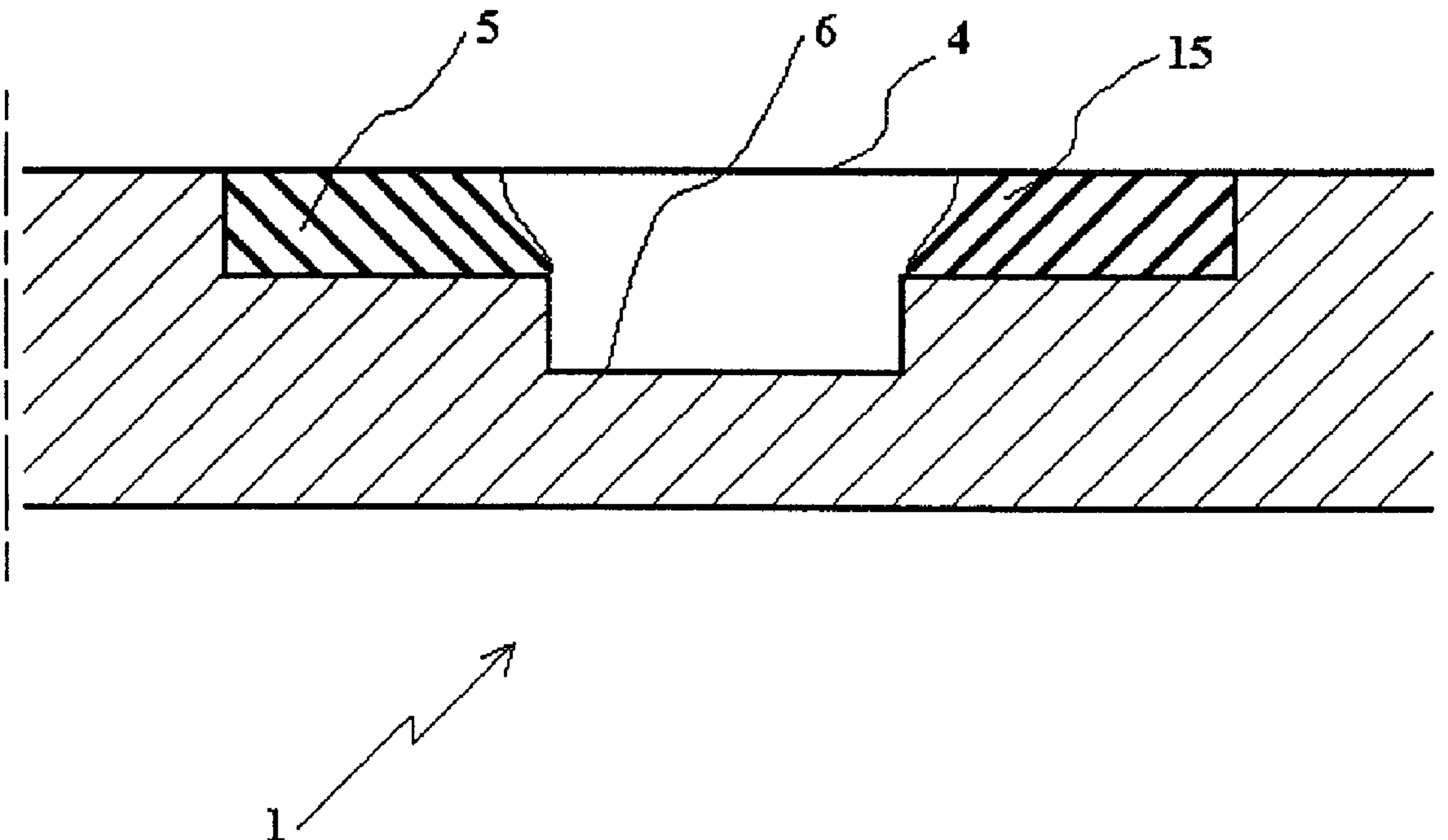




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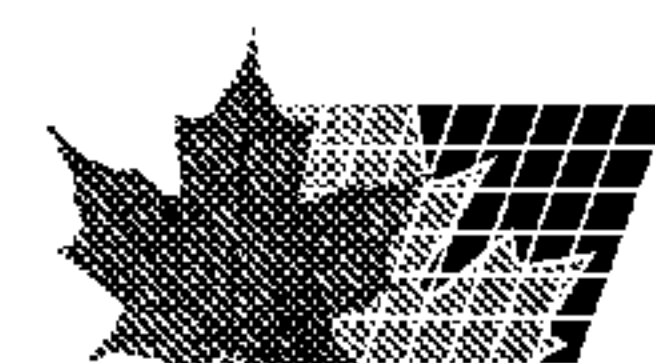
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(54) Titre : DISPOSITIF DE TEST POUR L'ANALYSE DES ANALYTES CHIMIQUES, BIOCHIMIQUES OU BIOLOGIQUES
(54) Title: TEST CARD FOR ANALYSIS OF CHEMICAL, BIOCHEMICAL OR BIOLOGICAL ANALYTES



(57) Abrégé/Abstract:

A test card comprising at least one planar surface wherein at least two compartments are made and defined by a partition comprising a film, the compartments creating a space which makes it possible to displace at least two liquid samples independently of one another, the compartments comprising at least two different types of grooves: at least one deep groove capable of partitioning samples from one another, the depth and the width of the deep groove in relation to the partition being such that capillary action of a sample is not enabled, and at least two shallow grooves, each of the shallow grooves being capable of receiving one of the samples, respectively, the depth of shallow grooves in relation to the partition being such that capillary action of the sample is enabled, each shallow groove being adjacent to the deep groove, along the entire length of the deep groove, wherein the test card is adapted for analysis of chemical, biochemical or biological analytes.



