBM	b0e2fed7-38bd-48d8-a786-ac574c9fa5be
TCGA-12-0618-01A-01D-1492-08	GBM	390fc5e9-787e-4a3f-86c8-e3e0e7e43824
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TCGA-12-0688-01A-02D-1492-08	GBM	143dc738-1694-4105-8115-9cc0902ef35b
TCGA-12-0692-01A-01W-0348-08	GBM	937fb2a6-3856-4086-a327-8d8e593b7b7b
TCGA-12-0821-01A-01W-0424-08	GBM	357e3a3c-cceb-4b38-bc35-6fe8f5be5ac8
TCGA-12-1597-01B-01D-1495-08	GBM	7d35c610-cc06-4aa5-8c96-2f7b7465069f
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TCGA-12-3650-01A-01D-1495-08	GBM	8b1d52e2-489b-4972-9bef-1690ccd2bac9
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TCGA-12-3653-01A-01D-1495-08	GBM	fdc52d48-828e-481f-ba1c-0264f1da38a5
TCGA-12-5295-01A-01D-1486-08	GBM	796f5741-3b2d-46e5-b74f-e5a76604a401
TCGA-12-5299-01A-02D-1486-08	GBM	a44954fc-49f2-489a-8593-7de98963e4f8
TCGA-12-5301-01A-01D-1486-08	GBM	891fc6bc-d0a7-4064-842c-43d500b4ef5d
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TCGA-14-0781-01B-01D-1696-08	GBM	13878ec6-fce7-423e-b545-6656145e9d2c
TCGA-14-0786-01B-01D-1492-08	GBM	75fa4de1-29fd-4b54-b63a-add459f1d69c
TCGA-14-0787-01A-01W-0424-08	GBM	184b240c-ebf1-4ecf-87eb-aae0718cd81f
TCGA-14-0789-01A-01W-0424-08	GBM	3462087f-f791-43b4-b9d9-b11cc48eaf9e
TCGA-14-0790-01B-01D-1494-08	GBM	d63d49a0-9413-4583-a7a5-cb2c202cc085
TCGA-14-0813-01A-01W-0424-08	GBM	754cd19e-a319-4ddf-887b-ddca4914cdf9
TCGA-14-0817-01A-01W-0424-08	GBM	a5f06dfc-e9b2-46a6-bee5-604d2839baad
TCGA-14-0862-01B-01D-1845-08	GBM	f0b7d451-8190-45a4-8242-bf698f05243d
TCGA-14-0871-01A-01W-0424-08	GBM	0cc45f48-0967-42dc-8035-e76c6bd0a3fd
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TCGA-14-1043-01B-11D-1845-08	GBM	a439c422-8728-42f5-8dda-6e9e1590478c
TCGA-14-1395-01B-11D-1845-08	GBM	8825b7a5-dfac-4e21-b4ec-05161b1341e9
TCGA-14-1450-01B-01D-1845-08	GBM	7ec7f174-13f6-44b1-83e3-6f35a244f00e
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TCGA-14-1823-01A-01W-0643-08	GBM	1c3ddf6a-e496-4b87-833b-084d814b6876
TCGA-14-1825-01A-01W-0643-08	GBM	f0d7cb8b-995c-419b-a366-aadb156879bc
TCGA-14-1829-01A-01W-0643-08	GBM	c69ca476-9e11-4f6e-a4f5-6952f792a580
TCGA-14-2554-01A-01D-1494-08	GBM	53dec97d-0464-4ffd-8e2e-95b2b9a03af0
TCGA-15-0742-01A-01W-0348-08	GBM	3c015456-02f0-4473-be25-b53166da41ea
TCGA-15-1444-01A-02D-1696-08	GBM	cbd4d4e7-f1c4-446c-8dbc-ce06c872ec14
TCGA-16-0846-01A-01W-0424-08	GBM	cf3eb226-36c2-4498-a5c1-3f161de6fa3f
TCGA-16-0861-01A-01W-0424-08	GBM	deab6efd-8213-4f35-a897-060c605ce58b
TCGA-16-1045-01B-01W-0611-08	GBM	c92c1d87-0df9-4c5a-baef-2dd26ad6d75a
TCGA-19-1390-01A-01D-1495-08	GBM	d7e8e408-0a8f-4177-ad38-08c5da484ed0
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TCGA-19-2629-01A-01D-1495-08	GBM	56ffaa35-814c-4c0b-b3c6-d4514d34fec2
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TCGA-19-5958-01A-11D-1696-08	GBM	fd385a8e-d6dc-4e65-a023-ce485793c410
TCGA-19-5959-01A-11D-1696-08	GBM	dd3e4733-7154-4162-9a61-a3a685e5f561
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TCGA-26-1442-01A-01D-1696-08	GBM	17e25583-886e-4dc9-802b-35e67971073d
TCGA-26-5132-01A-01D-1486-08	GBM	d1132127-1250-43af-9c16-425798a3d1a7
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TCGA-26-5136-01B-01D-1486-08	GBM	39e0587b-1b04-4c68-8ae4-3ae7781e8017
TCGA-26-5139-01A-01D-1486-08	GBM	8199001b-a3c9-47e1-97cf-943fa8030f46
TCGA-26-6173-01A-11D-1845-08	GBM	af373e42-cbbf-4a89-8479-bdd413011885
TCGA-26-6174-01A-21D-1845-08	GBM	3ba04f15-48f4-4851-a21f-8fa7cc9eac6b
TCGA-27-1830-01A-01W-0643-08	GBM	b391392a-9865-4bf4-b5f1-fa4fb2ad1343
TCGA-27-1831-01A-01D-1494-08	GBM	9880c3c9-5685-42a7-8fe9-7585ea1a1d37
TCGA-27-1832-01A-01W-0643-08	GBM	7ea7ee22-55a6-4748-9607-d93a6a367122
TCGA-27-1833-01A-01W-0643-08	GBM	4d8d34d9-7069-436c-84d6-ace5760c2aec
TCGA-27-1834-01A-01W-0643-08	GBM	a6c0824e-3d2a-498a-af77-44ea96ba5ce4
TCGA-27-1835-01A-01D-1494-08	GBM	6d5fd73b-4cad-44ae-8c79-67f2b9d30328
100/1-27-1000-01/1-01D-1494-00	TODIVI	0d51d750**TCdd**TTdC**0C77**0712U7U3U3U320

CGA-27-1836-01A-01D-1494-08		1	
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TCGA-27-2521-01A-01D-1494-08 GBM d60f54f5-b154-42c4-99fb-cea4e7a33dc7 TCGA-27-2523-01A-01D-1494-08 GBM d60f54f5-b154-42c4-99fb-cea4e7a33dc7 TCGA-27-2526-01A-01D-1494-08 GBM belabcb7-b4c9-4447-b0c5-0fc09401eec0 TCGA-27-2526-01A-01D-1494-08 GBM belabcb7-b4c9-4447-b0c5-0fc09401eec0 TCGA-27-2528-01A-01D-1494-08 GBM 374cbd87-428e-4509-85c1-b7d3302c30a0 TCGA-28-1747-01C-01D-1494-08 GBM 374cbd87-428e-4509-85c1-b7d3302c30a0 TCGA-28-1747-01C-01D-1494-08 GBM c7446081-ac14-dac2-9564-d67d5212627c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458e-4129-aa91-61b8e4b90e53 TCGA-28-2590-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2502-01B-01D-1494-08 GBM 70746668-138a-4e00-8806-6579464595cb TCGA-28-2502-01B-01D-1494-08 GBM 70746668-138a-4e00-8806-6579464595cb TCGA-28-2509-01A-01D-1696-08 GBM 5f2dc309-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1696-08 GBM 5f2dc309-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1494-08 GBM 5f2dc309-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1494-08 GBM 5f2dc309-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1494-08 GBM 5f2dc309-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1494-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1494-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2510-01A-01D-1486-08 GBM 6cef4a03fc-4529-8193-21b380d96344 TCGA-28-2510-01A-01D-1486-08 GBM 76209124-b3f0-4b62-8b2c-e268abdefc2b TCGA-28-5209-01A-01D-1486-08 GBM 76209124-b3f0-4b62-8b2c-e268abdefc2b TCGA-28-5209-01A-01D-1486-08 GBM 8666742-5604-d47d-b96c-5288f637142 TCGA-28-5213-01A-01D-1486-08 GBM 8666742-5604-d47d-b96c-5288f637142 TCGA-28-5213-01A-01D-1486-08 GBM 68068a98-3889-4402-bept-38605189448 TCGA-28-5213-01A-01D-1486-08 GBM 766071-6603-4150-bept-386058555 TCGA-28-5219-01A-01D-1486-08 GBM 766071-6603-4150-bept-386058555 TCGA-28-5219-01A-01D-1486-08 GBM 766071-6603-4150-bept-386058555 TCGA-28-5219-01A-01D-1486-08 GBM 766071-6603-4150-bept-386058555 TCGA-28-5219-01A-01D-1486-08 GBM 766071-6603-4150-bept-38605355 TCGA-28-5219-01A-01D-1486-08 GBM 766071-6603-670	TCGA-27-2518-01A-01D-1494-08	GBM	dae099ff-330f-492b-a06d-6f975e9e5aea
TCGA-27-2523-01A-01D-1494-08	TCGA-27-2519-01A-01D-1494-08	GBM	b0daafab-b783-4cfc-9f7d-8017d98e80bb
TCGA-27-2524-01A-01D-1494-08 GBM ce679bfd-fbf9-4c78-822e-37d2322d544b TCGA-27-2526-01A-01D-1494-08 GBM be1abcb7-b4e9-4447-b0c5-0fc09401eec0 TCGA-27-2527-01A-01D-1494-08 GBM b8b00995-ada6-493b-bafc-0f6c9def41c9 TCGA-27-2528-01A-01D-1494-08 GBM 374cbd87-428e-4509-85c1-b7d3302c30a0 TCGA-28-1753-01A-01D-1494-08 GBM 7c746081-ac14-4ac2-9564-d67d522fc27c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aag1-c1b8e4b90c53 TCGA-28-2750-01A-01D-1696-08 GBM 282853f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1696-08 GBM 707466c8-138a-4ed0-b80c-6579464595cb TCGA-28-2502-01B-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4ed98d3830df TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4ed98d38380df TCGA-28-2510-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-5214-01A-01D-1486-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM	TCGA-27-2521-01A-01D-1494-08	GBM	3678d5f3-9a29-4750-b0a9-20e971ff6aa4
TCGA-27-2526-01A-01D-1494-08 GBM bc1abcb7-b4e9-4447-b0c5-0fc09401eec0 TCGA-27-2527-01A-01D-1494-08 GBM b8b00995-ada6-493b-bafc-0f6c9def41c9 TCGA-27-2528-01A-01D-1494-08 GBM 374cbd87-428e-4509-85c1-b7d3302c30a0 TCGA-28-1747-01C-01D-1494-08 GBM 7c746081-ac14-4ae2-9564-d67d52f2627c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aa91-61b8c4b90e53 TCGA-28-2499-01A-01D-1494-08 GBM 28858f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2500-01A-01D-1696-08 GBM 282cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2500-01A-01D-1696-08 GBM 7074b668-138a-4edb-b80-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4cd98d8380df TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c53610c TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c53610c TCGA-28-2513-01A-01D-1486-08 GBM 6ecf4a0e-3fcf-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM 2d795a16-bd3-344f0-8c01-6eecc0e1adb1 TCGA-28-5209-01A-01D-1486-08 GBM 76209124-b3f0-4bc2-8b2-e268abdefc2b TCGA-28-5214-01A-01D-1486-08 GBM <t< td=""><td>TCGA-27-2523-01A-01D-1494-08</td><td>GBM</td><td>d60f54f5-b154-42c4-99fb-cea4e7a33dc7</td></t<>	TCGA-27-2523-01A-01D-1494-08	GBM	d60f54f5-b154-42c4-99fb-cea4e7a33dc7
TCGA-27-2527-01A-01D-1494-08	TCGA-27-2524-01A-01D-1494-08	GBM	ce679bfd-fbf9-4c78-822e-37d2322d544b
TCGA-27-2528-01A-01D-1494-08 GBM 374cbd87-428e-4509-85c1-b7d3302c30a0 TCGA-28-1747-01C-01D-1494-08 GBM 7c746081-ac14-4ac2-9564-d67d52f2627c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aap1-61b8e4b90e53 TCGA-28-2509-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1494-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4ed0-8806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4cd98d8380df TCGA-28-2510-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610e TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610e TCGA-28-2510-01A-01D-1486-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-860-6eec0e1a0b1 TCGA-28-5209-01A-01D-1486-08 GBM 7c209124-b3f0-4bbc2-8b2-e-2688bdefe2b TCGA-28-5213-01C-11D-1486-08 GBM 7c209124-b3f0-4bbc2-8b2-e-2688bdefe2b TCGA-28-5213-01A-01D-1486-08 GBM 68666742-5ed0-4d7d-b96c-5218f6f37142 TCGA-28-5213-01A-01D-1486-08 GBM 68666742-5ed0-4d7d-b96c-5218f6f37142 TCGA-28-5213-01A-01D-1486-08 GBM 68666742-5ed0-4d7d-b96c-5218f6f37142 TCGA-28-5218-01A-01D-1486-08 GBM 68668742-5ed0-4d7d-b96c-5218f6f37142 TCGA-28-5218-01A-01D-1486-08 GBM 68668742-5ed0-4d7d-b96c-5218f6f37142 TCGA-28-5218-01A-01D-1486-08 GBM 6808a98-3889-4dd2-bet9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM 6608a98-3889-4dd2-bet9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM 6708a98-3889-4dd2-bet9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM 6708a98-3889-4dd2-bet9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-0	TCGA-27-2526-01A-01D-1494-08	GBM	bc1abcb7-b4e9-4447-b0c5-0fc09401eec0
TCGA-28-1747-01C-01D-1494-08 GBM 7c746081-ac14-4ae2-9564-d67d52f2627c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aa91-61b8e4b90e53 TCGA-28-2499-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1696-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4ed0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-dc98d8380df TCGA-28-2510-01A-01D-1696-08 GBM 52dd303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2514-01A-02D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5204-01A-01D-1486-08 GBM 7c209124-b369-db2-8b2e-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM 7c209124-b369-db2-8b2-e2268abdefe2b TCGA-28-5213-01A-01D-1486-08 GBM ef8b63f3-b820-d6ac-a99c-3d401a6203d7 TCGA-28-5213-01A-01D-1486-08 GBM 6	TCGA-27-2527-01A-01D-1494-08	GBM	b8b00995-ada6-493b-bafc-0f6c9def41c9
TCGA-28-1747-01C-01D-1494-08 GBM 7c746081-ac14-4ae2-9564-d67d52f2627c TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aa91-61b8e4b90e53 TCGA-28-2499-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1696-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4ed0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-dc98d8380df TCGA-28-2510-01A-01D-1696-08 GBM 52dd303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2514-01A-02D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5204-01A-01D-1486-08 GBM 7c209124-b369-db2-8b2e-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM 7c209124-b369-db2-8b2-e2268abdefe2b TCGA-28-5213-01A-01D-1486-08 GBM ef8b63f3-b820-d6ac-a99c-3d401a6203d7 TCGA-28-5213-01A-01D-1486-08 GBM 6			
TCGA-28-1753-01A-01D-1494-08 GBM c7143f1e-458c-4129-aa91-61b8e4b90e53 TCGA-28-2499-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1696-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466e8-138a-4ed0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4cd98d8380df TCGA-28-2510-01A-01D-1696-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2514-01A-01D-1494-08 GBM 52dd150c-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0c-3fcf-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ce4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eecc01a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefc2b TCGA-28-5210-01A-01D-1486-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5210-01A-01D-1486-08 GBM 292e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5216-01A-01D-1486-08 GBM <t< td=""><td>TCGA-27-2528-01A-01D-1494-08</td><td>GBM</td><td>374cbd87-428e-4509-85c1-b7d3302c30a0</td></t<>	TCGA-27-2528-01A-01D-1494-08	GBM	374cbd87-428e-4509-85c1-b7d3302c30a0
TCGA-28-2499-01A-01D-1494-08 GBM 28583f40-c3fc-4213-91c1-99d7d536551e TCGA-28-2501-01A-01D-1696-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4cd0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cec2-487a-9a8a-4cd98d8380df TCGA-28-2510-01A-01D-1494-08 GBM 52dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2514-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-268abdefe2b TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-268abdefe2b TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5214-01A-01D-1486-08 GBM 6866e742-5ed0-4d7d-b96e-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM 292e603-30c9-4e30-a425-8050189d4f8 TCGA-28-5216-01A-01D-1486-08 GBM 34c77b5d-c3a6-483-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM 68	TCGA-28-1747-01C-01D-1494-08	GBM	7c746081-ac14-4ae2-9564-d67d52f2627c
TCGA-28-2501-01A-01D-1696-08 GBM 2a2cb25d-4069-4824-b09d-2d49634ed284 TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4ed0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cee2-487a-9a8a-4cd98d8380df TCGA-28-2519-01A-01D-1696-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-0942eSc5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM 2d795a16-bdc3-4ft9-98e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5209-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2-e268abdefe2b TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96e-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM 3dc77b5d-2a6-4e83-96f4-fad729362d9 TCGA-28-5216-01A-01D-1486-08 GBM 3dc77b5d-2a6-4e83-96f4-fad729362d9 TCGA-28-5210-01A-01D-1486-08 GBM 3dc77b5d-2a6-4e83-96f4-fad729362d9 TCGA-28-5210-01A-01D-1486-08 GBM 680	TCGA-28-1753-01A-01D-1494-08	GBM	c7143f1e-458c-4129-aa91-61b8e4b90e53
TCGA-28-2502-01B-01D-1494-08 GBM 707466c8-138a-4ed0-b806-6579464595cb TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cee2-487a-9a8a-4cd98d8380df TCGA-28-2510-01A-01D-1696-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6cef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5216-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5218-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM	TCGA-28-2499-01A-01D-1494-08	GBM	28583f40-c3fc-4213-91c1-99d7d536551e
TCGA-28-2509-01A-01D-1494-08 GBM f4a62fe0-cee2-487a-9a8a-4cd98d8380df TCGA-28-2510-01A-01D-1696-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eecc0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 3dc77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5219-01A-01D-1486-08 GBM	TCGA-28-2501-01A-01D-1696-08	GBM	2a2cb25d-4069-4824-b09d-2d49634ed284
TCGA-28-2510-01A-01D-1696-08 GBM 5f2dc303-9859-4b63-8aab-c387da4b2cc1 TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM f8bc3f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5216-01A-01D-1486-08 GBM 3dc77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5218-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5219-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1496-08 GBM <	TCGA-28-2502-01B-01D-1494-08	GBM	707466c8-138a-4ed0-b806-6579464595cb
TCGA-28-2513-01A-01D-1494-08 GBM 52dd150e-abd7-4fd2-abe9-09428c5a610c TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5213-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5219-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6bcbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-bpcd-58b1c8a955e9 TCGA-28-6450-01A-11D-1696-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-32-1970-01A-01D-1494-08 GBM	TCGA-28-2509-01A-01D-1494-08	GBM	f4a62fe0-cee2-487a-9a8a-4cd98d8380df
TCGA-28-2514-01A-02D-1494-08 GBM 6eef4a0e-3fef-4529-8193-21b380d96344 TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-32-1970-01A-01D-1494-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1696-08 GBM	TCGA-28-2510-01A-01D-1696-08	GBM	5f2dc303-9859-4b63-8aab-c387da4b2cc1
TCGA-28-5204-01A-01D-1486-08 GBM e9590ee4-92d8-4afb-908e-0c816d2b82f3 TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1494-08 GBM	TCGA-28-2513-01A-01D-1494-08	GBM	52dd150e-abd7-4fd2-abe9-09428c5a610c
TCGA-28-5207-01A-01D-1486-08 GBM 2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1 TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1980-01A-01D-1494-08 GBM	TCGA-28-2514-01A-02D-1494-08	GBM	6eef4a0e-3fef-4529-8193-21b380d96344
TCGA-28-5208-01A-01D-1486-08 GBM 76209124-b3f0-4bb2-8b2c-e268abdefe2b TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1980-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM	TCGA-28-5204-01A-01D-1486-08	GBM	e9590ee4-92d8-4afb-908e-0c816d2b82f3
TCGA-28-5209-01A-01D-1486-08 GBM ef8b63f3-b820-46ac-a99c-3d401a6203d7 TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1980-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5207-01A-01D-1486-08	GBM	2d795a16-bdc3-44f0-8c01-6eeec0e1a0b1
TCGA-28-5211-01C-11D-1845-08 GBM f8dc846b-1b17-4699-9dc5-3f79e21eee94 TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1980-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5208-01A-01D-1486-08	GBM	76209124-b3f0-4bb2-8b2c-e268abdefe2b
TCGA-28-5213-01A-01D-1486-08 GBM b866e742-5ed0-4d7d-b96c-52f8f6f37142 TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1980-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5209-01A-01D-1486-08	GBM	ef8b63f3-b820-46ac-a99c-3d401a6203d7
TCGA-28-5214-01A-01D-1486-08 GBM c992e603-30c9-4e30-a425-8050189db4f8 TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1980-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5211-01C-11D-1845-08	GBM	f8dc846b-1b17-4699-9dc5-3f79e21eee94
TCGA-28-5215-01A-01D-1486-08 GBM 34c77b5d-c3a6-4e83-96f4-fadd729362d9 TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5213-01A-01D-1486-08	GBM	b866e742-5ed0-4d7d-b96c-52f8f6f37142
TCGA-28-5216-01A-01D-1486-08 GBM cde8518a-ce8e-4b54-ab21-5ad4171ab1b3 TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5214-01A-01D-1486-08	GBM	c992e603-30c9-4e30-a425-8050189db4f8
TCGA-28-5218-01A-01D-1486-08 GBM 68008a98-3889-4dd2-bcf9-f1f6cbca6355 TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5215-01A-01D-1486-08	GBM	34c77b5d-c3a6-4e83-96f4-fadd729362d9
TCGA-28-5219-01A-01D-1486-08 GBM f016e9f7-66a3-4f50-b9cd-58b1c8a955e9 TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5216-01A-01D-1486-08	GBM	cde8518a-ce8e-4b54-ab21-5ad4171ab1b3
TCGA-28-5220-01A-01D-1486-08 GBM f7b80486-fa19-49c7-8ace-ea61338677d7 TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5218-01A-01D-1486-08	GBM	68008a98-3889-4dd2-bcf9-f1f6cbca6355
TCGA-28-6450-01A-11D-1696-08 GBM 5f10d0c5-05b8-44bb-98ce-bbea41820850 TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5219-01A-01D-1486-08	GBM	f016e9f7-66a3-4f50-b9cd-58b1c8a955e9
TCGA-32-1970-01A-01D-1494-08 GBM 65723119-bdfe-46f0-b629-c171023abd71 TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-5220-01A-01D-1486-08	GBM	f7b80486-fa19-49c7-8ace-ea61338677d7
TCGA-32-1979-01A-01D-1696-08 GBM 0c81ebb9-20a6-40c1-9be2-17b99517e988 TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-28-6450-01A-11D-1696-08	GBM	5f10d0c5-05b8-44bb-98ce-bbea41820850
TCGA-32-1980-01A-01D-1696-08 GBM 9b267205-1994-46ff-8d0f-56625dae7c1b TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-32-1970-01A-01D-1494-08	GBM	65723119-bdfe-46f0-b629-c171023abd71
TCGA-32-1982-01A-01D-1494-08 GBM 9cf7c4cb-ce19-4b79-9163-b74369603e22 TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-32-1979-01A-01D-1696-08	GBM	0c81ebb9-20a6-40c1-9be2-17b99517e988
TCGA-32-1986-01A-01D-1494-08 GBM 5afe3ffc-ba3a-49bb-9837-091b600cbb35	TCGA-32-1980-01A-01D-1696-08	GBM	9b267205-1994-46ff-8d0f-56625dae7c1b
	TCGA-32-1982-01A-01D-1494-08	GBM	9cf7c4cb-ce19-4b79-9163-b74369603e22
TCGA-32-2615-01A-01D-1495-08 GBM 65e3c804-b1a3-4e21-9407-90a6edc4e290	TCGA-32-1986-01A-01D-1494-08	GBM	5afe3ffc-ba3a-49bb-9837-091b600cbb35
	TCGA-32-2615-01A-01D-1495-08	GBM	65e3c804-b1a3-4e21-9407-90a6edc4e290

TCGA-32-2632-01A-01D-1495-08			
TCGA-32-2638-01A-01D-1495-08	TCGA-32-2632-01A-01D-1495-08	GBM	27203e18-af27-478c-a224-8bca77a81c90
TCGA-32-522-01A-01D-1495-08	TCGA-32-2634-01A-01D-1495-08	GBM	52b2a114-4f8c-4e02-af9d-24c4a05d4ca0
TCGA.41-2571-01A-01D-1495-08	TCGA-32-2638-01A-01D-1495-08	GBM	1e103221-ab46-4a5c-9b96-5e34f0d49fc2
TCGA-41-2573-01A-01D-1495-08	TCGA-32-5222-01A-01D-1486-08	GBM	f48abf4d-f1fb-48bf-97a1-0c38435b6af7
TCGA-41-2575-01A-01D-1495-08	TCGA-41-2571-01A-01D-1495-08	GBM	36349a22-17eb-48d8-9b69-1921ee7576ff
TCGA-41-3392-01A-01D-1495-08 GBM c08b37a5-9938-4ab0-8183-d73b01cb9a89 TCGA-41-6651-01A-01D-1696-08 GBM 5td77ba9-5015-4d8b-86a0-582e5c76bdd6 TCGA-41-6646-01A-11D-1845-08 GBM 6272bb0c-c47b-4cd2-9f59-398f1a75f020 TCGA-74-6573-01A-12D-1845-08 GBM 0941e50e1205-49ed-8735-1186eaf87718 TCGA-74-6577-01A-11D-1845-08 GBM f4ce96de-d7fc-4892-93a-68002f387a12 TCGA-74-6577-01A-11D-1845-08 GBM 5be142d5-b6f7-4e1e-ae75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-b66be58de8e0 TCGA-74-6584-01A-11D-1845-08 GBM ced2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b0702ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4929-01A-01D-1486-08 GBM 6230f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM 44a2774-ca69-4f54-9bce-ec33d8481fed TCGA-76-4931-01A-01D-1486-08 GBM 44a2774-ca69-4f54-9bce-ec33d8481fed TCGA-76-4931-01A-1D-1486-08 GBM 81656daa-af7c-430-afa3-ob-10eb965 TCGA-76-4934-01A-01D-1486-08 GBM 6296	TCGA-41-2573-01A-01D-1495-08	GBM	fadc9e2a-d97d-4e86-a814-4f32f8cfd7a5
TCGA-41-5651-01A-01D-1696-08 GBM 5fd77ba9-5015-448b-86a0-582e5c76bdd6 TCGA-41-6646-01A-11D-1845-08 GBM 6272bb0e-c47b-4cd2-9f59-398f1a75f020 TCGA-74-6573-01A-12D-1845-08 GBM 0941e50e-1205-49ed-8735-186eaf87718 TCGA-74-6575-01A-11D-1845-08 GBM 14ec96d6-47fc-4892-9a36-80802f387a12 TCGA-74-6578-01A-11D-1845-08 GBM 5bc142d3-5bf7-4c1e-ac75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ac2128-4d95-4261-a30d-bd6bc58de8c0 TCGA-74-6584-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aac-6a9bd38f2ceb TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4929-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4929-01A-01D-1486-08 GBM 6e30/277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM 34f38b89-837a-48b7-b0e7-12acc23fc285 TCGA-76-4931-01A-01D-1486-08 GBM 4da27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4935-01A-11D-1696-08 GBM <	TCGA-41-2575-01A-01D-1495-08	GBM	4943e80a-d098-49cd-8261-1d53d42f8223
TCGA-41-6646-01A-11D-1845-08 GBM 6272bb0c-c47b-4cd2-9f59-398f1a75f020 TCGA-74-6573-01A-12D-1845-08 GBM 0941e50e-1205-49ed-8735-1f86eaf87718 TCGA-74-6575-01A-11D-1845-08 GBM f4ec96d6-d7fc-4892-9a36-80802f387a12 TCGA-74-6577-01A-11D-1845-08 GBM 5be142d5-b6f7-4e1e-ae75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-74-658-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-74-658-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aac-6a9bd38f2ceb TCGA-76-4925-01A-01D-1486-08 GBM 3c93cb58-d39b-4a5e-907a-8b5438630d21 TCGA-76-4926-01B-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4929-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7e-0a5121cde6d1 TCGA-76-4931-01A-01D-1486-08 GBM 3f48889-837a-48b7-b067-12aec236c85 TCGA-76-4932-01A-01D-1486-08 GBM 3f48889-837a-48b7-b067-12aec23fc285 TCGA-76-4932-01A-01D-1486-08 GBM 8165daa-af7c-430c-af3a-0e10e9a695 TCGA-76-4932-01A-01D-1486-08 GBM 28d06abf-437d-4be9-804b-44345pf74f36 TCGA-76-6191-01A-11D-1696-08 GBM 28d0	TCGA-41-3392-01A-01D-1495-08	GBM	c08b37a5-9938-4ab0-8183-d73b01cb9a89
TCGA-74-6573-01A-12D-1845-08 GBM O941e50e-1205-49ed-8735-186eaf87718 TCGA-74-6575-01A-11D-1845-08 GBM f4ee96d6-d7fc-4892-9a36-80802f387a12 TCGA-74-6578-01A-11D-1845-08 GBM Sbe142d5-b6f7-4e1e-ae75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-74-6584-01A-11D-1845-08 GBM cedd2d3-371b-ab12-8aac-6a9bd38f2ccb TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 3c93cb58-d39b-4a5c-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5e445 TCGA-76-4929-01A-01D-1486-08 GBM de69425-dbfd-4228-ab78-541062b5e445 TCGA-76-4929-01A-01D-1486-08 GBM da487742-ca69-4554-9bce-ec33d8481fed TCGA-76-4929-01A-01D-1486-08 GBM db4a27742-ca69-4554-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM db4a27742-ca69-4554-9bce-ec33d8481fed TCGA-76-4934-01A-01D-1486-08 GBM db4a27742-ca69-4554-9bce-ec33d8481fed TCGA-76-4934-01A-01D-1486-08 GBM db4a27742-ca69-454-9bce-ec33d8481fed TCGA-76-4935-01A-01D-1486-08 GBM db66ef-4108-4385-a940-53a61fd96651 TCGA-76-6191-01A-12D-1696-08 GBM db66ef-4108-4886-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM db66ef-4108-4886-a8eb-6ba8cdefb4a2 TCGA-76-6193-01A-11D-1696-08 GBM db66ef-4108-4889-98a1-b8d69700f4a3 TCGA-76-6280-01A-21D-1845-08 GBM 1c7f63d2-a2a4-42c-3-928b-319695a66443 TCGA-76-6280-01A-21D-1845-08 GBM 1c7f63d2-a2a4-42c-3-928b-319695a66443 TCGA-76-6280-01A-11D-1845-08 GBM 1c7f63d2-a2a4-42c-3-928b-319695a66443 TCGA-76-6280-01A-11D-1845-08 GBM 1c6ff11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6656-01A-11D-1845-08 GBM 1c6ff11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM 1c6ff11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM 1c6661-1a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM 1c6661-1a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM 1c6661-1a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM 1c6661-1a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6666-01B-11D-1845-08 GBM 1c7f6-6666-016-11D-1845-08 GBM 1c76-6666-016-11D-1845-	TCGA-41-5651-01A-01D-1696-08	GBM	5fd77ba9-5015-4d8b-86a0-582e5c76bdd6
TCGA-74-6575-01A-11D-1845-08 GBM f4ee96d6-d7fc-4892-9a36-80802f387a12 TCGA-74-6577-01A-11D-1845-08 GBM 5be142d5-b6f7-4e1e-ae75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-74-6584-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aac-6a9bd38f2ccb TCGA-76-4925-01A-01D-1486-08 GBM ced2d2d9-371b-4b12-8aac-6a9bd38f2ccb TCGA-76-4926-01B-01D-1486-08 GBM ce32fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4928-01A-01D-1486-08 GBM af4f8b89-837a-448b7-b0c7-12acc23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-obe10eb9a695 TCGA-76-4932-01A-01D-1486-08 GBM c9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-6191-01A-12D-1696-08 GBM c8d06abf-437d-4bc9-804b-44345a74736 TCGA-76-6192-01A-11D-1696-08 GBM d4bf66ef-4108-488-4896-98a1-b8d69700f4a3 TCGA-76-6280-01A-21D-1845-08 GBM	TCGA-41-6646-01A-11D-1845-08	GBM	6272bb0c-c47b-4cd2-9f59-398f1a75f020
TCGA-74-6577-01A-11D-1845-08 GBM Sbe142d5-b6f7-4e1e-ae75-49b302b332a2 TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-76-6584-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aae-6a9bd38f2ecb TCGA-76-4925-01A-01D-1486-08 GBM cedd2d49-371b-4b12-8aae-6a9bd38f2ecb TCGA-76-4926-01B-01D-1486-08 GBM 3c93cb58-d39b-4a5e-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af48b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-6191-01A-12D-1696-08 GBM e8d06abf-437d-4be9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM c29754bc-44e8-4980-98a1-b8669700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM <	TCGA-74-6573-01A-12D-1845-08	GBM	0941e50e-1205-49ed-8735-1f86eaf87718
TCGA-74-6578-01A-11D-1845-08 GBM a2ae2128-4d95-4261-a30d-bd6be58de8e0 TCGA-74-6584-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aac-6a9bd38f2ecb TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 3c93cb58-d39b-4a5e-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875c-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af48b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-6191-01A-12D-1696-08 GBM d80f66ef-4108-af86-a8eb-6ba8cdefb4a2 TCGA-76-6191-01A-12D-1696-08 GBM d20754bc-44e8-4980-98a1-88d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-871b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-d47a-acab-ac64d26c52c3 TCGA-76-6280-01A-11D-1845-08 GBM <	TCGA-74-6575-01A-11D-1845-08	GBM	f4ec96d6-d7fc-4892-9a36-80802f387a12
TCGA-74-6584-01A-11D-1845-08 GBM cedd2d49-371b-4b12-8aac-6a9bd38f2ccb TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 3c93cb58-d39b-4a5e-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0c7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-6935-01A-01D-1486-08 GBM e8d06abf-437d-4bc9-804b-4345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM d4bf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6285-01A-11D-1845-08 GBM <	TCGA-74-6577-01A-11D-1845-08	GBM	5be142d5-b6f7-4e1e-ae75-49b302b332a2
TCGA-76-4925-01A-01D-1486-08 GBM ca2fa3da-18d6-4e8b-8081-b07022ead6a8 TCGA-76-4926-01B-01D-1486-08 GBM 3c93cb58-d39b-4a5e-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM e8d0abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM d4bf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fef-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 6a751d65-5fef-4c03-8253-8f1b8faccab2 TCGA-76-6283-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6285-01A-11D-1845-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6285-01A-11D-1845-08 GBM <	TCGA-74-6578-01A-11D-1845-08	GBM	a2ae2128-4d95-4261-a30d-bd6be58de8e0
TCGA-76-4926-01B-01D-1486-08 GBM 3c93c558-d39b-4a5e-907a-8b5438630d21 TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7e-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbff6eef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6686-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c2e-2e5f1a8854d3 TCGA-76-6666-01A-11D-1845-08 GBM	TCGA-74-6584-01A-11D-1845-08	GBM	cedd2d49-371b-4b12-8aac-6a9bd38f2ccb
TCGA-76-4927-01A-01D-1486-08 GBM 2dc69425-dbfd-4228-ab78-541062b5c445 TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM d4bf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-4448-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6280-01A-21D-1845-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6285-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-37b-b494caf84f8d TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe6f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6667-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-66661-01B-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6666-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76	TCGA-76-4925-01A-01D-1486-08	GBM	ca2fa3da-18d6-4e8b-8081-b07022ead6a8
TCGA-76-4928-01B-01D-1486-08 GBM 6e30f277-875e-4ab8-bc7c-0a5121cde6d1 TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6660-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM	TCGA-76-4926-01B-01D-1486-08	GBM	3c93cb58-d39b-4a5e-907a-8b5438630d21
TCGA-76-4929-01A-01D-1486-08 GBM af4f8b89-837a-48b7-b0e7-12aec23fc285 TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-ec33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM d4bf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6285-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-666-01A-11D-1845-08 GBM fe6ff1a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6660-01A-11D-1845-08 GBM <t< td=""><td>TCGA-76-4927-01A-01D-1486-08</td><td>GBM</td><td>2dc69425-dbfd-4228-ab78-541062b5c445</td></t<>	TCGA-76-4927-01A-01D-1486-08	GBM	2dc69425-dbfd-4228-ab78-541062b5c445
TCGA-76-4931-01A-01D-1486-08 GBM d4a27742-ca69-4f54-9bce-cc33d8481fed TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6666-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6667-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6666-01A-11D-1845-08 GBM	TCGA-76-4928-01B-01D-1486-08	GBM	6e30f277-875e-4ab8-bc7c-0a5121cde6d1
TCGA-76-4932-01A-01D-1486-08 GBM 81656daa-af7c-430c-afa3-0eb10eb9a695 TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6650-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 717c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6662-01A-11D-1845-08 GBM	TCGA-76-4929-01A-01D-1486-08	GBM	af4f8b89-837a-48b7-b0e7-12aec23fc285
TCGA-76-4934-01A-01D-1486-08 GBM e9bc4701-562e-4d35-a949-53a61fd96651 TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6660-01A-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6663-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM	TCGA-76-4931-01A-01D-1486-08	GBM	d4a27742-ca69-4f54-9bce-ec33d8481fed
TCGA-76-4935-01A-01D-1486-08 GBM c8d06abf-437d-4bc9-804b-44345af74f36 TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fef-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-66657-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6660-01A-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6663-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM	TCGA-76-4932-01A-01D-1486-08	GBM	81656daa-af7c-430c-afa3-0eb10eb9a695
TCGA-76-6191-01A-12D-1696-08 GBM 4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2 TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-76-6664-01A-11D-1845-08 GBM	TCGA-76-4934-01A-01D-1486-08	GBM	e9bc4701-562e-4d35-a949-53a61fd96651
TCGA-76-6192-01A-11D-1696-08 GBM c29754bc-44e8-4980-98a1-b8d69700f4a3 TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1845-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-66657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM	TCGA-76-4935-01A-01D-1486-08	GBM	c8d06abf-437d-4bc9-804b-44345af74f36
TCGA-76-6193-01A-11D-1696-08 GBM 6a751d65-5fcf-4c03-8253-8f1b8faccab2 TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM	TCGA-76-6191-01A-12D-1696-08	GBM	4dbf66ef-4108-4a86-a8eb-6ba8cdefb4a2
TCGA-76-6280-01A-21D-1845-08 GBM 9096e339-7730-4d7a-acab-a6c4d26c52c3 TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1696-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da	TCGA-76-6192-01A-11D-1696-08	GBM	c29754bc-44e8-4980-98a1-b8d69700f4a3
TCGA-76-6282-01A-11D-1696-08 GBM 1c7f63d2-a2a4-42c3-928b-319695a66443 TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6193-01A-11D-1696-08	GBM	6a751d65-5fcf-4c03-8253-8f1b8faccab2
TCGA-76-6283-01A-11D-1845-08 GBM a4083f8b-0c39-4d65-a372-b494caf84f8d TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6280-01A-21D-1845-08	GBM	9096e339-7730-4d7a-acab-a6c4d26c52c3
TCGA-76-6285-01A-11D-1696-08 GBM 28380a2f-d302-45fb-a4c5-31b2fd150bc3 TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6282-01A-11D-1696-08	GBM	1c7f63d2-a2a4-42c3-928b-319695a66443
TCGA-76-6286-01A-11D-1845-08 GBM 45d03116-6cff-4074-9c26-2e5f1a8854d3 TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6283-01A-11D-1845-08	GBM	a4083f8b-0c39-4d65-a372-b494caf84f8d
TCGA-76-6656-01A-11D-1845-08 GBM fe66f11a-e03d-49c5-befe-db74ef55ce61 TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6285-01A-11D-1696-08	GBM	28380a2f-d302-45fb-a4c5-31b2fd150bc3
TCGA-76-6657-01A-11D-1845-08 GBM 6ba47878-126c-420d-b3c1-ca7ea8c182d0 TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6286-01A-11D-1845-08	GBM	45d03116-6cff-4074-9c26-2e5f1a8854d3
TCGA-76-6660-01A-11D-1845-08 GBM f4960945-c464-49c2-8ad6-d73a6fa47b20 TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6656-01A-11D-1845-08	GBM	fe66f11a-e03d-49c5-befe-db74ef55ce61
TCGA-76-6661-01B-11D-1845-08 GBM 8329c910-7ccf-4e84-b468-bd6cf23327a2 TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6657-01A-11D-1845-08	GBM	6ba47878-126c-420d-b3c1-ca7ea8c182d0
TCGA-76-6662-01A-11D-1845-08 GBM 7f7c80ca-6ad9-4820-83ca-5248b3873eea TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6660-01A-11D-1845-08	GBM	f4960945-c464-49c2-8ad6-d73a6fa47b20
TCGA-76-6663-01A-11D-1845-08 GBM 624864ad-3178-4a6d-a0cf-7fa3e9bdf8da TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6661-01B-11D-1845-08	GBM	8329c910-7ccf-4e84-b468-bd6cf23327a2
TCGA-76-6664-01A-11D-1845-08 GBM 6a8f17c6-060d-492e-8a39-53d9ac7035a4 TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6662-01A-11D-1845-08	GBM	7f7c80ca-6ad9-4820-83ca-5248b3873eea
TCGA-81-5910-01A-11D-1696-08 GBM bcf79a66-30e6-4554-982e-38d8eab46114	TCGA-76-6663-01A-11D-1845-08	GBM	624864ad-3178-4a6d-a0cf-7fa3e9bdf8da
	TCGA-76-6664-01A-11D-1845-08	GBM	6a8f17c6-060d-492e-8a39-53d9ac7035a4
	TCGA-81-5910-01A-11D-1696-08	GBM	bcf79a66-30e6-4554-982e-38d8eab46114
TCGA-81-5911-01A-12D-1845-08 GBM a501e01b-249c-43cb-aee2-f355c3c697dd	TCGA-81-5911-01A-12D-1845-08	GBM	a501e01b-249c-43cb-aee2-f355c3c697dd

