



HU000031491T2

(19) **HU**(11) Lajstromszám: **E 031 491**(13) **T2****MAGYARORSZÁG**
Szellemi Tulajdon Nemzeti Hivatala**EURÓPAI SZABADALOM**
SZÖVEGÉNEK FORDÍTÁSA(21) Magyar ügyszám: **E 09 742109**(51) Int. Cl.: **C08J 11/08** (2006.01)(22) A bejelentés napja: **2009. 05. 07.**

(86) A nemzetközi (PCT) bejelentési szám:

PCT/EP 09/055508

(96) Az európai bejelentés bejelentési száma:

EP 20090742109

(87) A nemzetközi közzétételi szám:

WO 09135891

(97) Az európai bejelentés közzétételi adatai:

EP 2276801 A1 **2009. 11. 12.**

(97) Az európai szabadalom megadásának meghirdetési adatai:

EP 2276801 B1 **2017. 03. 08.**(30) Elsőbbségi adatok:
08155987 **2008. 05. 09.** **EP**(73) Jogosult(ak):
Serge Ferrari SAS, 38110 Saint Jean de Soudain (FR)(72) Feltalálók(k):
VAN WEYNBERGH, Jacques, B-1082 Bruxelles (BE)
NOEL, Stephane, B-1330 Rixensart (BE)(74) Képviselő:
SBGK Szabadalmi Ügyvivői Iroda, Budapest(54) **Eljárás szálerősítéses polimer alapú termékek újrafeldolgozására**

Az európai szabadalom ellen, megadásának az Európai Szabadalmi Közlönyben való meghirdetésétől számított kilenc hónapon belül, felszólalást lehet benyújtani az Európai Szabadalmi Hivatalnál. (Európai Szabadalmi Egyezmény 99. cikk(1))

A fordítást a szabadalmas az 1995. évi XXXIII. törvény 84/H. §-a szerint nyújtotta be. A fordítás tartalmi helyességét a Szellemi Tulajdon Nemzeti Hivatala nem vizsgálta.

(19)



(11)

EP 2 276 801 B1

(12)

EUROPEAN PATENT SPECIFICATION

(45) Date of publication and mention of the grant of the patent:
08.03.2017 Bulletin 2017/10

(21) Application number: **09742109.3**

(22) Date of filing: **07.05.2009**

(51) Int Cl.:
C08J 11/08 (2006.01)

(86) International application number:
PCT/EP2009/055508

(87) International publication number:
WO 2009/135891 (12.11.2009 Gazette 2009/46)

(54) **PROCESS FOR RECYCLING ARTICLES BASED ON A FIBRE REINFORCED POLYMER**
VERFAHREN ZUM RECYCELN VON ARTIKELN MIT EINEM FASERVERSTÄRKTEN POLYMER
PROCÉDÉ POUR RECYCLER DES ARTICLES BASÉS SUR UN POLYMÈRE RENFORCÉ À FIBRE

(84) Designated Contracting States:
AT BE BG CH CY CZ DE DK EE ES FI FR GB GR HR HU IE IS IT LI LT LU LV MC MK MT NL NO PL PT RO SE SI SK TR
Designated Extension States:
AL BA RS

(30) Priority: **09.05.2008 EP 08155987**

(43) Date of publication of application:
26.01.2011 Bulletin 2011/04

(73) Proprietor: **Serge Ferrari SAS**
38110 Saint Jean de Soudain (FR)

(72) Inventors:
• **VAN WEYNBERGH, Jacques**
B-1082 Bruxelles (BE)
• **NOEL, Stephane**
B-1330 Rixensart (BE)

(74) Representative: **Cabinet Laurent & Charras et al**
Le Contemporain
50 Chemin de la Bruyère
69574 Dardilly Cedex (FR)

(56) References cited:
US-A- 3 624 009 US-B1- 6 172 125

EP 2 276 801 B1

Note: Within nine months of the publication of the mention of the grant of the European patent in the European Patent Bulletin, any person may give notice to the European Patent Office of opposition to that patent, in accordance with the Implementing Regulations. Notice of opposition shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

Description

[0001] The present invention relates to a process for the recycling of articles, and in particular of sheets, based on a fibre reinforced polymer (preferably a vinyl chloride polymer).

[0002] Vinyl chloride polymers (PVC) are widely used for the manufacture of a variety of articles. They are used, for example, for the manufacture of sheets - generally reinforced with reinforcing fibres - intended for covering soils, for covering vehicles (trucks), for the concealing of buildings on which work is being carried out, for the construction of sets for entertainments or exhibitions, or for advertizing purposes. These sheets generally have a large surface area. Furthermore, in many applications, most particularly in advertising applications or in applications involving sets for entertainments or exhibitions, which at the present time are growing considerably, their lifetime is short, typically a few weeks or months. For these reasons, large quantities of such sheets are scrapped each year. The process of recycling them consequently constitutes a major ecological and economic problem.

[0003] The same situation applies with regard to other articles based on one or more, flexible or rigid, polymers (preferably PVC) for example with regard to conveyor belts, coated fabrics and other elements for the interior furnishing of vehicles, pipes and hoses, window frames or polymer-insulated power cables.

