- Astus *PATENT APPLICATION & GRM (CONVENTION AND NON-CONVENTION) COMMONWEALTH OF A GIRALIA 2 7 2 egulation 9 Patents Act 195 APPLICATION FOR A STANDARD PATENT OR A STANDARD PATENT OF ADDITION APPLICATION ACCEPTED AND AMENDMENTS 15.3.90 ALLOWED PATENT OFFICE ... SUB-OFFICE +/We (a) HOECHST AKTIENGESELLSCHAFT (a) Insert full name(s) of applicant(s) 9 Collector 6230 Frankfurt/Main 80, Germany Monevs (b) insert address(es) of applicant(s) of (b) 6 hereby apply for the grant of a (c) Standard Patent for an invention entitled (d) . . (c) Delete as "Decorative panel having improved surface properties" (c) Insert title which is described in the accompanying (c) provisione complete specification. (e) For a Convention application – details of basic application(s) – COUNTRY DATE OF APPLICATION NUMBER (e) for Convention cases only P 35 33 737.0 21st September 1985 Germany (f) For Patents of Addition (Section (f) For Patents of Addition only. I/We request that the Patent may be granted as a Patent of Addition LODGED AT SUB-OFFICE (g) Insert number of 'parent/main' application/or the Patent applied for or Application No. (g) 2 2 SEP 1986 tent as propriate. \ldots in the name of ^(h) . . Patent No. ^(g) Sydney application of 'parentinin application of patent as ap-propriate. e request that the term of the Patent of Addition be the same as that for the main invention or so much of the -My/Our address for service is ARTHUR S. CAVE & CO., Patent and Trade Mark Attorneys, 1 Alfred Street, Sydney, New South Wales, Australia 2000. (i))ns HOECHST AKTIENGESELLSCHAFT By Its Patent Attomeys (j) Signature of applicant or Australian attorney. ARTHUR S. CAVE & CO. (j) (Signature) (k) Seal, if any. (k) To: **Commissioner of Patents** JAMES G. SIELY, F.I.P.A.A. ARTHUR S. CAVE & CO. PATENT AND TRADE MARK ATTORNEYS SYDNEY

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Regulation 12(2)

PATENT DECLARATION FORM (CONVENTION) COMMONWEALTH OF AUSTRALIA

States.

Patents Act 1952

DECLARATION IN SUPPORT OF A CONVENTION APPLICATION FOR A PATENT

In support of the Convention application made for a patent for an invention entitled: DECORATIVE PANEL HAVING IMPROVED SURFACE PROPERTIES

I, Hector Cumming of Level 10, 10 Barrack Street, Sydney New South Wales 2000 Australia do solemnly and sincerely declare as follows:-

1. I am/We are authorised by HOECHST AKTIENGESELLSCHAFT the applicant for the patent to make this declaration on its behalf.

2. The basic Application(s) as defined by Section 141 of the Act was/were made in the following country or countries on the following date namely:in Germany on 21st Septebmer, 1985 by Hoechst Aktiengesellschaft

3. Johannes Christianus Willem <u>van der Hoeven</u> of Graaf Philippsstraat 15 NL-6085 CM Horn Netherlands is the actual inventor of the invention and the facts upon which the applicant(s) are entitled to make the application are as follows:

The Applicant is the Assignee of the said invention from the actual inventor.

4. The basic application(s) referred to in paragraph 2 of this Declaration was the first application(s) made in a Convention country in respect of the invention the subject of the application.

Declared at Sydney this gh day of flasch, 1990

To:

259595/JAW

The Commissioner of Patents

HECTOR CUMMING

ARTHUR S. CAVE & CO. PATENT AND TRADE MARK ATTORNEYS SYDNEY

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(12) PATENT ABRIDGMENT (11) Document No. AU-B-63036/86 (19) AUSTRALIAN PATENT OFFICE (10) Acceptance No. 597272

(54)	Title DECORATIVE PANEL HAVING IMPROVED SCRATCH RESISTANT SURFACE PROPERTIES
(51)⁴	International Patent Classification(s) B44C 003/02 B32B 027/30 B32B 031/20 B32B 031/28 B32B 033/00 E04C 002/24
(21)	Application No. : 63036/86 (22) Application Date : 22.09.86
(30)	Priority Data
(31)	Number (32) Date (33) Country 3533737 21.09.85 DE FEDERAL REPUBLIC OF GERMANY
(43)	Publication Date : 26.03.87
(44)	Publication Date of Accepted Application : 31.05.90
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(74)	Attorney or Agent ARTHUR S. CAVE & CO.
(56)	Prior Art Documents EP 166153 CA 1150178 US 4503115
(57)	Claim
1.	A decorative panel, consisting of:
	a core layer having the substrate function of the
decorative panel;	
	an outermost layer applied over said core layer, said
oute	ermost layer containing the radiation-polymerized product of
a ra	adiation-polymerizing mixture, said mixture containing
(i)	a radiation-polymerizable component selected from the
	group consisting of an epoxy-acrylate or -methacrylate
	oligomer, a silicone-acrylate or -methacrylate oligomer,

a polyester acrylate or -methacrylate oligomer or a urethane acrylate or -methacrylate oligomer,

(ii) a diacrylate or triacrylate or -methacrylate monomer,

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(11) AU-B-63036/86 (10) 597272

wherein said radiation-polymerized outermost layer is thermopressed under conditions of elevated temperature and pressure such that a scratch resistance of at least 1.5 Newtons is imparted to said outermost layer and having a reflectometer value no greater than 50 at an angle of incidence of 85°; and

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decoration means for forming a decoration covering said core layer.

