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Kucklick

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(54) **METHOD AND DEVICES FOR
IMPLANTATION OF BIOLOGIC
CONSTRUCTS**

(58) **Field of Classification Search**
CPC ... A61M 39/0606; A61M 29/00; A61M 25/04
See application file for complete search history.

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(56) **References Cited**

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U.S. PATENT DOCUMENTS

(73) Assignee: **Cannuflow, Inc.**, Campbell, CA (US)

(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 55 days.

5,250,033	A *	10/1993	Evans	A61M 25/0668
					604/160
6,663,599	B2 *	12/2003	Osbourne	A61M 39/0606
					604/167.04
7,169,130	B2 *	1/2007	Exline	A61B 17/3462
					604/167.01
7,435,236	B2 *	10/2008	Weaver	A61M 1/3653
					604/167.03
7,938,805	B2 *	5/2011	Harding	A61M 39/0606
					251/334
8,096,976	B2 *	1/2012	Sugiki	A61M 39/0606
					604/167.04
9,044,541	B2 *	6/2015	Blanchard	A61M 5/16881
2005/0049555	A1 *	3/2005	Moorehead	A61M 25/0075
					604/122

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Related U.S. Application Data

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(60) Provisional application No. 61/324,746, filed on Apr. 15, 2010.

(51) **Int. Cl.**
A61M 39/06 (2006.01)
A61F 2/00 (2006.01)
A61B 17/04 (2006.01)
A61B 17/06 (2006.01)

(52) **U.S. Cl.**
CPC *A61M 39/0606* (2013.01); *A61F 2/0063* (2013.01); *A61B 17/0469* (2013.01); *A61B 17/0483* (2013.01); *A61B 17/06061* (2013.01)

* cited by examiner

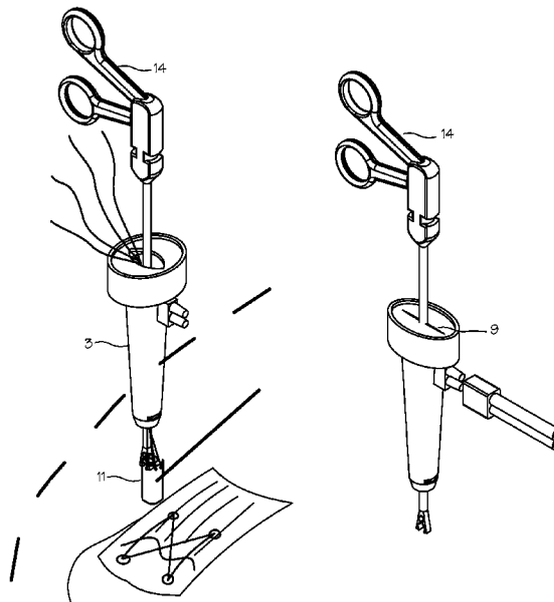
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(57) **ABSTRACT**

Seal systems and devices for delivering a biologic construct repair for shoulder and joint surgery into an arthroscopic workspace, including a pinch to open seal that uses fluid pressure to self-seal against a curved surface when closed.

