Title: SEALING VALVE ASSEMBLY FOR MEDICAL PRODUCTS

Abstract: A sealing valve assembly is provided for medical products. The valve assembly includes a valve member mountable within a passageway of a medical product body element, the valve member defining a peripheral portion spaced from a central axis and including two walls extending from the peripheral portion toward the central axis. The walls include ends that contact each other to preclude flow through the passageway. The valve member may be formed of a material having a durometer of less than about 20 Shore A. The valve member may also have a higher durometer material with other wall configurations, or may have a wall member having a varying durometer.
For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.
SEALING VALVE ASSEMBLY FOR MEDICAL PRODUCTS

CROSS-REFERENCE TO RELATED APPLICATIONS

The present application is a continuation-in-part (CIP) of U.S. Patent Application Serial No. 09/741,730, filed on December 19, 2000, entitled "Sealing Valve Assembly For Medical Products" whose inventors are Allan F. Willis and Alan Conlin. Application Serial No. 09/741,730 is incorporated by reference herein in its entirety for all purposes.

BACKGROUND OF THE INVENTION

Various types of valves incorporating sealing devices are known and widely used in the medical field. For example, mechanical sealing valves are required for various types of catheters, lavage devices, and endoscopy systems developed for a wide range of medical purposes. The valve sealing mechanisms typically preclude the flow of substances (gaseous or fluid) through the medical device in one direction, for example from the patient to the outside environment, while permitting the introduction or flow of desired substances (food, medication, etc.) in the other direction, for example into the patient through the medical device.

Enteral feeding devices utilizing a gastrostomy catheter ("feeding tube") are examples of conventional medical devices utilizing a sealing valve to prevent gastric fluids, gases, or particles from unintentionally exiting the patient's body via the catheter. U.S. Patent Nos. 5,997,503 and 5,997,546, both owned by Applicants' Assignee and incorporated by reference herein, disclose balloon catheters suitable for use as skin-level gastrostomy catheters for enteral feeding. Various other devices and catheters have also been proposed for gastrostomy feeding and are known to those skilled in the art.

Conventional sealing valves include duckbill valves, flap valves, slit seal valves, etc. Silicone rubber duckbill seals having durometer ratings of about 50 Shore and a wall thickness of about 0.040 inches have been employed in sealing gastrostomy catheters. Such sealing valves are typically kept closed via back pressure present on one side of the valve unless and until the seal is opened from the other side of the valve by insertion of a feeding tube or by a pressure differential across the seal.

Over time, conventional sealing valves may degrade or lose some of their flexibility (i.e., their ability to return to an initial sealing position after deflection) due
to repeated or extended periods of opening of the valve. Also, in certain bodily environments, such as the acidic environment of the gastro-intestinal tract, exposure to bodily fluids or stomach contents may have a corrosive or other negative effect on the sealing valve over time.

Thus, there is a need for an improved medical sealing valve for use in various medical devices, particularly gastrostomy catheter devices.

SUMMARY OF THE INVENTION

Objects and advantages of the invention will be set forth in part in the following description or may be apparent from the description or may be learned through practice of the invention.

It should be noted that any given range presented herein is intended to include any and all lesser included ranges. For example, a range of from 45 - 90 would also include 50-90, 45-80, 46-89, and the like.

According to the invention, an improved sealing valve assembly is provided for use in a wide array of medical products, including catheters, lavage devices, endoscopy systems, etc. The valve assembly according to the invention is not limited in its particular use and may be used in any medical product wherein it is necessary to permit fluid flow through the product in one direction while preventing backflow of fluids through the product in the opposite direction. For example, the valve assembly is particularly useful in a gastrostomy catheter. Accordingly, it should be appreciated that the present invention also includes medical products or devices incorporating the inventive valve assembly.

The valve assembly includes a valve housing defining a passageway therethrough. A valve member is seated in the valve housing within the passageway. The valve member includes a peripheral portion spaced from the central axis of the valve assembly and two walls extending from the peripheral portion towards the central axis. The walls include ends that contact and seal against each other to preclude flow through the passageway. The valve member may be formed of a material having a durometer of less than about 20 Shore.

