**Abstract:** The present invention relates to a medical implant which is vacuum-packed within an air tight cover material. Further, the present invention relates to a medical device for making a medical implant covered by an air tight cover material, wherein the medical device comprises at least an underpressure generating device and a sealing device, wherein the sealing device is arranged for air tight sealing of the cover material while the medical implant is located within the cover material, wherein the underpressure generating device is coupled or can be coupled with a space which is surrounded by the cover material, and wherein the underpressure generating device and the sealing device are controllable such that after generating a certain underpressure within the space surrounded by the cover material through the underpressure generating device the sealing device is activatable to seal the cover material in an air tight manner to form an air tight protection cover around the medical implant located therein. Further, the present invention relates to a method for making a medical implant covered by an air tight cover material, comprising the steps a) providing the medical implant, b) providing sterile cover material in the form of at least one or at least two pieces, c) placing the medical implant within a space surrounded by the at least one or at least two pieces of the cover material, d) creating an underpressure within the space surrounded by the at least one or at least two pieces of the cover material by evacuating air from this space, e) air tight sealing of the cover material.

![Diagram](Image)
Published:

— with international search report (Art. 21(3))
Medical implant, medical device and method for making a medical implant

Technical field of the invention

The present invention relates to medical implants of any kind for humans or animals. Such an implant can be e.g. a cardiac pacemaker, as an example for an implant which is widely used in surgery. Another example for such an implant is an endoprosthesis. The invention also relates to a device and a method for making such a medical implant.

Background of the invention

When implanting a medical implant into a living body, there is always a potential risk of infection. In particular relatively complex implants are difficult to disinfect, e.g. due to a complex shape or, in case of implants with an electrical function, due to cables, cords and so on.

It is therefore an object of the invention to reduce the likelihood of any infection caused by the implantation of a medical implant.

It is another object of the invention to provide for a medical implant, a medical device and a method for making a medical implant which is easy to use and does not require specific skills.

Prior art

Summary of the invention

The present invention provides for an improvement over the prior art proposals by means of the medical implant of claim 1, the medical device of claim 4 and the method for making a medical implant of claim 14. The dependent claims are directed to further embodiments of the invention.

The invention has the advantage that the likelihood of infections due to implantation of a medical implant can be significantly reduced. As a consequence, the need for corrective follow-up surgeries is reduced. By means of vacuum-packing of the medical implant the infection risk can be reduced both by using an appropriate cover material, like an aseptic or sterile material which can in addition be functionalized, e.g. by applying biocide substances to the cover material, e.g. anti-microbial and/or anti-inflammatory substances.

The cover material can also be functionalized by creating a certain surface structure, like a certain roughness or a certain relief. The infection risk is further reduced by removing air from the medical implant itself and from the space within the cover material where the medical implant is located in. By the removal of air any negative effects which such air could cause within a living body can be avoided.

As a consequence, the healing process of the patient and the acceptance of the implant within the patient are improved.
Further, the application of the invention is very easy for the user. For example, the packaging of the medical implant within the air tight cover material can be done directly in the operating room, e.g. while the surgery is performed. The medical device for making a medical implant, as defined in claim 4 and the embodiments of the dependent claims, supports the user by making the steps of vacuum-packing of the medical implant very simple.

The underpressure or vacuum to be generated by the underpressure generating device within the space which is surrounded by the cover material can be a vacuum or underpressure of any kind, which means of any pressure level below atmospheric pressure. Therefore, the terms underpressure and vacuum are used as synonyms. Generally it is advantageous to evacuate as much of the air as possible from the space surrounded by the cover material, in order to ensure that any germs and microbes are removed. In practice it is normally sufficient to create an underpressure which is e.g. 20 % less than the atmospheric pressure. Generally speaking, any vacuum whether it is low vacuum, medium vacuum, high vacuum, ultra-high vacuum or extremely high-vacuum, is usable for the present invention.

The cover material can be any air tight material which can be closed air tight. It is advantageous to use a cover material which has a sufficient elasticity to be sucked by the underpressure closely to the medical implant, in order to adapt the outer shape of the medical implant. The cover material can be e.g. in the form of a foil made of plastic, rubber, latex or any other biologically compatible material. The cover material can be a bio-resistant material which permanently resists within the patient after implantation. It is also possible to use a bio-degradable cover material. However, in such case it is advantageous to use a bio-degradable cover material which has a long degradation term, e.g. 2 months or more.