12 Claims, 9 Drawing Sheets



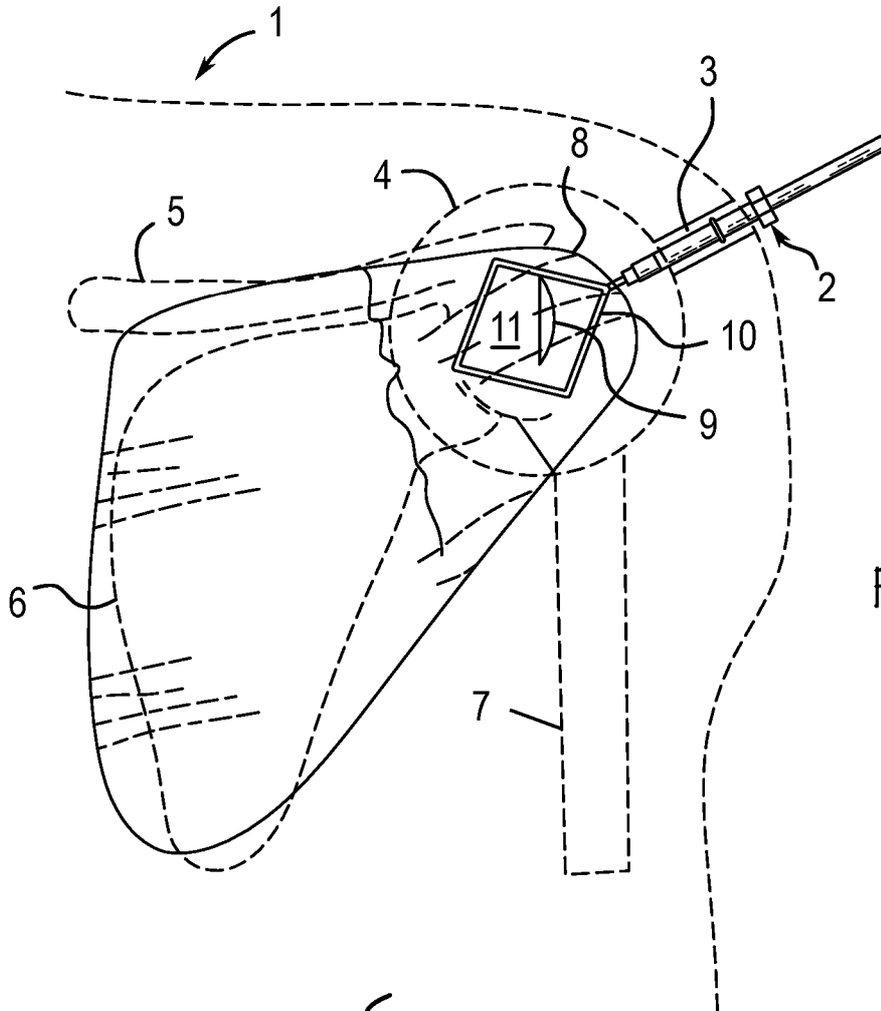


Fig. 1A

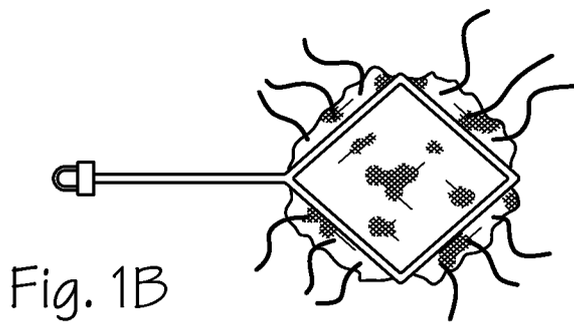


Fig. 1B

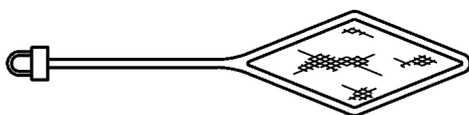


Fig. 1C

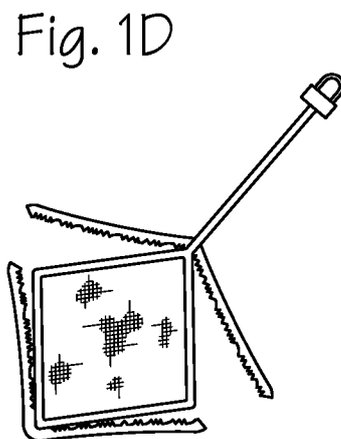


Fig. 1D

Fig. 2A

