(54) Titre : SYSTEME DE RESINE HYBRIDE (AQUEUX) A DEUX CONSTITUANTS, SON PROCEDE DE PRODUCTION ET SON UTILISATION

(55) Title: DUAL COMPONENT (AQUEOUS) HYBRID REACTIVE RESIN SYSTEM, METHOD FOR PRODUCTION AND USE THEREOF

(57) Abrégé/Abstract:
A dual component (aqueous) hybrid reactive resin system with improved working properties and improved quality profile is disclosed, obtained by a) the production of an epoxy-functional (aqueous) binding agent component (I) with an epoxide equivalent of 100 to 12500 g/eq, an average molecular mass of 200 to 25,000 Daltons and a viscosity of 1,000 to 150,000 mPa s (20 °C, Brookfield) and b) the production of a (latent) amino-functional hardener component (II). Said dual component (aqueous) hybrid reactive resin system can be used with excellent results in construction or industrial applications for production of chemical-, heat- and wear-resistant coating systems which are mechanically strong and easy to clean.
(19) Weltorganisation für geistiges Eigentum
Internationales Büro

(43) Internationales Veröffentlichungsdatum
18. Oktober 2007 (18.10.2007)

(51) Internationale Patentklassifikation:
C08G 18/28 (2006.01) C08G 18/42 (2006.01)


(22) Internationales Anmeldedatum:
3. April 2007 (03.04.2007)

(25) Einreichungssprache: Deutsch

(26) Veröffentlichungssprache: Deutsch

(30) Angaben zur Priorität:
10 2006 015 774.5 4. April 2006 (04.04.2006) DE

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(81) Bestimmungsstaaten (soweit nicht anders angegeben, für jede verfügbare nationale Schutzrechtsart): AE, AG, AL,


Erklärung gemäß Regel 4.17:
— Erfindererklärung (Regel 4.17 Ziffer iv)

Veröffentlicht:
— mit internationalen Recherchebericht
— vor Ablauf der für Änderungen der Ansprüche geltenden Frist, Veröffentlichung wird wiederholt, falls Änderungen eintreffen

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(54) Titel: DUAL COMPONENT (AQUEOUS) HYBRID REACTIVE RESIN SYSTEM, METHOD FOR PRODUCTION AND USE THEREOF

(54) Bezeichnung: ZWEIkomponentiges (wässriges) HYBRIDREAKTIVARZTSYSTEM, VERFAHREN ZU SEINER HERSTELLUNG SOWIE DIESSEN VERWENDUNG

(57) Abstract: A dual component (aqueous) hybrid reactive resin system with improved working properties and improved quality profile is disclosed, obtained by a) the production of an epoxy-functional (aqueous) binding agent component (I) with an epoxide equivalent of 100 to 12500 g/eq, an average molecular mass of 200 to 25,000 Daltons and a viscosity of 1,000 to 150,000 mPa·s (20 °C, Brookfield) and b) the production of a (latent) amino-functional hardener component (II). Said dual component (aqueous) hybrid reactive resin system can be used with excellent results in construction or industrial applications for production of chemical-, heat- and wear-resistant coating systems which are mechanically strong and easy to clean.

(57) Zusammenfassung: Es wird ein zweikomponentiges (wässriges) Hybridreaktivarztsystem mit verbesserten Verarbeitungsgegenschaften und verbessertem Eigenschaftsprofil beschrieben, welches erhärtet ist durch a) die Herstellung einer epoxyfunktionalen (wässrigen) Bindemittel-Komponente (I) mit einem Epoxid-Equivalent von 100 bis 12500 g/eq, einer mittleren Molekularmasse von 200 bis 25000 Dalton und einer Viskosität von 1000 bis 150000 mPa·s (20 °C, Brookfield) sowie durch b) die Herstellung einer (latent) aminofunktionalen Härterkomponente (II). Das erfindungsgemäße zweikomponentige (wässrige) Hybridreaktivarztsystem kann hervorragend im Bau- oder Industriebereich zur Herstellung von chemikalienbeständigen, hitzebeständigen, abriebbeständigen, mechanisch hoch belastbaren und einfach zu reinigenden Beschichtungssystemen verwendet werden.
Dual component (aqueous) hybrid reactive resin system, method for production and use thereof

Description

The present invention relates to a two-component (aqueous) hybrid reactive resin system having improved processing properties and improved property profile, process for its preparation and the use thereof in the construction sector or industrial sector.

Owing to their property profile, epoxy-modified polyurethanes are used especially in the area of adhesives and sealants, potting and coatings.

For use as adhesives and sealants, polyetherpolyols or polyesterpolyols are of course predominantly employed. "Synthesis and characterization of cryogenic adhesives based on epoxy-modified polyurethane resin systems" (Polymer International (1994), 35(4), 361-70) reports in detail on the synthesis of epoxy-modified polyurethanes. NCO-terminated polyurethane prepolymer are converted into epoxy-modified polyurethanes by reaction with glycidol. The property profiles of polyurethanes modified in this manner and based on PPG flexible segments of different molecular weight were emphasized with regard to mechanical properties, and adhesion at room temperature and low temperature in comparison with conventional, unmodified polyurethanes.

Epoxy-modified polyurethanes are distinguished not only by an improved property profile in comparison with conventional polyurethanes but they lead to improved properties even in mixtures with epoxy resins. Both the synthesis of epoxy-modified polyurethanes and the properties thereof in mixtures with epoxy resins with regard to adhesion, impact strength and fracture behaviour are described in Journal of Applied Polymer Science (1994), 52(8), 1137-51. Preliminary reaction of the synthesized polyurethanes with various curing agents gives systems which are distinguished by a lower degree of phase separation and improved mechanical properties.
A use of epoxy-modified polyurethanes for the finishing of textiles is described in US 3 814 578. Glycidol-modified polyurethanes, prepared by reaction of NCO-terminated polyurethane prepolymer with glycaldol, are used here. An application on wool and curing under acidic conditions leads to reduced shrinkage.

Further uses of epoxy-modified polyurethanes are explained, for example, in the following patents: Adhesives and laminated films (DE 3205733 A1), Polyurethanes (BE 620026) and Thermoplastic block polyamide-polyurethanes (DE 3504805 A1).

Common to all is the use of polyurethanes based on flexible polyols, such as, for example, polyethylene glycols, PPG, polybutadienes, polyesters, etc. A disadvantage of these materials is in particular poor resistance to chemicals and heat resistance, abrasion behaviour and mechanical load-bearing capacity. Uses as cementitious systems are not described.

Three- or four-component PCC coating systems known from the prior art and commercially available (e.g. UCrete®, Degussa Construction Chemicals GmbH) have, despite good resistance to chemicals, heat and abrasion, high mechanical load-bearing capacity and easy cleaning, a number of disadvantages in processing, in particular:

- application only by trained processors
- too short a processing time (about 10 min)
- too short an open time (about 20 min)
- excessively long curing (about 24 h)
- limited formulative range
- number of components
- VOC content

It was therefore the object of the present invention to develop a two-component (aqueous) hybrid reactive resin system having improved processing properties
and improved property profile for the preparation of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, which hybrid reactive resin system does not have said disadvantages of the prior art but has good performance characteristics and at the same time can be prepared taking into account ecological, economic and physiological aspects.