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TCGA-87-5896-01A-01D-1696-08	GBM	640c33a6-a7df-4dba-9c21-367a9a839f0f
TCGA-BA-4074-01A-01D-1434-08	HNSC	2c84e904-0cbc-4645-b7e5-94ec45e61268
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TCGA-BA-4076-01A-01D-1434-08	HNSC	93dda6a6-907d-4dc2-9391-36dd09c767c6
TCGA-BA-4077-01B-01D-1434-08	HNSC	9b37211a-2150-4d33-bc6a-9d6a0a429708
TCGA-BA-4078-01A-01D-1434-08	HNSC	f02d0332-d7c8-4d2a-98ca-dbe7826437ae
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TCGA-BA-5151-01A-01D-1434-08	HNSC	dac15d7e-3930-4fcb-b752-4a4f00449ddd
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TCGA-BA-5153-01A-01D-1434-08	HNSC	363ccc6f-dab0-413e-bc42-d738ee25abcd
TCGA-BA-5555-01A-01D-1512-08	HNSC	65dc1531-713b-41ba-a567-caa12340c0cf
TCGA-BA-5556-01A-01D-1512-08	HNSC	d31fda32-363b-44e4-8f2c-834a66f46b87
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TCGA-BA-6870-01A-11D-1870-08	HNSC	2fdd3f42-cb2f-4faf-8a47-b8bfee058265
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TCGA-BA-6873-01A-11D-1870-08	HNSC	f65b842c-257e-4ac7-a155-23d3ac12d41c
TCGA-BA-7269-01A-11D-2012-08	HNSC	2e8ffdfc-48f5-41e0-9192-d761f3b518ef
TCGA-BB-4217-01A-11D-2078-08	HNSC	5916ef19-7838-4621-a869-de8c2b34931c
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TCGA-BB-4224-01A-01D-1434-08	HNSC	cfa7d658-031d-4cd4-9ca3-ceaa201f702d
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TCGA-BB-4227-01A-01D-1870-08	HNSC	c1b315bb-773b-4fd0-88ec-d11044996adc
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TCGA-BB-7862-01A-21D-2229-08	HNSC	84c57a23-1428-488e-9275-9f2bc3673476
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TCGA-BB-7871-01A-11D-2229-08	HNSC	8e13f8a5-5d80-4e34-bffa-54ae808114e7
TCGA-BB-7872-01A-11D-2229-08	HNSC	c05cb0b5-b288-48fb-bdc0-ee9acd6643a8
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TCGA-CN-4726-01A-01D-1434-08	HNSC	2201e681-a727-4fd2-adec-cbcb543b2232
TCGA-CN-4727-01A-01D-1434-08	HNSC	b24fc60a-fe83-4743-a6d3-d90b807412e1
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TCGA-CN-4730-01A-01D-1434-08	HNSC	543bbfe3-4a11-49af-b445-303f0912bfc3
TCGA-CN-4731-01A-01D-1434-08	HNSC	31ffd2d8-ee97-4002-9737-08c044878ace
TCGA-CN-4733-01A-02D-1870-08	HNSC	12880a34-83d1-4075-b62a-9fc61d18ca09
TCGA-CN-4734-01A-01D-1434-08	HNSC	fd54bbfa-62a2-4d8b-88fb-b74b91e1b958
TCGA-CN-4735-01A-01D-1434-08	HNSC	369ebdf4-ee27-414d-978d-3698711fae98
TCGA-CN-4736-01A-01D-1434-08	HNSC	788337f5-722c-45d6-8ca4-8037c489cb64
TCGA-CN-4737-01A-01D-1434-08	HNSC	4c6857bb-f20f-4ac9-9c2c-cb83c5387a74
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TCGA-CN-5360-01A-01D-1434-08	HNSC	174f1ea8-abcf-44ee-b17b-9687b3ab6dae
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TCGA-CN-5363-01A-01D-1434-08	HNSC	203f8426-6ec5-427a-9ccf-ec2b4683504d
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TCGA-CN-5365-01A-01D-1434-08	HNSC	a419a54c-58b4-4682-aaca-ed85697dd2a0
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TCGA-CN-5367-01A-01D-1434-08	HNSC	57adb398-48c5-4a14-a43e-f79a19befbda
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TCGA-CN-5370-01A-01D-2012-08	HNSC	f4ca6755-68ca-4702-b08b-65005d31e9be
TCGA-CN-5373-01A-01D-1434-08	HNSC	00988676-1e9b-4e00-b4aa-a8f86c21b206
TCGA-CN-5374-01A-01D-1434-08	HNSC	28d5a97b-3f3d-4595-9034-8491999fcf40
TCGA-CN-6010-01A-11D-1683-08	HNSC	2d9693f3-0917-42be-97b8-4dc15cc4d3f6
TCGA-CN-6011-01A-11D-1683-08	HNSC	0e0aa5da-2cb2-47b8-b000-83a07d68ed29
TCGA-CN-6012-01A-11D-1683-08	HNSC	c5d99faa-ef68-4f08-af97-d722bcc383f5
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TCGA-CN-6016-01A-11D-1683-08	HNSC	fcb6e29c-864d-483f-a848-8a61202d9516
TCGA-CN-6017-01A-11D-1683-08	HNSC	7cd89cbe-6bd9-41a2-a042-345fa0a09866
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TCGA-CN-6019-01A-11D-1683-08	HNSC	00769a89-ffc5-46f5-a42e-25b3eae886c2
TCGA-CN-6020-01A-11D-1683-08	HNSC	1f33c4c7-4f08-44a2-91f5-7ed2d7da68f0
TCGA-CN-6021-01A-11D-1683-08	HNSC	e62a2c4d-18e3-4ec8-8d93-40e055e65be4
TCGA-CN-6022-01A-21D-1683-08	HNSC	90cd2296-7133-4cbe-99cb-84b084eb88cd
TCGA-CN-6023-01A-11D-1683-08	HNSC	d03b8f96-c932-4abf-b508-f4e1b50739ee
TCGA-CN-6024-01A-11D-1683-08	HNSC	0604584e-0654-4b00-94fc-45e76588000c

TCGA-CN-698-01A-11D-1912-08			
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TCGA-CN-6995-01A-31D-2012-08	TCGA-CN-6992-01A-11D-1912-08	HNSC	7a70356c-74a3-40c3-bd32-3049da642831
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TCGA-CN-6998-01A-23D-2012-08	TCGA-CN-6996-01A-11D-1912-08	HNSC	c063bec5-c716-4ea2-843a-e9f0bec3b540
TCGA-CQ-5323-01A-01D-1683-08	TCGA-CN-6997-01A-11D-2012-08	HNSC	11b531cc-d9d9-496a-8448-e654ba71c414
TCGA-CQ-5324-01A-01D-1683-08 HNSC 22b6abf5-aad8-46ab-9b87-e3c12309eb59 TCGA-CQ-5325-01A-01D-1870-08 HNSC 22b6abf5-aad8-46ab-9b87-e3c12309eb59 TCGA-CQ-5327-01A-01D-1683-08 HNSC 199249f9-808d-4565-bb6b-82724f61edaa TCGA-CQ-5327-01A-01D-1683-08 HNSC da19d7bc-9748-4cd4-bd54-4792894838f0 TCGA-CQ-5332-01A-01D-1683-08 HNSC da2ed7bc-9748-4cd4-bd54-4792894838f0 TCGA-CQ-5333-01A-01D-1683-08 HNSC da2ed3db-dbc0-44f1-b625-173f819c122 TCGA-CQ-5333-01A-01D-1683-08 HNSC d2c2d3db-dbc0-44f1-b625-173f819c122 TCGA-CQ-5333-01A-01D-1683-08 HNSC d2c2d3db-dbc0-44f1-b625-173f819c122 TCGA-CQ-5334-01A-01D-1683-08 HNSC d3978192-2119-4910-26f-53834a201bf2 TCGA-CQ-5334-01A-01D-1683-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6222-01A-11D-1912-08 HNSC d62e3492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC d62e3492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC d62e3492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC d62e3492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC c335130-8731-430d-a792-280e01629e8f TCGA-CQ-6222-01A-11D-1912-08 HNSC c335130-8731-430d-a792-280e01629e8f TCGA-CQ-6222-01A-11D-1912-08 HNSC c62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6222-01A-11D-1912-08 HNSC c62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6222-01A-11D-1912-08 HNSC c62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6223-01A-11D-1912-08 HNSC c62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6223-01A-11D-1912-08 HNSC c7e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-6228-01A-11D-1912-08 HNSC c62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC c64242bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-6228-01A-11D-1912-08 HNSC c64242bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-6228-01A-11D-1912-08 HNSC c64425b-a531-4636-8e68-bdaf212df6dc TCGA-CQ-6228-01A-01D-5151-08 HNSC c64425-b6446-8645-ba93352613f631	TCGA-CN-6998-01A-23D-2012-08	HNSC	9c364f7e-5b90-44ef-9f80-250e428989ef
TCGA-CQ-5325-01A-01D-1683-08 HNSC 22b6abt5-aad8-46ab-9b87-e3c12309cb59 TCGA-CQ-5326-01A-01D-1870-08 HNSC 199249f9-808d-4565-bb6b-82724f61edaa TCGA-CQ-5327-01A-01D-1683-08 HNSC da19d7bc-9748-4cd4-bd54-4792894838f0 TCGA-CQ-5329-01A-01D-1683-08 HNSC 5aa9b6fc-4169-4346-98ib-4c711d08d701 TCGA-CQ-5330-01A-01D-1683-08 HNSC 4ce7e702-9b62-459c-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC d2c2d3db-dbc0-44f1-b625-17f3819c122 TCGA-CQ-5331-01A-01D-1683-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-5334-01A-01D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC 65e67eda-16a4-4dfd-902-78c770362656 TCGA-CQ-6221-01A-11D-1912-08 HNSC 65e67eda-16a4-4dfd-949-546c76d94a02 TCGA-CQ-6222-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2e492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC d6263b94-0ffe-40e7-9e8bf2-a9e7abde575f TCGA-CQ-6223-01A-11D-1912-08 HNSC d2311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6223-01A-11D-1912-08 HNSC d311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6223-01A-11D-1912-08 HNSC d311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6223-01A-11D-1912-08 HNSC d311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6223-01A-11D-1912-08 HNSC d625509e-4f77-41ca-9bc2-3f20c2dd4c49 TCGA-CQ-6229-01A-11D-1912-08 HNSC d6242bb-a531-430d-a792-280e01629e8f TCGA-CQ-6229-01A-11D-1912-08 HNSC d55e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC d64-422bb-a531-4363-8e68-b-bdaf212dff6dc TCGA-CQ-7065-01A-11D-2078-08 HNSC d64-422bb-a531-4636-8e68-bdaf212dff6dc TCGA-CQ-7065-01A-11D-2078-08 HNSC d64-422bb-a531-4636-8e68-bdaf212dff6dc TCGA-CQ-7065-01A-11D-2078-08 HNSC d64-422bb-a531-4636-8e68-bdaf212dff6dc TCGA-CR-5243-01A-01D-1512-08 HNSC d58057b4-29ef-4a55-b6ab-39352613f631 TCGA-CR-5243-01A-01D-1512-08 HNSC d58057b4-29ef-4a55-b6ab-39352613f631 TCGA-CR-5243-01A-01D-1512-08 HNSC d58057b4-29ef-4a55-b6ab-39352613f631 TCGA-CR-5249-01A-01D-1512-08 HNSC d58057b4-29ef-4a55-	TCGA-CQ-5323-01A-01D-1683-08	HNSC	892067ef-c465-46ea-8f91-10636dd0081b
TCGA-CQ-5326-01A-01D-1870-08 HNSC 199249f9-808d-4565-bb6b-82724f61edaa TCGA-CQ-5327-01A-01D-1683-08 HNSC da19d7bc-9748-4cd4-bd54-4792894838f0 TCGA-CQ-5329-01A-01D-1683-08 HNSC 5aa9b6fc-4169-4346-98fb-4c711d08d701 TCGA-CQ-5330-01A-01D-1683-08 HNSC 4ce7e702-9b62-459e-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC 4fd4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5332-01A-01D-1683-08 HNSC 4fd4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC 65e6feda-16a4-4dfd-9409-546c76d94a02 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6166fdd-c0b5-44a3-814d-0e94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC d263b94-0ffe-4009-479c-8bf2-a9e7abde575f TCGA-CQ-6222-01A-11D-1912-08 HNSC d35130-8731-430d-a792-280e01629e8f TCGA-CQ-6222-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6222-01A-11D-1912-08 HNSC d55e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6222-01A-11D-1912-08 HNSC d55e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC d55e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC d5250be-4d7-4lca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC d5250be-4d7-alca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC d5250be-4d7-alca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC d64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC d64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2078-08 HNSC d64c42bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2078-08 HNSC d64c42bb-a531-4636-8e68-bdaf212df6dc TCGA-CR-5243-01A-01D-1512-08 HNSC d64c42bb-a531-4636-8e68-bdaf212df6dc TCGA-CR-5243-01A-01D-1512-08 HNSC d64c42bb-a531-4636-8e68-bdaf212df6dc TCGA-CR-5243-01A-01D-1512-08 HNSC d5663d7-8852-466-554808567 TCGA-CR-5243-01A-01D-1512-08 HNSC d5663d7-8852-466-554808567 TCGA-CR-524	TCGA-CQ-5324-01A-01D-1683-08	HNSC	67b184fe-c4f4-49f3-938e-5370eb6246b9
TCGA-CQ-5327-01A-01D-1683-08 HNSC da19d7bc-9748-4cd4-bd54-4792894838f0 TCGA-CQ-5332-01A-01D-1683-08 HNSC 5aa9b6fc-4169-4346-98fb-4c711d08d701 TCGA-CQ-5330-01A-01D-1683-08 HNSC 4ce7e702-9b62-459e-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC d2c2d3db-dbc0-44f1-b625-17f3f819c122 TCGA-CQ-5332-01A-01D-1683-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5332-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0e94c5be41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC d62e492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC be7eb54-1d09-479e-8bf2-ape7abde575f TCGA-CQ-6222-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6222-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-380e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d530-3669-3669-3669-3669-3669-3669-3669-36	TCGA-CQ-5325-01A-01D-1683-08	HNSC	22b6abf5-aad8-46ab-9b87-e3c12309cb59
TCGA-CQ-5329-01A-01D-1683-08 HNSC 5aa9b6fc-4169-4346-98fb-4c711d08d701 TCGA-CQ-5330-01A-01D-1683-08 HNSC 4ce7e702-9b62-459e-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC 4ce7e702-9b62-459e-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5332-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC 6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67eda-16a4-ddfd-94a9-546c76d94a02 TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67eda-16a4-ddfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0e94c5be41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6225-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6228-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6229-01A-11D-1912-08 HNSC c3d51a0-8731-430d-a792-320e01d6499 TCGA-CQ-6229-01A-11D-1912-08 HNSC 655e02b-1a6e-4eab-948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 655e02b-1a6e-4eab-948-4120d6c31c29 TCGA-CQ-6229-01A-11D-2229-08 HNSC 07c76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-6228-01A-11D-2229-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-5248-01A-01D-512-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-5248-01A-01D-1512-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-5249-01A-01D-1512-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-5249-01A-01D-1512-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-5249-01A-01D-1512-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-6470-01A-11D-1870-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf43 TCGA-CR-6470-01A-11D-1870-08 HNSC 97a96e61-f2de-4af4-807a-3925c1fibf4	TCGA-CQ-5326-01A-01D-1870-08	HNSC	199249f9-808d-4565-bb6b-82724f61edaa
TCGA-CQ-5330-01A-01D-1683-08 HNSC 4ce7e702-9b62-459e-b2b4-a26cabba3a93 TCGA-CQ-5331-01A-02D-1870-08 HNSC d2c2d3db-dbc0-44f1-b625-17i3i819c122 TCGA-CQ-5332-01A-01D-1683-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC d65e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0e94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6222-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c351a0-8731-430d-a792-280e01629e8f TCGA-CQ-6227-01A-11D-1912-08 HNSC c35509c-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC c65509c-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC c7e6152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7066-01A-11D-2078-08 HNSC 07e8b3-5b8b-4d5b-b812-86165f949a20 TCGA-CQ-7068-01A-11D-2078-08 HNSC 07e8b3-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNSC 07e76152-9e83-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bPca3-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bPca3-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bPca3-476-8b39-6ee652ad8e5f TCGA-CR-6467-01A-11D-1870-08 HNSC 42bPca3-476-8b39-6ee652ad8e5f TCGA-CR-6470-01A-11D-1870-08 HNSC 42bPca3-476-8b39-6ee652ad8e5f TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5	TCGA-CQ-5327-01A-01D-1683-08	HNSC	da19d7bc-9748-4cd4-bd54-4792894838f0
TCGA-CQ-5331-01A-02D-1870-08 HNSC d2c2d3db-dbc0-44f1-b625-17f3f819c122 TCGA-CQ-5332-01A-01D-1683-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC d65e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0e94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6222-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c3311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC c362509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC c362509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7066-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2012-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2012-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CR-5243-01A-01D-1512-08 HNSC 0297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNSC 02968b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNSC 04269-0446-045-045-045-045-045-045-045-045-045-045	TCGA-CQ-5329-01A-01D-1683-08	HNSC	5aa9b6fc-4169-4346-98fb-4c711d08d701
TCGA-CQ-5332-01A-01D-1683-08 HNSC 4fdf4f0d-0a55-4b5e-8545-65f1aad37c10 TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC 3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC 652679a4-16a4-4dfd-9018-4ceb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67ead-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-cob5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC bc7cb5b4-1d09-479c-8bf2-a9e7abdc575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd315590-ac69-4ff2-8bb-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC cd55502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 67c705b-3a-425-948-4120d6c31c29 TCGA-CQ-6229-01A-11D-2078-08 HNSC 67c962b-3a-425-948-4120d6c31c29 TCGA-CQ-7065-01A-11D-2078-08 HNSC	TCGA-CQ-5330-01A-01D-1683-08	HNSC	4ce7e702-9b62-459e-b2b4-a26cabba3a93
TCGA-CQ-5334-01A-01D-1683-08 HNSC 39978192-2119-4910-a2f6-53834a2b1bf2 TCGA-CQ-6218-01A-11D-1912-08 HNSC d3717097-7cdb-446f-a020-78c770362656 TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6221-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7c55b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6223-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c331590-3c69-4ff2-8fbd-cb5b0f21975c TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC c55e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7066-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5248-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC 42bf9ca3-4768-b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-476-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-476-8b39-6ee652ad8e5f TCGA-CR-6467-01A-11D-1870-08 HNSC 42bf9ca3-476-8b39-6ee652ad8e5f TCGA-CR-6467-01A-11D-1870-08 HNSC 297e330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 5087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-5331-01A-02D-1870-08	HNSC	d2c2d3db-dbc0-44f1-b625-17f3f819c122
TCGA-CQ-6218-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC d6166f0d-c0b5-4464-849-546c76d94a02 TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-c0b5-4443-814d-0e94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6222-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6223-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC c331590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC c331590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6228-01A-11D-1912-08 HNSC c32509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6229-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-becf-bda80af58d22 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-becf-bda80af58d22 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-becf-bda80af58d22 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-becf-bda80af58d22 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1eff1cdb5714 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1eff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-5332-01A-01D-1683-08	HNSC	4fdf4f0d-0a55-4b5e-8545-65f1aad37c10
TCGA-CQ-6219-01A-11D-1912-08 HNSC c6263b94-0ffe-40e7-9184-deb427c67802 TCGA-CQ-6220-01A-11D-1912-08 HNSC 55e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC d62c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-4lca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7066-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-2012-08 HNS	TCGA-CQ-5334-01A-01D-1683-08	HNSC	39978192-2119-4910-a2f6-53834a2b1bf2
TCGA-CQ-6220-01A-11D-1912-08 HNSC 65e67eda-16a4-4dfd-94a9-546c76d94a02 TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6222-01A-11D-1912-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc41b0 TCGA-CQ-6223-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-a6f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9c7abde575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-2012-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNS	TCGA-CQ-6218-01A-11D-1912-08	HNSC	d3717097-7cdb-446f-a020-78c770362656
TCGA-CQ-6221-01A-11D-2078-08 HNSC d6166f0d-c0b5-44a3-814d-0c94c5bc4lb0 TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-2012-08 HNSC 295e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-476-8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08	TCGA-CQ-6219-01A-11D-1912-08	HNSC	c6263b94-0ffe-40e7-9184-deb427c67802
TCGA-CQ-6222-01A-11D-1912-08 HNSC de2c492f-5cd8-4330-a5de-36f693ec31af TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2029-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5248-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNS	TCGA-CQ-6220-01A-11D-1912-08	HNSC	65e67eda-16a4-4dfd-94a9-546c76d94a02
TCGA-CQ-6223-01A-11D-1912-08 HNSC be7cb5b4-1d09-479c-8bf2-a9e7abde575f TCGA-CQ-6224-01A-11D-1912-08 HNSC c03d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7065-01A-11D-2078-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6477-01A-11D-1870-08 HNS	TCGA-CQ-6221-01A-11D-2078-08	HNSC	d6166f0d-c0b5-44a3-814d-0c94c5bc41b0
TCGA-CQ-6224-01A-11D-1912-08 HNSC cd3d51a0-8731-430d-a792-280e01629e8f TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-1512-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6222-01A-11D-1912-08	HNSC	de2c492f-5cd8-4330-a5de-36f693ec31af
TCGA-CQ-6225-01A-11D-1912-08 HNSC cd311590-3c69-4ff2-8fbd-cb5b0f21975e TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6223-01A-11D-1912-08	HNSC	be7cb5b4-1d09-479c-8bf2-a9e7abde575f
TCGA-CQ-6227-01A-11D-1912-08 HNSC ca62509e-d477-41ca-9bc2-3f20c2dd4e49 TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6471-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6224-01A-11D-1912-08	HNSC	c03d51a0-8731-430d-a792-280e01629e8f
TCGA-CQ-6228-01A-11D-1912-08 HNSC 655e502b-1a6e-4eab-a948-4120d6c31c29 TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6225-01A-11D-1912-08	HNSC	cd311590-3c69-4ff2-8fbd-cb5b0f21975e
TCGA-CQ-6229-01A-11D-1912-08 HNSC 07e76152-9e83-42a5-9111-c39a2310a2e4 TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6227-01A-11D-1912-08	HNSC	ca62509e-d477-41ca-9bc2-3f20c2dd4e49
TCGA-CQ-7065-01A-11D-2078-08 HNSC 64c422bb-a531-4636-8e68-bdaf212df6dc TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6228-01A-11D-1912-08	HNSC	655e502b-1a6e-4eab-a948-4120d6c31c29
TCGA-CQ-7067-01A-11D-2229-08 HNSC 01f46aa2-e15b-4544-add5-c783868b6c26 TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-6229-01A-11D-1912-08	HNSC	07e76152-9e83-42a5-9111-c39a2310a2e4
TCGA-CQ-7068-01A-11D-2078-08 HNSC 97a96e61-f2dc-4af4-807a-3925c1ffbf43 TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-7065-01A-11D-2078-08	HNSC	64c422bb-a531-4636-8e68-bdaf212df6dc
TCGA-CR-5243-01A-01D-1512-08 HNSC 297e8b35-5b8b-4d5b-b812-86165f949a20 TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-7067-01A-11D-2229-08	HNSC	01f46aa2-e15b-4544-add5-c783868b6c26
TCGA-CR-5247-01A-01D-2012-08 HNSC 3b5b07b4-29ef-4a55-b6ab-93352613f631 TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CQ-7068-01A-11D-2078-08	HNSC	97a96e61-f2dc-4af4-807a-3925c1ffbf43
TCGA-CR-5248-01A-01D-2012-08 HNSC e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CR-5243-01A-01D-1512-08	HNSC	297e8b35-5b8b-4d5b-b812-86165f949a20
TCGA-CR-5249-01A-01D-1512-08 HNSC 42bf9ca3-47d8-45ff-bccf-bda80af58d22 TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-a1eaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CR-5247-01A-01D-2012-08	HNSC	3b5b07b4-29ef-4a55-b6ab-93352613f631
TCGA-CR-5250-01A-01D-1512-08 HNSC 49e54f5a-9b3a-47ff-b6cc-aleaf54fd136 TCGA-CR-6467-01A-11D-1870-08 HNSC 2a7f5a16-9330-45a1-9024-1cff1cdb5714 TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CR-5248-01A-01D-2012-08	HNSC	e5af63d7-e8b2-4a76-8b39-6ee652ad8e5f
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TCGA-CR-6470-01A-11D-1870-08 HNSC 30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5 TCGA-CR-6471-01A-11D-1870-08 HNSC c087e87f-867c-45dd-8645-5ab774e4827c	TCGA-CR-5250-01A-01D-1512-08	HNSC	49e54f5a-9b3a-47ff-b6cc-aleaf54fd136
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	TCGA-CR-6470-01A-11D-1870-08	HNSC	30bc4d1e-f0cb-44c5-a32c-b4b690cd6cc5
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TCGA-CR-6478-01A-11D-1870-08	HNSC	c21f40c6-4260-4def-8cca-1c11895b35b0
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TCGA-CR-6481-01A-11D-1870-08	HNSC	5e7d2531-81c1-48bb-9c0a-1867d1f83f92
TCGA-CR-6482-01A-11D-1870-08	HNSC	684bcd80-30fb-49e5-b72a-09502a9d1468
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TCGA-CR-7365-01A-11D-2012-08	HNSC	ec114413-a950-4e74-abc8-98857af8b9ad
TCGA-CR-7367-01A-11D-2012-08	HNSC	b82e34db-7b0e-4bbd-bc42-ba063ac42409
TCGA-CR-7368-01A-11D-2129-08	HNSC	4b194ab3-d213-4a7a-be46-909b4f0c7291
TCGA-CR-7369-01A-11D-2129-08	HNSC	f16a5c08-c9f8-442e-ba13-45681cacda40
TCGA-CR-7370-01A-11D-2129-08	HNSC	9f8ec337-85f7-4b01-a2b6-5db9a9e62f30
TCGA-CR-7371-01A-11D-2012-08	HNSC	68201be8-a1a9-4c78-ad99-3c767ca8366b
TCGA-CR-7372-01A-11D-2012-08	HNSC	9032c525-9bed-47f9-b9f2-ecce4593ea37
TCGA-CR-7373-01A-11D-2012-08	HNSC	9b1f5f6d-503c-4933-944a-b4fd1cc3fa93
TCGA-CR-7374-01A-11D-2012-08	HNSC	2cf33b63-464e-49a0-88f0-6a6d5b0393c4
TCGA-CR-7376-01A-11D-2129-08	HNSC	a6b11f68-79da-4542-818d-f404116c0bf8
TCGA-CR-7377-01A-11D-2012-08	HNSC	93e4eb9a-7643-411b-be90-94b801f23566
TCGA-CR-7379-01A-11D-2012-08	HNSC	8cc45c01-a363-4151-9ea0-32c404b79da4
TCGA-CR-7380-01A-11D-2012-08	HNSC	ac968fdd-970b-41fc-99f7-5670c741bc06
TCGA-CR-7382-01A-11D-2129-08	HNSC	fdde6828-b9f4-4648-a86b-157c5d46abb2
TCGA-CR-7383-01A-11D-2129-08	HNSC	203629ed-2791-4e22-a9da-be647b0cdef5
TCGA-CR-7385-01A-11D-2012-08	HNSC	2c00b622-c4a4-4862-b14a-a97b7261f46f
TCGA-CR-7386-01A-11D-2012-08	HNSC	dac99486-00bc-41ad-92b4-8bed1a28b122
TCGA-CR-7388-01A-11D-2012-08	HNSC	3eddb2ad-6c75-4ae7-9d27-8ec0e7b4aa55
TCGA-CR-7389-01A-11D-2012-08	HNSC	37149937-8131-4dbf-916b-d599d203eba7
TCGA-CR-7390-01A-11D-2012-08	HNSC	714399af-e425-43bb-a82a-b62ca6fd735d
TCGA-CR-7391-01A-11D-2012-08	HNSC	7236609c-34dd-425a-b882-2dff36983f7b
TCGA-CR-7392-01A-11D-2012-08	HNSC	0616d3e5-9641-4329-a65a-19f4c6918e1c
TCGA-CR-7393-01A-11D-2012-08	HNSC	f59ef1d2-2fc0-44a0-9d2f-c4efd9e79f5d
TCGA-CR-7394-01A-11D-2012-08	HNSC	1fe9a612-4c9a-432d-b175-e1d8bdbc7c56
TCGA-CR-7395-01A-11D-2012-08	HNSC	bd0b1b16-ee20-48e5-be11-70eac9c15630
TCGA-CR-7397-01A-11D-2012-08	HNSC	b93863c2-4657-4ca2-8fce-094fe5df163a
TCGA-CR-7398-01A-11D-2012-08	HNSC	12c391dc-3138-4e73-bdc7-b06512dd0fa7

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TCGA-CV-5436-01A-01D-1512-08 HNSC 34dc613e-e4b4-4897-ac4b-13ff46e46d7e	
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TCGA-CV-5440-01A-01D-1512-08 HNSC 5f5ba5a9-8089-4fe7-92e3-6c31c5fb32d4	
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TCGA-CV-5443-01A-01D-1512-08 HNSC 9d279797-4464-4ef5-8858-640978ccc258	
TCGA-CV-5444-01A-02D-1512-08 HNSC cf975479-131b-4b37-927e-cacb1f13e62d	
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TCGA-CV-5970-01A-11D-1683-08 HNSC a52dc15f-d06d-46ed-a73e-aa004a2a736a	
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TCGA-CV-6003-01A-11D-1683-08 HNSC 9a040a5e-3d2b-433a-9786-7c26b433c0c2	
TCGA-CV-6433-01A-11D-1683-08 HNSC 16b220fa-a554-43c9-85b0-315331e5ba6e	
TCGA-CV-6436-01A-11D-1683-08 HNSC a5214457-3a86-4b29-b116-3baaa0aa5099	
TCGA-CV-6441-01A-11D-1683-08 HNSC 22b32736-3b91-4542-affa-46fa90819e69	
TCGA-CV-6933-01A-11D-1912-08 HNSC 8ef4b02e-4d34-4d58-aa2d-65a7f73982d5	
TCGA-CV-6934-01A-11D-1912-08 HNSC f5abf385-0372-4faa-9558-8bf02381b68b	
TCGA-CV-6935-01A-11D-1912-08 HNSC fdc0ebce-5ba2-4c18-b594-50b33ef6d116	
TCGA-CV-6936-01A-11D-1912-08 HNSC 2d4bdd75-d967-40b2-b55d-99e59cc7e125	
TCGA-CV-6937-01A-11D-2012-08 HNSC 1c78a20e-150f-4c12-8abe-b941f90e730f	
TCGA-CV-6938-01A-11D-1912-08 HNSC b1dcb76e-b98f-4989-90a2-885e50d8174c	
TCGA-CV-6939-01A-11D-1912-08 HNSC e2e84cc1-2944-489e-be1b-0018a4e723e4	
TCGA-CV-6940-01A-11D-1912-08 HNSC 39f2e005-79f9-4c63-a6d6-0b378481a3ba	
TCGA-CV-6941-01A-11D-1912-08 HNSC 87071681-0058-4081-91f3-f689a150fc94	
TCGA-CV-6942-01A-21D-2012-08 HNSC c5409f12-e438-4979-b40e-120899c1fa15	
TCGA-CV-6943-01A-11D-1912-08 HNSC 4fa37ade-3451-406d-b0bb-e135e1591b70	
TCGA-CV-6945-01A-11D-1912-08 HNSC fcfc9b74-5b8a-45b7-97ca-4e477e941e7c	
TCGA-CV-6948-01A-11D-1912-08 HNSC 03eb2650-4b9f-46d2-b09f-378d8e919ae2	