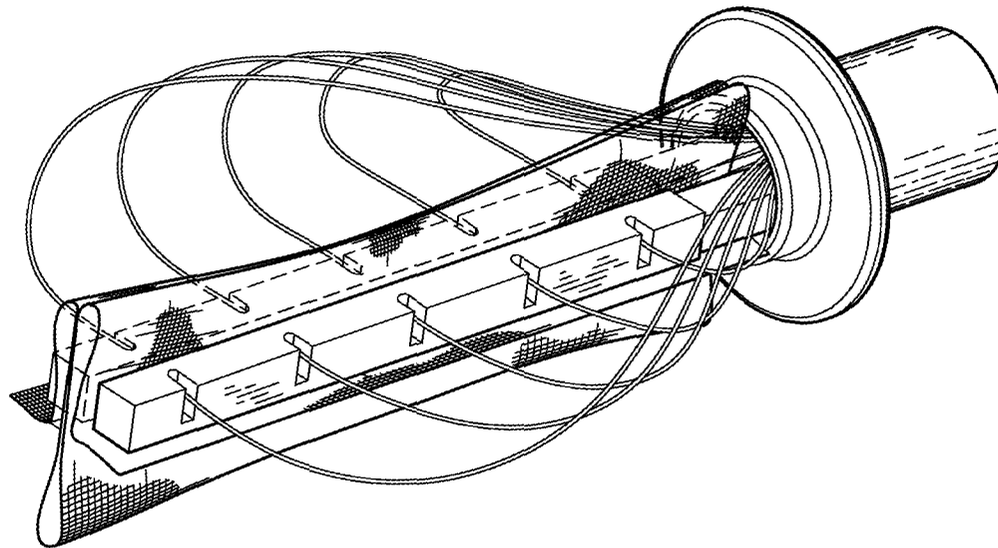


Fig. 2B

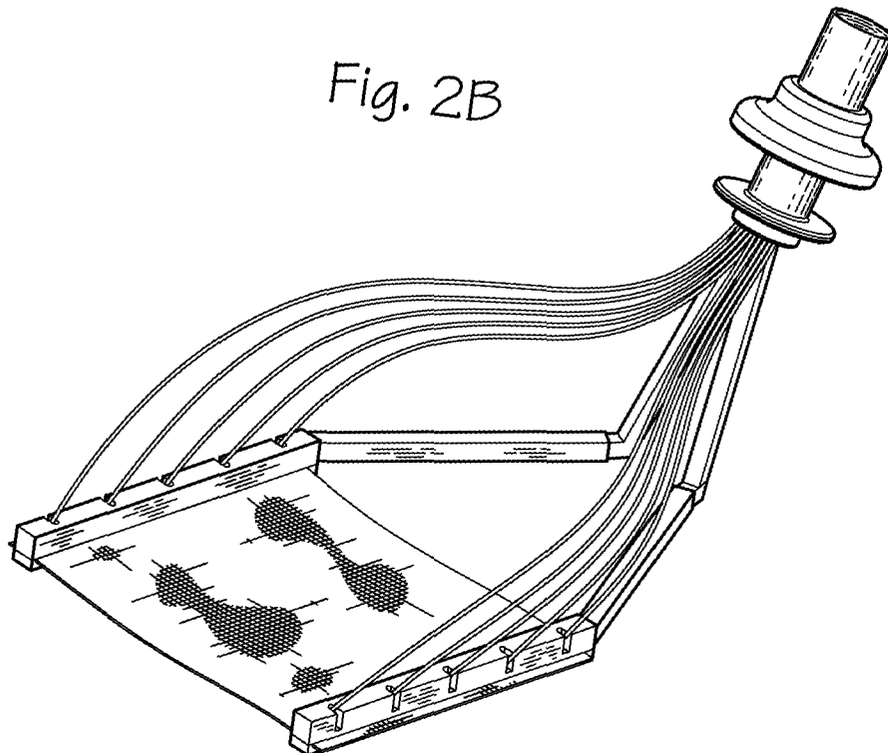


Fig. 3A

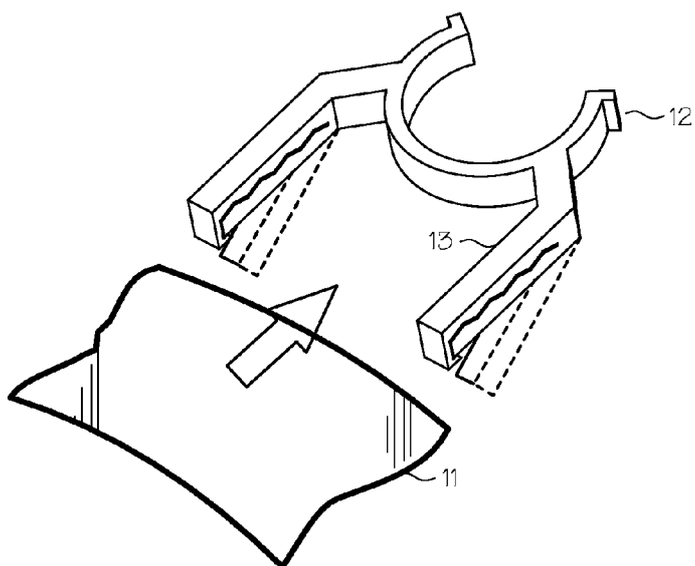


Fig. 3B

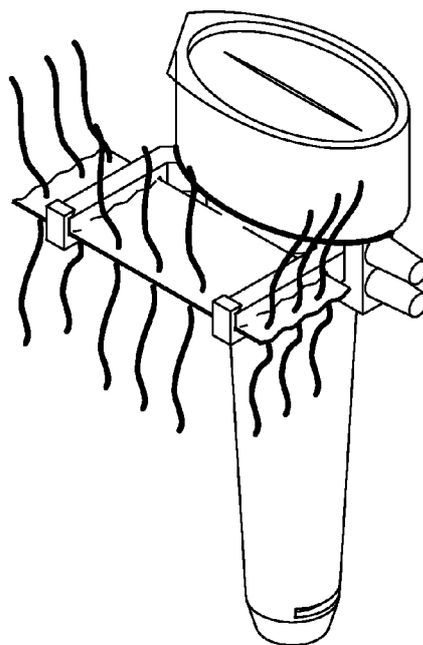


Fig. 4A

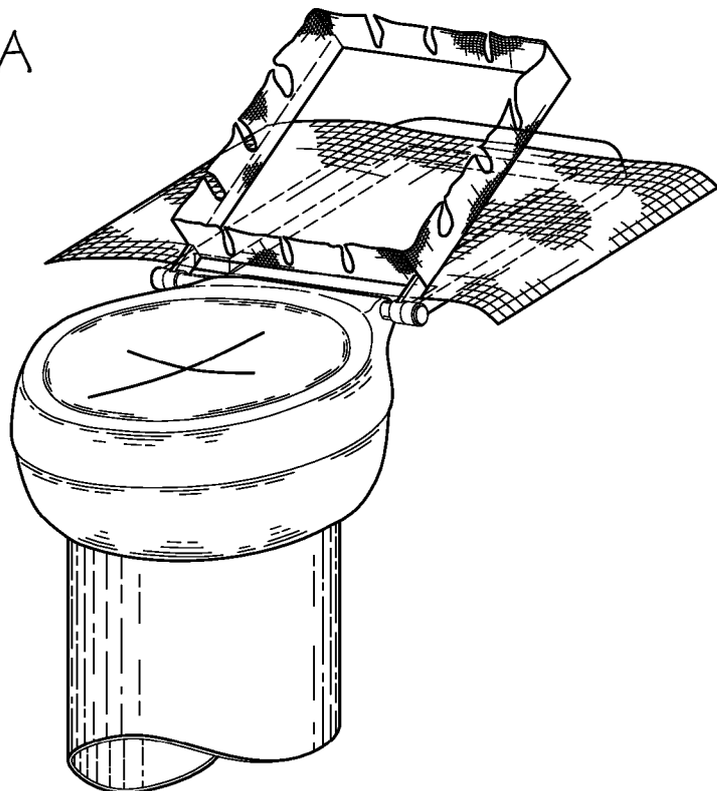


Fig. 4B

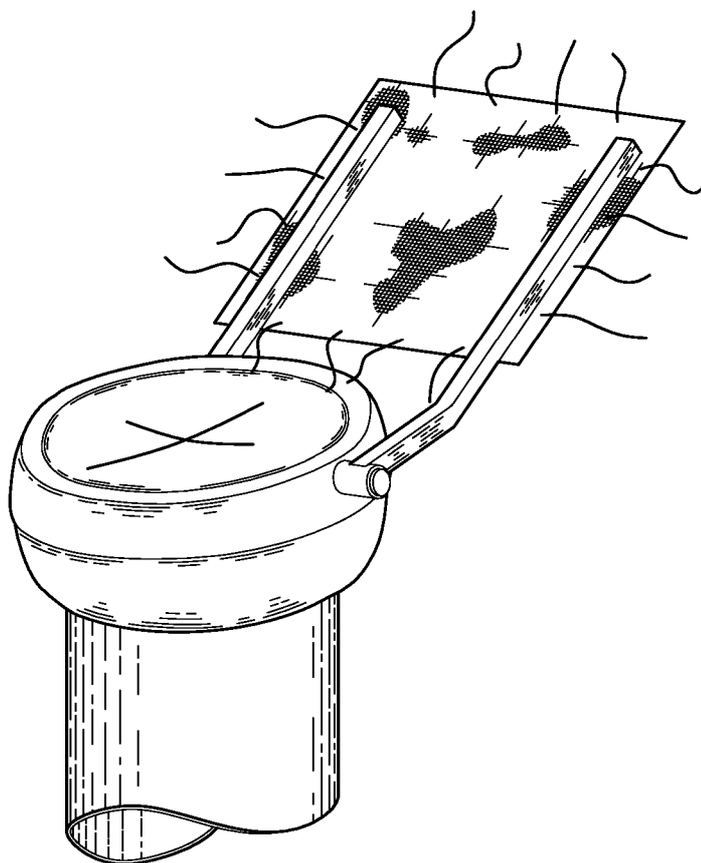


