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(54) Column for capillary chromatographic separations and method of manufacturing same
Kapillarsäule für chromatographische Trennungen und Verfahren zur Herstellung
Colonne capillaire pour séparations chromatographiques et procédé de fabrication

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(73) Proprietor: Agilent Technologies, Inc. (a Delaware corporation)
Palo Alto, CA 94303 (US)

(72) Inventors:
• Unger, Klaus K., Prof. Dr. 64342 Seeheim (DE)
• Adam, Thomas 66740 Saarlouis (DE)
• Rozing, Gerard, Dr. 76228 Karlsruhe (DE)
• Dittmann, Monika, Dr. 76359 Marxzell (DE)

(74) Representative: Harbach, Thomas
Agilent Technologies Deutschland GmbH
Patentabteilung
Herrenbergerstrasse 130
71034 Böblingen (DE)

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Description

FIELD OF THE INVENTION

[0001] The invention relates to a column for capillary chromatographic separations, for example for high performance liquid chromatography, or capillary electrophromatography, or supercritical chromatography. The invention also relates to a method for manufacturing such a column.

BACKGROUND OF THE INVENTION

[0002] Capillary chromatographic separation methods are preferably performed in fused silica (FS) tubing with internal diameters ranging from 5-530 μm. Such tubing consists of a silica (SiO2) glass drawn at high temperature (1300°C) from a quartz preform provided with a protective outside layer from polyimide or aluminum. Robustness, tensile strength, high pressure resistance, and bend stability are favorable mechanical properties of FS tubing. High chemical purity and well defined surface of the tubing provides in most cases low interaction with solutes and leads to optimum separation in many applications.

[0003] In US Patent 4,293,415 Dandeneau et al. describe the usage of a fused silica capillary, which may have wall coatings on the inside surface to stimulate specific interactions and/or further minimize secondary undesired solute/surface interactions, for open tubular capillary gas chromatography (CGC) and open tubular supercritical fluid chromatography (SFC). Jorgenson et al. (Anal. Chemistry, 1981, 53, p.1298) have demonstrated that such capillaries are also ideally suited for the new technique of capillary electrophoresis (CE).

[0004] It has been demonstrated that FS tubing can also be used for capillary separations performed in a packed bed, such as SFC, μ-HPLC and capillary electrophromatography (CEC). The mechanical properties of fused silica capillaries suffice to withstand the high pressure that occurs either when packing the tubing with small particles using a high pressure filtration technique or when operating the column especially in HPLC mode.

[0005] In FS (or other small i.d.) tubing the packing material in the column bed needs to be retained in the tubing; otherwise hydraulic or electrical forces drive the particles out of the capillary column. This is in most cases achieved by porous frits that are formed in the capillary by different processes.

[0006] In recent publications frits have been formed from the stationary phase particles directly by application of heat to a zone of the packed fused silica column where the terminating frit needs to be (e.g., Boughtflower et al., Chromatographia 40, 329 (1995), Smith et al., Chromatographia 38, 649 (1994), Rozing et al., LC-GC Magazine, October 1995). It is believed that under these conditions the particles are glued together by the fact that upon heating a small amount of silica dissolves in water forming silicic acid, and that upon cooling the re-polymerized silicic acid deposits between the particles. The advantage of this approach is that it does not substantially alter the chemical constitution of the zone that is fritted, that it can be done on the inlet and outlet side without problem, that the length of the fritted zone is well controlled by the dimension of the external heating source used and that the porosity of the bed is unaffected. Photographs e.g. by Boughtflower et al., show that the particle structure is not affected by this treatment and therefore inter-particle porosity is maintained.

[0007] The main problem with all these approaches is that although the packing is in principle retained between the frits, the stationary phase particles still have the ability to move or rearrange within the boundaries determined by the frits. It has been observed that the stationary phase particles in packed capillaries rearrange during standard operation of such a capillary, resulting in formation of voids (unpacked stretches) between the retaining frits. The reason for this are the electric and/or hydraulic forces that act upon the particles during column operation, which lead to changes in packing density. These voids lead to chromatographic artifacts like loss of efficiency, tailing peaks etc.

[0008] CH-A-427351 discloses a method for manufacturing a separation column for gas chromatography, wherein the inner wall of a capillary is coated with stationary phase material. During the manufacturing process, the stationary phase material is filled into a capillary, and the filled capillary is then drawn through a tube furnace.

[0009] DE-A-1673304 discloses a method for manufacturing a chromatographic column wherein silica gel and polyethylene powder are mixed and then sintered in a glass tube to produce a sintered column.

SUMMARY OF THE INVENTION

[0010] It is thus an object of the present invention to provide a method of manufacturing a packed column for capillary chromatography wherein the packing material is immobilized over the whole length of the column.