This object was achieved, according to the invention, by the provision of a two-component (aqueous) hybrid reactive resin system having improved processing properties and improved property profile, obtainable by

a) the preparation of an epoxyfunctional (aqueous) binder component (I) having an epoxide equivalent of 100 to 12 500 g/eq, an average molecular mass of 200 to 25 000 dalton and a viscosity of 1000 to 150 000 mPa·s (20°C, Brookfield),

a1) 5 to 300 parts by weight of a functionalized low molecular weight polyl component (A)(i), consisting of a hydroxyfunctional epoxyalcohol and/or glycidyl ether having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more epoxide group(s) substantially inert to isocyanate groups, having an epoxide equivalent of 100 to 500 g/eq and a molecular mass of 50 to 1000 dalton,

0 to 300 parts by weight of a functionalized higher molecular weight (polymeric) polyl component (A)(ii) having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more epoxide group(s) substantially inert to isocyanate groups, having an epoxide equivalent of 130 to 3000 g/eq and a molecular mass of 250 to 2500 dalton, being allowed to react with

5 to 500 parts by weight of a polyisocyanate component (B), consisting of at least one diisocyanate, polyisocyanate, polyisocyanate
derivative or polyisocyanate homologue having two or more (cyclo)aliphatic and/or aromatic isocyanate groups and a molecular mass of 100 to 2500 dalton, optionally in the presence of 0.01 to 0.5 part by weight of a catalyst component (K)(i) customary for polyaddition reactions with polyisocyanates, the mixture of the components (A)(i) and (B) being reacted either simultaneously or stepwise with the component (A)(ii), and optionally

0 to 200 parts by weight of a low molecular weight polyol component (A)(iii) having one or more hydroxyl group(s) reactive towards isocyanate groups and a molecular weight of 50 to 500 dalton,

0 to 500 parts by weight of a functionalized low molecular weight polyol component (A)(iv) having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more carboxyl and/or phosphonate and/or sulfonate group(s) inert to isocyanate groups and/or polyalkylene oxide group(s) and/or perfluoroalkyl group(s) and having a molecular mass of 50 to 2500 dalton, and

0 to 800 parts by weight of a higher molecular weight (polymeric) polyol component (A)(v) having one or more hydroxyl groups reactive towards isocyanate groups and a molecular mass of from 500 to 5000 dalton,

0 to 600 parts by weight of a reactive diluent component (C), consisting of at least one (aqueous) epoxy resin having one or more epoxide group(s) substantially inert to isocyanate groups, an epoxide equivalent of 130 to 400 g/eq and a molecular mass of 50 to 1000 dalton, and

0 to 50 parts by weight of a coalescence auxiliary component (D),

5 to 900 parts by weight of a formulation component (F)(i), consisting of reactive and/or inert fillers, pigments, carrier materials,
nanomaterials, nanocomposites, other additives, plasticizers, solvents and water

being added to the reaction mixture and

a2) optionally the prepolymer from stage a1) being emulsified or dispersed in 0 to 900 parts by weight of water and optionally the formulation component (F)(i) being added, and by

b) the preparation of a (latently) aminofunctional curing component (II),

10 to 900 parts by weight of a (polymeric) polyamine component (E), consisting of one or more (polymeric) polyamines having one or more (cyclo)aliphatic and/or aromatic primary and/or secondary amino group(s) reactive towards epoxide groups and optionally one or more hydroxyl group(s) and having a molecular mass of 60 to 5000 dalton, in the form of pure (polymeric) polyamines, polyaspartic acid esters, latent curing agents or reactive diluents based on aldimines and/or ketimines and/or enamines and/or oxazolidines, latent curing agents free of cleavage products and based on azetidines and/or diazepines and/or ammonium salts, commercially available liquid amine curing formulations or suitable combinations thereof,

10 to 900 parts by weight of a formulation component (F)(ii), consisting of reactive and/or inert fillers, pigments, carrier materials, nanomaterials, other additives, plasticizers, solvents and water, and

0.01 to 0.5 part by weight of an accelerator component (K)(ii) customary for polyaddition reactions with epoxy resins

being combined.
Surprisingly, it was found that, by using polyurethane-based two-component (aqueous) hybrid reactive resin systems with an epoxide/amine curing mechanism, not only are coating systems having substantially improved processing properties obtainable but these moreover have an improved property profile. In addition, it was not foreseeable that the viscosity of the epoxy-functional (aqueous) binder component would be very low.

In contrast to the three- or four-component PCC coating system (UCrete®) known from the prior art and containing a binder component, an isocyanate component, a cement-containing powder component and optionally also a pigment component, only two components are required in the case of the system according to the invention, with the result that mixing errors in the application can be avoided. The formulation constituents (powder component) can be directly integrated in the binder component and/or in the curing component. Since, in comparison with the prior art, no isocyanate/water reaction occurs and hence no carbon dioxide formation occurs, the use of calcium oxide and/or calcium hydroxide in the formulation component can be dispensed with in comparison with the prior art. For avoiding VOC emissions, the curing component can also be immobilized within the formulation component.

As suitable functionalized low molecular weight polyol component (A)(i), it is possible to use, for example, glycidol, glyceryl diglycidyl ether, (cyclo)aliphatic and/or aromatic polyols partly etherified with epichlorohydrin, such as butane-1,4-diol, p-tert-butylphenol, 1,4-cyclohexanediol, ethylene glycol, n-dodecanol, 2-ethylhexanol, glycerol and polyglycerol, hexane-1,6-diol, hydrogenated bisphenol A, hydrogenated bisphenol F, 2-methylpropane-1,3-diol, o-cresol, neopentylglycol, pentaerythritol, polyethylene glycols, polypropylene glycols, polyalkylene glycols, propane-1,2(3)-diol, n-tetradecanol, trimethylolpropane or hydroxyfunctional mono- and polyfunctional glycidyl ethers, epoxidized unsaturated fatty alcohols, reaction products of aliphatic and/or aromatic diepoxides and aliphatic and/or aromatic secondary monoamines, the reaction preferably being carried out in the molar ratio 1:1, reaction products of aliphatic and/or aromatic diepoxides and aliphatic and/or aromatic primary monoamines, the reaction preferably being carried out in the
molar ratio 2:1, reaction products of aliphatic and/or aromatic diepoxides and (un)saturated fatty acids and/or fatty alcohols and/or fatty amines, the reaction preferably being carried out in the molar ratio 1:1, or suitable combinations thereof. Glycidol and/or glyceryl diglycidyl ether and/or (cyclo)aliphatic and/or aromatic polyols partly etherified with epichlorohydrin or hydroxyfunctional mono- and polyfunctional glycidyl ethers are preferably used.

As suitable functionalized higher molecular weight (polymeric) polyol component (A)(ii), it is possible to use, for example, epoxidized and (partly) ring-opened (un)saturated triglycerides, dimer fatty acid diols, oleochemical polyols, the commercial products Sovermol® 45, 100, 320, 650 NS, 750, 760, 805, 810, 815, 818, 819, 820, 850, 860, 908, 912 Pearls, 920, 1005, 1012, 1014, 1052, 1055, 1058, 1059, 1066, 1068, 1080, 1083, 1090, 1095, 1102, 1106, 1111, 9155 and Speziol® C 10-2, C 18-2, C 36-2 from Cognis Deutschland GmbH & Co. KG, hydroxyfunctional epoxy resins or epoxy resin derivatives or suitable combinations thereof. (Un)saturated triglycerides which are epoxidized and (partly) ring-opened with alcohols are preferably used.

As suitable low molecular weight polyol component (A)(iii), it is possible to use, for example, 1,4-butanediol, 2-methyl-1,3-propanediol, 2,2-dimethyl-1,3-propanediol, 1,2-dihydroxyalkanediols having 5-50 carbon atoms of the general formula (I)

\[ C_nH_{2n+1} \text{CHOH-CH}_2\text{OH} \]

(\text{I})

where \( n = 3 \) to 48,

reaction products of alkylene 1-oxides of the general formula (II)

\[ C_nH_{2n+1} \text{-CH-CH}_2 \]

(\text{II})
where \( n = 3 \) to 48

with N-methylethanolamine or ethanolamine or diethanolamine or other compounds having a primary or secondary amino group and one or more hydroxyl group(s),

\( \alpha,\omega \)-dihydroxyalkanediols having 5 to 50 carbon atoms of the general formula (III)

\[
\text{HO--C}_n\text{H}_{2n-1}\text{OH}
\]

(III)

where \( n = 3 \) to 50

or suitable combinations thereof.