[0004] Thorough grinding of these articles usually results in a mixture of fine particles of heterogeneous composition, which is difficult to purify and reuse. Furthermore, in the case of fibre-reinforced articles (for example, polyester or nylon-fibre-reinforced articles), the fibres often form a kind of wadding which makes it much more complicated to reuse the ground particles.

[0005] Various processes based on dissolving, using organic solvents, have already been proposed ; however, they often cause safety and pollution problems.

[0006] Patents EP 945481, EP 1232204 and EP 1268628 to SOLVAY aim at solving that problem by providing a recycling process which is simple, economic and reliable, which does not produce much pollution and which allows plastics of high purity and advantageous morphology to be recovered, while substantially preventing any additives from being extracted therefrom. This process includes the dissolution of the polymer in an adequate solvent capable of dissolving it and susceptible of forming an azeotrope with water, and its precipitation by injecting steam in the solution thus obtained, which moreover leads to the stripping of the water-solvent azeotrope and so leaves behind a mixture essentially consisting of water and of solid polymer particles which are recovered, while the vapours resulting from the stripping are condensed and decanted so as to recover both a solvent phase for a subsequent dissolution and a "water" phase for a subsequent precipitation.

[0007] These patents specify that if the article is rein-

forced with fibres, said fibres may be recovered as well and in order to increase their purity, they may be subjected to a subsequent centrifuging and/or washing step, for example using the same solvent, for the purpose of removing any residual traces of polymer. According to their teaching, the solvent which would have been used for the washing may advantageously be mixed with the fresh solvent used for the dissolving step ; the fact that it contains traces of dissolved polymer does not in any way lessen the effectiveness of the dissolving operation. The fibres may be reused directly for the manufacture of plastic-based reinforced articles.

[0008] JP 20008-0621486 describes in its examples, a washing treatment according to which the recovered fibres are first washed by "filtration washing" (i.e. the fibres are recovered from the polymer solution on a filter through which solvent is passed, eventually several times) and then, they are either (1) merely dried with hot air, or (2) they are first dispersed again in solvent to form a slurry in which steam is injected to remove the solvent, after which the fibres are recovered by filtration and dried using hot air. However, such a process offers the disadvantage that either the residual content of solvent on the fibres is too high (alternative (1)) or the process is economically unattractive because much solvent has to be evaporated (alternative (2)) or because of the use of hot air (alternatives (1) and (2)). Additionally, in both alternatives, the residual polymer of vinyl chloride is removed by rinsing with solvent through the fibres retained on a filter, which is not very efficient.

[0009] The present invention aims at solving these problems by providing a process for recycling a fibre reinforced polymer article, according to which :

- said article, eventually cut in fragments, is dissolved in a solvent capable of dissolving the polymer but not the fibres and which is susceptible of forming an azeotrope with water, forming a solution ;
- the fibres are recovered from said solution ;
- said fibres are washed with solvent by putting them in the form of a slurry in said solvent and by agitating the slurry and/or by making the solvent circulate through it ;
- the washed fibres are recovered from said slurry ; and
- said recovered fibres are submitted to a stripping with steam in order to substantially remove the solvent.

[0010] The articles reinforced with fibres in question may be of any kind of polymer, although they preferably consist of one or more vinyl chloride (VC) polymers (PVC) reinforced with fibres. VC polymer (PVC) should be understood to mean any homopolymer or copolymer containing at least 50 % by weight of VC. Homopolymers of vinyl chloride are generally used. Apart from one or more polymers and fibres, the articles may also contain one or more of the usual additives such as, for example, plasti-

cizers, stabilizers, antioxidants, fire retardants, pigments, fillers, etc.

[0011] The articles may be in any form, for example in the form of flexible pipes or hoses or rigid pipes, containers, sheets for covering soils (carpet tiles for instance), tarpaulins, window frames, insulating sheaths of power cables, wall papers, etc. They may have been manufactured by any known technique : extrusion, coating, injection moulding, etc.

[0012] The term "sheet" should be understood to mean any thin, flexible or rigid, single-layer or multilayer article, which may or may not be reinforced by reinforcing fibres embedded in the plastic. These sheets have any thickness, however this is generally less than 10 mm ; their thickness is usually between 0.1 and 5 mm. The process is particularly advantageous for recycling carpet tiles or tarpaulins, i.e. fibre-reinforced sheets, intended especially for covering soils, for covering vehicles, for concealing buildings on which work is being carried out, for the construction of sets for entertainments or exhibitions, or for advertizing purposes. The sheets may have been manufactured by any technique, for example by calendering or by coating ; the reinforced sheets are often manufactured by coating a network of fibres by means of a plastisol and heating.

[0013] The reinforcing fibres may be of any kind - natural or synthetic ; glass fibres, cellulose fibres or plastic fibres may especially be used. Often these are plastic fibres, particularly polyester (like polyethylene terephthalate (PET)) or polyamide (nylon) fibres. The diameter of the fibres is usually of the order of 10 to 100 μm . They are often long fibres, the length of which may be up to several metres. However, they may also be shorter fibres - ranging from a few millimetres to a few centimetres in length - possibly forming a woven fabric, a nonwoven fabric or a felt. By way of illustration, the fibres may represent from 1 to 50 % of the weight of a reinforced sheet, generally from 10 to 40 %.

