

**(12) STANDARD PATENT**

**(11) Application No. AU 2005223418 B2**

**(19) AUSTRALIAN PATENT OFFICE**

(54) Title  
**Albumin-purification method comprising a nanofiltration step, solution, and composition for therapeutic use containing same**

(51) International Patent Classification(s)  
**A61K 38/00 (2006.01) C07K 14/765 (2006.01)**  
**C07K 14/76 (2006.01)**

(21) Application No: **2005223418** (22) Date of Filing: **2005.02.23**

(87) WIPO No: **WO05/090402**

(30) Priority Data

(31) Number **0402001** (32) Date **2004.02.27** (33) Country **FR**

(43) Publication Date: **2005.09.29**  
(44) Accepted Journal Date: **2008.12.18**

(71) Applicant(s)  
**Laboratoire Francais du Fractionnement et des Biotechnologies**

(72) Inventor(s)  
**Boyer, Stephane;Padrazzi, Bruno;Schmitthaeusler, Roland;Boulange, Paul;Chtourou, Sami**

(74) Agent / Attorney  
**Griffith Hack, Level 3 509 St Kilda Road, Melbourne, VIC, 3004**

(12) DEMANDE INTERNATIONALE PUBLIÉE EN VERTU DU TRAITÉ DE COOPÉRATION  
EN MATIÈRE DE BREVETS (PCT)

(19) Organisation Mondiale de la Propriété  
Intellectuelle  
Bureau international



(43) Date de la publication internationale  
29 septembre 2005 (29.09.2005)

PCT

(10) Numéro de publication internationale  
WO 2005/090402 A1

(51) Classification internationale des brevets<sup>7</sup> :  
C07K 14/76, 14/765

(74) Mandataire : CABINET LEPEUDRY; 43, rue de la  
Brèche aux Loups, F-75012 Paris (FR).

(21) Numéro de la demande internationale :  
PCT/FR2005/000416

(81) États désignés (sauf indication contraire, pour tout titre de protection nationale disponible) : AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, L.C, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW.

(22) Date de dépôt international :  
23 février 2005 (23.02.2005)

(84) États désignés (sauf indication contraire, pour tout titre de protection régionale disponible) : ARIPO (BW, GH, GM, KE, LS, MW, MZ, NA, SD, SL, SZ, TZ, UG, ZM, ZW), eurasien (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), européen (AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HU, IE, IS, IT, LT, LU, MC, NL, PL, PT, RO, SE, SI, SK, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG).

(25) Langue de dépôt : français  
(26) Langue de publication : français  
(30) Données relatives à la priorité :  
0402001 27 février 2004 (27.02.2004) FR

(71) Déposant (pour tous les États désignés sauf US) : LABORATOIRE FRANCAIS DU FRACTIONNEMENT  
ET DES BIOTECHNOLOGIES [FR/FR]; 3, avenue des  
Tropiques, ZA de Courtabœuf, F-91940 Les Ulis (FR).

Publiée :  
— avec rapport de recherche internationale

En ce qui concerne les codes à deux lettres et autres abréviations, se référer aux "Notes explicatives relatives aux codes et abréviations" figurant au début de chaque numéro ordinaire de la Gazette du PCT.

(54) Title: ALBUMIN-PURIFICATION METHOD COMPRISING A NANOFILTRATION STEP, SOLUTION, AND COMPOSITION FOR THERAPEUTIC USE CONTAINING SAME

(54) Titre : PROCÉDÉ DE PURIFICATION D'ALBUMINE COMPRENANT UNE ÉTAPE DE NANOFILTRATION, SOLUTION ET COMPOSITION À USAGE THÉRAPEUTIQUE LA CONTENANT

(57) Abstract: The invention relates to an albumin-purification method comprising a step consisting in subjecting an aqueous albumin solution, with a concentration of between 15 g/l and 80 g/l and a pH of not less than 7, to nanofiltration in a temperature range of between 15 °C and 55 °C. The invention also relates to: a virally-safe aqueous albumin solution which can be obtained using the inventive method and in which the sites for the transport and binding of the active therapeutic ingredients of the albumin are available; and an albumin composition for therapeutic use, which is obtained by adapting the albumin solution that is intended for clinical use.

(57) Abrégé : L'invention concerne un procédé de purification d'albumine comprenant une étape consistant à soumettre une solution aqueuse d'albumine, de concentration 15 g/l à 80 g/l et de pH non inférieur à (7), à une nanofiltration dans une plage de température allant de 15 °C à 55 °C. L'invention concerne également une solution aqueuse d'albumine viralement sécurisée susceptible d'être obtenue par la mise en œuvre du procédé, dans laquelle les sites de transport et de fixation de principes actifs thérapeutiques de l'albumine sont disponibles et une composition d'albumine à usage thérapeutique obtenue par un traitement d'adaptation de ladite solution d'albumine à un usage clinique.



A1

WO 2005/090402 A1

**ROBINTECH sarl**

Résidence Lahire A3; rue Henry de Montherlant; F33400 TALENCE  
0 / Fax 05 56 04 14 55  
SIRET 378 529 713 00034 / URSSAF 330 847 765 815  
contact@robintech.fr

---

**VERIFICATION OF TRANSLATION**

International Application NO.

WO 2005 /090402 A1  
PCT/FR2005/000416

We, Société ROBINTECH sarl sise à 33400 TALENCE rue Henry de Montherlant Résidence Lahire A3, are the translators of the document(s) attached and we state that the following is a true translation to the best of our knowledge and belief

TALENCE le 25 juillet 2006

**ROBINTECH sarl**

Rue H. de Montherlant - Résidence Lahire A3  
F 33400 TALENCE  
Tél : 05 56 04 14 55 - Fax : 05 56 04 14 61  
Siret : 378 529 713 00034 Bordeaux  
TVA FR 63 378 529 713

*H. Lahire, Sénart*  
*Salles*

ALBUMIN PURIFICATION METHOD COMPRISING A NANOFILTRATION  
STEP, SOLUTION AND COMPOSITION FOR THERAPEUTIC USE  
CONTAINING THE SAME

5 The present invention relates to an albumin purification method comprising a nanofiltration step, a solution and a composition for therapeutic use containing the same, obtainable by the method of the invention.

10 Albumin is a major protein of human or animal blood plasma. Clinical use of albumin, as an active ingredient, requires its extraction and purification, which is traditionally carried out by known methods, such as those of Cohn *et al.* (J. Am. Chem. Soc., 68, 459, 1946) and Kistler *et al.* (Vox Sang., 7, 1962, 414-424) which are 15 additionally applicable to an industrial scale.

EP-A-498 133, EP-A-1 329 461 and EP-A-1 329 460 describe unsafe albumine solutions. WO-A-9 600 237 and Biologicals 29(1), 2001, pages 17-21 describe albumine purification methods based on nanofiltration.

20 Albumin requirements amount to about 100-300 kg per million inhabitants according to country; for that reason, it is necessary, for clinical purposes, to provide an albumin free from pathogenic viruses and contaminants, which are sources of diseases. Thus, as 25 regards transfusion-transmissible viruses, safety is ensured by viral inactivation methods such as liquid-

state pasteurisation of an albumin composition at 60°C for 10 hours in the presence of a biologically compatible stabiliser (sodium caprylate and/or tryptophanate). This viral safety has been established for more than 60 years 5 and the literature does not report any proven case of virus transmission, e.g. hepatitis B, A or C, or HIV. However, some authors mentioned some cases correlated with the presence, after transfusions, of parvoviruses B 19 in albumin compositions or albumin 10 derivate products. Furthermore, a case has also been reported of an albumin composition that would have led to Creutzfeldt-Jakob disease in a patient having undergone liver transplantation (Créange A., 1995).

Some studies revealed a series of adverse effects 15 occurring after an albumin transfusion due to the presence of lipopolysaccharide substances (pyrogens) in amounts lower than the detection threshold of authorised methods, but capable of occurring during perfusions with large volumes of albumin (e.g. plasma exchanges).

20 Other side effects are related to the presence of albumin polymers occurring during albumin purification and especially during the above-mentioned pasteurisation step. Others are related to the presence of added stabilisers to prevent the thermal denaturation linked to 25 pasteurisation, especially in patients with allergic backgrounds.

In order to avoid the risk of transmissible 30 infectious agents being present, it has been suggested to produce a so-called "recombinant" albumin, according to US patent 6,210,683: the gene of albumin is introduced into a host cell, yeast or bacterium, having a high

proliferation potential. In turn, this host cell produces albumin in the culture medium or its cytoplasm. This albumin is then separated from the cells by extraction and purified. However, the presence of host cell proteins 5 is often detected and the purification methods must therefore have a very high resolution, which is generally detrimental to the yield. The production cost of a recombinant albumin may then prove to be too high in comparison to that of an albumin generated from plasma.