As suitable functionalized low molecular weight polyol component (A)(iv), it is possible to use, for example,

(i) bishydroxyalkanecarboxylic acids, such as dimethylolpropionic acid,

and/or

(ii) dihydroxyfunctional reaction products of monofunctional alkyl/cycloalkyl/arylpolylalkylene glycols, diisocyanates and dialkanolamines

and/or

(iii) amino- and/or hydroxy- and/or mercaptofunctional fluoromodified macromonomers or telechelic structures having a polymer-bound fluorine content of 1 to 99\% by weight and a molecular mass of 100 to 10 000 dalton, containing, arranged intrachenally in the main chain and/or side chain and/or laterally and/or terminally, the structural elements of the general formula (IV)
\(-(\text{CF}_2\text{CF}_2)\text{n}_-\)  
(IV)

where \(n \geq 3\)

and/or of the general formula (V)

\(-(\text{CF}_2\text{CFR-O})\text{n}_-\)  
(V)

where \(n \geq 3\) and \(R = F, \text{CF}_3\),

having in each case one or more (cyclo)aliphatic and/or aromatic primary and/or secondary amino group(s) and/or hydroxyl group(s) and/or mercapto group(s), preferably dihydroxyfunctional reaction products of perfluoroalkyl alcohols, diisocyanates and dialkanolamines, or suitable combinations thereof.

As suitable higher molecular weight (polymeric) polyol component (A)(v), it was possible to use, for example, (hydrophobically modified) polyalkylene glycols, (un)saturated aliphatic and/or aromatic polyesters, polycaprolactones, polycarbonates, \(\alpha,\omega\)-polybutadiene polyols, \(\alpha,\omega\)-polymethacrylatediols, \(\alpha,\omega\)-polysulphidediols, \(\alpha,\omega\)-dihydroxyalkyldimethylsiloxanes, hydroxyfunctional epoxy resins, hydroxyfunctional ketone resins, alkyd resins, dimer fatty acid dialcohols, reaction products based on bisepoxides and (un)saturated fatty acids, further hydroxyfunctional macromonomers and telechelic structures, mono- and/or di- and/or triesters of glycerol and saturated and/or unsaturated and optionally hydroxyfunctional fatty acids having 1 to 30 carbon atoms and having a functionality of \(f_{OH} \geq 2\) or suitable combinations, such as blends or hybrid polymers thereof, or suitable combinations thereof.

As suitable polyisocyanate component (B), it is possible to use, for example, polyisocyanates, polyisocyanate derivatives or polyisocyanate homologues having two or more aliphatic and/or aromatic isocyanate groups of identical or different reactivity or suitable combinations thereof. In particular, the polyisocyanates sufficiently well known in polyurethane chemistry or
combinations thereof are suitable. As suitable aliphatic polyisocyanates, it is possible to use, for example, 1,6-diisocyanatohexane (HDI), 1-isocyanato-5-isocyanatomethyl-3,3,5-trimethylcyclohexane or isophorone diisocyanate (IPDI, commercial product VESTANAT® IPDI from Degussa AG), bis(4-isocyanatocyclohexyl)methane (H12MDI, commercial product VESTANAT® H12MDI from Degussa AG), 1,3-bis(1-isocyanato-1-methylene)benzene (m-TMXDI), 2,2,4-trimethyl-1,6-diisocyanatohexane or 2,4,4-trimethyl-1,6-diisocyanatohexane (TMDI, commercial product VESTANAT® TMDI from Degussa AG) or industrial isomer mixtures of the individual aliphatic polyisocyanates or suitable combinations thereof. As suitable aromatic polyisocyanates, it is possible to use, for example, 2,4-diisocyanatotoluene or toluene diisocyanate (TDI), bis(4-isocyanatophenyl)methane (MDI) and optionally higher homologues thereof (polymeric MDI) or industrial isomer mixtures of the individual aromatic polyisocyanates or suitable combinations thereof. Furthermore, the so-called "coating polyisocyanates" based on bis(4-isocyanatocyclohexyl)methane (H12MDI), 1,6-diisocyanatohexane (HDI), 1-isocyanato-5-isocyanatomethyl-3,3,5-trimethylcyclohexane (IPDI) or suitable combinations thereof are in principle also suitable. The term "coating polyisocyanates" designates those derivatives of these diisocyanates which have allophanate, biuret, carbodiimide, isocyanurate, oxadiazinetriene, uretdione or urethane groups and in which the residual content of monomeric diisocyanates was reduced to a minimum in accordance with the prior art. In addition, it is also possible to use modified polyisocyanates which are obtainable, for example, by hydrophilic modification of "coating polyisocyanates" with monohydroxyfunctional polyethylene glycols or aminosulphonic acids. For example, the commercial products VESTANAT® T 1890 E, VESTANAT® T 1890 L, VESTANAT® T 1890 M, VESTANAT® T 1890 SV, VESTANAT® T 1890/100 (polyisocyanates based on IPDI trimer), VESTANAT® HB 2640 MX, VESTANAT® HB 2640/100, VESTANAT® HB 2640/LV (polyisocyanates based on HDI biuret), VESTANAT® HT 2500 L, VESTANAT® HB 2500/100, VESTANAT® HB 2500/LV (polyisocyanates based on HDI isocyanurate) from Degussa AG or suitable combinations thereof can be used as suitable "coating polyisocyanates". Bis(4-isocyanatophenyl)methane (MDI) and higher homologues thereof (polymeric MDI) and derivatives and/or 2,4-toluene diisocyanate and/or 2,6-toluene diisocyanate and/or isophorone
diisocyanate or industrial isomer mixtures of the individual aliphatic and/or aromatic polyisocyanates and/or (hydrophilically modified) "coating polyisocyanates" having allophanate, biuret, carbodiimide, isocyanurate, oxadiazinetrione, uretidone or urethane groups and based on bis(4-isocyanatocyclohexyl)methane (H₄MDI), 1,6-diisocyanatohexane (HDI), 1-isocyanato-5-isocyanatomethyl-3,3,5-trimethylcyclohexane (IPDI) are preferably used.

As suitable reactive diluent component (C), it is possible to use, for example, (cyclo)aliphatic and/or aromatic polyols completely etherified with epichlorohydrin, such as butane-1,4-diol, p-tert-butylphenol, 1,4-cyclohexanedimethanol, ethylene glycol, n-dodecanol, 2-ethylhexanol, glycerol and polyglycerol, hexane-1,6-diol, hydrogenated bisphenol-A, hydrogenated bisphenol-F, 2-methylpropane-1,3-diol, o-cresol, neopentyl glycol, pentaerythritol, polyethylene glycols, polypropylene glycols, polyalkylene glycols, propane-1,2(3)-diols, n-tetradecanol, trimethylolpropane or mono- and polyfunctional glycidyl ethers, bisphenol-A diglycidyl ether and higher homologues thereof, bisphenol-F diglycidyl ether and higher homologues thereof, phenol novolak resins, epoxy resin dispersions, the commercial products Polyox® E 064, E 150, E 152, E 221, E 227, E 237, E 253, E 254, E 260, E 270, E 270/700, E 270/500, E 280, E 280/700, E 280/500, E 375, E 395, E 403, E 411, E 442, E 492, E 630 (epoxy resin (solvent-free)), E 2400/75, E 2401.80, E 1001x75 (epoxy resins (solvent-containing)), E 260 W, E 2500/60 W (epoxy resins (for aqueous systems)), R 3, R 6, R 7, R 9, R 11, R 12, R 14, R 16, R 17, R 18, R 19, R 20, R 24 (glycidyl ether) from UPPC AG or suitable combinations thereof, industrial products also containing (cyclo)aliphatic and/or aromatic polyols partly etherified with epichlorohydrin, according to (A)(i). (Cyclo)aliphatic and/or aromatic polyols completely etherified with epichlorohydrin or hydroxyfunctional mono- and polyfunctional glycidyl ethers are preferably used. For example, alcohols, such as, for example, benzyl alcohol, or a suitable combination thereof can be used as suitable extenders.