COMMONWEALTH OF AUSTRALIA 5972.7.2

PATENTS ACT, 1952

COMPLETE SPECIFICATION

(ORIGINAL)

FOR OFFICE USE

Short Title:

Int. Cl:

Application Number: 63036/86. Lodged:

Complete Specification-Lodged: Accepted: Lapsed: Published: '.'.' Priority: '.'.' Related Art:

This document contains the amendments made under Section 49 and is correct for printing.

Regulation 13(2)

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TO BE COMPLETED BY APPLICANT

•• Name of Applicant: HOECHST AKTIENGESELLSCHAFT

Address of Applicant: 6230 Frankfurt/Main 80, Germany

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ARTHUR S. CAVE & CO., Patent and Trade Mark Attorneys, 1 Alfred Street, Sydney, New South Wales, Australia, 2000.

Complete Specification for the invention entitled: "Decorative panel having improved surface properties"

The following statement is a full description of this invention, including the best method of performing it known to me:-

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DECORATIVE PANEL HAVING IMPROVED SURFACE PROPERTIES

BACKGROUND OF THE INVENTION

The invention relates to a decorative panel 5 composed of a core layer and a layer decorated on one or both sides and to a process for its production. Panels of this type are employed for interior or exterior uses in the building industry, being used as cladding panels or self-supporting units, depending on 10 their thickness.

The decorative panels used hitherto are, for example, high pressure decorative laminates (H.P.D.L.) (DIN 19,926), as

they are called. They are composed of a stack of paper webs impregnated with resin and compressed under hot conditions, as the core layer, and of a top layer made of resin-impregnated decorative paper. These panels have the disadvantage that they are attacked by mineral acids, especially at concentrations above 10% and at an exposure time longer than 10 minutes. In addition, in the standard embodiment, these panels are not adequately resistant to weathering, since the type of resin used in the top layer is sensitive to hydrolysis. Panels of this type can, therefore, only be used to a limited extent as work benches in chemical laboratories or for the production of wet cells which have to be

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cleaned with acids. If they are used for exterior purposes, additional expensive measures are necessary to improve their resistance to the effects of weathering.

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Laminates and panels based on plastics, such as polyester or acrylate panels, are, however, particularly sensitive to scratching and are not adequately resistant to organic solvents. For this reason they are also not very suitable for these 10 applications.

Application 42562/85, ation -85,105,851.1, which does not constitute a prior publication, relates to a decorative panel which is particularly suitable for exterior uses, for interior construction and for the 15 production of special furniture, and which has a surface which is not sensitive to hydrolysis and is adequately resistant to the effects of weathering, mineral acids and organic solvents, and which also has a high surface hardness. This is composed of a core 20 layer and a decorative layer on one or both sides. At least the outermost layer of the panel on at least one of the two panel surfaces is composed predominantly of a synthetic resin formed from one or more components, polymerized by radiation and selected from the group 25 comprising unsaturated acrylates and methacrylates. This layer has a particularly high surface hardness. It is still scratch-resistant at a scratch loading of at least 1.5 N, preferably 2 to 7 N (DIN 53,799, part 10). In the process for the production of this panel, 30 a liquid surface layer comprising the components which can be polymerized by radiation is applied to a substrate and is subsequently polymerized by radiation. The panel surface does not have the desired properties until after a further stage in which the surface layer

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which has been polymerized by radiation is compressed together with the substrate at an elevated temperature. However, this decorative panel has the property, which is frequently undesirable, of exhibiting a certain degree of gloss. If texturized 5 separating media are used in the final compression under heat, it is admittedly possible to give the panel surface a surface structure which is texturized in conformity with the surface of the separating medium, for example a surface structure similar to orange peel, 10 but its surface gloss is, as before, very high. The addition of known delustering agents, such as silicon dioxide pigments, to the outermost surface layer of the panel also does not, for practical purposes, reduce the gloss, since the pigment-containing surface, which is 15 initially still satin-frosted after the radiation polymerization, unaccountably becomes glossy again as soon as the panel is subsequently subjected to compression under heat.

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SUMMARY OF THE INVENTION

It is therefore an object of the invention to provide a decorative panel which is resistant to weathering, acids and solvents and which has a high surface hardness and only a slight surface gloss.

This object is achieved by a decorative panel which comprises a core layer; an outermost layer applied over the core layer, the outermost layer being a synthetic resin polymerized by radiation and comprising an unsaturated acrylate or an unsaturated methacrylate, and further being scratch resistant at a scratch loading of at least about 1.5 Newtons as defined by DIN 53,799 part 10 and having a reflectometer value no greater than about 50 at an

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angle of incidence of 85° as defined by DIN 67,530; and decoration means for forming a decoration covering the core layer.

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The object of the invention is also achieved by a process for the production of a 5 decorative panel which comprises applying a liquid synthetic resin layer to a substrate; applying a film having a surface roughness over the resin layer; transferring the surface roughness to the resin layer; polymerizing the resin layer by exposure to radiation; 10 and compressing the resin layer and the substrate at an elevated temperature.

The object of the invention is further achieved by a process for the production of a decorative panel which comprises applying a first layer containing decoration means to a substrate; applying a second layer over the first layer, the second layer being a synthetic resin; applying a film having a surface roughness over the second layer; transferring 20 the surface roughness to the second layer; polymerizing the resin layer by exposure to radiation; and compressing the first and second layers and the substrate at an elevated temperature.