[0011] In particular, it is an object of the invention to avoid or reduce the above mentioned problems associated with rearrangement of particles in the packing and subsequent formation of voids or unpacked stretches.

[0012] According to the invention these objects are achieved by a method according to claim 1. A column for capillary chromatographic separations in high performance liquid chromatography or capillary electrophromatography manufactured according to the invention comprises a column bed of packing material arranged in the inner bore of the column wherein the packed bed is completely immobilized by a thermal treatment described below.

[0013] According to a preferred embodiment of the present invention, the mentioned prior art problems are circumvented in the following way. The fused silica tub-
ing that is used for preparation of a micro high performance liquid chromatography or CEC is packed in the usual way (see e.g. Boughtflower et al., Rozing et al.). After the column packing is finished, instead of preparing the retaining frits, the complete packing bed is immobilized by a thermal treatment. This is done the following way: A coiled heating wire with the capillary to be treated in the center of the coil is slowly moved along the column. During this process pressure is applied to the inlet side of the column to generate a flow through the packed bed. Under these conditions the immobilization takes place due to the following mechanism. At the elevated temperatures induced by the heating coil a small amount of silica is dissolved to form silicic acid. The silicic acid molecules are then transported by the hydraulic flow to a colder region of the packing where they repolymerize and deposit between the stationary phase particles acting as a glue.

BRIEF DESCRIPTION OF THE DRAWINGS

[0014] Figure 1 is a schematic diagram of an embodiment of the invention.

[0015] Figure 1 shows a packed column 1 according to a first embodiment of the invention. The column is made of fused silica and typically has a length of about 5-200 cm. The column has an interior diameter in the range between about 5-530 micrometers. The interior of the column is filled over the major part of its length with a packed bed 2 which serves for separating the sample substances passing through the column. The immobilization of the packing bed is achieved by moving the heating coil 3 which is connected to a power supply 4 along the column 1 with an appropriate speed. During this process a liquid 5 (preferably water) is pumped through the column using a high pressure pump 6. The velocity with which the heating coil is moved along the column and the liquid flow rate through the packing bed depend on the nature of the packing material to be immobilized and have to be determined experimentally. A typical velocity for the heating coil would be 2 mm/s and a typical liquid flow velocity would be 1-2 mm/s. The temperature of the heating coil is chosen such that the temperature inside the capillary is about 300 - 400 °C. In the process described the polyimide coating on the outside of the capillary is not removed or destroyed. This is important to retain the mechanical strength of the capillary.

[0016] Figure 2 shows an electromagnetic picture of the immobilized zone demonstrating that the structure of the packing particles is unaltered and that the permeability of the packed bed is unchanged. This factor is important for the chromatographic performance of the column. Chromatographic experiments have shown that the retention properties of the stationary phase are not altered by the immobilization process.

[0017] Figure 3 shows chromatograms of a standard sample on a packed capillary before (a) and after (b) immobilization.

[0018] According to the invention it is thus possible to generate an immobilized packed bed in which the packing particles are “glued” together without altering the mechanical, chemical and chromatographic properties of the packing. This results in better long term stability of the chromatographic bed.

Claims

1. A method of manufacturing a column for capillary chromatographic separations in high performance liquid chromatography or capillary electrochromatography, comprising the steps of:

- filling stationary phase packing material into an inner bore of a capillary for forming a packed column bed of a determined length in the capillary,
- applying a thermal treatment to the stationary phase packing material, characterized in that

- the thermal treatment is applied locally, moving a heating element along the entire length of the packed column bed in the capillary to immobilize all of the stationary phase packing material (2), and
- during the thermal treatment liquid is pumped through the packed column bed by applying pressure to an inlet side of the column (1).

2. Method as in claim 1, wherein the heating element is a heating wise and the thermal treatment comprises moving said heating wire (3) along the column (1).

3. Method as in claim 2, wherein

- the heating wire is an electrical heating coil (3) with the column (1) being arranged in the center of the coil, and
- the heating coil is moving at velocities of 0.5-5
mm/s opposite to the direction of the applied liquid flow moving at 0.5-5 mm/s under a linear pressure gradient over the bed ranging from 100-1000 bar to atmospheric pressure.

4. Method as in any of the preceding claims, wherein the packing material (2) consists of silica based particles with a coating suited for high pressure liquid chromatography or capillary electrophoresis.

5. Method as in any of the preceding claims, wherein the inner bore of the capillary (1) has a diameter between approximately 5 and 530 micrometers, and wherein the length of the capillary is between about 5 and 200 cm.