[0014] While the process according to the present invention allows to recycle fibre reinforced polymer articles, this process can also be used for recycling polymer (preferably PVC) / metal composites containing preferably from 5 to 99 % by weight of metal. Any metal may be present in such composites. Preferably metals such as copper, aluminium, iron alloys or steels are present in the composites. The composites may be in any form, for example in the form of plates, cables, rods or threads. They may have been manufactured by any known technique. All the characteristics and preferences detailed hereafter for the process according to the invention apply for the process for recycling such composites except that preference is particularly given to the use of a rotating tumbler.

[0015] The first step of the process according to the invention consists, if necessary, in cutting the articles so as to reduce them to fragments which are easy to handle. It is clear that if the article is already in the form of fragments of suitable dimensions, the cutting step is super-

fluous.

[0016] The fragments of articles thus obtained are then subjected to the action of a solvent having several specific characteristics. This operation may be carried out in any suitable apparatus, especially one which takes into account the safety and environmental requirements, for example in a closed reactor exhibiting sufficient chemical resistance. The apparatus is preferably a cylindrical closed reactor which can be placed horizontally or vertically. Among such apparatus which can be used, one can cite the dissolver-filter which is a cylindrical closed reactor placed vertically in which a screen is incorporated as a bottom filtration plate, the rotating tumbler which is a cylindrical closed reactor placed horizontally containing a cylindrical perforated basket and other cylindrical closed reactors placed vertically containing a cylindrical perforated basket.

[0017] The reaction mixture is preferably stirred (this is particularly the case with the dissolver-filter which is preferably equipped with an axial pumping impeller), placed in rotation in only one direction or in each direction alternatively (this is particularly the case with the rotating tumbler containing a cylindrical perforated basket, the rotation of which is forced by an engine) and/or a pump is used to circulate the solvent inside the apparatus (this is particularly the case with the other cylindrical closed reactors placed vertically containing a cylindrical perforated basket).

[0018] The solvent used is a substance - or a mixture of substances - capable of dissolving the polymer, preferably vinyl chloride polymer, contained in the treated article. However the solvent must not dissolve the reinforcing fibres. It is also necessary, in the context of the process according to the invention, for the solvent used to be advantageously at least partially miscible with water, and to form an azeotrope with water.

[0019] Particularly if the polymer is PVC, the solvent is advantageously chosen from methyl ethyl ketone (MEK), methyl isobutyl ketone (MIBK), n-methylpyrrolidone (NMP) and tetrahydrofuran (THF). Especially if the polymer is PVC, it is preferred to use MEK (preferably both for dissolving the article and for the washing of the fibres) which forms, with water, an azeotrope containing (at atmospheric pressure) 11 % water and 89 % MEK (by weight). This solvent does not dissolve glass, polyester or nylon fibres and gives quite good results.

[0020] In fact, in the context of an industrial process which is either a batch process functioning in loop, or a continuous process, where the reactives (water and solvent) are recycled and where additives are used (like a phase separation agent (PSA) and an alcohol as described in patent application FR 2921372 in the name of SOLVAY), the solvent is generally not pure but does comprise a few percents (in weight) of these additives (for instance up to 10 %, even 31 %). The term "solvent" is therefore to be understood to mean both single substances and mixtures of substances.

[0021] Examples of PSA are aliphatic hydrocarbon

having 5 to 7 carbon atoms. In particular with MEK as solvent, excellent results have been obtained by choosing n-hexane or isohexane (2-methylpentane) as a phase separation agent. Examples of alcohols are linear aliphatic alcohols apart from methanol. In particular, C2-C6 linear aliphatic alcohols are especially suitable, and most particularly C3-C5 linear aliphatic alcohols. Isopropanol and *tert*-butyl alcohol (2-methyl-2-propanol) are therefore particularly suitable. The solvent may also contain a certain amount of water.

[0022] When the polymer is PVC, good results have been obtained when the solvent used both for dissolving the article and for washing of the fibres contains between 2 % and 8 % by weight of alcohol (preferably isopropanol), between 13 % and 17 % of a PSA (preferably isohexane) and between 4 % and 6 % of water, the remainder (namely between 69 % and 81 % by weight) being composed of MEK.

[0023] The dissolving operation is carried out at any temperature, but the solvent is advantageously liquid at this temperature, of course. It is preferred to operate under auto-generated pressure and at a temperature of advantageously between 20 and 120°C, preferably from 50 to 100°C.

[0024] The dissolving operation may be carried out at any pressure. However, it is preferred to carry out the operation at a pressure of from 2 to 10 barg (relative pressure, that is the gauge pressure measured on the pressure gauge), preferably from 2 to 4 bar.

[0025] The time during which the dissolving operation is carried is advantageously of at least 5, preferably of at least 10 minutes. This time is advantageously of at most 120, preferably of at most 40 minutes.

[0026] According to a preferred embodiment of the present invention, prior to being dissolved, the (fragments of the) article are put either directly into the dissolver-filter above the screen incorporated as a bottom filtration plate, or into the cylindrical perforated basket contained in the rotating tumbler or in the other cylindrical closed reactors. The holes of the screen or of the cylindrical perforated basket are preferably chosen of a size such that they keep most the fibres inside the recipient, while still allowing the solvent to circulate so that efficient dissolution can occur.