10 Another possibility to get round the above-mentioned drawbacks is to implement filtration methods extensively used for the retention of viruses, and chemical or bacteriological contaminants, etc., using qualified filters with variable porosities. One can mention for 15 instance ultrafiltration that has been used from time to time in the absence of other means of virus elimination to ensure the biological safety of an extraction protein such as growth hormone (hGH). The method requires a tangential flow and is only intended for protein 20 solutions or polypeptides having molecular weights equal to or lower than 65 kDa and low concentrations. This leaves chance for viruses to pass through due to the manufacturing imprecision of filtration membranes. Moreover, filter clogging occurs in front operation, 25 resulting in a blockage of the filtrate flow. Filter clogging is all the faster as protein or polypeptide concentration increases.

To increase the viral safety of biological solutions further, it is contemplated to implement nanofiltration, 30 the preferred method for the retention of particles with sizes equal to or greater than about 15-20 nm. This

particle retention on qualified filters is easily achieved on a large scale with aqueous solutions or solutions containing small-sized solutes, peptides, amino-acids, mineral ions or organic compounds smaller than about 5 kDa.

The nanofiltration of large-sized solutes has been achieved, for example, with blood coagulation factors, factor IX and factor XI. The article by Burnouf-Radosevich *et al.* (Vox Sang., 67, p. 132-138, 1994 in reference with Burnouf *et al.*, Vox Sang., 57, p. 225-232, 1989 and Burnouf-Radosevich *et al.*, Transfusion, 22, p. 861-867, 1992) shows that the working conditions of the nanofiltration method imply small volumes of factor IX and factor XI concentrates (3-4 litres), with protein concentrations not exceeding 1 g/L (0.21 g/L and 0.75 g/L, respectively).

Actually, during the nanofiltration process, the large-sized solutes accumulate on the filter, which slows down the filtrate flow until the filter is completely clogged.

The technical solutions to this clogging which consist of implementing a tangential flow nanofiltration, increasing the pressure and changing the directions of the flows do not prove to be efficient because the clogging is progressive and irreversible. An industrial development is therefore greatly compromised because it requires the use of very numerous filters and very large volumes of solutions to be filtered, which generates a prohibitive cost and abnormally increased manufacturing times.

To avoid the clogging phenomenon, a solution consists of adjusting the physicochemical parameters influencing the recovery yield of solutes, while avoiding the passage of contaminants through the filter. Varying 5 these parameters -such as ionic strength, the nature of the solute to be filtered and the pH of the solution to be filtered- as well as the working conditions of the filtration -such as flow rate and pressure- has been the subject of many studies. For instance, the scientific 10 publications by C. Wallis et al., Ann. Rev. Microbiol., 33, p. 413-437, 1979 and S. Jacob, Methods of Biochemical Analysis, 22, p. 307-350, 1974, show that the effect of each parameter can individually result in increased or decreased efficiency of virus retention and recovery 15 yield of solutes, and that combining several parameters does not systematically favour a synergy of the effects of improvement of the filtration conditions.

Considering the clinical significance of albumin and the ever-increasing needs in albumin, it appears 20 necessary to supply patients with an albumin showing degrees of safety and tolerance higher than those of the currently available albumins, while optimising the conditions of a large-scale production. For that purpose, the Applicant therefore strived to develop an albumin 25 purification method comprising a step of nanofiltering concentrated albumin solutions satisfying these objectives. Therefore, the object of the present invention is to provide a solution corresponding to a good compromise between two criteria, the retention 30 efficiency of viruses and/or other macromolecules likely to induce diseases or side effects in patients, and the

recovery yield of albumin, the relative importance of these two criteria depending on the desired application.

Accordingly, the invention relates to a method for purifying albumin comprising a step of submitting an aqueous solution of albumin, with a concentration of 5 15 g/L to 80 g/L and a pH not lower than 7, to a nanofiltration in a temperature range of 15°C to 55°C.

Thus, the Applicant surprisingly found that using a judicious combination of pH, albumin concentration and 10 temperature (and consequently viscosity) values, in the aqueous albumin solution submitted to the nanofiltration step, makes it possible to reach an efficient optimisation of the albumin recovery yield, and rates of 15 reduction of viruses and other undesirable macromolecules higher than the limits set by the control authorities (4 log). It has been brought to light that a 20 nanofiltration carried out under the conditions of the invention makes it possible to filter amounts higher than 5 kg albumin per m<sup>2</sup> filter, thus defining the protein load, while optimising the duration of the operation and the filtrate flow rate. The nanofiltered albumin 25 solutions show a very high degree of safety in respect of particulate contaminants with a size of e.g. at least about 13 nm, e.g. viruses such as non-enveloped viruses, prions, albumin polymers (tetramers-decamers) generated 30 during steps of albumin manufacture or during the pasteurisation at 60°C, micelle-like lipopolysaccharides, nucleic acids and aggregated proteins. The aqueous albumin solutions thus obtained are also an intermediate product capable of being processed into pharmaceutical formulations for clinical use (see further on).

In the context of the invention, the aqueous albumin solutions are solutions free from any reagent employed during various classical steps of albumin manufacture or purification, such as e.g. polyethyleneglycol (PEG), 5 ethanol, organic salts (sodium caprylate, etc.) and inorganic salts. These various reagents are removed from the albumin solution by known processes, such as diafiltration, ultrafiltration, dialysis, etc. As explained above, the presence of such reagents, or 10 variations in their respective concentrations from one sample to another, can result in unfavourable efficiencies in terms of virus retention and albumin recovery.

Thus, surprisingly, the Applicant has further found 15 that a decreasing ionic strength is correlated with a better viral reduction.

It should be made clear that, in the context of the invention, the aqueous albumin solutions submitted to the nanofiltration may have an ionic strength lower than a 20 maximum threshold connected, in this case, with the maximum allowable basic pH value. This value typically corresponds to about 11.5. The pH must be controlled in particular so that the variation in ionic strength caused by the addition of a pH control agent to the aqueous 25 albumin solution is only very small, if not insignificant. Such pH controllers will preferably be selected among strong alkali metal bases such as NaOH and KOH, and among strong acids such as HCl, while respecting the above-mentioned criteria. Furthermore, it is 30 preferable to avoid pH controllers based on organic compounds, such as organic bases.

Thus, as an example, when aqueous albumin solutions require pH adjustment up to the value of about 11.5, which is achieved by adding 0.5 M NaOH, the solutions obtained will therefore have an ionic strength of about 5 0.0032.

The method of the invention may be implemented with all types of frontal or tangential nanofiltration devices, in particular frontal nanofiltration devices, known to those skilled in the art. Currently used filters 10 have pore sizes smaller than 100 nm.

The nanofiltration step according to the invention is preferably carried out on qualified filters having porosities of at least 13 nm, for example the nanometric filters with porosities of  $15 \pm 2$  or  $20 \pm 2$  nm which are 15 commercially available as pleated membranes, flat membranes or hollow fibres. Regenerated cellulose nanofilters such as PLANNOVA® having a porosities of 15 nm and surface areas of  $0.01 \text{ m}^2$  (from Asahi, Japan), or PALL virus filters (U.S.A.) with porosities of 20 or 50 nm, 20 may be mentioned by way of example.

Any source of albumin starting material (freeze-dried pure product, concentrate, etc.) may be used, in particular those resulting from the fractionation of human or animal blood plasma according to the methods of 25 Cohn et al. (J. Am. Chem. Soc., 68, 459, 1946) or Kistler et al. (Vox Sang., 7, 1962, 414-424).

The pH of the aqueous albumin solution is preferably in the range from about 7.8 to about 11.5, and more preferably, from 9 to 10.5. Albumin is irreversibly 30 altered by pH values higher than approximately 11.5.

The method of the invention is preferably implemented with aqueous albumin solutions having concentrations in the range of 40 to 60 g/L and in a temperature range of 30 to 55°C.

5 According to the invention, the method may further comprise a step of adding a pharmaceutically acceptable salt or salt mixture to the aqueous albumin solution to provide a solution with an ionic strength higher than 0.0032, and preferably, in the range of 0.01 to 0.55, 10 more preferably, of 0.01 to 0.3, even more preferably, of 0.05 to 0.15, and in particular, of 0.1 to 0.13. Alkali metal salts are preferably used, in particular sodium chloride present in an amount imparting to the albumin solution an ionic strength of 0.15.