TCGA-CV-695-01A-11D-1912-08			
TCGA-CV-6952-01A-11D-1912-08	TCGA-CV-6950-01A-11D-1912-08	HNSC	4a341860-44fb-493e-bd46-aeb6610842de
TCGA-CV-6953-01A-11D-1912-08	TCGA-CV-6951-01A-11D-1912-08	HNSC	9e1bf26c-6a68-44d2-aaa8-9af2f67828aa
TCGA-CV-6954-01A-11D-1912-08	TCGA-CV-6952-01A-11D-1912-08	HNSC	2d859062-3655-471e-b3dd-e6ff0671c076
TCGA-CV-6955-01A-11D-2012-08	TCGA-CV-6953-01A-11D-1912-08	HNSC	fb79f2be-3dec-4b5a-b5f3-e29e0fb05a98
TCGA-CV-6959-01A-11D-1912-08	TCGA-CV-6954-01A-11D-1912-08	HNSC	08f56645-763e-4864-a145-c0136dacd4f5
TCGA-CV-6959-01A-11D-1912-08	TCGA-CV-6955-01A-11D-2012-08	HNSC	f2c7fbe1-af36-4c42-b5ae-b9bf1e88fe36
TCGA-CV-6960-01A-41D-2012-08	TCGA-CV-6956-01A-21D-2012-08	HNSC	9ccee056-124e-40d5-a07d-c208765d8640
TCGA-CV-7099-01A-11D-2012-08 HNSC D52767d9-10b4-4ec4-9437-5a5186e284ca TCGA-CV-7099-01A-11D-2012-08 HNSC 125ccb76-bf8d-4ce7-a04c-4424d6da0322 TCGA-CV-7090-01A-11D-2012-08 HNSC 5c636c2d-f426-43a9-984d-b4455e4388e5 TCGA-CV-7091-01A-11D-2012-08 HNSC 5c636c2d-f426-43a9-984d-b4455e4388e5 TCGA-CV-7095-01A-21D-2012-08 HNSC 1888-1888-1888-1888-1888-1888-1888-188	TCGA-CV-6959-01A-11D-1912-08	HNSC	ff4cc4f1-9897-4d04-a3f6-c28a9b928b7a
TCGA-CV-7099-01A-11D-2012-08 HNSC 125ccb76-bf8d-4ce7-a04c-4424d6da0322 TCGA-CV-7090-01A-11D-2012-08 HNSC 5c636c2d-f426-43a9-984d-b4455e4388e5 TCGA-CV-7091-01A-11D-2012-08 HNSC 5c636c2d-f426-43a9-984d-b4455e4388e5 TCGA-CV-7097-01A-11D-2012-08 HNSC e4aba107-a048-46e5-b0aa-901076b6f61 TCGA-CV-7097-01A-11D-2012-08 HNSC 23336d44-b79y-4361-b661-ce26eae06692 TCGA-CV-7099-01A-41D-2012-08 HNSC 12a3de68-c814-4a18-a469-d7edc76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC f21a5e1f-84b8-4e6f-8230-03d31cc7e431 TCGA-CV-7100-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-lab46be0ddbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-be03-1057ae308329 TCGA-CV-7177-01A-11D-2012-08 HNSC e984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01a-21D-2012-08 HNSC e984165c-88ea-4840-a980-be818db16820 TCGA-CV-7180-01A-11D-2012-08 HNSC af30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC af30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7183-01A-11D-2012-08 HNSC af30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7236-01A-11D-2012-08 HNSC af306-4040-2976e-4971ed1724a9 TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-116-4fe3-8196-d837a9091038 TCGA-CV-7238-01A-11D-2012-08 HNSC dc220a9d-116-4fe3-8196-d837a9091038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e3-7185-4158-9e8-454446960b60 TCGA-CV-7238-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7247-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2	TCGA-CV-6960-01A-41D-2012-08	HNSC	750da72e-cabd-4b97-8160-8c4e39272b8b
TCGA-CV-7090-01A-11D-2012-08 HNSC 5c636c2d-f426-43a9-984d-b4455e4388e5 TCGA-CV-7091-01A-11D-2012-08 HNSC 563c5a89-6dad-467e-b2ea-e07677574a08 TCGA-CV-7095-01A-21D-2012-08 HNSC e4aba107-a048-46e5-b0aa-901f076b6f61 TCGA-CV-7099-01A-41D-2012-08 HNSC 12a04e68-c814-4a18-a469-d7ede76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC 12a6468-c814-4a18-a469-d7ede76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC 511a5a8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7101-01A-11D-2012-08 HNSC 511a5a8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7103-01A-21D-2012-08 HNSC eda5514f-3aa1-447e-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC eda5514f-3aa1-447e-ad07-55ec307c26e3 TCGA-CV-7104-01A-11D-2012-08 HNSC e04f3556-ac16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC e94f3556-ac16-410d-bc03-1057ae308329 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8-4057-abd1-a9dd1e49ec78 TCGA-CV-718-01A-21D-2012-08 HNSC 2984165c-88ea-4840-a980-be818db16820 TCGA-CV-718-01A-21D-2012-08 HNSC 3f30774f-2b8-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 2020a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7235-01A-11D-2012-08 HNSC 9e0f3e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7235-01A-11D-2012-08 HNSC 9e0f3e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e0f3eb-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f3eb-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f3eb-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f3eb-2-97e-492-a0da-270ba8-37a55a7a2c3 TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f3eb-2-97-e39-a38-a264-9037a48401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f3eb-2-97-a92-a0da-270ba8-37a55a7a2c3 TCGA-CV-7245-01A-11D-2012-08 HNSC 9e0f2eb-2-e97-a92-a0da-270ba8-37ba5a7a2c3 TCGA-CV-7255-01A-11D-2012-08 HNSC 9e0f2eb-2-e97-a92-a0da-270ba9-4e6e TCGA-CV-7255-01A-11D-2012-08 HNSC 9f3eb-2-438-a266-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 9f3eb-2-438-a56-b128e07d1963	TCGA-CV-6962-01A-11D-1912-08	HNSC	0b2767d9-10b4-4ec4-9437-5a5186e284ca
TCGA-CV-7091-01A-11D-2012-08 HNSC 563c5a89-6dad-467e-b2ea-e07677574a08 TCGA-CV-7095-01A-21D-2012-08 HNSC e4aba107-a048-46e5-b0aa-901f076b6f61 TCGA-CV-7099-01A-41D-2012-08 HNSC 23336d44-bb79-4361-b661-ce26eae06692 TCGA-CV-7099-01A-41D-2012-08 HNSC 12a604e8-e814-4a18-a469-d7edc76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC 12a5e1f-84b8-4e6f-8230-03d31cc7c431 TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC e94165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-11D-2012-08 HNSC e94165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9ddle49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7180-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7235-01A-11D-2012-08 HNSC 20209d-11f16-4fe3-8196-d837a909f038 TCGA-CV-7234-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC 9e07a1bc-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c484-8551-8d1db9072ee2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c484-8551-8d1db9072ee2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c948-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c948-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c98-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c98-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c98-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c98-8551-8d1db9072ee2 TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-4c98-8551-8d168-8d5-8d58d4451b0e TCGA-CV-7255-01A-11D-2012-08 HNSC bc6a2brc-8a6-bc1a-8abc-8	TCGA-CV-7089-01A-11D-2012-08	HNSC	125ccb76-bf8d-4ce7-a04c-4424d6da0322
TCGA-CV-7095-01A-21D-2012-08 HNSC e4aba107-a048-46e5-b0aa-901f076b6f61 TCGA-CV-7097-01A-11D-2012-08 HNSC 23336d44-bb79-4361-b661-ce26eae06692 TCGA-CV-7099-01A-41D-2012-08 HNSC 12a04e68-c814-4a18-a469-d7edc76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC f21a5e1f-84b8-4e6f-8230-03d31cc7c431 TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e043556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC e043556-ae16-410d-bc03-1057ae308329 TCGA-CV-7178-01A-11D-2012-08 HNSC e984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-718-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-718-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8bc-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC d220a9d-1f16-4fc3-8196-d837a909f038 TCGA-CV-7242-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7243-01A-11D-2012-08 HNSC 9e07albc-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC 9e07albc-17c7-4cb4-b3b1-92162a79de0e TCGA-CV-7240-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a8401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6b-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6b-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6b-4084-8551-8d1db9072ec2 TCGA-CV-7250-01A-11D-2012-08 HNSC bc6a2b7-8a6b-4084-8551-8d1db9072ec2 TCGA	TCGA-CV-7090-01A-11D-2012-08	HNSC	5c636c2d-f426-43a9-984d-b4455e4388e5
TCGA-CV-7097-01A-11D-2012-08 HNSC 23336d44-bb79-4361-b661-ce26cae06692 TCGA-CV-7099-01A-41D-2012-08 HNSC 12a04e68-c814-4a18-a469-d7edc76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC f21a5e1f-84b8-4e6f-8230-03d31cc7c431 TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ac308329 TCGA-CV-7104-01A-11D-2012-08 HNSC e984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-11D-2012-08 HNSC 2984165c-88ea-4840-a980-be818db16820 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7180-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9607a1bc-77c-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC	TCGA-CV-7091-01A-11D-2012-08	HNSC	563c5a89-6dad-467e-b2ea-e07677574a08
TCGA-CV-7099-01A-41D-2012-08 HNSC 12a04e68-e814-4a18-a469-d7edc76e362d TCGA-CV-7100-01A-11D-2012-08 HNSC f21a5e1f-84b8-4e6f-8230-03d31cc7c431 TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC 4f429401-f71e-4908-9663-2e66bacbebdd TCGA-CV-7177-01A-11D-2012-08 HNSC c984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7183-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7250-01A-11D-2012-08 HNS	TCGA-CV-7095-01A-21D-2012-08	HNSC	e4aba107-a048-46e5-b0aa-901f076b6f61
TCGA-CV-7100-01A-11D-2012-08 HNSC f21a5e1f-84b8-4e6f-8230-03d31cc7c431 TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7177-01A-11D-2012-08 HNSC e984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc62b1b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7248-01A-11D-2012-08 HN	TCGA-CV-7097-01A-11D-2012-08	HNSC	23336d44-bb79-4361-b661-ce26eae06692
TCGA-CV-7101-01A-11D-2012-08 HNSC 511c3fa8-476b-4ee8-8e93-1ab46bc40dbe TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC c984165c-88ea-4840-a980-be818db16820 TCGA-CV-7177-01A-11D-2012-08 HNSC d429401-f71e-4908-9663-2e66bacbebdd TCGA-CV-7178-01A-1D-2012-08 HNSC d233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7180-01A-11D-2012-08 HNSC d233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7236-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d44966b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87669 TCGA-CV-7245-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7245-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87669 TCGA-CV-7250-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87669 TCGA-CV-7250-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87669 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47de-4768-977b-bc3c1fe93722 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47de-4882b6-b128e07d1963 TCGA-CV-7250-01A-11D-2012-08 HNSC 1464ba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7250-01A-11D-2012-08 HNSC 19a07472-e8b9-4a34-b2cb-11ace3	TCGA-CV-7099-01A-41D-2012-08	HNSC	12a04e68-c814-4a18-a469-d7edc76e362d
TCGA-CV-7102-01A-11D-2012-08 HNSC eda5514f-3aa1-447c-ad07-55ec307c26e3 TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC 4f429401-f71e-4908-9663-2e66bacbebdd TCGA-CV-7177-01A-11D-2012-08 HNSC c984165c-88ea-4480-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9ddle49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7243-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7253-01A-11D-2012-08 HNS	TCGA-CV-7100-01A-11D-2012-08	HNSC	f21a5e1f-84b8-4e6f-8230-03d31cc7c431
TCGA-CV-7103-01A-21D-2012-08 HNSC e04f3556-ae16-410d-bc03-1057ae308329 TCGA-CV-7104-01A-11D-2012-08 HNSC 4f429401-f71e-4908-9663-2e66bacbebdd TCGA-CV-7177-01A-11D-2012-08 HNSC c984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7253-01A-11D-2012-08 HNS	TCGA-CV-7101-01A-11D-2012-08	HNSC	511c3fa8-476b-4ee8-8e93-1ab46bc40dbe
TCGA-CV-7104-01A-11D-2012-08 HNSC 4f429401-f71e-4908-9663-2e66bacbebdd TCGA-CV-7177-01A-11D-2012-08 HNSC c984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7247-01A-11D-2012-08 HNSC bc6e2b62-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC bcce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7255-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7253-01A-11D-2012-08 HNS	TCGA-CV-7102-01A-11D-2012-08	HNSC	eda5514f-3aa1-447c-ad07-55ec307c26e3
TCGA-CV-7177-01A-11D-2012-08 HNSC c984165c-88ea-4840-a980-be818db16820 TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fc3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7247-01A-11D-2012-08 HNSC bce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC bce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 4501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC<	TCGA-CV-7103-01A-21D-2012-08	HNSC	e04f3556-ae16-410d-bc03-1057ae308329
TCGA-CV-7178-01A-21D-2012-08 HNSC 3f30774f-2b8c-4057-abd1-a9dd1e49ec78 TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-frc7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7l9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7253-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7255-01A-11D-2012-08 HNS	TCGA-CV-7104-01A-11D-2012-08	HNSC	4f429401-f71e-4908-9663-2e66bacbebdd
TCGA-CV-7180-01A-11D-2012-08 HNSC 4233a363-ba28-495c-8590-644199c33d64 TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 3ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7253-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7254-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7255-01A-11D-2012-08 HNSC ddedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7256-01A-11D-2012-08 HNS	TCGA-CV-7177-01A-11D-2012-08	HNSC	c984165c-88ea-4840-a980-be818db16820
TCGA-CV-7183-01A-11D-2012-08 HNSC 172e7b30-829e-40b2-976e-4971cd1724a9 TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7248-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7250-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7261-01A-11D-2012-08 HNS	TCGA-CV-7178-01A-21D-2012-08	HNSC	3f30774f-2b8c-4057-abd1-a9dd1e49ec78
TCGA-CV-7235-01A-11D-2012-08 HNSC 1758147b-cb09-430b-a8cb-6a144744a79f TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7255-01A-11D-2012-08 HNSC ddedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7180-01A-11D-2012-08	HNSC	4233a363-ba28-495c-8590-644199c33d64
TCGA-CV-7236-01A-11D-2012-08 HNSC dc220a9d-1f16-4fe3-8196-d837a909f038 TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7183-01A-11D-2012-08	HNSC	172e7b30-829e-40b2-976e-4971cd1724a9
TCGA-CV-7238-01A-11D-2012-08 HNSC e9619e49-7185-4158-9e8b-45d446960b60 TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7235-01A-11D-2012-08	HNSC	1758147b-cb09-430b-a8cb-6a144744a79f
TCGA-CV-7242-01A-11D-2012-08 HNSC 9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7236-01A-11D-2012-08	HNSC	dc220a9d-1f16-4fe3-8196-d837a909f038
TCGA-CV-7243-01A-11D-2012-08 HNSC bc6a2b7c-8a6c-4084-8551-8d1db9072ec2 TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7238-01A-11D-2012-08	HNSC	e9619e49-7185-4158-9e8b-45d446960b60
TCGA-CV-7245-01A-11D-2012-08 HNSC 56291b3c-595c-4388-a264-9037a48401d8 TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7242-01A-11D-2012-08	HNSC	9e07a1bc-f7c7-4cb4-b3b1-92162a79de0e
TCGA-CV-7247-01A-11D-2012-08 HNSC b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3 TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7243-01A-11D-2012-08	HNSC	bc6a2b7c-8a6c-4084-8551-8d1db9072ec2
TCGA-CV-7248-01A-11D-2012-08 HNSC 8ffc7f9d-16da-4cff-b845-f2ff8df87569 TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7245-01A-11D-2012-08	HNSC	56291b3c-595c-4388-a264-9037a48401d8
TCGA-CV-7250-01A-11D-2012-08 HNSC 14516d2b-47dc-4768-977b-bc3c1fe93722 TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7247-01A-11D-2012-08	HNSC	b0ce56d2-8e2b-42b4-ac59-d37ba5a7a2c3
TCGA-CV-7252-01A-11D-2012-08 HNSC 9692c6b2-ce97-4c92-a0dd-f27d01a94e6e TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7248-01A-11D-2012-08	HNSC	8ffc7f9d-16da-4cff-b845-f2ff8df87569
TCGA-CV-7253-01A-11D-2012-08 HNSC d501a7e5-70e7-4f80-851a-efe8859d603a TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7250-01A-11D-2012-08	HNSC	14516d2b-47dc-4768-977b-bc3c1fe93722
TCGA-CV-7254-01A-11D-2012-08 HNSC fd22e861-571e-44da-82b6-b128e07d1963 TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7252-01A-11D-2012-08	HNSC	9692c6b2-ce97-4c92-a0dd-f27d01a94e6e
TCGA-CV-7255-01A-11D-2012-08 HNSC 4dedba61-e137-4ae4-8312-94231e3b1d16 TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7253-01A-11D-2012-08	HNSC	d501a7e5-70e7-4f80-851a-efe8859d603a
TCGA-CV-7261-01A-11D-2012-08 HNSC 9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7254-01A-11D-2012-08	HNSC	fd22e861-571e-44da-82b6-b128e07d1963
TCGA-CV-7263-01A-11D-2012-08 HNSC 19a07472-c8b9-4a34-b2cb-11ace35e7903	TCGA-CV-7255-01A-11D-2012-08	HNSC	4dedba61-e137-4ae4-8312-94231e3b1d16
	TCGA-CV-7261-01A-11D-2012-08	HNSC	9fa7bc79-d05b-41da-8bcc-8d5ad4451b0c
TCGA-CV-7406-01A-11D-2078-08 HNSC 8c9effa8-acb6-4db0-874a-8f0df386924c	TCGA-CV-7263-01A-11D-2012-08	HNSC	19a07472-c8b9-4a34-b2cb-11ace35e7903
	TCGA-CV-7406-01A-11D-2078-08	HNSC	8c9effa8-acb6-4db0-874a-8f0df386924c

TCGA-CV-7407-01A-11D-2078-08	HNSC	94631dc8-6dcb-49ed-bb68-e1a57a65f1cb
TCGA-CV-7409-01A-31D-2229-08	HNSC	47fa56f1-0802-403a-a644-913f1a0fdeca
TCGA-CV-7410-01A-21D-2078-08	HNSC	b89c4f94-b07c-485b-95ba-ffe815616d78
TCGA-CV-7411-01A-11D-2078-08	HNSC	790e387e-9e87-48d0-bc9d-2bc92f20abc5
TCGA-CV-7413-01A-11D-2078-08	HNSC	be482a19-0de0-4e60-a831-9ebe8545a6f3
TCGA-CV-7414-01A-11D-2078-08	HNSC	7137f980-5301-4b18-9664-d887eaced75e
TCGA-CV-7415-01A-11D-2078-08	HNSC	bb1e4188-130c-4206-8671-d7ce3eb8ee74
TCGA-CV-7418-01A-11D-2078-08	HNSC	25a70d04-f533-4e60-b9fc-e74d600db296
TCGA-CV-7421-01A-11D-2078-08	HNSC	ee675976-b447-48c8-bc67-6878a0d35e07
TCGA-CV-7422-01A-21D-2078-08	HNSC	5eb3f291-082c-48a8-b653-09264342adee
TCGA-CV-7423-01A-11D-2078-08	HNSC	a99653e0-2751-4423-93f7-abcf258c9868
TCGA-CV-7424-01A-11D-2078-08	HNSC	76d5fc22-fd06-43f6-94a8-943a09db5fd6
TCGA-CV-7425-01A-11D-2078-08	HNSC	f8cc6696-91d0-4eba-a765-ef7d044238ce
TCGA-CV-7427-01A-11D-2078-08	HNSC	3fdb4698-4a38-4a81-a403-d1ce5568c225
TCGA-CV-7429-01A-11D-2129-08	HNSC	14b42e59-e519-4efc-8105-6f6b83d33353
TCGA-CV-7430-01A-11D-2129-08	HNSC	29a4027f-4d4f-4133-b40a-3bfab6d2ac9e
TCGA-CV-7432-01A-11D-2129-08	HNSC	60da7e3f-4d9c-4cb3-856d-6cc02e381028
TCGA-CV-7433-01A-11D-2129-08	HNSC	15380da5-6a0b-4649-b21b-ce1ed7d61b67
TCGA-CV-7434-01A-11D-2129-08	HNSC	d64e4e80-e6c6-42c8-8bc6-0fafb6475c51
TCGA-CV-7435-01A-11D-2129-08	HNSC	16b7fd85-3664-4c4a-9a43-48b107dbcf7f
TCGA-CV-7437-01A-21D-2129-08	HNSC	53413980-80cc-4c73-8bb6-31a01d6df86e
TCGA-CV-7438-01A-21D-2129-08	HNSC	6fd3ecf3-c87c-46c3-81f0-11e2f8936d61
TCGA-CV-7440-01A-11D-2129-08	HNSC	901c2ed5-8348-4dd9-a84c-6c0b18d6525e
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TCGA-CX-7085-01A-21D-2012-08	HNSC	4f6ee10b-246d-49cd-8b60-01dcb175e634
TCGA-CX-7086-01A-11D-2078-08	HNSC	dfcb7c6e-b0f4-4557-9669-4c580d1093a0
TCGA-CX-7219-01A-11D-2012-08	HNSC	83f92af6-60ab-402e-8990-e1060ca3cc4c
TCGA-D6-6515-01A-21D-1870-08	HNSC	15c4d640-884c-4d55-897e-2f68314423fe
TCGA-D6-6516-01A-11D-1870-08	HNSC	5ab94b24-1a1f-4df7-a5c6-b1dce8ee9be5
TCGA-D6-6517-01A-11D-1870-08	HNSC	c553e4a2-cbea-43d6-8937-a48836856b5a
TCGA-D6-6823-01A-11D-1912-08	HNSC	e1f4d8ef-f24a-417b-bf22-c03cdb6b5275
TCGA-D6-6824-01A-11D-1912-08	HNSC	b658aa3f-0812-4812-8254-816d9a4d7c04
TCGA-D6-6825-01A-21D-1912-08	HNSC	01f44db3-84dc-4f96-888d-b0370bf582a5
TCGA-D6-6826-01A-11D-1912-08	HNSC	368030ac-f855-452a-a3d3-3698ab9a00dd
TCGA-D6-6827-01A-11D-1912-08	HNSC	059be8f9-9536-40c0-a751-5fe529a2f01f
TCGA-DQ-5624-01A-01D-1870-08	HNSC	01282192-5bb6-44d6-bbc7-33a42eba416b
TCGA-DQ-5625-01A-01D-1870-08	HNSC	4e042e1d-8604-484a-b229-94b85745a478
TCGA-DQ-5629-01A-01D-1870-08	HNSC	e748f828-0b80-47f3-aa92-fb3b2be0dcc2
TCGA-DQ-5630-01A-01D-1870-08	HNSC	5aa7ff44-d4ff-4163-81db-9f09bec8d5b0
TCGA-DQ-5631-01A-01D-1870-08	HNSC	e389975a-e588-48d4-9ed3-548e8ed9de1c
TCGA-DQ-7588-01A-11D-2078-08	HNSC	6aad9b01-6a99-4f21-955f-7938af25a188

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TCGA-DQ-7591-01A-11D-2078-08 HNSC 4068a2fc-452d-4b2c-88d8-72d30097527b	
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TCGA-F7-7848-01A-11D-2129-08 HNSC ba8a3e47-ee55-4c88-b29f-6d161ffae1d0	
TCGA-H7-7774-01A-21D-2078-08 HNSC 0eb5b79a-e3be-4b19-aef6-74247986aaf6	
TCGA-HD-7229-01A-11D-2012-08 HNSC 26b27991-540f-47f4-95f3-a59a493da593	
TCGA-HD-7753-01A-11D-2078-08 HNSC 7dc33525-6f57-4b12-9b72-c9c845296ae3	
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TCGA-HD-7831-01A-11D-2129-08 HNSC ae914215-3b1a-4edb-9f5a-ce4a17154178	
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TCGA-HD-7917-01A-11D-2229-08 HNSC 451948c9-3d16-4771-b006-28b98580db2c	
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TCGA-A3-3320-01A-01D-0966-08 RCC 5c4cc718-d7b5-453c-89d8-186ab0869e68	
TCGA-A3-3322-01A RCC 6f329d07-3308-4c84-9113-2bf000e9be3b	
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TCGA-A3-3326-01A-01D-0966-08 RCC 60ed222b-cd0c-4bc5-acd0-39f207be3289	
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TCGA-A3-3347-01A-02D-1386-10 RCC 2f4a6bd7-16ff-4689-b41d-c5fabb87823b	
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TCGA-A3-3370-01A-02D-1421-08 RCC 21ce7121-87b4-4686-9bf6-aff71d8b2223	
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TCGA-A3-3378-01A-01D-0966-08 RCC f04f3a00-e743-4fed-a0b0-e6a81bdd6ddd	
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TCGA-AK-3444-01A-01D-0966-08 RCC ea794170-156d-4251-b899-abfd60b213b0 TCGA-AK-3451-01A RCC 242777f6-a875-4072-9696-8d7f7d718906 TCGA-AK-3455-01A-01D-0966-08 RCC 3fbeeda4-a6c4-45a4-a963-dc6ca3f7e0ba TCGA-AK-3456-01A-02D-1386-10 RCC d36fe1be-96a5-4001-a95e-d499a6087146 TCGA-AK-3458-01A-01D-1501-10 RCC 0198f3c3-78f2-4c19-90d5-c77b74044ca2 TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4818-01A RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-AK-3451-01A RCC 242777f6-a875-4072-9696-8d7f7d718906 TCGA-AK-3455-01A-01D-0966-08 RCC 3fbeeda4-a6c4-45a4-a963-dc6ca3f7e0ba TCGA-AK-3456-01A-02D-1386-10 RCC d36fe1be-96a5-4001-a95e-d499a6087146 TCGA-AK-3458-01A-01D-1501-10 RCC 0198f3c3-78f2-4c19-90d5-c77b74044ca2 TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fc091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-AK-3455-01A-01D-0966-08 RCC 3fbeeda4-a6c4-45a4-a963-dc6ca3f7e0ba TCGA-AK-3456-01A-02D-1386-10 RCC d36fe1be-96a5-4001-a95e-d499a6087146 TCGA-AK-3458-01A-01D-1501-10 RCC 0198f3c3-78f2-4c19-90d5-c77b74044ca2 TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-AK-3456-01A-02D-1386-10 RCC d36fe1be-96a5-4001-a95e-d499a6087146 TCGA-AK-3458-01A-01D-1501-10 RCC 0198f3c3-78f2-4c19-90d5-c77b74044ca2 TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-AK-3458-01A-01D-1501-10 RCC 0198f3c3-78f2-4c19-90d5-c77b74044ca2 TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-AS-3778-01A-01D-0966-08 RCC 7b56e923-2bc5-4368-8e28-42649d3bf169 TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-B0-4700-01A-02D-1534-10 RCC 32cb433f-359c-44c3-b2df-d2a64df90175 TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-B0-4706-01A-01D-1501-10 RCC 040fdd9b-db76-4357-9aed-77a8cbde058d TCGA-B0-4710-01A RCC 6fc8cb4b-1dc0-46b8-ae80-7dbd022c9431 TCGA-B0-4712-01A-01D-1501-10 RCC 032b33f8-ff79-47de-8cb2-d744eab8bd1a TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
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TCGA-B0-4810-01A-01D-1501-10 RCC e014eeeb-c48e-42bb-a683-93299087a3cf TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
TCGA-B0-4811-01A-01D-1501-10 RCC a46182dc-2481-4911-9f6b-9532666f9f8c TCGA-B0-4815-01A-01D-1501-10 RCC fe091054-41d3-44fa-86a2-fad3ae58423f TCGA-B0-4816-01A RCC d05c3419-4164-4a69-8b11-ce1f5c29b5d4 TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
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TCGA-B0-4818-01A-01D-1501-10 RCC 213bf382-c2ca-45d4-95ae-329e6653620f	
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TCGA-B0-4827-01A-02D-1421-08 RCC 02f83f9a-4e4d-44f3-8d67-b4fc2d35102b	
TCGA-B0-4842-01A-02D-1421-08 RCC ae765ade-6a06-439c-a1cd-67222a70f44e	
TCGA-B0-4852-01A-01D-1501-10 RCC 28dbeb57-c919-4f91-aa3c-7b8f3809011e	
TCGA-B0-4945-01A-01D-1421-08 RCC 9fae377f-6c63-4f47-a769-a1396fb15f56	
TCGA-B0-5075-01A RCC 200819c3-826e-49a1-8824-6d4752e6eb6f	
TCGA-B0-5077-01A-01D-1462-08 RCC 587f2bd8-952a-4f31-98e7-7654c80b8a99	
TCGA-B0-5080-01A-01D-1501-10 RCC 9adf0a63-1d5c-403a-9e78-cb9d62a249a4	
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TCGA-B0-5096-01A-01D-1421-08 RCC 261de0a2-6006-4b3b-aac0-37d9b33840aa	
TCGA-B0-5097-01A-01D-1421-08 RCC 3af2978e-b892-4817-be05-39f020c06b5e	
TCGA-B0-5099-01A-01D-1421-08 RCC c3150136-ae55-49d0-9212-86728464167d	
TCGA-B0-5100-01A-01D-1421-08 RCC b20bd619-59c9-4e2a-8e64-7bb44eaa75ce	
TCGA-B0-5102-01A-01D-1421-08 RCC abea5e3e-705a-4d2c-b207-1ab43767a19b	