ABSTRACT

A test card comprising at least one planar surface wherein at least two compartments are made and defined by a partition comprising a film, the compartments creating a space
5 which makes it possible to displace at least two liquid samples independently of one another, the compartments comprising at least two different types of grooves: at least one deep groove capable of partitioning samples from one another, the depth and the width of the deep groove in relation to the partition being such that capillary action of a sample is not enabled, and at least two shallow grooves, each of the shallow grooves being capable
10 of receiving one of the samples, respectively, the depth of shallow grooves in relation to the partition being such that capillary action of the sample is enabled, each shallow groove being adjacent to the deep groove, along the entire length of the deep groove, wherein the test card is adapted for analysis of chemical, biochemical or biological analytes.

TEST CARD FOR ANALYSIS OF CHEMICAL, BIOCHEMICAL OR BIOLOGICAL ANALYTES

Field of the invention

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This invention concerns an apparatus wherein compartments are defined by a partition, thus creating a space in which at least one liquid sample can be displaced in a directed and independent fashion. When there are at least two liquid samples, they can both be displaced in an independent way and brought together so that they can react with one
10 another.

Background of the invention

Many documents in the background art deal with exploiting capillary action in fluid
15 micromanipulation applications. Thus, document GB-A-2.261.284 pertains to an apparatus for transferring liquids for the purposes of diagnostic testing. This apparatus is based on channels made of a porous material.

In this embodiment, the capillary action of a porous material is used. This requires the
20 incorporation of this porous material and it also necessitates having an impermeable separation between the two porous channels which both contain different liquids. In consequence, this method is fairly expensive to implement.

Patent U.S. Pat. No. 5,842,787 relates to fluid micromanipulation systems which include
25 channels of varying dimensions. It is essentially the depth of the channels which can be modified although such variation also affects width so that the deeper the channel, the smaller its width (and vice versa). Unfortunately, these channels are not open; in other words, the liquids which are to be transferred inside the channels normally occupy the entire cross-sectional volume. As a result there are strong retention forces which inhibit
30 the displacement of the liquids and therefore mean that sophisticated transfer systems are required (e.g. powerful pumps, the use of a vacuum, etc.).

In patent U.S. Pat. No. 5,660,993, capillary action is used to create a valve where two capillary channels meet. Apart from this novel function of opening up and shutting down

liquid flow, exactly the same problems are encountered as with the previous document-
because the channels are closed, retention is a problem.

According to documents EP-A-0.075.605 and WO-A-99/55852, shallow and deep
5 grooves are combined to direct liquids. However, there is no description of the use of any
physical property (neither capillarity nor any other physical phenomenon) in association
with the deep and shallow grooves, and no such association is obvious to those skilled in
the art.

10 **Summary of the invention**

In accordance with this invention, the apparatus proposed resolves all the problems
mentioned in that it uses capillary action to move liquids while, at the same time, it
minimizes retention phenomena. This makes for perfectly effective directed
15 displacement, even in the presence of a free space which means that the transferred liquid
is not physically confined.

The invention therefore provides a test card comprising at least one planar surface
wherein at least two compartments are made and defined by a partition comprising a film,
20 the compartments creating a space which makes it possible to displace at least two liquid
samples independently of one another, the compartments comprising at least two different
types of grooves: at least one deep groove capable of partitioning samples from one
another, the depth and the width of the deep groove in relation to the partition being such
that capillary action of a sample is not enabled, and at least two shallow grooves, each of
25 the shallow grooves being capable of receiving one of the samples, respectively, the depth
of shallow grooves in relation to the partition being such that capillary action of the
sample is enabled, each shallow groove being adjacent to the deep groove, along the
entire length of the deep groove, wherein the test card is adapted for analysis of chemical,
biochemical or biological analytes.

30

One deep groove may be located between two shallow grooves. In this case, the deep
groove has an end where the two shallow grooves meet to create a reaction zone.
According to a first embodiment, the distance between the reaction zone and the partition
or the partitioning film is such that capillary action is enabled.

According to a second embodiment, the distance between the reaction zone and the partition or the partitioning film is such that capillary action is not enabled.

Description of the figures

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The figures herewith are given by way of example and are not to be taken as in any way limiting. They are intended to make the invention easier to understand.

FIG. 1 shows an overhead view of the side of the apparatus with the compartment
10 according to the invention.