15 Although any albumin may be suitably used as starting material, its origin may be a factor affecting the nanofiltration yield, depending on whether it contains thermal stabilisers or was heat-treated (thermal shock or pasteurisation). Thus, in the method according 20 to the invention, using an albumin obtained by ethanol extraction according to Cohn et al. or Kistler et al. as mentioned above, and purified by ion-exchange or affinity chromatography, can result in an increased albumin recovery and/or a reduced filtration duration.

25 The nanofiltration of the aqueous albumin solution can be carried out in two successive steps on two filters with decreasing porosities, respectively. Advantageously, these two successive steps are carried out on filters with porosities of 23 to 50 nm and 15 or 20 nm, 30 respectively.

When PLANOV<sup>A</sup> filters with porosities of 15 nm and surface areas of 0.01 m<sup>2</sup> are used, the method of the invention is implemented at pressures not exceeding 1 bar, preferably in the range of 0.2 to 0.8 bar.

5 The method may comprise a subsequent step of known specific process intended to make the aqueous albumin solutions suitable to various therapeutic uses in accordance with the European Pharmacopoeia, such as their adjustment to the physiological pH if appropriate, to the 10 isotonicity in the case of an intravenous injection and to a physiologically acceptable salt concentration, and/or conditioning, for example in freeze-dried or liquid form.

15 The invention also relates to a virally safe aqueous albumin solution obtainable by implementing the method of the invention, in which the transport and binding sites of therapeutically active ingredients are available in the albumin. This albumin solution is further characterised in that it is free from macromolecules 20 having a sedimentation constant, according to Svedberg Ph. et al. ("Ultracentrifüg", 7th edition, Ed. Steinkopff, Dresden, 1940) above 7 S (i.e. a molecular weight of about 160 kDa). In particular, this albumin solution contains at most 1% albumin polymers with a size 25 smaller than 100 nm, preferably smaller than 20 nm.

The invention further relates to an albumin composition for therapeutic use obtained by processing said albumin solution according to the invention to make it suitable to a clinical use.

30 The safety achieved with these albumin compositions for therapeutic use makes it possible to suppress the

pasteurisation step, a source of drawbacks as mentioned above, and therefore, to add usual protection stabilisers against thermal effects, which also bind on the albumin sites, thus preventing albumin from binding the relevant 5 molecules. The albumin of these compositions according to the invention retains its binding and transport potential of various active ingredients, and through this binding, reduces their toxicity or increases the bioavailability by a depot effect.

10 One can also mention other applications in which the albumin compositions according to the invention may be used:

- stabilisation against denaturation and oxidation of proteins in low concentrations (lower than 1 g/L) 15 and with high specific activities, such as factor VIII, the von Willebrand factor and their recombinant equivalents, specific immunoglobulins, monoclonal antibodies, vaccines (viral, bacterial, proteinic, DNA vaccines), allergens for the 20 detection of allergies in sensitive patients, cytokines or peptidic hormones;
- excipient for proliferation and incubation media for *in-vitro* fertilisation of human oocytes; and
- control standard proteins acting as placebos in 25 clinical studies.

The invention will become more apparent upon review of the following examples, with reference to the accompanying drawings in which:

- Figure 1 shows a device for implementing the method 30 of the invention, and

- Figures 2 to 6 are plots showing the variations of the instantaneous filtration flow rate (mL/min) versus the nanofiltration duration, obtained on the basis of various parameters.

5

#### Example 1

##### 1.1 Methodology for implementing the nanofiltration method (Figure 1)

A filtration device 1 containing a PLANNOVA® 15-nm filter with a surface area of 0.01 m<sup>2</sup> (from Asahi, Japan) is equipped with tubes 10, 11 at the retentate (pre-filtration solution) outlet and at the filtrate (post-filtration solution) inlet and outlet, made of pharmaceutically compatible materials, with diameters of about 5 mm, closed by clamps. The device is arranged on a stand (not shown) in vertical position with the help of graspers.

The inlet of the 15-nm filter is connected, through the tube 10, to a pressure vessel 12 the pressure of which is measured with a digital pressure gauge 13 connected to the upstream filter circuit.

Before use, an integrity test is carried out on the 15-nm filter in accordance with the manufacturer's procedure: "Air leakage test during the integrity pre- and post-filtration test of PLANNOVA® 15, 35, 75-nm filters".

Then, the system is rinsed. For that purpose, the compressed air inlet 14 is connected to the pressure vessel 12 filled with a volume of about 100 mL NaCl at 30 9 g/L. The flask is pressurised progressively until a pressure of 0.5 bar is reached at the inlet of the 15-nm

filter. The 15-nm filter is filled while air is released at the retentate outlet, without filling the outside of the filter fibres. The lower outlet is open and connected to an optical density measurement detection cell 15 5 connected to a recording device 16, the top filtrate outlet remaining clamped. The rinse filtrate is collected in a container 17 on a weighing scale 18 connected to and controlled by a microcomputer 19 that records the filtrate weight increase, which makes it possible to 10 monitor the instantaneous filtration flow rate. The necessary time to filter a minimum volume of 40 mL at 0.5 bar is measured, and the average flow rate in mL/min is deduced. The system is progressively depressurised and 15 all the outlets are clamped. Depending on the temperature required for the nanofiltration, the system can be at room temperature (about 20°C) or placed in an oven in which the working temperature is between 25 and 60°C.

### 1.2 Albumin solutions

20 As reference aqueous albumin solutions, solutions at 20 g/L, pH 7, containing 9 g/L NaCl are used. The albumin starting material used, noted A, except for Examples 6 and 9 below, is the one obtained by fractionation of 25 human plasma according to Kistler et al. and having undergone alcohol elimination process, diafiltration and concentration by ultrafiltration. It should be noted that the above fractionation of various sources of plasma generally results in albumin with variable characteristics due to the biological nature of the 30 starting material. This albumin may therefore influence the duration of the nanofiltration, the yield, etc.

Albumin aqueous solutions to be filtered are prepared from the reference solutions above, by modification of characteristics thereof such as concentration, pH and ionic strength. Their volumes are 5 adjusted so as to provide the desired protein loads. Depending on the particular example, the albumin concentrations vary from 15 g/L to 80 g/L. The pH is adjusted by adding 0.5 M NaOH or HCl and the ionic strength is adjusted by adding sodium chloride. Then, the 10 albumin solutions are all prefiltered on commercially available 0.2- $\mu\text{m}$  filters.

### 1.3 Nanofiltration of an albumin solution

Once the filtration of a predetermined volume of 15 albumin is achieved, the filter inlet is closed. For rinsing, one volume of purified water for injection (PWI) is introduced into the repressurised vessel 12, and the filter is flushed out. The filtrate is collected until a perceptible reduction in optical density which is 20 monitored on the recording device 16. The downstream circuit is then flushed out to collect the finished filtrate. Samples are taken from the filtrate and submitted to subsequent analytic tests such as polymer assays, virus titrations, etc. The duration of filtration 25 of one volume of albumin solution required to provide a fixed protein load is also measured, and the filtration yield is determined as the quotient between the albumin amount contained in the filtrate and in the retentate.

After use, an integrity test is carried out on the 30 15-nm filter in accordance with the manufacturer's procedure: "Air leakage test during the integrity pre-

and post-filtration test of PLANANOVA® 15, 35, 75-nm filters".

Example 2

5 Two preliminary tests are carried out to assess the feasibility of the nanofiltration in terms of protein load, duration and yield, when the nanofiltration is implemented with albumin solutions A (Example 1) filtered at 20°C and under 0.5-bar pressure. Volumes of 10 solutions A are prepared with protein loads of 0.5 kg/m<sup>2</sup> and 1 kg/m<sup>2</sup>. The results are shown in Table 1.

Table 1			
Solutions	Protein load (kg/m <sup>2</sup> )	Filtration duration (min.)	Protein recovery (%)
A	0.5	190	98
A	1	282	99

15 The above results show that, depending on the selected conditions, the nanofiltration of albumin solutions A provides a protein load of 1 kg/m<sup>2</sup> without filter clogging and with an excellent yield. It also appears that doubling the protein load does not result in 20 a nanofiltration duration twice as long.

The results of the tests have therefore prompted the Applicant to consider the nanofiltration of albumin solutions with a view to provide higher protein loads, while optimising some influencing parameters if needed.