CCGA-B0-5104-01A			
TCGA-B0-5107-01A	TCGA-B0-5104-01A	RCC	ac2cfbde-9d62-49db-9a07-e8166003f10f
TCGA-B0-5108-01A-01D-1421-08 RCC d1d37af8-d2c3-4825-8e47-1a2e52e3aebb TCGA-B0-5110-01A RCC 38041aeb-60fe-4784-a548-fd04b5c0c5f8 TCGA-B0-5110-01A RCC 38041aeb-60fe-4784-a548-fd04b5c0c5f8 TCGA-B0-5113-01A-01D-1421-08 RCC 64b234e0-74fe-4537-4556-280e01fba09b TCGA-B0-5115-01A-01D-1421-08 RCC 672246b-5137-4456-84e8-54e541eec531 TCGA-B0-5115-01A-01D-1421-08 RCC 97421d06-b199-4246-b2da-8do9ba313335 TCGA-B0-5119-01A-02D-1421-08 RCC 414d47e7-41bb-4e83-8edf-703fa0a46f01 TCGA-B0-5120-01A-01D-1421-08 RCC 6ce58fbe-6742-4ade-84b0-cd025266e030 TCGA-B0-5120-01A-01D-1421-08 RCC a7751eb2-8545-490e-92d9-edb97755d288 TCGA-B0-5399-01A RCC a1dddbed-780-412a-b563-914f71e5e75d TCGA-B0-5309-01A RCC a1dddbed-780-412a-b563-914f71e5e75d TCGA-B0-5400-01A-01D-1501-10 RCC ea62bea0-a008-481e-8a91-a0f3a-9589255 TCGA-B0-5691-01A-11D-1534-10 RCC ac62bea0-a008-481e-8a91-a0f3a-9589255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb TCGA-B0-5693-01A T1D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb TCGA-B0-5693-01A T1D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb TCGA-B0-5693-01A T1D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb TCGA-B0-5695-01A RCC 86e4862-7405-105-1731-be06e52eadd TCGA-B0-5695-01A RCC 86e4862-7405-05-1731-be06e52eadd TCGA-B0-5695-01A RCC 86e4862-7405-05-1731-be06e52eadd TCGA-B0-5695-01A RCC 86e4862-7405-05-1731-be06e52eadd TCGA-B0-5695-01A RCC 86e4862-7405-405-5731-be06e52eadd TCGA-B0-5703-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfe4-257e8ea8fa66 TCGA-B0-5703-01A-11D-1534-10 RCC 80e5534-9-2172-43a7-9f52-ab7d0888429 TCGA-B0-5703-01A-11D-1534-10 RCC 80e55349-2172-43a7-9f52-ab7d0888429 TCGA-B0-5703-01A-11D-1534-10 RCC 80e5634-ee60-478b-9266-e390e7561892 TCGA-B0-5703-01A-11D-1534-10 RCC 80e5634-ee60-478b-9266-e390e7561892 TCGA-B0-5703-01A-11D-1534-10 RCC 80e5038-660-478b-9266-e390e7561892 TCGA-B0-5703-01A-11D-1534-10 RCC 80e50	TCGA-B0-5106-01A-01D-1421-08	RCC	c0e28603-7204-416d-ba3d-5377a38f677d
TCGA-B0-5109-01A-02D-1421-08 RCC 38041aeb-60fe-4784-a5d8-fd04b5c0c5f8 RCGA-B0-5113-01A-01D-1421-08 RCC 64b234e0-74f6-453f4-b5cb-286e01fba09b RCGA-B0-5113-01A-01D-1421-08 RCC f122b61c-4537-4456-84e8-54e541eec531 RCGA-B0-5115-01A-01D-1421-08 RCC f122b61c-4537-4456-84e8-54e541eec531 RCGA-B0-5119-01A-02D-1421-08 RCC 414447c7-41bb-4c83-8cdf-703fa0a4d6f01 RCGA-B0-5190-01A-02D-1421-08 RCC 414447c7-41bb-4c83-8cdf-703fa0a4d6f01 RCGA-B0-5120-01A-01D-1421-08 RCC a2751cb2-8545-490c-92d9-edb9775d32b8 RCGA-B0-5121-01A-02D-1421-08 RCC a1dddbed-780-412a-8563-91471e5c75d RCGA-B0-5399-01A RCC a1dddbed-780-412a-8563-91471e5c75d RCGA-B0-5399-01A RCC a1dddbed-780-412a-8563-91471e5c75d RCGA-B0-5400-01A-01D-1501-10 RCC ca62bea0-3008-818-8a91-405a9598255 RCGA-B0-5602-01A-01D-1501-10 RCC ca62bea0-3008-818-8a91-405a9598255 RCGA-B0-5692-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb RCGA-B0-5693-01A-11D-1534-10 RCC ar40135-8357-40b7-b711-478633a70997 RCGA-B0-5693-01A-11D-1534-10 RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5693-01A-11D-1534-10 RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5695-01A RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5696-01A-11D-1534-10 RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5696-01A-11D-1534-10 RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5696-01A-11D-1534-10 RCC debba05-b935-482-b070-8fc80ea6b609 RCGA-B0-5699-01A RCC debba05-b935-482-b076-8fc86841ed RCGA-B0-5699-01A RCC debba05-730-4b5-ace2-e2fofffaced9 RCGA-B0-5703-01A-11D-1534-10 RCC debba05-730-4b5-ace2-e2fofffaced9 RCGA-B0-5703-01A-11D-1534-10 RCC debba05-730-4b5-ace2-e2fofffaced9 RCGA-B0-5703-01A-11D-1534-10 RCC debba05-370-4b5-ace2-e2fofffaced9 RCC debba05-370-501A-11D-1534-10 RCC debba05-370-601A-11D-1534-10 RCC debba05-370-601A-11D-1534-10 RCC debba05-370-601A-11D-1534-10 RCC debba05-370-601A-38-408-8048-8096-808-808-808-808-808-808-808-808-808-80	TCGA-B0-5107-01A	RCC	4c6f4edb-9a29-48e6-8521-9c5fd2572e2d
TCGA-B0-5110-01A RCC 38041aeb-60fe-4784-a5d8-fd04b5c0c5f8 TCGA-B0-5113-01A-01D-1421-08 RCC 64b234e0-74fe-453f-b5cb-280e01fba09b TCGA-B0-5115-01A-01D-1421-08 RCC f122b61e-d537-4456-84e8-54e54leec531 TCGA-B0-5119-01A-02D-1421-08 RCC 97421d06-b199-4246-b2da-80abba313335 TCGA-B0-5119-01A-02D-1421-08 RCC 414d47c7-41bb-4c83-8cdf-703fa0a46f01 TCGA-B0-5120-01A-01D-1421-08 RCC 6cc88tbc-6742-4ade-84b0-cd02566c030 TCGA-B0-5121-01A-02D-1421-08 RCC ar751cb2-8545-490c-92d9-edb9775d32b8 TCGA-B0-5329-01A RCC ar1dddbed-c780-412a-b563-914f71e5c75d TCGA-B0-5402-01A-01D-1501-10 RCC c7128330-77b1-48bc-b970-be986aa63ea8 TCGA-B0-5402-01A-01D-1501-10 RCC cac2bea0-a008-481e-8a91-d07as9598255 TCGA-B0-569-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-7rc8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC be92ce16-6288-46c0-a3a7-7a27020cd7ca TCGA-B0-5693-01A-11D-1534-10 RCC dedba05-b935-4832-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ca6d TCGA-B0-5698-01A-11D-1534-10 RCC 48b270a76-b78-40b5-b73f-be0c6c52ca	TCGA-B0-5108-01A-01D-1421-08	RCC	d1d37af8-d2c3-4825-8e47-1a2e52e3acbb
TCGA-B0-5113-01A-01D-1421-08 RCC 64b234e0-74f6-453f-b5cb-280e01fba09b TCGA-B0-5115-01A-01D-1421-08 RCC f122b61c-d537-4456-84e8-54e54leec531 TCGA-B0-5119-01A-02D-1421-08 RCC 97421d06-b199-4246-b2da-80a9ba313335 TCGA-B0-5119-01A-02D-1421-08 RCC 414447c7-41bb-4c83-8cdf-703fa0a46f01 TCGA-B0-5121-01A-02D-1421-08 RCC 6ce58fbc-6742-4ade-84b0-cd025266e030 TCGA-B0-5121-01A-02D-1421-08 RCC a2751cb2-8545-490c-92d9-edb9775d32b8 TCGA-B0-5121-01A-02D-1421-08 RCC a1dddbed-780-412a-b563-91477le5c75d TCGA-B0-5399-01A RCC a1dddbed-780-412a-b563-91477le5c75d TCGA-B0-5400-01A-01D-1501-10 RCC ca62bea0-a008-481e-8a91-d078a9598255 TCGA-B0-5400-01A-01D-1501-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5693-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77a27020cd7ca TCGA-B0-5693-01A-11D-1534-10 RCC dedbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862e-7405-405-b73f-be0e6c52ea6d TCGA-B0-5699-01A-11D-1534-10 RCC 86e4862e-7405-405-b73f-be0e6c52ea6d TCGA-B0-5699-01A-11D-1534-10 RCC 9ca4e38-595-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5699-01A-11D-1534-10 RCC 9ca4e38-595-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5699-01A RCC 9ca4e38-395-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5700-01A-11D-1534-10 RCC 9ca4e38-995-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5700-01A-11D-1534-10 RCC 9ca4e38-995-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5700-01A-11D-1534-10 RCC 9ca4e38-995-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5700-01A-11D-1534-10 RCC 9ca4e38-840-998-986-ed90c7561892 TCGA-B0-5700-01A-11D-1534-10 RCC 9ca4e38-840-	TCGA-B0-5109-01A-02D-1421-08	RCC	58d6e408-ed00-4e1f-bffa-e73250cfe4a0
TCGA-B0-5115-01A-01D-1421-08 RCC	TCGA-B0-5110-01A	RCC	38041aeb-60fe-4784-a5d8-fd04b5c0c5f8
TCGA-B0-5116-01A	TCGA-B0-5113-01A-01D-1421-08	RCC	64b234e0-74f6-453f-b5cb-280e01fba09b
TCGA-B0-5119-01A-02D-1421-08 RCC 414447c7-41bb-4c83-8cdf-703fa0a46f01 TCGA-B0-5120-01A-01D-1421-08 RCC 6ce58fbc-6742-4ade-84b0-cd025266e030 TCGA-B0-5121-01A-02D-1421-08 RCC a2751cb2-8545-490c-92d9-edb9775d32b8 TCGA-B0-5399-01A RCC a1dddbed-c780-412a-b563-914f71e5c75d TCGA-B0-5400-01A-01D-1501-10 RCC ca62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5402-01A-01D-1501-10 RCC ac62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5692-01A-11D-1534-10 RCC ac62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5692-01A-11D-1534-10 RCC be92ce16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5692-01A-11D-1534-10 RCC be92ce16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6cdbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5694-01A-11D-1534-10 RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaccd9 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1534-10 RCC 9ca4e638-5a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5699-01A RCC 3c86554a9-2172-4a37-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 9c34e638-5a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5701-01A-11D-1534-10 RCC 9c34e638-3a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5701-01A-11D-1534-10 RCC 9c34e638-3a95-4ceb-bfc4-257e8ea8fa66 TCGA-B0-5701-01A-11D-1534-10 RCC 9c3400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5707-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5707-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5701-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5701-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5701-01A-11D-1669-08 RCC 12f1a370-c269-4b95-a89b-a1f3ac42e876 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1a370-c269-4b95-a89b-a1f3ac42e876 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1a370-c269-4b95-a89b-a1f3ac42e876 TCGA-B0-5710-01A-11D-1669-08 RCC 215275	TCGA-B0-5115-01A-01D-1421-08	RCC	f122b61c-d537-4456-84e8-54e541eec531
TCGA-B0-5120-01A-01D-1421-08 RCC 6ce58fbc-6742-4ade-84b0-cd025266e030 TCGA-B0-5121-01A-02D-1421-08 RCC a2751cb2-8545-490c-92d9-edb9775d32b8 TCGA-B0-5399-01A RCC a1dddbed-c780-412a-b563-91471e5c75d TCGA-B0-5400-01A-01D-1501-10 RCC e7128330-77b1-48be-b9f0-be986aa63ea8 TCGA-B0-5691-01A-11D-1530-10 RCC ea62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a7097 TCGA-B0-5693-01A-11D-1534-10 RCC deb92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5695-01A RCC 48b270af-07f2-4cb5-acc2-e2676ffaced9 TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-acc2-e2676ffaced9 TCGA-B0-5697-01A-11D-1534-10 RCC 2cddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5698-01A-11D-1534-10 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5701-01A-11D-1534-10 RCC 06553a-ee60-478b-9286-ed90c7561892 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d393-41a5-8c4	TCGA-B0-5116-01A	RCC	97421d06-b199-4246-b2da-80a9ba313335
TCGA-B0-5121-01A-02D-1421-08 RCC a2751cb2-8545-490c-92d9-edb9775d32b8 TCGA-B0-5399-01A RCC a1dddbed-c780-412a-b563-914t71e5c75d TCGA-B0-5400-01A-01D-1501-10 RCC e7128330-77b1-48be-b9f0-be986aa63ea8 TCGA-B0-5402-01A-01D-1501-10 RCC ca62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a7097 TCGA-B0-5693-01A-11D-1534-10 RCC de92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-acce-2e2foffaccd9 TCGA-B0-5697-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2dd12fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 9c34c638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5701-01A-11D-1534-10 RCC 06553a-ee60-478b-9286-ed90c7561892 TCGA-B0-5770-1A-11D-1534-10 RCC 963400a2-d939-41a5-8e2-9fc3a04b8362<	TCGA-B0-5119-01A-02D-1421-08	RCC	414d47c7-41bb-4c83-8cdf-703fa0a46f01
TCGA-B0-5399-01A RCC aldddbed-c780-412a-b563-914f71e5c75d TCGA-B0-5400-01A-01D-1501-10 RCC e7128330-77b1-48be-b9f0-be986aa63ea8 TCGA-B0-5402-01A-01D-1501-10 RCC e62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77e8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a70f97 TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-dcb5-acc2-e26f6ffaced9 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-2578ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2e-8fce6f8e41ed TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5706-01A-11D-1534-10 RCC b60c910-2d2e-483a-49be	TCGA-B0-5120-01A-01D-1421-08	RCC	6ce58fbc-6742-4ade-84b0-cd025266e030
TCGA-B0-5400-01A-01D-1501-10 RCC e7128330-77b1-48be-b9f0-be986aa63ea8 TCGA-B0-5402-01A-01D-1501-10 RCC ca62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a70f97 TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-br3f-be0c6c52ea6d TCGA-B0-5697-01A-11D-1534-10 RCC 9ca4e638-5a95-aeeb-bfc4-2578ea8fa66 TCGA-B0-5699-01A RCC 2ddf2fa6-7871-49fb-be2e-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC d3095df5-5466-49b8-9f6d-8a8916cca TCGA-B0-5700-01A-11D-1534-10 RCC b60cp10-242e-483a-a9de-c	TCGA-B0-5121-01A-02D-1421-08	RCC	a2751cb2-8545-490c-92d9-edb9775d32b8
TCGA-B0-5402-01A-01D-1501-10 RCC ca62bea0-a008-481e-8a91-d0f3a9598255 TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970eb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a7097 TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5695-01A RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaced9 TCGA-B0-5698-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1534-10 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 061c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC b76926-4b88-9f6d-f8ae891-6cca <td>TCGA-B0-5399-01A</td> <td>RCC</td> <td>a1dddbed-c780-412a-b563-914f71e5c75d</td>	TCGA-B0-5399-01A	RCC	a1dddbed-c780-412a-b563-914f71e5c75d
TCGA-B0-5691-01A-11D-1534-10 RCC ac2e1d29-e239-4dab-9d81-77c8d45970cb TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a70f97 TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaced9 TCGA-B0-5698-01A-11D-1534-10 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5698-01A-11D-1534-10 RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC b2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9	TCGA-B0-5400-01A-01D-1501-10	RCC	e7128330-77b1-48be-b9f0-be986aa63ea8
TCGA-B0-5692-01A-11D-1534-10 RCC laf40135-8357-40b7-b711-478633a70f97 TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaced9 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90c7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1534-10 RCC b60cf910-2de-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1669-08 RCC 12f1a370-c269-4895-a89	TCGA-B0-5402-01A-01D-1501-10	RCC	ca62bea0-a008-481e-8a91-d0f3a9598255
TCGA-B0-5693-01A-11D-1534-10 RCC be92ee16-6288-46c0-aaa7-7a27020cd7ca TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaced9 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90c7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-498-9f6d-f8ae8916cca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1669-08 RCC 121f370-c269-4995-a89b	TCGA-B0-5691-01A-11D-1534-10	RCC	ac2e1d29-e239-4dab-9d81-77c8d45970eb
TCGA-B0-5694-01A-11D-1534-10 RCC 6edbaa05-b935-4f82-b070-8fc80ea6b609 TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5695-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaccd9 TCGA-B0-5696-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-9	TCGA-B0-5692-01A-11D-1534-10	RCC	1af40135-8357-40b7-b711-478633a70f97
TCGA-B0-5695-01A RCC 86e4862c-7405-40b5-b73f-be0c6c52ea6d TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaccd9 TCGA-B0-5697-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5699-01A RCC 08c554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90c7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5709-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeacebe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30e	TCGA-B0-5693-01A-11D-1534-10	RCC	be92ee16-6288-46c0-aaa7-7a27020cd7ca
TCGA-B0-5696-01A-11D-1534-10 RCC 48b270af-07f2-4cb5-ace2-e2676ffaccd9 TCGA-B0-5697-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fee6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110dac0	TCGA-B0-5694-01A-11D-1534-10	RCC	6edbaa05-b935-4f82-b070-8fc80ea6b609
TCGA-B0-5697-01A-11D-1534-10 RCC 9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66 TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5705-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5812-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 21527594-ed75-4654-9caf-83d31f24	TCGA-B0-5695-01A	RCC	86e4862c-7405-40b5-b73f-be0c6c52ea6d
TCGA-B0-5698-01A-11D-1669-08 RCC 2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC 2f35dbf4-3223-494e-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B2-3924-01A RCC 6327ce2e-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC 6242adb8-db67-475e-a0e4-52a622666b12	TCGA-B0-5696-01A-11D-1534-10	RCC	48b270af-07f2-4cb5-ace2-e2676ffaccd9
TCGA-B0-5699-01A RCC 086554a9-2172-43a7-9f52-aab7d0888429 TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110dac0c7e TCGA-B2-4099-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973	TCGA-B0-5697-01A-11D-1534-10	RCC	9ca4e638-5a95-4eeb-bfc4-257e8ea8fa66
TCGA-B0-5701-01A-11D-1534-10 RCC 0e1c563a-ee60-478b-9286-ed90e7561892 TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5709-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5698-01A-11D-1669-08	RCC	2ddf2fa6-7871-49fb-be2c-8fce6f8e41ed
TCGA-B0-5702-01A-11D-1534-10 RCC 780b3f3e-1c49-40de-9131-65c4df9ebba6 TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5709-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de	TCGA-B0-5699-01A	RCC	086554a9-2172-43a7-9f52-aab7d0888429
TCGA-B0-5703-01A-11D-1534-10 RCC 963400a2-d939-41a5-8c42-9fc3a04b8362 TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5709-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5701-01A-11D-1534-10	RCC	0e1c563a-ee60-478b-9286-ed90e7561892
TCGA-B0-5705-01A-11D-1534-10 RCC d3095df5-5466-4b98-9f6d-f8ae8916ccca TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5702-01A-11D-1534-10	RCC	780b3f3e-1c49-40de-9131-65c4df9ebba6
TCGA-B0-5706-01A-11D-1534-10 RCC b60cf910-2d2e-483a-a9de-ce1e5f8d3825 TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5703-01A-11D-1534-10	RCC	963400a2-d939-41a5-8c42-9fc3a04b8362
TCGA-B0-5707-01A-11D-1534-10 RCC eb2f9f38-bce2-4746-a3c8-40abc3379b32 TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5705-01A-11D-1534-10	RCC	d3095df5-5466-4b98-9f6d-f8ae8916ccca
TCGA-B0-5709-01A-11D-1534-10 RCC bfeaecbe-7148-4642-b69a-b908a248f328 TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5706-01A-11D-1534-10	RCC	b60cf910-2d2e-483a-a9de-ce1e5f8d3825
TCGA-B0-5710-01A-11D-1669-08 RCC 12f1e370-c269-4b95-a89b-a1f3ae42e876 TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5707-01A-11D-1534-10	RCC	eb2f9f38-bce2-4746-a3c8-40abc3379b32
TCGA-B0-5711-01A-11D-1669-08 RCC cf09ae91-5523-494c-8f30-c26f6ba37624 TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5709-01A-11D-1534-10	RCC	bfeaecbe-7148-4642-b69a-b908a248f328
TCGA-B0-5713-01A-11D-1669-08 RCC 2f35dbf4-3223-4550-951b-1409a30ece68 TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5710-01A-11D-1669-08	RCC	12f1e370-c269-4b95-a89b-a1f3ae42e876
TCGA-B0-5812-01A-11D-1669-08 RCC 6327ce2c-8a24-45b9-9577-7b7d7b603e68 TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5711-01A-11D-1669-08	RCC	cf09ae91-5523-494c-8f30-c26f6ba37624
TCGA-B2-3924-01A RCC 21527594-ed75-4654-9caf-83d31f248e67 TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5713-01A-11D-1669-08	RCC	2f35dbf4-3223-4550-951b-1409a30ece68
TCGA-B2-4098-01A RCC 6463ae73-a885-4d69-9345-7110ddac0c7e TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B0-5812-01A-11D-1669-08	RCC	6327ce2c-8a24-45b9-9577-7b7d7b603e68
TCGA-B2-4099-01A RCC e242adb8-db67-475e-a0e4-52a622666b12 TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B2-3924-01A	RCC	21527594-ed75-4654-9caf-83d31f248e67
TCGA-B2-4101-01A RCC a9947b6c-dbc7-4ba5-af61-7647e11e2973 TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B2-4098-01A	RCC	6463ae73-a885-4d69-9345-7110ddac0c7e
TCGA-B4-5377-01A-01D-1501-10 RCC a615b02d-fd18-47ef-bd66-6dba56de6981	TCGA-B2-4099-01A	RCC	e242adb8-db67-475e-a0e4-52a622666b12
	TCGA-B2-4101-01A	RCC	a9947b6c-dbc7-4ba5-af61-7647e11e2973
TCGA-B8-4143-01A-01D-1806-10 RCC bb186c78-1052-48ec-97f4-c94bddf0df72	TCGA-B4-5377-01A-01D-1501-10	RCC	a615b02d-fd18-47ef-bd66-6dba56de6981
	TCGA-B8-4143-01A-01D-1806-10	RCC	bb186c78-1052-48ec-97f4-c94bddf0df72

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TCGA-B8-4148-01A-02D-1386-10	RCC	fe752e2b-e694-4fa9-99d6-46d5bff9e8cf
TCGA-B8-4151-01A-01D-1806-10	RCC	3f847558-8bc7-49b0-899d-2a7b8f0e3d1a
TCGA-B8-4153-01B-11D-1669-08	RCC	a66078d8-a6b2-4dc4-bfa3-def5a2e4504f
TCGA-B8-4154-01A-01D-1251-10	RCC	e48f5c14-4b64-4d4b-8273-bebc74182181
TCGA-B8-4620-01A	RCC	e4ec1484-4f77-4520-9ff5-bc4dc8a0fb15
TCGA-B8-4621-01A	RCC	242a72ad-5968-4bbf-936d-75b398a61b96
TCGA-B8-4622-01A	RCC	1c86e0f6-a019-47a5-8325-bbb82f76488c
TCGA-B8-5158-01A-01D-1421-08	RCC	9d730534-98e7-464e-945c-5964cec5362a
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TCGA-B8-5164-01A	RCC	471ce542-e85b-4bdb-b365-4562a93ef1e5
TCGA-B8-5165-01A-01D-1421-08	RCC	d1579785-5c42-4bda-9825-15ead235f7f4
TCGA-B8-5545-01A-01D-1669-08	RCC	514d2342-64ba-4c9f-9866-63bdbc26fda3
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TCGA-B8-5552-01B-11D-1669-08	RCC	13b52e49-20df-4e39-9dc9-cf8f7c157bd7
TCGA-B8-5553-01A-01D-1534-10	RCC	7c19e63c-770b-4289-aa47-9b2cf261b4ca
TCGA-BP-4161-01A	RCC	154de511-2bba-4959-970b-6a8429f29793
TCGA-BP-4162-01A	RCC	ca4eac28-22c9-48d8-8139-7cda2cfe4ae2
TCGA-BP-4163-01A	RCC	e44de28c-bce0-471d-bd4c-bea710f7c3cc
TCGA-BP-4164-01A	RCC	a8fab76e-ae69-43d6-972b-5837aec668fd
TCGA-BP-4167-01A-02D-1386-10	RCC	79b810e1-4de4-496d-9f70-ab62246e781b
TCGA-BP-4770-01A-01D-1501-10	RCC	aecbc5db-f75a-42d0-a84d-aa0369b08eec
TCGA-BP-4782-01A	RCC	a6c21bf2-dd9b-4243-863e-9d53b056666f
TCGA-BP-4801-01A-02D-1421-08	RCC	d3e62cb1-5ced-42cb-a360-479ee01877aa
TCGA-BP-4960-01A-01D-1462-08	RCC	36d21be3-2f46-47af-84aa-2305f2513aa1
TCGA-BP-4961-01A	RCC	f207131d-8db7-464b-a3e5-44218da1cafc
TCGA-BP-4962-01A-01D-1462-08	RCC	3454a6fe-2547-4531-a0be-cb27c1879e72
TCGA-BP-4963-01A-01D-1462-08	RCC	154bfa5d-0d9a-40c6-a2a5-bde1054702c3
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TCGA-BP-4967-01A-01D-1462-08	RCC	75866d14-47d5-4560-a5a0-32ba3e15ac63
TCGA-BP-4968-01A-01D-1462-08	RCC	d777d5ec-4632-446e-aeac-8ae3e5273fe2
TCGA-BP-4970-01A-01D-1462-08	RCC	205e81c6-235a-450f-b1f8-80c518eb3478
TCGA-BP-4971-01A-01D-1462-08	RCC	c07945e8-8133-4237-9d1f-18c023bc9d2c
TCGA-BP-4972-01A-01D-1462-08	RCC	b2da5d39-33f6-4807-9d1d-92b7cef2a8df
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TCGA-BP-4974-01A	RCC	a75c92b2-c67b-42b5-a8c2-7eea1b567ed0
TCGA-BP-4975-01A-01D-1462-08	RCC	109d2752-17f8-4b00-a61f-dfd8e2e3ca81
TCGA-BP-4976-01A-01D-1462-08	RCC	95bd81ec-3c06-4c4d-9915-5cc3dd7a7155
TCGA-BP-4977-01A-01D-1462-08	RCC	7c3bf7c1-07d9-4540-9a5e-614fd60b63ec
TCGA-BP-4981-01A-01D-1462-08	RCC	64a1f085-50cc-4129-a617-e0f691a58039

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TCGA.BP-4987-01A-01D-1462-08 RCC 7924f8if-8e78-4910-9dc5-db14d5ee7011 TCGA.BP-4988-01A-01D-1462-08 RCC 79269867-ecea-4520-bbb7-5dabc290664f TCGA.BP-4989-01A-01D-1462-08 RCC 7966085b-ed5b-ed61-8957-a6adcf7e818a TCGA.BP-4989-01A-01D-1462-08 RCC d54c714e-b1e4-4669-986d-5e13d2fc3cc3 TCGA.BP-4993-01A-01D-1462-08 RCC 212717dd-25f1-4c76-a648-b8a7d65eaecf TCGA.BP-4993-01A-01D-1462-08 RCC 34315bea-6ef2-42ec-b17e-c73eed40647f TCGA.BP-4993-01A-01D-1462-08 RCC 93b9afae-e12e-4942-96ae-274da6581d76 TCGA.BP-4998-01A-01D-1462-08 RCC e646f930-967b-43a3-bd70-184e5c38efe5 TCGA.BP-4999-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322e4d01 TCGA.BP-5000-01A-01D-1462-08 RCC b8816eaa-3c60-4fbf-abd6-6d869ca9eca7 TCGA.BP-5000-01A-01D-1462-08 RCC e368d35-0382-4979-b599-033a06a1f50b TCGA.BP-5000-01A-01D-1462-08 RCC e368d35-0382-4979-b599-033a06a1f50b TCGA.BP-5000-01A-01D-1462-08 RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA.BP-5000-01A-01D-1462-08 RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA.BP-5009-01A-01D-1462-08 RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA.BP-5009-01A-01D-1462-08 RCC 3baa3edc-c63e-4556-baf1-c3b03175b0fa TCGA.BP-5009-01A-01D-1462-08 RCC 3baa3edc-c63e-4556-baf1-c3b03175b0fa TCGA.BP-5168-01A-01D-1421-08 RCC 3baa3edc-c63e-4556-baf1-c3b03175b0fa TCGA.BP-5168-01A-01D-1421-08 RCC 3525be18-66d3-4b3-4-576-96a6dd2e6043 TCGA.BP-5169-01A-01D-1429-08 RCC 3525be18-66b9-4adf-9b60-955f79ed0f11 TCGA.BP-5173-01A-01D-1429-08 RCC 36263fc-094e-4128-8a59-560994433aa TCGA.BP-5173-01A-01D-1429-08 RCC 36263fc-094e-4128-8a59-60907d1c13f TCGA.BP-5173-01A-01D-1429-08 RCC 36263fc-094e-4128-8a59-60907d1c13f TCGA.BP-5173-01A-01D-1429-08 RCC 36263fc-094e-4128-8a59-60907d1c13f TCGA.BP-5173-01A-01D-1429-08 RCC 36263fc-094e-4128-8a59-60907d1c13f TCGA.BP-5182-01A-01D-1429-08 RCC 676648-716-748-55-6609-3347dadaaae TCGA.BP-5184-01A-01D-1429-08	TCGA-BP-4985-01A-01D-1462-08	RCC	e56acfea-aec6-4102-8fe0-25df396c10ae
TCGA-BP-4988-01A-01D-1462-08 RCC 792e9867-ceea-4520-bbb7-5dabe290664f TCGA-BP-4989-01A-01D-1462-08 RCC 7096085b-ed5b-4cd1-8957-a6adct7c818a TCGA-BP-4991-01A-01D-1462-08 RCC d54c714c-b1c4-4669-986d-5c13d2fc3cc3 TCGA-BP-4992-01A RCC 212717da-25f1-4c76-a648-88a7d65seacef TCGA-BP-4993-01A-01D-1462-08 RCC 34315bea-6ef2-42ec-b17e-c73ec4d0647f TCGA-BP-4993-01A-01D-1462-08 RCC 93b9afac-e12e-49d2-96ac-274da6581d76 TCGA-BP-4999-01A-01D-1462-08 RCC 86fb814-7c65-426b-67b5-7250322c4d01 TCGA-BP-4999-01A-01D-1462-08 RCC 86fb814-7c65-426b-67b5-7250322c4d01 TCGA-BP-5000-01A-01D-1462-08 RCC 863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC 863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-38730994f3 TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-38730994f3 TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-38730994f3 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1420-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 353cbal8-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5173-01A-01D-1420-08 RCC 3527b21e-972b-4231-356-8394ce0c500 TCGA-BP-5173-01A-01D-1420-08 RCC 3527b21e-972b-4231-356-8394ce0c500 TCGA-BP-5173-01A-01D-1420-08 RCC 356586-6b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-099-4af1-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-099-4af1-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-040-4af0-960-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-040-4af0-960-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-040-4af0-960-955f79ed0f11 TCGA-BP-5173-01A-01D-1420-08 RCC 36256-040-4af0-960-955f79ed0f11 TCGA-BP-5180-01A-01D-1420-08 RCC 36256-040-3	TCGA-BP-4986-01A-01D-1462-08	RCC	4465171a-d048-4078-b1ae-021b2c635ff4
TCGA-BP-4989-01A-01D-1462-08 RCC T096085b-cd5b-4cd1-8957-a6adc47c818a TCGA-BP-4991-01A-01D-1462-08 RCC d54c714e-b1c4-4669-986d-5e13d2fc3cc3 TCGA-BP-4992-01A RCC 212717dd-25f1-4c76-a648-b8a7d65casecf TCGA-BP-4993-01A-02D-1421-08 RCC 34315bea-6ef2-42ec-b17e-c73ecd40647f TCGA-BP-4993-01A-01D-1462-08 RCC 93b9afac-e12e-49d2-96ac-274da655k1d76 TCGA-BP-4998-01A-01D-1462-08 RCC e646f930-967b-43a3-bd70-184e5c38efe5 TCGA-BP-4999-01A-01D-1462-08 RCC e646f930-967b-43a3-bd70-184e5c38efe5 TCGA-BP-4999-01A-01D-1462-08 RCC 86fb814-7c65-426b-b7b5-7250322c4d01 TCGA-BP-5000-01A-01D-1462-08 RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5000-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5004-01A-01D-1462-08 RCC a4deb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5006-01A-01D-1462-08 RCC a4deb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC a4deb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC a4deb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-e63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-e63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 3baa3cdc-e63e-4556-baf1-c3b03175b0fa TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5170-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0500 TCGA-BP-5173-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c39b-6c9500 TCGA-BP-5173-01A-01D-1429-08 RCC 35058f8d-f3d-fac-fac-9e-fac-b0c907d1c13f TCGA-BP-5177-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5170-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5170-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-518-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-518-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-518-01A-01D-1429-08 RCC 30e58fa1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-518-01A-01D-1	TCGA-BP-4987-01A-01D-1462-08	RCC	7924f8ff-8e78-4910-9dc5-db14d5ee7011
TCGA-BP-4991-01A-01D-1462-08 RCC d54c714e-b1c4-d669-986d-5c13d2fc3cc3 TCGA-BP-4992-01A RCC 212717dd-25f1-4c76-a648-b8a7d65caccf TCGA-BP-4993-01A-02D-1421-08 RCC 34315bca-6cf2-42cc-b17e-c73ecd40647f TCGA-BP-4998-01A-01D-1462-08 RCC 93b9afac-e12e-49d2-96ac-274da6581d76 TCGA-BP-4998-01A-01D-1462-08 RCC e646930-967b-43a3-bd70-184e5c38efc5 TCGA-BP-5090-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322c4d01 TCGA-BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA-BP-5000-01A RCC e3d82fc4-b491-4172-86da-429cf16508de TCGA-BP-5000-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde3-387309e94f3 TCGA-BP-5000-01A-01D-1462-08 RCC 11fb962b-b4b8-4fc4-bde3-387309e94f3 TCGA-BP-5000-01A RCC a4deb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-1462-08 RCC 3ba3a5dc-628-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-142-08 RCC 3ba3a5dc-628-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-142-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-142-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 <td>TCGA-BP-4988-01A-01D-1462-08</td> <td>RCC</td> <td>792c9867-ceea-4520-bbb7-5dabe290664f</td>	TCGA-BP-4988-01A-01D-1462-08	RCC	792c9867-ceea-4520-bbb7-5dabe290664f
TCGA-BP-4992-01A	TCGA-BP-4989-01A-01D-1462-08	RCC	7096085b-cd5b-4cd1-8957-a6adcf7e818a
TCGA-BP-4993-01A-02D-1421-08 RCC 34315bea-6ef2-42ec-b17e-c73eed40647f TCGA-BP-4995-01A-01D-1462-08 RCC 93b9afac-e12e-49d2-96ac-274da6581d76 TCGA-BP-4998-01A-01D-1462-08 RCC e646930-967b-43a3-bd70-184e5c38efe5 TCGA-BP-4999-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322e4d01 TCGA-BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA-BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429ef16508de TCGA-BP-5004-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3i87309e94f3 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8490-338b66efefc1 TCGA-BP-5008-01A RCC 3ba32cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1420-08 RCC 3ba32cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1421-08 RCC 553cbe18-6dd3-4934-h7fe-96a6dd2e6943 TCGA-BP-5169-01A-01D-1421-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5179-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955179ed0f11 TCGA-BP-5178-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-9570rde13	TCGA-BP-4991-01A-01D-1462-08	RCC	d54c714e-b1c4-4669-986d-5e13d2fc3cc3
TCGA-BP-4995-01A-01D-1462-08 RCC 93b9afac-e12e-49d2-96ac-274da6581d76 TCGA-BP-4998-01A-01D-1462-08 RCC e646f930-967b-43a3-bd70-184e5c38efe5 TCGA-BP-4999-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322c4d01 TCGA-BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA-BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5004-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-1462-08 RCC 3ba32dc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1421-08 RCC 3ba32dc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5168-01A-01D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5179-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-9557ped0f11 TCGA-BP-5175-01A-01D-1429-08 RCC 30c58a1e-c7db-43ce-a7e-8-a1d21fd148	TCGA-BP-4992-01A	RCC	212717dd-25f1-4c76-a648-b8a7d65caecf
TCGA-BP-4998-01A-01D-1462-08 RCC e646f930-967b-43a3-bd70-184e5c38efe5 TCGA-BP-4999-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322c4d01 TCGA-BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA-BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 34aeb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1421-08 RCC 3baa3cdc-c63e-44556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6d3d-4b34-b7fc-96a6d2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 3527b21e-972b-4c31-b3de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b3de-8c394ce0e500 TCGA-BP-5175-01A-01D-1429-08 RCC 3687b2-66b9-4adf-9b60-95579ed0f11 TCGA-BP-5175-01A-01D-1429-08 RCC 30c9a5fc-09ae-412a-8a5b-56d9-44f433	TCGA-BP-4993-01A-02D-1421-08	RCC	34315bea-6ef2-42ec-b17e-c73eed40647f
TCGA.BP-4999-01A-01D-1462-08 RCC 86ffb814-7c65-426b-b7b5-7250322c4d01 TCGA.BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA.BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA.BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA.BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA.BP-5007-01A RCC 41c094e9-6c23-4993-8d90-338b66efefe1 TCGA.BP-5008-01A RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA.BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA.BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7ic-96a6dd2e6943 TCGA.BP-5168-01A-01D-1421-08 RCC 930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA.BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-dc31-b5de-8c394ce0c500 TCGA.BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f1 TCGA.BP-5173-01A-01D-1429-08 RCC 3b5c8a1e-e7db-43ce-a7e8-a1fd21f4438a TCGA-BP-5175-01A-01D-1429-08 RCC 30c58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5175-01A-01D-1429-08 RCC 30c58a1e-e7db-43ce-a7e8-a1fd21f443	TCGA-BP-4995-01A-01D-1462-08	RCC	93b9afac-e12e-49d2-96ac-274da6581d76
TCGA-BP-5000-01A-01D-1462-08 RCC b9816eaa-3c60-4fbf-abd6-6d869ca9cca7 TCGA-BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5009-01A-01D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 930560d-22e6-43aa-a6f0-02515f7a8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b56e-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 3ce0a5fe-09ac-412a-8a5b-56d9a44433aa TCGA-BP-5173-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-9lec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30c58a1e-e7db-43ce-a7e-a1fd21f4438e TCGA-BP-5175-01A-01D-1429-08 RCC 607eb48b-1647-4a35-ac60-f650341e304 TCGA-BP-5178-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f650341e304<	TCGA-BP-4998-01A-01D-1462-08	RCC	e646f930-967b-43a3-bd70-184e5c38efe5
TCGA-BP-5001-01A RCC e863bd35-0382-4979-b599-033a06a1f50b TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0e907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5178-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5178-01A-01D-1429-08 RCC 60888d5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 60888d5-1408-4bfb-bf27-f3e22f5488	TCGA-BP-4999-01A-01D-1462-08	RCC	86ffb814-7c65-426b-b7b5-7250322c4d01
TCGA-BP-5004-01A-01D-1462-08 RCC e3d82fe4-b491-4172-86da-429cf16508de TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-b	TCGA-BP-5000-01A-01D-1462-08	RCC	b9816eaa-3c60-4fbf-abd6-6d869ca9cca7
TCGA-BP-5006-01A-01D-1462-08 RCC 11fb962b-b4b8-46f4-bde4-3f87309e94f3 TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ac-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 30e58a1e-c7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-c7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5178-01A-01D-1429-08 RCC 608e8adcs-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 608e8adcs-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 608e8adcs-1408-4bf	TCGA-BP-5001-01A	RCC	e863bd35-0382-4979-b599-033a06a1f50b
TCGA-BP-5007-01A RCC a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5 TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5173-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c503414304 TCGA-BP-5177-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c503414304 TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4b1b-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4b1b-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 00523547-da1c-4b1a-a	TCGA-BP-5004-01A-01D-1462-08	RCC	e3d82fe4-b491-4172-86da-429cf16508de
TCGA-BP-5008-01A RCC 41c094e9-6c23-4993-8d90-338b66efefc1 TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5183-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC ddebed14	TCGA-BP-5006-01A-01D-1462-08	RCC	11fb962b-b4b8-46f4-bde4-3f87309e94f3
TCGA-BP-5009-01A-01D-1462-08 RCC 3baa3cdc-c63e-4556-baf1-c3b03175b0fa TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-ac6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5173-01A-01D-1429-08 RCC 30c58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5175-01A-01D-1429-08 RCC 607eb48b-1647-4c35-ac60-f6c50341e304 TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4c35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a7f6bde5-7503-459e-8419-dc0d744a651e TCGA-BP-5183-01A-01D-1429-08 RCC ddebd14-f47f-de6-ac39-c74ed3363211 TCGA-BP-518-01A-01D-1429-08 RCC ddebd14-f47f-de6-ac39-c74ed3363211 TCGA-BP-5180-01A-01D-1429-08 RCC	TCGA-BP-5007-01A	RCC	a44eb1d6-3b5c-42e8-b17a-d71ffc0503d5
TCGA-BP-5010-01A-02D-1421-08 RCC 553cbe18-6dd3-4b34-b7fe-96a6dd2e6943 TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5178-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5180-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5183-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5186-01A-01D-1429-08 RCC	TCGA-BP-5008-01A	RCC	41c094e9-6c23-4993-8d90-338b66efefc1
TCGA-BP-5168-01A-01D-1421-08 RCC 9930560d-22e6-43aa-a6f0-02515f7af8f0 TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690	TCGA-BP-5009-01A-01D-1462-08	RCC	3baa3cdc-c63e-4556-baf1-c3b03175b0fa
TCGA-BP-5169-01A-01D-1429-08 RCC 3527b21e-972b-4c31-b5de-8c394ce0e500 TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC d2bed6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5189-01A-02D-1429-08 RCC	TCGA-BP-5010-01A-02D-1421-08	RCC	553cbe18-6dd3-4b34-b7fe-96a6dd2e6943
TCGA-BP-5170-01A-01D-1429-08 RCC 68761b2c-66b9-4adf-9b60-955f79ed0f11 TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5189-01A-02D-1429-08 RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5190-01A-01D-1429-08 RCC	TCGA-BP-5168-01A-01D-1421-08	RCC	9930560d-22e6-43aa-a6f0-02515f7af8f0
TCGA-BP-5173-01A-01D-1429-08 RCC 3ce0a5fc-09ae-412a-8a5b-56d9a44433aa TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0e907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5189-01A-02D-1429-08 RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5190-01A-01D-1429-08 RCC	TCGA-BP-5169-01A-01D-1429-08	RCC	3527b21e-972b-4c31-b5de-8c394ce0e500
TCGA-BP-5174-01A-01D-1429-08 RCC 53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5199-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08	TCGA-BP-5170-01A-01D-1429-08	RCC	68761b2c-66b9-4adf-9b60-955f79ed0f11
TCGA-BP-5175-01A-01D-1429-08 RCC 30e58a1e-e7db-43ce-a7e8-a1fd21f4438e TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5173-01A-01D-1429-08	RCC	3ce0a5fc-09ae-412a-8a5b-56d9a44433aa
TCGA-BP-5176-01A-01D-1429-08 RCC 607eb48b-1647-4e35-ac60-f6c50341e304 TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5189-01A-02D-1429-08 RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5174-01A-01D-1429-08	RCC	53b5cf8d-f3cf-4e7e-91ec-b0c907d1c13f
TCGA-BP-5177-01A-01D-1429-08 RCC ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5175-01A-01D-1429-08	RCC	30e58a1e-e7db-43ce-a7e8-a1fd21f4438e
TCGA-BP-5178-01A-01D-1429-08 RCC 60888dc5-1408-4bfb-bf27-f3e22f5488e4 TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5176-01A-01D-1429-08	RCC	607eb48b-1647-4e35-ac60-f6c50341e304
TCGA-BP-5180-01A-01D-1429-08 RCC a776bde5-7503-459c-8419-dc0d744a651e TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5177-01A-01D-1429-08	RCC	ad4cc7e3-c4d1-4cc0-9c93-33b47dadaaae
TCGA-BP-5182-01A-01D-1429-08 RCC 00523547-da1c-4bb1-a627-c0946849b376 TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5178-01A-01D-1429-08	RCC	60888dc5-1408-4bfb-bf27-f3e22f5488e4
TCGA-BP-5183-01A-01D-1429-08 RCC cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4 TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5180-01A-01D-1429-08	RCC	a776bde5-7503-459c-8419-dc0d744a651e
TCGA-BP-5184-01A-01D-1429-08 RCC ddebed14-f47f-46e6-ac39-c74ed3363211 TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5182-01A-01D-1429-08	RCC	00523547-da1c-4bb1-a627-c0946849b376
TCGA-BP-5185-01A-01D-1429-08 RCC 42dc6d82-f52a-4b13-b3bc-c63002b47e98 TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5183-01A-01D-1429-08	RCC	cd4c37c3-95f2-4612-b6a8-9d6d1dfb5fd4
TCGA-BP-5186-01A-01D-1429-08 RCC 02b98f85-07df-4fb2-b27e-efd368c84ec8 TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5184-01A-01D-1429-08	RCC	ddebed14-f47f-46e6-ac39-c74ed3363211
TCGA-BP-5187-01A RCC 3257e690-9306-434f-b6ac-17da58ab1243 TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5185-01A-01D-1429-08	RCC	42dc6d82-f52a-4b13-b3bc-c63002b47e98
TCGA-BP-5189-01A-02D-1429-08 RCC ca98342a-65ec-468a-9cc1-44c7d31a67d6 TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5186-01A-01D-1429-08	RCC	02b98f85-07df-4fb2-b27e-efd368c84ec8
TCGA-BP-5190-01A-01D-1429-08 RCC 5491645b-552c-47a9-b081-e8e508d1df3d TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5187-01A	RCC	3257e690-9306-434f-b6ac-17da58ab1243
TCGA-BP-5191-01A-01D-1429-08 RCC 64dd8a08-483e-4dce-90b0-64a751fdbebd	TCGA-BP-5189-01A-02D-1429-08	RCC	ca98342a-65ec-468a-9cc1-44c7d31a67d6
	TCGA-BP-5190-01A-01D-1429-08	RCC	5491645b-552c-47a9-b081-e8e508d1df3d
TCGA-BP-5192-01A-01D-1429-08 RCC 4db23b76-46dd-4ed9-a168-fee43b2fc7d7	TCGA-BP-5191-01A-01D-1429-08	RCC	64dd8a08-483e-4dce-90b0-64a751fdbebd
	TCGA-BP-5192-01A-01D-1429-08	RCC	4db23b76-46dd-4ed9-a168-fee43b2fc7d7

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TCGA-BP-5194-01A-02D-1429-08	RCC	5b52c97e-fdd2-4ae2-b036-297feeb1c7e2
TCGA-BP-5195-01A-02D-1429-08	RCC	c2ab2f01-3744-434a-b5b6-0f22599c9a17
TCGA-BP-5196-01A-01D-1429-08	RCC	201bf07d-0be9-442f-ad66-15ea8c7e812d
TCGA-BP-5198-01A-01D-1429-08	RCC	ac66d658-97d4-416b-8028-0077a1c8a01d
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TCGA-CJ-4902-01A-01D-1429-08	RCC	3ef9ea62-85c4-4261-af23-ecb86f192cdf
TCGA-CJ-4903-01A-01D-1429-08	RCC	3b685193-f1fa-4c1b-949b-bcdb2d1b934c
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123.1 23 3000 0171 112 1334 10	1 1100	20.10011 /020 1/00 0100 201/0/100000