FIG. 2 shows a partial, transverse cross-section through A--A in FIG. 1.

FIG. 3 shows exactly the same view as FIG. 2 but with a liquid sample present.

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FIG. 4 shows exactly the same view as FIGS. 2 and 3 but with two different liquid samples present.

FIG. 5 shows a cross-section exactly like that in FIG. 2, but of a second embodiment
20 containing a liquid sample.

Finally, FIG. 6 shows a cross-section exactly like that in FIG. 2, but of a third embodiment of this invention containing a liquid sample.

25 Description of the embodiments of the invention

This invention relates to an apparatus (1) which is clearly illustrated in FIGS. 2 through 6 which are partial, transverse cross-sections through three different embodiments. Such an apparatus (1) can be used for the analysis of one or more different liquid samples to
30 identify one or more analytes, using any method, be it a simple or complex method and be it based on one or more different reagents, depending on the chemical, physical or biological nature of the analyte being tested. The technical principles defined hereafter are not restricted to any single, specific analyte; the only required condition is that the

analyte must either be dissolved or in suspension in the test sample. In particular, the test process being used can be performed on a homogenous, heterogeneous or mixed form.

One particular, non limited mode of such a device, concerns biological tests for the
5 detection and/or quantitative determination of one or more ligands, in which the assay involves one or more anti-ligands. The word ligand is taken to mean any biological species, e.g. an antigen, a fragment of an antigen, a hapten, a nucleic acid, a fragment of nucleic acid, a hormone or a vitamin. One example of an application of the test methods concerns immunoassays, whatever their particulars and whether the assay is direct or
10 based on competition. Another example of an application concerns the detection and/or quantitative determination of nucleic acids, including all operations required for such detection and/or quantitation in any kind of sample containing the target nucleic acid species. Among such diverse operations, the following could be specified: lysis, melting, concentration, enzyme-mediated nucleic acid amplification, and any detection modalities
15 which include a hybridization step using, for example, a DNA chip or a labeled probe. Patent application WO-A-97/02357 stipulates the various stages involved in the case of nucleic acid analysis.

In a particularly interesting embodiment shown in FIGS. 1 to 4, it can be seen that the
20 apparatus (1) actually consists of a card with two sides, an upper and a lower side which are parallel to one another. Of course, it does not necessarily have to be used in a horizontal position-it can also be used in a vertical position or on a slope.

In the figures, both sides are planar but it is the upper side which is of greater interest for
25 this invention. Thus, the upper planar surface (2) of the apparatus (1) includes cavities which create the compartments (3). The compartments are partitioned off with respect to the surfaces that are flush with the surface (2) by means of a film or partition (4). This compartment (3) thus isolated actually consists of a set of different forms. On the sides, there are two shallow grooves (16) and in the middle there is one deep groove (6). The
30 view in FIG. 2 corresponds to a partial cross-section through A--A in FIG. 1. From FIG. 1, it can be seen that the two shallow grooves (16) are parallel to one another for the entire length of the deep groove (6). However, one of the ends of the deep groove (6) has an end (7) and the two shallow grooves (16) meet there to create a reaction zone (8).

It is possible to isolate a first liquid sample (5) in one of the shallow grooves (16), as shown in FIG. 3. Similarly, it is possible to isolate another liquid sample (15) in the other shallow groove (16), as shown in FIG. 4. In practice, to insure that liquids (5) and (15) remain in position in the shallow grooves (16) without mixing, the distance separating the bottom of the groove (16) and the partitioning film (4) should be small enough for capillary action to take place. The ideal distance between the film (4) and the bottom of the groove (16) for capillary action is between 50 and 800 micrometers (μm) (preferably between 300 and 500 μm). In the case of an apparatus consisting of a card made of impact polystyrene and a BOPP film being used to transfer an aqueous solution containing 9 g/liter NaCl, 1 g/liter NaN_3 and 1 ml/l of either Tween 20 (registered trademark) or Triton X100 (registered trademark), the distance between the film (4) and the bottom of the groove (16) might be set at 400 μm . This dimension is actually typical for the kinds of liquid (5 and/or 15) which are likely to be used in this apparatus (1), given the materials used to make the apparatus (1). This distance may have to be varied for various reasons, e.g. depending on the viscosity, density, wetting activity and surface tension of the liquids being used, and on the hydrophilic/hydrophobic properties of the materials used to make the film and the card.

In contrast, the distance separating the film (4) from the bottom of the deep groove (6) must be great enough to insure that capillary action does not lead to the retention of liquid (5 or 15) here. Of course, it is obvious that the width value of this deep groove must be such that capillary action cannot take place.

The nature of the flexible film may vary according to the nature of the test card and of the fluids being tested, especially when compatibility is at issue. For example, TPX (polymethyl pentene copolymer) or BOPP (bi-oriented polypropylene) films are suitable for biological assays. These films can be fixed in place either using an adhesive (with the adhesive applied to the film, e.g. a silicon-based adhesive) or by heat-sealing. An example of an adhesive is available from BioMerieux Inc. (St. Louis, Mo., USA) (reference: 022004-2184).