According to an advantageous embodiment to the invention, the air tight cover
material is a flexible material which rests again the outer surface of the medical implant due to the vacuum within the cover material. By means of the vacuum the cover material can be sucked onto the outer surface of the medical implant and adapts to the outer shape of the medical implant. Compared with prior art proposal, the invention saves space which is required for the medical implant in the body of the patient.

The sealing of the cover can be done one or more of welding, gluing, vulcanizing or any other bonding technology. The sealing can also be done by any kind of form fitting connection technologies, e.g. by providing means for a form fit connection between parts of the cover material, for example through producing a connection like a zip lock, a tongue and groove, a key and slot or a dovetail connection. In any case, the sealing has to be air tight.

The sealing device may comprise at least one welding unit or is a welding unit. Then a welding process is used for sealing the cover material. The sealing is normally done in the form of a weld seam. The weld seam can be produced e.g. by thermal-welding and/or chemical welding.

According to an advantageous embodiment of the invention, the medical device comprises at least a control device which is arranged for automatic control at least of the sealing device in dependence of the underpressure generated by the underpressure generating device. This has the advantage that the user is unburdened from controlling and/or activating the sealing device. For example, the sealing device can be automatically activated in case a certain value of the underpressure is sensed by the control device.

According to an advantageous embodiment of the invention, the medical device comprises exchangeable adapter inserts, wherein the medical implant located within the cover material can be put between the adapter inserts for further modification with the medical device. For example, the medical implant can be
located between an upper and a lower adapter insert. This has the advantage that the adapter inserts can be disinfected before use, e.g. by performing a disinfection step or by using prefabricated sterile adapter inserts. As a result, it is not necessary to disinfect the whole medical device. This saves time and effort and makes it easier to use the medical device e.g. during a surgery.

According to an advantageous embodiment of the invention, the adapter inserts are shaped parts which are adapted to the outer shape of the medical implant located within the cover material. This has the advantage that the shaping of the vacuum-packed medical device can be performed and supported by the adapter inserts.

According to an advantageous embodiment of the invention the sealing device comprises at least one welding unit or is a welding unit, wherein the welding unit comprises at least a weld seam producing device for producing a weld seam at the cover material with a defined outline.

According to an advantageous embodiment of the invention the welding unit is arranged for selective creation of heat required for thermal welding of the cover material only in one or more edge areas of the cover material to be welded and is arranged for substantially avoiding heating of the cover material in other areas. This has the advantage that other areas of the cover material as well as the medical implant located therein are not unnecessarily subjected to heat.

According to an advantageous embodiment of the invention the sealing device is arranged for creation of a circumferential sealing seam only on the outer edges of the cover material, so that the cover material is to be provided as a single-layer, non-bag form material, e.g. in the form of at least two pieces, which are connectable to each other by the sealing seam to form a bag.

According to a further embodiment of the invention the sealing device
comprises at least a first and a second sealing seam creation device which are separately controllable, whereby different edge areas of the sealing material to be sealed are sealable separately from each other by the first and second sealing seam creation device, in particular at different points in time. The sealing seam produced by the first and/or second sealing seam creation device can be a weld seam.

According to a further embodiment of the invention the medical device comprises at least one control device which is arranged for automatic control at least of sealing device, wherein the control device is arranged for creating a first part of the sealing seam by activating the first sealing seam creation device at a first point in time and for creating a second part of the sealing seam by activating the second sealing seam creation device at a second point in time which is different from the first point in time. This has the advantage that vacuum-packed medical implant can be created in two steps using raw cover material which is provided in non-bag form. In a first sealing step the cover material can be modified into a bag form. Then the vacuum can be applied. The vacuum can be applied permanently until the cover material is completely sealed in an air tight manner in a second sealing step. This avoids that any air could enter again the space which is surrounded by the cover material.

According to a further embodiment of the invention a functionalized cover material is provided as the cover material, in particular in the form of a foil made of plastic, rubber, latex or any other biologically compatible material. The functionalized cover can provide for one or more of the following functions and advantages: supporting the healing process of the patient, supporting the acceptance of the implant within the patient, providing anti-microbial and/or anti-inflammatory support.