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BRIEF DESCRIPTION OF THE DRAWINGS

The invention is illustrated in greater detail by means of Figs. 1 to 3a and the examples below.

Fig. 1 shows the sequence of the process variant described in Example 1;

Fig. la shows a partial cross-section through a panel according to Fig. 1;

Fig. 2 shows the sequence of the process variant described in Example 2;

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Fig. 3 shows the sequence of the process variant described in Example 3; and

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Fig. 3a shows a partial cross-section through a panel according to Figs. 2 and 3.

DETAILED DESCRIPTION OF THE PREFERRED EMBODIMENTS

The panel of the invention is a sheet-like unit having a surface shape and surface structure suited to the intended use and which can, for example, also have a bent shape. Within the scope of the invention, a panel is also to be understood as sheeting.

In the measurement of scratch resistance carried out as specified in DIN 53,799, part 10, the force with which a diamond needle produces a visible scratch on the panel surface is determined. This assessment is made immediately after the action of the diamond needle, since, as a result of the elasticity of the surface layer, a slow recovery of the surface deformation can take place after the surface has been subjected to scratching. The scratch resistance is a measure of the surface hardness.

It has been found, surprisingly, that this decorative panel having the special layer of synthetic resin polymerized by radiation on at least one of its external surfaces not only has an excellent resistance to weathering, compared with the panels hitherto known, but, surprisingly, has an improved surface hardness. Additionally, it is appreciably less sensitive to acids and organic solvents.

The reflectometer value is determined using a type RB/Dr. Lange reflectometer as specified in DIN 67,530 as a measurable indication of the gloss of the panel surface. A reflectometer value determined as

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specified in this standard constitutes an optical characteristic value for the surface of a test specimen and bears a relation to the gloss of the surface. It must be borne in mind in this respect that the gloss is not a purely physical quantity, but is also a quantity conditioned by physiological and psychological considerations. Direct measurement of gloss is, therefore, not possible, but the "gloss power", that is to say the contribution made by the surface, by virtue of its reflectance properties, to the creation of the impression of gloss, can, in principle, be measured in a suitable manner. The reflectometer value can be used as a measure of the gloss power, since it is essentially determined by the reflectance properties of the surface.

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The reflectometer system defined by this standard is modeled on the layout described in ASTM D 523-67. The 20°, 60° and 85° angles of incidence are chosen arbitrarily.

The 20° geometry of measurement is used for test specimens having a 60° reflectometer value of over 70, while the 85° geometry of measurement is used for test specimens having a 60° reflectometer value of less than 30.

In the reflectometer, a light source is formed into an image centrally in the aperture of a diaphragm. The light rays strike the panel surface at the prescribed angle of incidence (20°, 60° or 85°) and are reflected in a scattered manner. The light flux passing through the diaphragm is measured by means of a photoelectronic detector located behind the diaphragm.

The core layer has the substrate function of the panel. It is composed, for example, of wood. Panels or sheeting made of plastics, for example based on polyvinyl chloride, polyethylene or polystyrene, or

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made of metal, for example steel, aluminum, copper, brass or other alloys, are also suitable as the core layer. The layer of synthetic resin polymerized by radiation is located right on the surface of these core layers or is attached to the core layer by means of film glues or glued joints, or preferably by means of adhesion-promoting synthetic resins, such as, for example, a phenol-formaldehyde or resorcinolformaldehyde precondensate. Glued joints are merely layers of adhesive, while film glues are carrier layers which are coated or impregnated with adhesive. Adhesion promoters are substances which, without themselves being an adhesive, promote the attachment of two different materials.

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The core layer can additionally be composed of the sheets of paper, especially sulfate kraft paper, which are customary in H.P.L. sheets and are impregnated with heat-curable synthetic resin, in particular phenol-formaldehyde resin, and which are compressed under hot conditions. Depending on the panel thickness desired, 1 to about 100 sheets on top of one another are compressed under hot conditions.

The core layer can also be composed of a nonwoven fabric or mats, consolidated under pressure and composed of mineral fibers, glass fibers, plastic 25 fibers or a mixture of fibers, preferably cellulose. Cellulose-containing fiber layers are, for example, wood fibers or wood chips distributed in a random manner. The nonwoven fabric or the mat composed of wood and/or cellulose fibers is prepared by applying a 30 synthetic resin to the fibers, drying the resin-coated fibers, shaping the product into a fiber mat and subjecting this mat to preliminary compression under the influence of pressure (European patent application No. 35 0,081,147).

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If appropriate, there is an underlay containing a heat-curable aminoplast or phenoplast resin located on the external surface(s) of this fibercontaining core layer. This underlay is composed, for example, of a pigmented or non-pigmented nonwoven fabric or paper.

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Immediately upon the fiber-containing core layer or on this underlay layer there follows, in a preferred embodiment, a layer of synthetic resin polymerized by radiation, which is decorative, i.e., 10 exhibits a particular optical effect or a decorative effect as a result of added dyestuffs. On the decorative layer of synthetic resin polymerized by radiation there can also be a clear, i.e., transparent and dyestuff-free, layer of synthetic resin polymerized 15 by radiation, which form(s) the outermost surface(s) of the panel; however, it is entirely possible to omit this clear layer of synthetic resin, so that the decorative layer(s) of synthetic resin then form(s) the outermost layer(s). 20

Instead of the decorative layer of synthetic resin, it is also possible to use a decorative layer based on a dyed and/or printed plastics film or based on paper; this is usually composed of pigmented, dyed and/or printed decorative 25 paper. The layer, in this case transparent and free from dyestuffs, of synthetic resin polymerized by radiation is located on the plastics film or on the decorative paper. The decorative paper contains a heat-curable synthetic resin which is customary for 30 this purpose, in particular an aminoplast resin, and is located on core layers which have been built up from the sulfate kraft paper typical for H.P.L panels or from wood or cellulose fibers distributed in a random 35 manner and treated with phenolic resins.

