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DESCRIPTION

[0001] This application claims the benefit of United States Application No. 61/259,403, filed November 9, 2009.

FIELD OF THE INVENTION

[0002] The present invention relates to oral pharmaceutical formulations of neratinib provided in the form of coated tablets prepared by fluid bed granulation or by wet granulation, and improved methods of making these coated tablets.

BACKGROUND OF THE INVENTION

[0003] Protein kinases are important in the transmission of biochemical signals, which initiate cell replication. Protein kinases are enzymes that catalyze the transfer of a phosphate group from ATP to an amino acid residue, such as tyrosine, serine, threonine, or histidine on a protein. Regulation of these protein kinases is essential for the control of a wide variety of cellular events including proliferation and migration. Specific protein kinases have been implicated in adverse conditions including cancer [Traxler, P. M., Exp. Opin. Ther. Patents, 8, 1599 (1998); Bridges, A. J., Emerging Drugs, 3, 279 (1998)], restenosis [Mattsson, E., Trends Cardiovas. Med. 5, 200 (1995); Shaw, Trends Pharmacol. Sci. 16, 401 (1995)], atherosclerosis [Raines, E. W., Bioessays, 18, 271 (1996)], angiogenesis [Shawver, L. K., Drug Discovery Today, 2, 50 (1997); Folkman, J., Nature Medicine, 1, 27 (1995)] and osteoporosis [Boyce, J. Clin. Invest., 90,1622 (1992)]. Compounds capable of inhibiting the activity of receptor tyrosine kinases are knwn to be useful in the treatment of cancers, including but not limited to for example, non-small cell lung cancer (NSCLC), breast cancer, polycystic kidney disease, colonic polyps, and stroke in mammals. A specific kinase inhibitor is (E)-N-(4-(3-chloro-4-(pyridin-2-ylmethoxy)phenylamino)-3-cyano-7-ethoxyquinolin-6-yl)-4-(dimethylamino)but-2enamide, also known as neratinib. Nerartinib is a weak base having low bioavailability and low solubility in both water and alcohol.

[0004] Neratinib maleate particles exhibit very high surface free energy, (work of cohesion = 45.62 mN / m). This property renders the primary particles very cohesive and prone to aggregation as described by B. Janczuk and T.Bialopiotrowicz, "Surface Free-Energy Components of Liquids and Low Energy Solids and Contact Angles," in J. Colloid Interf. Sci. 127 (1989), p.189-204; W.R. Good, "A Comparison of Contact Angle Interpretations," in J. Colloid Interf. Sci. 44 (1973), p. 63; M.D. Lechner (Ed.), Landolt Börnstein, New Series, Vol. IV/16, "Surface Tension of Pure Liquids and Binary Liquid Mixture," Springer Verlag, 1998.; and J.J. Jasper, "The Surface Tension of Pure Liquid Compounds," in J. Phys. Chem. Ref. Data, Vol.1, No. 4, 1972, p. 859. As a consequence of cohesiveness, neratinib maleate powder does

not lend itself easily to pharmaceutical operations such as mixing, flow or fluidization especially when it constitutes a high proportion in a composition. Due to these limitations, it was not possible to develop a neratinib maleate formulation comprising a capsule or tablet of higher strength employing a direct compression or roller compaction processes successfully. A formulation using a conventional wet granulation method led to chemical degradation and stability issues.

SUMMARY OF THE INVENTION

[0005] It is desirable to provide a neratinib maleate formulation, where the surface property of the active ingredient is modified by spraying or otherwise applying a substance, such as a polymer like povidone, of low surface energy (for instance about 38 mN/m) on the surface of neratinib maleate particles.

[0006] The present invention provides pharmaceutically acceptable solid compositions suitable for oral administration comprising the active ingredient neratinib maleate as claimed. In certain embodiments, such solid compositions are provided in the form of coated tablets prepared by fluid bed granulation. In some embodiments, the present invention provides a unit dosage form comprising neratinib maleate.

[0007] Disclosed is a pharmaceutically acceptable composition comprising: a granulation comprising intragranular components: (a) 10-70 weight percent neratinib maleate; (b) 15-65 weight percent of one or more fillers; (c) 0-8 or 0.5-8 weight percent of one or more disintegrants; and (d) 0.2-8 weight percent, in certain embodiments 0.2-6 weight percent, of one or more glidants; and (e) 5-15 weight percent of one or more surface modifying agents. The granulation is combined with extragranular components (f) 1-25 or 4-25 weight percent of one or more fillers; (g) 1-8 or 0-8 weight percent of one or more disintegrants and (h) 0.1-3 or 0.5-3 weight percent of one or more lubricants, and then compressed into tablets or dry-filled into capsules.