[0027] The introduction of the (fragments of the) article into the apparatus is preferably followed by oxygen removal and replacement by an inert gas like nitrogen, for safety reasons.

[0028] Available after the dissolving step is advantageously a solution comprising, on the one hand, a liquid phase consisting of the solvent in which the polymer is dissolved and, on the other hand, any undissolved constituents including the reinforcing fibres. According to the invention, these are recovered from said solution by being separated from the liquid phase, preferably by filtration, more preferably through the screen of the dissolver-filter whose apertures have dimensions of the order of 0.1 to 10 mm, or through the cylindrical perforated basket

contained in the rotating tumbler or in the other cylindrical closed reactors whose apertures have dimensions of the order of 0.1 to 10 cm.

[0029] In a first preferred embodiment of the present invention, the dissolution takes place in a dissolver-filter in which a screen is incorporated into the dissolver as a bottom filtration plate so that at the end of the dissolution step, the polymer solution is discharged through said plate to be treated separately, while the fibres and all eventual other insolubles remain inside the dissolver to be further treated therein as claimed i.e. by washing them first, recovering them substantially dry again and then, stripping them with steam. Hence, in that embodiment, the dissolver is called a dissolver-filter because dissolution, separation, washing and steam stripping steps all occur therein.

[0030] In a second preferred embodiment of the present invention, the dissolution takes place in a rotating tumbler containing a cylindrical perforated basket so that at the end of the dissolution step, the polymer solution is discharged through said perforated basket to be treated separately, while the fibres and all eventual other insolubles remain inside the basket to be further treated inside said basket and tumbler as claimed i.e. by washing them first, recovering them substantially dry again and then, stripping them with steam. Hence, in that embodiment, dissolution, separation, washing and steam stripping steps all occur inside the basket and tumbler.

[0031] If the stripping step is performed on the fibres as they are after their separation from the polymer solution, the residual content of polymer (preferably PVC) on the fibres is such that they will stick together. Hence, the residual polymer content must be very low and/or said polymer must be precipitated on the fibres before stripping them.

[0032] Hence, according to the invention, the fibres are washed with solvent by putting them in the form of a slurry in said solvent (preferably hot, more preferably the same as the one used for the dissolution of the article), advantageously at least once, before steam stripping them. Preferably, they are washed at least twice and even more preferably, at least 3 times before steam stripping them. The solvent resulting from each washing step is preferably stored in a buffer tank so that it can be re-used in a subsequent batch, for instance for washing and/or dissolving. During washing, the slurry is preferably agitated and/or the solvent circulates through it.

[0033] Alternatively or additionally, after having been washed (eventually several times) with solvent, the fibres may be washed with (preferably hot) water by putting them in the form of a slurry in water, before being steam stripped (so as to precipitate the polymer onto the fibres). During washing, the slurry is preferably agitated and/or water circulates through it.

[0034] Preferably, the recovering of the fibres after the/each washing step occurs as after the dissolution i.e. the fibres are retained on the bottom filtration plate of the dissolver-filter or inside the cylindrical perforated basket

of the rotating tumbler and the steam stripping step occurs while the fibres are still retained on/in these items, by blowing steam therein. In the particular case of the dissolver-filter, steam can be either introduced by the bottom or by the top, preferably by the top. In the particular case of the rotating tumbler, the way steam is introduced is not critical. The evaporated solvent is advantageously sent to a condensation sector and recovered.

[0035] After the stripping, either the fibres are advantageously placed in suspension under stirring by addition of cold water and discharged from the dissolver-filter to the atmosphere by by-passing the bottom filtration plate or the tumbler is advantageously opened then the basket extracted from the tumbler by a mechanical device and opened, so that the fibres retained therein are discharged to the atmosphere.

[0036] On the other hand, the polymer solution is advantageously sent to the precipitator. In this vessel, the solvent is advantageously evaporated under vacuum and agitation by direct steam injection. At a certain moment, the organic phase is no longer a solvent for the polymer so that said polymer will advantageously precipitate in small particles with the fillers and the plasticizer. The particle size and the bulk density of the polymer particles are advantageously controlled by the injection of water and surfactant agent during this process. The organic phase is advantageously replaced by the aqueous phase and a solvent-free slurry of recycled polymer particles into water is obtained at the end of the process. This slurry is advantageously discharged to a slurry tank and continuously centrifuged. The water is advantageously sent to the waste water treatment and the recycled polymer is advantageously dried in a fluidized bed, sieved and packaged in big-bags.

[0037] The evaporated solvent is advantageously sent to the condensation sector, decanted from aqueous phase and stored, ready for use. The non condensed gas, containing nitrogen and solvent is advantageously burned.

[0038] The invention will further be described in more in detail based on one best mode of carrying out the first preferred embodiment of the present invention and which concerns the treatment of tarpaulins made of plasticized PVC containing among others DOP (dioctyl phthalate) as plasticizer, polyester as well as CaCO_3 as inert filler.

[0039] The tarpaulins are first cut into fragments, the dimensions of which are of maximum 10 to 15 cm and then fed into a cylindrical dissolver-filter, placed vertically, equipped with an axial pumping impeller and with a screen incorporated as a bottom filtration plate. This dissolver-filter is closed and the oxygen is removed and replaced by nitrogen. This step is necessary for safety reasons because the vapour of solvent mixed with oxygen is explosive.