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The compounds envisaged for the preparation of the uppermost layer of synthetic resin, polymerized by radiation, embrace acrylic acid esters or methacrylic acid esters which can be polymerized by a 5 free radical mechanism by actinic radiation and which are present, on their own or together, in a polymerizable mixture. The preferred component is a polyfunctional prepolymer, i.e., a polyunsaturated prepolymer. In addition to this predominant component, 10 the copolymerizable mixture contains, if appropriate, a further component having a diluting action, which is described as a diluent monomer or diluent oligomer. The proportion in the mixture of the polyfunctional prepolymer is 50 to 100%, in particular 60 to 90%, by 15 weight of the total weight of copolymerizable components. Prepolymers of a low viscosity (less than 100 poise at 20°C) are employed without the monomers or oligomers producing dilution.

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The components used have a strong tendency 20 to polymerize by a free radical mechanism under the action of actinic radiation. Suitable actinic radiation is a light in the near UV region or highenergy radiation, for example electron, particle

or x-ray radiation. The prepolymer which can 25 be polymerized by free radical mechanism is a polyfunctional, unsaturated aliphatic or aromatic acrylate or methacrylate, preferably an unsaturated polyester acrylate oligomer and especially an aliphatic urethane acrylate oligomer. Although aromatic urethane 30 acrylate oligomers also produce scratch-resistant surface coatings, these yellow after some time in exterior applications.

In addition to the prepolymer, a mono-, di-, tri-, tetra-, penta- or hexaacrylate or -meth-35 acrylate, preferably a diacrylate or triacrylate, is

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used as an additional suitable monomer or oligomer in the mixture which can be copolymerized by a free radical mechanism. These mono- to hexaacrylates or mono- to hexamethacrylates are esters of polyols having 5 1 to 6 OH groups with acrylic acid or methacrylic acid, respectively, and are therefore also known as polyol acrylates or polyol methacrylates, respectively. Suitable diacrylates are esters of acrylic acid with aliphatic, dihydric alcohols, in particular ethylene 10 glycol, 1,2-propylene glycol, 1,3-propyleneglycol, butane diols, 1,6-hexane diol or neopentylglycol, with aliphatic ether-alcohols, in particular diethylene glycol, dipropylene glycol, dibutylene glycol, polyethylene glycols or polypropylene glycols, with 15 oxyalkylated compounds of the above-mentioned aliphatic alcohols and ether-alcohols or with aromatic dihydroxy compounds, in particular bisphenol A, pyrocatechol, resorcinol, hydroquinone, p-xylyleneqlycol or phydroxybenzyl alcohol. Preferred diacrylates are 1,6-20 hexanediol diacrylate, tripropylene glycol diacrylate and 1,4-butanediol diacrylate. Preferred triacrylates

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In addition to the urethane acrylate 25 oligomers and unsaturated polyester acrylate oligomers already mentioned, suitable polyfunctional prepolymers are also epoxy-acrylate and silicone-acrylate oligomers, which are preferably used together with the diacrylates or triacrylates mentioned in the mixture 30 which can be copolymerized by a free radical mechanism. The prepolymers are compounds known <u>per se</u> and are prepared, for example, from hydroxylated copolymers in which the hydroxyl groups are distributed statistically along the copolymer chain. Statistically 35 unsaturated acrylic copolymers are obtained from this

are trimethylolpropane triacrylate and pentaerythritol

triacrylate.

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copolymer by esterifying the hydroxyl groups with acrylic acid. Semi-terminal unsaturated acrylic copolymers are prepared by having the hydroxyl group at the end of the chain in the preparation of the 5 hydroxylated copolymers. Urethane acrylate oligomers are prepared by reacting (meth)acrylic acid esters containing hydroxyl groups, for example, hydroxyethyl methacrylate, with polyfunctional isocyanates, preferably diisocyanates. The diisocyanates or 10 polyisocyanates can preferably be reaction products of diols, polyether-diols or polyester-diols containing a stoichiometric excess of monomeric diisocyanate or polyisocyanate.

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If the polyfunctional prepolymer 15 preponderates in the polymerizable mixture, as the base resin, by virtue of its chemical nature, it determines the properties of the cured surface layer. The monoacrylate to hexaacrylate or monomethacrylate to hexamethacrylate added makes it possible as a diluent 20 monomer or oligomer, to adjust the viscosity of the mixture to be cured, which is normally within a viscosity range from 20 to 100 poise (20°C), and entirely takes part in the free radical polymerization. When irradiated, the coating is cured through free 25 radical polymerization between the double bonds of the prepolymer and of the diluent monomer or oligomer which may be present.

If curing is carried out under the influence of UV-light, it is necessary to add 30 photoinitiators which facilitate, by the formation of radicals, the initiation of the free radical polymerization. If curing is carried out with electron beams, no photoinitiators are required, however. Most of the photoinitiators contain at least 35 one carbonyl group conjugated with an aromatic ring. A

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photoinitiator system composed of several components is usually employed.

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In addition, the synthetic resin which is polymerized by radiation contains, if appropriate, 5 suitable additives such as plasticizers, fillers, dyestuff pigments, agents for improving abrasion resistance and stabilizers, in order to achieve the desired decorative, mechanical and physical surface properties. These substances include, for example, 10 barium sulfate, silica, aluminum oxide and light-stable pigments.