6. Method as in any of the preceding claims wherein the capillary (1) is made of fused silica.

Patentansprüche

1. Verfahren zur Herstellung einer Säule für kapillar-chromatographische Trennungen in der Hochleistungs-Flüssigkeitschromatographie oder der Kapillar-Elektrochromatographie, welches die folgenden Schritte umfasst:
   - Einfüllen von Packungsmaterial der stationären Phase in eine Innenbohrung einer Kapillare, um ein gepacktes Säulenbett mit einer bestimmten Länge in der Kapillare zu bilden,
   - Einwirken einer Wärmebehandlung auf das Packungsmaterial der stationären Phase, dadurch gekennzeichnet, dass
     - die Wärmebehandlung örtlich erfolgt, indem ein Heizelement über die gesamte Länge des gepackten Säulenbettes in der Kapillare geführt wird, um das gesamte Packungsmaterial der stationären Phase (2) zu immobilisieren, und
     - während der Wärmebehandlung durch Ausüben von Druck auf eine Einlassseite der Säule (1) Flüssigkeit durch das gepackte Säulenbett gepumpt wird.

2. Verfahren nach Anspruch 1, wobei das Heizelement ein Heizdraht ist und die Wärmebehandlung die Bewegung des Heizdrahtes (3) entlang der Säule (1) umfasst.

3. Verfahren nach Anspruch 2, wobei
   - der Heizdraht eine elektrische Heizspule (3) ist und die Säule (1) in der Mitte der Spule angeordnet ist, und
   - sich die Heizspule mit Geschwindigkeiten von 0,5 bis 5 mm/s entgegen der Richtung des mit einer Geschwindigkeit von 0,5 bis 5 mm/s unter einem linearen Druckgradienten von 100 bis 1000 bar bis zu Atmosphärendruck über das Bett strömenden Flüssigkeitsstroms bewegt.

4. Verfahren nach einem der vorangehenden Ansprüche, wobei das Packungsmaterial (2) aus siliciumdioxidbasierten Partikeln mit einer Beschichtung besteht, welche für die Hochdruck-Flüssigkeitschromatographie oder die Kapillar-Elektrochromatographie geeignet ist.

5. Verfahren nach einem der vorangehenden Ansprüche, wobei die Innenbohrung der Kapillare (1) einen Durchmesser zwischen ungefähr 5 und 530 µm aufweist und die Länge der Kapillare ungefähr zwischen 5 und 200 cm beträgt.

6. Verfahren nach einem der vorangehenden Ansprüche, wobei die Kapillare (1) aus Quarzglas besteht.

Revendications

1. Procédé de fabrication d'une colonne pour séparations par chromatographie capillaire en chromatographie liquide haute performance ou en électrochromatographie capillaire, comprenant les étapes consistant à :
   - remplir l'orifice de passage interne d'un capillaire d'un matériau de garnissage pour phase stationnaire afin de former dans le capillaire un lit de colonne garnie de longueur déterminée,
   - appliquer un traitement thermique au matériau de garnissage pour phase stationnaire,
   caractérisé en ce que
   - le traitement thermique est appliqué localement, grâce au déplacement d'un élément chauffant le long de toute la longueur du lit de la colonne garnie dans le capillaire afin d'immobiliser tout le matériau de garnissage pour phase stationnaire (2), et
   - pendant le traitement thermique, on pompe du liquide à travers le lit de la colonne garnie en appliquant une pression du côté entrée de la colonne (1).

2. Procédé selon la revendication 1, dans lequel l'élément chauffant est un fil chauffant et le traitement thermique comprend le fait de déplacer ledit fil chauffant (3) le long de la colonne (1).

3. Procédé selon la revendication 2, dans lequel
le fil chauffant est un serpentin électrique chauffant (3), la colonne (1) étant placée au centre du serpentin, et

- le serpentin chauffant se déplace à des vitesses de 0,5-5 mm/s dans le sens opposé au sens de l'écoulement du liquide appliqué qui se déplace dans le lit à 0,5-5 mm/s sous un gradient de pression linéaire allant de 100-1 000 bars à la pression atmosphérique.

4. Procédé selon l'une quelconque des revendications précédentes, dans lequel le matériau de garnissage (2) se compose de particules à base de silice portant un enrobage approprié pour la chromatographie liquide haute pression ou l'électrochromatographie capillaire.

5. Procédé selon l'une quelconque des revendications précédentes, dans lequel l'orifice de passage interne du capillaire (1) a un diamètre compris entre environ 5 et 530 micromètres et dans lequel la longueur du capillaire est comprise entre environ 5 et 200 cm.

6. Procédé selon l'une quelconque des revendications précédentes, dans lequel le capillaire (1) est fait de silice fondue.
Fig. 1
Fig. 3

Chromatograms of a standard sample before and after immobilization of the packed bed.
1 thiourea, 2 methylparabene, 3 ethylparabene, 4 propylparabene, 5 butylparabene, 6 naphthalene,
7 fluorene, 8 phenanthrene, 9 anthracene, 10 fluoranthene