According to a further embodiment of the invention a connection stud area for connecting the underpressure creating device is provided on the cover
material, and the connection stud area is sealed in an air tight manner after creation of a certain underpressure within the space surrounded by the cover material while the evacuation suction of the underpressure creating device is maintained.

According to a further embodiment of the invention the cover material is provided as at least two separate pieces, the at least two separate pieces are coupled to each other in a first sealing step, which is executed before an underpressure is created within the space surrounded by the cover material, then the underpressure is created and then the cover material is sealed in an air tight manner in a second sealing step. This has the advantage that the raw cover material can be supplied in any suitable form or packaging, like in the form of an endless material on a reel. The cover material can be cut into the required shape by the end user, e.g. by using scissors.

A further advantage is that storage of the raw materials for the cover material is simplified. By using certain standardized pieces of the cover material which are designed for certain kinds of implants and/or diseases, the invention provides for a high flexibility at low efforts (cost and space) for storage of the raw materials. In addition, the flexibility for the end user is maximized. As another advantage, application of the invention does not require complicate or costly devices. The medical device for making the medical implant can be produced at low to medium costs.

The invention can be used for any kind of medical implant. In particular, it is possible to use the invention for medical implants which have one or more cords, like electrical lines or hoses. In such case, according to a further embodiment of the invention one or more cords of the medical implant extend through a sealing seam area of the cover material, wherein a sealing seam is created on the cover material at least in the sealing seam area through which the one or more cords extend.
An advantageous method for making a medical implant covered by an air tight cover material comprises the steps:

a) providing the medical implant,
b) providing sterile cover material in the form of at least one or at least two pieces,
c) placing the medical implant within a space surrounded by the at least one or at least two pieces of the cover material,
d) creating an underpressure within the space surrounded by the at least one or at least two pieces of the cover material by evacuating air from this space,

e) air tight sealing of the cover material.

According to a further embodiment of the invention the method is executed less than 30 minutes before implantation of the medical implant. This means that it is possible to execute the method during a surgery.

According to a further embodiment of the invention the cover material is provided in the form of one or more tailored pieces according to the physical dimensions and/or outer shape of the medical implant at least in one viewing direction on the medical implant.

According to a further embodiment of the invention the medical implant located within the space surrounded by the cover material is placed for further modification between at least two exchangeable adapter inserts of the medical device.

According to a further embodiment of the invention the method is executed using the medical device of claim 4.

Brief description of the drawings
Various examples of embodiments of the invention will be described in detail with reference to the following figures, wherein:

figure 1 is an elevational view of a medical implant and its cover material,

figure 2 shows the medical implant within the cover material in an elevational view and a sectional side view,

figure 3 shows a vacuumizing step applied to the medical implant located within the cover material in similar views as figure 2,

figure 4 shows a finalizing step in making a vacuum-packed medical implant in an elevational view,

figure 5 shows the final vacuum-packed medical implant in an elevational view,

figure 6 shows a medical device for making a medical implant covered by an air tight cover material.

It should be understood that the drawings are not necessarily to scale. In certain instances, details which are not necessary to the understanding of the invention or render other details difficult to perceive may have been omitted. It should be understood, of course, that the invention is not necessarily limited to the particular embodiments illustrated herein. Same reference numerals are used throughout the drawings.

Detailed description of the illustrated embodiments

Figure 1 shows a medical device 1, e.g. a cardiac pacemaker, having two cords 2 which are for example electrical lines to be connected to a patient's
heart. Figure 1 further shows two pieces 3, 4 of an air tight functionalized cover material. The pieces 3, 4 were cut from a storage reel of cover material from which the pieces 3, 4 were tailored into the shape shown in figure 1.

In the figures 2 and 3, in the upper part an elevational view on the devices is shown and in the lower part a sectional view (section A-A) is shown.

In a next step shown in figure 2, the medical implant 1 is placed between the two pieces 3, 4 of the cover material. This creates a space 8 surrounded by the cover material where the medical implant 1 is located in. Now, using a medical device for making a medical implant, a sealing seam 6 is created by a first sealing seam creation device 9, e.g. a thermal welding unit. As can be seen, the sealing seam 6 circumferentially surrounds the medical implant 1 on the outer edges of the pieces 3, 4, thereby creating a stud 7 which is still open.