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TCGA-05-4384-01A-01D-1753-08	LUAD	4c71b66b-813f-472b-b866-b34b5b9199e7
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TCGA-05-4417-01A-22D-1855-08	LUAD	57e3657d-7a3c-4d80-a2c2-2de0293f5f05
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	LUAD	8c5a3460-c1fa-4b7b-9b31-11f9c7b03255
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TCGA-17-Z016-01A-01W-0746-08	LUAD	39bbd67b-52fd-46e5-98cf-b5632400216d
TCGA-17-Z017-01A-01W-0746-08	LUAD	37049bf1-55cb-44d3-b673-1e270ea835f7
TCGA-17-Z018-01A-01W-0746-08	LUAD	dd1a61eb-8362-41a9-952d-b7e6887457ad
TCGA-17-Z020-01A-01W-0746-08	LUAD	7ea20aa3-68cf-4389-9ace-99d6149d16c1
TCGA-17-Z021-01A-01W-0746-08	LUAD	9394b536-cd08-414b-86a3-c6491f967709
TCGA-17-Z022-01A-01W-0746-08	LUAD	7f07e5b3-bf70-4690-84ba-a9eace798a24
TCGA-17-Z023-01A-01W-0746-08	LUAD	bd72330a-463f-471b-9eba-2f188524e74c
TCGA-17-Z025-01A-01W-0746-08	LUAD	99eab29e-32d3-49d5-aa30-56de8be556e7
TCGA-17-Z026-01A-01W-0746-08	LUAD	bb048ffc-de00-4706-85bb-d052c0fb6496
TCGA-17-Z027-01A-01W-0746-08	LUAD	880452fe-00ed-4732-bbcf-14b55c235e61
TCGA-17-Z028-01A-01W-0746-08	LUAD	1f55fb6e-342a-41e0-9a8e-7c5156c95eaa
TCGA-17-Z030-01A-01W-0746-08	LUAD	e35e27e8-6cc5-495b-9ae8-89f65d94ebed
TCGA-17-Z031-01A-01W-0746-08	LUAD	6516244a-dfd8-4568-a2d2-7556cbea52b1
TCGA-17-Z032-01A-01W-0746-08	LUAD	92bc438b-02c1-4b81-a90a-4a1302786a81
TCGA-17-Z033-01A-01W-0746-08	LUAD	639aea7c-5a38-4641-bf0d-90a9ce8e2980
TCGA-17-Z035-01A-01W-0746-08	LUAD	a4bcbb2e-594f-4a89-8b72-8c922a64cdef
TCGA-17-Z036-01A-01W-0746-08	LUAD	374b881a-dbe2-4b4b-bfc0-8431f1aec06c
TCGA-17-Z037-01A-01W-0746-08	LUAD	bffe237d-31b0-4950-a7ab-4ac7047aa3c0
TCGA-17-Z038-01A-01W-0746-08	LUAD	8785c362-1c4d-41da-a29e-5cff21dc2a2e
TCGA-17-Z040-01A-01W-0746-08	LUAD	62d2ca54-b8e0-4907-b75e-cb9786069b52
TCGA-17-Z041-01A-01W-0746-08	LUAD	c0ead7c7-169e-4932-a987-5461611c95e6
TCGA-17-Z042-01A-01W-0746-08	LUAD	3c303c9d-6cda-490d-a64d-21bc40b064f3
TCGA-17-Z043-01A-01W-0746-08	LUAD	577f7267-c568-4002-a153-26c09d1eca97
TCGA-17-Z044-01A-01W-0746-08	LUAD	cb1aaeb8-0c6f-4266-968c-38a3823d85f6
TCGA-17-Z045-01A-01W-0746-08	LUAD	ec7e65c5-7158-427f-8034-8616077da50b
TCGA-17-Z046-01A-01W-0746-08	LUAD	7aac0e3f-39fe-4c9a-9482-50f02f1b919d
TCGA-17-Z047-01A-01W-0747-08	LUAD	2c04cfa8-6e99-46fa-82ad-36fb96e5ffef
TCGA-17-Z048-01A-01W-0746-08	LUAD	8495e150-796b-4e15-9fa6-1fba558d7b10
TCGA-17-Z049-01A-01W-0746-08	LUAD	ac31bcc6-6ccc-43b7-96f2-3ab47050be76
TCGA-17-Z050-01A-01W-0747-08	LUAD	d086dd38-a9e0-466c-b1e5-9a4a879abd55
TCGA-17-Z051-01A-01W-0747-08	LUAD	5584878f-0608-45d9-8e28-29c277bf655f
TCGA-17-Z052-01A-01W-0747-08	LUAD	afdf7c82-2a17-4c73-980c-74ec822dc803
TCGA-17-Z053-01A-01W-0747-08	LUAD	2d422986-6e91-4299-b6cf-4076f3706c83
TCGA-17-Z054-01A-01W-0747-08	LUAD	409dd077-dab9-4f79-9c33-2c3b75b63125
TCGA-17-Z055-01A-01W-0747-08	LUAD	de78326d-3afc-4f29-af7f-1750da544826
TCGA-17-Z056-01A-01W-0747-08	LUAD	e6cb3d63-5a55-4eba-84d2-a25917c7b18e
TCGA-17-Z057-01A-01W-0747-08	LUAD	4236905d-1549-4cbc-b3c6-e62db9ea598b

TCGA-17-Z058-01A-01W-0747-08	LUAD	6b0b1fca-efce-49d6-9f7b-a2c34bb343e9
TCGA-17-Z059-01A-01W-0747-08	LUAD	88ec6fb4-1b81-422e-8204-ef9e8dbf260c
TCGA-17-Z060-01A-01W-0747-08	LUAD	f834dfa4-8d9c-4e0b-861f-a3cc31245237
TCGA-17-Z061-01A-01W-0747-08	LUAD	1eb07e6e-6cf8-45e4-9b5c-1a9a5d38d117
TCGA-17-Z062-01A-01W-0747-08	LUAD	f3280e5f-7d6e-4a18-a5a5-e84b805c9e66
TCGA-35-3615-01A	LUAD	7407d705-6ec6-4143-93d2-eedcf5a22399
TCGA-35-3621-01A-01D-0969-08	LUAD	4a0cc41a-562c-4aea-a7c3-b1186d46cda8
	LUAD	408e1cb4-64a8-4801-bf58-3b8183ede851
TCGA-35-4122-01A-01D-1105-08 TCGA-35-4123-01A-01D-1105-08	LUAD	7ceccaee-df27-4f7f-bfcd-e1c59b365711
TCGA-35-5375-01A-01D-1625-08	LUAD	63e76bef-3ef1-445f-b591-649d774729cd
TCGA-38-4625-01A-01D-1553-08		6f317d31-c9a4-4345-b5b1-b75776536402
	LUAD	85b56ce7-b420-433e-a77d-43ef628d685c
TCGA 38 4627 01A 01D 1553 08		
TCGA 38 4627-01A-01D-1553-08	LUAD LUAD	abef97da-d7db-495f-b594-fa66577becd6
TCGA-38-4628-01A-01D-1265-08 TCGA-38-4629-01A-02D-1265-08	LUAD	67bc44b7-92cf-4e8f-a7f6-c53bf34a17c6
		4797f969-5f4d-4681-9fc5-68f25ba8f4d8
TCGA-38-4630-01A-01D-1265-08	LUAD	2b139bb4-5a29-4684-901d-8d966ff79ac2
TCGA-38-4631-01A-01D-1753-08	LUAD	b3ffc36d-b0b8-4ada-a00a-b48890c0162c
TCGA 28 (178 01A 11D 1753 08	LUAD	83519ed1-29e2-4f1b-922c-5779f64178bc
TCGA-38-6178-01A-11D-1753-08	LUAD	7fa467f1-d928-4d81-bd0b-68d67a5c18cf
TCGA-44-2655-01A	LUAD	9fcabda1-ea79-4188-8b3f-7d0fd060a819
TCGA-44-2656-01A	LUAD	5593f581-3d45-4a4a-a525-bfae1f4753a0
TCGA-44-2657-01A-01D-1105-08	LUAD	e3aa9b45-13b9-4b61-a30f-ae3f88466040
TCGA-44-2661-01A-01D-1105-08	LUAD	3c3a2e7c-9aa0-495e-95c7-87f661b9ed92
TCGA 44-2662-01A	LUAD	d2198941-e96f-40bd-9fbe-82886217d5db
TCGA-44-2665-01A	LUAD	a0863fa6-515c-44fa-825f-f9e243f945f1
TCGA-44-2666-01A	LUAD	27a64f32-69c5-4c49-86b4-c8fc923cae08
TCGA-44-2668-01A	LUAD	dd9a6c68-b8b4-4168-9ff9-72a45f20c44f
TCGA 44-3396-01A-01D-1265-08	LUAD	d68b216c-b304-4b30-9af7-eb3a9a1a55ae
TCGA 44-3398-01A-01D-1105-08	LUAD	82284bb3-2dfa-4016-a908-3b5994e00d31
TCGA-44-3918-01A-01D-1105-08	LUAD	7f456c3f-58e3-43f1-9f76-4422451528a5
TCGA 44 4112 01A	LUAD	9de8d353-3442-41d8-8bfe-a08c4975eaca 6c206676-e511-4281-91f5-bfe91b3279a4
TCGA 44 5642 OLA OLD 1625 08	LUAD	
TCGA 44 5645 01A 01D 1625 08	LUAD	44286013-ae97-4890-86d3-1163285ac0cd
TCGA-44-5645-01A-01D-1625-08	LUAD	dac33765-0c88-4a51-8389-c042ccb78c83
TCGA 44-6144-01A-11D-1753-08	LUAD	f19575fd-eb9d-429f-96ce-c0e8f4bbc593
TCGA 44-6145-01A-11D-1753-08	LUAD	220dc947-4afc-4485-bcc7-cea046100b4b
TCGA-44-6146-01A-11D-1753-08	LUAD	d5e90162-d7d2-4a7c-89f0-51c2b32c9ef0
TCGA 44-6147-01A-11D-1753-08	LUAD	7b6daa70-492e-4283-b3d2-b26f4e26a8d4
TCGA 44 6774 01A 21D 1855 08	LUAD	9c7b3ac8-1352-49cd-8a8c-df6b19f6fd64
TCGA-44-6774-01A-21D-1855-08	LUAD	f9ccld71-bece-4693-b953-3e73d1b6c11c
TCGA-44-6775-01A-11D-1855-08	LUAD	7a70a44f-84f3-440a-b898-dc3a0eff748e

TCGA.44-6776-01A-11D-1855-08		1	T
TCGA-44-6778-01A-11D-1855-08	TCGA-44-6776-01A-11D-1855-08	LUAD	7d3c5101-fae2-4320-a8a2-a93753375368
TCGA-44-6779-01A-11D-1855-08	TCGA-44-6777-01A-11D-1855-08	LUAD	32a0f0f3-3879-4b96-b9bb-eeab87827f6e
TCGA-49-4486-01A-01D-1265-08	TCGA-44-6778-01A-11D-1855-08	LUAD	903182ad-3145-4fa3-869e-62774aedf86c
TCGA-49-4487-01A-21D-1855-08	TCGA-44-6779-01A-11D-1855-08	LUAD	d6990a90-6a99-490b-a476-5298f0c4e4f2
TCGA-49-4488-01A-01D-1753-08	TCGA-49-4486-01A-01D-1265-08	LUAD	3ac132c3-4889-4dc3-8b3d-0ef98065a858
TCGA-49-4490-01A-21D-1855-08	TCGA-49-4487-01A-21D-1855-08	LUAD	9bd8e303-a81e-4ff8-882b-d46a2f7c55d2
TCGA-49-4494-01A-01D-1265-08	TCGA-49-4488-01A-01D-1753-08	LUAD	3635bb9c-a332-4445-ad81-83cec426dd02
TCGA-49-4501-01A-01D-1265-08 LUAD 0c53bb1b-5e6f-44a8-97a0-i89d43e0e789 TCGA-49-4505-01A-01D-1265-08 LUAD d707f8ad-5ea5-493a-a745-9b5dba64f213 TCGA-49-4507-01A-01D-1265-08 LUAD 562a09a1-b491-45c8-a87d-3c2471353c0d TCGA-49-4510-01A-01D-1265-08 LUAD 562a09a1-b491-45c8-a87d-3c2471353c0d TCGA-49-4510-01A-01D-1265-08 LUAD 562a09a1-b491-45c8-a87d-3c2471353c0d TCGA-49-4510-01A-21D-1855-08 LUAD fa6a6015-8949-4e01-9435-d3117601627f TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475e-8ec-5-66d76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 5459d29-a8e0-4d2d-8552-d27946f96070 TCGA-49-6743-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768e96 TCGA-49-6767-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768e96 TCGA-49-6767-01A-11D-1855-08 LUAD bf8p7048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD bf8p7048-977b-4722-be8f-3dd37370ba30 TCGA-50-5044-01A-21D-1855-08 LUAD b673498-64bf7-4554-b635-ca6d9-30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b67349-46742-04-04-01D-1625-08 LUAD b67349-6767-01A-11D-1855-08 LUAD b67349-6767-01A-1D-1625-08 LUAD b67349-6767-01A-2D-1855-08 LUAD b67349-6767-01A-2D-1855-08 LUAD b6747-49-80e-0879-66068e TCGA-50-5051-01A-21D-1855-08 LUAD b6747-49-80e-0879-66068e TCGA-50-5051-01A-21D-1855-08 LUAD b6747-49-80e-0879-66068e TCGA-50-5051-01A-21D-1855-08 LUAD b6747-49-80e-0879-66068e TCGA-50-5091-01D-1625-08 LUAD b6747-49-80e-0879-66068e TCGA-50-5093-01A-11D-1753-08 LUAD b6748-664-664-461-8761-8862-990-77688-3718735 TCGA-50-5931-01A-11D-1753-08 LUAD b6748-664-664-461-8701-0488-3718735 TCGA-50-5933-01A-11D-1753-08 LUAD b66615-6944-461-8701-0488-3718735 TCGA-50-5930-01A-11D-1753-08 LUAD b66615-6944-461-8701-4688-570-5936-6917b4 TCGA-50-5930-01A-11D-1753-08 LUAD b66615-6946-4648-8572-ff18bda4765 TCGA-50-59	TCGA-49-4490-01A-21D-1855-08	LUAD	940455cf-aa91-432a-bc39-9dfba206e32b
TCGA-49-4505-01A-01D-1265-08 LUAD e773a2fe-1d80-492d-bba8-105036a14a92 TCGA-49-4506-01A-01D-1265-08 LUAD d707f8ad-5ea5-493a-a745-9b5dba64f213 TCGA-49-4507-01A-01D-1265-08 LUAD 562a09a1-b491-45c8-a87d-3c2471353c0d TCGA-49-4510-01A-01D-1265-08 LUAD b2c12bff-addd-45a2-ada4-c30ac935809c TCGA-49-4512-01A-21D-1855-08 LUAD fa6a60f5-8949-4e01-9435-d3117601627f TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475c-8ec-5.6dd76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82be2e608890 TCGA-49-6742-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bf6b7048-977b-4722-be8f-3dd37370ba30 TCGA-9-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5049-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5051-01A-21D-1855-08 LUAD b150bc27-fb18-4eec-8785-b88b69bcb6 TCGA-50-5055-01A-01D-1625-08 LUAD<	TCGA-49-4494-01A-01D-1265-08	LUAD	136bc973-1908-4767-9b22-d43d522b7c71
TCGA-49-4506-01A-01D-1265-08 LUAD TCGA-49-4507-01A-01D-1265-08 LUAD TCGA-49-4510-01A-01D-1265-08 LUAD TCGA-49-4510-01A-01D-1265-08 LUAD TCGA-49-4510-01A-01D-1265-08 LUAD TCGA-49-4512-01A-21D-1855-08 LUAD TCGA-49-4514-01A-21D-1855-08 LUAD TCGA-49-4514-01A-21D-1855-08 LUAD TCGA-49-4514-01A-21D-1855-08 LUAD TCGA-49-4514-01A-11D-1855-08 LUAD TCGA-49-6742-01A-11D-1855-08 LUAD TCGA-49-6743-01A-11D-1855-08 LUAD TCGA-49-6743-01A-11D-1855-08 LUAD TCGA-49-6743-01A-11D-1855-08 LUAD TCGA-49-6745-01A-11D-1855-08 LUAD TCGA-49-6745-01A-11D-1855-08 LUAD TCGA-49-6745-01A-11D-1855-08 LUAD TCGA-49-6745-01A-11D-1855-08 LUAD TCGA-49-6767-01A-11D-1855-08 LUAD TCGA-49-6767-01A-11D-1855-08 LUAD TCGA-49-6767-01A-11D-1855-08 LUAD TCGA-50-5044-01A-21D-1855-08 LUAD TCGA-50-5045-01A-01D-1625-08 LUAD TCGA-50-5045-01A-01D-1625-08 LUAD TCGA-50-5051-01A-21D-1855-08 LUAD TCGA-50-5051-01A-21D-1855-08 LUAD TCGA-50-5055-01A-01D-1625-08 LUAD TCGA-50-5056-01A-01D-1625-08 LUAD TCGA-50-5066-01A-01D-1625-08 LUAD TCGA-50-5068-01A-01D-1625-08 LUAD TCGA-50-5068-01A-01D-1625-08 LUAD TCGA-50-5068-01A-01D-1625-08 LUAD TCGA-50-5068-01A-01D-1625-08 LUAD TCGA-50-5093-01A-11D-1753-08 LUAD TCGA-50-5094-01A-11D-1753-08 LUAD TCGA-50-5094-01A	TCGA-49-4501-01A-01D-1265-08	LUAD	0c53bb1b-5e6f-44a8-97a0-f89d43e0e789
TCGA-49-4507-01A-01D-1265-08 LUAD 562a09a1-b491-45c8-a87d-3c2471353c0d TCGA-49-4510-01A-01D-1265-08 LUAD b2c12bff-addd-45a2-ada4-c30ac935809c TCGA-49-4512-01A-21D-1855-08 LUAD fa6a60f5-8949-4e01-9435-d3117601627f TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475e-8ec5-66d76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d71-b076-24ac2f768696 TCGA-49-674-01A-11D-1855-08 LUAD bf8b97048-9775-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 982f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD 982f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD 96358297-0735-4ab-a01c-a6be530-0da28 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4ab-a01c-a6be5366a3de TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4ab-a01c-a6be5866be586a3de TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-90c-f680e2311cf6 TCGA-50-5055-01A-01D-1625-08 LUA	TCGA-49-4505-01A-01D-1265-08	LUAD	e773a2fe-1d80-492d-bba8-105036a14a92
TCGA-49-4510-01A-01D-1265-08 LUAD b2c12bff-addd-45a2-ada4-c30ac935809c TCGA-49-4512-01A-21D-1855-08 LUAD fa6a60f5-8949-4e01-9435-d3117601627f TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475e-8ec5-66d76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6767-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5045-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5366a3de TCGA-50-5051-01A-21D-1855-08 LUAD b550bc27-fb18-4eec-8785-b8e8b69bcbe6 TCGA-50-5050-01A-01D-1625-08 LUAD 12fe153c-a8f7-49ec-96c-f680e2311cf6 TCGA-50-5055-01A-01D-1625-08 LUAD 15a97315-1906-4774-980c-0879c6ad368e TCGA-50-5066-01A-01D-1625-08 LUAD	TCGA-49-4506-01A-01D-1265-08	LUAD	d707f8ad-5ea5-493a-a745-9b5dba64f213
TCGA-49-4512-01A-21D-1855-08 LUAD fa6a60f5-8949-4e01-9435-d3117601627f TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475e-8ec5-66d76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 545e9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6745-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD e034986-4bf7-4554-b635-ca6d9e30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5050-01A-01D-1625-08 LUAD 96358297-0735-4cab-a01c-a6be5d86a3de TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD 15a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD 15a97315-1906-4774-980e-0879c6ad368e TCGA-50-5093-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-077d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD<	TCGA-49-4507-01A-01D-1265-08	LUAD	562a09a1-b491-45c8-a87d-3c2471353c0d
TCGA-49-4514-01A-21D-1855-08 LUAD 7751af67-1415-475e-8ec5-66d76f515014 TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6745-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD e034986-4bf7-4554-b635-ca6d9e30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD 15a97315-1906-4774-980e-0879c6ad368e TCGA-50-5072-01A-21D-1855-08 LUAD 16a48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5930-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5931-01A-11D-1753-08 LUAD <td>TCGA-49-4510-01A-01D-1265-08</td> <td>LUAD</td> <td>b2c12bff-addd-45a2-ada4-c30ac935809c</td>	TCGA-49-4510-01A-01D-1265-08	LUAD	b2c12bff-addd-45a2-ada4-c30ac935809c
TCGA-49-6742-01A-11D-1855-08 LUAD 49dec0c2-8e75-4f44-a253-82b2ea605890 TCGA-49-6743-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa255298 TCGA-50-5044-01A-21D-1855-08 LUAD ec334986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6bc5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5056-01A-01D-1625-08 LUAD 12fe153a-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-50568-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-50568-01A-01D-1625-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5931-01A-11D-1753-08 LUAD 6726c157-f688-491d-856-35628645d189 TCGA-50-5933-01A-11D-1753-08 LU	TCGA-49-4512-01A-21D-1855-08	LUAD	fa6a60f5-8949-4e01-9435-d3117601627f
TCGA-49-6743-01A-11D-1855-08 LUAD 545c9d29-a8e0-4d2d-8552-d27b46f96070 TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD ec034986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6bc5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD b50bc27-fb18-4ece-8785-b8e8b69bcb6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-50566-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD d64c48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-bs89-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-s70f-0488e3718f35 TCGA-50-5931-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD </td <td>TCGA-49-4514-01A-21D-1855-08</td> <td>LUAD</td> <td>7751af67-1415-475e-8ec5-66d76f515014</td>	TCGA-49-4514-01A-21D-1855-08	LUAD	7751af67-1415-475e-8ec5-66d76f515014
TCGA-49-6744-01A-11D-1855-08 LUAD bf6ba698-7154-4d7f-b076-24ac2f768696 TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD ec034986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-90c-f680c2311cf6 TCGA-50-5056-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5068-01A-01D-1625-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5935-01A-11D-1753-08 LUAD	TCGA-49-6742-01A-11D-1855-08	LUAD	49dec0c2-8e75-4f44-a253-82b2ea605890
TCGA-49-6745-01A-11D-1855-08 LUAD bfb97048-977b-4722-be8f-3dd37370ba30 TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD ec034986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5044-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eec-8785-b8e8b69bcb6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5939-01A-11D-1625-08 LUAD	TCGA-49-6743-01A-11D-1855-08	LUAD	545c9d29-a8e0-4d2d-8552-d27b46f96070
TCGA-49-6767-01A-11D-1855-08 LUAD 9f82f494-042a-4f00-954c-4761fa25b298 TCGA-50-5044-01A-21D-1855-08 LUAD ec034986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcb6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5931-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5930-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5930-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUA	TCGA-49-6744-01A-11D-1855-08	LUAD	bf6ba698-7154-4d7f-b076-24ac2f768696
TCGA-50-5044-01A-21D-1855-08 LUAD ec034986-4bf7-4554-b635-ca6d9c30da28 TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5941-01A-11D-1753-08 LUA	TCGA-49-6745-01A-11D-1855-08	LUAD	bfb97048-977b-4722-be8f-3dd37370ba30
TCGA-50-5045-01A-01D-1625-08 LUAD b0d734ad-1222-4bc0-b02b-1d2262b8ac35 TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD clefdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5944-01A-21D-1753-08 LUA	TCGA-49-6767-01A-11D-1855-08	LUAD	9f82f494-042a-4f00-954c-4761fa25b298
TCGA-50-5049-01A-01D-1625-08 LUAD 96358297-0735-4eab-a01c-a6be5d86a3de TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ee-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1753-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUA	TCGA-50-5044-01A-21D-1855-08	LUAD	ec034986-4bf7-4554-b635-ca6d9c30da28
TCGA-50-5051-01A-21D-1855-08 LUAD bb50bc27-fb18-4eee-8785-b8e8b69bcbe6 TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5935-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5946-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUA	TCGA-50-5045-01A-01D-1625-08	LUAD	b0d734ad-1222-4bc0-b02b-1d2262b8ac35
TCGA-50-5055-01A-01D-1625-08 LUAD 12fe153e-a8f7-49ec-9e0c-f680e2311cf6 TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5935-01A-11D-1753-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD 4314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5049-01A-01D-1625-08	LUAD	96358297-0735-4eab-a01c-a6be5d86a3de
TCGA-50-5066-01A-01D-1625-08 LUAD f5a97315-1906-4774-980e-0879c6ad368e TCGA-50-5068-01A-01D-1625-08 LUAD clefdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5051-01A-21D-1855-08	LUAD	bb50bc27-fb18-4eee-8785-b8e8b69bcbe6
TCGA-50-5068-01A-01D-1625-08 LUAD clefdc48-6ea5-45f0-9fa3-94c42ecf3ab4 TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5055-01A-01D-1625-08	LUAD	12fe153e-a8f7-49ec-9e0c-f680e2311cf6
TCGA-50-5072-01A-21D-1855-08 LUAD 3c6dcba5-1312-40ca-b589-07f7d88b3477 TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5066-01A-01D-1625-08	LUAD	f5a97315-1906-4774-980e-0879c6ad368e
TCGA-50-5930-01A-11D-1753-08 LUAD bd3e88b3-b37c-4641-85fa-d8125ba324ca TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5068-01A-01D-1625-08	LUAD	c1efdc48-6ea5-45f0-9fa3-94c42ecf3ab4
TCGA-50-5931-01A-11D-1753-08 LUAD 290847c6-c9d4-4a16-a70f-0488e3718f35 TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5072-01A-21D-1855-08	LUAD	3c6dcba5-1312-40ca-b589-07f7d88b3477
TCGA-50-5932-01A-11D-1753-08 LUAD 6726c157-f688-491d-8b56-35628645df89 TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5930-01A-11D-1753-08	LUAD	bd3e88b3-b37c-4641-85fa-d8125ba324ca
TCGA-50-5933-01A-11D-1753-08 LUAD cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1 TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5931-01A-11D-1753-08	LUAD	290847c6-c9d4-4a16-a70f-0488e3718f35
TCGA-50-5935-01A-11D-1753-08 LUAD 9570cd02-3339-4805-855a-74ebe429df96 TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5932-01A-11D-1753-08	LUAD	6726c157-f688-491d-8b56-35628645df89
TCGA-50-5936-01A-11D-1625-08 LUAD 82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08 TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5933-01A-11D-1753-08	LUAD	cc3a9cfe-8a14-4fb4-a60f-3ec795c5d7a1
TCGA-50-5939-01A-11D-1625-08 LUAD aa9108d7-5036-4059-ad82-dc64161d5bc3 TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5935-01A-11D-1753-08	LUAD	9570cd02-3339-4805-855a-74ebe429df96
TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5936-01A-11D-1625-08	LUAD	82d380d5-4c07-4cf0-a6e9-7ca9e3fc9a08
TCGA-50-5941-01A-11D-1753-08 LUAD 86ef12c0-d5fc-4852-9960-593366e717b4 TCGA-50-5942-01A-21D-1753-08 LUAD 95475c1b-086d-4e09-a871-47d8f76c1a07 TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5939-01A-11D-1625-08	LUAD	aa9108d7-5036-4059-ad82-dc64161d5bc3
TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122		LUAD	
TCGA-50-5944-01A-11D-1753-08 LUAD a314ee0c-694b-4ac8-b572-ff1fbbda4765 TCGA-50-5946-01A-11D-1753-08 LUAD 142d43e8-10e1-4945-a37c-f2824d53b122	TCGA-50-5942-01A-21D-1753-08	LUAD	95475c1b-086d-4e09-a871-47d8f76c1a07
	TCGA-50-5944-01A-11D-1753-08	LUAD	a314ee0c-694b-4ac8-b572-ff1fbbda4765
TCGA-50-6590-01A-12D-1855-08 LUAD 85de182b-f4ae-41e6-b3fb-f60f46c072e4	TCGA-50-5946-01A-11D-1753-08	LUAD	142d43e8-10e1-4945-a37c-f2824d53b122
	TCGA-50-6590-01A-12D-1855-08	LUAD	85de182b-f4ae-41e6-b3fb-f60f46c072e4

TCGA-50-6591-01A-11D-1753-08			
TCGA-50-6593-01A-11D-1753-08	TCGA-50-6591-01A-11D-1753-08	LUAD	bf7462a2-394f-4838-bcb6-4d0126fa48b1
TCGA-50-6594-01A-11D-1753-08	TCGA-50-6592-01A-11D-1753-08	LUAD	d0303d05-a937-4a7d-9934-ffa93cc1c5de
TCGA-50-6595-01A-12D-1855-08	TCGA-50-6593-01A-11D-1753-08	LUAD	10e03053-f6e3-42b7-8638-ce58c6e7dfaa
TCGA-50-6597-01A-11D-1855-08	TCGA-50-6594-01A-11D-1753-08	LUAD	e1365c7d-e93e-4478-a8e9-ae2d7ca30bc6
TCGA-55-1592-01A	TCGA-50-6595-01A-12D-1855-08	LUAD	9913e506-fc98-467d-8601-89595d0475e8
TCGA-55-1594-01A	TCGA-50-6597-01A-11D-1855-08	LUAD	cd0aeed5-93a1-4287-8a88-fe6b7b5e3983
TCGA-55-1595-01A-01D-0969-08 LUAD f1be8e08-5201-49bb-abf7-cede0eff06d6 TCGA-55-1596-01A LUAD 9a7a1b22-9df6-438F-ad00-54755c7dbc7c TCGA-55-1596-01A LUAD 9a7a1b22-9df6-438F-ad00-54755c7dbc7c TCGA-55-1596-01A LUAD ac7ab3b3-eb76-4da9-bfb3-82b90c8d79d6 TCGA-55-6543-01A-11D-1855-08 LUAD ac7ab3b3-eb76-4da9-bfb3-82b90c8d79d6 TCGA-55-6642-01A-11D-1855-08 LUAD ac7ab3b3-eb76-4da9-bfb3-82b90c8d79d6 TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1676-01A LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1678-01A-01W-0928-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1678-01A-01W-0928-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1678-01A-01W-0928-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 Dc6-64-1678-01A-01D-1625-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 Dc6-64-5778-01A-01D-1625-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 Dc6-64-5778-01A-01D-1625-08 LUAD bc6eaf2b-9cc-4ac7-9b18-01c4-01D-1625-08 LUAD bc74bc6e-9c2a-92c0612fce33 TCGA-64-5781-01A-01D-1625-08 LUAD bc74bc74-8ca-4ad9-ace2-2b779336e557 TCGA-67-3770-01A LUAD bc74bc72d5-f642-423e-bd96-b2de1boc1778 TCGA-67-3771-01A LUAD bc74bc72d5-f642-423e-bd96-b2de1boc1778 TCGA-67-3774-01A LUAD bc74bc74bc74bc74bc74bc74bc74bc74bc74bc74	TCGA-55-1592-01A	LUAD	e190a9e4-10ae-4060-a071-4b8b73479023
TCGA-55-1596-01A LUAD 9a7a1b22-9di6-438f-ad00-54755c7dbc7c TCGA-55-5899-01A-11D-1625-08 LUAD ddaf36f7-7503-4ab4-b7f5-9777c0c1518c TCGA-55-6543-01A-11D-1753-08 LUAD ac7ab3b3-eb76-4da9-bib3-82b90c8d79d6 TCGA-55-6642-01A-11D-1855-08 LUAD 3c756f7c-d1f0-4ab1-9c9f-41d2282af3bf TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0f420a3 TCGA-64-1676-01A LUAD dbdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1670-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD dbdbe623-cf95-f4b18-8i62-e7b0a9c0f1db TCGA-64-1680-01A LUAD dbdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD dif5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5781-01A-01D-1625-08 LUAD ff9cf849-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5781-01A-01D-1625-08 LUAD ff9cf849-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5781-01A-01D-1625-08 LUAD ff9cf849-99cf-4f49-8f3d-e25e762eb3ce TCGA-67-3771-01A LUAD bd41cd6-693d-41d6-9dad-d1b1c30f5cb TCGA-67-3772-01A-01W-0928-08 LUAD db466f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3772-01A-01W-0928-08 LUAD db46c6f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3772-01A-01W-0928-08 LUAD bd41cd6-693d-41d6-9dad-d1b1c30f5cb TCGA-67-3772-01A-01W-0928-08 LUAD db46c6f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3772-01A-01D-1753-08 LUAD db46c6f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3772-01A-11D-1753-08 LUAD db46c6f4-e847-40bf-af14-20a3867a1c35 TCGA-67-6215-01A-11D-1753-08 LUAD bd40cd6-693d-41d6-9dad-d1b1c30f5cb TCGA-67-6215-01A-11D-1753-08 LUAD db46c6f4-e847-40bf-af14-20a3867a1c35 TCGA-67-6215-01A-11D-1753-08	TCGA-55-1594-01A	LUAD	2885d4b3-34a6-421d-b20c-eedad721d10a
TCGA-55-5899-01A-11D-1625-08 LUAD ddaf36f7-7503-4ab4-b7f5-9777c0c1518c TCGA-55-6543-01A-11D-1753-08 LUAD ac7ab3b3-eb76-4da9-bfb3-82b90e8d79d6 TCGA-55-6642-01A-11D-1855-08 LUAD 3c756f7c-d1f0-4ab1-9c9f-41d2282af3bf TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1676-01A LUAD dbdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1678-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 42e3b592-b57f-4b18-8f62-e7b0a9c011db TCGA-64-1678-01A-01W-0928-08 LUAD 0bdbe623-ct95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD 0bdbe623-ct95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD 3c340f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5778-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fce33 TCGA-64-5781-01A-01D-1625-08 LUAD f99cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD 9226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD 9226bc4-0202-4405-b3c9-208e8ff	TCGA-55-1595-01A-01D-0969-08	LUAD	f1be8e08-5201-49bb-abf7-cedc0eff06d6
TCGA-55-6543-01A-11D-1753-08 LUAD ac7ab3b3-eb76-4da9-bfb3-82b90c8d79d6 TCGA-55-6642-01A-11D-1855-08 LUAD 3c756f7c-d1f0-4ab1-9c9f-41d2282af3bf TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1676-01A LUAD 4bdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 42e3b592-b57f-4b18-8f62-e7b0a9c0f1db TCGA-64-1678-01A-01D-1625-08 LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD dfs957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5778-01A-01D-1625-08 LUAD 3c340f87-5981-db7a-b1ab-30c2056c785e TCGA-64-5778-01A-01D-1625-08 LUAD 5734711b-52c4-de6e-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD 5734711b-52c4-de6e-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f9-8f3d-e25e-f2eb3ce TCGA-67-3771-01A LUAD 4bd10cd6-69-4d-49d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 7bd10cd6-69-4d-40d-bd-bd-bde1bcf778 TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-	TCGA-55-1596-01A	LUAD	9a7a1b22-9df6-438f-ad00-54755c7dbc7c
TCGA-55-6642-01A-11D-1855-08 LUAD 3c756f7c-d1f0-4ab1-9e9f-41d2282af3bf TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1676-01A LUAD 4bdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 42e3b592-b57f-4b18-8f62-c7b0a9c0f1db TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD 209d392-7d3a-481c-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5781-01A-01D-1625-08 LUAD 5734711b-32cd-46e6-92ca-92c0612fec33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-63-5770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3772-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3773-01A LUAD 3585415-9ab9-4614-8b15-9a92-208e8ffb7408 <td>TCGA-55-5899-01A-11D-1625-08</td> <td>LUAD</td> <td>ddaf36f7-7503-4ab4-b7f5-9777c0c1518c</td>	TCGA-55-5899-01A-11D-1625-08	LUAD	ddaf36f7-7503-4ab4-b7f5-9777c0c1518c
TCGA-55-6712-01A-11D-1855-08 LUAD bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3 TCGA-64-1676-01A LUAD 4bdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1680-01A LUAD 42e3b592-b57f-4b18-8f62-e7b0a9c0f1db TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD 209d392-7d3a-481e-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0c1778 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0c1778 TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3772-01A-01W-0928-08 LUAD 93585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD 68c2a355-862c-4657-b296-5776ed8447b0	TCGA-55-6543-01A-11D-1753-08	LUAD	ac7ab3b3-eb76-4da9-bfb3-82b90c8d79d6
TCGA-64-1676-01A LUAD 4bdf77d2-33cc-46e0-af34-1e66a90a213a TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 42e3b592-b57f-4b18-8f62-e7b0a9c0f1db TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-481e-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-acc2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3770-01A LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-6216-01A-11D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c	TCGA-55-6642-01A-11D-1855-08	LUAD	3c756f7c-d1f0-4ab1-9c9f-41d2282af3bf
TCGA-64-1677-01A-01W-0928-08 LUAD 559017d8-4b22-4313-abdd-d3526c889d7f TCGA-64-1678-01A-01W-0928-08 LUAD 42a3b592-b57f-4b18-8f62-e7b0a9e0f1db TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-481c-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-acc2-2b779336e557 TCGA-67-3770-01A LUAD b6410c46-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3771-01A LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66fd1dc TCGA-67-67-2901B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6216-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed	TCGA-55-6712-01A-11D-1855-08	LUAD	bc6eaf2b-9ccc-4ac7-9b19-204b0ff420a3
TCGA-64-1678-01A-01W-0928-08 LUAD 42e3b592-b57f-4b18-8f62-e7b0a9c0f1db TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-483e-990b-d6369fb990b8 TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762e3ce TCGA-64-5781-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336c557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3770-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30f5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-61-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8e-2ecf-412f-b2c4-74529adb7c0f	TCGA-64-1676-01A	LUAD	4bdf77d2-33cc-46e0-af34-1e66a90a213a
TCGA-64-1680-01A LUAD 0bdbe623-cf95-465a-917d-87dfb6a8618e TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-481e-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5778-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-9c20612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3770-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c	TCGA-64-1677-01A-01W-0928-08	LUAD	559017d8-4b22-4313-abdd-d3526c889d7f
TCGA-64-5774-01A-01D-1625-08 LUAD df5957d5-20d3-483e-990b-d6369fb990b8 TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-481c-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-acc2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3770-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-73-4658-01A-01D-1753-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55	TCGA-64-1678-01A-01W-0928-08	LUAD	42e3b592-b57f-4b18-8f62-e7b0a9c0f1db
TCGA-64-5775-01A-01D-1625-08 LUAD c209d392-7d3a-481c-8cc7-398a6b90290a TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6215-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-71-6725-01A-11D-1855-08 LUAD 3146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11	TCGA-64-1680-01A	LUAD	0bdbe623-cf95-465a-917d-87dfb6a8618e
TCGA-64-5778-01A-01D-1625-08 LUAD 3c540f87-5981-4b7a-b1ab-30c2056c785e TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-71-6725-01A-11D-1855-08 LUAD 3146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4666-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7	TCGA-64-5774-01A-01D-1625-08	LUAD	df5957d5-20d3-483e-990b-d6369fb990b8
TCGA-64-5779-01A-01D-1625-08 LUAD 5734711b-52cd-46e6-9c2a-92c0612fee33 TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6217-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4666-01A-01D-1265-08 LUAD 48262c89-eac-44c6-9a06-7170b7b41058 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037	TCGA-64-5775-01A-01D-1625-08	LUAD	c209d392-7d3a-481c-8cc7-398a6b90290a
TCGA-64-5781-01A-01D-1625-08 LUAD fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-49103	TCGA-64-5778-01A-01D-1625-08	LUAD	3c540f87-5981-4b7a-b1ab-30c2056c785e
TCGA-64-5815-01A-01D-1625-08 LUAD e800c8d4-786a-4a9d-ace2-2b779336e557 TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4668-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-49103	TCGA-64-5779-01A-01D-1625-08	LUAD	5734711b-52cd-46e6-9c2a-92c0612fee33
TCGA-67-3770-01A LUAD 74bcf2d5-fd42-423e-bd96-b2de1b0cf778 TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4666-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-64-5781-01A-01D-1625-08	LUAD	fb9cfb49-99cf-4f49-8f3d-e25e762eb3ce
TCGA-67-3771-01A LUAD b0410cd6-693d-41d6-9dad-d1b1c30bf5cb TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4658-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb1956850 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-64-5815-01A-01D-1625-08	LUAD	e800c8d4-786a-4a9d-ace2-2b779336e557
TCGA-67-3772-01A-01W-0928-08 LUAD 09226bc4-0202-4405-b3c9-208e8ffb7408 TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-3770-01A	LUAD	74bcf2d5-fd42-423e-bd96-b2de1b0cf778
TCGA-67-3773-01A LUAD e4cb66f4-e847-40bf-af14-20a3867a1c35 TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-3771-01A	LUAD	b0410cd6-693d-41d6-9dad-d1b1c30bf5cb
TCGA-67-3774-01A LUAD b3585415-9ab9-4614-8b15-8edb66efd1dc TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4668-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-3772-01A-01W-0928-08	LUAD	09226bc4-0202-4405-b3c9-208e8ffb7408
TCGA-67-4679-01B-01D-1753-08 LUAD 341bf21e-abd5-498e-8c49-111782af842c TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-3773-01A	LUAD	e4cb66f4-e847-40bf-af14-20a3867a1c35
TCGA-67-6215-01A-11D-1753-08 LUAD 68c2a355-862c-4657-b296-5776ed8447b0 TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-3774-01A	LUAD	b3585415-9ab9-4614-8b15-8edb66efd1dc
TCGA-67-6216-01A-11D-1753-08 LUAD 6dc6da8c-2ecf-412f-b2c4-74529adb7c0f TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-4679-01B-01D-1753-08	LUAD	341bf21e-abd5-498e-8c49-111782af842c
TCGA-67-6217-01A-11D-1753-08 LUAD cb98d825-668f-4b16-a05e-501e1c94f3fe TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-6215-01A-11D-1753-08	LUAD	68c2a355-862c-4657-b296-5776ed8447b0
TCGA-71-6725-01A-11D-1855-08 LUAD 3a146eb4-7b9b-4834-b3d0-eac80f9173ec TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-6216-01A-11D-1753-08	LUAD	6dc6da8c-2ecf-412f-b2c4-74529adb7c0f
TCGA-73-4658-01A-01D-1753-08 LUAD b11151cf-6976-4812-a77e-1a12f9d1245c TCGA-73-4659-01A-01D-1265-08 LUAD 13989aec-b1a3-47c2-bc8e-ccf55f8e0c11 TCGA-73-4662-01A-01D-1265-08 LUAD 48262c89-ecac-44c6-9a06-7170b7b41058 TCGA-73-4666-01A-01D-1265-08 LUAD f49fc77e-03cd-423c-b3e1-18bb19568650 TCGA-73-4668-01A-01D-1265-08 LUAD 0fdcb5e9-ada2-4755-ae02-491037ee9c10	TCGA-67-6217-01A-11D-1753-08	LUAD	cb98d825-668f-4b16-a05e-501e1c94f3fe
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TCGA-18-3408-01A-01D-0983-08 LUSC cab7a425-e081-4bae-b666-6cdf8ba4dd70
TCGA-18-3409-01A-01D-0983-08 LUSC aa733cb0-37a9-4fef-8d40-d57596ce9e51
TCGA-18-3410-01A-01D-0983-08 LUSC 7e6382c3-368a-43a5-9812-c58f54ceba3f
TCGA-18-3411-01A-01D-0983-08 LUSC 6a9cc303-c7fd-4f40-8933-1636dea99252
TCGA-18-3412-01A-01D-0983-08 LUSC 84aca315-8380-4625-887f-a8b3c704c0a9
TCGA-18-3414-01A-01D-0983-08 LUSC 239deee9-2791-4163-b777-fdf8c49c9e33
TCGA-18-3415-01A-01D-0983-08 LUSC ad0365d1-10b1-41e6-b838-9c5794b9ad42
TCGA-18-3416-01A-01D-0983-08 LUSC e03577e7-37be-460b-96e8-5f6e0b49b3aa
TCGA-18-3417-01A-01D-1441-08 LUSC 024d8a82-06c5-4b82-9a27-c52bc4fd450a
TCGA-18-3419-01A-01D-0983-08 LUSC c75ed357-d845-4443-8c9e-a2afa8ed30df
TCGA-18-3421-01A-01D-0983-08 LUSC 9f0e482e-e72d-4c57-b4f7-4580edabd390
TCGA-18-4083-01A-01D-1352-08 LUSC 0b87a82d-096c-4dd7-80c4-b4054fc1eba2
TCGA-18-4086-01A-01D-1352-08 LUSC 9bbdf36b-6804-416f-977d-fce772972bcc
TCGA-18-4721-01A-01D-1441-08 LUSC d2ab2555-7288-47a4-a80c-bf62d65b67b8
TCGA-18-5592-01A-01D-1632-08 LUSC 1a6da454-8faf-4725-a702-55d29da461a5
TCGA-18-5595-01A-01D-1632-08 LUSC 973b8ed8-2295-4fb0-b857-f4433dfc785a
TCGA-21-1070-01A-01D-1521-08 LUSC 9e300205-b16d-4f40-bf1b-f47410678f6d
TCGA-21-1071-01A-01D-1521-08 LUSC e01302f9-c5d6-4745-9c5d-d8bb8d278a77
TCGA-21-1076-01A-02D-1521-08 LUSC 504d4cb0-d2dd-420d-82e6-9ec14434a0fc