In terms of production, the test cards are manufactured by the machining of special plastic material, e.g. impact polystyrene (reference: R540E from the Goodfellow company) which is compatible with the liquids being processed. For industrial-scale production, the

card could be manufactured by precision molding, but any other manufacturing method (including those used in the semi-conductor industry as stipulated in patent application WO-A-97/02357) may be used for test card production.

5 Of course, a number of other embodiments can be imagined and two of these are shown in FIGS. 5 and 6. That in FIG. 5 corresponds to a substantially reversed configuration of the first embodiment shown in FIGS. 1 to 4. Thus, in FIG. 5, one shallow groove (16) occupies the central position between two deep grooves (6). The liquid sample (5) is only in contact with the bottom of the shallow groove (16).

10

In another embodiment shown in FIG. 6, it is possible to have a single shallow groove (16) and a single deep groove (6).

Of course, all permutations are possible and can be imagined. For example, there might
15 be a whole series of deep grooves (6) or shallow grooves (16). The only prerequisite condition is that the deep grooves (6) be located between the shallow grooves, (16) or vice versa. Liquids (5 and/or 15) can be introduced by means of valves, pumps, and/or channels, as described in the patent applications submitted by the applicant on the same day with the following titles:

20

"A device and a method for positioning a liquid", for the first document,

"A pumping device for transferring at least one fluid into a consumable," for the second document, and finally

25

"A test sample card with improved filling" for the third document.

The liquids (5 and 15) can be moved in different ways, e.g. the card (1) could be made to vibrate or it could be placed in a substantially vertical position so that the liquids are
30 driven by the force of gravity; alternatively, centrifugal force could be used. Pumping systems could be used, either located inside or outside the card; these could be based on diaphragm pumps (U.S. Pat. No. 5,277,556), piezoelectric peristaltic pumps (U.S. Pat. No. 5,126,022), ferrofluid transport systems, or electric and hydrodynamic pumps

(Richter et al., Sensors and Actuators, 29, p159 165, 1991). Combinations of more than one of these types of system could also be used.

REFERENCES

1. Apparatus
- 5 2. Planar surface of the apparatus (1)
3. Compartments
4. Partition or partitioning film
5. First liquid sample
6. First type of groove, said to be deep
- 10 7. End of the groove (6)
8. Reaction zone
15. Second liquid sample
16. Second type of groove, said to be shallow

CLAIMS

1. A test card comprising at least one planar surface wherein at least two compartments are made and defined by a partition comprising a film, the
5 compartments creating a space which makes it possible to displace at least two liquid samples independently of one another, the compartments comprising at least two different types of grooves: at least one deep groove capable of partitioning samples from one another, the depth and the width of the deep groove in relation to the partition being such that capillary action of a sample is not
10 enabled, and at least two shallow grooves, each of the shallow grooves being capable of receiving one of the samples, respectively, the depth of shallow grooves in relation to the partition being such that capillary action of the sample is enabled, each shallow groove being adjacent to the deep groove, along the entire length of the deep groove, wherein the test card is adapted for analysis of
15 chemical, biochemical or biological analytes.
2. The test card of claim 1, wherein a deep groove is positioned between two shallow grooves.
- 20 3. The test card of claim 2, wherein the two shallow grooves meet at one of the ends of the deep groove to create a reaction zone there, where the two liquid samples may be brought together.
4. The test card of claim 3, wherein the distance between the reaction zone and the
25 partition is such that capillary action is enabled.
5. The test card of claim 3, wherein the distance between the reaction zone and the partition is such that capillary action is not enabled.