Example 3

The dependence of the filtration durations and yields to the pH of the albumin solutions in a such a way as a protein load of 4 kg/m<sup>2</sup> is provided is determined 5 using a batch of albumin A as starting material. Five solutions of albumin at 40 g/L, A1 to A5, are prepared in a solution of NaCl at 9 g/L, and adjusted to pH 5, 7, 9, 9.5 and 10 respectively, with 0.5 M solutions of HCl or NaOH, and submitted to a nanofiltration. The 10 nanofiltration tests are carried out at 20°C at a pressure of 0.5 bar. Figure 2 shows the plots of filtration flow rate (mL/min) versus nanofiltration duration for each of the solutions considered.

An analysis of these plots shows, with solution A1 15 (pH 5), a rapid reduction in flow rate and, as a consequence, filter clogging. With solutions A2 and A3, there is a reduction in the initial flow rate which is faster with solution A3, but with quite acceptable variations, no filter clogging being observed after more 20 than 25 hours of filtration. The best results are obtained with solutions A4 and A5, i.e. at pH 9.5 and 10, with higher respective initial flow rates and shorter filtration durations than with solutions A2 and A3, solution A4 showing the most optimal initial flow rate.

25

Example 4

In this example, the joint effect of pH and temperature which provides a protein load of 4 kg/m<sup>2</sup> is tested on solutions A3 and A4, defined in Example 3, at 30 20°C and 30°C. The nanofiltration tests are carried out at a pressure of 0.5 bar. The filtration durations thus

obtained are listed in Table 2. Figure 3 compiles the plots of filtration flow rate (mL/min) versus nanofiltration duration for each of the solutions considered.

5

Table 2			
Solution	Temperature (°C)	Filtration duration (h)	Yield (%)
A3	20	11	98
A3	30	9.5	98
A4*	20	12.75 ± 0.75	98
A4	30	9	98

\*: Two nanofiltration tests

The shortest filtration durations are observed at 30°C with almost superimposable flow rate plots (solutions A3 and A4) without any progressive filter 10 clogging. At 20°C, the filtration flow rates are slightly lower than at 30°C.

#### Example 5

This example is intended to show the joint influence 15 of albumin concentration and temperature on nanofiltration in such a way as a protein load of 8 kg/m<sup>2</sup> is provided, by analysing the filtration durations obtained (Table 3) and the flow rate plots (Figure 4). In this series of tests, solutions A4 (40 g/L), A6 (60 g/L) 20 and A7 (80 g/L) are used, all of them containing 9 g/L NaCl, pH 9.5, and being submitted to a nanofiltration at a pressure of 0.5 bar.

Table 3			
Solutions	Temperature (°C)	Filtration duration (h)	Yield (%)
A4	20	24.6	98
A4	30	19.5	98
A6	20	22.5	98
A6	30	28.4	98
A7	20	31	98

An analysis of the results from Table 3 and Figure 4 shows that the filter can bear a load of 8 kg/m<sup>2</sup> without any clogging. In particular, optimal filtration conditions are obtained with solution A4 at 30°C. With solution A6, the initial filtration flow rate is higher at 30°C than at 20°C with, in addition, a longer filtration duration. These results have therefore prompted the Applicant to consider a nanofiltration temperature of 20°C for the most concentrated solution A7. Under these conditions, the filtration duration is the longest duration observed and the initial flow rate is the lowest flow rate observed.

15 Example 6

Three nanofiltration tests are carried out with solutions prepared from two different albumin starting materials. The first starting material is albumin A and the second one is albumin A purified by column ion-exchange chromatography (named A'). In this series of tests, solutions A4 and A6 from Example 5, and a solution A'1 (60 g/L, NaCl at 9 g/L, pH of 9.5), which are then submitted to a nanofiltration at 20 and 30°C at a

pressure of 0.5 bar, are used in order to provide a protein load of  $8 \text{ kg/m}^2$ . Table 4 shows the filtration durations and yields. Figure 5 shows the plots of filtration flow rate (mL/min) versus nanofiltration duration for each of the solutions considered.

Table 4			
Solutions	Temperature (°C)	Filtration duration (min)	Yield (%)
A4	30	18.5	98
A6	20	27.5	98
A'1	20	17	98

The shortest nanofiltration duration is that observed with A'1. However, the most optimal initial flow rate is observed with solution A4.

#### Example 7

The effect of a variation in ionic strength in the albumin solutions on the filtration durations and yields is studied with three albumin solutions A at 40 g/L, prepared in purified water for injection (solution A8), in a NaCl solution at 9 g/L (solution A2; ionic strength: 0.15) and in a NaCl solution at 30 g/L (solution A9; ionic strength: 0.5), respectively, all three being at pH 7. The nanofiltration tests are carried out at 20°C at a pressure of 0.5 bar so as to provide a constant protein load of  $4 \text{ kg/m}^2$ . The results are compiled in Table 5.

Table 5

Solutions	Concentration of NaCl (g/L)	Filtration duration (h)	Yield (%)
A8	0	24.5	98
A2	9	18.7	98
A9	30	22	98

The best result, in terms of optimised filtration duration and yield, is obtained with an ionic strength of 0.15, close to that corresponding to isotonicity. This 5 result is confirmed by monitoring the filtrate weight increase, which makes it possible to plot the variations in the filtration flow rate (mL/min) versus the nanofiltration duration for the studied solutions (Figure 6). In this example, it appears that increasing 10 the NaCl concentration to 30 g/L does not reduce the filtration duration.

Example 8

This example is intended to demonstrate the 15 influence of variations in protein load and nanofiltration pressure on the filtration durations and yields in the products obtained. For that purpose, two solutions of albumins A (Example 2) and A2 (pH 7) are used. The results are shown in Table 6.

Table 6				
Solutions (g/L)	Protein load (kg/m <sup>2</sup> )	Pressure (bar)	Filtration duration (min)	Yield (%)
A2	2	0.5	355	100
A	2	0.8	457	98
A2	4	0.5	1 350	98
A2	4	0.8	1 730	98

These results demonstrate that, with solutions A and A2, a pressure increase gives longer filtration durations in spite of the fact that solution A is less concentrated 5 than solution A2. The results obtained with solutions A2 show that the PLANOV<sup>A</sup> 15-nm filters have a protein load or capacity of 4 kg/m<sup>2</sup> and that a pressure increase does not decrease the filtration duration.

10 Example 9

This example is intended to demonstrate the possibility of implementing nanofiltration on an industrial scale by using PLANOV<sup>A</sup> 15-nm filters with surface areas of 1 m<sup>2</sup>. The albumin used is obtained by 15 fractionation according to the method of Kistler *et al.* and is set to therapeutic quality by heating at 60°C for 10 hours in the presence of a suitable stabiliser, in accordance with the requirements of the European Pharmacopoeia. The albumin B thus obtained is then mixed 20 with a buffer comprising: 0.01 M trisodium citrate, 0.12 M glycine, 0.016 M L-Lysine, 0.001 M calcium chloride and 0.17 M sodium chloride, pH 7-7.5. The final concentration of the albumin solutions is 20 g/L. After being prefILTERED on 0.2-μm filters, various solutions

thus obtained are submitted to a nanofiltration as described in Example 1, at room temperature and at a pressure of 0.5 bar. The volumes of solution B percolating through the filter are determined so as to provide a protein load of 120 g/m<sup>2</sup>. The same nanofiltration tests are carried out using filters of 0.01 m<sup>2</sup>. Table 7 compiles the results of filtration durations and yields obtained with both filters, as average values of 3 tests.

10

Table 7		
Solutions B	Filtration duration (h)	Yield (%)
Filter of 0.01 m <sup>2</sup>	3.1 ± 0.1	87 ± 4
Filter of 1 m <sup>2</sup>	2.5 ± 0.1	83 ± 1

15 The results obtained show that using filters with surface areas of 1 m<sup>2</sup> results in a shorter filtration duration compared to that obtained with filters having surface areas of 0.01 m<sup>2</sup>, but with a very slight decrease in filtration yield.

#### Example 10

20 The same solutions B and working conditions as in Example 9 are used to measure the reduction in polymer content after nanofiltration (Table 8).

Table 8

Solutions B	Polymer content before filtration (%)	Polymer content after filtration (%)
Filter of 0.01 m <sup>2</sup>	3.1	0.47 ± 0.13
Filter of 1 m <sup>2</sup>	4.3 ± 0.7	0.38 ± 0.05

It should be noted that the pasteurisation of albumin gives polymers in a percentage that, however, always remains lower than the 5% limit recommended by the 5 European Pharmacopoeia (4th edition, chapter "solution of human albumin", page 642). The nanofiltration tests carried out on the two filters surfaces considered provide an approximately 10-fold reduction in polymer content in both cases.