In another embodiment, the valve member walls may define a cross-section that tapers in thickness from about 0.005 to about 0.010 inches at their respective sealing end to about 0.040 inches at a location spaced from the sealing end. The walls may taper continuously or discontinuously.
In accordance with another aspect of the invention, at least one of the valve member walls may define an overlapping portion extending at an angle with respect to a plane of the remaining portion of the wall. This overlapping portion overlaps and seals against a portion of the other wall. The overlapping portion may be parallel to the other wall.

In accordance with yet another aspect of the invention, the valve member walls may include a first portion including a material having a durometer of about 50 Shore A and a second portion having a durometer of less than about 20 Shore A. The second portion is disposed at the sealing end of the wall. The first portion is disposed spaced from the sealing end.

In accordance with another exemplary embodiment a medical product is provided that includes a valve housing that defines an opening with a central axis. The opening extends through the valve housing so that the opening forms part of a fluid path through the medical product. A valve member is disposed within the opening and has a single seal interface defined by at least two opposing flexible walls. The valve member includes a peripheral portion that has opposing flexible walls that extend from the peripheral portion towards the central axis. The opposing flexible walls include ends that contact each other along the single seal interface. A stiffening member is also included and is located on one of the opposing flexible walls. The stiffening member remains free from contact with the valve housing when the valve member is in the closed position and also remains free from contact with the valve housing when the valve member is in the open position.

The invention will be described in greater detail below with reference to the figures.

**BRIEF DESCRIPTION OF THE DRAWINGS**

Fig. 1 is a perspective view of a medical product, in particular a gastrostomy catheter, according to the present invention;

Fig. 2 is a cross-sectional view of the gastrostomy catheter of Fig. 1 taken along line A-A of Fig. 1 particularly illustrating a valve assembly disposed within a catheter body element;

Fig. 3 is an enlarged cross-sectional view of the valve assembly of Fig. 2 taken along line A-A of Fig. 1;
Fig. 4 is a side view of a first embodiment of the valve member of the valve assembly from the perspective taken in Fig. 3;

Fig. 5 is a side view of the valve member of Fig. 4 rotated 90 degrees;

Fig. 6 is a top view of the valve member of Fig. 4;

Fig. 7 is a cross-sectional view of the assembly of Fig. 3 with the end of a feeding tube inserted therein;

Fig. 8 is a cross-sectional view of an alternate embodiment of a valve member;

Fig. 9 is a cross-sectional view of a second alternate embodiment of a valve member;

Fig. 10 is a cross-sectional view of a third alternate embodiment of a valve member;

Fig. 11 is a cross-sectional view of an exemplary embodiment of a valve member that includes a pair of stiffening members;

Fig. 12 is a perspective view of the valve member of Fig. 11;

Fig. 13 is a cross-sectional view of a valve housing assembly configured with a valve member in the closed position that includes a stiffening member in accordance with one exemplary embodiment; and

Fig. 14 is a cross-sectional view of the valve housing assembly and valve member of Fig. 13 in which the valve member is in the open position.

**DETAILED DESCRIPTION**

Embodyments of the invention will now be described in detail with reference to the examples shown in the figures. Each example is provided by way of explaining the invention, and not as a limitation of the invention. Various modifications and variations can be made in the invention without departing from the scope and spirit of the invention. For example, features illustrated or described with respect to one embodiment may be used in another embodiment to yield still a further embodiment.

In the following description, the improved sealing device is described with reference to a gastrostomy catheter for purposes of explaining and illustrating the sealing valve assembly. This is not meant as a limitation of the invention. It should be appreciated that the improved sealing device according to the present invention has wide application in the medical field and can be used in any medical device
that incorporates a seal to prevent the flow of substances in one direction while permitting the flow of substances in the opposite direction.

A gastrostomy catheter 10 is shown in Figs. 1 and 2 as an example of a medical product incorporating the improved valve sealing assembly according to the present invention. The use and operation of such gastrostomy catheters is known to those skilled in the art and need not be described in detail herein. Reference is made to U.S. Patent Nos. 5,997,546 and 5,997,503 incorporated herein by reference for a detailed explanation of gastrostomy catheters.

As used herein, the term "distal" refers to the direction of the patient and the term "proximal" refers to the direction of the clinician.