[0040] Hot solvent (coming from a buffer tank containing the solvent of the first washing of the last batch) is introduced into the dissolver-filter. The dissolution occurs typically at 95°C, 3 barg and takes 10 minutes under ag-

itation.

[0041] Typically the solvent contains 75 % MEK, 15 % isohexane, 5 % water and 5 % isopropanol.

[0042] This solvent will selectively dissolve the PVC resin and the plasticizer. The filler (CaCO_3) is not dissolved but is in suspension in the solution. The polyester and the nylon fibres are not dissolved.

[0043] After complete dissolution of the PVC, the solution is discharged from the dissolver-filter to a precipitator through the bottom filtration plate. This filtration plate is perforated with holes small enough to keep the fibres and big enough to avoid plugging and have a high filtration rate.

[0044] Once the filtration is completed, the fibres are still wet and contain a certain amount of solution containing PVC resin.

[0045] Depending on the maximum allowed concentration of PVC resin in the fibres, some washing steps may be required. This (these) washing step(s) is performed by putting the fibres in suspension into the dissolver-filter with hot solvent. The hot solvent is coming from buffer tank(s) containing the washing solvent of the previous batch. Generally, it is only after three washing steps that the PVC content into the fibres is low enough to minimize the risk of sticking the fibres during the steam stripping step. Preferably, only the last washing step is performed with clean hot solvent. For every step, the washing solvent is stored in a separate buffer and used for the next batch.

[0046] The washed fibres are wet and still contain solvent. This solvent has to be removed in order to discharge the fibres safely to the atmosphere. The removal of the solvent is done in the dissolver-filter by injecting steam therein through the top of the dissolver. The evaporated solvent is sent to a condensation sector and recovered. After the stripping, the fibres are placed in suspension under stirring by addition of cold water and discharged from the dissolver-filter to the atmosphere near the fibres treatment facilities.

[0047] The dissolver filter is then ready for the next batch.

[0048] On the other hand, the PVC solution is sent to the precipitator. In this vessel, the solvent is evaporated under vacuum and agitation by direct steam injection. At a certain moment, the organic phase is no longer a solvent for the PVC so that said PVC will precipitate in small particles with the fillers and the plasticizer. The particle size and the bulk density of the PVC particles are controlled by the injection of water and surfactant agent during this process. The organic phase is replaced by the aqueous phase and a solvent-free slurry of recycled PVC particles into water is obtained at the end of the process. This slurry is discharged to a slurry tank and continuously centrifuged. The water is sent to the waste water treatment and the recycled PVC is dried in a fluidized bed, sieved and packaged in big-bags.

[0049] The evaporated solvent is sent to the condensation sector, decanted from aqueous phase and stored,

ready for use. The non condensed gas, containing nitrogen and solvent is burned.

[0050] The invention will further be described in more in detail based on one best mode of carrying out the second preferred embodiment of the present invention and which concerns the treatment of carpet tiles made of plasticized PVC containing among others DOP (dioctyl phthalate) as plasticizer, polyester and/or nylon fibres, as well as CaCO_3 as inert filler.

[0051] These tiles, the dimensions of which are 50 cm x 50 cm square, are fed, without being cut beforehand, into a cylindrical perforated basket which is introduced with a mechanical device into a tumbler, the rotation of which can be forced by an engine. This tumbler is closed and the oxygen is removed and replaced by nitrogen. This step is necessary for safety reasons because the vapour of solvent mixed with oxygen is explosive.

[0052] Hot solvent (coming from a buffer tank containing the solvent of the first washing of the last batch) is introduced into the tumbler. The dissolution occurs typically at 95°C, 3 barg and takes 10 minutes under rotation of the tumbler and the basket.

[0053] Typically the solvent contains 75 % MEK, 15 % isohexane, 5 % water and 5 % isopropanol.

[0054] This solvent will selectively dissolve the PVC resin and the plasticizer. The filler (CaCO_3) is not dissolved but is in suspension in the solution. The polyester and the nylon fibres are not dissolved.

[0055] After complete dissolution of the PVC, the solution is discharged from the tumbler to a precipitator through the cylindrical perforated basket. This basket is perforated with holes small enough to keep the fibres and big enough to avoid plugging and have a high filtration rate.

[0056] Once the filtration is completed, the fibres are still wet and contain a certain amount of solution containing PVC resin.

[0057] Depending on the maximum allowed concentration of PVC resin in the fibres, some washing steps may be required. This (these) washing step(s) is performed by putting the fibres in suspension into the tumbler with hot solvent. The hot solvent is coming from buffer tank(s) containing the washing solvent of the previous batch. Generally, it is only after three washing steps that the PVC content into the fibres is low enough to minimize the risk of sticking the fibres during the steam stripping step. Preferably, only the last washing step is performed with clean hot solvent. For every step, the washing solvent is stored in a separate buffer and used for the next batch.

[0058] The washed fibres are wet and still contain solvent. This solvent has to be removed in order to discharge the fibres safely to the atmosphere. The removal of the solvent is done in the tumbler by injecting steam therein. The evaporated solvent is sent to a condensation sector and recovered. After the stripping, the tumbler is opened, then the basket is extracted from the tumbler by a mechanical device and opened, so that the fibres retained

therein are discharged to the atmosphere near the fibres treatment facilities. The tumbler is then ready for the next batch (another basket full of carpet tiles should be ready at this moment).