[0008] The present invention provides a tablet comprising a pharmaceutically acceptable composition comprising: a granulation comprising intragranular components (a) 35 or 41 weight percent of neratinib maleate; (b) 15-65 weight percent of mannitol and microcrystalline cellulose; (c) 0.5-8 weight percent of crospovidone or croscarmellose sodium; and (d) 0.2-8 weight percent, in certain embodiments 0.2-6 weight percent, of colloidal silicon dioxide, and (e) 5-15 weight percent of povidone. The granulation is combined with extragranular components (f) 1-25 weight percent of microcrystalline cellulose; (g) 1-8 weight percent of crospovidone or croscarmellose sodium and (h) 0.5-3 weight percent of magnesium stearate, and then compressed into tablets or dry-filled into capsules.

[0009] The present invention also provides methods of preparing stable, pharmaceutically acceptable neratinib-maleate formulations for oral administration comprising components described above and herein, which allow for improved processing characteristics while

maintaining acceptable pharmokinetic properties.

BRIEF DESCRIPTION OF THE DRAWINGS

[0010] Figure 1 summarizes mean pharmacokinetic parameters versus time for neratinib maleate following the administration of immediate release tablet formulations of neratinib with different dissolution rates. TR refers to a rapid dissolving tablet, while SR dissolve relatively slowly. The presented data shows plasma concentration levels following administration of a single oral dose (240-mg tablet) in subjects.

DETAILED DESCRIPTION OF CERTAIN EMBODIMENTS OF THE INVENTION

1. Definitions:

[0011] As used herein, an "effective amount" of a compound or pharmaceutically acceptable composition can achieve a desired therapeutic and/or prophylactic effect. In some embodiments, an "effective amount" is at least a minimal amount of a compound, or composition containing a compound, which is sufficient for treating one or more symptoms of a disorder or condition associated with modulation of protein tyrosine kinases. In certain embodiments, an "effective amount" of a compound, or composition containing a compound, is sufficient for treating symptoms associated with, a disease associated with an aberrant tyrosine kinase receptor (e.g. cancer, including malignant and benign tumor growths).

[0012] The term "subject", as used herein, means a mammal and includes human and animal subjects, such as domestic animals (e.g., horses, dogs, cats, etc.).

[0013] The terms "suffer" or "suffering" as used herein refers to one or more conditions that a patient has been diagnosed with, or is suspected to have.

[0014] The terms "treat" or "treating," as used herein, refers to partially or completely alleviating, inhibiting, delaying onset of, preventing, ameliorating and/or relieving a disorder or condition, or one or more symptoms of the disorder or condition.

[0015] "Therapeutically active agent" or "active agent" refers to a substance, including a biologically active substance, that is useful for therapy (e.g., human therapy, veterinary therapy), including prophylactic and therapeutic treatment. Therapeutically active agents include organic molecules that are drug compounds, peptides, proteins, carbohydrates, monosaccharides, oligosaccharides, polysaccharides, nucleoprotein, mucoprotein, lipoprotein, synthetic polypeptide or protein, small molecules linked to a protein, glycoprotein, steroid, nucleic acid, DNA, RNA, nucleotide, nucleoside, oligonucleotides, antisense oligonucleotides,

lipid, hormone, and vitamin. Therapeutically active agents include any substance used as a medicine for treatment, prevention, delay, reduction or amelioration of a disease, condition, or disorder. Among therapeutically active agents useful in formulations are opioid receptor antagonist compounds, opioid analgesic compounds, and the like. Further detailed description of compounds useful as therapeutically active agents is provided below. A therapeutically active agent includes a compound that increases the effect or effectiveness of a second compound, for example, by enhancing potency or reducing adverse effects of a second compound.