The decorative panel is produced by applying the liquid compounds which can be polymerized by radiation to the substrate to be coated, for 15 example, by spraying, casting, a doctor system, a roller or screen printing. If it has been applied to a decorative layer, the layer applied is then transparent. It can, however, also be decorative itself and it is then colored and is located on a non-20 decorative paper layer or right on the core layer. In a further embodiment, an additional layer which can be polymerized by radiation but which, however, is not decorative but transparent, is also applied to this decorative layer of synthetic resin after the curing by 25 radiation.

The substrate used for the application of the compounds which can be polymerized by radiation is thus a paper layer, a decorative paper layer or the above-mentioned core layers based on wood, plastics, 30 metal or a stack of further fiber-containing layers which forms the core of the panel subsequently obtained. The fiber-containing layers of the stack, which are preferably composed of sulfate kraft paper or of a nonwoven fabric made of wood and/or cellulose 35 fibers, contain the heat-curable, precured resins

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customary for H.P.L. panels, in particular phenolformaldehyde resins, whereas the paper sheets which may optionally additionally be present on the stack contain an aminoplast resin or, especially, a phenoplast resin. 5 The content of heat-curable resins is 20 to 250% by weight, relative to the particular layer.

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The immersion or impregnation of the fibercontaining layers or the paper layers is carried out, for example, by immersion in a bath of a solution or 10 dispersion containing the heat-curable resin or by application or spraying by means of a metering system. The solvent or dispersing agent is aqueous alcohol, aqueous acetone or water, depending on the synthetic resin used. It can also contain up to 20% by weight of 15 fire-retardant agents. The desired amount of resin is then distributed by scraping off or squeezing out, for example, by means of rollers.

The heat-curable resins of the substrate are precured and dried in a customary manner before the 20 application to the intended substrate of the compounds which can be polymerized by radiation.

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During the radiation polymerization, the outermost, still liquid, layer composed of compounds which can be polymerized by radiation is covered by a 25 sheet or panel of plastics or paper or a composite film made from several layers of plastics or layers of plastics and paper, which has a rough surface structure and which must be sufficiently transparent for the actinic radiation. The film or panel intended for 30 covering must not have a very porous surface, since otherwise there is a risk that the liquid compounds still capable of polymerization by radiation will penetrate into the surface. In this event it is no longer possible to remove the film or panel after 35 polymerization. This outermost liquid layer can itself

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be decorative and can contain a dyestuff or can be nondecorative, i.e., transparent, and can then be located on a decorative layer or on a decorative layer of polymerized by synthetic resin which will be It is preferable to employ films having a 5 radiation. thickness of up to 0.1 mm, since thicker coverings are not adequately transparent for electron beams or UV radiation, or require relatively long exposure times. In general, films of a thickness from 20 to 60 /um are 10 used, since, on the one hand, they are adequately transparent for the radiation and, on the other hand, they also have an adequate mechanical strength. For the sake of simplicity, they will be described as films or covering films in the following text.

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15 The plastics film is composed, in particular, of a polyester or polypropylene film which has been oriented by biaxial stretching. The rough structure of the film intended for covering is produced, for example, by the addition of pigments, at 20 least in the neighborhood of its outer surface. This surface roughness depends on elevations in the surface of the film, the height of which is, however, only slight in comparison with the thickness of the film and is within the range of a few micrometers at the most. 25 The pigments are composed, for example, of inorganic particles, in particular of aluminum oxide, aluminum sulfate, barium sulfate, calcium carbonate, magnesium carbonate, kaolin, talc, silicon dioxide, titanium dioxide or glass microbeads, or organic plastics 30 particles which are incompatible with the plastic of the film and which are dispersed as particles in the The pigments usually have a particle size from film. 0.1 to 20 /um, the average particle size being within the range from 0.1 to 4 /um. Their concentration is 35 0.01 to 10% by weight, relative to the weight of the

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film. The concentration of the pigments in the film and their size is adjusted to suit the desired surface roughness of the plastics film.

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The application of the covering film to the 55 liquid layer still capable of polymerization by radiation is effected by first applying this liquid layer to the substrate in the manner described above and then providing it with the covering film, the rough side of the surface of the covering film coming into 10 contact with the liquid, polymerizable layer. It is also possible, however, first to apply the liquid, polymerizable layer to the rough side of the surface of the covering film and then to apply the covering film together with this liquid layer to the substrate.

15 The roughness of the surface is transferred to the surface of the layer to be polymerized by radiation, which then assumes the surface structure of the covering film and obtains a mat appearance. This result is all the more surprising because the surface gloss of the end produce can, for practical 20 purposes, no longer be reduced by using texturized separating media in the final compression process.

The polymerization caused by radiation can be initiated by employing a customary source for the formation of free radicals, such as, for example, a 25 photoinitiator, or heat alone is supplied. If the photopolymerizable layer contains photoinitiators, the polymerization is initiated by passage under mercury The absence of oxygen is not necessary vapor lamps. for curing by means of UV radiation. It is appropriate 30 for the electron beams used for curing the polymerizable compounds to have an energy corresponding to 150 to 350 KeV. The energy of the electron accelerator is determined by the thickness of the synthetic resin layer to be formed, the dose of 35

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radiation required and the time of treatment or the speed with which it is carried out.