5 **[0059]** On the other hand, the PVC solution is sent to the precipitator. In this vessel, the solvent is evaporated under vacuum and agitation by direct steam injection. At a certain moment, the organic phase is no longer a solvent for the PVC so that said PVC will precipitate in small particles with the fillers and the plasticizer. The particle size and the bulk density of the PVC particles are controlled by the injection of water and surfactant agent during this process. The organic phase is replaced by the aqueous phase and a solvent-free slurry of recycled PVC particles into water is obtained at the end of the process. This slurry is discharged to a slurry tank and continuously centrifuged. The water is sent to the waste water treatment and the recycled PVC is dried in a fluidized bed, sieved and packaged in big-bags.

10 **[0060]** The evaporated solvent is sent to the condensation sector, decanted from aqueous phase and stored, ready for use. The non condensed gas, containing nitrogen and solvent is burned.

25

Claims

1. - Process for recycling a fibre reinforced polymer article, according to which:
- 30
- said article, eventually cut in fragments, is dissolved in a solvent capable of dissolving the polymer but not the fibres and which is susceptible of forming an azeotrope with water, forming a solution ;
 - the fibres are recovered from said solution ;
 - said fibres are washed with solvent by putting them in the form of a slurry in said solvent and by agitating the slurry and/or by making the solvent circulate through it ;
 - the washed fibres are recovered from said slurry ; and
 - said recovered fibres are submitted to a stripping with steam in order to substantially remove the solvent.
- 35
2. - Process according to Claim 1, in which the solvent used for dissolving the article and for the washing of the fibres is the same.
- 40
3. - Process according to any one of Claims 1 or 2, in which the polymer is a vinyl chloride polymer and the solvent used both for dissolving the article and for the washing of the fibres contains between 2 % and 8 % by weight of alcohol, between 13 % and 17 % of a phase separation agent and between 4 % and 6 % of water, the remainder (namely between 69 % and 81 % by weight) being composed of methyl ethyl
- 45
- 50
- 55

ketone.

4. - Process according to any one of Claims 1 to 3, in which the dissolution takes place in a dissolver-filter in which a screen is incorporated into the dissolver as a bottom filtration plate so that at the end of the dissolution step, the polymer solution is discharged through said plate to be treated separately while the fibres and all eventual other insolubles remain inside the dissolver to be further treated therein. 5
5. - Process according to any one of Claims 1 to 3, in which the dissolution takes place in a rotating tumbler containing a cylindrical perforated basket so that at the end of the dissolution step, the polymer solution is discharged through said perforated basket to be treated separately while the fibres and all eventual other insolubles remain inside the basket to be further treated inside said basket and tumbler. 10 15
6. - Process according to any one of Claims 1 to 5, in which the fibers are washed at least once before steam stripping them. 20
7. - Process according to any one of Claims 1 to 6, in which the fibers are washed at least twice before steam stripping them. 25
8. - Process according to any one of Claims 6 or 7, in which the solvent resulting from each washing step is stored in a buffer tank and re-used in a subsequent batch, for instance for washing and/or dissolving. 30
9. - Process according to any one of Claims 1 to 8, in which after having been washed with solvent and before being steam stripped, the fibres are washed with water so as to precipitate the polymer onto the fibres. 35
10. - Process according to any one of Claims 4 to 9, in which after the washing step, the fibres are retained on the bottom filtration plate of the dissolver-filter or inside the cylindrical perforated basket of the rotating tumbler and the steam stripping step occurs while the fibres are still retained on/in these items, by blowing steam therein. 40 45

Patentansprüche

1. Verfahren zum Recyceln von faserverstärkten Polymerartikeln, demzufolge:
- dieser Artikel, eventuell in Teile geschnitten, in einem Lösungsmittel aufgelöst wird, das in der Lage ist, Polymer, aber nicht die Fasern aufzulösen und welches geeignet ist, ein Azeotrop mit Wasser zu bilden, durch Bildung einer Lösung;

- diese Fasern werden aus der Lösung zurückgewonnen;

- diese Fasern werden mit einem Lösungsmittel gewaschen, indem sie in der Form einer Aufschlämmung in das Lösungsmittel gegeben und die Aufschlämmung gerührt wird und/oder indem das Lösungsmittel dazu gebracht wird sie zu durchströmen;

- die gewaschenen Fasern werden aus der Aufschlämmung zurückgewonnen;

und

- diese recycelten Fasern werden einem Dampfstripping unterzogen, um das Lösungsmittel größtenteils zu entfernen.