[0016] "Unit dosage form" as used herein refers to a physically discrete unit of inventive formulation appropriate for the subject to be treated. It will be understood, however, that the total daily usage of the compositions of the present invention will be decided by the attending physician within the scope of sound medical judgment. The specific effective dose level for any particular subject or organism will depend upon a variety of factors including the disorder being treated and the severity of the disorder; activity of specific active agent employed; specific composition employed; age, body weight, general health, sex and diet of the subject; time of administration, and rate of excretion of the specific active agent employed; duration of the treatment; drugs and/or additional therapies used in combination or coincidental with specific compound(s) employed, and like factors well known in the medical arts.

[0017] In dry granulation (slugging or roller compaction) intragranular materials are blended to prepare slugs or roller compaction. Material is milled and blended with extragranular materials followed by capsule filling or tablet 20 compression. Wet granulation entails blending intragranular materials. Wet granulate the blend with water, with or without a binder, (using high sheer, low sheer granulators) and dry (using temperatures up to 100° C). Material is milled and blended with extragranular materials followed by capsule filling or tablet compression. See, 25 Handbook of Pharmaceutical Granulation Technology, 1997, Dilip Parikh, Marcel Dekker, Inc. ISBN 0-8247-9882-1, pages 338-368.

2. Pharmaceutically Acceptable Compositions and Formulations:

[0018] In certain embodiments, the present invention provides a pharmaceutically acceptable composition for intravenous administration comprising: neratinib maleate. Neratinib and other 4-amino-3-cyanoquinoline compounds are disclosed in U.S. Pat. Nos. 6,002,008, 6,288,082, 6,297,258, 6,384,051 and 7,399,865. Neratinib has the following chemical structure:

and is isolated as a free base or prepared as a pharmaceutically acceptable salt, such as a

maleate salt. Neratinib is a weak base with an intrinsic low solubility in water.

[0019] In certain embodiments, solid pharmaceutically acceptable compositions of neratinib maleate are provided in the form of tablets prepared by fluid bed granulation. Intragranular particulate components comprising an active ingredient, namely neratinib maleate, one or more fillers, a disintegrant and a glidant are sprayed with, or otherwise wholly or partially covered with, a surface modifying agent, such as povidone, to lower the surface energy of the particles. The fluid bed process is employed to effectively modify the surface behavior of the particulate active ingredient, so that any water instantly dries and does not bring about any polymorphic or chemical change in active during the process. The surface property of the active ingredient is modified by spraying a polymer, for instance povidione, having a low surface energy (for instance of about 38 mN/m) on the surface of the intragranular particles. After the modification of surface properties, the intragranular particles are no longer cohesive, or are significantly less cohesive, and are easily rendered to all pharmaceutical operations. The surface-modified intragranular particles are then further processed, typically by combining with extragranular components typically comprising a filler, a disintegrant and a lubricant, and further processed into dry-filled capsules or tablets for oral administration. The surface modified intragranular components can also be used directly to make dosage forms without combination with extragranular components, for instance in connection with dry-filled capsules.

[0020] In certain embodiments, solid pharmaceutically acceptable compositions of neratinib maleate are provided in the form of tablets prepared by wet granulation. Increasing the glidant and lubrication levels provided an improved neratinib maleate formulation that flowed without aggregation of granules, as compared to a wet granulated neratinib maleate formulation used in clinical trials. The glidant was increased from 0.5% to 2.0 % to improve the flow of the preblended material. The picking and sticking issues observed during compression were eliminated by increasing the lubrication level from 0.5% to 3.0%, in certain embodiments 0.5% to 2.0%. In certain examples the amount of lubricant such as magnesium stearate needed is as low as 0.2% or even 0.1%. The increase in amount of glidant and lubricant was compensated by a corresponding reduction in the amount of fillers added to the formulation.

[0021] In some cases, the active ingredient comprises a 4-amino-3-cyanoquinoline compound such as neratinib, specicially neratinib maleate, or a pharmaceutically acceptable salt thereof. Suitable examples of 4-amino-3-cyanoquinoline compounds are disclosed in U.S. Pat. Nos. 6,002,008, 6,288,082, 6,297,258, 6,384,051 and 7,399,865. According to one embodiment, neratinib maleate is the active ingredient. The active ingredient comprises 35 weight % or 41 weight %, based upon total weight of the formulation.

[0022] According to one embodiment, a surface modifying agent is sprayed onto particulate intragranular components before further processing with extragranular components. Suitable surface modifying agents include, but are not limited to for example, povidone, gelatin, starch, hydroxy propyl methyl cellulose and hydroxy propyl cellulose. In one embodiment, povidone is the surface modifying agent. The surface modifying agent in the claimed composition comprises from about 5 weight % to about 15 weight %, including from 5-10 weight %, based

upon total weight of the formulation.

