Title: INTRAOCULAR LENS MANUFACTURING PROCESS

Abstract: A method of cast molding an intraocular implant from two or more dissimilar materials using disposable molds. The subject cast molding method is particularly useful in the production of intraocular lenses manufactured from dissimilar materials. It is desirable to produce intraocular lenses from dissimilar materials in order to optimize the optical characteristics of the intraocular lens optic portion and to optimize stability and flexibility characteristics of the intraocular lens haptic elements.
INTRAOCULAR LENS MANUFACTURING PROCESS

Field of the Invention:

The present invention relates to a method of cast molding a surgical implant produced from two or more dissimilar materials, implants so produced and molds useful therefor. More particularly, the present invention relates to a method of cast molding intraocular lenses produced from two or more dissimilar materials using disposable plastic molds.

Background of the Invention:

The use of intraocular lenses (IOLs) to improve vision through the replacement of damaged or diseased natural lenses or to work in conjunction with a natural lens has obtained wide acceptance since the early 1980s. Accordingly, a wide variety of IOLs has been developed for surgical implantation into the posterior and/or anterior chamber of an eye. Commercially available IOLs generally comprise an optic portion and one or more haptic elements or plates to maintain proper positioning of the optic portion within the eye. The optic portions of such IOLs are commonly manufactured from relatively hard or rigid materials such as, for example, polymethylmethacrylate (PMMA), or from relatively soft, resilient polymeric materials such as, for example, hydrogels,
acrylics or silicones. More resilient polymeric materials are advantageous in the production of IOLs in that such materials are deformable and foldable to allow for implantation of the IOL through a smaller incision than that possible if implanting a more rigid IOL.

To manufacture a biocompatible IOL using known molding techniques, a polished stainless steel mold, having a mold cavity formed in a shape to achieve the desired refraction of light for the particular material utilized, is first selected. In the case of silicone for example, the uncured silicone polymer is introduced into the mold cavity and then cured. Several methods of molding IOLs are known such as injection molding, liquid injection molding, compression molding and transfer molding.

Several significant problems have been associated with known IOL molding techniques. The first problem is that current molding processes are labor intensive. Many elastomers used to mold IOLs, such as for example silicone elastomers, often times leave a residue in the stainless steel molds. Due to this residue, the molds must be cleaned between each molding cycle. In addition to being labor intensive, the cleaning requirements result in significant downtime for the equipment, which further increases production costs. A second problem associated with current known molding techniques is that of frequent tool damage and wear due to the repeated cleanings. Accordingly, molds must be replaced often resulting in increased production costs. A third problem associated with such molding techniques is one of quality control with respect to
the molded lenses. Intraocular implants such as IOLs must have smooth polished edges for implantation within an eye. Improperly finished edges on implants may result in damage to interior structures of the eye. In the case of improperly finished edges on IOLs, abrasions of the iris and tearing of the trabecular meshwork may result. Unfortunately, steel molds typically leave minute gaps between mold halves during the molding operation due to construction tolerances. Consequently, material flows out through the gaps during the molding of the IOL resulting in a phenomenon known as “flash”. Flash is unwanted material attached at the mold parting line on the molded implant. This flash material must be ground and/or polished off the implant, which is again labor intensive and increases production costs.

**Summary of the Invention:**

The present invention is a process for cast molding surgical implants, such as but not limited to corneal inlays, shunts and intraocular lenses (IOLs), but most preferably IOLs, wherein in the case of IOLs, the optic portion and haptic elements are produced using two or more dissimilar biocompatible materials. However, if desired, the subject molds and molding techniques are likewise useful in the manufacture of surgical implants such as in the case of IOLs having an optic portion and haptic elements produced using the same or similar biocompatible materials. The present cast molding process avoids the problems noted with regard to known molding techniques through the use of
disposable plastic molds, which are less expensive and less labor intensive to make and use.

The cast molding process of the present invention utilizes a multi-part, but preferably in the case of more customary IOLs a four-part, disposable plastic mold system. The first mold part of the subject mold system is a female base mold having a positioning wall formed along the periphery of an interior surface thereof and a molding surface on the interior surface. The molding surface is comprised of a center cavity used to form one surface of an IOL optic portion, one or more but preferably two junction cavities and two or more haptic element cavities. The center cavity is in fluid connection with each junction cavity. Also, each haptic element cavity is in fluid connection with at least one junction cavity.