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The devices used for accelerating the electron beams are commercially available. These are 5 the accelerators known as the "scanner type" and the "linear cathode type". Free radicals are formed by interaction with the components of the polymerizable layer. This curing process is usually carried out at room temperature. For curing by means of electron 10 beams too, it is not necessary for this process to be carried out in an inert atmosphere, i.e., a substantially oxygen-free atmosphere, since the polymerizable surface layer is protected by the plastics film resting on it.

15 The covering film can be removed after the polymerization effected by radiation. It is also possible, however, not to remove the covering film until the completion of the process, i.e., after the compression under heat, or to use it as a casing for 20 the finished panel. If the substrates are sufficiently flexible, they are rolled up for storage or are immediately cut to the desired size. If the substrate containing the resin polymerized by radiation is only composed of a paper layer, it is laid on a stack of 25 fiber-containing layers forming the core layer. It is also possible additionally to provide the lower face of the stack with a substrate of this type.

The resulting bundle of layers composed of a fiber-containing core layer and surface layer(s) 30 polymerized by radiation and, if appropriate, intermediately placed layers of paper or decorative paper is compressed under hot conditions, as is customary in the production of H.P.L panels, to give a decorative panel, in the course of which the heat-35 curable resins ar cured. The temperature is preferably

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120 to 210°C, the pressure is within the range from 10 to 100 bar and the treatment time is 1 to 30 minutes. If, however, the core layer is composed of a wooden, plastics or metal panel, the temperature and the pressure can usually be reduced to values of 80°C and 5 bar.

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Compression is carried out in a daylight, s. .: cycle or continuous press.

The number and chickness of fiber-containing layers in 10 the core layer or the thickness of the core layer is selected depending on the use of the panel, panel thicknesses of 3 to 25 mm being required for exterior applications, depending on the intended use. If a large number of panels containing layers of synthetic 15 resin polymerized by radiation are stacked one on top of the other in the press, which is of economic advantage if the core layer has a low thickness, the individual panels are separated from one another in each case by means of a separating medium. The 20 separating medium is, for example, a paper layer, a plastics film or a metal plate. If the separating medium has a coarse surface structure, i.e., elevations or indentations, this coarse structure is imparted to the adjacent external layer of the particular panel, 25 the mat fine structure already present being retained. The degree of gloss on the panel surface, already established by the radiation process, cannot, for practical purposes, be modified further by means of the separating medium.

The decorative panels produced are, surprisingly, particularly resistant to weathering and extremely scratch-resistant, which could possibly be due to an unforeseeable interaction between the various resins or perhaps also to subsequent crosslinking of the compounds polymerizable by free radicals when

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compressed under heat. The scratch resistance and the resistance to chemicals are, surprisingly, substantially higher than in the case of a panel which receive a coating of the same compounds which can be polyme__zed by radiation, this coating -- without the compression under heat -- only having been polymerized by a free radical mechanism by means of radiation. In addition, the surface of the panel has a considerably reduced gloss, such as cannot be achieved even if delustering agents are used in the surface layer.

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Components having identical functions are marked with identical numbers in the figures. In Fig. 1, the sulfate kraft paper 1 containing partly cured synthetic resin is provided with a liquid layer 2 which contains dyestuff and can be polymerized by radiation. The layer 2 is covered by the plastics film 3 and is cured by means of electron beams in the apparatus 4. In Fig. 2, the dyestuff-containing layer 5 is already partly cured by radiation when it is provided with the 20 transparent liquid layer 6 which can be polymerized by radiation. The plastics film 3 is applied to the layer б. Fig. 3 differs from Fig. 1 only in that there is also a transparent liquid layer 6 which can be polymerized by radiation located on the plastics film In the figures, the guide rollers are marked 7, 8, 3. 9 and 10 and the coating devices are marked 11 and 12. Compressed panels 14 according to the layer arrangements of Figs. 1, 2 and 3 can be seen in Figs. la and 3a. The stack forming the core is marked 13.

In the following examples, the percentages are percent by weight.

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Example 1

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As shown in Fig. 1, a pasty liquid 2 (viscosity 50 poise at 25°C) composed of a mixture, polymerizable by radiation, of 85 parts by weight of an 5 aliphatic urethane acrylate oligomer as prepolymer, 15 parts by weight of hexane diol diacrylate as diluent monomer and 10 parts by weight of organic dyestuff pigments is applied to a sulfate kraft paper 1 which has first been impregnated with a heat-curable phenol-10 formaldehyde resin (amount of resin applied 70%), after the resin has been partly cured by means of rollers, whereupon a continuous film (layer thickness about Immediately afterwards, a 80/um) is formed. delustered, biaxially stretched polypropylene singleply film 3 containing 8% by weight of calcium carbonate 15 of average particle size 3 /um is applied to this film composed of compounds polymerizable by radiation, and the film is crosslinked in a substantially homogeneous manner by means of electron beams at room temperature 20 without the application of pressure. The dose of radiation absorbed is 60 KGy.

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After the plastics film 3 has been removed, the paper 1 with the external layer 2 of synthetic resin copolymerized by radiation is placed in each case on the external face of a stack 13 composed of 50 superposed sheets of paper. The sheets of paper have previously been impregnated with a heat-curable phenolformaldehyde resin, and the resin has been partly cured. The bundle of layers is compressed at 150°C and 80 bar for 10 minutes in a press between two texturized metal sheets. A bundle has the following composition:

A first decorative layer 2 (synthetic resin containing organic dyestuff pigments and polymerized by radiation) as the outer layer on a first previously

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impregnated paper layer 1, 50 paper webs (impregnated with a phenol-formaldehyde resin) as the core layer 13, and a second decorative layer 2 (synthetic resin containing organic dyestuff pigments and polymerized by radiation) as the outer layer on a previously impregnated second paper layer 1.