As suitable coalescence auxiliary component (D), it is possible to use, for example, low-boiling, aprotic solvents, such as acetone or propanone,
butanone, 4-methyl-2-pentanone, ethyl acetate, n-butyl acetate, or high-boiling aprotic solvents, such as N-methyl-2-pyrrolidone, N-ethyl-2-pyrrolidone, diethylene glycol dimethyl ether, dipropylene glycol dimethyl ether, ethylene glycol monoalkyl ether acetates, diethylene glycol monoalkyl ether acetates, dialkyl adipates, cyclic alkylene carbonates or suitable combinations thereof. High-boiling solvents, such as N-ethylpyrrolidone and/or N-methylpyrrolidone and/or dipropylene glycol dimethyl ether and/or dialkyl adipates and/or cyclic alkylene carbonates are preferably used.

As suitable (polymeric) polyamine component (E) it is possible to use, for example, polyamines having two or more aliphatic and/or aromatic, primary and/or secondary amino groups of identical or different reactivity or suitable combinations thereof. As suitable aliphatic polyamines, it is possible to use, for example, 1,3-pentanediamine (DAMP), 2-methylpentamethylenediamine (MPMDA), benzylationpropylamine (BAPA), bisaminomethylcyclohexane or 1,3-bis(aminomethyl)cyclohexane (1,3-BAC), cyclohexylaminepropylamine (NAPCHA), diaminocyclohexane or 1,2-diaminocyclohexane (DAC or DCH), diethylaminopropylamine (DEAPA), diethylenetriamine or 1,4,7-triazaheptane (DETA), dimethyl-PACM or bis(4,4'-amino-3,3'-methylcyclohexyl)ethane (DM-PACM), dipropylenetriamine or 1,5,9-triazanane, ethylenediamine or 1,2-diaminoethane (EDA), hexamethylenediamine or 1,8-diazaoctane (HMDA), isophoronediamine or 3-methylamino-3,5,5-trimethylaminocyclohexane (IPD or IPDA), methylpentamethylenediamine or 2-methyl-1,7-diazaheptane, N3-amine or 1,4,8-triazaoctane, N4-amine or 1,5,8,12-tetraazadecane, N-aminopropylhexamethane or 1-(2-aminoethyl)-1,4-diazacyclohexane (NAEP), N-aminopropylcyclohexylamine (NAPCH), p-aminocyclohexylmethane or bis(4,4'-aminocyclohexyl)methane (PACM), pentaethylenehexamine or 1,4,7,10,13,16-hexazaazadecane (PEHA), propylenediamine or 1,5-diazapentane (PDA), tetraethylpentamethane or 1,4,7,10,13-pentaazatridecane (TEPA), tricyclododecanediamine or 3(4),8(9)-bis(aminomethyl)tricyclo[5,2,1,0²,6]decane (TCD), triethylenetetramine or 1,4,7,10-tetraazadecane (TETA), trimethylhexamethylenediamine or 2,2,4-trimethyl-1,8-diazaoctane and/or 2,4,4-trimethyl-1,8-diazaoctane or suitable combinations thereof. As suitable aromatic polyamines, it is possible to use, for example, diaminodi-
phenylmethane or bis(4,4'-aminophenyl)methane (DDM), diaminodiphenyl sulphone or bis(4,4'-aminophenyl) sulphone (DDS), diethylaminodiphenylmethane (DEDMD), diethyltoluenediamine (DETDA), m-xylylenediamine or 1,3-bis(aminomethyl)benzene (mXDA) or suitable combinations thereof. As suitable aromatic polyoxyalkyleneamines, it is possible to use, for example, polyoxyethylenepolyamines, polyyxypropylenepolyamines, polyytetrahydrofuranpolyamines, other polyyxalkylenepolyamines based on any desired alkylene oxide or mixtures thereof (co, block, random), butanediol ether diamine or 1,14-diaza-5,10-dioxotetracane (BDA) or suitable combinations thereof. In addition, polyaminoamides, Mannich bases, epoxide adducts, such as EDA adduct, DETA adduct, type 100, type 115, type 125, type 140, type 250 (genamide), PAA adduct, the commercial products Polypox® IH 7001, IH 7002, IH 7003, IH 7004, H 013, H 014, H 015, H 016, H 030, H 038, H 043, H 043 S, H 043 L, H 051, H 060, H 100, H 129, H 147, H 160, H 205, H 206, H 229, H 244, H 262, H 269, H 276/90, H 300, H 300 S, H 300 SL, H 310, H 333, H 354, H 354 L, H 415, H 445, H 445 L, H 480, H 483, H 488, H 488 L, H 489, H 490, H 497, H 501, H 503, H 610, H 611 (epoxy resin curing agents (polyamines)), IH 7005W, IH 7006W, W 800, W 802, W 804, W 810, W 860 (epoxy resin curing agents (aqueous)), P 215x70, P 225, P 240, P 245, P 250, P 350, P 370, P 450, P 450 S, P 499, P 502 (epoxy resin curing agents (polyaminoamides/polyaminoimidazolidines)) from UPPC AG or suitable combinations thereof can be used. Ethylenediamine and/or liquid epoxy resin curing agents formulated ready for use and based on aliphatic and/or aromatic polyamines and/or polyamidoamines are preferably used.

The component (E) may be present in coated and/or microencapsulated and/or carrier-fixed and/or hydrophilized and/or solvent-containing form and optionally may have sustained-release properties.

As suitable formulation components (F)(i) and (F)(ii), it is possible to use, for example, reactive inorganic fillers selected from the group consisting of cement, calcium oxide, calcium hydroxide or calcium sulphate or suitable combinations thereof.
As suitable formulation components (F)(i) and (F)(ii), it was also possible to use, for example, (functionalized) inorganic and/or organic fillers and/or light fillers inert to water, (functionalized) inorganic and/or organic pigments, (functionalized) inorganic and/or organic carrier materials, (functionalized) inorganic and/or organic nanomaterials, (functionalized) inorganic and/or organic nanocomposites, inorganic and/or organic fibres, graphite, carbon black, carbon fibres, carbon nanotubes, metal fibres and metal powders, conductive organic polymers, redispersible polymer powders or superabsorbers and suitable combinations thereof.

As suitable formulation components (F)(i) or (F)(ii), it is also possible to use, for example, other additives selected from the group consisting of antifoaming agents, deaerators, lubricating and levelling additives, substrate wetting additives, wetting and dispersing additives, water repellants, rheology additives, coalescence auxiliaries, dulling agents, adhesion promoters, antifreezes, antioxidants, UV stabilizers, biocides and suitable combinations thereof.

As suitable formulation components (F)(i) or (F)(ii), it was furthermore possible to use, for example, plasticizers selected from the group consisting of dialkyl phthalate, dialkyl adipate, biodiesel, rapeseed oil methyl ester, fatty acid derivatives, triglyceride derivatives or suitable combinations thereof.

As suitable catalyst component (K)(i), it was possible to use, for example, dibutylin oxide, dibutylin dilaurate (DBTL), triethylamine, tin(II) octanoate, 1,4-diazaabicyclo[2,2,2]octane (DABCO), 1,4-diazaabicyclo[3,2,0]-5-nonene (DBN), 1,5-diazaabicyclo[5,4,0]-7-undecene (DBU), morpholine derivatives, such as, for example, JEFFCAT® amine catalysts or suitable combinations thereof.

As suitable accelerator component (K)(ii), it was possible to use, for example, benzylidimethylamine, 4-N,N-dimethylaminophenol, 2,4,6-tris(N,N-dimethylaminomethyl)phenol, 2-methylimidazole, 2-phenylimidazole, other suitable tertiary amines or suitable combinations thereof.
The present invention furthermore relates to a process for the preparation of the two-component (aqueous) hybrid reactive resin system according to the invention, characterized in that

a) an epoxyfunctional (aqueous) binder component (I) is prepared by

a₁) allowing the components (A)(i), (A)(ii) and (B) to react, optionally in the presence of the component (K)(i), the mixture of the components (A)(i) and (B) being reacted either simultaneously or stepwise with the component (A)(ii), and optionally the components (A)(iii), (A)(iv), (A)(v), (C) and (D) also being added to the reaction mixture, and

a₂) optionally the prepolymer from stage a₁) being emulsified or dispersed in water and optionally the formulation component (F)(i) being added, and

b) preparing a (latently) aminofunctional curing component (II) by combining the components (E), (F)(ii) and (K)(ii) in any desired sequence.