2. Verfahren gemäß Anspruch 1, bei dem das gleiche Lösungsmittel zum Auflösen des Artikels und zum Waschen der Fasern verwendet wird. 20
3. Verfahren nach einem der Ansprüche 1 oder 2, bei dem es sich bei dem Polymer um ein Vinylchlorid-Polymer handelt und das Lösungsmittel, das sowohl zum Auflösen des Artikels und zum Waschen der Fasern verwendet wird, zwischen 2 und 8 Gewichts-% Alkohol, zwischen 13 % und 17 % eines Phasentrennungsmittels und zwischen 4 % und 6 % Wasser enthält, der Rest (namentlich zwischen 69 und 81 Gewichts-%) besteht aus Methylethylketon 25
4. Verfahren nach einem der Ansprüche 1 bis 3, bei dem die Auflösung in einem Dissolver-Filter stattfindet, bei dem im Dissolver eine Zwischenwand integriert ist, als Boden - Filtrationsplatte, so dass am Ende des Auflösungs-schrittes die Polymerlösung durch diese Platte abgeleitet wird, um getrennt behandelt zu werden, während die Fasern und alle verbleibenden sonstigen, nicht lösliche Stoffe im Dissolver zurückbleiben, um darin weiter behandelt zu werden. 30
5. Verfahren nach einem der Ansprüche 1 bis 3, bei dem die Auflösung in einem rotierenden Tumbler stattfindet, der einen zylindrischen, perforierten Korb enthält, so dass am Ende des Auflösungs-schrittes die Polymerlösung durch diesen perforierten Korb abgeleitet wird, um getrennt behandelt zu werden, während die Fasern und alle verbleibenden sonstigen, nicht lösliche Stoffe im Korb zurückbleiben um in diesem Korb und Tumbler weiter behandelt zu werden. 35 40 45
6. Verfahren nach einem der Ansprüche 1 bis 5, in dem die Fasern vor dem Dampfstripping mindestens einmal gewaschen werden. 50
7. Verfahren nach einem der Ansprüche 1 bis 6, in dem die Fasern vor dem Dampfstripping mindestens

zweimal gewaschen werden.

8. Verfahren nach einem der Ansprüche 6 oder 7, bei dem das Lösungsmittel, das für den Waschen Schritt verwendet wird, in einem Puffertank gelagert wird um bei einer folgenden Charge, zum Beispiel zum Waschen und/ oder Auflösen eingesetzt zu werden.
9. Verfahren nach einem der Ansprüche 1 bis 8, in dem, nachdem die Fasern gewaschen und vor dem Dampfstripping, die Fasern mit Wasser gewaschen werden, um das Polymer auf die Fasern abzuscheiden.
10. Verfahren nach einem der Ansprüche 4 bis 9, bei dem nach dem Waschen die Fasern auf einer Boden- Filtrationsplatte des Dissolver- Filters oder innerhalb des zylindrischen, perforierten Korbes des rotierenden Tumblers zurückgehalten werden und das Dampfstripping stattfindet, während die Fasern noch auf/ in diesen Vorrichtungen zurückgehalten werden, indem Dampf darüber geleitet wird.

Revendications

1. Procédé pour recycler un article basé sur un polymère renforcé à fibre, dans lequel :
 - ledit article, finalement découpé en fragments, est dissous dans un solvant capable de dissoudre le polymère mais pas les fibres et susceptible de former un azéotrope avec de l'eau, formant une solution ;
 - les fibres sont recouvertes de ladite solution ;
 - lesdites fibres sont lavées avec un solvant en les mettant sous la forme d'une bouillie dans ledit solvant et en agitant la bouillie et/ou en y faisant circuler le solvant ;
 - les fibres lavées sont récupérées de ladite bouillie ; et
 - lesdites fibres récupérées sont soumises à un décapage à la vapeur afin de retirer l'essentiel du solvant.
2. Procédé selon la revendication 1, dans lequel le solvant utilisé pour dissoudre l'article et pour le décapage des fibres est le même.
3. Procédé selon l'une quelconque des revendications 1 ou 2, dans lequel le polymère est un polymère de chlorure de vinyle et le solvant utilisé tant pour dissoudre l'article que pour décapager les fibres contient entre 2% et 8% en poids d'alcool, entre 13% et 17% d'un agent de séparation de phase et entre 4% et 6% d'eau, le reste (à savoir entre 69% et 81% en poids) étant composé de méthyle éthyle cétone.

4. Procédé selon l'une quelconque des revendications 1 à 3, dans lequel la dissolution intervient dans un filtre dissolvant dans lequel un tamis est incorporé dans le dissolvant comme plaque de filtration inférieure de façon qu'à la fin de l'étape de dissolution, la solution de polymère soit évacuée au travers de ladite plaque pour être traitée séparément cependant que les fibres et toutes les autres particules insolubles subsistantes restent dans le dissolvant pour continuer à y être traitées.
5. Procédé selon l'une quelconque des revendications 1 à 3, dans lequel la dissolution intervient dans un tambour culbuteur rotatif contenant un panier perforé cylindrique de façon qu'à la fin de l'étape de dissolution, la solution de polymère soit évacuée au travers dudit panier perforé pour être traitée séparément cependant que les fibres et tous les autres débris insolubles subsistants restent à l'intérieur du panier pour continuer à être traités dans lesdits panier et tambour culbuteur.
6. Procédé selon l'une quelconque des revendications 1 à 5, dans lequel les fibres sont lavées au moins une fois avant que la vapeur ne les décape.
7. Procédé selon l'une quelconque des revendications 1 à 6, dans lequel les fibres sont lavées au moins deux fois avant que la vapeur ne les décape.
8. Procédé selon l'une quelconque des revendications 6 à 7, dans lequel le solvant résultant de chaque étape de lavage est stocké dans une cuve tampon et réutilisé dans un bain ultérieur, par exemple pour le lavage et/ou la dissolution.
9. Procédé selon l'une quelconque des revendications 1 à 8, dans lequel après avoir été lavées avec le solvant et avant d'être décapées à la vapeur, les fibres sont lavées à l'eau de façon à faire précipiter le polymère sur les fibres.
10. Procédé selon l'une quelconque des revendications 4 à 9, dans lequel après l'étape de lavage, les fibres sont maintenues sur la plaque de filtration inférieure du dissolvant/filtre ou à l'intérieur du panier cylindrique perforé du tambour culbuteur rotatif et l'étape de décapage à la vapeur intervient alors que les fibres sont encore maintenues sur/dans ces éléments, en y soufflant la vapeur.