TCGA-21-1077-01A-01D-1521-08	LUSC	a71d74cb-5b10-4787-a654-7049cbb49a92
TCGA-21-1078-01A-01D-1521-08	LUSC	8cf9b32d-3d6f-4898-8c7a-89511b754021
TCGA-21-1081-01A-01D-1521-08	LUSC	811f7a11-635c-4606-91fd-3729b97ffd8e
TCGA-21-5782-01A-01D-1632-08	LUSC	4c2ad4a0-5d57-4e27-9f35-058b2f205f50
TCGA-21-5784-01A-01D-1632-08	LUSC	f79285af-c364-4ec3-97d3-70a7d9b5800b
TCGA-21-5786-01A-01D-1632-08	LUSC	d7404e0f-d171-419b-97d3-807570aba129
TCGA-21-5787-01A-01D-1632-08	LUSC	7cb79e4b-c1f1-434d-b13b-6c2eb7760ee8
TCGA-22-0944-01A-01D-1521-08	LUSC	818a6f09-a7fd-4cce-8373-adb4bcb5bc8c
TCGA-22-1002-01A-01D-1521-08	LUSC	7c7604fe-8321-46cb-ac34-0e7994b8853b
TCGA-22-1011-01A-01D-1521-08	LUSC	c9924f9f-fd86-434c-a83d-393d65272e64
TCGA-22-1012-01A-01D-1521-08	LUSC	3b75368a-d57f-4787-a0ef-3f478c7d22bc
TCGA-22-1016-01A-01D-1521-08	LUSC	935b113e-f5ed-4a07-8e1d-1603daba7f40
TCGA-22-4591-01A-01D-1267-08	LUSC	bcfb93d4-8653-477b-b5d2-c2832a0e3d92
TCGA-22-4593-01A-21D-1817-08	LUSC	b4a48075-92fd-43ab-95f3-476bcea88d7b
TCGA-22-4595-01A-01D-1267-08	LUSC	7fcf5123-2d1b-4666-9d39-a1aaf63cf954
TCGA-22-4599-01A-01D-1441-08	LUSC	08732b51-8ec8-4888-b0c8-a0cb83181cb9
TCGA-22-4601-01A-01D-1441-08	LUSC	6c05b3f5-65e9-4e7d-9f99-a694006f2ed0
TCGA-22-4604-01A-01D-1267-08	LUSC	db2614fb-109c-4ce1-af4c-f648a0d417fb
TCGA-22-4607-01A-01D-1267-08	LUSC	d8c6bb83-ebdd-4547-9077-3eba5c8bb9f0
TCGA-22-4613-01A-01D-1441-08	LUSC	5d1d538a-57d3-42ec-9fa3-0fad10b0f52f
TCGA-22-5471-01A-01D-1632-08	LUSC	665e98bf-6163-4d18-9665-ba93df9ecf6d
TCGA-22-5472-01A-01D-1632-08	LUSC	be780766-483f-42f5-b0d0-11d23a940156
TCGA-22-5473-01A-01D-1632-08	LUSC	c107ca1d-5e35-470a-8c39-80dc7624e306
TCGA-22-5474-01A-01D-1632-08	LUSC	leda33fc-80e5-4c5f-8c61-43976ca0106f
TCGA-22-5477-01A-01D-1632-08	LUSC	e7ebc6fb-0926-4c8a-a67b-0c6b9c1ffaba
TCGA-22-5478-01A-01D-1632-08	LUSC	0ac704eb-d722-4c27-bfb4-fea6ca7af240
TCGA-22-5480-01A-01D-1632-08	LUSC	24e426fb-219a-4a4d-a45c-c9b0896d0e88
TCGA-22-5482-01A-01D-1632-08	LUSC	b57c316e-1cae-4286-bdbb-8b65c020b3fa
TCGA-22-5485-01A-01D-1632-08	LUSC	448af8b4-e071-48b0-a65b-b4ad17afdc0c
TCGA-22-5489-01A-01D-1632-08	LUSC	c4eb6681-7ec3-4688-b06a-c47a0043f3fb
TCGA-22-5491-01A-01D-1632-08	LUSC	ed4b5a8c-1dae-41a3-8a2a-f54fa51be4b8
TCGA-22-5492-01A-01D-1632-08	LUSC	abc94013-71f5-4ac6-88a4-01b4ef9f9d2f
TCGA-33-4532-01A-01D-1267-08	LUSC	c8baeba2-2a73-41d7-9226-b89a8f42e18f
TCGA-33-4533-01A-01D-1267-08	LUSC	52b8c7c1-2cfe-410d-a738-1dec43109e24
TCGA-33-4538-01A-01D-1267-08	LUSC	e04814f8-a51f-4b6b-a4e9-bd8d2291817c
TCGA-33-4547-01A-01D-1267-08	LUSC	7e622fc2-06c5-4686-a885-e407725c2f08
TCGA-33-4566-01A-01D-1441-08	LUSC	ddd84ea3-dd5e-4f95-97c3-84c107c19cad
TCGA-33-4582-01A-01D-1441-08	LUSC	4cb06585-62f9-4aae-969a-2085b4d514c3
TCGA-33-4583-01A-01D-1441-08	LUSC	fb901997-6e46-436f-ad34-74aadc344245
TCGA-33-4586-01A-01D-1441-08	LUSC	e6bf4288-9fdd-4c56-b6d2-fa2f5ee542b6
TCGA-33-6737-01A-11D-1817-08	LUSC	3b21ce38-16c6-4c68-9104-fa11f1b619b1

TCGA-34-2596-01A-01D-1522-08	LUSC	66e35f68-f4db-46ee-876e-e770ea616ef3
TCGA-34-2600-01A-01D-1522-08	LUSC	167e0f4e-e7d3-4942-885a-cf06419bbe6d
TCGA-34-2608-01A-02D-1522-08	LUSC	3c90209b-b6f6-40b2-a374-6cd37d6d3895
TCGA-34-5231-01A-21D-1817-08	LUSC	c9862ed2-4ba6-434d-a205-b1bda292d218
TCGA-34-5232-01A-21D-1817-08	LUSC	f32fff2f-0bbf-475f-b088-3f1699203c31
TCGA-34-5234-01A-01D-1632-08	LUSC	7b19ae84-2cab-47e7-87df-46c497da17e0
TCGA-34-5236-01A-21D-1817-08	LUSC	46cb2de7-bbe1-4444-b17e-4c5677a05249
TCGA-34-5239-01A-21D-1817-08	LUSC	6e596912-2146-4c4f-97b6-70b610f5d4b4
TCGA-34-5240-01A-01D-1441-08	LUSC	4c3840df-9824-40db-879e-6d24adc8c155
TCGA-34-5241-01A-01D-1441-08	LUSC	0bcdbc37-cde8-47df-9184-621b2b47da5b
TCGA-34-5927-01A-11D-1817-08	LUSC	d717b13a-e487-4cad-9aae-4b0d649236c4
TCGA-34-5928-01A-11D-1817-08	LUSC	9e2d032e-f982-44fc-b6e0-3be82f029689
TCGA-34-5929-01A-11D-1817-08	LUSC	a25de54e-c13d-4973-864a-e307fbe7324a
TCGA-37-3783-01A-01D-1267-08	LUSC	711e9b21-bd8c-4058-a0ce-5ff4dc23b527
TCGA-37-3789-01A-01D-0983-08	LUSC	d732196f-ef85-43ea-aac7-7c9060bf19c5
TCGA-37-4133-01A-01D-1352-08	LUSC	a678cc49-9009-4027-826f-e17f4533538d
TCGA-37-4135-01A-01D-1352-08	LUSC	754dda66-fceb-4f63-bc99-c98aaa86b0c2
TCGA-37-4141-01A-02D-1352-08	LUSC	3d4f4555-d71a-4c7d-8667-c42dcc20c076
TCGA-37-5819-01A-01D-1632-08	LUSC	edf2a2c0-3829-4da2-8960-598fbd5c4c07
TCGA-39-5016-01A-01D-1441-08	LUSC	d63a0a46-7676-40f5-8e03-b8317d243c73
TCGA-39-5019-01A-01D-1817-08	LUSC	6aecd71e-84f1-4b4d-bff6-ede33026f58b
TCGA-39-5021-01A-01D-1441-08	LUSC	4d8b4c6f-e6eb-4799-b64d-119afc691e3d
TCGA-39-5022-01A-21D-1817-08	LUSC	f60928ab-0cb1-4483-8d61-48a5333defbf
TCGA-39-5024-01A-21D-1817-08	LUSC	388478e9-8c1f-43f8-88c4-811bf3cc2500
TCGA-39-5027-01A-21D-1817-08	LUSC	32c14926-b510-4714-90b2-b0bd68569cd4
TCGA-39-5028-01A-01D-1441-08	LUSC	015b9329-ecf2-4410-b7b6-f9313b5d2adb
TCGA-39-5029-01A-01D-1441-08	LUSC	aa02c83c-7ef0-400d-bd8d-729dacda6352
TCGA-39-5030-01A-01D-1441-08	LUSC	9e7b63f2-6080-4bb0-b45d-a0d40dffcbe0
TCGA-39-5031-01A-01D-1441-08	LUSC	3eab4096-8e8e-459d-a2bb-6ef03f414315
TCGA-39-5035-01A-01D-1441-08	LUSC	035fe73e-56b4-4afe-b70e-dd3c34027f2d
TCGA-39-5036-01A-01D-1441-08	LUSC	a1aa5fba-f179-4777-8d49-345a366d12fa
TCGA-39-5037-01A-01D-1441-08	LUSC	825bd82c-f8f8-4776-a7f5-713b3a574955
TCGA-39-5039-01A-01D-1441-08	LUSC	0c14e914-abd4-4406-be82-a810b10a1320
TCGA-43-2578-01A-01D-1522-08	LUSC	7ce90b30-d372-4edb-9807-b71cb5eb4cb7
TCGA-43-3394-01A-01D-0983-08	LUSC	bb72e789-f8ad-4ab5-805b-a9ac21cef0e3
TCGA-43-3920-01A-01D-0983-08	LUSC	a97333f4-d289-493f-8dff-88e52719fa86
TCGA-43-5668-01A-01D-1632-08	LUSC	f01dfe80-aee9-44f6-b32d-3591fbc3c0f5
TCGA-43-6143-01A-11D-1817-08	LUSC	3874253f-7168-4cd6-b1d6-f426fa207313
TCGA-43-6647-01A-11D-1817-08	LUSC	90b97948-26f7-4431-be89-af8c432baae0
TCGA-43-6770-01A-11D-1817-08	LUSC	404ca8c2-f1bb-4749-8abd-87f491a8111c
TCGA-43-6771-01A-11D-1817-08	LUSC	20735861-1f84-4141-a467-f598108e1e41

TCGA-46-3765-01A-01D-0983-08			
TCGA-46-3767-01A-01D-0983-08	TCGA-46-3765-01A-01D-0983-08	LUSC	6c4bb09f-46c8-4a42-bf4f-8bad5316603d
TCGA-46-3768-01A-01D-0983-08	TCGA-46-3766-01A-01D-0983-08	LUSC	0a691892-2209-4f3c-ab16-c2560e4928b4
TCGA.46-3769-01A-01D-0983-08	TCGA-46-3767-01A-01D-0983-08	LUSC	db4ea3ec-e926-4e75-a97b-a527c101b3b9
TCGA-46-6025-01A-11D-1817-08	TCGA-46-3768-01A-01D-0983-08	LUSC	30666313-cc29-4fce-8308-b04fb932083c
TCGA-46-6026-01A-11D-1817-08	TCGA-46-3769-01A-01D-0983-08	LUSC	108a1360-a545-4573-a775-49b3420814e2
TCGA-51-4079-01A-01D-1458-08	TCGA-46-6025-01A-11D-1817-08	LUSC	767a9ae0-2aa4-467b-b9c3-fb3bf701b642
TCGA-51-4080-01A-01D-1458-08	TCGA-46-6026-01A-11D-1817-08	LUSC	42a4a60c-257e-4bf6-a9ba-6f162dbca94a
TCGA-51-4081-01A-01D-1521-08	TCGA-51-4079-01A-01D-1458-08	LUSC	0a43aade-225c-4a29-b1d8-6b930eb8a1db
TCGA-56-1622-01A-01D-1521-08	TCGA-51-4080-01A-01D-1458-08	LUSC	2498ada2-b8d3-4220-8283-45af67a8119a
TCGA-56-5897-01A-11D-1632-08 LUSC 056acb55-f3ba-4ce0-9735-3cfe6516df55 TCGA-56-5898-01A-11D-1632-08 LUSC aaf47efe-4a0a-40d1-b70f-9c9168cbdae0 TCGA-56-6545-01A-11D-1817-08 LUSC 16756a08-8308-4ad3-9c21-2cea0cd7028e TCGA-56-6546-01A-11D-1817-08 LUSC 87e71949-5bd9-458c-9517-4b19882c2b4f TCGA-60-2698-01A-01D-1522-08 LUSC 2045c788-9ea8-4ea5-a5e3-65(c16a62adb TCGA-60-2707-01A-01D-1522-08 LUSC 3371189b-5808-4408-824e-8dacce925cc5 TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacce925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2709-01A-21D-1817-08 LUSC 4321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e45fa TCGA-60-2711-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2720-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2721-01A-01D-1522-08 LUSC </td <td>TCGA-51-4081-01A-01D-1458-08</td> <td>LUSC</td> <td>1492c429-1041-4d86-9358-c9b9babd1401</td>	TCGA-51-4081-01A-01D-1458-08	LUSC	1492c429-1041-4d86-9358-c9b9babd1401
TCGA-56-5898-01A-11D-1632-08 LUSC aaf47efe-4a0a-40d1-b70f-9c9168cbdae0 TCGA-56-6545-01A-11D-1817-08 LUSC 16756a08-8308-4ad3-9c21-2cea0cd7028e TCGA-56-6546-01A-11D-1817-08 LUSC 87e71949-5bd9-458c-95f7-4b19882c2b4f TCGA-60-2698-01A-01D-1522-08 LUSC 2045c788-9ea8-4ea5-a5c3-65fc16a62adb TCGA-60-2708-01A-01D-1522-08 LUSC 5d1fa470-2789-4576-9743-0362af682c1d TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacec925cc5 TCGA-60-2710-01A-01D-1522-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2711-01A-01D-1522-08 LUSC faecb1fc-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85ce9-31bc-dcea-9c54-13b7c0c645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e06a68-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC	TCGA-56-1622-01A-01D-1521-08	LUSC	0bbc7ede-5022-4084-925c-d65baaf7abc2
TCGA-56-6545-01A-11D-1817-08 LUSC 16756a08-8308-4ad3-9e21-2cea0cd7028e TCGA-56-6546-01A-11D-1817-08 LUSC 87e71949-5bd9-458c-95f7-4b19882c2b4f TCGA-60-2698-01A-01D-1522-08 LUSC 2045c788-9ea8-4ea5-a5e3-65fc16a62adb TCGA-60-2707-01A-01D-1522-08 LUSC 5d1fa470-2789-4576-9743-0362af682c1d TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacec925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faceb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9c54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2721-01A-01D-1522-08 LUSC 8e4fff62-9395-47cb-b919-4b8487dea8e TCGA-60-2721-01A-01D-1522-08 LUSC 8deffff62-9395-47cb-b19-4b8487dea8e TCGA-60-2721-01A-01D-1522-08 LUSC 8deffff62-9395-47cb-b19-4b8487dea8e TCGA-60-2722-01A-01D-1522-08 LUSC </td <td>TCGA-56-5897-01A-11D-1632-08</td> <td>LUSC</td> <td>056acb55-f3ba-4ce0-9735-3cfe6516df55</td>	TCGA-56-5897-01A-11D-1632-08	LUSC	056acb55-f3ba-4ce0-9735-3cfe6516df55
TCGA-56-6546-01A-11D-1817-08 LUSC 87e71949-5bd9-458c-95f7-4b19882c2b4f TCGA-60-2698-01A-01D-1522-08 LUSC 2045c788-9ea8-4ea5-a5e3-65fc16a62adb TCGA-60-2707-01A-01D-1522-08 LUSC 5d1fa470-2789-4576-9743-0362af682c1d TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacec925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faceb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e65a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2720-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2725-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-0078052a5732 TCGA-60-2725-01A-01D-1522-08 LUS	TCGA-56-5898-01A-11D-1632-08	LUSC	aaf47efe-4a0a-40d1-b70f-9c9168cbdae0
TCGA-60-2698-01A-01D-1522-08 LUSC 2045c788-9ea8-4ea5-a5e3-e5fc16a62adb TCGA-60-2707-01A-01D-1522-08 LUSC 5d1fa470-2789-4576-9743-0362af682c1d TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8daece925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faecb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9c54-13b7c0c645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC a8c6c68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2725-01A-01D-1522-08 LUS	TCGA-56-6545-01A-11D-1817-08	LUSC	16756a08-8308-4ad3-9e21-2cea0cd7028e
TCGA-60-2707-01A-01D-1522-08 LUSC 5d1fa470-2789-4576-9743-0362af682c1d TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacec925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faecb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6622dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2721-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-403-5r9-e15240dc6dd4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-403-5r9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c651p-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1670-08 LUSC <td>TCGA-56-6546-01A-11D-1817-08</td> <td>LUSC</td> <td>87e71949-5bd9-458c-95f7-4b19882c2b4f</td>	TCGA-56-6546-01A-11D-1817-08	LUSC	87e71949-5bd9-458c-95f7-4b19882c2b4f
TCGA-60-2708-01A-01D-1522-08 LUSC a371189b-5808-4408-824e-8dacec925cc5 TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faccb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2713-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05ca30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8deffff2-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4002-82e8-6b10391aae5d TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2725-01A-01D-152-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2726-01A-01D-152-08 LUSC	TCGA-60-2698-01A-01D-1522-08	LUSC	2045c788-9ea8-4ea5-a5e3-65fc16a62adb
TCGA-60-2709-01A-21D-1817-08 LUSC 4f321c92-ae27-4253-bd8b-4505ba8c7dc4 TCGA-60-2710-01A-01D-1522-08 LUSC faecb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4003-b7c9-e15240dc6dd4 TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2723-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2726-01A-01D-1522-08 LUS	TCGA-60-2707-01A-01D-1522-08	LUSC	5d1fa470-2789-4576-9743-0362af682c1d
TCGA-60-2710-01A-01D-1522-08 LUSC faecb1fe-b4ef-434d-818c-81ad2167dd25 TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2715-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC 38rc6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1522-08 LUSC 396eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC 396eddfc-3afb-4bf8-a440-c91778113fbd TCGA-66-2726-01A-01D-1817-08 LUS	TCGA-60-2708-01A-01D-1522-08	LUSC	a371189b-5808-4408-824e-8dacec925cc5
TCGA-60-2711-01A-01D-1522-08 LUSC 2ed85cc9-31bc-4cea-9e54-13b7c0e645fa TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2715-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-66-2726-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-66-2727-01A-01D-0983-08 LUS	TCGA-60-2709-01A-21D-1817-08	LUSC	4f321c92-ae27-4253-bd8b-4505ba8c7dc4
TCGA-60-2712-01A-01D-1522-08 LUSC 6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30 TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2715-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391ae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ea35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-66-2727-01A-01D-0983-08 LUSC c3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2734-01A-01D-0983-08 LUSC	TCGA-60-2710-01A-01D-1522-08	LUSC	faecb1fe-b4ef-434d-818c-81ad2167dd25
TCGA-60-2713-01A-01D-1522-08 LUSC 79eb7bba-f0d8-462c-add7-20a2fb7843e1 TCGA-60-2715-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ea35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2744-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2744-01A-01D-0983-08 LUS	TCGA-60-2711-01A-01D-1522-08	LUSC	2ed85cc9-31bc-4cea-9e54-13b7c0e645fa
TCGA-60-2715-01A-01D-1522-08 LUSC 8e05a30d-2177-45e0-90fd-8c5961268c39 TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1522-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2744-01A-01D-0983-08 LUS	TCGA-60-2712-01A-01D-1522-08	LUSC	6662dd1b-3e4f-4b7a-b603-cfa7fd92fc30
TCGA-60-2719-01A-01D-1522-08 LUSC ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259 TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUS	TCGA-60-2713-01A-01D-1522-08	LUSC	79eb7bba-f0d8-462c-add7-20a2fb7843e1
TCGA-60-2720-01A-01D-1522-08 LUSC 3b435ddf-a496-40a2-82e8-6b10391aae5d TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2715-01A-01D-1522-08	LUSC	8e05a30d-2177-45e0-90fd-8c5961268c39
TCGA-60-2721-01A-01D-1522-08 LUSC 8defff62-9395-47cb-bb19-4b8487d9ea8e TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2719-01A-01D-1522-08	LUSC	ee6cc68e-8d2a-41ee-82c6-0fecdf7e6259
TCGA-60-2722-01A-01D-1522-08 LUSC eb955f72-83bf-4635-a7ed-89e4d66e08f4 TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2720-01A-01D-1522-08	LUSC	3b435ddf-a496-40a2-82e8-6b10391aae5d
TCGA-60-2723-01A-01D-1522-08 LUSC 8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4 TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2721-01A-01D-1522-08	LUSC	8defff62-9395-47cb-bb19-4b8487d9ea8e
TCGA-60-2724-01A-01D-1522-08 LUSC 387c6519-6529-4074-a5ab-00f8052a5732 TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2722-01A-01D-1522-08	LUSC	eb955f72-83bf-4635-a7ed-89e4d66e08f4
TCGA-60-2725-01A-01D-1267-08 LUSC f3ed705b-e5aa-4756-9794-e4b85303693a TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2723-01A-01D-1522-08	LUSC	8a6aa45a-ef6d-4005-b7c9-e15240dc6dd4
TCGA-60-2726-01A-01D-1522-08 LUSC a96eddfc-3afb-4bf8-a440-c91778113fbd TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2724-01A-01D-1522-08	LUSC	387c6519-6529-4074-a5ab-00f8052a5732
TCGA-63-5128-01A-01D-1441-08 LUSC d3b9b51e-eeea-4355-829d-ee35bdd2cf5b TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2725-01A-01D-1267-08	LUSC	f3ed705b-e5aa-4756-9794-e4b85303693a
TCGA-63-5131-01A-01D-1441-08 LUSC b290a86e-22da-4f10-a421-2616bb47bc1b TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-60-2726-01A-01D-1522-08	LUSC	a96eddfc-3afb-4bf8-a440-c91778113fbd
TCGA-63-6202-01A-11D-1817-08 LUSC a3c568a6-0c43-47a7-a35a-3225fedeeb44 TCGA-66-2727-01A-01D-0983-08 LUSC c2b2c909-1461-42ce-8fd9-736147dcacd8 TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-63-5128-01A-01D-1441-08	LUSC	d3b9b51e-eeea-4355-829d-ee35bdd2cf5b
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TCGA-66-2734-01A-01D-0983-08 LUSC 9f7a24a2-10e2-4039-ad27-13d7ec28ff36 TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43be1a37-b18e-4e96-89e6-ed6ee1d8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-63-6202-01A-11D-1817-08	LUSC	a3c568a6-0c43-47a7-a35a-3225fedeeb44
TCGA-66-2742-01A-01D-0983-08 LUSC 07047a99-45bd-4df6-ad6f-934a48e8e213 TCGA-66-2744-01A-01D-0983-08 LUSC 43bela37-bl8e-4e96-89e6-ed6eeld8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-66-2727-01A-01D-0983-08	LUSC	c2b2c909-1461-42ce-8fd9-736147dcacd8
TCGA-66-2744-01A-01D-0983-08 LUSC 43bela37-b18e-4e96-89e6-ed6eeld8e65a TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-66-2734-01A-01D-0983-08	LUSC	9f7a24a2-10e2-4039-ad27-13d7ec28ff36
TCGA-66-2754-01A-01D-0983-08 LUSC c34a64c8-3746-44f8-a7ee-77f502b6256c	TCGA-66-2742-01A-01D-0983-08	LUSC	07047a99-45bd-4df6-ad6f-934a48e8e213
	TCGA-66-2744-01A-01D-0983-08	LUSC	43be1a37-b18e-4e96-89e6-ed6ee1d8e65a
TCGA-66-2755-01A-01D-1522-08 LUSC 177d64a9-65dc-4aa1-8774-bd8208e40f04	TCGA-66-2754-01A-01D-0983-08	LUSC	c34a64c8-3746-44f8-a7ee-77f502b6256c
	TCGA-66-2755-01A-01D-1522-08	LUSC	177d64a9-65dc-4aa1-8774-bd8208e40f04

TCGA-66-2756-01A-01D-1522-08	LUSC	472c95e6-eccb-4988-be16-fdace73b2ed8
TCGA-66-2757-01A-01D-1522-08	LUSC	1886dba0-4662-4342-84ac-96af0beb2393
TCGA-66-2758-01A-02D-1522-08	LUSC	71c4e854-a704-4787-a37a-fa6642ca5dac
TCGA-66-2759-01A-01D-1522-08	LUSC	fecd0a2b-d176-438a-be95-306f453fde40
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TCGA-66-2766-01A-01D-1522-08	LUSC	452b75d0-1818-46aa-8804-9cfc0bd66449
TCGA-66-2767-01A-01D-1522-08	LUSC	ca748128-272c-4fad-9a1f-01328b93b3f4
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TCGA-66-2770-01A-01D-1522-08	LUSC	e417903d-ab76-44f0-aae9-3a91fa9a8d3c
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TCGA-66-2773-01A-01D-1267-08	LUSC	fb0b515b-afc4-40c3-abe6-e90c442f0249
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TCGA-66-2780-01A-01D-1522-08	LUSC	d088bd17-a1a0-4bd9-bfe1-d57b5725c53b
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TCGA-66-2782-01A-01D-1522-08	LUSC	640ff507-203c-45aa-8bc1-030ee8639b5d
TCGA-66-2783-01A-01D-1267-08	LUSC	f574d3b7-4ae4-49bc-9e05-f965fbc86119
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TCGA-66-2786-01A-01D-1522-08	LUSC	999a6582-33cf-47ca-b268-9b2da102e99b
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TCGA-66-2788-01A-01D-0983-08	LUSC	2466d424-98bb-4380-9967-36abaa0e69d7
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TCGA-85-6561-01A-11D-1817-08	LUSC	f5aa0f1c-da19-4c04-b695-01ed5b20e79e
TCGA-04-1332-01A-01W-0488-09	ov	b52e5d90-dc57-438c-9c38-e043308c24ac
TCGA-04-1336-01A-01W-0488-09	OV	586101df-93c9-4d0b-ba0e-58df7a2f9598
TCGA-04-1343-01A-01W-0488-09	ov	fbbc3d80-aff2-463e-8eb3-c4361ad7cb98
TCGA-04-1346-01A-01W-0488-09	ov	9f494df7-f64f-4935-ae42-eeb0b94624dc
TCGA-04-1347-01A-01W-0488-09	OV	21b50b8c-781a-4e15-a4ad-715f416f0fa2
TCGA-04-1348-01A-01W-0494-09	ov	1f4dee42-8f3d-4307-b6e5-3381d77d201c
TCGA-04-1349-01A-01W-0494-09	OV	e456f707-f0a0-4624-98bc-e9dfe779182b

TCGA-04-1361-01A-01W-0494-09	OV	0fc567bd-2201-4f3d-820e-2c0dbe58da6f
TCGA-04-1362-01A-01W-0494-09	OV	830e207f-458e-4628-b7bc-287c2f2e12e5
TCGA-04-1542-01A-01W-0553-09	OV	317a63af-e862-43df-8ef5-7c555b2cb678
TCGA-09-0366-01A-01W-0372-09	OV	62269d21-50dc-42b0-b1e4-75ed8010080a
TCGA-09-0369-01A-01W-0372-09	OV	633f5c4d-c224-404c-9f68-24daafd1fc84
TCGA-10-0930-01A-02W-0421-09	OV	ec98ed86-1d2f-4e54-b2d4-5976469bf0b8
TCGA-10-0933-01A-01W-0421-09	ov	3ec4215f-b57d-4ae7-b247-55ea1f7e97d3
TCGA-10-0935-01A-03W-0421-09	OV	af0edbf4-9d90-4373-a9ce-0875ebbe1d04
TCGA-13-0723-01A-02W-0372-09	OV	6f9e5a76-5d2a-4bb0-babf-3f365a177236
TCGA-13-0724-01A-01W-0372-09	OV	2b6aa1c8-5150-4d8f-af59-d5a826321308
TCGA-13-0726-01A-01W-0372-09	OV	201415c2-5b5a-4bb8-8005-bf2c78d4d88e
TCGA-13-0755-01A-01W-0372-09	OV	9bd227fa-e52a-4805-bd04-ad63df0930af
TCGA-13-0760-01A-01W-0372-09	OV	5181630f-246a-4cb4-88c2-1534b5fb8e37
TCGA-13-0765-01A-01W-0372-09	OV	5bcfe3ea-d95e-47ff-9718-6b123d3acaef
TCGA-13-0791-01A-01W-0372-09	OV	70f63e2f-9bc6-4ed9-8d91-f1889287d7b7
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TCGA-13-0800-01A-01W-0372-09	OV	757862e3-0392-4e05-a242-25e3d2094ee8
TCGA-13-0804-01A-01W-0372-09	OV	7f39610d-45b8-45ae-806e-16b7acebafa6
TCGA-13-0807-01B-02W-0421-09	OV	f80466d9-6cc8-461b-acc2-addee22bd42a
TCGA-13-0884-01B-01W-0494-09	OV	c5f0aa38-556b-401c-b4da-ac82cdc2e637
TCGA-13-0885-01A-02W-0421-09	OV	a530d9a9-b21e-47be-b4d8-1707b71f360a
TCGA-13-0887-01A-01W-0421-09	OV	e05146f2-688d-416b-a992-e2c7a2b7b244
TCGA-13-0890-01A-01W-0421-09	OV	15b867fb-7a7b-4158-9abd-91870ba77eb7
TCGA-13-0893-01B-01W-0494-09	OV	a335ab49-84b7-4d3b-a03d-9c3931904ca5
TCGA-13-0894-01B-01W-0494-09	OV	eb57990e-702f-4fac-9ef5-7447ecb45cec
TCGA-13-0897-01A-01W-0421-09	OV	f48ed68f-a833-4b78-971a-3c746c563d24
TCGA-13-0903-01A-01W-0421-09	OV	854167b5-03ab-4867-af34-9c92e385822e
TCGA-13-0910-01A-01W-0421-09	ov	26cebe0b-b7a7-431e-bc12-7fda22af72f3
TCGA-13-0912-01A-01W-0421-09	ov	517f4d7f-c962-414f-8824-f2a7ae19cb6d
TCGA-13-0920-01A-01W-0421-09	OV	2e28969b-c9a9-41ec-80bf-f583197b7f92
TCGA-13-0924-01A-01W-0421-09	OV	510dda3c-6a1f-4781-972f-c9c270608c72
TCGA-13-1403-01A-01W-0494-09	OV	acbc77ba-7cc0-4af2-9ab6-0c835ce33998
TCGA-13-1404-01A-01W-0494-09	OV	692e4b24-daf0-4771-b4a6-b0599f122ad8
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TCGA-13-1411-01A-01W-0494-09	OV	e254d7f4-1edf-4054-9ca6-9fe058a05484
TCGA-13-1412-01A-01W-0494-09	OV	f7edafe2-3eab-4bac-9d25-ed5c223b4aee
TCGA-13-1481-01A-01W-0549-09	OV	f9eab025-5518-4240-b1a8-19f8ff8354f0
TCGA-13-1482-01A-01W-0549-09	OV	a68927d4-e827-49c9-9c3a-23ce0543261b
TCGA-13-1483-01A-01W-0549-09	ov	52280c07-44f5-4e9c-8601-7455b5b0de7a
TCGA-13-1488-01A-01W-0549-09	ov	886a8c10-63cf-4cb2-83d2-5a99bbda193d
TCGA-13-1489-01A-01W-0549-09	OV	395c1d93-7216-4c9d-bfad-26ff95fb8afe