[0023] Suitable fillers (also referred to as "diluents") are known in the art. For example, suitable fillers include but are not limited to starch, dextrin, sucrose, Sorbitol, Sodium Saccharin, Acesulfame potassium, Xylitol, Aspartame, Mannitol, starch, PVP (polyvinyl pyrrolidone), low molecular weight HPC (hydroxypropyl cellulose), microcrystalline cellulose (MCC), low molecular weight HPMC (hydroxypropyl methylcellulose), low molecular weight carboxymethyl cellulose, ethylcellulose, dicalcium phosphate, silicified microcrystalline cellulose, alginates, gelatin, polyethylene oxide, acacia, dextrin, sucrose, magnesium aluminum silicate, and polymethacrylates. Fillers include agents selected from the group consisting of microcrystalline cellulose, starch, lactitol, lactose, a suitable inorganic calcium salt, sucrose, glucose, mannitol, silicic acid, or a combination thereof. The fillers, as an intragranular component in the claimed composition, comprise from about 15 weight % to about 65 weight %, based upon total weight of the formulation. In one embodiment, the intragranular filler is a combination of mannitol and microcrystalline cellulose. The fillers, as an extragranular component in the claimed composition, comprise from about 1 weight % to about 25 weight %, based upon total weight of the formulation. In one embodiment, the extragranular filler is microcrystalline cellulose.

[0024] Suitable disintegrants are known in the art and include but are not limited to, agar, calcium carbonate, potato or tapioca starch, alginic acid, certain silicates, sodium carbonate, crospovidone (cross-linked PVP), sodium carboxymethyl starch (sodium starch glycolate), cross-linked sodium carboxymethyl cellulose (croscarmellose), pregelatinized starch (starch 1500), microcrystalline starch, water insoluble starch, sodium starch glycolate, potassium polacrilin, sodium alginate, calcium carboxymethyl cellulose, magnesium aluminum silicate (Veegum) or a combination thereof. In some embodiments, a disintegrant is crospovidone. The disintegrant, as an intragranular component in the claimed composition, comprises from about 0.5 weight % to about 8 weight %, including from 0.5-6 weight %, based upon total weight of the formulation. The disintegrant, as an extragranular component in the claimed comprises from about 1 weight % to about 8 weight %, based upon total weight of the formulation.

[0025] A glidant is used as an intragranular component of the formulation. Suitable glidants include, without limitation, colloidal silicon dioxide, talc, magnesium carbonate, calcium silicate, fumed silicon dioxide combinations thereof. The glidant is colloidal silicon dioxide. The amount of glidants used in the claimed composition is 0.2-8 weight percent, or 0.2-5 weight percent, including 0.5-2 weight %, based on the total weight of the formulation.

[0026] A lubricant is used as an extragranular component of the formulation. Suitable lubricants or glidants include for example stearates, sodium stearyl fumarate, magnesium salts and magnesium stearate. The lubricant is magnesium stearate. The amount of lubricants used in the claimed composition is 0.5-3 weight %, based on the total weight of the formulation.

[0027] Provided compositions may be formulated into a unit dosage form. Such formulations are well known to one of ordinary skill in the art. In certain embodiments, the present invention

provides a formulation comprising a solid dosage form as a tablet. In other embodiments, the present invention provides a solution for oral administration. In some embodiments, a unit dosage form contains 5, 10, 20, 25, 40, 50 mg, 75 mg, 100 mg, 125 mg, 150 mg, 175 mg, 200 mg, 225 mg, 250 mg, 275 mg, 300 mg, 325 mg, 350 mg, 375 mg, 400 mg, 425 mg, 450 mg, 475 mg, or 500 mg, 525 mg, 550 mg, 575 mg, 600 mg, 625 mg, 650 mg, 675 mg, 700 mg, 725 mg, 750 mg, 775 mg, 800 mg, 825 mg, 850 mg, 875 mg, 900 mg, 925 mg, 950 mg, 975 mg, 1000 mg, 1025 mg, 1050 mg, 1075 mg, 1100 mg, 1125 mg, 1150 mg, 1175 mg, 1200 mg, 1225 mg, 1250 mg, 1275 mg, 1300 mg, 1325 mg, 1350 mg, 1375 mg, 1400 mg, 1425 mg, 1450 mg, 1475 mg, 1500 mg of neratinib In some embodiments, a unit dosage form contains between 5 mg and 500 mg, inclusive, or between 10 mg and 450 mg, inclusive, of neratinib. In some embodiments, a unit dosage form contains 40 mg, 80 mg, 100 mg, 120 mg, 240 mg, 360 mg, or 480 mg. In some embodiments, a unit dosage form contains more than 500 mg of neratinib.