Referring briefly to Figs. 1 and 2, the gastrostomy device 10 includes a catheter body element 12 having a head 20, an elastomeric sleeve forming a balloon 16, and a catheter segment 22. The catheter segment 22 includes a shaft 23 having a dispensing tip 38 attached to its distal end. A first central opening or port 27 in the head 20 incorporates a valve housing assembly 14 that enables the injection of nutrients, enteral feeding solution, medications, etc., into the patient through the head 20 and into a first lumen 34 of the catheter segment 22. The valve housing assembly 14 includes the improved sealing valve according to the invention, as described in greater detail below. A plug 32 may be provided at the end of a lanyard 30 attached to the catheter head 20. The plug 32 can be inserted into the first port 27 when the port is not being used to administer fluids into the patient.

A second port 29 in the head 20 serves as an opening through which fluid such as air or saline solution may be injected into or removed from the balloon 16 through a second lumen 35 having a distal opening 37. An inflation valve assembly 18 is disposed in the second port 29 for permitting the clinician to control inflation and deflation of the balloon 16, as is understood by those skilled in the art.

The valve housing assembly 14 is shown in greater detail in Fig. 3. The valve housing assembly 14 includes a valve housing 40 and the improved sealing valve member 42. The valve housing 40 has an annular member 44 and a relatively flat cap member 46 connected to the annular member 44, for example by a press-fit or an adhesive. The cap member 46 may have a ridge 48 that fits within a groove 50 in the annular member 44 for securing the two parts together. The
valve housing 40 includes a flange 54 that extends into a corresponding groove in the catheter body element 12 to seat and secure the valve housing therein. The flange 54 extends radially from a lower tubular portion 58 of the valve housing 40. The tubular portion 58 may have inner walls that angle towards a centerline axis 69 of the valve housing assembly 14. The walls of the tubular portion 58 parallel to the plane of Fig. 3 may be parallel to the axis 69.

A passage 52 is defined through both portions of the valve housing assembly 14. The passage 52 is in fluid communication with the feeding lumen 34 through the valve member 42 for introducing nutrients, feeding solutions, medication, etc., into the patient's stomach, intestine, or other body cavity. The valve member 42 is disposed within the passage 52 for selectively blocking the fluid path through feeding lumen 34.

The valve member 42 incorporates novel features of the present invention and various embodiments of the valve member 42 are partially shown in Figs. 3-10. Referring to Fig. 3, the valve member 42 includes a proximal flange portion 68 disposed between facing surfaces 64, 66 of the annular member 44 and the cap member 46. Small circular ridges 60, 62 on the facing surfaces 64, 66 engage the flange portion 68 and help to seat and hold the valve member 42 in place. The valve member 42 includes two walls 70, 72 on opposite sides of the central axis 69 that extend from the peripheral portion 68 and are angled towards the central axis 69. The walls 70, 72 meet at a single seal interface 74 defined by ends 76, 78 of the walls. Parts 68, 70, and 72 form what is known in the art as a duckbill valve. As explained in greater detail below, to achieve satisfactory sealing, the thickness and profile of the walls 70, 72 may be designed in various ways according to the present invention depending on the durometer of the material used for the valve member 42.

The seal interface 74 is biased to a closed position and maintained in the closed position by body cavity pressure unless forced open by an object inserted into the valve member 42 through the cap member 46 or by a significant pressure differential between the patient cavity and the external environment. An object useful to open the valve member 42 could be a feeding tube, a feeding tube adapter, or a pressure relief tube. As an example, Fig. 7 shows the valve housing assembly 14 of Fig. 3 with the end of a feeding tube 108 inserted therein. Note
that the tube end 108 pushes open the walls 70,72 of valve member to allow
communication with the patient's body cavity through the valve housing 14 and
feeding lumen 34.

The various components of medical device 10 are preferably formed from
bio-compatible materials such as medical grade silicone. More particularly, the
valve member 42 may be made of a liquid injection molding silicone elastomer
such as MED-4810 (10 Shore Durometer) or MED-4820 (20 Shore Durometer),
available from NuSil Technology of Carpintera, California. The valve member is
molded, and the sealing interface 74 is created by slitting or cutting apart the
respective walls after molding.