REFERENCES CITED IN THE DESCRIPTION

This list of references cited by the applicant is for the reader's convenience only. It does not form part of the European patent document. Even though great care has been taken in compiling the references, errors or omissions cannot be excluded and the EPO disclaims all liability in this regard.

Patent documents cited in the description

- EP 945481 A [0006]
- EP 1232204 A [0006]
- EP 1268628 A [0006]
- JP 200080621486 B [0008]
- FR 2921372 [0020]

Eljárás szálerősítéssel polimer alapú termékek újrafeldolgozására

SZABADALMI IGÉNYPONTOK



1. Eljárás szálerősítéssel polimer termékek újrafeldolgozására, mely szerint:
 - az említett terméket, adott esetben darabokra vágva, olyan oldószerben oldjuk, mely oldószer képes a polimer oldására, de nem oldja a szálakat, és amely alkalmas vízzel azeotróp képzésére, oldatot alkotva;
 - a szálakat az említett oldatból kinyerjük;
 - a szálakat oldószerrel mossuk, úgy, hogy behelyezzük őket az említett oldószerbe zagy formában, a zagyot keverjük, és/vagy az oldószert keresztülcirkuláltatjuk rajta;
 - a kimosott szálakat a zagyból kinyerjük; és
 - az említett kinyert szálakat gőzzel sztrippeljük, annak érdekében, hogy az oldószert lényegében eltávolítsuk belőlük.
2. Az 1. igénypont szerinti eljárás, ahol a termék oldására és a szálak mosására használt oldószer azonos.
3. Az 1. vagy 2. igénypontok bármelyike szerinti eljárás, ahol a polimer egy vinilklorid polimer és a termék oldásához, valamint a szálak mosásához használt oldószer egyaránt 2 tömeg% és 8 tömeg% közötti mennyiségű alkoholt, 13 tömeg% és 17 tömeg% közötti mennyiségű fázis elválasztó szert és 4 tömeg% és 6 tömeg% közötti mennyiségű vizet tartalmaz, fennmaradó mennyiségét (nevezetesen 69 tömeg% és 81 tömeg% közötti mennyiségét) pedig metiletilketon alkotja.
4. Az 1-3. igénypontok bármelyike szerinti eljárás, ahol az oldást egy disszolvenszűrőben végezzük, ahol egy elválasztó van beépítve a disszolvenszűrőbe fenék szűrőlemezként, úgy, hogy az oldási lépés végén a polimer oldatot az említett lemezen keresztül távolítjuk el külön kezeléshez, míg a szálak és minden fennmaradó egyéb oldhatatlan anyag visszamarad a disszolvenszűrőben, abban történő további kezelésre.
5. Az 1-3. igénypontok bármelyike szerinti eljárás, ahol az oldást hengeres perforált kosarat tartalmazó forgó dobban végezzük, úgy, hogy az oldási lépés végén a polimer oldat távozik a perforált kosáron keresztül külön történő kezelésre, míg a szálak és minden

fennmaradó egyéb oldhatatlan anyag visszamarad az említett kosár és dob belsejében, abban történő további kezelésre.

6. Az 1-5. igénypontok bármelyike szerinti eljárás, ahol a szálakat legalább egyszer mossuk mielőtt gőzzel sztrippelnénk őket.

7. Az 1-6. igénypontok bármelyike szerinti eljárás, ahol a szálakat legalább kétszer mossuk, mielőtt gőzzel sztrippelnénk őket.

8. A 6. vagy 7. igénypontok bármelyike szerinti eljárás, ahol a mosási lépésekből visszamaradó oldószert egy puffertankban tároljuk és egy következő fürdőben újra felhasználjuk, például mosásra és/vagy oldásra.

9. Az 1-8. igénypontok bármelyike szerinti eljárás, ahol az oldószerezrel való mosás után és a gőzzel való sztrippelés előtt a szálakat vízzel mossuk, a szálakon lévő polimer kicsapására.

10. A 4-9. igénypontok bármelyike szerinti eljárás, ahol a mosási lépést követően, és a disszolver szűrő fenék szűrőlemezén vagy a forgó dob hengeres perforált kosarában a szálak visszamaradnak, és a gőzzel való sztrippelési lépést úgy végezzük, hogy a szálakat még visszatartjuk ezeken/ezekben a berendezésekben, miközben gőzt fúvatunk be.

A meghatalmazott

~~Déry Zoltán~~
szakszolgálati igazgató
SBGK Szabványügyi Ügyviteli Iroda
H-1002 Budapest, Andrássy út 113.
Telefon: 461-1000 Fax: 461-1008
Email: derzsi@sbgk.hu