[0028] In some embodiments, satisfactory results are obtained when the compounds of the invention are administered at a daily dosage of from about 0.5 to about 1000 mg/kg of body weight, optionally given in divided doses two to four times a day, or in sustained release form. The total daily dosage is projected to be from about 1 to 1000 mg, preferably from about 2 to 500 mg. Dosage forms suitable for internal use comprise from about 0.5 to 1000 mg of the active compound in intimate admixture with a solid or liquid pharmaceutically acceptable carrier. This dosage regimen may be adjusted to provide the optimal therapeutic response. For example, several divided doses may be administered daily or the dose may be proportionally reduced as indicated by the exigencies of the therapeutic situation.

[0029] For the treatment of cancer, the inventive formulations of this invention can be administered in combination with other anti-tumor substances or with radiation therapy. These other substances or radiation treatments can be given at the same or at different times as the compounds of this invention. These combined therapies may effect synergy and result in improved efficacy. For example, the compounds of this invention can be used in combination with mitotic inhibitors such as taxol or vinblastine, alkylating agents such as cisplatin or cyclophosamide, anti-metabolites such as 5-fluorouracil or hydroxyurea, DNA intercalators such as adriamycin or bleomycin, topoisomerase inhibitors such as etoposide or camptothecin, antiangiogenic agents such as angiostatin, and antiestrogens such as tamoxifen.

[0030] Based on the results disclosed for neratinib and other 4-amino-3-cyanoquinoline compounds in U.S. Pat. No. 6,297,258, the invented formulations are useful antineoplastic agents of significant efficacy, which are useful in treating, inhibiting the growth of, or eradicating neoplasms. In particular, the compounds of this invention are useful in treating, inhibiting the growth of, or eradicating neoplasms that express the receptor protein produced by the erbB2 (Her2) oncogene.

3. Combined Administration:

[0031] In certain embodiments, inventive compositions, and formulations thereof, may be administered alone to treat one or more disorders as described herein, or alternatively may be administered in combination with (whether simultaneously or sequentially) one or more other active agents useful to treat one or more disorders as described herein. Thus, an inventive composition, or formulation thereof, can be administered concurrently with, prior to, or subsequent to, one or more active agents.

[0032] In certain described aspects, compositions include one or more other active agents in addition to neratinib that is not neratinib. In some described aspects, formulations comprise both another anticancer compound and neratinib.

[0033] The amount of additional active agent(s) present in combination compositions will typically be no more than the amount that would normally be administered in a composition comprising that active agent as the only therapeutic agent. In certain described aspects, the amount of additional active agent will range from about 50% to 100% of the amount normally present in a composition comprising that compound as the only therapeutic agent.

4. Uses and Kits of Inventive Compositions:

[0034] Provided compositions, and formulations thereof, are also useful in treatment of conditions including cancers.

[0035] In still further embodiments, veterinary applications (e.g., treatment of domestic animals, e.g. horse, dogs, cats, etc.) of use of inventive compositions, and formulations thereof, are provided. Thus, use of provided formulations in veterinary applications analogous to those discussed above for human subjects is contemplated.

[0036] It will also be appreciated that inventive compositions, and formulations thereof, can be employed in combination therapies, that is, an inventive composition, or formulation thereof, can be administered concurrently with, prior to, or subsequent to, one or more other desired therapeutics or medical procedures. Particular combination therapies (therapeutics or procedures) to employ in a combination regimen will take into account compatibility of the desired therapeutics and/or procedures and the desired therapeutic effect to be achieved. It will also be appreciated that therapies employed may achieve a desired effect for the same disorder (for example, a formulation may be administered concurrently with another compound used to treat the same disorder), or they may achieve different effects (e.g., control of any adverse effects). As used herein, additional therapeutic compounds which are normally administered to treat or prevent a particular disease, or condition, are known as "appropriate for the disease, or condition, being treated".

