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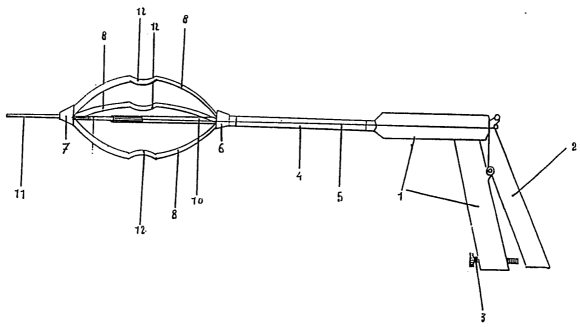
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(54) Title: PERCUTANEOUS MECHANICAL DILATING CATHETER FOR CARDIAC VALVES AND BLOOD VESSELS



(57) Abstract

This invention is a mechanical dilating catheter that serves to dilate stenotic cardiac valves and blood vessels. It is composed of four parts: 1) the handle (1); 2) the sheath (4); 3) the wire (5); 4) the dilating body. This has a concave area (12) in the central section of the expansion blades (8) which firmly anchors the body to the edges of stenotic cardiac valves and to stenotic segments of blood vessels.

^{*} See back of page

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Percutaneous mechanical dilating catheter for cardiac valves and blood vessels.

Great progress has been made in interventional cardiology over the last decade. The therapeutic use of balloon catheters has been extended to stenotic cardiac valves with the aim of increasing the flow of blood by dilating the valve. However, balloon catheters have certain limitations. This is due partly to their structure and conformation and partly to the fact that they can have a dilating effect on the valve only at the moment of maximum inflation of the balloon. Below, we outline some of the most common drawbacks associated with this type of catheter.

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The first of these is serious hypotension with cerebral ischemia which occurs when the balloon is fully inflated and temporarily obstructs the flow of blood inside the valve undergoing dilation.

The second is the onset of cardiac valve insufficiency, which aggravates the criginal illness and necessitates valve transplantation within a fairly short timescale. This is due to the fact that the balloon must be inflated to its maximum capacity relatively quickly, in addition to the fact that it can only assume one shape regardless of the type of valve involved. A third drawback is the damage that can be caused by a balloon larger in diameter that the valve ring or the stenotic blood vessel being treated. Yet another drawback is the instability of the balloon within stenotic valves. It can slip on the surface of such structures, producing little or no dilation.

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The aim of the percutaneous mechanical dilating catheter is to eliminate the serious drawbacks mentioned above. This new catheter solves the problem of how to dilate stenotic cardiac valves and blood vessels.

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This catheter is undoubtedly an improvement on the existing balloon catheter for the following reasons;

- a) unlike the balloon catheter, it allows an improved flow of blood during the process of dilation itself, thus avoiding the risk of serious systemic hypotension and ischemia.
- b) the new catheter allows the dilating body to be positioned in the most anatomically favourable way inside the stenotic valve or vessel.
- c) the cardiologist can regulate the dimensions of the body as required during the procedure.
- d) the dilation precedure can be carried cut in a much less traumatic manner without limitations of time and with maximum accuracy of pressure on the stenotic valve cusps and blood vessels.
- e) the body of the new catheter can be firmly anchored to the edges of stenotic valves and to the stenotic segments of blood vessels.

From observation obtained during experiments on dead subjects, the inventors deduced that only the concavity on the central section of the expansion blades allows to firmly anchor the dilating body to the edges of stenotic valves and to the tracts of blood vessels. Used in vivo without a firm anchorage, the dilating body would frequently lose contact with the stenotic structure, possibly providing little or no dilating effect.

DESCRIPTION

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Fig. 1 shows the percutaneous mechanical dilating catheter. It comprises four parts; the handle, the sheath, the wire. and the dilating body. The diagram shows the catheter in the act of dilation. The handle 1 is pistol-shaped and is furnished with a lever 2. This transmits commands via a wire 5 to open and close the dilating body. At the bottom (lower extremity) of the handle there is a screw to regulate expansion 3. The flexible sheath 4 has a lenght and diameter which can be adjusted to suit the age and the surface area of the patient. The wire runs through the sheath and is attached at one end to the handle and at the other to a head 7 of the body. One end of the sheath is connected to the handle and the other to a head 6 of the body. The body, which is shown in the act of dilation, is made up as follows The two heads 6, 7 are located at each extremity of the body. Each head has three sites for the expansion blades 8. The proximal head 6 has a hole for the wire 5; the distal head 7 has an anchor point for the wire 5. Each of the heads is fitted with a tube 9, 10 which fit into each other and serve to regulate the movement of the heads. The wire runs through these tubes. At the end of the body there is a flexible guide 11 attached to the head 7. There is a concave area 12 located in the central section of the expansion blades 8.

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MODE OF EMPLOYMENT

Pressure applied on the lever 2 modifies the position of the wire. This brings the two heads closer together, thus opening (flexing) the three blades. During this expansion, the blades come into contact with stenotic valves or blood vessels, which they then dilate as required. After dilation, the pressure on the heads is released allowing the body to return to its initial closed position.