The cords 2 extend through the sealed area between the pieces 3, 4 which is closed by the sealing seam 6.

In a next step shown in figure 3, an underpressure generating device 10, e.g. in the form of a pump or any other form of suction device, is coupled e.g. via a hose 11 to the stud 7. Through the underpressure generating device 10 air is evacuated from the space 8, which results in the flexible cover material resting on the outer surface of the medical implant 1.

In a next step shown in figure 4 the stud 7 is closed by creating another sealing seam 13 by means of a second sealing seam creation device 12, e.g. again by thermal welding. During the step, the underpressure created by the underpressure generating device 10 is maintained. Once the sealing seam 13 is created, the medical implant 1 is completely enclosed in an air tight manner and therefore vacuum-packed within the cover material 5.

The result is shown in figure 5. The vacuum-packed medical implant of figure 5
can now be implanted into a patient.

Figure 6 shows in a sectional side view of a medical device for making a medical implant covered by an air tight cover material. The medical device comprises an upper frame 15 and a lower frame 17. In the upper frame 15 an exchangeable upper insert 14 is located. In the lower frame 17 an exchangeable lower insert 16 is located. The two inserts 14, 16 comprise inner openings which are shaped according to the outer shape of the medical implant and/or its surrounding cover material. For performing the steps explained before, the upper frame 15 is moved to the lower frame 17, in order to enclose the medical implant and its surrounding cover material between the inserts 14, 16. Then the first part of the sealing seam can be produced by the first sealing seam creation device 9. Then air is evacuated through the underpressure generating device 10. Then the second sealing seam 13 is produced by the second sealing seam creation device 12. Then the underpressure generating device 10 and its hose 11 can be removed. Then the vacuum-packed medical implant can be taken from the medical device after opening the frames 15, 17. In such a way, the medical device for making a medical implant covered by an air tight cover material can be designed similarly to a waffle iron.

Those reviewing this disclosure will appreciate that various exemplary embodiments have been shown and described, and that according to various exemplary embodiments, features associated with one exemplary embodiment may be used with features included in other exemplary embodiments.

As utilized herein, the terms "approximately," "about," "substantially," and similar terms are intended to have a broad meaning in harmony with the common and accepted usage by those of ordinary skill in the art to which the subject matter of this disclosure pertains. It should be understood by those of skill in the art who review this disclosure that these terms are intended to allow a description of certain features described and claimed without restricting the
scope of these features to the precise numerical ranges provided. Accordingly, these terms should be interpreted as indicating that insubstantial or inconsequential modifications or alterations of the subject matter described and claimed are considered to be within the scope of the invention as recited in the appended claims.

It should be noted that the term "exemplary" as used herein to describe various embodiments is intended to indicate that such embodiments are possible examples, representations, and/or illustrations of possible embodiments (and such term is not intended to connote that such embodiments are necessarily extraordinary or superlative examples).

The terms "coupled," "connected," and the like as used herein mean the joining of two members directly or indirectly to one another. Such joining may be stationary (e.g., permanent) or moveable (e.g., removable or releasable). Such joining may be achieved with the two members or the two members and any additional intermediate members being integrally formed as a single unitary body with one another or with the two members or the two members and any additional intermediate members being attached to one another.

References herein to the positions of elements (e.g., "top," "bottom," "above," "below," etc.) are merely used to describe the orientation of various elements in the FIGURES. It should be noted that the orientation of various elements may differ according to other exemplary embodiments, and that such variations are intended to be encompassed by the present disclosure.

It is important to note that the construction and arrangement of the battery module having electrochemical cells with integrally formed terminals as shown in the various exemplary embodiments is illustrative only. Although only a few embodiments of the present inventions have been described in detail in this disclosure, those skilled in the art who review this disclosure will readily
appreciate that many modifications are possible (e.g., variations in sizes, dimensions, structures, shapes and proportions of the various elements, values of parameters, mounting arrangements, use of materials, colors, orientations, etc.) without materially departing from the novel teachings and advantages of the subject matter recited in the claims. For example, the battery may be non-cylindrical (e.g., oval, rectangular, etc.), the position of elements may be reversed or otherwise varied (e.g., orientation of terminals), and the battery could be a number of different types (e.g., nickel metal hydride, lithium ion, lithium polymer, etc.). Accordingly, all such modifications are intended to be included within the scope of the present inventions. The order or sequence of any process or method steps may be varied or re-sequenced according to exemplary embodiments. Other substitutions, modifications, changes and omissions may be made in the design, operating conditions and arrangement of the various exemplary embodiments without departing from the scope of the present invention.
Claims