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PATENT AGENTS

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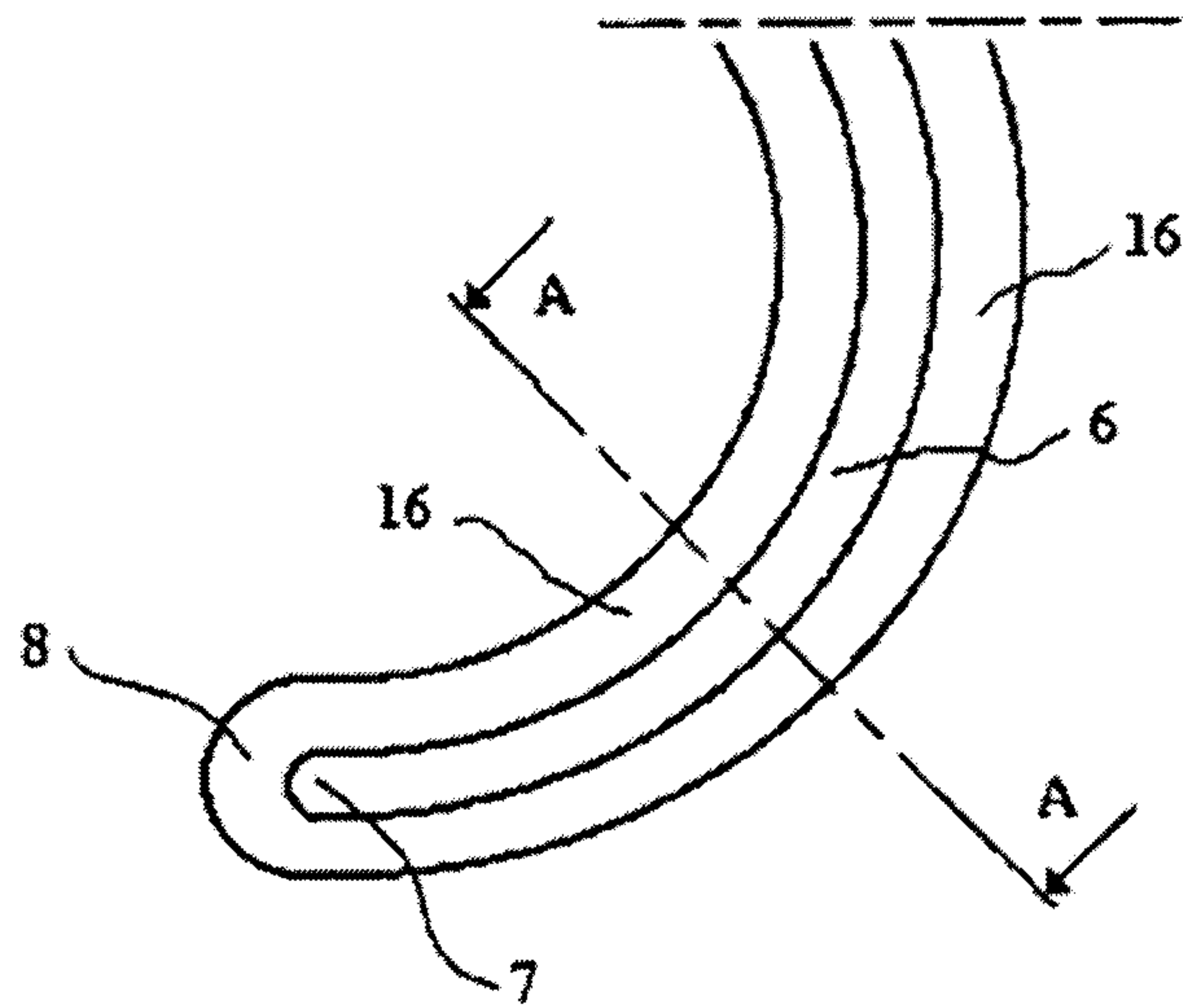