TCGA-13-1491-01A-01W-0549-09	OV	fb7d1c2b-3e87-4d05-a58b-92d0e1016986
TCGA-13-1497-01A-01W-0549-09	OV	04e814c6-ea28-4ade-bc8f-a618552943da
TCGA-13-1498-01A-01W-0549-09	OV	b00d9680-4099-43fe-87de-b3cc8b9e70c8
TCGA-13-1499-01A-01W-0549-09	OV	b4ce07b1-677e-4a9c-8f8e-2b7762487692
TCGA-13-1506-01A-01W-0549-09	OV	7534b542-88f8-445c-ae4a-9f44fb6798a8
TCGA-13-1507-01A-01W-0549-09	OV	5423db1a-5b59-4a5b-a676-00a54570b04a
TCGA-13-1509-01A-01W-0549-09	OV	4d3fab96-bc22-48d0-a3ef-1844ad894d0f
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TCGA-23-1022-01A-02W-0488-09	OV	160a0e7d-315e-4de3-a7d4-928412fd909c
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TCGA-23-1123-01A-01W-0488-09	OV	22cfe2c8-5e1f-4b64-854d-2a7a02bf10fe
TCGA-23-1124-01A-01W-0488-09	OV	8a4061a0-77f2-4bb4-a3da-9b3d9f0314b9
TCGA-24-0966-01A-01W-0977-09	OV	dc069342-661a-4012-9bda-0c67469e117d
TCGA-24-0980-01A-01W-0421-09	OV	87d32a92-a8d2-4656-a100-798328338486
TCGA-24-0982-01A-01W-0488-09	OV	7667c0e6-e44a-448f-b118-6e2171a99b6c
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TCGA-24-1413-01A-01W-0494-09	OV	1b2d2cde-4553-472e-82f1-8224745ac1eb
TCGA-24-1416-01A-01W-0549-09	OV	21f5e805-c0b4-487b-9ccd-02963e2369ff
TCGA-24-1417-01A-01W-0549-09	OV	f6f43d04-a9e3-48c8-a276-3bebcaf416d7
TCGA-24-1418-01A-01W-0549-09	OV	6093bcb5-4889-4cb9-9b01-e4e4278e72aa
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TCGA-24-1425-01A-02W-0553-09	OV	f8d4c37d-5b4d-4f5a-8022-7da2b32cc1b0
TCGA-24-1426-01A-01W-0549-09	OV	063f8696-2c9d-4af4-a863-df10c42a5ea8
TCGA-24-1427-01A-01W-0549-09	OV	6511d3d4-722c-4702-a644-29bb98e5e5c3
TCGA-24-1428-01A-01W-0549-09	OV	52866517-eddf-4d63-a121-a296d6b2d264
TCGA-24-1435-01A-01W-0549-09	OV	28d236f6-dddc-48c2-be30-b1568a4d6055
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TCGA-24-1463-01A-01W-0549-09	OV	c01ca9e7-ee9b-4698-8e4d-920ad7bfbe5f
TCGA-24-1464-01A-01W-0549-09	OV	01ec3cbb-c68a-4874-b396-f5e34876e04a
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TCGA-24-1563-01A-01W-0553-09	OV	b6c46b53-f94d-4936-9005-518c8f1c1449
TCGA-24-1616-01A-01W-0553-09	OV	c464b2f6-9cfe-463a-b5e3-9a76cd4480c5
TCGA-25-1315-01A-01W-0494-09	OV	52f45b5e-af86-454c-be63-a56c6c21b730
TCGA-25-1316-01A-01W-0494-09	OV	d75a0b16-04e4-4ba3-a695-132c5ace698b
TCGA-25-1322-01A-01W-0494-09	OV	626f1798-fb15-4b01-8d8f-db19777d72e9
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TCGA-AF-3913-01A-02W-1073-09 READ 4ebe7cf9-ce4f-485d-9332-ca9b536c38e2 TCGA-AG-3887-01A-01W-1073-09 READ 6d2de0f5-e812-4d3f-903b-7febdcfed2f7 TCGA-AG-3890-01A-01W-1073-09 READ 042e984f-e106-4b23-9908-5abataf07c694 TCGA-AG-3892-01A-01W-1073-09 READ 26acdac6-b01a-4dbd-b0b8-f6d97fe01808 TCGA-AG-3893-01A-01W-1073-09 READ 0faa6d28-c01c-4847-9552-912733485610 TCGA-AG-3894-01A-01W-1073-09 READ 2508d0c8-cdaf-463f-bb03-47af1bc41866 TCGA-AG-3898-01A-01W-1073-09 READ 22c7d09a-e69b-44be-8d8e-0a0cc9adf57c TCGA-AG-3898-01A-01W-1073-09 READ c3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-3999-01A-01W-1073-09 READ 55075176-07a4-4183-9f8r-9f472b15a6b4 TCGA-AG-4001-01A-02W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4004-01A-01W-1073-09 READ 8cfctdc8f-22c0-dc3a-9e46-58c0a68e818e TCGA-AG-4004-01A-01W-1073-09 RE			
TCGA-AG-3890-01A-01W-1073-09 READ 042e984f-c106-4b23-9908-5abaf407e694 TCGA-AG-3892-01A-01W-1073-09 READ 26acdae6-b01a-4dbd-b0b8-f6d97fe01808 TCGA-AG-3893-01A-01W-1073-09 READ 0faa6d28-c01c-4847-9552-912733485610 TCGA-AG-3894-01A-01W-1073-09 READ e508d0c8-cdaf-463f-bb03-47af1bc41866 TCGA-AG-3898-01A-01W-1073-09 READ 22c7d09a-e69b-44be-8d8e-0a0cc9adf57c TCGA-AG-3898-01A-01W-1073-09 READ cc3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-4f88-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ 0445426d-b9c-4ce5-b1cc-cb236d4381cf TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ 6fcdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-400-01A-01W-1073-09 READ 6fcdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-400-01A-01W-1073-09 READ 3cdsc15-8eab-4d6-9ga2-36ec719f6774 TCGA-AG-4008-01A-01W-0073-09 READ	TCGA-AF-3913-01A-02W-1073-09	READ	4ebe7cf9-ce4f-485d-9332-ea9b536e38e2
TCGA-AG-3892-01A-01W-1073-09 READ 26acdae6-b01a-4dbd-b0b8-f6d97fe01808 TCGA-AG-3893-01A-01W-1073-09 READ 0faa6d28-c01c-4847-9552-912733485610 TCGA-AG-3894-01A-01W-1073-09 READ e508d0c8-cdaf-463f-bb03-47af1bc41866 TCGA-AG-3896-01A-01W-1073-09 READ 22c7d09a-e69b-44be-8d8e-0a0cc9adf57c TCGA-AG-3898-01A-01W-1073-09 READ cc3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-3909-01A-01W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4001-01A-02W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 82cd3c15-8eab-4dd6-b9a2-36ee719f6774 TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4dd6-b9a2-36ee719f6774 TCGA-AG-A008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0c7-507181b3fb86 TCGA-AG-AOOC-01A-01W-A005-10 REA	TCGA-AG-3887-01A-01W-1073-09	READ	6d2de0f5-e812-4d3f-903b-7febdcfcd2f7
TCGA-AG-3893-01A-01W-1073-09 READ Ofaa6d28-c01c-4847-9552-912733485610 TCGA-AG-3894-01A-01W-1073-09 READ e50840c8-cdaf-463f-bb03-47af1bc41866 TCGA-AG-3896-01A-01W-1073-09 READ 22c7d09a-e69b-44be-848e-0a0cc9adf57c TCGA-AG-3898-01A-01W-1073-09 READ c3516ba-2941-4cfa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f2-ab75-b27a4cf27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ece3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4007-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-0005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1d648a-cfc2-4801-ac94-94c5d8058a9f TCGA-AG-A00H-01A READ <	TCGA-AG-3890-01A-01W-1073-09	READ	042e984f-c106-4b23-9908-5abaf407e694
TCGA-AG-3894-01A-01W-1073-09 READ e508d0c8-cdaf-463f-bb03-47af1bc41866 TCGA-AG-3896-01A-01W-1073-09 READ 22c7d09a-e69b-44be-8d8e-0a0ce9adf57c TCGA-AG-3898-01A-01W-1073-09 READ cc3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ecc3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ec5-b1c-c-b236d4381cf TCGA-AG-3099-01A-01W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4001-01A-02W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4005-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-AOOC-01A-01W-A005-10 READ b50ae1df-eeff-4a5e-ba4b-e962d740ab22 TCGA-AG-AOOH-01A-01W-A005-10 READ b50ae1df-eeff-4a5e-ba4b-e962d740ab22 TCGA-AG-AO11-01A READ	TCGA-AG-3892-01A-01W-1073-09	READ	26acdae6-b01a-4dbd-b0b8-f6d97fe01808
TCGA-AG-3896-01A-01W-1073-09 READ 22c7d09a-e69b-44be-8d8e-0a0ce9adf57c TCGA-AG-3898-01A-01W-1073-09 READ cc3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ecc3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ec5-b1c-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4005-01A-01W-1073-09 READ 6fefdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 2c16f8e0f-04bf-4a0d-93a-36ee719f6774 TCGA-AG-4008-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-93a-36e804ba6c1607 TCGA-AG-AO08-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-AO04-01A-01W-A005-10 READ 15dd4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-AO0H-01A-01W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-e962d740ab22 TCGA-AG-AO11-01A READ	TCGA-AG-3893-01A-01W-1073-09	READ	0faa6d28-c01c-4847-9552-912733485610
TCGA-AG-3898-01A-01W-1073-09 READ cc3516ba-2941-4efa-80fc-7b5041194d52 TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ece3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4005-01A-01W-1073-09 READ 6fefde8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A0H-01A-01W-A005-09 READ b50ae1d-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A01-01A-01W-A005-10 READ b50ae1d-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A01-01A-01W-A005-10 READ <td>TCGA-AG-3894-01A-01W-1073-09</td> <td>READ</td> <td>e508d0c8-cdaf-463f-bb03-47af1bc41866</td>	TCGA-AG-3894-01A-01W-1073-09	READ	e508d0c8-cdaf-463f-bb03-47af1bc41866
TCGA-AG-3901-01A-01W-1073-09 READ 84859471-1136-4f42-ab75-b27a4ef27199 TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ece3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-3999-01A-01W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4001-01A-02W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfd8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A000-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ f6c4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A014-01A READ fb7a61	TCGA-AG-3896-01A-01W-1073-09	READ	22c7d09a-e69b-44be-8d8e-0a0cc9adf57c
TCGA-AG-3902-01A-01W-1073-09 READ b679f02d-f48d-49eb-b245-65f341e4c181 TCGA-AG-3909-01A-01W-1073-09 READ f5ece3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4008-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-93ae-8034ba6c1607 TCGA-AG-A008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-AO0Y-01A-02W-A005-10 READ b50ae1df-ee6f-445e-ba4b-c962d740ab22 TCGA-AG-AO11-01A READ b5dd8f49-26fc-4849-a964-d8ebdcca9e19 TCGA-AG-AO14-01A READ fb6af1fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-AO15-01A-01W-A005-10 READ f20ae	TCGA-AG-3898-01A-01W-1073-09	READ	cc3516ba-2941-4efa-80fc-7b5041194d52
TCGA-AG-3909-01A-01W-1073-09 READ f5ece3cf-39eb-4277-8975-986e548bc1ea TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00W-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-AOH-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-AO0Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-AO11-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-AO14-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-AO15-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-AO16-01A-01W-A005-10 READ b034c9	TCGA-AG-3901-01A-01W-1073-09	READ	84859471-1136-4f42-ab75-b27a4ef27199
TCGA-AG-3999-01A-01W-1073-09 READ 0445426d-b9c0-4ce5-b1cc-cb236d4381cf TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00C-01A-01W-A005-10 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00H-01A-01W-A00E-09 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5d8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-AO25-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-AO25-01A-01W-A00E-09 READ 7b5a3c	TCGA-AG-3902-01A-01W-1073-09	READ	b679f02d-f48d-49eb-b245-65f341e4c181
TCGA-AG-4001-01A-02W-1073-09 READ 55075176-07a4-4183-9f8f-9f472b15a6b4 TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-1073-09 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A008-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A005-10 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00H-01A-01W-A005-09 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A01L-01A READ f02ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02C-01A-01W-A00E-09 READ 954527dc-8a7d-474	TCGA-AG-3909-01A-01W-1073-09	READ	f5ece3cf-39eb-4277-8975-986e548bc1ea
TCGA-AG-4005-01A-01W-1073-09 READ be1d3bda-de1a-4768-a2e4-22c07326ddc3 TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-AO0Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02C-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40b	TCGA-AG-3999-01A-01W-1073-09	READ	0445426d-b9c0-4ce5-b1cc-cb236d4381cf
TCGA-AG-4007-01A-01W-1073-09 READ 6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A01L-01A READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A02S-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-4001-01A-02W-1073-09	READ	55075176-07a4-4183-9f8f-9f472b15a6b4
TCGA-AG-4008-01A-01W-1073-09 READ 83cd3c15-8eab-4d46-b9a2-36ee719f6774 TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-4005-01A-01W-1073-09	READ	be1d3bda-de1a-4768-a2e4-22c07326ddc3
TCGA-AG-4015-01A-01W-1073-09 READ cf6f8e0f-04bf-4a0d-933e-8034ba6c1607 TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-4007-01A-01W-1073-09	READ	6fcfdc8f-22c0-4c3a-9e46-58c0a68e818e
TCGA-AG-A008-01A-01W-A005-10 READ 2221cfc4-b324-4329-ad37-3dd9a5adf36e TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-4008-01A-01W-1073-09	READ	83cd3c15-8eab-4d46-b9a2-36ee719f6774
TCGA-AG-A00C-01A-01W-A005-10 READ 1a4f95be-32d3-4202-a0e7-507181b3fb86 TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-4015-01A-01W-1073-09	READ	cf6f8e0f-04bf-4a0d-933e-8034ba6c1607
TCGA-AG-A00H-01A-01W-A00E-09 READ fdc4c8ac-fee2-4801-ae94-94c5d8058a9f TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A008-01A-01W-A005-10	READ	2221cfc4-b324-4329-ad37-3dd9a5adf36e
TCGA-AG-A00Y-01A-02W-A005-10 READ b50ae1df-ee6f-4a5e-ba4b-c962d740ab22 TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A00C-01A-01W-A005-10	READ	1a4f95be-32d3-4202-a0e7-507181b3fb86
TCGA-AG-A011-01A READ b5dd8f49-26fc-48d9-a964-d8ebdcca9e19 TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A00H-01A-01W-A00E-09	READ	fdc4c8ac-fee2-4801-ae94-94c5d8058a9f
TCGA-AG-A014-01A READ fbfa61fe-4fb7-4b2a-9bf0-33140fd41873 TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A00Y-01A-02W-A005-10	READ	b50ae1df-ee6f-4a5e-ba4b-c962d740ab22
TCGA-AG-A015-01A-01W-A005-10 READ abb751f0-c4df-4556-ac9b-ad1e1971cccf TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A011-01A	READ	b5dd8f49-26fc-48d9-a964-d8ebdcca9e19
TCGA-AG-A016-01A-01W-A005-10 READ f20ae301-b10b-4dfa-9169-04bc6c3d103a TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A014-01A	READ	fbfa61fe-4fb7-4b2a-9bf0-33140fd41873
TCGA-AG-A01L-01A READ b034c90b-d0bd-466a-88ba-b61efd36c6e4 TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A015-01A-01W-A005-10	READ	abb751f0-c4df-4556-ac9b-ad1e1971cccf
TCGA-AG-A025-01A-01W-A00E-09 READ 7b5a3c33-cd13-4e4d-a1f8-3405dab5998f TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A016-01A-01W-A005-10	READ	f20ae301-b10b-4dfa-9169-04bc6c3d103a
TCGA-AG-A02G-01A-01W-A00E-09 READ 954527dc-8a7d-474d-b580-82199e86cb5a TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A01L-01A	READ	b034c90b-d0bd-466a-88ba-b61efd36c6e4
TCGA-AG-A02X-01A-01W-A00E-09 READ 9ffb8919-a98c-40bd-bdad-146b1ccc14ef	TCGA-AG-A025-01A-01W-A00E-09	READ	7b5a3c33-cd13-4e4d-a1f8-3405dab5998f
	TCGA-AG-A02G-01A-01W-A00E-09	READ	954527dc-8a7d-474d-b580-82199e86cb5a
TCGA-AG-A032-01A-01W-A00E-09 READ 7522eb6b-797a-4964-8aca-6d70590b5f9f	TCGA-AG-A02X-01A-01W-A00E-09	READ	9ffb8919-a98c-40bd-bdad-146b1ccc14ef
	TCGA-AG-A032-01A-01W-A00E-09	READ	7522eb6b-797a-4964-8aca-6d70590b5f9f

Pipeline for prediction of peptides derived from gene mutations with binding to personal HLA alleles: MHC-binding affinity was predicted across all possible 9-mer and 10-mer peptides generated from each somatic mutation and the corresponding wildtype peptide using NetMHCpan (version 2.4). These tiled peptides were analyzed for their binding affinities (IC50 nM) to each class I alleles in the patients' HLA profile. An IC50 value of less than 150 nM was considered a predicted strong to intermediate binder, an IC50 of 150-500 nM was

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considered a predicted weak binder, while an IC50 > 500 nM was considered a non-binder. Experimental confirmation of predicted peptides binding to HLA molecules (IC50 < 500 nM) was performed using a competitive MHC class I allele-binding assay and has been described in detail elsewhere (Cai et al. 28 and Sidney et al. 2001).

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Sources of antigen: Peptides were synthesized to >95% purity (confirmed by high performance liquid chromatography) from New England Peptide (Gardner, MA); or RS Synthesis, (Louisville, KY). Peptides were reconstituted in DMSO (10 mg/ml) and stored at – 80°C until use. Minigenes comprised of a sequence of 300 bp encompassing mut or wt *FNDC3B* were PCR-cloned from Pt 2's tumor into the expression vector pcDNA3.1 using the following primers: 5'primer: GACGTCGGATCCCACCATGGGTCCCGGAATTAAGAAAACAGAG; 3' primer:

CCCGGGGCCGCCTAATGGTGATGGTGATGGTGACATTCTAATTCTTCTCCACTG TAAA. Minigenes were expressed in antigen-presenting target cells by introducing 20 µg of the plasmid into 2 million K562 cells (ATCC) stably transfected with HLA-A2 by Amaxa nucleofection (Solution V, Program T16, Lonza Inc; Walkersville, MD). Cells were incubated in RPMI media (Cellgro; Manassas, VA), supplemented with 10% fetal bovine serum (Cellgro), 1% HEPES buffer (Cellgro), and 1% L-glutamine (Cellgro). The cells were harvested 2 days following nucleofection for immune assays.

Analysis of gene expression in CLL cases: previously reported microarray data (NCI Gene Expression Omnibus accession GSE37168) was reanalyzed. Affymetrix CEL files were processed using the affy package in R. The Robust Multichip Analysis (RMA) algorithm was used for background correction which models the observed intensities as a mixture of

exponentially distributed signal and normally distributed noise. This was followed by quantile normalization across arrays to facilitate comparison of gene expression under different conditions. The individual probe-level was finally summarized using the median polish approach to get robust probeset-level values. Gene-level values were obtained by selecting the probe with the maximal average expression for each gene. Batch effects in the data were removed by using the Combat program.

Generation and detection of antigen specific T cells from patient PBMCs:

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Autologous dendritic cells (DCs) were generated from immunomagnetically-isolated CD14⁺ cells (Miltenyi, Auburn CA) that were cultured in RPMI (Cellgro) supplemented with 3% fetal bovine serum, 1% penicillin-streptomycin (Cellgro), 1% L-glutamine and 1% HEPES buffer in the presence of 120 ng/ml GM-CSF and 70 ng/ml IL-4 (R&D Systems, Minneapolis, MN). On days three and five, additional GM-CSF and IL-4 were added. On day six, cells were exposed to 30μg/ml Poly I:C (Sigma Aldrich, St Louis, MO) to undergo maturation (for 48 hours), in addition to adding IL-4 and GM-CSF. CD19⁺ B cells were isolated from patient PBMCs by immunomagnetic selection (CD19⁺ microbeads; Miltenyi, Auburn, CA), and seeded at Ix10⁶ cells/well in a 24-well plate. B cells were cultured in *B cell media* (Iscoves modified Dulbecco medium (IMDM; Life Technologies, Woburn, MA), supplemented with 10% human AB serum (GemCell, Sacramento, CA), 5μg/mL insulin (Sigma Chemical, St Louis, MO), 15 μg/mL gentamicin, IL-4 (2ng/ml, R&D Systems, Minneapolis, MN) and CD40L-Tri (1μg/ml). CD40L-Tri was replenished every 3-4 days. For some experiments, CD40L-Tri activated and expanded CD19⁺ B cells were used as APCs.

Generation of antigen-specific T cells from patient PBMCs: To generate peptide-reactive T cells from CLL patients, immunomagnetically selected CD8⁺ T cells (5x10⁶/well) from pre- and post-transplant PBMCs (CD8+ Microbeads, Miltenyi, Auburn, CA) were cultured with autologous peptide pool-pulsed DCs (at 40:1 ratio) or CD40L-Tri-activated irradiated B cells (at 4:1 ratio) respectively, in complete medium supplemented with 10% FBS and 5-10 ng/mL IL-7, IL-12 and IL-15. APCs were pulsed for 3 hours with peptide pools (10 μM/ peptide/pool). CD8⁺ T cells were re-stimulated weekly (for 1-3 weeks, starting on day 7) with APCs.

Detection of antigen-specific T cells: T cell specificity against peptide pools was tested by IFN- γ ELISPOT assay, 10 days following 2nd and 4th stimulations. IFN- γ release was detected using test and control peptide-pulsed CD40L-activated B cells (50,000 cells/well) co-incubated with 50,000 CD8⁺ T cells/well (Millipore, Billerica, MA) for 24 hours on ELISPOT plate. IFN- γ was detected using capture and detection antibodies, as directed (Mabtech AB, Mariemont, OH), and imaged (ImmunoSpot Series Analyzer; Cellular Technology, Cleveland, OH). To test T cell reactivity dependence on MHC class I, ELISPOT plates were first coated with APCs co-incubated with class I blocking antibody (W6/32) for 2 hours at 37°C, prior to introduction of T cells into the wells. MHC class I tetramer was used to test specificity of T cells where indicated (Emory University, Atlanta GA). For tetramer staining, 5x10⁵ cells were incubated for 60 minutes at 4°C with 1µg/mL PE-labeled tetramer, and then incubated with the addition of anti-CD3-FITC and anti-CD8-APC antibodies (BD Biosciences, San Diego CA) for another 30 minutes at 4°C. A minimum of 100,000 events were acquired per sample. Secretion of GM-CSF and IL-2 from cultured CD8⁺ T cells was detected by analysis of culture supernatants using a Luminex multiplex bead-based technology, per the manufacturer's recommendations (EMD

Millipore, Billerica, MA). In brief, fluorescent-labeled microspheres were coated with specific cytokine capture antibodies. After incubation with the culture supernatant sample, captured cytokines were detected by a biotinylated detection antibody followed by a streptavidin-PE conjugate and median fluorescence intensity (MFI) was measured (Luminex 200 Bead Array instrument; Luminex Corporation, Austin TX). Based on a standard curve, cytokine levels were calculated in the Bead View Software program (Upstate, EMD Millipore, Billerica, MA). For detection and quantitation of TCR V β clonotypes, *mut-FNDC3B* specific T cells were enriched from Pt 2's T cell lines using the IFN- γ secretion assay (Miltenyi, Auburn, CA) according to the manufacturer's instructions and as previously described.

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Statistical considerations: Two-way ANOVA models were constructed for T cells reactivity against mut vs wt peptide in the form of IFN-gamma, GM-CSF, and IL-2 release and included concentration and mutational status as fixed effects along with an interaction term as appropriate. *P*-values for these models were adjusted for multiple comparisons post-hoc using the Tukey method. For normalized comparisons of IFN-gamma, a *t*-test was performed to test the hypothesis that the normalized ratio equaled one. For other comparisons of continuous measures between groups, a Welch *t*-test was used. All *P*-values reported are two-sided and considered significant at the 0.05 level with appropriate adjustment for multiple comparisons. Analysis was performed in SAS v9.2.

Detection and quantitation of TCR Vβ clonotypes: To detect mut-*FNDC3B* specific TCR Vβ, a two-step nested PCR from peptide-specific IFN- γ enriched T cell populations was performed. In short, the dominant Vβ subfamily was identified among the 24 known Vβ subfamilies. First, 5 pools of Vβ forward primers (pool 1: Vβ 1–5.1; pool 2: Vβ 5.2–9; pool 3:

Vβ 10–13.2; pool 4: Vβ 14–19; and pool 5: Vβ 20, 22–25) were generated. RNA extracted from the T cell clones (QIAamp RNA Blood Mini-kit; Qiagen, Valencia, CA), was reverse transcribed into cDNA (Superscript, GIBCO BRL, Gaithersburg, MD) using random hexamers, and PCR-amplified in five separate 20 μl volume reactions. Second, T cell clone-derived cDNA was reamplified, with each of the 5 individual primers contained within a positive pool together with a FAM-conjugated Cβ reverse (internal) primer. Subsequently, 4 μl of this PCR product was amplified with 1 μl of the clone CDR3 region-specific primer and probe, and 10 μl of Taqman Fast Universal PCR Master Mix (Applied Biosystems, Foster City, CA) in a total volume of 20 μl. The PCR amplification conditions were: 95°C for 20 minutes × 1 cycle, and 40 cycles of 95°C for 3 seconds followed by 60°C for 30 seconds (7500 Fast Real-time PCR cycler; Applied Biosystems, Foster City, CA). Test transcripts were quantified relative to *S18* ribosomal RNA transcripts by calculating 2^(S18 rRNA CT-target CT) as described previously.

Detection of molecular tumor burden: The clonotypic IgH sequence of Pt 2 was identified using a panel of VH-specific PCR primers, as previously described. Based on this sequence, a quantitative Taqman PCR assay was designed such that a sequence-specific probe was located in the region of junctional diversity (Applied Biosystems; Foster City, CA). This Taqman assay was applied to cDNA from tumor. All PCR reactions consisted of: 50°C for 1 minute x1 cycle, 95°C for 10 minutes x1 cycle, and 40 cycles of 95°C for 15 seconds followed by 60°C for 1 minute. All reactions were performed using a 7500 Fast Real-time PCR cycler (Applied Biosystems, Foster City, CA). Test transcripts were quantified relative to GAPDH.

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Other Embodiments

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From the foregoing description, it will be apparent that variations and modifications may be made to the invention described herein to adopt it to various usages and conditions. Such embodiments are also within the scope of the following claims.

The recitation of a listing of elements in any definition of a variable herein includes definitions of that variable as any single element or combination (or sub-combination) of listed elements. The recitation of an embodiment herein includes that embodiment as any single embodiment or in combination with any other embodiments or portions thereof.

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Incorporation by Reference

All patents and publications mentioned in this specification are herein incorporated by reference to the same extent as if each independent patent and publication was specifically and individually indicated to be incorporated by reference.

The Claims Defining the Invention are as Follows:

1. A method of making a personalized neoplasia vaccine for a subject diagnosed as having a neoplasia, comprising:

identifying a plurality of sequences comprising mutations in the neoplasia;

analyzing the plurality of sequences comprising mutations to identify a subset of at least five neoantigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising

identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and

ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising:

- (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation;
- (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\geq 1000 \text{ nM}$;
- (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\leq 150 \text{ nM}$;
- (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation;
- (v) a polypeptide that binds to the HLA of the subject with a Kd of $150 \le 500$ nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $150 - \le 500$ nM; and

producing, based on the identified subset, a personalized neoplasia vaccine.

2. A personalized neoplasia vaccine when used in a method for treating a subject diagnosed as having a neoplasia, said method comprising:

identifying a plurality of sequences comprising mutations in the neoplasia;

analyzing the plurality of sequences comprising mutations to identify a subset of at least five neoantigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic

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mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising

identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and

ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising:

- (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation;
- (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\geq 1000 \text{ nM}$;
- (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\leq 150 \text{ nM}$;
- (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation;
- (v) a polypeptide that binds to the HLA of the subject with a Kd of $150 \le 500$ nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of 150- \leq 500 nM;

producing, based on the identified subset, a personalized neoplasia vaccine; and administering the personalized neoplasia vaccine to the subject, thereby treating the neoplasia.

- 3. The method of claim 1 or the personalized neoplasia vaccine of claim 2, wherein identifying a plurality of sequences comprising mutations in the neoplasia comprises:
- 25 sequencing the genome, transcriptome, or proteome of the neoplasia.
 - 4. The method of claim 1 or claim 3, or the personalized neoplasia vaccine of claim 2 or claim 3, wherein the ranking comprises:

determining one or more characteristics associated with the neo-antigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the characteristics selected from the group consisting of molecular weight, cysteine content, hydrophilicity, hydrophobicity, charge, and binding affinity; and

basing the said ranking, on the determined characteristics.

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- 5. The method of any one of claims 1, 3 and 4, or the personalized neoplasia vaccine of any one of claims 2 to 4, wherein the personalized neoplasia vaccine comprises at least about 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least about 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations.
- 6. The method of any one of claims 1, 3 and 4, or the personalized neoplasia vaccine of any one of claims 2 to 4, wherein the personalized neoplasia vaccine comprises at least about 20 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least about 20 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 20 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations.
- 7. The method of any one of claims 1, 3 and 4, or the personalized neoplasia vaccine of any one of claims 2 to 4, wherein the personalized neoplasia vaccine comprises at least about 20 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, and wherein:

the at least 20 neo-antigenic peptides range from about 5 to about 50 amino acids in length, or the at least 20 neo-antigenic peptides range from about 15 to about 35 amino acids in length, or the at least 20 neo-antigenic peptides range from about 18 to about 30 amino acids in length, or the at least about 20 neo-antigenic peptides range from about 6 to about 15 amino acids in length, or the at least about 20 neoantigenic peptides are 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 amino acids in length.

- 8. The method of claim 1, wherein the personalized neoplasia vaccine further comprises an adjuvant.
- 9. The method of claim 8, wherein the adjuvant is selected from the group consisting of poly-ICLC, 1018 ISS, aluminum salts, Amplivax®, AS15, BCG, CP-870,893, CpG7909, CyaA, dSLIM®, GM-CSF, IC30, IC31, Imiquimod, ImuFact IMP321, IS Patch, ISS, ISCOMATRIX®, Juvlmmune, LipoVac, MF59®, monophosphoryl lipid A, Montanide® IMS 1312, Montanide® ISA 206, Montanide® ISA 50V, Montanide® ISA-51, OK-432, OM-174, OM-197-MP-EC, ONTAK®, PepTel RTM, vector system, PLGA microparticles, resiguimod, SRL172, Virosomes and other Virus-like particles, YF-17D, VEGF trap, R848, beta-glucan, Pam3Cys, Aquila's QS21 stimulon®, vadimezan, and AsA404 (DMXAA).

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- 10. The method of claim 9, wherein the adjuvant is poly-ICLC.
- 11. A method of treating a subject diagnosed as having a neoplasia with a personalized neoplasia vaccine, comprising:

identifying a plurality of sequences comprising mutations in the neoplasia;

analyzing the plurality of sequences comprising mutations to identify a subset of at least five neoantigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising

identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and

ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising:

- (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation;
- (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\ge 1000 \text{ nM}$;
- (iii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM;
- (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation,
- (v) a polypeptide that binds to the HLA of the subject with a Kd of $150 \le 500$ nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $150 - \le 500 \text{ nM}$;

producing, based on the identified subset, a personalized neoplasia vaccine; and administering the personalized neoplasia vaccine to the subject, thereby treating the neoplasia.

- 30 12. The method of claim 11, wherein identifying a plurality of sequences comprising mutations in the neoplasia comprises: sequencing the genome, transcriptome, or proteome of the neoplasia.
 - 13. The method of claim 11 or claim 12, wherein the ranking comprises:

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determining one or more characteristics associated with the neo-antigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the characteristics selected from the group consisting of molecular weight, cysteine content, hydrophilicity, hydrophobicity charge, and binding affinity; and

basing the said ranking, on the determined characteristics.

- 14. The method of any one of claims 11 to 13, wherein the personalized neoplasia vaccine comprises at least about 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least about 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations, or the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 10 neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations.
- 15. The method of any one of claims 11 to 13, wherein the personalized neoplasia vaccine comprises at least 20 neo-antigenic peptides corresponding to the neo-antigenic mutations.
 - 16. The method of any one of claims 11 to 13, wherein the personalized neoplasia vaccine comprises one or more DNA molecules capable of expressing at least 20 neo-antigenic peptides corresponding to the neo-antigenic mutations.
 - 17. The method of any one of claims 11 to 13, wherein the personalized neoplasia vaccine comprises one or more RNA molecules capable of expressing at least 20 neo-antigenic peptides corresponding to the neo-antigenic mutations.
- 25 18. The method of claim 15, wherein the at least 20 neo-antigenic peptides range from about 5 to about 50 amino acids in length.
 - 19. The method of claim 15, wherein the at least 20 neo-antigenic peptides range from about 15 to about 35 amino acids in length.
 - 20. The method of claim 15, wherein the at least 20 neo-antigenic peptides range from about 18 to about 30 amino acids in length.

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- 21. The method of claim 15, wherein the at least 20 neo-antigenic peptides range from about 6 to about 15 amino acids in length.
- 22. The method of claim 15, wherein the at least 20 neo-antigenic peptides are 15, 16, 17, 18, 19, 20, 21, 22, 23, 24, or 25 amino acids in length.
- 23. The personalized neoplasia vaccine of any one of claims 2 to 7, wherein administering further comprises:

dividing the produced vaccine into two or more sub-pools; and injecting each of the sub-pools into a different location of the patient.

- 24. The personalized neoplasia vaccine of claim 23, wherein each of the sub-pools injected into a different location comprises neo-antigenic peptides such that a number of individual peptides in the subpool targeting any single patient HLA is one, or as few above one as possible.
- 25. The personalized neoplasia vaccine of any one of claims 2 to 4, wherein the personalized neoplasia vaccine comprises at least 20 neo-antigenic peptides encoded by the sequences comprising the neoantigenic mutations, wherein the at least 20 neo-antigenic peptides range from about 18 to about 30 amino acids in length, wherein administering further comprises dividing the produced vaccine into two or more sub-pools, wherein each sub-pool comprises at least five neo-antigenic peptides selected to optimize intrapool interactions.
- 26. The personalized neoplasia vaccine of claim 25, wherein optimizing comprises reducing negative interaction among the neo-antigenic peptides in the same pool.
- 27. The personalized neoplasia vaccine of any one of claims 2 to 4, wherein administering further comprises delivering a dendritic cell (DC) vaccine, wherein the DC is loaded with one or more of the at least five neo-antigenic sequences comprising mutations predicted to encode expressed neo-antigenic peptides.
- 28. A personalized neoplasia vaccine prepared according to the method of any one of claims 1 and 3 to 7.
- 29. Use of a personalized neoplasia vaccine in manufacture of a medicament for treating a subject diagnosed as having a neoplasia, wherein the personalized neoplasia vaccine is produced by the steps of:

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analyzing the plurality of sequences comprising mutations to identify a subset of at least five neoantigenic sequences comprising mutations predicted to encode neo-antigenic peptides, the neo-antigenic mutations selected from the group consisting of missense mutations, neoORF mutations, and any combination thereof; said analyzing comprising

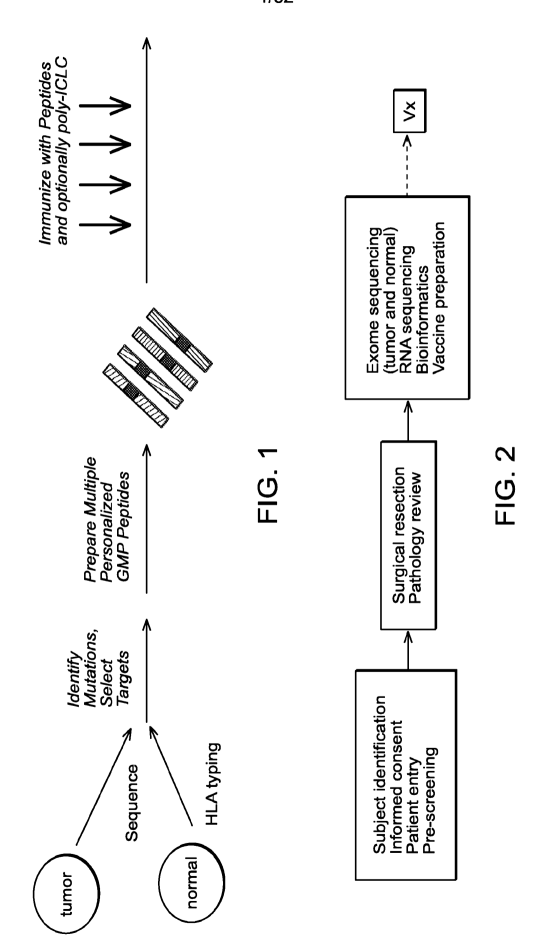
identifying sequences comprising neo-antigenic mutations from the plurality of sequences comprising mutations identified in the neoplasia and

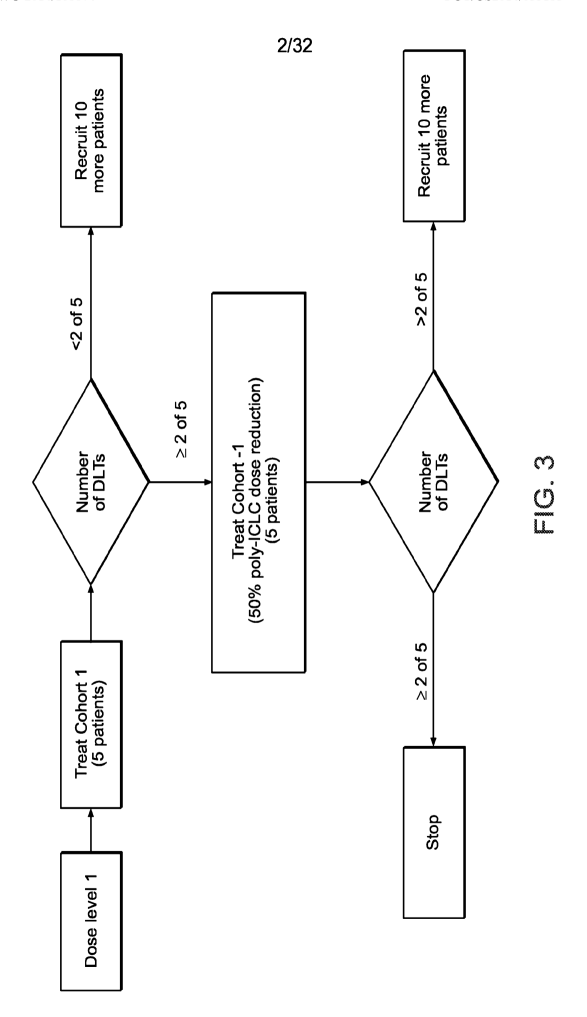
ranking the neo-antigenic peptides encoded by the sequences comprising the neo-antigenic mutations in an order of decreasing priority, the order comprising:

- (i) a neoORF polypeptide that binds to the HLA of the subject with a Kd of ≤ 500 nM encoded by a sequence comprising a neoORF mutation;
- (ii) a polypeptide that binds to the HLA of the subject with a Kd of ≤ 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $\ge 1000 \text{ nM}$;
- (iii) a polypeptide that binds to the HLA of the subject with a Kd of \leq 150 nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of ≤ 150 nM;
- (iv) a neoORF polypeptide that binds to the HLA of the subject with a Kd of > 500 nM encoded by a sequence comprising a neoORF mutation,
- (v) a polypeptide that binds to the HLA of the subject with a Kd of $150 \le 500$ nM encoded by a sequence comprising a missense mutation, wherein the native cognate protein has a Kd of $150 - \le 500 \text{ nM}$.