1. Medical implant which is vacuum-packed within an air tight cover material.

2. The medical implant of claim 1, wherein the air tight cover material is a flexible material which rests against the outer surface of the medical implant due to the vacuum within the cover material.

3. The medical implant of claim 1, wherein the cover material is a functionalized cover material, in particular in the form of a foil made of plastic, rubber, latex or any other biologically compatible material.

4. Medical device for making a medical implant covered by an air tight cover material, wherein the medical device comprises at least an underpressure generating device and a sealing device, wherein the sealing device is arranged for air tight sealing of the cover material while the medical implant is located within the cover material, wherein the underpressure generating device is coupled or can be coupled with a space which is surrounded by the cover material, and wherein the underpressure generating device and the sealing device are controllable such that after generating a certain underpressure within the space surrounded by the cover material through the underpressure generating device the sealing device is activatable to seal the cover material in an air tight manner to form an air tight protection cover around the medical implant located therein.
5. The medical device of claim 4, wherein the cover material is a functionalized cover material, in particular in the form of a foil made of plastic, rubber, latex or any other biologically compatible material.

5 6. The medical device of claim 4, wherein the medical device comprises at least a control device which is arranged for automatic control at least of the sealing device in dependence of the underpressure generated by the underpressure generating device.

10 7. The medical device of claim 4, wherein the medical device comprises exchangeable adapter inserts, wherein the medical implant located within the cover material can be put between the adapter inserts for further modification within the medical device.

15 8. The medical device of claim 7, wherein the adapter inserts are shaped parts which are adapted to the outer shape of the medical implant located within the cover material.

9. The medical device of claim 4, wherein the sealing device comprises at least one welding unit or is a welding unit, wherein the welding unit comprises at least a weld seam producing device for producing a weld seam at the cover material with a defined outline.

10. The medical device of claim 9, wherein the welding unit is arranged for selective creation of heat required for thermal welding of the cover material only in one or more edge areas of the cover material to be welded and is arranged for substantially avoiding heating of the cover material in other areas.

30 11. The medical device of claim 4, wherein the sealing device is arranged for creation of a circumferential sealing seam only on the outer edges of the
cover material, so that the cover material is to be provided as a single-layer, non-bag form material, e.g. in the form of at least two pieces, which are connectable to each other by the sealing seam to a bag.

5  12. The medical device of claim 4, wherein the sealing device comprises at least a first and a second sealing seam creation device which are separately controllable, whereby different edge areas of the sealing material to be sealed are sealable separately from each other by the first and second sealing seam creation device, in particular at different points in time.

13. The medical device of claim 12, wherein the medical device comprises at least one control device which is arranged for automatic control at least of the sealing device, wherein the control device is arranged for creating a first part of the sealing seam by activating the first sealing seam creation device at a first point in time and for creating a second part of the sealing seam by activating the second sealing seam creation device at a second point in time which is different from the first point in time.

14. Method for making a medical implant covered by an air tight cover material, comprising the steps
   a) providing the medical implant,
   b) providing sterile cover material in the form of at least one or at least two pieces,
   c) placing the medical implant within a space surrounded by the at least one or at least two pieces of the cover material,
   d) creating an underpressure within the space surrounded by the at least one or at least two pieces of the cover material by evacuating air from this space,
   e) air tight sealing of the cover material.
15. The method of claim 14, wherein a functionalized cover material is provided as the cover material, in particular in the form of a foil made of plastic, rubber, latex or any other biologically compatible material.

16. The method of claim 14, wherein a connection stud area for connecting the underpressure creating device is provided on the cover material, and the connection stud area is sealed in an air tight manner after creation of a certain underpressure within the space surrounded by the cover material while the evacuation suction of the underpressure creating device is maintained.