Fig. 5A

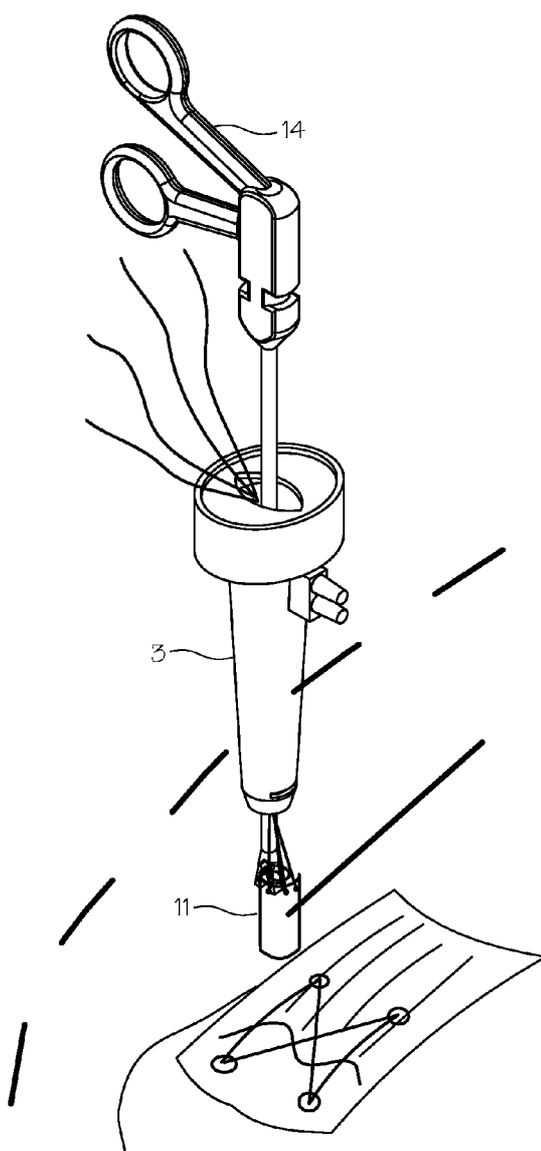
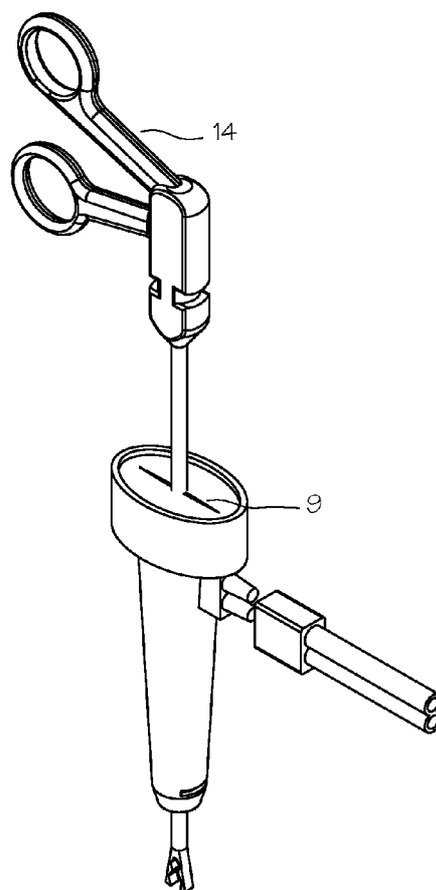


Fig. 5B



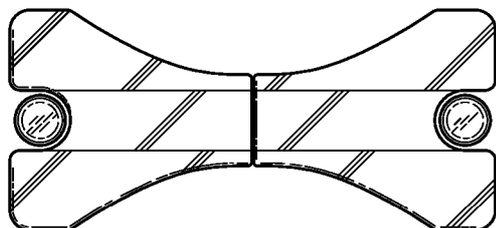


Fig. 6A

Fig. 6B

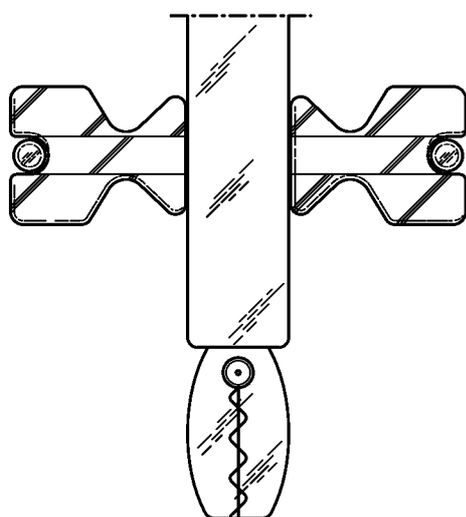
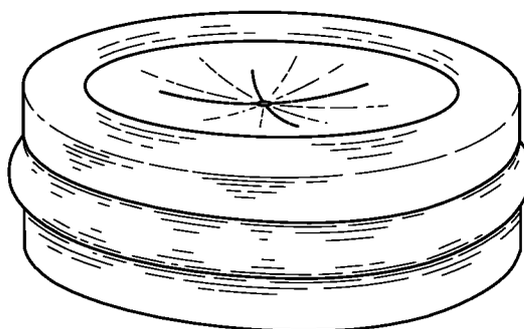