The second mold part of the subject mold system is a center male mold having a molding surface on an interior surface comprised of an optic cavity used to form the second surface of the IOL optic portion. The center male mold is sized to be fully received within the positioning walls of the female base mold and may be shaped to ensure axial and rotational alignment.

The third and fourth mold parts of the subject mold system are haptic molds. Each haptic mold likewise has a molding surface on an interior surface comprised of one or more junction cavities and at least one haptic element cavity. When the haptic molds are placed in an interlocked relationship with center male mold, junction cavities and haptic element cavities are in fluid
connection with optic cavity. Haptic molds are also male molds sized to be fully received within the positioning walls of the female base mold and preferably shaped to ensure axial and rotational alignment. Each haptic mold is also formed to have material guides or ports extending from the haptic element cavity and/or junction cavity through to the exterior surface of the mold. Optionally, center male mold and haptic molds may be formed as a unitary mold.

The subject preferably four-part mold is used to cast mold a surgical implant, preferably an IOL, using two or more dissimilar biocompatible materials. An IOL having an optic portion of one preferably more resilient biocompatible material and haptic elements of a dissimilar preferably more rigid biocompatible material is produced by filling the base mold center cavity with the desired more resilient IOL optic material. The center male mold is then inserted into the female base mold allowing excess molding material to pass into one or more overflow reservoirs. During this process, some molding material flows into fluidly connected junction cavities so as to only partially fill the same. Shields are then positioned over the partially filled junction cavities and the molding material in the center cavity is polymerized using methods of polymerization known to those skilled in the art. Due to shielding, or any suitable method of protection, the molding material in the junction cavities is not polymerized. After removing the shields, the haptic molds are then inserted into the female base mold and a second dissimilar relatively rigid biocompatible molding material is provided through the material guides or ports to completely fill the junction cavities and the
haptic element cavities. The remaining unpolymerized first molding material and
the second molding material are then polymerized using methods of
polymerization known to those skilled in the art. Following polymerization, all
three male molds are removed from the female mold. The IOL is removed from
the female mold through the use of solvents or vibration.

Accordingly, it is an object of the present invention to provide a cast
molding system to produce IOLs having an optic portion and haptic elements
produced from dissimilar biocompatible materials.

Another object of the present invention is to provide a method for cast
molding IOLs having an optic portion and haptic elements produced from
dissimilar materials which is less labor intensive.

Another object of the present invention is to provide a method for cast
molding IOLs having an optic portion and haptic elements produced from
dissimilar materials with lower production costs.

Another object of the present invention is to provide molds for cast
molding IOLs having an optic portion and haptic elements produced from
dissimilar materials.

Still another object of the present invention is to provide a method for cast
molding IOLs having an optic portion and haptic elements produced from
dissimilar materials suitable for high volume production.

These and other objectives and advantages of the present invention,
some of which are specifically described and others that are not, will become
apparent from the detailed description, drawings and claims that follow wherein like features are designated by like numerals.

**Brief Description of the Drawings:**

Figure 1 is a plan view of an intraocular lens having an optic portion and two haptic elements;

Figure 2 is a plan view of an intraocular lens having an optic portion and three looped haptic elements;

Figure 3 is a plan view of an intraocular lens having an optic portion and two plate haptic elements;

Figure 4 is an exploded view in perspective of a disposable mold system for molding an IOL having an optic portion and haptic elements produced from dissimilar materials, constructed in accordance with the teachings of the present invention;

Figure 5 is a plan side view of the female base mold of the mold system of Figure 4;

Figure 6 is a cross-sectional side view of the female base mold of Figure 5 taken along lines 6—6;

Figure 7 is a plan side view of the center male mold of the mold system of Figure 4;

Figure 8 is a plan bottom view of the center male mold of Figure 7;
Figure 9 is a plan side view of the haptic mold of the mold system of
Figure 4;

Figure 10 is a cross-sectional side view of the haptic mold of Figure 9
taken along lines 10—10;

Figure 11 is an exploded view in perspective of a disposable mold system
for molding an IOL having an optic portion and haptic elements produced from
dissimilar materials, constructed in accordance with the teachings of the present
invention;