17. The method of claim 14, wherein the cover material is provided as at least two separate pieces, the at least two separate pieces are coupled to each other in a first sealing step, which is executed before an underpressure is created within the space surrounded by the cover material, then the underpressure is created and then the cover material is sealed in an air tight manner in a second sealing step.

18. The method of claim 14, wherein one or more cords of the medical implant extend through a sealing seam area of the cover material, wherein a sealing seam is created on the cover material at least in the sealing seam area through which the one or more cords extend.

19. The method of claim 14, wherein the method is executed less than 30 minutes before implantation of the medical implant.

20. The method of claim 14, wherein the cover material is provided in the form of one or more tailored pieces according to the physical dimensions and/or outer shape of the medical implant at least in one viewing direction on the medical implant.
21. The method of claim 14, wherein the medical implant located within the space surrounded by the cover material is placed for further modification between at least two exchangeable adapter inserts of the medical device.

22. The method of claim 14, wherein the method is executed using the medical device of claim 4.
INTERNATIONAL SEARCH REPORT

Box No. II  Observations where certain claims were found unsearchable (Continuation of item 2 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 No. III  Observations where unity of invention is lacking (Continuation of item 3 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 fees, this Authority did not invite payment of additional fees.

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 and, where applicable, the payment of a protest fee.

☐ The additional search fees were accompanied by the applicant’s protest but the applicable protest fee was not paid within the time limit specified in the invitation.

☐ No protest accompanied the payment of additional search fees.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER

INV. A61N1/375 A61F2/00 B65B31/02 B65B31/04 B65D81/20
B65B11/52 B65D75/30
ADD. A61L31/04 A61L31/16 A61L31/14 A61B19/02

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

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)
A61N A61F B65B B65D A61L A61B

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 practicable, search terms used)
EPO-Internal, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>US 2012/158136 AI (VOISARD CYRI L [CH] ET AL) 21 June 2012 (2012-06-21) paragraphs [0022], [0023], [0038] - [0042], [0056] - [0031]; claim 1; figures 1, 2, 11, 12</td>
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Further documents are listed in the continuation of Box C. See patent family annex.

* 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 application or patent 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 principle or theory underlying the invention
"X" document of particular relevance; the claimed invention cannot be considered novel or 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
"A" document member of the same patent family

Date of the actual completion of the international search
8 May 2015

Date of mailing of the international search report
19/05/2015

Name and mailing address of the ISA/
European Patent Office, P.B. 5818 Patentlaan 2
NL-2280 HV Rijswijk
Tel. (+31-70) 340-2040, Fax: (+31-70) 340-3016

Authorized officer
Fischer, Olivier

Form PCT/ISA/210 (second sheet) (April 2005)
### DOCUMENTS CONSIDERED TO BE RELEVANT

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<td>wo 2011/151641 A2 (LANCER UK LTD [GB]; BAKER SIMON [GB]) 8 December 2011 (2011-12-08) page 4, line 7 - page 7, line 21; figure 1A page 10, line 32 - page 12, line 22; figures 4A, 4B, 6</td>
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International Searching Authority found multiple (groups of) inventions in this international application, as follows:

1. claims: 1-22

Medical implant, medical device for making a medical implant and method for making a medical implant

1.1. claims: 1-3

Medical implant which is vacuum-packed within an airtight cover material.

1.2. claims: 4-13

Medical device for making a medical implant covered by an airtight cover material, wherein the medical device comprises at least an underpressure generating device and a sealing device, wherein the sealing device is arranged for airtight sealing of the cover material while the medical implant is located within the cover material, wherein the underpressure generating device is coupled or can be coupled with a space which is surrounded by the cover material, and wherein the underpressure generating device and the sealing device are controllable such that after generating a certain underpressure within the space surrounded by the cover material through the underpressure generating device the sealing device is actutable to seal the cover material in an airtight manner to form an airtight protecting cover around the medical implant located therein.

1.3. claims: 14-22

Method for making a medical implant covered by an airtight cover material, comprising the steps: a) providing the medical implant, b) providing sterile cover material in the form of at least one or at least two pieces, c) placing the medical implant within a space surrounded by at least one or at least two pieces of the cover material, d) creating an underpressure within the space surrounded by the at least one or at least two pieces of the cover material by evacuating air from this space, e) airtight sealing of the cover material.

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