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Out-of-Frame InDels and Gene Fusions ...LTYSGRKTA...

Splice SiteESVANGHPVLT...

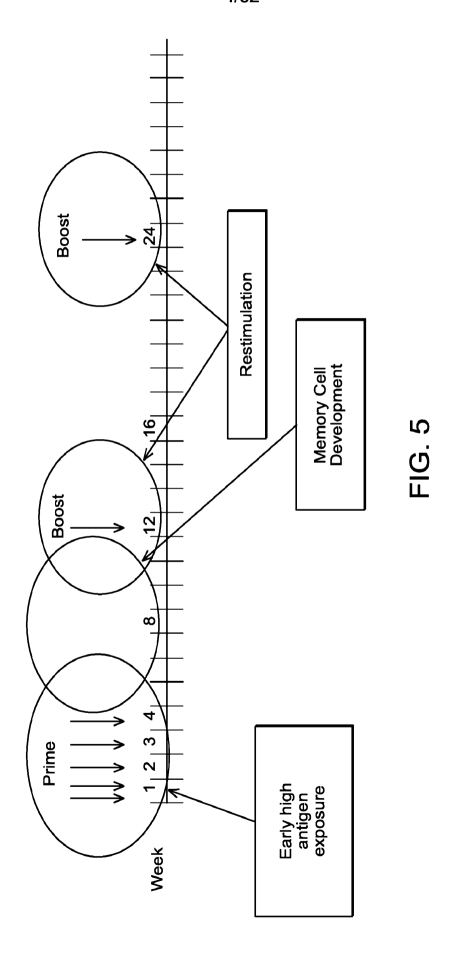
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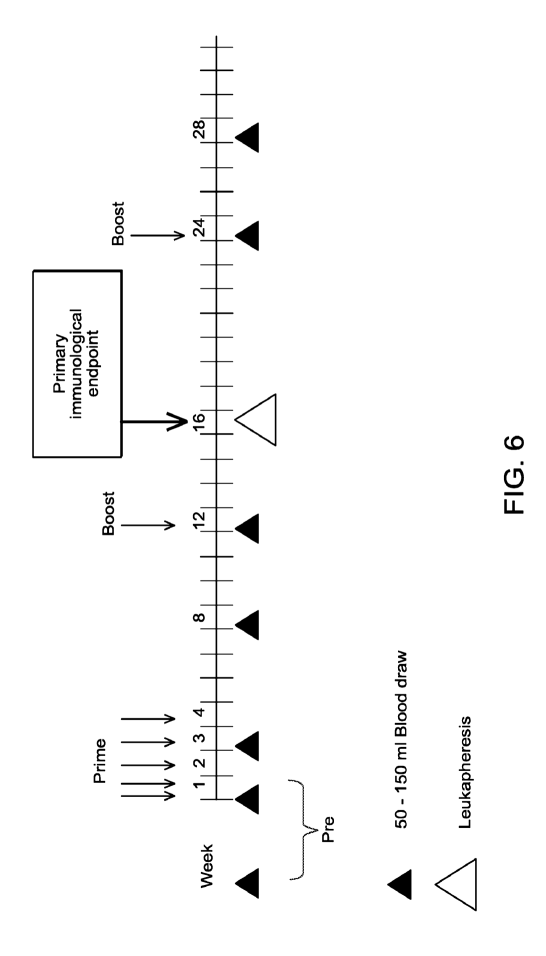
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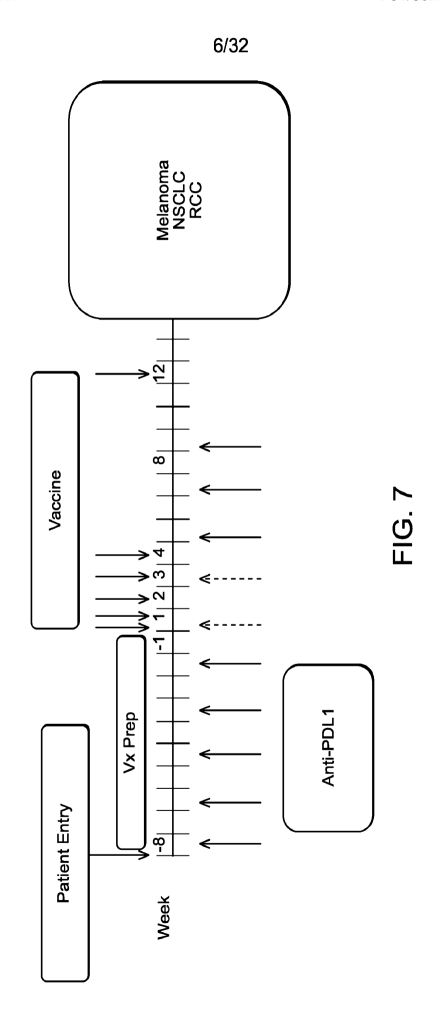
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FIG. 4B

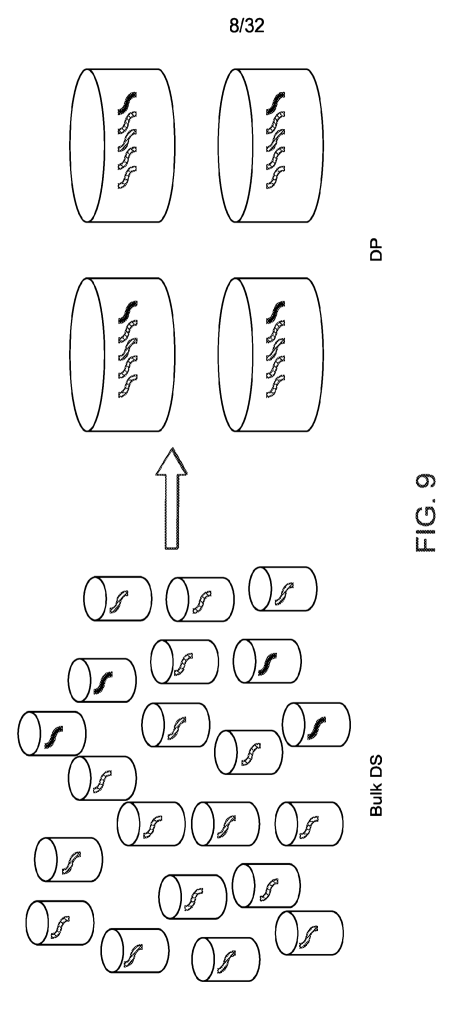




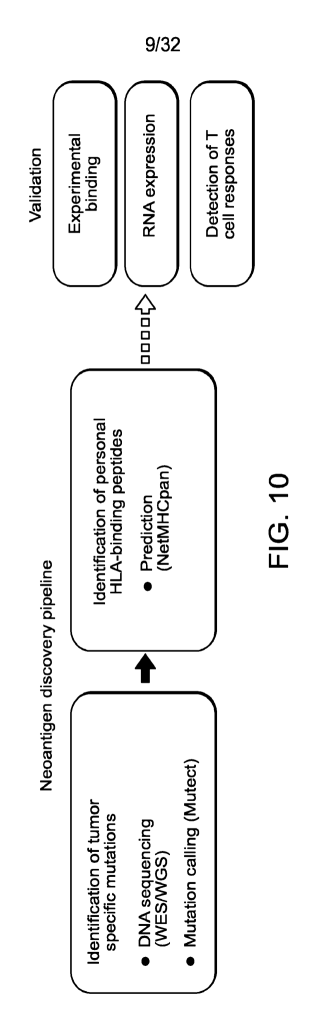


CATEGORY	NUTATON	Mutant peptide Ka	Native peptide Kd	rnalevel
	NeoORE	SOO nM	A A	~/]/M/H
N	Missense	< 150 nM	$\geq 1000 \mathrm{nM}$	H/M/L
r")	Missense	< 150 nM	< 150 nM	H/M/L
4	NeoORF	>500 nM	NA	H/M/L
\$ 4 .3	Missense	150 - < 500 nM	150 - < 500 nM	H/M/L

o D



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Missense mutation

DNA WT: CTTGATCGGATCGTAGCTACG
Mut: CTTGAACGGATCGTAGCTACG

a.a. WT: L**D**RIVAT
Mut: L<u>E</u>RIVAT

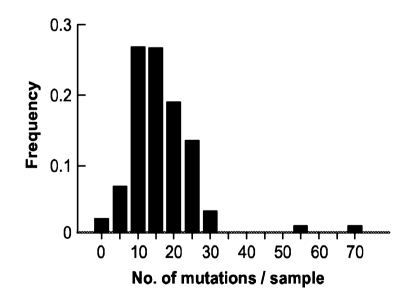
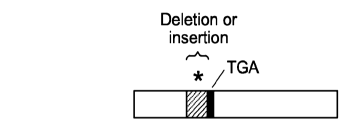


FIG. 11A

Frameshift mutations





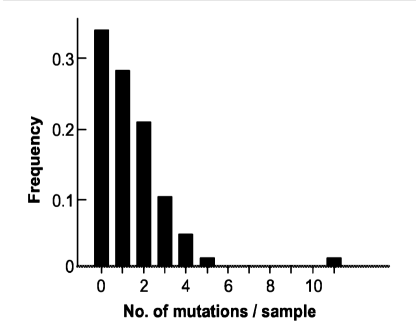
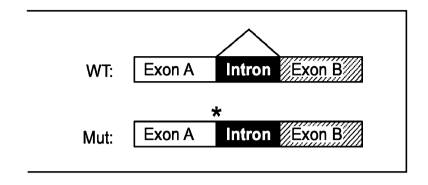


FIG. 11B

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Splice site mutations



WT: ESVAN GHPULT

Mut: ESVAN GFTLSNQR

Neo ORF

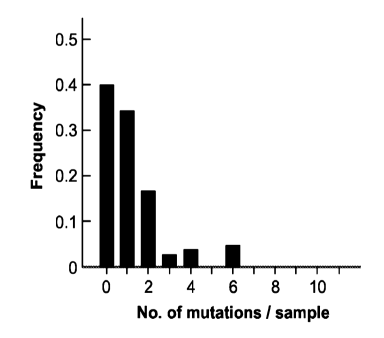


FIG. 11C

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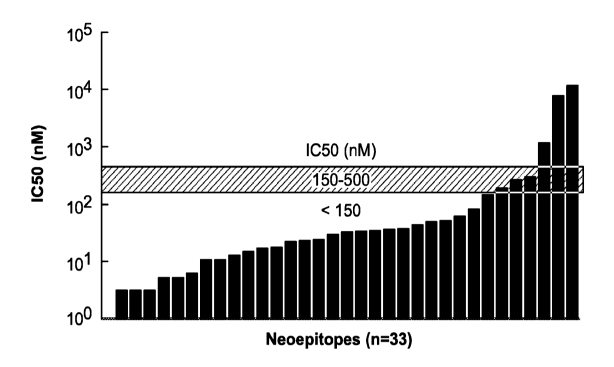


FIG. 12A

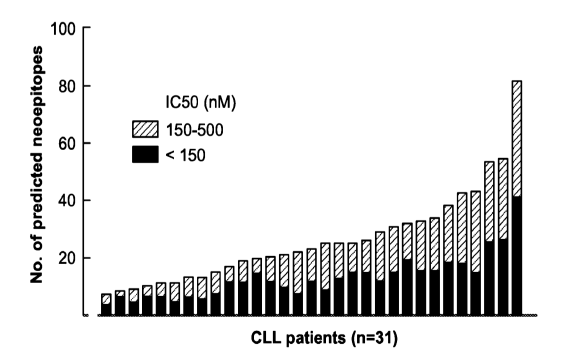


FIG. 12B

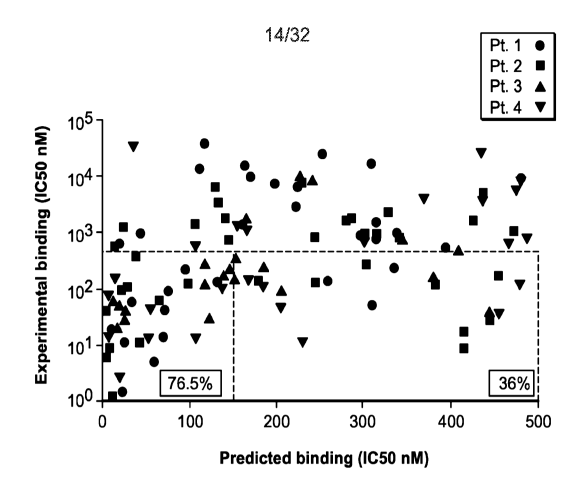
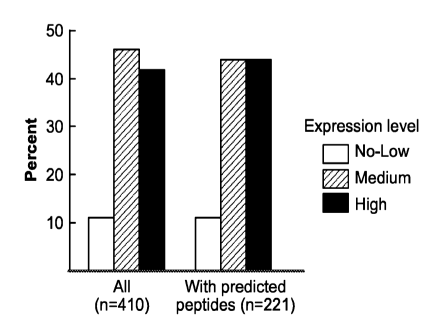


FIG. 12C



Genes with missense mutations from 28 CLLs

FIG. 12D

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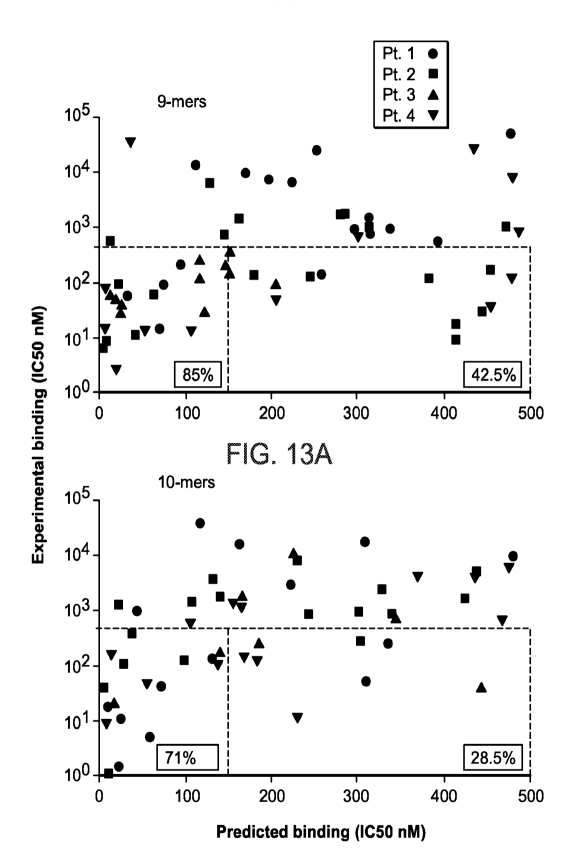
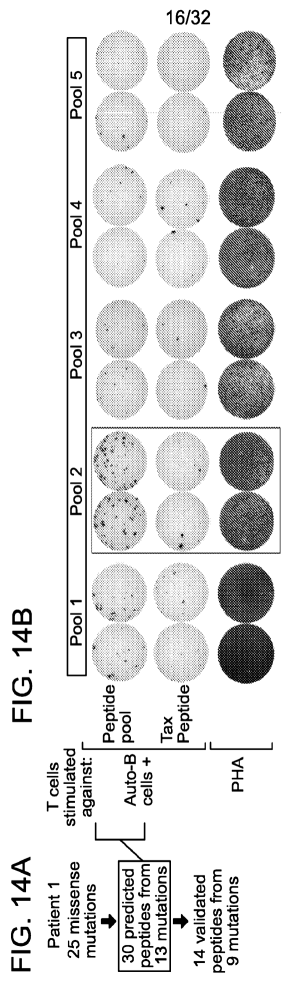
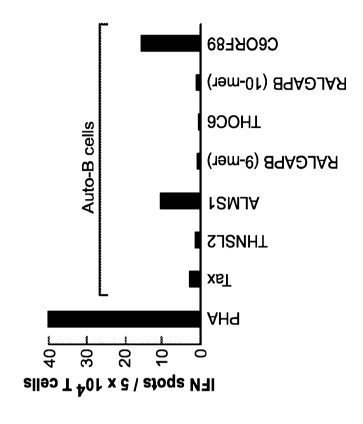
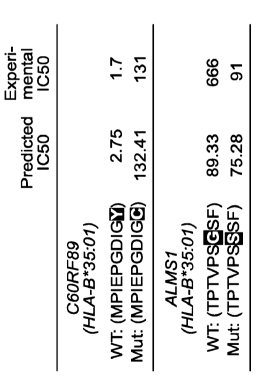


FIG. 13B

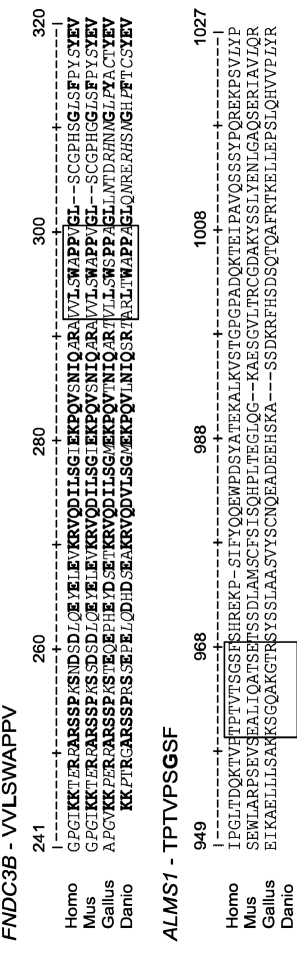


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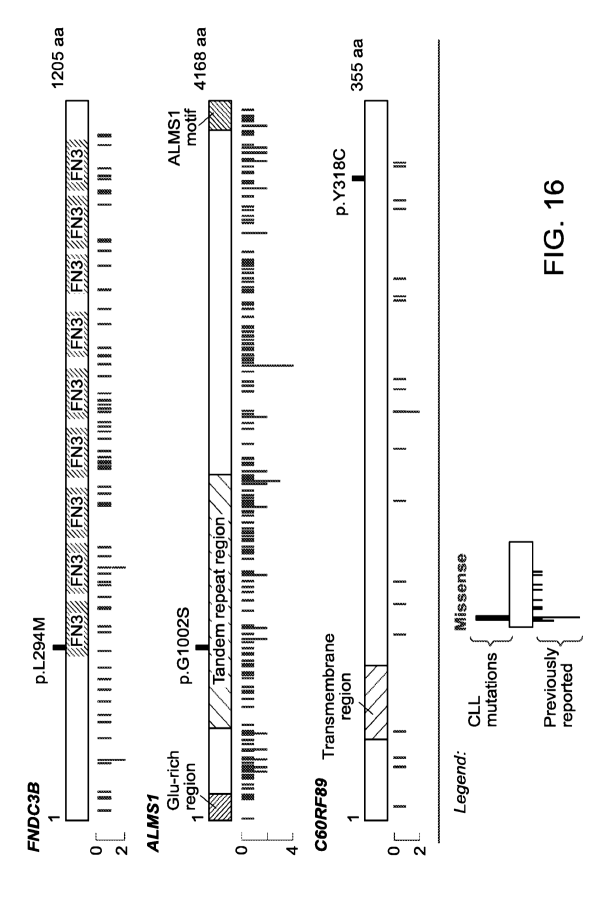
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LPFPK--D*SS***ln**K¢fliqpe**p**vvgSkmhkvhd*lf*tl*gS*¢eamlqli**p**Pfq**cr**t**hc**qsvampi**e**S**gd**ig**y**adaah**w**kvyiv LS*FPK*--TV*S*INNCFLIR-HPDLGNKSYSLHS*LFVVGS*GHLTLTVAPLDKCRGHCEM*FK*VD*L*EAGDLGYASMDY**W**MMSFV CPLLEIWSSTLQRCRLSSRRPQ--PSRVQVLGWMVVADGSPDVRLLPVQRCRKHCRSFSLR*LEPG*DMVFADSQIWLMELS *LPFPK*--DA*S*INKCSFLHPE**P**VVGSKMHKMPD*LFIIGSG*EAMLQLI**P**PFQCRRHCQSVAMPIE*PG*DIGYVDTTHWKVYVI 300 280 260 241 Gallus Homo Danio Mus

C6ORF89 - MPIEPGDIGY

TG. 15



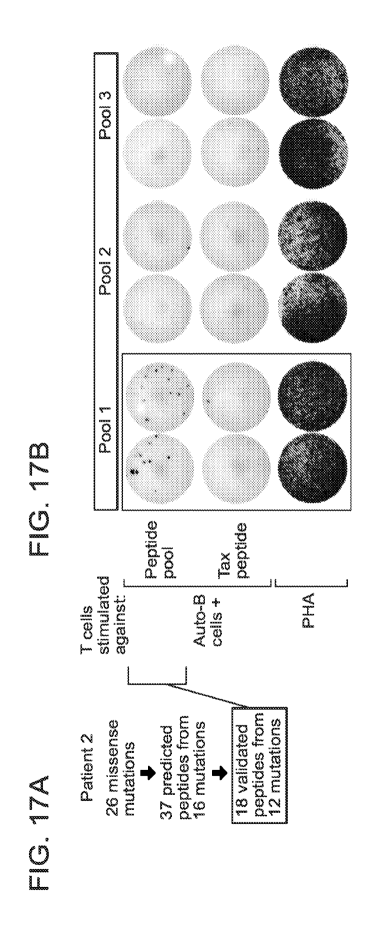
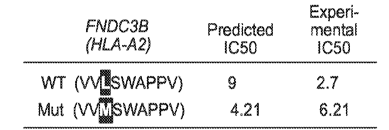
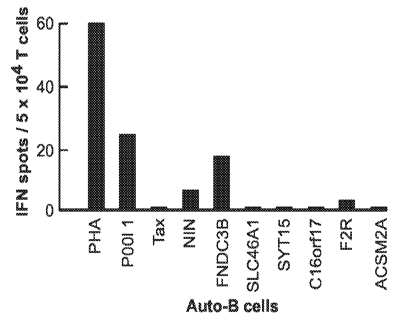
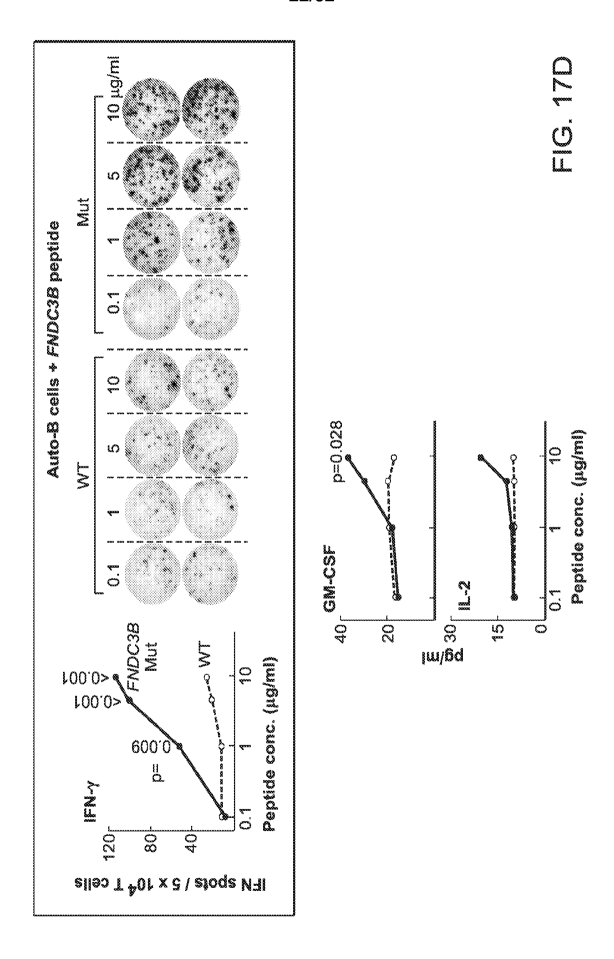


FIG. 17C







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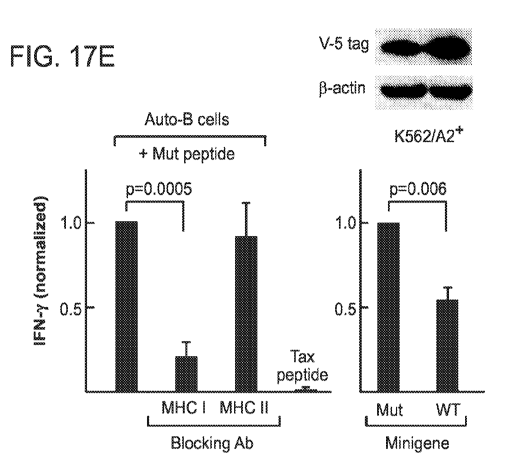


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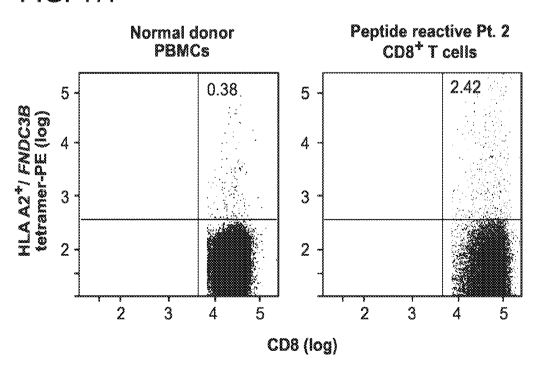
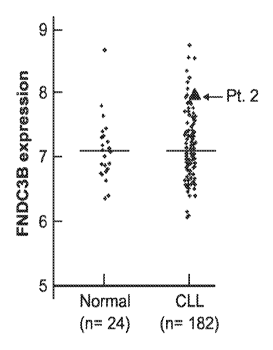
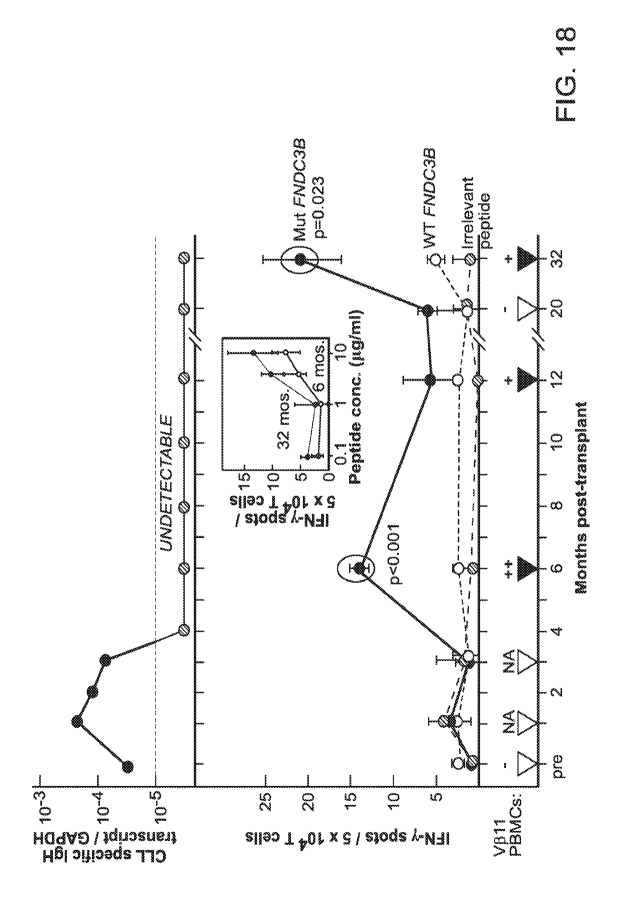


FIG. 17G





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FIG. 19A

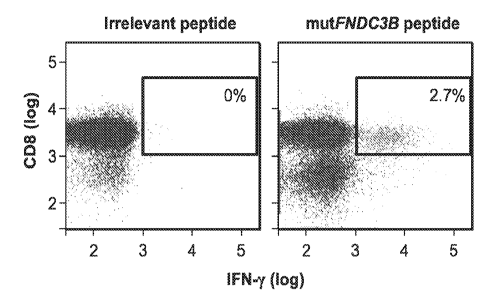


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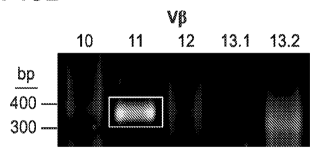


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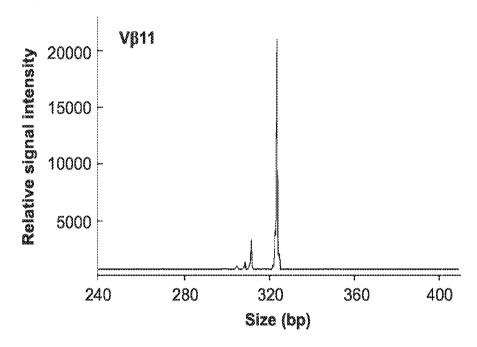
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FIG. 19D



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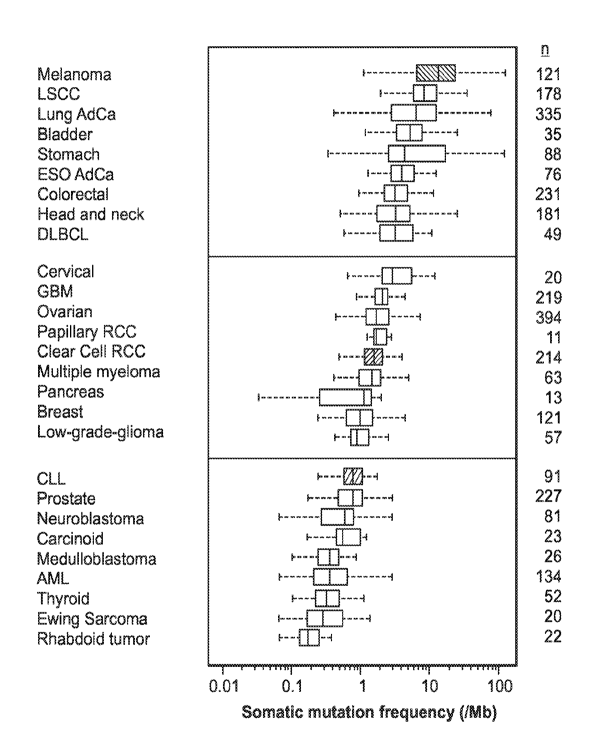
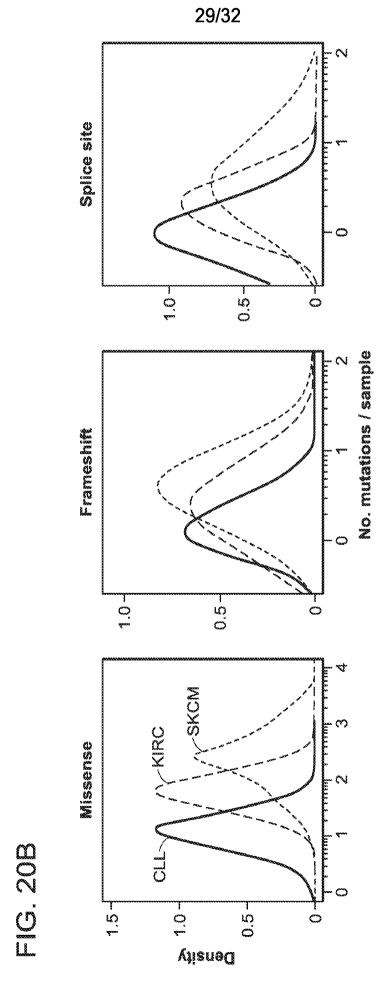


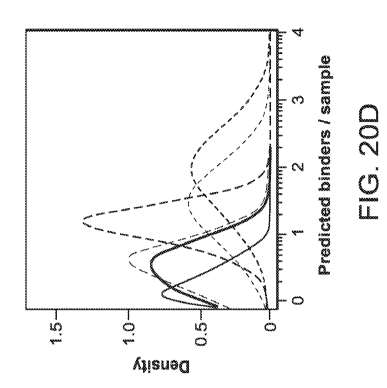
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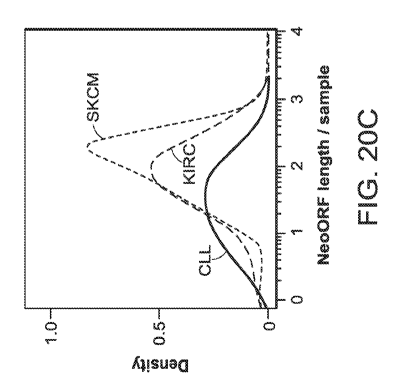


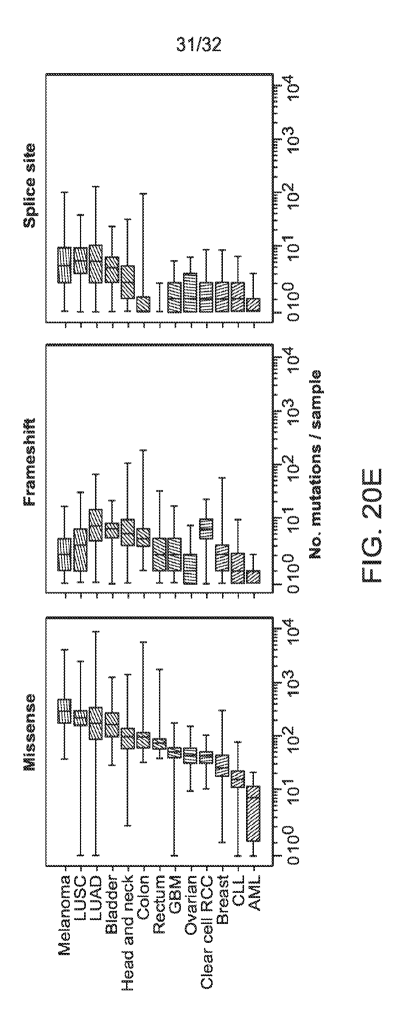
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PCT/US2014/033185

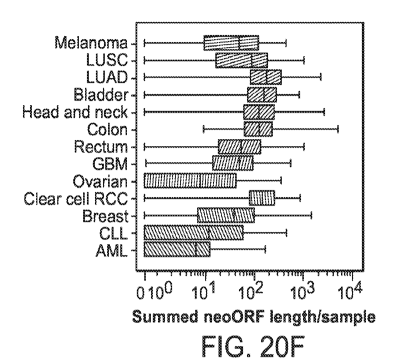






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IC50 <150 nM IC50 <500 nM Melanoma-LUSC-LUAD-Bladder-Head and neck-Colon-Rectum-GBM-Ovarian-Clear cell RCC-Breast-CLL **AML** 103 104 0100 103 102 102 101 101 104 0100 No. predicted binders / sample

FIG. 20G

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Asn Gln Glu Ala Asp Glu Glu His Ser Lys Ala Ser Ser Asp Lys Arg 35 40 45

Phe His Ser Asp Ser Gln Thr Gln Ala Phe Arg Thr Lys Glu Leu Leu 50 60

Glu Pro Ser Leu Gln His Val Val Pro Leu Tyr Arg
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<213> Homo sapiens

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Leu Pro Phe Pro Lys Asp Ala Ser Leu Asn Lys Cys Ser Phe Leu His 10 15

Pro Glu Pro Val Val Gly Ser Lys Met His Lys Met Pro Asp Leu Phe 20 25 30

Ile Ile Gly Ser Gly Glu Ala Met Leu Gln Leu Ile Pro Pro Phe Gln 35 40 45

Cys Arg Arg His Cys Gln Ser Val Ala Met Pro Ile Glu Pro Gly Asp 50 60

Ile Gly Tyr Val Asp Thr Thr His Trp Lys Val Tyr Val Ile 70 75

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Pro Glu Pro Val Val Gly Ser Lys Met His Lys Val His Asp Leu Phe 20 25 30

Thr Leu Gly Ser Gly Glu Ala Met Leu Gln Leu Ile Pro Pro Phe Gln 35 40

Cys Arg Thr His Cys Gln Ser Val Ala Met Pro Ile Glu Ser Gly Asp 50 60

Ile Gly Tyr Ala Asp Ala Ala His Trp Lys Val Tyr Ile Val

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His Pro Asp Leu Gly Asn Lys Ser Tyr Ser Leu His Ser Leu Phe Val 20 25 30

Val Gly Ser Gly His Leu Thr Leu Thr Val Ala Pro Leu Asp Lys Cys 35 40 45

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Gly Tyr Ala Ser Met Asp Tyr Trp Met Met Ser Phe Val 65 70 75

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Cys Pro Leu Leu Glu Ile Trp Ser Ser Thr Leu Gln Arg Cys Arg Leu

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Val Val Ala Asp Gly Ser Pro Asp Val Arg Leu Leu Pro Val Gln Arg 35 40 45

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