Figure 12 is a plan side view of the female base mold of the mold system
of Figure 11;

Figure 13 is a cross-sectional side view of the female base mold of Figure
12 taken along lines 13—13;

Figure 14 is a plan side view of the center male mold of the mold system
of Figure 11;

Figure 15 is a plan bottom view of the center male mold of Figure 14;

Figure 16 is a plan side view of the haptic mold of the mold system of
Figure 11;

Figure 17 is a cross-sectional side view of the haptic mold of Figure 16
taken along lines 17—17; and

Figure 18 is a plan side view of a shield useful with the mold system of
Figure 4 or Figure 11.
**Detailed Description of the Invention:**

The present invention is a method of cast molding surgical implants, such as most preferably intraocular lenses (IOLs), from dissimilar biocompatible materials, disposable molds useful for such cast molding method and surgical implants produced using such cast molding method. Figures 1 through 3 illustrate various IOLs that may be molded using the cast molding method of the present invention. Various IOLs are illustrated herein for purposes of example only and are not intended to be in any way limiting to the scope of the present invention. Figure 1 illustrates an IOL 10 having an optic portion 12 and two haptics 18. Figure 2 illustrates an IOL 10 having an optic portion 12 and three looped haptics 18. Figure 3 illustrates an IOL 10 having an optic portion 12 and two plate haptics 18.

The present cast molding method is useful in the manufacture of surgical implants, such as IOLs 10 as illustrated in Figures 1 through 3, wherein the optic portion 12 and haptic elements 18 are produced using two or more dissimilar biocompatible materials. It is desirable to produce IOLs 10 from dissimilar materials to optimize optical characteristics of the IOL 10 optic portion 12, optimize support and flexibility characteristics of the IOL 10 haptic elements 18, and, if desired, to also optimize support and flexibility characteristics of the IOL 10 haptic attachment area 13. However, if desired, the subject molds may be used in the manufacture of IOLs 10 having an optic portion 12 and haptic elements 18 produced using the same or similar biocompatible materials.
Suitable biocompatible optic molding materials include but are not limited to most preferably materials having a refractive index of 1.25 or higher and a level of resiliency that enables the material to return to its original shape after having been folded or compressed in small incision implantation. Examples of such materials include but are not limited to silicone polymers, hydrocarbon and fluorocarbon polymers, hydrogels, soft acrylic polymers, polyesters, polyamides, polyurethane, silicone polymers with hydrophilic monomer units, fluorine-containing polysiloxane elastomers and combinations thereof. The preferred material for the production of optic portion 12 due to desirable characteristics is a hydrogel made from 2-hydroxyethyl methacrylate (HEMA) and 6-hydroxyhexyl methacrylate (HOHEXMA), i.e., poly(HEMA-co-HOHEXMA). Suitable haptic molding materials include but are not limited to materials capable of providing support without excessive fragility. Examples of such materials include but are not limited to methacrylates, acrylates, more rigid hydrogels, silicone polymers and combinations thereof. The preferred material for the production of haptic elements 18 due to desirable characteristics is polymethyl methacrylate (PMMA).

Materials having characteristics suitable for the manufacture of other ophthalmic implants formed of dissimilar materials utilizing the method and molds of the present invention become obvious to those skilled in the art in light of the present disclosure.

The cast molding method or process of the present invention utilizes a multi-part, but preferably a four-part for manageability, disposable plastic mold
system 24 as best illustrated in Figure 4. The first part of mold system 24 is female base mold 26 having a positioning wall 52 along periphery 56 of interior surface 50 and a molding surface 54 comprised of a center cavity 28, used to form one anterior surface 14 or posterior surface 16 of optic portion 12, one or more but preferably two junction cavities 30 in fluid connection with center cavity 28 and at least one haptic element cavity 32 in fluid connection with junction cavities 30. Surrounding center cavity 28, junction cavities 30 and haptic element cavity 32 is extended edge 84 to eliminate material flow between molds to prevent flash. Base mold 26 is sized as needed according to the article to be manufactured using the same. For the manufacture of IOLs 10, base mold 26 is approximately 20 to 35 mm but preferably approximately 25 to 35 mm in length for ease in handling, approximately 10 to 20 mm but preferably approximately 15 to 20 mm in width for ease in handling and approximately 10 to 20 mm but preferably 15 to 20 mm in height for ease of handling. Each of the molding cavities of base mold 26 may be sized to be slightly larger in size than the final IOL product desired to accommodate for material shrinkage, often times as high as 15 percent, during polymerization thereof. Alternatively, overflow reservoirs 66 may be shielded from polymerization whereby nonpolymerized material therein may flow into adjacent cavities upon shrinkage of polymerized material in the adjacent cavities and be polymerized. Polymerization of material to form an
implant is preferably carried out under pressure within the range of approximately 50,000 pounds per square inch at extended edge 84 and edge recess 86 to eliminate cosmetic defects in the product formed.