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The resulting panel 14, which is 10 mm thick and is decorated on both sides, has a scratch resistance higher than 3.0 N (DIN 53,799, part 10). It is insensitive to hydrolysis and shows no changes after being boiled in water for 100 hours. Its surface is not attacked by concentrated mineral acid during an exposure time of 6 hours (DIN 53,230). The fastness to light of this panel is given rating 8 (DIN 54,004). The resistance of the panel to the effects of weathering is determined as specified in ASTM G 53-84, in which a time cycle of 4 hours UV/4 h CON (condensation period) is maintained for 1500 hours at a test temperature of 50°C. The decorative surfaces exhibit a low surface gloss corresponding to a reflectometer value of 20-22, 60° angle of incidence or to a reflector value 44-45, 85° angle of incidence (DIN After weathering, the panel exhibits no 67,530). discoloration or change in gloss.

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The viscous, radiation-polymerizable, dyestuff-containing liquid 2 of Example 1 is applied, as described in Example 1, to a sulfate kraft paper 1 containing precured phenolformaldehyde resin, and is crosslinked in an essentially homogeneous manner by means of electron beams. The dose absorbed is 5 to 10 KGy. A further layer 6 composed of transparent - i.e., dyestuff-free - radiation-polymerizable liquid which,

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apart from the dyestuff, contains the same compounds as the layer first applied is applied as shown in Fig. 2, by means of rollers or rotary screen printing, to the surface of the paper on which the decorative layer 5 of synthetic resin polymerized by radiation is located. This layer 6 forms a continuous film having a layer thickness of about 20 /um. Immediately after the application of this second layer, a biaxially oriented polyethylene terphthalate single-ply film 3 which has been delustered by stretching is placed on the wet layer 6. Curing is carried out analogously to Example 1 by means of electron beams. The dose of radiation absorbed is 60 KGy. After the plastics film 3 has been removed, the paper 1 containing the external layer 6 of synthetic resin copolymerized by radiation is laid in each case on the outer face of a stack 13 composed of 50 superposed sheets of paper. The sheets of paper have previously been impregnated with a heat-curable phenol-formaldehyde resin and the resin has been partly cured. The bundle of layers is compressed at 150°C and 80 bar for 10 minutes in a press between two sheets of

It has the following composition:

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Transparent layer 6 (synthetic resin polymerized by radiation) as the outermost layer,

Decorative layer 5 (synthetic resin containing organic dyestuff pigments and polymerized by radiation), both layers on a previously impregnated paper layer 1,

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metal.

50 paper webs (impregnated with a phenolformaldehyde resin) as the core layer 13,

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Decorative layer 5 (synthetic resin containing organic dyestuff pigments and polymerized by radiation), and

Transparent layer 6 (synthetic resin polymerized by radiation) as the outermost layer, both layers on a previously impregnated paper layer 1.

The resulting decorative panel 14 has a scratch resistance higher than 2.0 N (DIN 53,799, part 10 10). Its surface is not attacked by concentrated mineral acid during a treatment time of 6 hours. The fastness to light of this panel is given the rating 8 (DIN 54,004). It has a surface gloss corresponding to a reflectometer value of 22-24, 60° angle of incidence, 15 and a reflectometer value of 44-45, 85° angle of incidence (DIN 67,530).

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<u>Example 3</u>

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The pasty, dyestuff-containing liquid 2, composed of radiation-polymerizable compounds, of Example 1 is applied to a sulfate kraft paper 1 which has first been impregnated with a heat-curable phenolformaldehyde resin (amount of resin applied 70%) of which the resin has been partially cured, and a continuous film (layer thickness about 80 /um) is formed (cf. Fig. 3).

A transparent layer 6 composed of a pasty, dyestuff-free liquid composed of the same radiationpolymerizable compounds is applied (layer thickness about 20 to 40 /um) to a delustered polypropylene plastics film 3. The paper 1 and the plastics film 3 are then placed one on top of the other in sheet or web

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form, so that the two liquid layers 2 and 6 come into contact with one another. In so doing, care must be taken that inclusions of air are not formed. The polymerizable compounds are crosslinked by means of electron beams which impinge on the liquid layers 2 and 6 through the plastics film 3. The dose absorbed is 60 kGy. After the plastics film 3 has been removed, the paper 1 containing the polymerized surface layer 2 and 6 is processed further to give a decorative panel 14 by compression under heat together with a paper stack 13, as described in Example 2.

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The layers of synthetic resin polymerized by radiation in the examples still exhibit relatively low values of scratch resistance within the range from about 0.7 to 0.9 Newtons before the compression under heat. A substantially higher surface hardness in the panel is only obtained, surprisingly, after the layer of synthetic resin polymerized by radiation has been subjected, in accordance with the invention, to compression under heat.