Fig. 6C

Fig. 7A

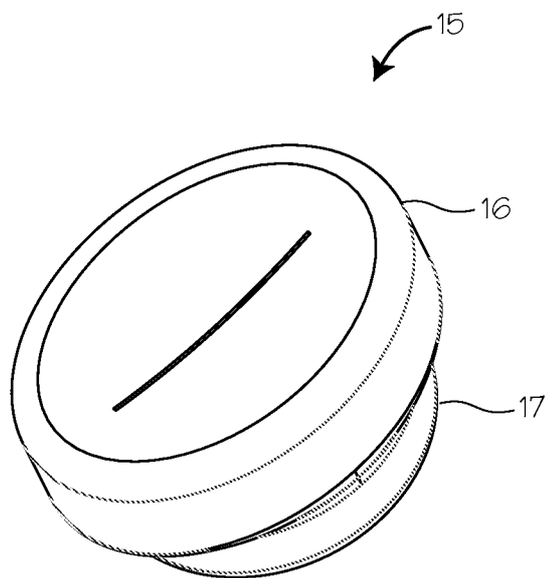


Fig. 7B

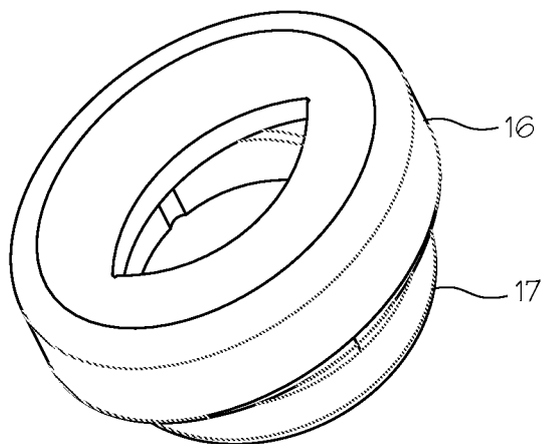


Fig. 8A

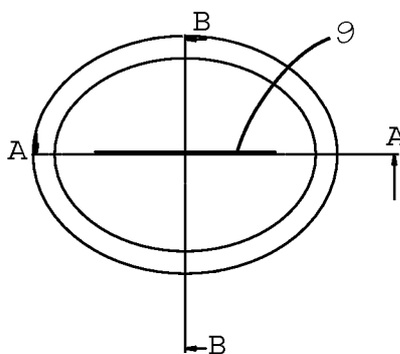
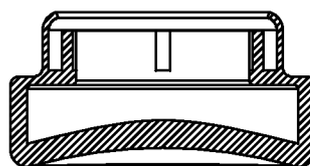
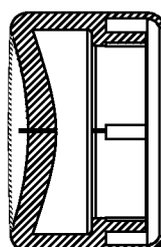


Fig. 8B



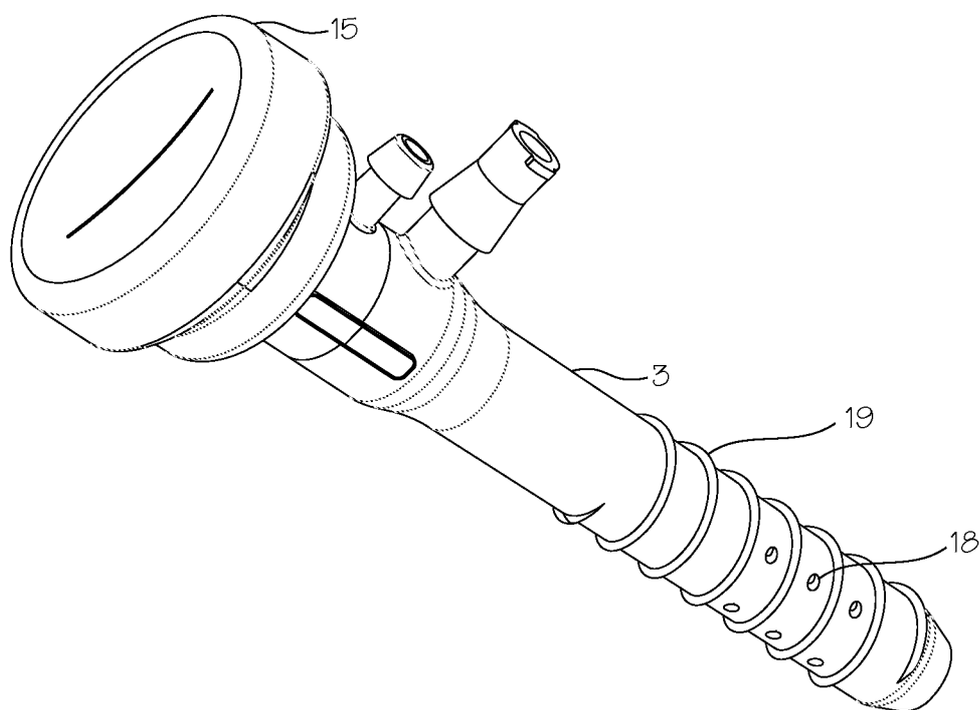
Section A-A

Fig. 8C



Section B-B

Fig. 9



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METHOD AND DEVICES FOR IMPLANTATION OF BIOLOGIC CONSTRUCTS

RELATED APPLICATIONS

This application is a continuation in part of application Ser. No. 13/088,327, filed Apr. 15, 2011, now U.S. Pat. No. 8,585,773, which claims priority to U.S. Provisional Patent Application 61/324,746 filed Apr. 15, 2010.

FIELD OF THE INVENTIONS

The inventions described below relate to the field of arthroscopic surgery and more specifically to implantation of biologic constructs.

BACKGROUND OF THE INVENTIONS

Biologic constructs, which include “regenerative tissue matrices” “acellular tissue matrices” or “collagen scaffolds” and other such substances, are small sheets or disks of flexible, engineered biomaterial, such as collagen or fibrin, decellularized dermal matrix crosslinked and sterilized xenograft tissues, and “platelet-rich-plasma” patches sometimes totally devoid of living cell material and sometimes loaded with active adjuncts such as biologic growth factors. Biologic constructs may also be tissue-engineered from mesenchymal stem cells. For the purposes of this application, the term biologic construct refers to any sheet-like or disc-like configuration of regenerative tissue matrix, acellular tissue matrices or collagen scaffolds, or similar materials suitable for placement within the body to promote healing.