10

Example 11

To show the influence of variations in the working conditions of the nanofiltration (filter surface area, pressure and temperature) and of the physicochemical 15 parameters of the albumin solutions (pH, buffer composition), on the rate of viral reduction, viruses are introduced into different solutions of albumin A (20 g/L) which are submitted to nanofiltration tests. The tests are carried out by infecting the albumin solutions with 20 bacteriophage virus Phi-X 174, suspensions of which are obtained in accordance with AFNOR Standard NFT 72-181 (December 1989). This virus is a good marker for the 15- nm filter as its size is between 25 and 30 nm, which corresponds to the non-enveloped viruses that are 25 transmissible to humans, such as parvovirus B19, the

inactivation by pasteurisation of which is not satisfactory.

The results obtained are shown in Table 9.

Table 9

Filter surface area (m <sup>2</sup> )	Pressure (bar)	T (°C)	Osmolality (mosm)	Buffer composition	pH	Viral reduction (Δ log)
0.01	0.5	35	280	NaCl: 0.15 M	7.2	5.9
0.001	0.5	35	280	NaCl: 0.15 M	7.2	6.5
0.001	0.2	35	280	NaCl: 0.15 M	7.2	> 5.0
0.001	1.0	35	280	NaCl: 0.15 M	7.2	4.9
0.001	0.5	26	280	NaCl: 0.15 M	7.2	> 5.5
0.001	0.5	35	280	NaCl: 0.15 M	4.1	> 6.0
0.001	0.5	35	555	NaCl: 0.30 M	7.2	4.2
0.001	0.5	35	610	NaCl: 0.15 M; Sucrose: 100 g/L	7.2	> 6.3
0.001	0.5	35	335	Na <sub>2</sub> SO <sub>4</sub> : 0.15 M	7.2	3.7
0.001	0.5	35	280	NaCl: 0.15 M Tween <sup>®</sup> 80: 0.06%	7.2	> 6.5

An analysis of the results shows that, with variations in the working conditions:

- the sole variation in the filter surface area and temperature, whichever albumin solution considered, does not even nearly have an influence on the rate of viral reduction;
- a pressure increase from 0.5 bar to 1.0 bar has an unfavourable influence on the rate of viral reduction which, however, remains high (> 4 log).

Variations in the physicochemical parameters of the albumin A solutions show that:

- the presence of a detergent (Tween® 80) and a non-ionic compound (sucrose), and a variation in pH, do not even nearly have an influence on the rate of viral reduction;
- on the other hand, replacing NaCl with a divalent salt, sodium sulfate, has a negative influence on this rate which becomes lower than the threshold allowed by the control authorities. The same phenomenon is observed when the NaCl concentration is increased (from 0.15 M to 0.3 M), although the rate of reduction remains satisfactory.

#### Example 12

The properties of albumin A, nanofiltered without any stabiliser, in the transport and binding of medicines, are studied by comparing them with those obtained with two different albumin A batches pasteurised in the presence of sodium caprylate.

For that purpose, three solutions A4 (Example 3) are provided, one of which has been nanofiltered under the

conditions of the invention. These solutions are then processed into pharmaceutical formulations in a 0.07 M phosphate buffer, pH 7.4, to give albumin compositions, A'4, A"4 and A'''4, with concentrations of 2.5 g/L. The 5 albumin compositions A"4 and A'''4, not nanofiltered, are pasteurised in the presence of sodium caprylate, respectively. A sample of 1 mL is taken from each of the compositions, and a volume of between 10  $\mu$ L and 1 000  $\mu$ L of a parent alcohol-based solution of [ $^{14}$ C]warfarine and 10 [ $^{14}$ C]diazepam at 0.1 M respectively, two active ingredients of the respective class of anticoagulants and neurotropes, is added to the albumin compositions considered in order to provide variable concentrations in these active ingredients. The mixture is then 15 homogenised.

The study of the properties of albumin in binding the two above active ingredients is based on the methodology of equilibrium dialysis, known to those skilled in the art, and whose principle is summarised 20 below. A dialysis device including a cell with two compartments separated by a suitable dialysis membrane is obtained. One volume V of an albumin/active ingredient mixture is introduced into compartment 1 and the same volume V of dialysis buffer (phosphate buffer, as defined 25 above) is introduced into compartment 2. Once the equilibrium is reached, after a few hours, compartment 1 contains albumin-bound and unbound active ingredient and compartment 2 contains unbound active ingredient. Some samples are taken from each compartment 1 and 2 and 30 radioactivity is measured by liquid scintillation. These

measurements determine the respective concentrations in albumin-bound and unbound active ingredients.

Table 10 gives the percentages of albumin-bound  $[^{14}\text{C}]$ diazepam obtained with compositions A'4, A"4 and 5 A'''4 respectively, in which increasing volumes of active ingredient have been added. Table 11 shows the results obtained with  $[^{14}\text{C}]$ warfarine instead of  $[^{14}\text{C}]$ diazepam. The results are shown as the average value of 5 tests.

Table 10

Albumin compositions	Total concentration in $[^{14}\text{C}]$ diazepam ( $\mu\text{m}$ )	Percentage of albumin-bound $[^{14}\text{C}]$ diazepam
A'4	1.68 $\pm$ 0.06	80.9 $\pm$ 1.46
	4.17 $\pm$ 0.28	82.34 $\pm$ 1.92
	8.04 $\pm$ 0.24	78.08 $\pm$ 1.83
	40.09 $\pm$ 1.28	57.18 $\pm$ 2.14
	65.27 $\pm$ 3.46	48.58 $\pm$ 1.49
	115.02 $\pm$ 4.83	36.38 $\pm$ 1.69
A"4	1.09 $\pm$ 0.23	34.46 $\pm$ 2.04
	5.52 $\pm$ 0.53	32.26 $\pm$ 1.34
	21.95 $\pm$ 0.42	31.00 $\pm$ 0.93
	42.84 $\pm$ 5.67	31.85 $\pm$ 2.99
	85.00 $\pm$ 13.6	30.12 $\pm$ 2.00
	109.79 $\pm$ 10.96	29.12 $\pm$ 2.67
A'''4	1.23 $\pm$ 0.09	38.47 $\pm$ 1.88
	2.91 $\pm$ 0.21	40.93 $\pm$ 4.83
	11.55 $\pm$ 0.19	37.85 $\pm$ 1.25
	44.98 $\pm$ 3.14	34.96 $\pm$ 0.6
	86.09 $\pm$ 5.01	33.04 $\pm$ 1.17
	107.97 $\pm$ 9.43	30.74 $\pm$ 1.78

The results show that the higher the concentration in  $[^{14}\text{C}]$ diazepam in the albumin composition A'4, the lower the percentage of albumin-bound  $[^{14}\text{C}]$ diazepam. This demonstrates the presence of one saturable site or bond 5 and one non-saturable bond in albumin.

Increasing the concentration in  $[^{14}\text{C}]$ diazepam in the albumin compositions A"4 and A'''4 results in appreciably constant percentages of albumin-bound  $[^{14}\text{C}]$ diazepam. Therefore, the binding site identified in the albumin A, 10 nanofiltered without any stabiliser, in composition A'4 is no more functional since it is occupied by the stabiliser.

Table 11

Albumin compositions	Total concentration of [ $^{14}\text{C}$ ]warfarine ( $\mu\text{m}$ )	Percentage of albumin-bound [ $^{14}\text{C}$ ]warfarine
A'4	1.99 $\pm$ 0.19	93.32 $\pm$ 0.49
	10.12 $\pm$ 0.71	91.75 $\pm$ 0.75
	34.29 $\pm$ 2.28	83.43 $\pm$ 1.11
	62.73 $\pm$ 3.55	71.74 $\pm$ 1.20
	103.66 $\pm$ 4.24	55.34 $\pm$ 2.94
	123.43 $\pm$ 8.07	47.83 $\pm$ 2.03
A"4	1.71 $\pm$ 0.19	88.18 $\pm$ 0.82
	8.35 $\pm$ 1.13	87.61 $\pm$ 0.84
	30.97 $\pm$ 2.98	81.14 $\pm$ 2.07
	54.40 $\pm$ 10.02	73.24 $\pm$ 3.46
	102.08 $\pm$ 14.70	57.02 $\pm$ 4.36
	119.90 $\pm$ 6.96	53.11 $\pm$ 3.46
A'''4	1.69 $\pm$ 0.08	88.58 $\pm$ 0.32
	8.33 $\pm$ 0.30	87.66 $\pm$ 0.28
	31.65 $\pm$ 0.72	80.39 $\pm$ 0.89
	57.17 $\pm$ 3.81	71.01 $\pm$ 1.68
	104.64 $\pm$ 6.28	55.04 $\pm$ 0.68
	127.98 $\pm$ 4.83	49.77 $\pm$ 2.66

The results from this table show that the percentages of albumin-bound [ $^{14}\text{C}$ ]warfarine vary with the increase in [ $^{14}\text{C}$ ]warfarine concentration in the three albumin compositions considered. Therefore, the binding site in the albumin is not saturated by the stabiliser.