The reflectometer value measured with a Dr. Lange type RB reflectometer as specified in DIN 67,530 is 45 to 47, 20° angle of incidence, and about 83, 60° angle of incidence, if the delustered plastics film is absent when curing with electron beams is carried out. It falls to values of 37 to 41, 20° angle of incidence, and about 79, 60° angle of incidence, if texturized compression elements which impart to the outermost layer of the panel, for example, a surface structure similar to that of orange peel are used in the compression under heat. Even lower reflectometer values of about 30 to 36, 20° angle of incidence, and about 75, 60° angle of incidence, can be achieved by adding delustering agents to the surface layer. The particularly low gloss values according to the

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invention can, however, only be achieved by means of the special measures used in the polymerization of the synthetic resin layer. The claims defining the invention are as follows:
 A decorative panel, consisting of:

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a core layer having the substrate function of the decorative panel;

an outermost layer applied over said core layer, said outermost layer containing the radiation-polymerized product of a radiation-polymerizing mixture, said mixture containing

 (i) a radiation-polymerizable component selected from the group consisting of an epoxy-acrylate or -methacrylate oligomer, a silicone-acrylate or -methacrylate oligomer, a polyester acrylate or -methacrylate oligomer or a urethane acrylate or -methacrylate oligomer,

(ii) a diacrylate or triacrylate or -methacrylate monomer, wherein said radiation-polymerized outermost layer is thermopressed under conditions of elevated temperature and pressure such that a scratch resistance of at least 1.5 Newtons is imparted to said outermost layer and having a reflectometer value no greater than 50 at an angle of incidence of 85°; and

decoration means for forming a decoration covering said core layer.

2. A decorative panel as claimed in Claim 1, wherein said outermost layer is scratch resistant at a scratch loading of from 2 to 7 Newtons.

3. A decorative panel as claimed in Claim 1, wherein said synthetic resin comprises an epoxyacrylate or methacrylate oligomer, a siliconeacrylate or methacrylate oligomer, a polyester acrylate or methacrylate oligomer or a urethane acrylate or methacrylate oligomer.

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4. A decorative panel as claimed in Claim 3, wherein said outermost layer includes mono-, di-, tri-, tetra-, penta-, or hexaacrylate or -methacrylate polymerized with said synthetic resin.

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5. A decorative panel as claimed in Claim 4, wherein said outermost layer includes a

di- or triacrylate or methacrylate.



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6. A decorative panel as claimed in Claim
5, wherein said synthetic resin comprises an epoxy3 acrylate oligomer, a silicone-acrylate oligomer
or a polyester-acrylate oligomer.

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7. A decorative panel as claimed in Claim 4, wherein said synthetic resin comprises said aliphatic urethane acrylate oligomer which has been polymerized by radiation together with a diacrylate or triacrylate.

8. A decorative panel as claimed in Claim l, wherein said decorative panel includes an underlay between said core and said outermost layer, said underlay comprising a pigmented or non-pigmented 5 nonwoven fabric or paper and containing a heat curable aminoplast or phenoplast resin.

9. A decorative panel as claimed in Claim l, wherein said decoration means is incorporated in said outermost layer.

10. A decorative panel as claimed in Claim 9, wherein said core layer comprises wood, a plurality of paper layers, plastic or metal, and wherein a layer of paper is positioned between said core layer and said outermost layer.

II. A decorative panel as claimed in Claim l, wherein said decorative panel includes at least one additional layer between said core layer and said outermost layer, said decoration means being incorporated in said additional layer, and wherein said outermost layer is clear.

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12. A decorative panel as claimed in Claim 1, wherein said decoration means comprises a decorative layer between said core layer and said outermost layer, said decorative layer comprising a dyed or printed plastic film or paper.

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13. A decorative panel as claimed in Claim1, wherein a separate outermost layer is applied on3 opposite sides of said core layer.

14. A process for the production of a decorative panel as defined in claim 1, comprising:

applying a liquid synthetic resin layer to a substrate;

applying a film having a surface roughness over said resin layer;

transferring said surface roughness to said resin layer;

polymerizing said resin layer by 10 exposure to radiation; and

compressing said resin layer and said substrate at an elevated temperature.

15. A process as claimed in Claim 14,including the step of removing said film from saidresin layer prior to said compressing step.

16. A process as claimed in Claim 14,2 wherein said resin layer includes decoration means.

17. A process as claimed in Claim 14, wherein said substrate comprises a paper or a decorative paper layer, and wherein said compressing step includes compressing said resin layer and said paper or decorative paper layer to a core layer.

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18. A process as claimed in Claim 17, wherein said compressing step includes compressing a separate resin layer and paper or decorative paper layer on opposite sides of said core layer, said paper or decorative paper layers being adjacent said core layer.

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19. A process as claimed in Claim 14, wherein said compressing step includes compressing said resin layer at a temperature of from 80 to 220°C and at a pressure of from 5 to 100 bar.

20. A process for the production of a decorative panel as defined in Claim 1 comprising:

applying a first layer containing decoration means to a substrate;

applying a second layer over said first layer, said second layer being a synthetic resin;

applying a film having a surface roughness over said second layer;

transferring said surface roughness to said second layer; polymerizing said resin layer by exposure to radiation;

and

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compressing said first and second layers and said substrate at an elevated temperature.

21. A process as claimed in Claim 20, wherein said first layer is a synthetic resin and is partially cured on said substrate prior to application of said second layer.

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22. A process as claimed in Claim 20, wherein said film is applied over said second layer prior to applying said second layer over said first layer.

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23. A process as claimed in Claim 20, wherein said substrate layer comprises paper, and wherein said compressing step includes compressing said paper layer and said first and second layers onto a core layer.

24. A process as claimed in Claim 23, wherein said compressing step includes compressing separate paper layers and first and second layers on opposite sides of said core layer, said paper layers being adjacent said core layer.

25. A process as claimed in Claim 20, wherein said compressing step includes compressing at least said second layer at a temperature of from about 80 to about 220°C and at a pressure of from about 5 to about 100 bar.

DATED this 17th day of January, 1990.

<u>HOECHST AKTIENGESELLSCHAFT</u> By Its Patent Attorneys <u>ARTHUR S. CAVE & CO.</u>

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