[0037] In other embodiments, inventive compositions, and formulations thereof, and unit dose forms are useful in preparation of medicaments, including, but not limited to medicaments useful in the treatment of cancer.

[0038] Still further encompassed by the invention are pharmaceutical packs and/or kits comprising inventive compositions, and formulations thereof, and a container (e.g., a foil or plastic package, or other suitable container). Optionally instructions for use are additionally provided in such kits.

[0039] In order that the invention described herein may be more fully understood, the following examples are set forth. It should be understood that these examples are for illustrative purposes only and are not to be construed as limiting this invention in any manner.

[0040] All features of each of the aspects of the invention apply to all other aspects *mutatis mutandis*.

EXAMPLES

Example 1. Preparation of Coated Tablets of a Neratinib Maleate Formulation by Fluid Bed Wet Granulation Process (as reference)

[0041] A pharmaceutically acceptable formulation of neratinib mealate is prepared: a granulation comprising intragranular components (a) 10-70 weight percent of neratinib maleate; (b) 15-65 weight percent of mannitol and microcrystalline cellulose; (c) 0.5-8 weight percent of crospovidone or crosscarmellose sodium; (d) 0.2-8 weight percent of colloidal silicon dioxide, and (e) 5-15 weight percent of povidone. The granulation is combined with extragranular components (f) 4-25 weight percent of microcrystalline cellulose; (g) 1-8 weight percent of crospovidone and (h) 0.5-3 weight percent of magnesium stearate, and then compressed into tablets or dry-filled into capsules. This and certain preferred ranges of materials are shown below in Table 1.

[0042] The formulation was prepared according to the following procedure:

1. Mixed neratinib maleate, mannitol, microcrystalline cellulose and crospovidone and silicon dioxide. Any diffusive or convective mixer may be used.

Table 1

Component	Weight %	Weight % range	Weight % range
Intraganular Components			
HKI-272 Maleate, anhydrous	41	20-50	10-70
Mannitol			15-65
Microcrystalline Cellulose			5-35
Crospovidone/croscarmellose sodium	0.5-5.00	2-6	0.5-8
Povidone	5-10	3-12	1-15

Component	Weight %	Weight % range	Weight % range
Intraganular Components		***************************************	***************************************
Colloidal Silicon Dioxide	0.5-2.00	0.2-3.0	0.2-5
Extraganular Components			
Microcystalline Cellulose	4-25	4-25	4-25
Crospovidone	1-8	1-8	1-8
Magnesium Stearate	0.5-3.0	0.2-3.5	0.2-4.0
Total (Core Tablet)	1.1.1	1.1.2	1.1.3
Film Coat:			
Opadry [®]			

- 2. Dissolved povidone in purified water.
- 3. Fluidized the powder blend in step 1 and sprayed it with solution prepared in step 2 in a suitable fluid bed granulator and dryer.
- 4. Dried the granulation.
- 5. Milled the granulation.
- 6. Added microcrystalline cellulose and crospovidone to the granulation in step 5 and mixed.
- 7. Added magnesium stearate to the mixture in step 6 and mixed.
- 8. Compressed the blend in step 7 into tablets of desired strength.
- 9. Applied film-coat to compressed tablets using Opadry II of desired color.
- 10. Alternatively the powder blend can be filled in capsules shells.

Example 2. Unit dosage forms of an Exemplary Neratinib Maleate Formulation (as reference)

[0043] Using the fluid bed process described in Example 1, different unit dosages of neratinib maleate were prepared from an exemplary formulation, as summarized in Table 2.

Table 2

Ingredient	Function	Granulation wt/wt (%)	40 mg mg/tablet	80 mg mg/tablet	240 mg mg/tablet
Intragranular Components					
HKI-272 Maleate ^a	Active	35.00	40.00	80.00	240.00
Mannitol (Pearlitol 200 SD)	Diluent	38.94	44.50	89.01	267.02
Macrocrystalline (Avicel PH 101)	Diluent	10.56	12.07	24.14	72.41
Crospovidone	Disintegrant	3.00	3.43	6.86	20.57
Povidone USP/K-	Surface	5.00	5.71	11.43	34.29