As mentioned, the thickness and profile of the walls 70, 72 will depend on
the durometer of the material used to form the valve member 42. For example, for
a wall thickness of about 0.040 inches, the durometer of the valve member may be
about 20 Shore A or less, and may be in the range of from about 8 Shore A to
about 15 Shore A, and more particularly about 10 Shore A. Applicants have
discovered that use of a lower durometer material than had been used previously
in the art for the valve member unexpectedly provides an improved seal and
greater seal flexibility without sacrificing the structural integrity of the valve
member.

However, if the durometer of the valve member 42 is higher, such as up to
about 50 Shore A as has been used previously, the wall cross sections of the valve
member can be altered to provide improved seal flexibility. For example, alternate
embodiments of the valve member 42 are shown in Figs. 8-10.

The valve members in Figs. 8 and 9 have walls with varying cross-section
dimensions. The valve member 142 in Fig. 8 includes two walls 170,172 that are
continuously tapered, narrowing toward the sealing interface 174. The walls may
taper up to about 15% of their upper wall thickness. For example, the walls 170,
172 may taper from about 0.040 inches to about 0.005 to about 0.010 inches at the
seal interface. The valve member 242 in Fig. 9 includes two walls 270, 272 having
a constant cross-sectional thickness along their upper portions 274, 276 of about
0.040 inches and reduced thickness edge portions 278, 280 having a thickness of
about 0.005 to about 0.010 inches. The valve member 342 in Fig. 10 has walls
370, 372 which may have a thickness throughout of about 0.040 inches. The wall
372 has an edge portion angled with respect to the remaining portion of the wall 372. The angled portion 380 is parallel to and overlaps a portion 378 of the wall 370. The overlapping edge portions 378, 380 may have a thickness and length of about 0.040 inches.

Each of the valve members of Figs. 8-10 may be molded of relatively high durometer materials, with the modified wall structure providing improved sealing and wall flexibility.

As another option, a valve member 42 having a shape as shown in Figs. 3-7 could be molded having a varying durometer that decreases in the direction of the seal interface. In such a valve member, the peripheral portion 68 and the upper portion of a wall or the walls 70,72 may be made from a material of relatively higher durometer (for example in the range of about 50 Shore A), and the remaining portion may be made of a lower or a gradually decreasing durometer material (for example, decreasing to the range of about 20 Shore A or less, more particularly about 8 Shore A to about 15 Shore A, and even more particularly to about 10 Shore A). The softer tip of the walls 70,72 would thus provide improved sealing while the harder upper portion would maintain the shape of the valve member.

Figures 11 and 12 show another exemplary embodiment of the valve member 42. Here, the valve member 42 is provided with a stiffening member 400 on the wall 70. Another stiffening member 402 is located on wall 72. Although shown as incorporating a pair of stiffening members 400 and 402, it is to be understood that any number of stiffening members may be employed in accordance with other exemplary embodiments. For instance, only a single stiffening member 400 may be employed in certain exemplary embodiments. Alternatively, the valve member 42 may be provided with three stiffening members on wall 70 and two stiffening members on wall two in accordance with other exemplary embodiments. The stiffening members 400 and 402 may be integrally formed with the valve member 42 or may be separate components that are attached thereon in accordance with various exemplary embodiments.

The stiffening members 400 and 402 act to strengthen the walls 70 and 72. Additional strength may be desired, for instance, in preventing unwanted opening of the valve member 42 due to an increased pressure inside of the body cavity into
which the gastrostomy device 10 is employed. Additionally or alternatively, added strength to the walls 70 and 72 of the valve member 42 may be desired in order to ensure that the valve member 42 opens upon application of a desired force when inserting a feeding tube, feeding tube adapter, pressure relief tube, or the like. By acting as a strengthening component, the stiffening members 400 and 402 may act to urge the walls 70 and 72 towards one another so as to create a tighter seal interface 74. Alternatively, the stiffening members 400 and 402 may be configured so that their presence does not act to urge the legs 70 and 72 towards one another so as to increase the strength of the seal at the seal interface 74.

The stiffening members 400 and 402 may also be employed in order to help the walls 70 and 72 retain their original shape when the feeding tube, feeding tube adapter, pressure relief tube, or the like is removed. It is sometimes the case that valve members will become deformed if they are placed into a certain position for an extended amount of time. For example, if a feeding tube adapter were inserted into the valve member 42 so as to keep the valve member 42 in an open position for an extended period of time, the walls 70 and 72 may become plastically deformed such that removal of the feeding tube adapter may cause the walls 70 and 72 to only partially retake their closed position shape. In this instance, the walls 70 and 72 may not completely return to their original shape so that seal interface 74 is not created and hence the valve member 42 will not be fully closed. The stiffening members 400 and 402 may therefore act as “memory” components such that they urge the walls 70 and 72 into their original, closed shape after the valve member 42 is opened.