Fig. 1

Section A-A

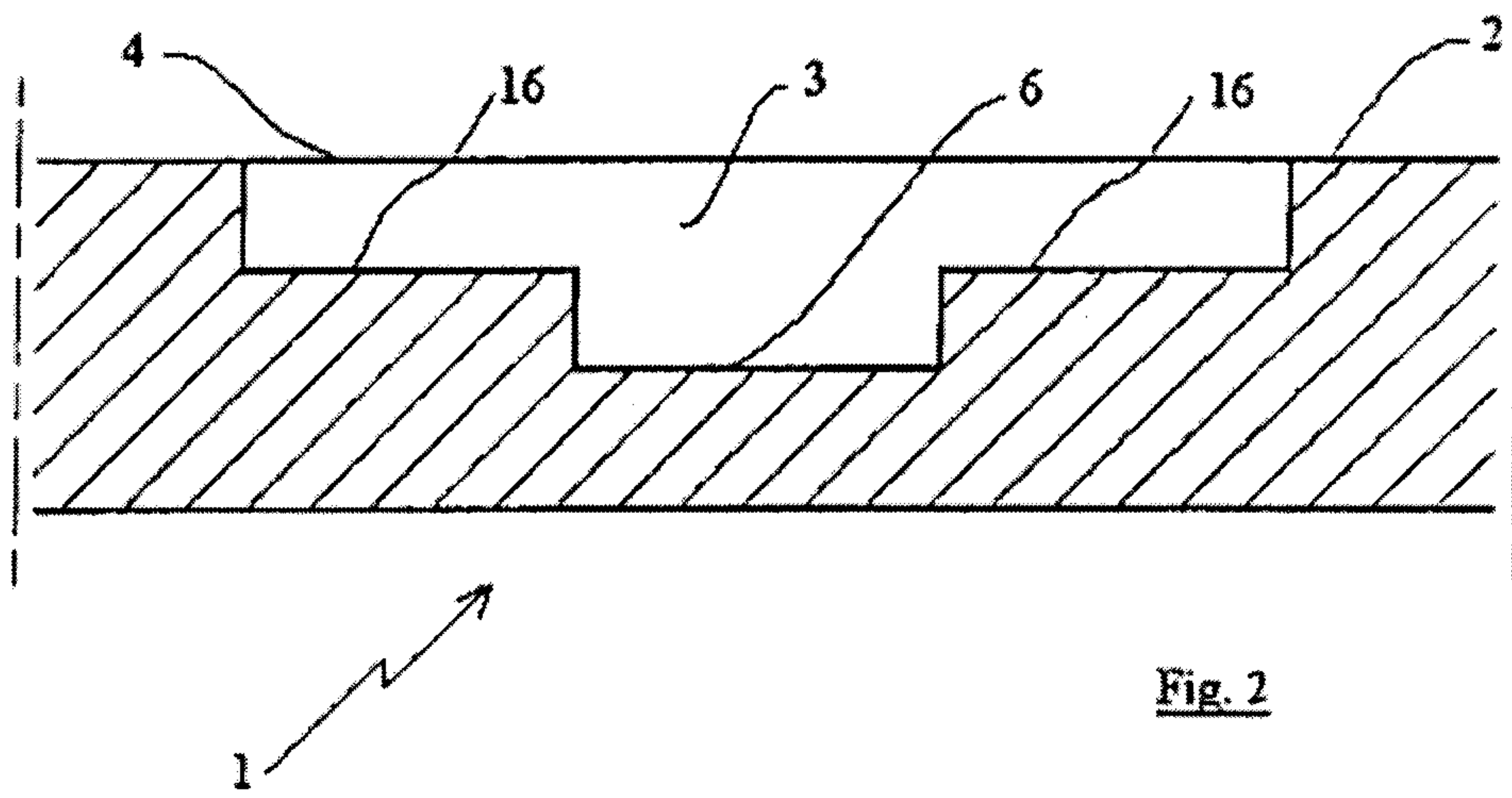
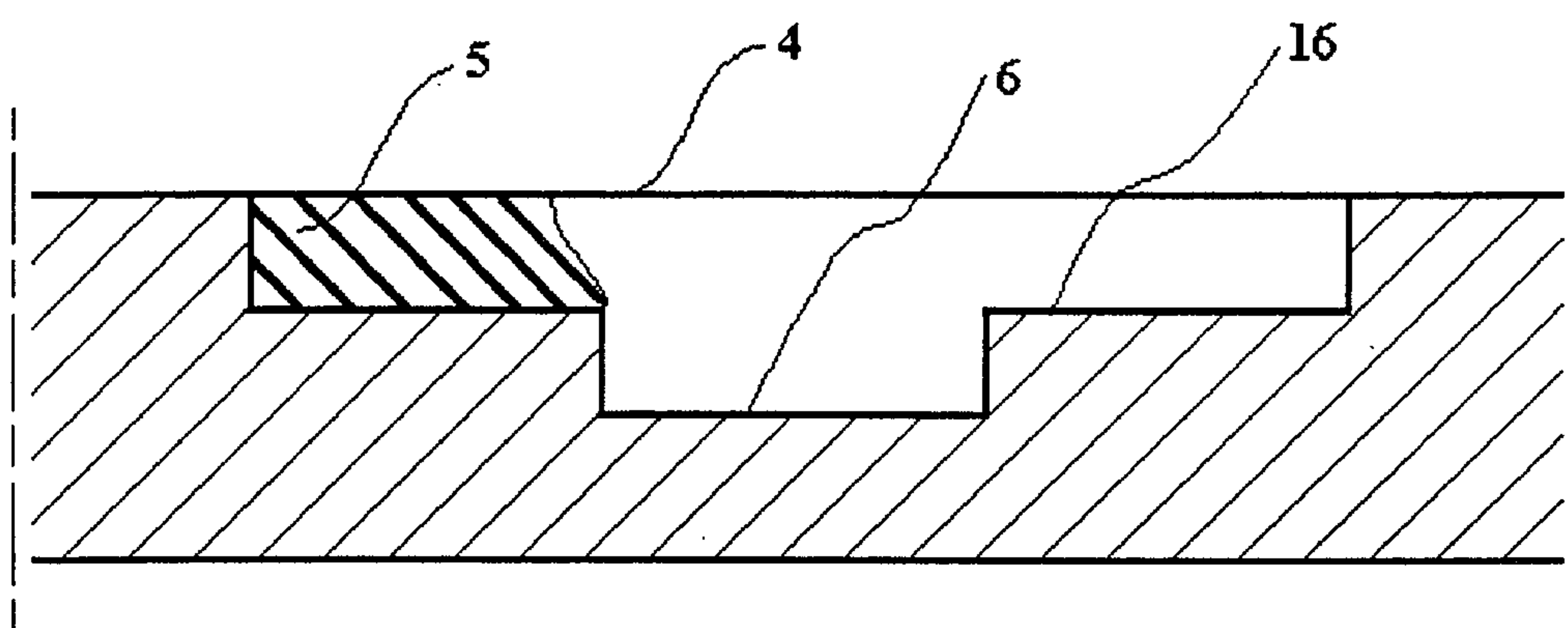