The second part of mold system 24 is center male mold 34 having an interior surface 50 with a molding surface 58 comprised of an optic cavity 60 used to form the second surface, either anterior surface 14 or posterior surface 16 of optic portion 12, surrounded by edge recess 86. Edge recess 86 is sized to accept and work in conjunction with extended edge 84 of female base mold 26. Center male mold 34 is sized to be fully received snugly within positioning walls 52 of female base mold 26 and shaped to interlock for axial and rotational alignment. To ensure axial and rotational alignment, any of a variety of means may be used such as for example but not limited to providing base mold 26 and center male mold 34 with aligning pins 80 and pin recesses 82 on interior surface 50 thereof, providing shape specific forms to base mold 26 and center male mold 34 such as illustrated in Figure 4 and/or providing base mold 26 and center male mold 34 with tab 20 and groove 22 alignment means as illustrated in Figure 11. For the manufacture of IOLs 10, center male mold 34 is approximately 5 to 12 mm but preferably approximately 6 to 10 mm in length to allow optic cavity 60 to be slightly larger than the final desired optic diameter, approximately 8 to 18 mm but preferably approximately 13 to 18 mm in width for proper fit within positioning walls 52 and approximately 10 to 20 mm but preferably 15 to 20 mm in height for ease of handling. As noted above, optic
cavity 60 of center male mold 34 may be sized to be slightly larger in size than the final optic portion 12 desired due to material shrinkage during polymerization thereof.

The third and fourth mold parts of mold system 24 are haptic molds 62. Each haptic mold 62 likewise has an interior surface 50 with molding surface 64 comprised of one or more junction cavities 40 and at least one haptic element cavity 42 in fluid connection with optic cavity 60 surrounded by edge recess 86. Haptic molds 62 are male molds sized to be fully received within positioning walls 52 of the female base mold 26 and shaped to interlock for axial and rotational alignment as described above. Haptic molds 62 are also formed to have material guides or ports 46 extending from haptic element cavity 42 and/or junction cavity 40 through to the exterior surface 48 of the haptic mold 62. For the manufacture of IOLs 10, haptic mold 62 is approximately 14 to 22 mm but preferably approximately 18 to 24 mm in length to allow proper fit within positioning walls 52, approximately 8 to 18 mm but preferably approximately 13 to 18 mm in width to allow proper fit within positioning walls 52 and approximately 10 to 20 mm but preferably 15 to 20 mm in height for ease of handling. As noted above, each cavity of haptic mold 62 may be sized to be slightly larger in size than the final corresponding product part desired due to material shrinkage during polymerization thereof.
Suitable materials from which mold system 24 may be manufactured include for example but are not limited to polyurethanes, polypropylene, polyvinyl chloride or acrylates. In the case of ultraviolet light curing of the IOL 10 materials, a transparent or translucent material such as polypropylene is preferred.