Biologic constructs and implants are growing rapidly in popularity for the repair of joint pathologies. Biologic constructs may be a new alternative for carpal arthroplasty in patients with clinical conditions such as osteonecrosis. Biologic constructs are a family of biologically derived implants to promote tissue growth or to patch and repair tissue defects and tears. These include the repair of arthritic cartilage, the joining of tendons to bone and the bridging of degenerated rotator cuff in the shoulder. Biologic constructs, and graft material such as platelet rich fibrin membrane, acellular dermal allograft, (MTF) and xenograft materials (Pegasus Biologics) and graft patches (Wright Medical Graft jacket) have enabled the reconstruction and treatment of previously untreatable and irreparable musculoskeletal injuries and pathologies. Biologic constructs now occupy an increasingly important place in the orthopedic surgeons armamentarium.

One of the key problems with biologic constructs is that the delivery instrumentation has not kept pace with advances in these implants. For example, fluid seals effectively hold fluid, but do not allow passage of sutures and metal instruments through the biologic constructs without tearing and damage. This can render the construct useless, and add significantly to the cost of the case, as these implants can be fragile as well as expensive. A damaged implant can result in several hundred dollars of added expense.

In addition to these constructs being soft, floppy, and prone to damage from tearing, suture management is very difficult, both during the preparation of the implant outside the patient as well as managing the implant and sutures when the implant is manipulated and attached in the joint. Currently an implant construct is hand-held with hemostats by an assistant while the graft is prepared with sutures.

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An important clinical need exists to make these biologic constructs easier to prepare, handle, deliver, and implant so that the potential of these important advances in biologics can be fully realized.

SUMMARY

The present invention solves the most important problems with biologic constructs, that is, preparation of the graft prior to insertion into the patient, the ability to pass the fragile implant into the joint space without damaging or tearing the implant, and being able to manipulate the soft and floppy implant within the joint space. All of these improvements to the delivery instrumentation will reduce costs, reduce risks to the patient, and greatly improve the speed and success of the procedure.

There are four basic components to the system: 1) having a frame to prepare the graft outside the body, 2) having a seal system that allows passage of a large, soft construct without damage, and has the ability to be fluid-tight 3) a means of deploying and orienting the construct within the body and 4) a means of managing the sutures.

There are two basic procedures being addressed: one is to bridge a massively retracted and irreparable rotator cuff tear, (with a sheet of repair material) and the other is to augment a tendon to bone repair in order to promote healing of tendon to bone (platelet rich patch).

BRIEF DESCRIPTION OF THE DRAWINGS

FIGS. 1A, 1B, 1C, and 1D illustrate the kite.

FIGS. 2A and 2B illustrate the cage.

FIGS. 3A and 3B illustrate the implant frame and clamp.

FIGS. 4A and 4B illustrate the suture management accessory.

FIGS. 5A and 5B illustrates the coin purse cannula.

FIGS. 6A, 6B, and 6C illustrate the sphincter seal.

FIGS. 7A and 7B illustrate the features of the coin purse cannula illustrated in FIGS. 5A and 5B.

FIGS. 8A, 8B and 8C illustrate the specific features of the seal of FIGS. 7A and 7B in greater detail.

FIG. 9 illustrates a seal system that allows passage of an instrument or a large biological construct through the seal.

DETAILED DESCRIPTION OF THE INVENTIONS

The methods and devices described below provide for convenient prepping and delivering a biologic construct repair for shoulder and joint surgery into an arthroscopic workspace. There are four basic components to the system: 1) a frame to prepare the graft outside the body, 2) a seal system that allows passage of a large, soft construct without damage, and has the ability to be fluid-tight 3) a means of deploying and orienting the construct within the body and 4) a means of managing the sutures.

FIGS. 1B, 1C, 1D, and 1A show a “kite” graft repair and delivery system. The “kite” is a loop of superelastic or shape memory nickel-titanium with clamps that hold the construct, and is delivered through a tube and deployed inside the body. This design calls for a nitinol or hinged frame with integrated clamps that allow the biologic construct sheet to be folded, and delivered either through a surgical wound portal, or a cannula device to a site to be repaired, such as a torn rotator cuff in a shoulder. This design allows the graft to be prepped outside the body with sutures, and held in a collapsible frame for handling, and to be deployed within a

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body. The frame has a feature to allow it to be releasably attached to a handle for insertion through a surgical wound or cannula.

As shown in FIG. 1A, in use, the surgeon delivers the biologic construct to a joint within the body of a patient **1** by creating an arthroscopic workspace **4** around the joint and inserting a cannula **3** through the skin of the patient proximate the arthroscopic workspace. A sheet of biologic construct **11** is provided. A collapsible first frame **10** is configured to hold the sheet of biologic construct **11**. The surgeon attaches the sheet of biologic construct to the collapsible first frame, collapses the collapsible first frame with the sheet attached such that the frame and sheet fit, in a collapsed configuration, through the cannula. The surgeon inserts the collapsible first frame and sheet, in the collapsed configuration, through the cannula **3** and into the arthroscopic workspace; expanding the collapsible first frame and sheet within the workspace and positioning the sheet proximate an intended site of implantation; releasing the sheet from the collapsible first frame, collapsing the collapsible first frame and removing the collapsible first frame from the workspace; and securing the sheet to body tissue within the workspace.