The stage a₁) is carried out with avoidance of epoxide/isocyanate secondary reactions (e.g. cycloaddition with formation of cyclic urethanes).

The metering of the components (A), (B), (C), (D), (F)(i), (K)(i) used in the stages a) and b) can be effected in any desired manner.

The NCO/OH equivalent ratio of the components (A) and (B) in stage a) is preferably adjusted to 1.2 to 2.5, in particular to 1.3 to 2.0.

The stage a₁) is carried out at a preferred temperature of 40 to 90°C, in particular at 65 to 85°C.

The stage a₂) is carried out at a preferred temperature of 30 to 60°C, in particular at 40 to 50°C.
The stage b) is carried out at a preferred temperature of 10 to 40°C, in particular at 20 to 30°C.

According to a preferred embodiment, an epoxyfunctional binder component (I) which is self-emulsifying even without additional anionic and/or nonionic hydrophilization is used in stage a).

The solids content of the epoxyfunctional aqueous binder component (I) consisting of the components (A), (B) and (C) in stage a) is preferably adjusted to 10 to 100% by weight, in particular to 25 to 75% by weight.

The solids content of the two-component coating system consisting of the components (I) and (II) is preferably adjusted to 10 to 100% by weight, in particular to 25 to 75% by weight.

The present invention furthermore relates to the use of the two-component (aqueous) hybrid reactive resin system according to the invention for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for mineral and nonmineral surfaces based on concrete, cement, lime, gypsum, anhydrite, geopolymers, glass, wood and wood-based materials, composite materials, artificial and natural stone, plastic and glass fibre-reinforced plastic (GRP), metal and polymers.

The two-component (aqueous) hybrid reactive resin system according to the invention is suitable in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant, and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for the applications

- antigraffiti coatings
- antisoiling coatings
- seals
- antislip coverings
• dischargeable floor coating systems (ESD/AS)
• balcony coatings
• easy-to-clean coatings
• levelling and priming of concrete
• fresh concrete coatings
• floor coatings
• garage coatings
• water protection coating systems according to § 19 WHG
• high-bay warehouse coatings according to DIN 15185
• parking floor coatings
• PCC coating systems
• pipeline coatings
• crack-bridging coating systems
• hopper coatings
• sport floor covering systems
• wall coatings.

The two-component (aqueous) hybrid reactive resin system according to the invention is suitable in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for the following fields of use

• wastewater treatment
• chemical industry
• printing industry
• disposal
• beverage industry
• commercial kitchens and restaurants
• hygiene applications
• cold halls and cold stores
• storage halls and warehouses
• agricultural
• food industry
• paper industry
• pharmaceutical industry
• pipelines
• private households
• refineries
• clean-room areas (e.g. chip and wafer production)

The two-component (aqueous) hybrid reactive resin system according to the invention is suitable in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean and optionally consist of a primer and at least one primer coat which is not lightfast and is optionally sanded and optionally of a topcoat which is lightfast and optionally fluoromodified and optionally sanded.

The two-component (aqueous) hybrid reactive resin system according to the invention can be used in any desired combination with conventional three-component PCC coating systems (UCrete®) and/or aqueous and/or reactive polyurethane coating systems and/or aqueous and/or reactive epoxy resin coating systems in the applications

• repair
• retopping
• mixed system structure.

The two-component (aqueous) hybrid reactive resin system according to the invention can be used in the applications

• job-mix concrete
• concrete products (precast concrete parts, concrete ware, cast stones)
• poured-in-place concrete
• air-placed concrete
• ready-mixed concrete.
The epoxyfunctional (aqueous) binder component (I) and the (latently) amino-functional curing component (II) are mixed in the preferred epoxide/amino equivalent ratio of 0.8 to 1.2, in particular 0.9 to 1.1, to give a two-component coating system.

The coating system is preferably applied in coats having a total thickness of 0.1 to 50 mm to elastic or rigid substrates, it being used in particular in an amount of 0.1 to 10.0 kg per m² of the area to be coated and per operation.

The coating system can be applied here horizontally and vertically and without a primer (and without bubble formation) to (moist) fresh concrete.

The hybrid reactive resin system according to the invention can be used in particular for crack-bridging and cavity-filling coatings.

The application of the coating system is effected by the methods known from painting and coating technology, such as, for example, flooding, pouring, application with a doctor blade, roller-coating with a soft roller, spraying, brushing, immersion or roller-coating with a hard roller.

The following examples are intended to illustrate the invention in more detail.
Examples

Example 1

Synthesis of an epoxyfunctional hybrid resin (1)

In a 250 ml three-necked round-bottomed flask with a KPG stirrer, internal thermometer and air condenser, a mixture of 45.00 g of IPDI (Degussa AG) and 14.62 g of glycidol was cooled to 17°C and DBTL was added as a catalyst. Thereafter, the reaction mixture was stirred at a temperature of 60°C until the theoretical NCO value of 14.23% by weight was reached. After addition of 45.21 g of Sovernol 818 (Cognis GmbH), stirring was effected for a further 4 h at 80°C until complete conversion of the NCO groups, and a resin viscosity of about 140 000 mPa·s was established by dilution with 34.99 g of Polypox R 18 (UPPC AG).

The product obtained was a homogeneous, slightly yellow resin having an epoxide value of 3.03 mol/kg.

For curing of the reactive resin, 8.19 g of the curing agent Polypox VH 01198/10 (UPPC AG) were added to 45.00 g of (1) and homogenization was effected. After a potlife of about 40 min, a Shore D hardness of about 71 resulted.
Example 2
Synthesis of an epoxyfunctional hybrid resin (2)

In a 500 ml three-necked round-bottomed flask with a KPG stirrer, internal thermometer and air condenser, a mixture of 134.99 g of Polypox R 18 (UPPC AG), 22.14 g of glycidol and 108.00 g of IPDI (Degussa AG) was heated to a temperature of 33°C with stirring and 0.15 g of DBTL was added as a catalyst. After the very strongly exothermic reaction had died down, the reaction mixture was stirred for a further 30 min at 80°C until a theoretical NCO value of 7.70% by weight had been reached. After addition of 108.51 g of Sovermol 818 (Cognis GmbH), stirring was effected for a further 4 h at 60°C for complete conversion of all NCO groups and a total proportion of reactive diluent of 45% by weight was established by dilution with 60.42 g of Polypox R 18 (UPPC AG).

The product obtained was a homogeneous, slightly yellow resin having an epoxide value of 276.73 g/eq.

For curing of the reactive resin, 3.79 g of the curing agent Polypox H 503 (UPPC AG) were added to 30.00 g of (2) and homogenization was effected. After curing for 48 h at 50°C, a casting having a Shore D hardness of 74 was obtained.
Example 3

Synthesis of an epoxyfunctional hybrid resin (3)

In a 250 ml three-necked round-bottomed flask with a KPG stirrer, internal thermometer and air condenser, a mixture of 45.00 g of IPDI (Degussa AG) and 14.62 g of glycidol was initially introduced and 0.15 g of DBTL was added as a catalyst while cooling with a waterbath. After the exothermic reaction had died down, the reaction mixture was stirred for a further 60 min at 60°C until the theoretical NCO value of 14.23% by weight was reached. After addition of 43.92 g of Desmophen VP LS 2328 (Bayer MaterialScience AG), stirring was effected for a further 3 h at 80°C for complete conversion of the NCO groups and, after dilution with 34.56 g of Polypox R 18 (UPPC AG), a reactive resin having a viscosity of 91 000 mPa·s and an epoxide value of 327.60 g/eq was obtained.