The subject preferably four-part mold system 24 is useful to cast mold intraocular implants such as preferably an IOL 10 having an optic portion 12 and haptics 18 preferably manufactured from dissimilar biocompatible materials. An IOL 10 is cast molded in accordance with the present invention by providing a predetermined quantity of a suitable optic portion 12 molding material into the center cavity 28 of base mold 26. The center male mold 34 is then inserted within positioning walls 52 of female base mold 26 allowing excess molding material to pass into one or more overflow reservoirs 66. During this process, some molding material flows into junction cavities 30. Shields 68 are then placed within positioning walls 52 using handles 70 to shield interior surface 50 of base mold 26. Base 72 of shield 68 is dimensioned to be the same as that of interior surface 50 of haptic mold 62 to ensure proper fit within positioning walls 52. Shields 68 are preferably fabricated from the same material as mold system 24 with the addition of an ultraviolet light absorber such as for example but not limited to 2-hydroxy-5-acryloyloxyphenyl-2H-benzotriazoles or vinylsilylalkoxyarylbenzotriazoles. The molding material between center cavity 28 and optic cavity 60 is then polymerized using methods of polymerization.
known to those skilled in the art such as but not limited to ultraviolet light or heat curing. Due to shields 68, the molding material in junction cavities 30 is not polymerized. Shields 68 are then removed from base mold 26 and haptic molds 62 are then placed within positioning walls 52 of female base mold 26. Suitable haptic molding material is provided through material guides 46 of haptic molds 62 to fill junction cavities 30 and 40 and haptic element cavities 32 and 42 with any excess material flowing into reservoir 66. The haptic molding material is then polymerized using methods of polymerization known to those skilled in the art such as but not limited to ultraviolet light or heat curing. Following polymerization, all three male molds, i.e., the center male mold 34 and both haptic molds 62, are removed from female base mold 26. IOL 10 is removed from female base mold 26 through the use of solvents or vibration. IOL 10 is then optionally polished as needed, sterilized and packaged as customary in the art.

As described in detail above, the method of cast molding intraocular implants such as but not limited to IOLs, the molds suitable for such cast molding and the IOLs so produced in accordance with the present invention provides a relatively inexpensive method of manufacturing implants produced from two or more dissimilar materials. The present description is provided for purposes of illustration and explanation. It will be apparent to those skilled in the art that modifications and changes may be made to the preferred embodiment described herein without departing from its scope and spirit.
I claim:

1. A method for cast molding an ophthalmic implant from two or more dissimilar materials comprising:
   filling a center cavity and allowing partial fill of junction cavities of a mold with a first biocompatible material;
   polymerizing said first biocompatible material in said center cavity while shielding and not polymerizing said first biocompatible material in said junction cavities;
   filling partially filled junction cavities and remaining cavities of said mold with a second biocompatible material dissimilar to said first biocompatible material; and
   polymerizing said first and second biocompatible materials in said junction cavities and said remaining cavities.
2. A method for cast molding an intraocular lens from two or more dissimilar materials comprising:

filling a center cavity and allowing partial fill of junction cavities of a mold with a first biocompatible material;

polymerizing said first biocompatible material in said center cavity while shielding and not polymerizing said first biocompatible material in said junction cavities;

filling partially filled junction cavities and remaining cavities of said mold with a second biocompatible material dissimilar to said first biocompatible material; and

polymerizing said first and second biocompatible materials in said junction cavities and said remaining cavities.

3. The method of claim 1 or 2 wherein said first biocompatible material is selected from the group consisting of silicone polymers, hydrocarbon and fluorocarbon polymers, hydrogels, soft acrylic polymers, polyesters, polyamides, polyurethane, silicone polymers with hydrophilic monomer units, fluorine-containing polysiloxane elastomers and combinations thereof.
4. The method of claim 1 or 2 wherein said first biocompatible material is 2-hydroxyethyl methacrylate (HEMA) and 6-hydroxyhexyl methacrylate (HOHEXMA), i.e., poly(HEMA-co-HOHEXMA).

5. The method of claim 1 or 2 wherein said second biocompatible material is selected from the group consisting of methacrylates, acrylates, hydrogels, silicone polymers and combinations thereof.

6. The method of claim 1 or 2 wherein said second biocompatible material is polymethyl methacrylate.

7. The method of claim 1 or 2 wherein said first and second biocompatible materials are polymerized using ultraviolet light.

8. The method of claim 1 or 2 wherein said first and second biocompatible materials are polymerized using heat.

9. The method of claim 1 or 2 wherein said first and second biocompatible materials are both hydrogel materials possessing dissimilar characteristics.

10. The method of claim 1 or 2 wherein said method includes polishing said ophthalmic implant or intraocular lens following removal from said mold.
11. The method of claim 1 or 2 wherein said method includes sterilizing said ophthalmic implant or intraocular lens following removal from said mold.