CLAIMS

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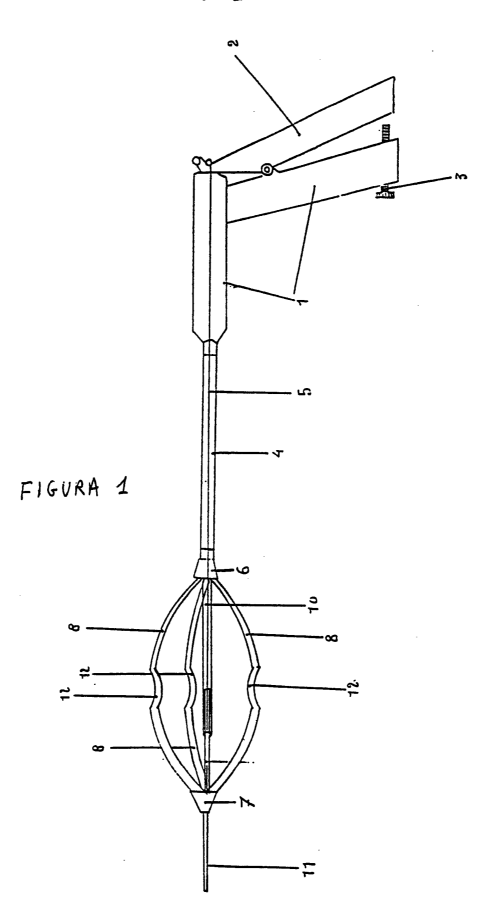
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- 1. the "Percutaneous mechanical dilating catheter for cardiac valves and blood vessels" characterized by being composed of four parts; a handle, a sheath, a wire, a dilating body.
- 5 2. the "Percutaneous mechanical dilating catheter for cardiac valves and blood vessels", as preceding, having a dilating body.
 - 3. the "Percutaneous mechanical dilating catheter for cardiac valves and blood vessels", as preceding, having a dilating body preferably made of stainless steel, but not excluding any other suitable material.
 - 4. the "Percutaneous mechanical dilating catheter for cardiac valves and blood vessels", as preceding, having a dilating body comprising three flexible, elastic blades 8 preferably made of stainless steel, but not excluding any other suitable material.
 - 5. the "Percutaneous mechanical dilating catheter for cardiac valves and blood vessels", as preceding, of which the blades 8 have a convave area 12.
- 6. the "Percutaneous mechanical dilating catheter for cardiac valves and blocd vessels", as preceding, of which the concave area 12 is located in the central section of the blades 8.

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INTERNATIONAL SEARCH REPORT

International Application No PCT/IT 90/00072

I. CLASS	IFICATION OF SUBJECT MATTER (if several classific	ation symbols apply, indicate all) ⁶	
	to International Patent Classification (IPC) or to both Nation	nal Classification and IPC	
IPC ⁵ :	A 61 M 29/00		
II. FIELDS	SEARCHED		
	Minimum Documenta		
Classification	on System CI	lassification Symbols	
IPC ⁵	A 61 M, A 61 B		
	Documentation Searched other that to the Extent that such Documents a	an Minimum Documentation are Included in the Fields Searched *	
			Relevant to Claim No. 13 1-3 1-4 1-4 1-4 1-4 Inter the international filing date conflict with the application but inciple or theory underlying the levance; the claimed invention el or cannot be considered to levance; the claimed invention who are inventional to the considered to levance; the claimed invention who are inventional to the claim obvious to a person skilled tame patent family
III. DOCL	MENTS CONSIDERED TO BE RELEVANT		
Category •	Citation of Document, 11 with Indication, where appro	opriate, of the relevant passages 12	Relevant to Claim No. 13
X	DE, A, 1963316 (INSTITUT SOSUDISTOI) 24 June 1971 see claims 1-3; figu		1-3
X	US, A, 3517128 (HINES) 23 June 1970 see claim 1; figures	1,4	1-4
X	US, A, 4648402 (SANTOS) 10 March 1987 see claim 1; figures	1-6	1-4
X	US, A, 1677671 (COUNCILL 17 July 1928 see claims 1-6; figu		1-4
"A" do co "E" ea fili "L" do wh cit "O" do do "P" do lat	al categories of cited documents: 10 cument defining the general state of the art which is not natidered to be of particular relevance flier document but published on or after the international ing date cument which may throw doubts on priority claim(s) or nich is cited to establish the publication date of another ation or other special reason (as specified) incument referring to an oral disclosure, use, exhibition or her means incument published prior to the international filling date but are than the priority date claimed TIFICATION THE Actual Completion of the International Search 22nd November 1990	cited to understand the print invention "X" document of particular relevant to considered novel involve an inventive step "Y" document of particular relevant to the considered to involve an inventive step	nflict with the application but siple or theory underlying the siple or theory underlying the sance; the claimed invention or cannot be considered to rance; the claimed invention we an inventive step when the one or more other such docume obvious to a person skilled me patent family
Internation	enal Searching Authority EUROPEAN PATENT OFFICE	Signature of Authorites Since	MISS T. TA 7FL ASSE

ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

IT 9000072 SA 38820

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 04/12/90
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Patent document cited in search report	Publication date	Patent family member(s)	Publication date
DE-A- 1963316	24-06-71	None	
US-A- 3517128	23-06-70	None	
US-A- 4648402	10-03-87	None	
US-A- 1677671		None	