A comparison of the results shown in Tables 10 and 11 demonstrates that albumin behaves differently in respect of the considered medicines. This may be due to the structures of the medicines considered, compared with

the configurations of the albumin binding sites. These studies also determined the various binding constants ( $K_a$ ) of both medicines on the different albumins A'4, A"4 and A'"4, as shown in Table 12.

5

Table 12		
Albumin composition	$K_a$ [ $^{14}\text{C}$ ]diazepam (mM $^{-1}$ )	$K_a$ [ $^{14}\text{C}$ ]warfarine (mM $^{-1}$ )
A'4	187	215
A"4	4.2	135
A'"4	3.35	118

Example 13

The power of the nanofiltered albumin to stabilise von Willebrand factor (vWF) was assessed using dry heating tests of vWF at 80°C for 72 hours. On the basis of preliminary tests, it was found that adding a mixture of judiciously selected excipients, such as glycine and arginine, to a solution of vWF, provided its stabilisation during the dry heating test (80°C, 10 72 hours). Although the activity of vWF (vWF:Ag) was retained (> 82% compared to the non-heated vWF), its multimeric profile was not maintained, which was revealed by the disappearance of multimers with high molecular weights (> 10), responsible for the therapeutic effect of 15 vWF.

20

To overcome this drawback, three solutions of 1 g/L of vWF were prepared in the presence of glycine and arginine, and of nanofiltered albumin A for therapeutic use. The characteristics of the solutions thus obtained, 25 A10, A11 and A12, are shown in Table 13.

Table 13

Solutions	Albumin (g/L)	Glycine (g/L)	Arginine (g/L)
A10	15	5	40
A11	15	10	40
A12	15	5	30

These solutions are then submitted to freeze-drying (45 ± 3 hours) and dry heating as defined above. After 5 heating, the addition of purified water for injection (PWI) to the dry residues reconstitutes solutions A11-A12. The following parameters are analysed: appearance of the solution, dissolution time, activity of vWF and multimeric profile, as measured by electrophoresis. 10 Table 14 shows the results obtained.

Table 14

Solutions	Solution appearance	Dissolution time (min)	vWF:Ag (%)**	Multimeric profile
A10	L*	< 1	91	C***
A11	L*	< 1	89	C***
A12	L*	< 1	87	C***

\*: clear, without visible particles

\*\*: ratio between the activity of heated vWF and non-heated vWF

\*\*\*: conform to the original product (before heating)

These results demonstrate on the one hand that the presence of albumin does not have any negative effect on 15 the stabilisation by heating of vWF, which is revealed by a clear solution appearance (without visible particles),

a very short solubilisation time, and in particular, the preservation of vWF activity, and on the other hand, that this presence also maintains the integrity of its high molecular weight multimers.

CLAIMS

1 A method for purifying albumin, characterised in that it comprises a step consisting in subjecting an aqueous solution of albumin, having a concentration of from 15 g/L to 80 g/L and a pH not less than 7, to nanofiltration in a temperature range from 15°C to 55°C.

2. A method according to Claim 1, characterised in that the nanofiltration is carried out on a filter defined for a porosity of at least 13 nm.

3. A method according to either of Claims 1 and 2, characterised in that the pH of the aqueous solution of albumin is within the range of values from 7.8 to 11.5, and preferably from 9 to 10.5.

4. A method according to any of Claims 1 to 3, characterised in that it also comprises a step of addition of a pharmaceutically acceptable salt or mixture of pharmaceutically acceptable salts to the aqueous solution of albumin so as to obtain a solution with an ionic strength within the range of from 0.01 to 0.55.

5. A method according to Claim 4, characterised in that the pharmaceutically acceptable salt is a salt of an alkali metal.

6. A method according to Claim 5, characterised in that the salt of an alkali metal is sodium chloride

present in an amount that confers on the solution of albumin an ionic strength of 0.15.

7. A method according to any one of Claims 1 to 6, 5 characterised in that the concentration of the aqueous solution of albumin is within the range of from 40 g/L to 60 g/L.

8. A method according to any one of Claims 1 to 7, 10 characterised in that the temperature of the aqueous solution of albumin is from 30°C and 55°C.

9. A method according to any one of Claims 1 to 8, 15 characterised in that the nanofiltration of the aqueous solution of albumin is carried out in two successive steps on respectively two filters of decreasing porosities.

10. A method according to Claim 9, characterised in 20 that the two successive nanofiltration steps are carried out respectively on filters with a porosity of from 23 to 50 nm and 15 or 20 nm.

11. A method according to any one of Claims 1 to 25 10, characterised in that it is carried out with 15 nm regenerated cellulose filters having a surface area of 0,01 m<sup>2</sup>, at a pressure no greater than 1 bar.

12. A method according to Claim 11, characterised 30 in that the pressure is within the range of from 0.2 to 0.8 bar.

13. A method according to any one of Claims 1 to 12, characterised in that the albumin is obtained by ethanolic extraction and/or by purification by ion-exchange or affinity chromatography.

14. A method according to any one of Claims 1 to 13, characterised in that it comprises a subsequent step of treatment for adapting the aqueous solution of albumin to therapeutic use.

15. A virally safe aqueous solution of albumin that can be obtained by carrying out the method according to any one of Claims 1 to 14, in which the sites for transport and binding of therapeutic active ingredients of the albumin are available.

16. An aqueous solution of albumin according to Claim 15, characterised in that it contains at most 1% of albumin polymers less than 100 nm in size.

17. An aqueous solution of albumin according to Claim 15 or 16, characterised in that it contains at most 1% of albumin polymers less than 20 nm in size.

25  
18. An albumin composition for therapeutic use, obtained by a treatment according to Claim 14, for adapting an aqueous solution of albumin, according to any one of Claims 15 to 17, to clinical use.

19. The use of an albumin composition for therapeutic use according to Claim 18, for stabilising at least one member chosen from the group consisting of proteins in low concentrations and high specific 5 activity, specific immunoglobulins, monoclonal antibodies, vaccines, allergens, cytokines and peptide hormones.

20. The use according to Claim 19, characterised in 10 that the proteins represent factor VIII or von Willebrand factor, and their recombinant equivalent.

21. The use of an albumin composition for therapeutic use according to Claim 18, for the transport 15 and binding of therapeutic active ingredients.

22. The use of an albumin composition for therapeutic use according to Claim 18, as an excipient for an incubation medium for "in-vitro" fertilisation of 20 human oocytes.