12. An intraocular implant manufactured by the method of claim 1.

13. An intraocular lens manufactured by the method of claim 1 or 2.

14. An intraocular lens with two or more haptics manufactured by the method of claim 1 or 2.

15. An intraocular lens with two or more looped haptics manufactured by the method of claim 1 or 2.

16. An intraocular lens with two or more plate haptics manufactured by the method of claim 1 or 2.

17. Disposable molds for cast molding an intraocular implant from two or more dissimilar materials comprising a female base mold with a molding surface on an interior surface thereof, a center male mold with a molding surface on an interior surface thereof and one or more secondary male molds with a molding surface on an interior surface thereof.
18. The disposable molds of claim 8 wherein said molds are formed from the same or different materials selected from the group consisting of polyurethanes, polypropylene, polyvinyl chloride and acrylates.

19. The disposable molds of claim 8 wherein said molds are formed from polyurethane.

20. The disposable molds of claim 8 wherein said molding surfaces have mold cavities surrounded by an extended edge or a recessed edge to prevent flash.
### INTERNATIONAL SEARCH REPORT

**PCT/US 01/05223**

**A. CLASSIFICATION OF SUBJECT MATTER**

**IPC 7** B29D11/02

According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**

Minimum documentation searched: (classification system followed by classification symbols)

**IPC 7** B29D B29C

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

EPO-Internal, PAJ, WPI Data

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
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<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
<td>X</td>
<td>EP 0 328 246 A (SHEPHERD THOMAS H) 16 August 1989 (1989-08-16) figures IA-1F</td>
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<td>A</td>
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<td>X</td>
<td>EP 0 920 980 A (MENICON CO LTD) 9 June 1999 (1999-06-09) claim 1</td>
<td>12-14</td>
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<tr>
<td>X</td>
<td>EP 0 552 528 A (MENICON CO LTD) 28 July 1993 (1993-07-28) column 1, line 5 - line 10</td>
<td>12-14</td>
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<tr>
<td>A</td>
<td>US 5 762 836 A (BOS GILLES ET AL) 9 June 1998 (1998-06-09) abstract; figure 3</td>
<td>17-20</td>
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</table>

Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

- **X** Special categories of cited documents:
  - *A* document defining the general state of the art which is not considered to be of particular relevance
  - *E* earlier document but published on or after the international filing date
  - *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
  - *O* document referring to an oral disclosure, use, exhibition or other means
  - *P* document published prior to the international filing date but later than the priority date claimed

- **T** Later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principles or theory underlying the invention

- **X** Document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is taken alone

- **Y** Document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

- **Q** Document member of the same patent family

**Date of the actual completion of the international search**

26 July 2001

**Date of mailing of the international search report**

09.08.2001

**Name and mailing address of the ISA**

European Patent Office, P.B. 6816 Patentlaan 2 NL - 2280 HV Rijswijk
Tel: (+31-70) 340-2060, Tx. 31 651 eep nl, Fax: (+31-70) 240-3916

**Authorized officer**

Roberts, P

File PCT/ISA/210 (second draft) (July 1999)
<table>
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<th>Relevant to claim No.</th>
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| A        | EP 0 592 228 A (MENICON CO LTD)  
13 April 1994 (1994-04-13)  
figures 5,14,15 | 17-20                |
INTERNATIONAL SEARCH REPORT

Box I  Observations where certain claims were found unsearchable (Continuation of item 1 of first sheet)

This International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:

1. □ Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:

2. □ Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:

3. □ Claims Nos.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).

Box II  Observations where unity of invention is lacking (Continuation of item 2 of first sheet)

This International Searching Authority found multiple inventions in this international application, as follows:

see additional sheet

1. □ As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.

2. □ As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.

3. □ As only some of the required additional search fees were timely paid by the applicant, this International Search Report covers only those claims for which fees were paid, specifically claims Nos.:

4. □ No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:

Remark on Protest

□ The additional search fees were accompanied by the applicant's protest.

X No protest accompanied the payment of additional search fees.
This International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. Claims: 1-16
   Intraocular implant and method for its manufacture in two steps

2. Claims: 17-20
   Mold for casting intraocular implants comprising secondary male mold parts
<table>
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<th>Patent document cited in search report</th>
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