For curing of the resin, 8.24 g of the curing agent Polypox VH 01198/10 (UPPC AG) were added to 45.00 g of (3) and homogenization was effected. After a potlife of 45 min, a casting having a Shore D hardness of 67 was obtained.
Example 4

Synthesis of an epoxyfunctional hybrid resin (4)

In a 250 ml three-necked round-bottomed flask with a KPG stirrer, internal thermometer and air condenser, a mixture of 45.00 g of IPDI (Degussa AG) and 15.00 g of glycidol was initially introduced and 0.15 g of DBTL was added as a catalyst while cooling with a waterbath. After the exothermic reaction had died down, the reaction mixture was stirred for a further 30 min at 80°C until the theoretical NCO value of 14.14% by weight was reached. After addition of 99.01 g of Oxyester T 1136 (Degussa AG), stirring was effected for a further 3.5 h at 80°C for complete conversion of the NCO groups and a total proportion of reactive diluent of 25% by weight was established by dilution with 53.05 g of Polytopx R 18 (UPPC AG).

The product obtained was a homogeneous, slightly yellow resin having an epoxide value of 390.69 g/eq.

For curing of the resin, 6.91 g of the curing agent Polytopx VH 01198/10 (UPPC AG) were added to 45.00 g of (3) and homogenization was effected. After a potlife of about 90 min, a casting having a Shore D hardness of 26 was obtained.
Example 5

Synthesis of an epoxyfunctional hybrid resin (5)

In a 500 ml three-necked round-bottomed flask with a KPG stirrer, internal thermometer and air condenser, a mixture of 150.83 g of Polypox R 18 (UPPC AG), 24.60 g of glycidol and 120.00 g of IPDI was initially introduced and, after heating to 38°C, 0.15 g of DBTL was added as a catalyst. After the exothermic reaction had died down, stirring was effected for about 30 min at 80°C until a theoretical NCO value of 7.67% by weight was reached. After addition of 264.03 g of Oxyester T 1136 (Degussa AG), stirring was effected for a further 6 h at 80°C for complete conversion of the NCO groups.

The product obtained was a homogeneous, slightly yellow resin having an epoxide value of 426.37 g/eq and a viscosity of 50 000 mPa·s. The total proportion of reactive diluent is 26.95% by weight.

For curing of the resin, 2.46 g of the curing agent Polypox H 503 (UPPC AG) were added to 30.00 g of (3) and homogenization was effected. After curing for 24 h at 70°C, a casting having a Shore D hardness of 36 was obtained.
Mechanical properties of various reactive resins

<table>
<thead>
<tr>
<th>Reactive resin</th>
<th>Reactive diluent [% by wt.]</th>
<th>Viscosity [mPa·s]</th>
<th>Shore D**</th>
<th>Elongation at break [%]**</th>
<th>Tensile strength [MPa]**</th>
</tr>
</thead>
<tbody>
<tr>
<td>(2)*</td>
<td>45</td>
<td>8800</td>
<td>67-74</td>
<td>34-38</td>
<td>20-23</td>
</tr>
<tr>
<td>(6)*</td>
<td>50</td>
<td>4000</td>
<td>66</td>
<td>35-49</td>
<td>15-20</td>
</tr>
<tr>
<td>(7)*</td>
<td>55</td>
<td>2500</td>
<td>60</td>
<td>45-57</td>
<td>15-22</td>
</tr>
<tr>
<td>(8)*</td>
<td>60</td>
<td>1200</td>
<td>52-57</td>
<td>48-55</td>
<td>16-24</td>
</tr>
<tr>
<td>(9)*</td>
<td>65</td>
<td>670</td>
<td>41</td>
<td>45-52</td>
<td>11-16</td>
</tr>
</tbody>
</table>

* The syntheses of the reactive resins (6)-(9) take place analogously to the synthesis of resin (2) but with an increased proportion of reactive diluent.

** Polypox H503 was used for curing all resins.
Example 6
Guide recipe for the formulation of epoxyfunctional hybrid reactive resins

<table>
<thead>
<tr>
<th>No.</th>
<th>Formulation constituents</th>
<th>% by weight</th>
<th>Weight taken (g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Reactive resin component (cf. Examples 1-5)</td>
<td>10.00</td>
<td>8.00</td>
</tr>
<tr>
<td>2</td>
<td>SR-POX 2500 (Maeder GmbH)</td>
<td>37.00</td>
<td>29.60</td>
</tr>
<tr>
<td>3</td>
<td>PERENOL E5 (Cognis GmbH)</td>
<td>1.00</td>
<td>0.80</td>
</tr>
<tr>
<td>4</td>
<td>TEXAPHOR P63 (Cognis GmbH)</td>
<td>1.00</td>
<td>0.80</td>
</tr>
<tr>
<td>5</td>
<td>PHOTOMER 4094 (Cognis GmbH)</td>
<td>7.50</td>
<td>6.00</td>
</tr>
<tr>
<td>6</td>
<td>TALKUM AT1</td>
<td>5.00</td>
<td>4.00</td>
</tr>
<tr>
<td>7</td>
<td>ALBAWHITE 60</td>
<td>27.00</td>
<td>21.60</td>
</tr>
<tr>
<td>8</td>
<td>RILANIT SPEZIAL MICRO W</td>
<td>1.50</td>
<td>1.20</td>
</tr>
<tr>
<td>9</td>
<td>HEUCOSIN-SPEZ. Light grey G 1039</td>
<td>10.00</td>
<td>8.00</td>
</tr>
<tr>
<td></td>
<td></td>
<td>100.00</td>
<td>80.00</td>
</tr>
</tbody>
</table>

No. 1-5 were initially introduced into a mixing beaker and homogenized. No. 6-9 were then weighed into a further beaker and added to the mixture of 1-5. The total mixture thus obtained was dispersed in a Speedmixer until a temperature of 50°C was reached (temperature monitoring). For deaeration, the paint thus obtained was allowed to stand overnight.
For curing, 5.84 g of the curing agent Polypox H 503 (UPPC AG) or 10.01 g of the curing agent Polypox VH 01198/10 (UPPC AG) were added in each case to 80.00 g of the formulated binder (epoxide content: 2.09 mol/kg) and homogenization was effected. After a potlife of about 3 h (Polypox H 503) or 1.5 h (Polypox VH 01198/10), Shore D hardnesses of 70-75 were obtained in each case.
Patent claims

1. Two-component (aqueous) hybrid reactive resin system having improved processing properties and improved property profile, obtainable by

a) the preparation of an epoxyfunctional (aqueous) binder component (I) having an epoxide equivalent of 100 to 12 500 g/eq, an average molecular mass of 200 to 25 000 dalton and a viscosity of 1000 to 150 000 mPa·s (20°C, Brookfield),

a₁) 5 to 300 parts by weight of a functionalized low molecular weight polyl component (A)(i), consisting of a hydroxyfunctional epoxy-alcohol and/or glycidyl ether having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more epoxide group(s) substantially inert to isocyanate groups, having an epoxide equivalent of 100 to 500 g/eq and a molecular mass of 50 to 1000 dalton,

0 to 300 parts by weight of a functionalized higher molecular weight (polymeric) polyl component (A)(ii) having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more epoxide group(s) substantially inert to isocyanate groups, having an epoxide equivalent of 130 to 3000 g/eq and a molecular mass of 250 to 2500 dalton, being allowed to react with

5 to 500 parts by weight of a polyisocyanate component (B), consisting of at least one diisocyanate, polyisocyanate, polyisocyanate derivative or polyisocyanate homologue having two or more (cyclo)aliphatic and/or aromatic isocyanate groups and a molecular mass of 100 to 2500 dalton, optionally in the presence of 0.01 to 0.5 part by weight of a catalyst component (K)(i) customary for polyaddition reactions with polyisocyanates, the mixture of the components (A)(i) and (B) being reacted either
simultaneously or stepwise with the component (A)(ii), and optionally

0 to 200 parts by weight of a low molecular weight polyol component (A)(iii) having one or more hydroxyl group(s) reactive towards isocyanate groups and a molecular weight of 50 to 500 dalton,