23. The use of an albumin composition for therapeutic use according to Claim 18, as a control standard protein.

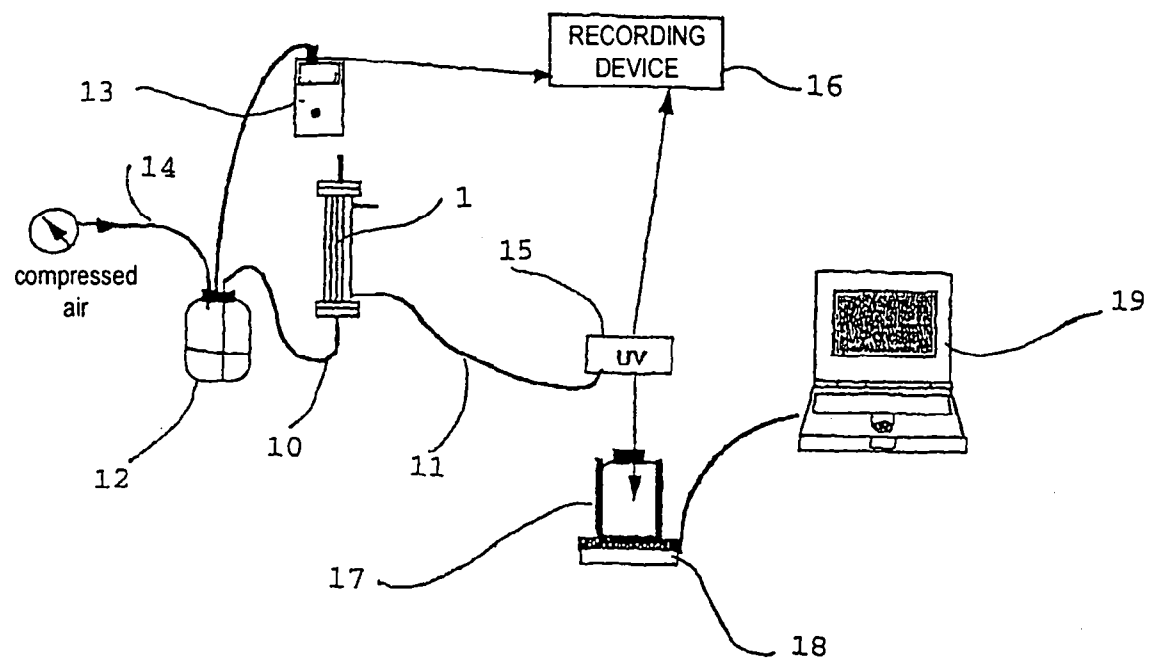


FIGURE 1

FIGURE 2

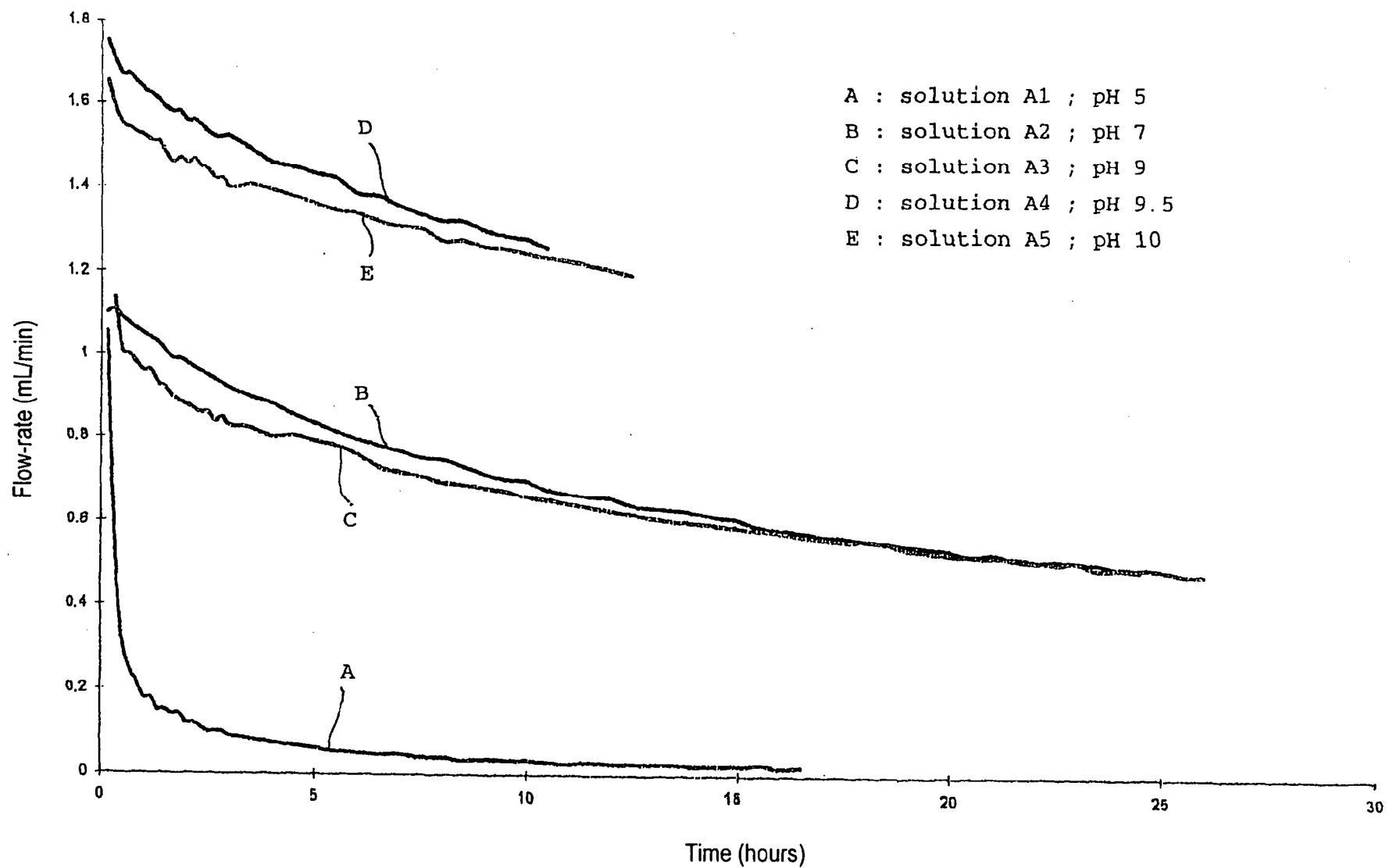


FIGURE 3

A : solution A4 ; 30°C  
B : solution A3 ; 30°C  
C : solution A3 ; 20°C  
D : solution A4 ; 20°C

3/6

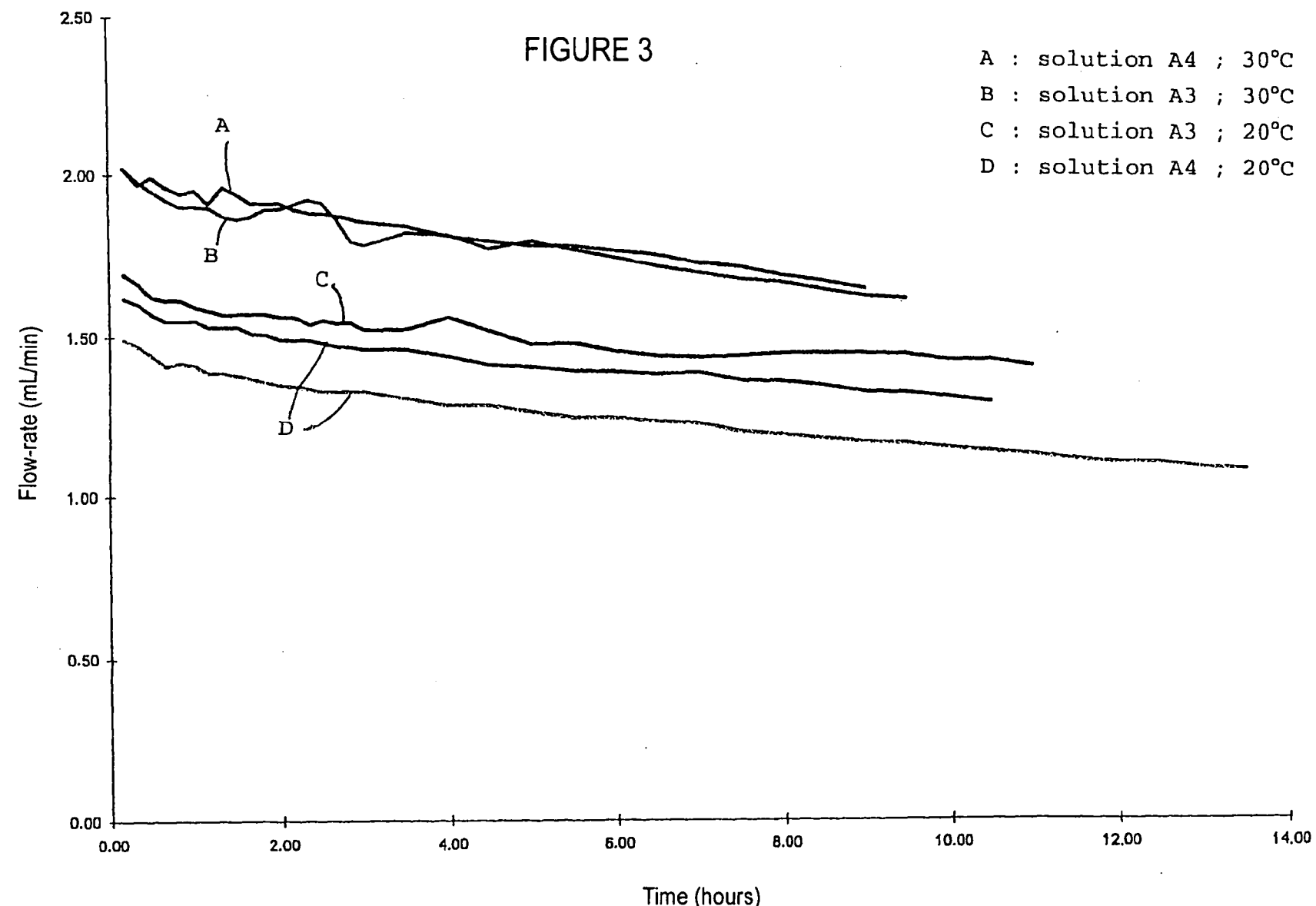


FIGURE 4

A : solution A4 ; 20°C  
B : solution A4 ; 30°C  
C : solution A6 ; 20°C  
D : solution A6 ; 30°C  
E : solution A7 ; 20°C