Figure 13 shows the valve housing 14 configured with the valve member 42 along with a feeding tube 108 that is positioned therein but does not contact the valve member 42 so as to open the valve member 42. As such, the valve member 42 is in the closed position so that the seal interface 74 is formed. The valve member 42 includes a pair of stiffening members 400 and 402 on the walls 70 and 72. The stiffening members 400 and 402 do not contact the valve housing 40.

Figure 14 shows the valve housing assembly 14 and valve member 42 of Fig. 13 with the feeding tube 108 inserted therein so as to open the valve member 42. In this instance, the feeding tube 108 pushes the walls 70 and 72 towards the valve housing assembly 14. The stiffening members 400 and 402 remain free from
contact with the valve housing assembly 14 in the open position. As can be seen, a space remains between the stiffening members 400 and 402 and the inner walls of the tubular portion 58. As such, the stiffening members 400 and 402 remain free from contact with the valve housing assembly 14 during times in which the valve member 42 is both in the open and closed positions.

The stiffening members 400 and 402 may be of any shape or any thickness in accordance with various exemplary embodiments. For example, in accordance with one exemplary embodiment the stiffening members 400 and 402 have a thickness of about 0.05 inches. The stiffening member 400 may have an arcuate shaped outer surface 404. The arcuate shaped outer surface 404 may extend along the entire angled length of leg 70. The other stiffening member 402 may also be provided with an arcuate shaped outer surface 406 that likewise extends along the entire outer length of the angled portion of leg 72. It is to be understood, however, that in accordance with other exemplary embodiments one or more of the surfaces of the stiffening members 400 and 402 need not be completely or partially arcuate shaped.

In accordance with one exemplary embodiment the valve member 42 may be formed at least in part by a material that has a durometer of less than 10 Shore A. Additionally or alternatively, at least one of the walls 70 or 72 may define a first portion that includes a material with a durometer of about 50 Shore A and a second portion with a durometer of less than about 20 Shore A. Here, the second portion is disposed at the respective end of at least one of the walls 70 or 72, and the first portion is disposed spaced from the respective end. It is to be understood that various exemplary embodiments exist in which one or more of the stiffening members 400 and 402 may be provided on any of the valve members as previously discussed in accordance with prior exemplary embodiments. The stiffening members 400 and 402 so provided may be configured so that they remain free from contact with the valve housing assembly 14 at times in which the valve member 42 is in both the open and closed positions.

Thus, applicants have disclosed above numerous designs and/or material selections that provide for improved sealing across the seal interface of a valve member as well as improved flexibility of the valve member. Such improved sealing flexibility may provide a longer lifespan to medical devices such as
gastrotomy catheters and to the valve members and related valve assemblies used therein. Such improved sealing may also provide a more reliable product in terms of reducing or eliminating undesired leakage.

It should be appreciated by those skilled in the art that various modifications and variations can be made in the present invention without departing from the scope and spirit of the invention. It is intended that the invention include such modifications and variations as come within the full scope of the appended claims and any equivalents thereof.
WHAT IS CLAIMED IS:

1. A medical product defining a fluid path, the medical product comprising a valve assembly disposed in the path, the valve assembly comprising:

   a valve housing defining an opening having a central axis, the opening extending through the valve housing such that the opening forms part of the fluid path through the medical product;

   a valve member disposed within the opening, the valve member having a single seal interface defined by at least two opposing flexible walls, the valve member having a peripheral portion with the opposing flexible walls extending from the peripheral portion toward the central axis, the opposing flexible walls including ends that contact each other along the single seal interface; and

   a stiffening member located on one of the opposing flexible walls, wherein the stiffening member remains free from contact with the valve housing when the valve member is in the closed position, and wherein the stiffening member remains free from contact with the valve housing when the valve member is in the open position.