0 to 500 parts by weight of a functionalized low molecular weight polyol component (A)(iv) having one or more hydroxyl group(s) reactive towards isocyanate groups and one or more carboxyl and/or phosphonate and/or sulfonate group(s) inert to isocyanate groups and/or polyalkylene oxide group(s) and/or perfluoroalkyl group(s) and having a molecular mass of 50 to 2500 dalton, and

0 to 800 parts by weight of a higher molecular weight (polymeric) polyol component (A)(v) having one or more hydroxyl groups reactive towards isocyanate groups and a molecular mass of from 500 to 5000 dalton,

0 to 600 parts by weight of a reactive diluent component (C), consisting of at least one (aqueous) epoxy resin having one or more epoxide group(s) substantially inert to isocyanate groups, an epoxide equivalent of 130 to 400 g/eq and a molecular mass of 50 to 1000 dalton, and

0 to 50 parts by weight of a coalescence auxiliary component (D),

5 to 900 parts by weight of a formulation component (F)(i), consisting of reactive and/or inert fillers, pigments, carrier materials, nanomaterials, nanocomposites, other additives, plasticizers, solvents and water

being added to the reaction mixture and
a₂) optionally the prepolymer from stage a₁) being emulsified or
dispersed in 0 to 900 parts by weight of water and optionally the
formulation component (F)(i) being added,

and by

b) the preparation of a (latently) aminofunctional curing component (II),

10 to 900 parts by weight of a (polymeric) polyamine component (E),
consisting of one or more (polymeric) polyamines having one or more
(cyclo)aliphatic and/or aromatic primary and/or secondary amino
group(s) reactive towards epoxide groups and optionally one or more
hydroxyl group(s) and having a molecular mass of 60 to 5000 dalton,
in the form of pure (polymeric) polyamines, polyaspartic acid esters,
latent curing agents or reactive diluents based on aldimines and/or
ketimines and/or enamines and/or oxazolidines, latent curing agents
free of cleavage products and based on azetidines and/or diazepines
and/or ammonium salts, commercially available liquid amine curing
formulations or suitable combinations thereof,

10 to 900 parts by weight of a formulation component (F)(ii),
consisting of reactive and/or inert fillers, pigments, carrier materials,
nanomaterials, other additives, plasticizers, solvents and water, and

0.01 to 0.5 part by weight of an accelerator component (K)(ii)
customary for polyaddition reactions with epoxy resins

being combined.

2. Hybrid reactive resin system according to Claim 1, characterized in that
the component (A)(i) comprises glycidol and/or glyceryl diglycidyl ether
and/or (cyclo)aliphatic and/or aromatic polyols partly etherified with
epichlorohydrin or mono- and polyfunctional glycidyl ethers.
3. Hybrid reactive resin system according to either of Claim 1 or 2, characterized in that the component (A)(ii) comprises (un)saturated triglycerides epoxidized and (partly) ring-opened with alcohols.

4. Hybrid reactive resin system according to any of Claims 1 to 3, characterized in that the component (A)(iii) comprises 1,4-butanediol and/or 2-methyl-1,3-propanediol and/or 2,2-dimethyl-1,3-propanediol and/or 1,2-dihydroxyalkanediols having 5 to 50 carbon atoms of the general formula (I)

\[ C_{n}H_{2n+1} \text{-CHOH-CH}_{2}\text{OH} \]

(I)

where \( n = 3 \) to 48

and/or reaction products of alkylene 1-oxides of the general formula (II)

\[ C_{n}H_{2n+1} \text{-CH-CH}_{2} \]

(II)

where \( n = 3 \) to 48

with N-methylethanolamine or ethanolamine or diethanolamine or other compounds having a primary or secondary amino group and one or more hydroxyl group(s)

and/or \( \alpha,\omega \)-dihydroxyalkanediols having 5 to 50 carbon atoms of the general formula (III)

\[ \text{HO-C}_{n}H_{2n-1}\text{-OH} \]

(III)

where \( n = 3 \) to 50.

5. Hybrid reactive resin system according to any of Claims 1 to 4, characterized in that the component (A)(iv) comprises
(i) bishydroxyalkanecarboxylic acids, such as dimethylolpropionic acid and/or

(ii) dihydroxyfunctional reaction products of monofunctional alkyl/cycloalkyl/arylpolyalkylene glycols, diisocyanates and dialkanolamines and/or

(iii) amino- and/or hydroxy- and/or mercaptofunctional fluoromodified macromonomers or telechelic structures having a polymer-bound fluorine content of 1 to 99% by weight and a molecular mass of 100 to 10 000 dalton, containing, arranged intrachenally in the main chain and/or side chain and/or laterally and/or terminally, the structural elements of the general formula (IV)

$$-(CF_2-CF_2)_n-$$

(IV)

where \( n \geq 3 \)

and/or of the general formula V

$$-(CF_2-CFR-O)_n-$$

(V)

where \( n \geq 3 \) and \( R = F, CF_3 \),

having in each case one or more (cyclo)aliphatic and/or aromatic primary and/or secondary amino group(s) and/or hydroxyl group(s) and/or mercapto group(s), preferably dihydroxyfunctional reaction products of perfluoroalkyl alcohols, diisocynates and dialkanolamines.

6. Hybrid reactive resin system according to any of Claims 1 to 5, characterized in that the component (A)(v) comprises (hydrophobically
modified) polyalkylene glycols, (un)saturated aliphatic and/or aromatic polyesters, polycaprolactones, polycarbonates, α,ω-polybutadienepolyols, α,ω-polyacrylate diols, α,ω-polysulphidediols, α,ω-diethoxyalkylpolydimethylsiloxanes, hydroxyfunctional epoxy resins, hydroxyfunctional ketone resins, alkyd resins, dimer fatty acid dialcohols, reaction products based on bisepoxides and (un)saturated fatty acids, further hydroxyfunctional macromonomers and telechelic structures, mono- and/or di- and/or triesters of glycerol and saturated and/or unsaturated and optionally hydroxyfunctional fatty acids having 1 to 30 carbon atoms and having a functionality of $f_{OH} \geq 2$ or suitable combinations, such as blends or hybrid polymers thereof.

7. Hybrid reactive resin system according to any of Claims 1 to 6, characterized in that the component (B) comprises bis(4-isocyanatophenyl)methane (MDI) and higher homologues thereof (polymeric MDI) and derivatives and/or 2,4-toluene diisocyanate and/or 2,6-toluene diisocyanate and/or isophorone diisocyanate or industrial isomer mixtures of the individual aliphatic and/or aromatic polyisocyanates and/or (hydrophilically modified) "coating polyisocyanates" having allophanate, biuret, carbodiimide, isocyanurate, oxadiazinetrione, uretdione or urethane groups and based on bis(4-isocyanatocyclohexyl)methane (H₂₃MDI), 1,6-diisocyanatohexane (HDI), 1-isocyanato-5-isocyanatomethyl-3,3,5-trimethylcyclohexane (IPDI).

8. Hybrid reactive resin system according to any of Claims 1 to 7, characterized in that the component (C) comprises (cyclo)aliphatic and/or aromatic polyols completely etherified with epichlorohydrin or mono- and polyfunctional glycidyl ethers, industrial products also containing polyols partially etherified with epichlorohydrin or hydroxyfunctional mono- and polyfunctional glycidyl ethers.

9. Hybrid reactive resin system according to any of Claims 1 to 8, characterized in that the component (D) comprises high-boiling solvents,
such as N-ethylpyrrolidone and/or N-methylpyrrolidone and/or dipropylene glycol dimethyl ether and/or dialkyl adipates and/or cyclic alkylene carbonates.

10. Hybrid reactive resin system according to any of Claims 1 to 9, characterized in that the component (E) comprises ethylenediamine and/or liquid epoxy resin curing agents formulated ready for use and based on aliphatic and/or aromatic polyamines and/or polyamidoamines.

11. Hybrid reactive resin system according to any of Claims 1 to 10, characterized in that the component (E) is present in coated and/or microencapsulated and/or carrier-fixed and/or hydrophilized and/or solvent-containing form and optionally may have sustained-release properties.