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Ingredient	Function	Granulation wt/wt (%)	40 mg mg/tablet	80 mg mg/tablet	240 mg mg/tablet
Intragranular Components					
25	Modifying Agent				
Colloidal SiO ₂	Glidant	2.00	2.29	4.57	13.71
Extragranular Components					
Avicel PH 101	Diluent	1.50	1.71	3.43	10.29
Crospovidone	Disintegrant	2.00	2.29	4.57	13.71
Mg Stearate	Lubricant	2.00	2.29	4.57	13.71
Total Weight		100.00	114.29	228.57	685.71
Film Coating:					
Opadry II (85F15443) Red	Film Coat	-	3.429 (3%)	-	-
Opadry II (85F92177) Yellow	Film Coat	_	-	6.86 (3%)	-
Opadry II (85F94211) Pink	Film Coat	-	-	-	20.57 (3%)
Total Tablet Wt		_	117.714	235.43	706.28
^a Weight percent as HKI-272 free base					

Example 3. Coated Tablets of Targeted Release Neratinib Maleate Manufactured by Spraying Povidone on Intragranular Components in a Fluid Bed.

[0044] An exemplary targeted release (TR) neratinib maleate formulation is summarized in Table 3.

Example 4. Coated Tablets of Slow Release Neratinib Maleate Manufactured by Spraying Povidone on Intragranular Components in a Fluid Bed.

[0045] An exemplary slow release	(SR) neratinib	maleate	formulation i	is summarized in	Tables
4A and 4B.					

Table 3

Ingredient	% Wt/Wt	40 mg Tablet (mg)	240 mg Tablet (mg)	Function
Intragranular Components				
HKI-272 Maleate, anhydrous	35.00	40.00	240.00	Active
Mannitol	38.25	43.79	262.72	Filler
Microcrystalline Cellulose	12.75	14.50	86.99	Filler
Crospovidone	3.00	3.43	20.57	Disintegrant
Povidone	5.00	5.71	34.29	Surface Modifying Agent
Colloidal Silicon Dioxide	0.50	0.57	3.43	Glidant
Extragranular Components				
Microcystalline Cellulose	3.00	3.43	20.57	Filler
Crospovidone	2.00	2.29	13.71	Disintegrant
Magnesium Stearate	0.50	0.57	3.43	Lubricant
Total (Core Tablet)	100.00	114.29	685.71	
Film Coat:				
Opadry [®]	-	3.43	20.57	Film Coat

Table 4A (as reference)

Ingredient Name	Function	& Wt/Wt	mg/tablet240mg Tablet
Intragranular Components	***************************************		
HKI-272 Maleate	Active	35.00	240.00
Mannitol USP	Filler	41.81	286.69
Microcrystalline Cellulose NF	Filler	9.19	63.02
Crospovidone NF	Disintegrant	0	
Colloidal Silicon Dioxide, NF	Glidant	0.50	3.43
Povidone USP/K-25	Binder	10.00	68.57
Purified Water, EP/BP/USP	Solvent	-	
Extragranular Components			
Microcystalline Cellulose NF	Filler	-	
Crospovidone NF	Disintregrant	0.50	3.43
Magnesium Stearate NF/Ph/EU,	Lubricant	3.00	20.57

Extragranular Components	***************************************	
(Vegetable grade)		

Table 4B

Function	Granulation wt/wt (%)	40 mg mg/tablet	80 mg mg/tablet	240 mg mg/tablet
Active	35.00	40.00	80.00	240.00
Diluent	38.94	44.50	89.01	267.02
Diluent	10.56	12.07	24.14	72.41
Disintegrant	3.00	3.43	6.86	20.57
Surface Modifying Agent	5.00	5.71	11.43	34.29
Glidant	2.00	2.29	4.57	13.71
Diluent	1.50	1.71	3.43	10.29
Disintegrant	2.00	2.29	4.57	13.71
Lubricant	2.00	2.29	4.57	13.71
	100.00	114.29	228.57	685.71
Film Coat	-	3.429 (3%)	-	_
Film Coat	-	-	6.86 (3%)	-
Film Coat	-	-	-	20.57 (3%)
	-	117.714	235.43	706.28
	Active Diluent Disintegrant Surface Modifying Agent Glidant Diluent Disintegrant Lubricant Film Coat Film Coat	Active 35.00 Diluent 38.94 Diluent 10.56 Disintegrant 3.00 Surface Modifying Agent Clidant 2.00 Diluent 1.50 Disintegrant 2.00 Lubricant 2.00 Film Coat - Film Coat -	Active 35.00 40.00 Diluent 38.94 44.50 Diluent 10.56 12.07 Disintegrant 3.00 3.43 Surface Modifying Agent 5.00 5.71 Glidant 2.00 2.29 Diluent 1.50 1.71 Disintegrant 2.00 2.29 Lubricant 2.00 2.29 Lubricant 2.00 114.29 Film Coat - 3.429 (3%) Film Coat - - Film Coat - - Film Coat - -	wt/wt (%) mg/tablet mg/tablet Active 35.00 40.00 80.00 Diluent 38.94 44.50 89.01 Diluent 10.56 12.07 24.14 Disintegrant 3.00 3.43 6.86 Surface Modifying Agent 5.00 5.71 11.43 Glidant 2.00 2.29 4.57 Diluent 1.50 1.71 3.43 Disintegrant 2.00 2.29 4.57 Lubricant 2.00 2.29 4.57 Lubricant 2.00 114.29 228.57 Film Coat - 3.429 (3%) - Film Coat - 6.86 (3%) Film Coat - - 6.86 (3%)