2. The medical product as in claim 1, wherein a pair of stiffening members are present on the valve member such that one stiffening member is located on one of the opposing flexible walls and the other stiffening member is located on the other one of the opposing flexible walls, wherein the stiffening members remain free from contact with the valve housing when the valve member is in the closed position, and wherein the stiffening members remain free from contact with the valve housing when the valve member is in the open position.
3. The medical product as in claim 1 or 2, wherein the valve member and the stiffening member are integrally formed with one another.

4. The medical product as in any one of claims 1 to 3, wherein the stiffening member has an arcuate surface.

5. The medical product as in any one of claims 1 to 4, wherein the valve member is at least partially made from a material that has a durometer value of less than 10 Shore A.

6. The medical product as in claim 5, wherein the valve member is at least partially made from a material that has a durometer value between about 8 to less than 10 Shore A.

7. The medical product as in any one of the preceding claims, wherein the medical product is a gastrostomy catheter, the fluid path defined between a patient’s body cavity and an exterior of the body.
### INTERNATIONAL SEARCH REPORT

**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**

INV. A61M39/24 A61J15/00

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)

A61M A61J A61F A61B F16K

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

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>X</td>
<td>US 6 702 255 B2 (DEHNASHTIANN MARK) 9 March 2004 (2004-03-09) column 4, line 62 - column 6, line 34 column 7, line 1 - line 8 figures 1-4,9</td>
<td>1-4</td>
</tr>
<tr>
<td>Y</td>
<td></td>
<td>7</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Further documents are listed in the continuation of Box C.

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

21 April 2006

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

28/04/2006

**Name and mailing address of the ISA/EP**

Schönleben, J

European Patent Office, P.B. 5816 Patentlaan 2 NL – 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Fax. (+31-70) 340-3016

**Authorized officer**

Schönleben, J

Form PCT/ISA/210 (second half) (April 2006)
<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
</table>
| X         | US 3 941 149 A (MITTLEMAN ET AL) 
2 March 1976 (1976-03-02) 
column 2, line 54 - column 4, line 6; 
figures 1-9 | 1,3 |
| X         | US 5 261 459 A (ATKINSON ET AL) 
16 November 1993 (1993-11-16) 
column 3, line 64 - column 4, line 17 
column 5, line 35 - column 8, line 10 
column 10, line 25 - line 42 
figures 1-7 | 1-3 |
| A         | US 4 222 126 A (BORETOS ET AL) 
16 September 1980 (1980-09-16) 
figures 1-6 | 1-4 |
<table>
<thead>
<tr>
<th>Patent document cited in search report</th>
<th>Publication date</th>
<th>Patent family member(s)</th>
<th>Publication date</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>US 2005010176 A1</td>
<td>13-01-2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2003085373 A1</td>
<td>08-05-2003</td>
</tr>
<tr>
<td>US 2002077603 A1</td>
<td>20-06-2002</td>
<td>BR 0115781 A</td>
<td>07-12-2004</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2429367 A1</td>
<td>12-09-2002</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 1343555 A2</td>
<td>17-09-2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2005503838 T</td>
<td>10-02-2005</td>
</tr>
<tr>
<td></td>
<td></td>
<td>MX PA030055264 A</td>
<td>25-09-2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NO 20032757 A</td>
<td>17-06-2003</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 02070060 A2</td>
<td>12-09-2002</td>
</tr>
<tr>
<td>US 3941149 A</td>
<td>02-03-1976</td>
<td>AU 8570975 A</td>
<td>21-04-1977</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 1021664 A1</td>
<td>29-11-1977</td>
</tr>
<tr>
<td></td>
<td></td>
<td>FR 2290618 A1</td>
<td>04-06-1976</td>
</tr>
<tr>
<td></td>
<td></td>
<td>GB 1514237 A</td>
<td>14-06-1978</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 51070530 A</td>
<td>18-06-1976</td>
</tr>
<tr>
<td></td>
<td></td>
<td>ZA 7506362 A</td>
<td>29-09-1976</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 5669294 A</td>
<td>04-07-1994</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2151135 A1</td>
<td>23-06-1994</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 0725907 A1</td>
<td>14-08-1994</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 8506881 T</td>
<td>23-07-1996</td>
</tr>
<tr>
<td>US 4222126 A</td>
<td>16-09-1980</td>
<td>NONE</td>
<td></td>
</tr>
</tbody>
</table>