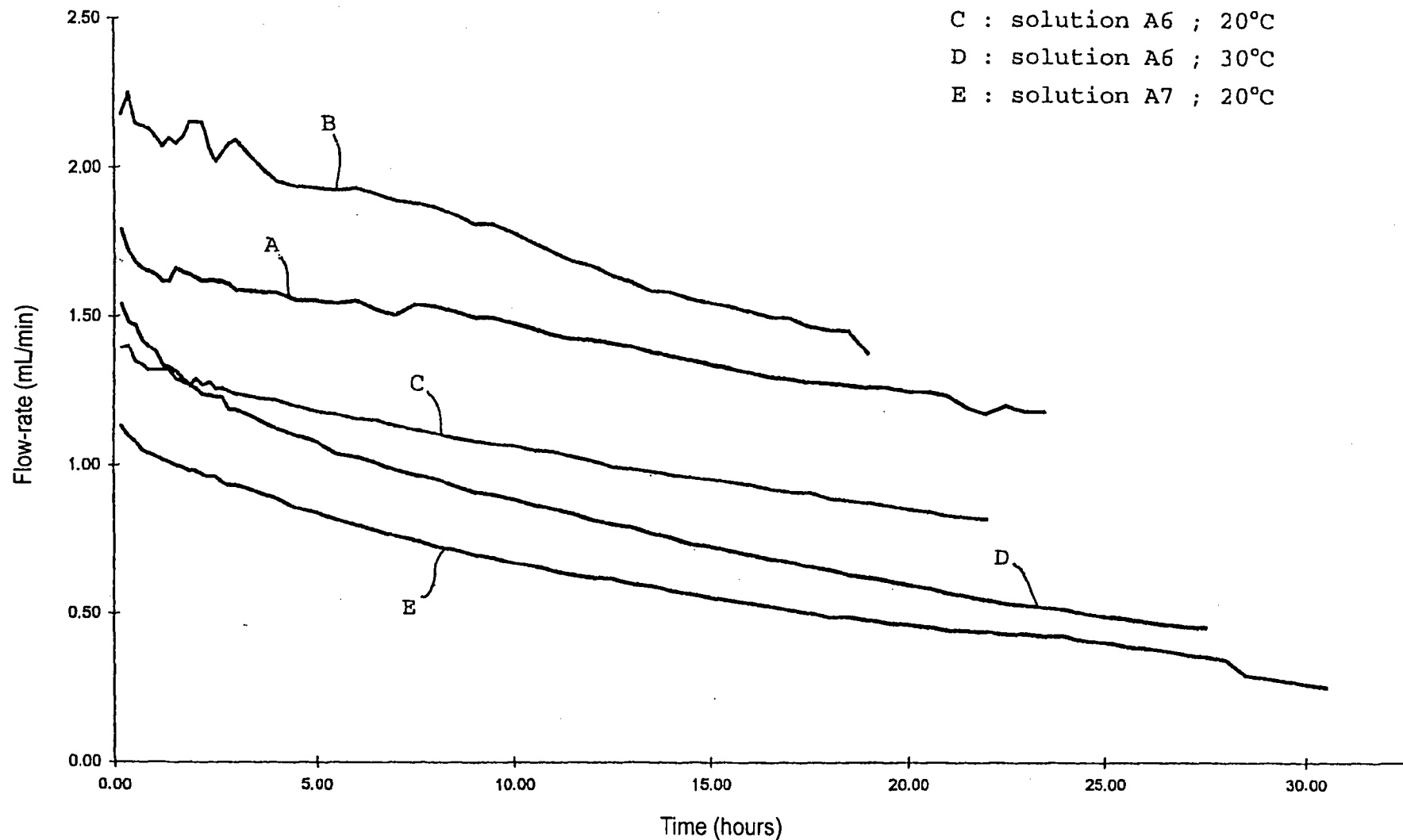


FIGURE 5

A : solution A4 ; 30°C  
B : solution A6 ; 20°C  
C : solution A'1 ; 20°C

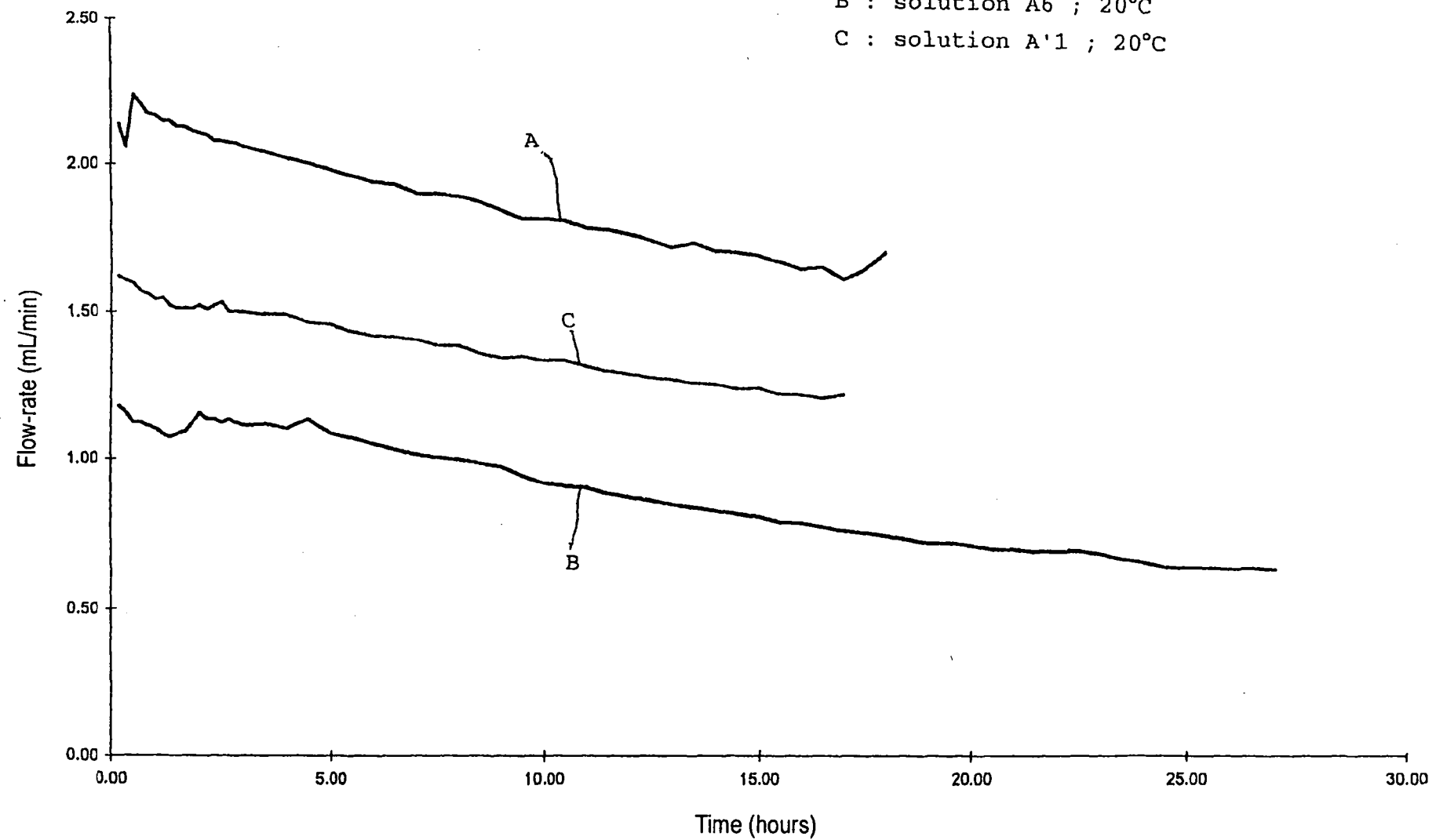


FIGURE 6