Example 5: Drug Release Data

and 4, as summarized in Table 5. Dissolution of tablets was carried out employing 900ml of 0.1N HCl as dissolution medium in USP dissolution apparatus 2, and a paddle speed of 50 ± 1 rpm at 37 ± 0.5 °C. Samples were taken at specific time points and were analyzed by UV spectrometer at 266 nm.

[0047] Mean pharmacokinetic parameters for neratinib maleate in targeted release and slow release formulations following administration of a single oral dose (240-mg tablet) in subjects were evaluated and summarized in Table 6. The mean concentration versus time profiles for targeted release and slow release formulations are summarized in Figure 1.

Table 6. Summary of Mean Pharmacokinetic Parameters for Neratinib Maleate Formulations Following Single Oral Dose (240-mg tablet) in Healthy Subjects Under Fed Conditions

Mean ± SD (CV%)	240 mg TR	240 mg SR
[Geometric Mean]	(n=26)	(n=26)
C _{max} (ng/mL)	81.38 ± 27.49 (34)	82.21 ± 29.73 (36)
and the same of th	[77.13]	[77.12]
t _{max} (hr)	5.00	6.00
	(3.00, 12.00)	(3.00, 12.00)
AUC	1432 ± 450 (31)	1474 ± 467 (32)
	[1358]	[1393]

Table 5

Time (minutes)	% Dissolved		
and the same of th	TR Tablet Formulation	SR Tablet Formulation	
15	47	24	
30	83	43	
45	99	64	
60	99	82	

REFERENCES CITED IN THE DESCRIPTION

Cited references

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Patentkray

- **1.** Tablet omfattende en farmaceutisk acceptabel sammensætning omfattende intragranulære komponenter:
 - (a) 35 eller 41 vægtprocent neratinibmaleat;
- 5 (b) 15-65 vægtprocent mannitol, mikrokrystallinsk cellulose eller en kombination af begge;
 - (c) 0,5-8 vægtprocent crospovidon, croscarmellosenatrium eller en kombination af begge;
 - (d) 0,2-8 vægtprocent kolloid siliciumdioxid; og
- 10 (e) 5-15 vægtprocent povidon;

og ekstragranulære komponenter:

- (f) 1-25 vægtprocent mikrokrystallinsk cellulose;
- (g) 1-8 vægtprocent crospovidon; og
- (h) 0,5-3 vægtprocent magnesiumstearat.

15

- **2.** Tablet ifølge krav 1, hvor mængden af mannitol og mikrokrystallinsk cellulose, som en intragranulær komponent, er omkring 50 vægtprocent.
- 3. Tablet ifølge krav 1, hvor mængden af crospovidon, som en intragranulærkomponent, er omkring 3 vægtprocent; og/eller hvor mængden af povidon, som en intragranulær komponent, er omkring 5 vægtprocent.
 - **4.** Tablet ifølge et hvilket som helst af kravene 1 til 3, i form af en oral dosis på 40 mg, 80 mg eller 240 mg.

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- **5.** Tablet ifølge krav 1, hvor sammensætning omfatter
 - (b) mannitol og mikrokrystallinsk cellulose og
 - (c) crospovidon.
- 30 **6.** Tablet ifølge et hvilket som helst af kravene 1 til 5 til anvendelse i en fremgangsmåde til behandling af cancer.

DRAWINGS