12. Hybrid reactive resin system according to any of Claims 1 to 11, characterized in that the formulation components (F)(i) and (F)(ii) consist of reactive inorganic fillers selected from the group consisting of cement, calcium oxide, calcium hydroxide or calcium sulphate.

13. Hybrid reactive resin system according to any of Claims 1 to 12, characterized in that (functionalized) inorganic and/or organic fillers and/or light fillers inert to water, (functionalized) inorganic and/or organic pigments, (functionalized) inorganic and/or organic carrier materials, (functionalized) inorganic and/or organic nanomaterials, (functionalized) inorganic and/or organic nanocomposites, inorganic and/or organic fibres, graphite, carbon black, carbon fibres, carbon nanotubes, metal fibres and metal powders, conductive organic polymers, redispersible polymer powders or superabsorbers are used as formulation components (F)(i) or (F)(ii).

14. Hybrid reactive resin system according to any of Claims 1 to 13, characterized in that the formulation components (F)(i) or (F)(ii) consist of other additives selected from the group consisting of antifoaming agents,
deaerators, lubricating and levelling additives, substrate wetting additives, wetting and dispersing additives, water repellents, rheology additives, coalescence auxiliaries, dulling agents, adhesion promoters, antifreezes, antioxidants, UV stabilizers and biocides.

15. Hybrid reactive resin system according to any of Claims 1 to 14, characterized in that plasticizers selected from the group consisting of dialkyl phthalate, dialkyl adipate, biodiesel and rapeseed oil methyl ester are used as formulation components (F)(i) or (F)(ii).

16. Hybrid reactive resin system according to any of Claims 1 to 14, characterized in that dibutyltin oxide, dibutyltin dilaurate (DBTL), triethylamine, tin(II) octanoate, 1,4-diazabicyclo[2.2.2]octane (DABCO), 1,4-diazabicyclo[3.2.0]-5-nonene (DBN), 1,5-diazabicyclo[5.4.0]-7-undecene (DBU), morpholine derivatives, such as, for example, JEFFCAT® amine catalysts are used as component (K)(i).

17. Hybrid reactive resin system according to any of Claims 1 to 16, characterized in that the component (K)(ii) comprises benzylidimethylamine and/or 4-N,N-dimethylaminophenol and/or 2,4,6-tris(N,N-dimethylaminomethyl)phenol and/or 2-methylimidazole and/or 2-phenylimidazole and/or other suitable tertiary amines.

18. Process for the preparation of the two-component (aqueous) hybrid reactive resin system according to any of Claims 1 to 17, characterized in that

a) an epoxyfunctional (aqueous) binder component (I) is prepared by

a1) allowing the components (A)(i), (A)(ii) and (B) to react, optionally in the presence of the component (K)(i), the mixture of the components (A)(i) and (B) being reacted either simultaneously or stepwise with the component (A)(ii), and optionally the
components (A)(iii), (A)(iv), (A)(v), (C) and (D) also being added to the reaction mixture, and

a2) optionally the prepolymer from stage a1) being emulsified or dispersed in water and optionally the formulation component (F)(i) being added, and

b) preparing a (latently) aminofunctional curing component (II) by combining the components (E), (F)(ii) and (K)(ii) in any desired sequence.

19. Process according to Claim 18, characterized in that the metering of the components (A), (B), (C), (D), (F)(i), (K)(i) used in the stages a) and b) can be effected in any desired manner.

20. Process according to either of Claims 18 and 19, characterized in that the NCO/OH equivalent ratio of the components (A) and (B) in stage a) is adjusted to 1.2 to 2.5, preferably to 1.3 to 2.0.

21. Process according to any of Claims 18 to 20, characterized in that the stage a1) is carried out at a temperature of 40 to 90°C, preferably at 65 to 85°C.

22. Process according to any of Claims 18 to 21, characterized in that the stage a2) is carried out at a temperature of 30 to 60°C, preferably at 40 to 50°C.

23. Process according to any of Claims 18 to 22, characterized in that the stage b) is carried out at a temperature of 10 to 40°C, preferably at 20 to 30°C.

24. Process according to any of Claims 18 to 23, characterized in that an epoxyfunctional binder component (I) which is self-emulsifying without additional anionic and/or nonionic hydrophilization is used in stage a1).
25. Process according to any of Claims 18 to 24, characterized in that the solids content of the epoxyfunctional aqueous binder component (I) consisting of the components (A), (B) and (C) in stage a) is adjusted to 10 to 100% by weight, preferably 25 to 75% by weight.

26. Process according to any of Claims 16 to 25, characterized in that the solids content of the two-component coating system consisting of the components (I) and (II) is adjusted to 10 to 100% by weight, preferably to 25 to 75% by weight.

27. Use of the two-component (aqueous) hybrid reactive resin system according to any of Claims 1 to 17 in the construction sector or industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for mineral and nonmineral surfaces based on concrete, cement, lime, gypsum, anhydrite, geopolymers, glass, wood and wood-based materials, composite materials, artificial and natural stone, plastic and glass fibre-reinforced plastic (GRP), metal and polymers.

28. Use according to Claim 27 in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for the applications

- antigraffiti coatings
- antisoiling coatings
- seals
- antislip coverings
- dischargeable floor coating systems (ESD/AS)
- balcony coatings
- easy-to-clean coatings
- levelling and priming of concrete
- fresh concrete coatings
- floor coatings
- garage coatings
- water protection coating systems according to § 19 WHG
- high-bay warehouse coatings according to DIN 15185
- parking floor coatings
- PCC coating systems
- pipeline coatings
- crack-bridging coating systems
- hopper coatings
- sport floor covering systems
- wall coatings.

29. Use according to either of Claims 27 and 28 in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean, for the following fields of use

- wastewater treatment
- chemical industry
- printing industry
- disposal
- beverage industry
- commercial kitchens and restaurants
- hygiene applications
- cold halls and cold stores
- storage halls and warehouses
- agricultural
- food industry
- paper industry
- pharmaceutical industry
- pipelines
- private households
• refineries
• clean-room areas (e.g. chip and wafer production)

30. Use according to any of Claims 27 to 29 in the construction sector and industrial sector for the production of coating systems which are resistant to chemicals, heat-resistant and abrasion-resistant, have a high mechanical load-bearing capacity and are easy to clean and optionally consist of a primer and at least one primer coat which is not lightfast and is optionally sanded and optionally of a topcoat which is lightfast and is optionally fluoromodified and optionally sanded.

31. Use according to any of Claims 27 to 30 in any desired combination with conventional three-component PCC coating systems (UCrete®) and/or aqueous and/or reactive polyurethane coating systems and/or aqueous and/or reactive epoxy resin coating systems in the applications

• repair
• retopping
• mixed system structure.

32. Use according to any of Claims 27 to 31 in the applications

• job-mix concrete
• concrete products (precast concrete parts, concrete ware, cast stones)
• poured-in-place concrete
• air-placed concrete
• ready-mixed concrete.

33. Use according to any of Claims 27 to 32, characterized in that the epoxy-functional (aqueous) binder component (I) and the (latently) amino-functional curing component (II) are mixed in the epoxide/amino equivalent ratio 0.8 to 1.2, preferably 0.9 to 1.1, to give a two-component coating system.
34. Use according to any of Claims 27 to 33, characterized in that the coating system is applied in coats having a total thickness of 0.1 to 50 mm to elastic or rigid substrates.

35. Use according to any of Claims 27 to 34, characterized in that the coating system is used in an amount of from 0.1 to 10.0 kg per m² of the area to be coated and per operation.

36. Use according to any of Claims 27 to 35, characterized in that the coating system is applied horizontally and vertically.

37. Use according to any of Claims 27 to 36, characterized in that the coating system is applied without a primer to (moist) fresh concrete.

38. Use according to any of Claims 27 to 37, characterized in that the hybrid reactive resin system is used for crack-bridging and cavity-filling coatings.