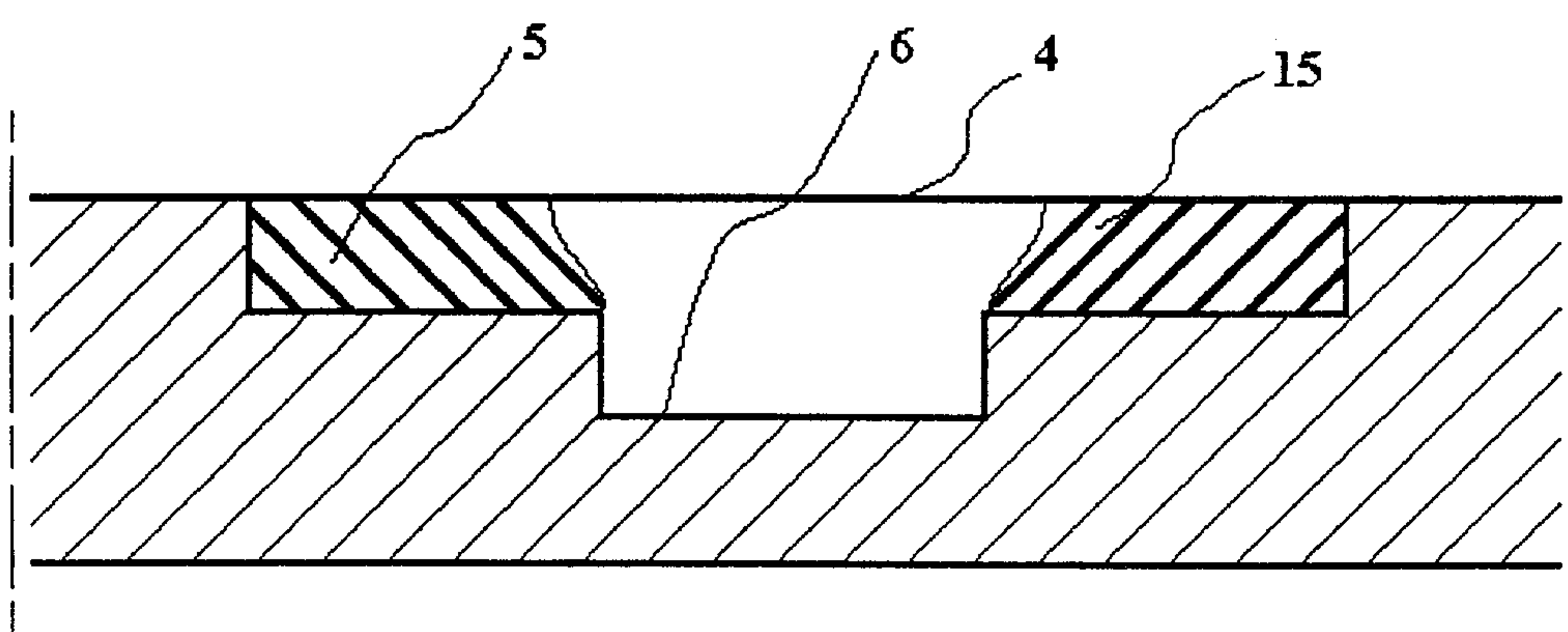
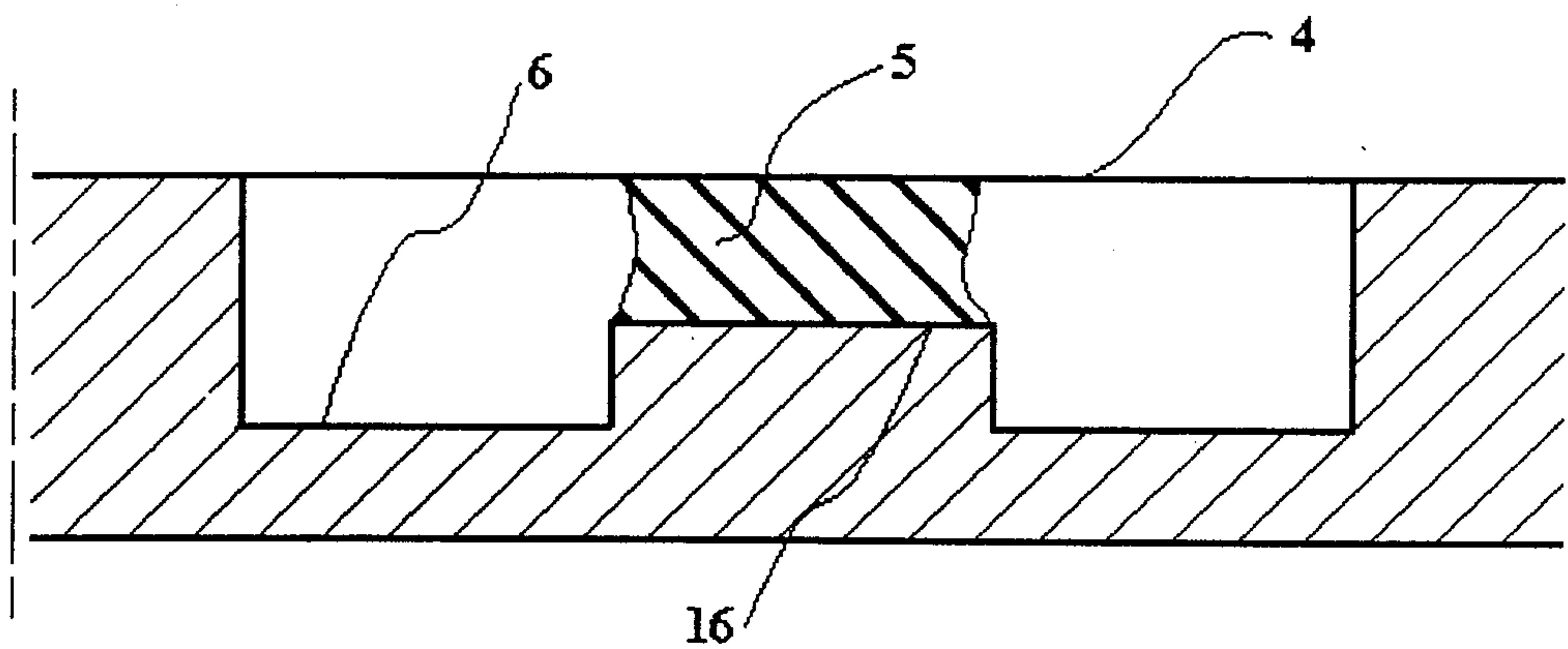