The delivery device shown in FIG. 1A through 1D comprises an elongate insertion portion **2** having a proximal end and a distal end adapted for insertion into an arthroscopic workspace through a cannula or small incision proximate the workspace; a grasping member disposed on the distal end of the elongate insertion portion, wherein the grasping member comprises a collapsible frame comprising a first frame member which is collapsible from a wide configuration which tensions the sheet into a substantially flat configuration to a narrow configuration adapted for insertion into arthroscopic workspace through the cannula or small incision proximate the workspace. The grasping member further comprise a clamping member adapted to trap the sheet between the clamping member and the first frame.

The delivery device may also comprise an elongate insertion portion **2** having a proximal end and a distal end adapted for insertion into an arthroscopic workspace through a cannula or small incision proximate the workspace and a grasping member disposed on the distal end of the elongate insertion portion, with the grasping member comprising a collapsible frame comprising a first frame member which is collapsible from a wide configuration which tensions the sheet into a substantially flat configuration to a narrow configuration adapted for insertion into arthroscopic workspace through the cannula or small incision proximate the workspace. The grasping member further comprises a clamping member adapted to trap the sheet between the clamping member and the first frame.

The first frame member can be rectangular, and attached to the elongate insertion portion at a corner of frame. The first frame member can be rectangular, and attached to the elongate member at a corner of frame, and the clamping member also comprises an articulating arm that may be selectively apposed to a side of the rectangular frame to trap the sheet between the clamping member and the frame. The first frame member can be rectangular and attached to the elongate member at a corner of frame and be hinged at the remaining three corners of the frame. The first frame member can be rectangular comprising a superelastic material. The first frame member can also be circular or elliptical.

Another embodiment is where the holders for the construct are "legs" made of a preshaped spring material. This embodiment is shown in 2A and 2B. Inserted in the tube in a closed position, implant is folded up on the end. At the

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surgical site, spreads open inside, for example, the shoulder, at the surgical site. The legs spring open, which puts the implant in an open unfolded position. This device comprises an elongate insertion portion having a proximal end and a distal end adapted for insertion into an arthroscopic workspace through a cannula or small incision proximate the workspace and a pair of grasping jaws disposed on the distal end of the insertion portion. The pair of grasping jaws pivotably engaged with the elongate insertion portion such that they may be moved in away from each other to an open configuration and tension the biologic construct to a flat configuration, and closed in apposition to each other to fold the biologic construct between the pair of grasping jaws. Also, each of the pair of grasping jaws is operable to grasp the biologic construct and release the biologic construct. At least one of the grasping jaws includes a plurality of slots adapted to accept a suture segment attached to the biologic construct, as shown in FIG. 2A.

The grasping jaws can be made of a superelastic or resilient material biased to the open configuration, such that the grasping jaws may be forced in apposition to each other to fit into a cannula and thereafter superelastically or resiliently open upon exit from a distal end of the cannula.

FIGS. 3A and 3B show an implant graft holder. The graft holder holds a tissue construct patch **11** in place for suturing. Currently, an assistant has to hold the patch in the air with two hemostats. Instead of requiring assistance, the present implant graft holder includes a clamp **12** and frame **13** that are detachable from the cannula for graft prep, and easily removable when the suturing procedure is complete. The implant frame holds the implant grafts for the surgeon while the surgeon places the sutures in the graft. The clamp removably clips to cannula, as shown in FIG. 3B. The implant clamps into the frame. The frame is flexible and generally made of injection moldable plastic or other suitable material. The frame holds the implant in place for easy suture placement.

A second frame is adapted to hold the sheet to the cannula, secure the sheet to the second frame, secure a number of suture segments to the sheet while the sheet is secured to the second frame, and thereafter transfer the sheet to the collapsible first frame and perform the inserting step. The sheet can also be secured to body tissue within the workspace with the suture segments with an interrupted suture technique.

FIGS. 4A and 4B illustrate an accessory that allows a biologic construct sheet to be secured in a flat form, to be prepared with sutures for sewing into a defect, and a means for these sutures to be sorted and managed to enable efficient knot-tying and prevent tangling of the sutures as the prepared construct is delivered into the surgical space and secured into the tissue defect. Detachable allows back table prep and transfer.

FIGS. 5A and 5B show a cannula and seal system that provides for a way to pass a larger object through a cannula. Traditional septum seals are too tight for this. The seal opening at the end of the cannula is oval in shape and made of a flexible material that when pinched, opens the slit **9**. When the surgeon lets go, the slit seals back up again. This provides for a way to pass larger or fragile items like biologic graft patches and gelatinous platelet blob implants into the surgical site. As shown in FIG. 5A, the grasper **14** is inserted into the seal at the top of the cannula, with the implant **11** attached with the grasper, the grasper is inserted into the cannula **3** and the graft patch is delivered to the rotator cuff **6** at the repair site.

The "coin purse" seal shown in FIGS. 5A and 5B are part of a system that allows a large, floppy biologic construct to

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be prepared outside the patient, passed through a cannula, and arranged in place inside a body arthroscopically. The “coin purse” cannula is one embodiment that allows passage of a large soft object like a biologic sheet, and still allows the seal to close and prevent the leakage of fluid from the joint. The coin purse cannula seal allows for passage of larger objects into the joint, e.g. biologics and tissue implants. When the seal is closed, it creates a fluid seal and distension. The septum opens to allow passage of large implant constructs that do not fit thru conventional cannulae and allows access to endoscopic tissue repair sites, e.g. rotator cuff repair 6.