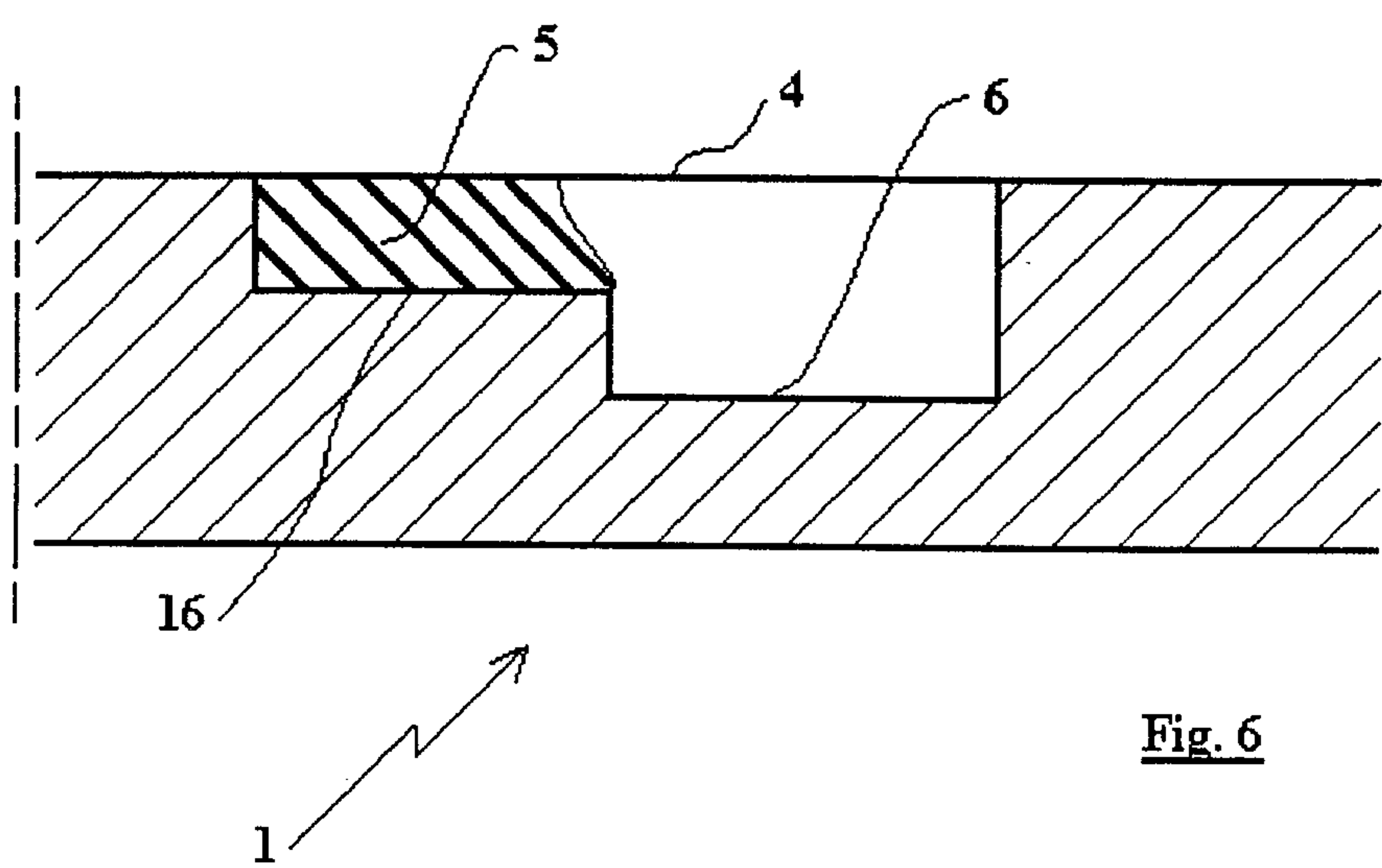


Fig. 2

2 / 3

Fig. 3Fig. 4

3 / 3

Fig. 5Fig. 6