Details of the coin purse seal are illustrated in FIGS. 7A and 7B and FIGS. 8A, 8B and 8C. The seal 15 is a single unitary piece having a first section 16 with an oval or elliptical top face. The seal surface is inwardly curved as illustrated in FIG. 7A. The first section has a central inwardly curving or parabola shape that curves towards a slit on the top face. The seal also has a second section 17 integrally formed to the first section that includes a narrower diameter than the first section. The second section of the seal is shaped to allow the seal to be secured to the proximal end of a cannula. The seal is made of an elastomeric material that opens along the slit upon application of pressure, or pinching of the seal, as illustrated in FIG. 7B. The seal defaults to a closed position when the seal is no longer pinched, as illustrated in FIG. 7A. Pressure applied to the seal along the major axis of the ellipse, or perpendicular to and along the top and bottom of the slit, allows the seal to be squeezed into an open position to allow passage of instrument or biological constructs. The configuration of the seal allows the seal to act as a valve so that when in a closed position, hydrostatic pressure on the convex inner surface of the seal pushes the seal closed. The fluid pressure against the inner convex seal surface maintains the seal in a closed position. The curved surface causes the lips of the seal to open inward and allow larger objects to pass through and then close when the seal is not being pinched open. Water pressure acts on the inner curved surface to force the seal lips together and form a tighter seal as the water pressure increases. The seal may also include a circular wiper seal distal to the pinch open seal.

FIGS. 8A, 8B and 8C illustrate the specific features of the seal of FIGS. 7A and 7B in greater detail. FIG. 8A shows a top view of the seal having a top surface in the shape of an oval or ellipse. FIG. 8B illustrates cross section A-A of FIG. 8A along the major axis of the ellipse, also showing a central inwardly curved or parabola shape that curves towards the slit on the top face. FIG. 8C illustrates the cross section B-B of FIG. 8A along the minor axis of the ellipse, also showing the parabola surface that curves towards the slit in the central top face of the seal. Water pressure exerted on the convex surface of the parabola maintains the seal in a closed position and maintains the seal in a fluid-tight position so that no fluid passes through the seal until pressure is applied and the seal is opened by the user.

Another seal design is shown in FIGS. 6A, 6B, and 6C. This seal acts like a sphincter. The seal has a low durometer of 10-30 A with a stiffer o-ring of 40-80 A. The center seal stretches while the O-ring pulls the seal closed when the instrument is removed. This dual-compliance system allows easy passage of instrument and sealing of septum in “sphincter-like” manner. It is a “dual state” seal that is tight and leak resistant when the O-ring is in place, and more compliant when the O-ring is slipped out of its groove. The O-ring may be slipped back in the groove for a tighter seal once the implant is passed through. This also allows passage of large,

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soft implants without damaging the implant, and allows returning to a fluid-tight seal state.

FIG. 9 illustrates a seal system that allows passage of an instrument or a large biological construct through the seal. The seal 15, detailed in FIGS. 7 and 8, is removably attached to a cannula 3. The cannula includes openings 18 to allow for fluid flow through the cannula and out of the openings. Windings 19 on the outside of the cannula also allow for securement of the cannula in position.

In use, the seal 15 is affixed to the proximal end of a cannula and whereupon the distal end of the cannula is delivered through a surgical wound portal to a site to be repaired such as a torn rotator cuff in a shoulder. When an arthroscopic instrument or biological construct needs to be introduced to the repair site, pressure is applied along the major axis of the seal (or perpendicular to the slit) to open the lips or slit on the top face of the seal. The applied pressure breaks the hydrostatic pressure exerted under the slit and allows access to the repair site for introduction of instruments or biologic constructs to the repair site. When access to the repair site is no longer needed, pressure is removed from the first section, the slit or lips close and remain closed under the hydrostatic pressure applied to the parabola surface of the first surface.

While the preferred embodiments of the devices and methods have been described in reference to the environment in which they were developed, they are merely illustrative of the principles of the inventions. The elements of the various embodiments may be incorporated into each of the other species to obtain the benefits of those elements in combination with such other species, and the various beneficial features may be employed in embodiments alone or in combination with each other. Other embodiments and configurations may be devised without departing from the spirit of the inventions and the scope of the appended claims.

I claim:

1. A seal system for use with an arthroscopic instrument comprising:

a seal having a first section with a top face, the top face having a central inwardly curved or parabola shape that curves towards a slit in the top face, said slit characterized by a longitudinal axis, said seal having a second section integrally formed with the first section, said second section having a narrower outer diameter than the first section; and

a cannula for conducting fluid therethrough, said cannula having a proximal end;

said seal second section releasably attached to the cannula proximal end; and

wherein the seal first section is moveable from a first closed configuration to a second open configuration said seal operable to attain said open configuration upon application of compressive force along the longitudinal axis of the slit, and operable to revert to the closed configuration upon removal of said compressive force.

2. The seal system of claim 1 further including a circular wiper seal.

3. The seal system of claim 1 wherein the top face is in the shape of an ellipse, wherein the slit is aligned along a major axis of the ellipse.

4. The seal system of claim 1 wherein the top face is in the shape of an oval, wherein the slit is aligned along a major axis of the oval.

5. The seal of claim 4 further including a circular wiper seal.

6. The seal of claim 4 wherein the top face is in the shape of an ellipse, wherein the slit is aligned along a major axis of the ellipse.

7. The seal of claim 4 wherein the top face is in the shape of an oval, wherein the slit is aligned along a major axis of the oval. 5

8. The seal system of claim 1 wherein the seal is hollow.

9. The seal system of claim 1 wherein the first section has an inner diameter and wherein the second section has an inner diameter, wherein the second section inner diameter is narrower than the first section inner diameter. 10

10. A seal for use with a cannula, said seal comprising: a seal having a first section with a top face, the top face having a central inwardly curved or parabola shape that curves towards a slit in the top face, said slit characterized by a longitudinal axis, said seal having a second section integrally formed with the first section, said second section having a narrower outer diameter than the first section; and 15

wherein the seal first section is moveable from a first closed configuration to a second open configuration said seal operable to attain said open configuration upon application of compressive force along the longitudinal axis of the slit, and operable to revert to the closed configuration upon removal of said compressive force. 20 25

11. The seal of claim 10 wherein the seal is hollow.

12. The seal of claim 10 wherein the first section has an inner diameter and wherein the second section has an inner diameter, wherein the second section inner diameter is narrower than the first section inner diameter. 30

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