UNITED STATES PATENT AND TRADEMARK OFFICE

BEFORE THE PATENT TRIAL AND APPEAL BOARD

ARTHREX, INC.,
Petitioner,

v.

KFX MEDICAL, LLC.,
Patent Owner.

Case IPR2016-01698
Patent 8,926,663 B2


MARSCHALL, Administrative Patent Judge.

DECISION
Instituting of Inter Partes Review
37 C.F.R. § 42.108
I. INTRODUCTION

Arthrex, Inc. (“Petitioner”) filed a Petition for *inter partes* review of claims 1–18 of U.S. Patent No. 8,926,663 B2 (Ex. 1003, “the ’663 patent”). Paper 1 (“Pet.”). KFx Medical, LLC (“Patent Owner”) filed a Preliminary Response. Paper 6 (“Prelim. Resp.”). Institution of an *inter partes* review is authorized by statute only when “the information presented in the petition . . . and any response . . . shows that there is a reasonable likelihood that the petitioner would prevail with respect to at least 1 of the claims challenged in the petition.” 35 U.S.C. § 314(a); see 37 C.F.R. § 42.108. For the reasons set forth below, we conclude that the information presented in the Petition establishes a reasonable likelihood that Petitioner will prevail in showing the unpatentability of claims 1–18. Accordingly, we institute an *inter partes* review as to those claims.

A. Related Matters


B. The ’663 Patent

The ’663 patent is titled “System and Method for Attaching Soft Tissue to Bone.” Ex. 1001, Title. The ’663 patent summarizes its disclosed invention as follows:

The present invention is particularly suited for use in arthroscopic procedures, including but not limited to rotator cuff surgery. More broadly, it can be used in any procedure in which it is desired to fix a suture to a solid object without tying of knots,
including not only arthroscopic procedures, but also open surgery, and can be used for such diverse purposes as bladder neck suspension, tendon and ligament affixation or repair, prosthetic attachment, and rotator cuff repair.

*Id.* at 1:56–63.

Figures 11–13 of the ’663 patent are reproduced below:

![Figure 11](image1.png)  
**FIG. 11**

![Figure 12](image2.png)  
**FIG. 12**

Figure 11 shows a piercing bone anchor tip; Figure 12 shows an inserter for inserting a piercing bone anchor into bone; and Figure 13 shows the interface between a piercing bone anchor and an anchor inserter. *Id.* at 3:40–44. A piercing bone anchor includes: (1) anchor inserter attachment structure 674, which forms a proximal portion of the anchor; (2) structure 670 with suture aperture 672, which forms a medial portion; and (3) anchor tip 656, which constitutes a distal portion. *Id.* at 10:45–11:5. Piercing bone anchor 704 also includes hollow cylinder 650, which facilitates attachment of the bone anchor to outer sleeve 702 of the inserter. *Id.* at 11:19–30. Hollow cylinder 650 surrounds inner tube 720 of the inserter, which provides structural reinforcement of anchor 704 and pushes against anchor
tip 656 when driving the anchor into bone. *Id.* at 11:32–37. After driving the bone anchor into bone, pressing deployment lever 706 of the insert deploys and detaches anchor 704. *Id.* at 11:10–12.

The ‘663 patent further explains the following regarding inner tube 720:

The inner tube 720 may be fixed relative to the handle 712 and outer sleeve 702 during insertion, however, during deployment of the anchor 704, the inner tube 720 may be released by switching safety switch 708 so that the inner tube 720 can move axially relative to the outer sleeve 702 while the anchor cylinder 650 collapses.

*Id.* at 11:37–43.

Figure 10B of the ‘663 patent is reproduced below:

FIG. 10B

Figure 10B depicts a piercing bone anchor in a deployed state. *Id.* at 3:38–39. In that deployed state, cylinder 650 (not numbered in Figure 10B) collapses into a configuration in which lateral wings 660 of the cylinder prevent removal of the anchor from the bone. *Id.* at 10:27–32.
C. Claims

Claims 1, 8, and 13 are independent. Claims 1 and 8 claim a “bone implant” and an “inserter and bone implant combination,” respectively, and claim 13 claims a “method of fixing a bone implant within bone.” Claims 2–7, 9–12, and 14–18 ultimately depend from one of those independent claims. Claims 1 and 8 are illustrative and are reproduced below:

1. A bone implant, comprising:
   a substantially hollow cylinder comprising bone-engaging protrusions configured to prevent removal of the implant from bone; and
   a bone-piercing tip that comprises
      a distal portion comprising a pointed tip configured to pierce through bone and having a maximum diameter at its proximal end,
      a medial portion comprising a transverse suture aperture and having a constant diameter along its length that is smaller than the distal portion maximum diameter, and
      a proximal portion having a constant diameter along its length that is smaller than the medial portion diameter and configured to couple to an inserter, wherein the proximal portion of the bone-piercing tip is configured to be positioned within a distal portion of the substantially hollow cylinder and not extend to a proximal end of the substantially hollow cylinder, wherein the bone-piercing tip has a length that is shorter than the length of the substantially hollow cylinder, and
      wherein the substantially hollow cylinder has an inner diameter that is smaller than the bone-piercing tip distal portion maximum diameter.

8. An inserter and bone implant combination, comprising:
   an inserter, comprising:
      a handle,
an outer sleeve coupled to the handle and fixed relative to the handle, and
an inner tube disposed within the outer sleeve; and

a bone implant coupled to the inserter, the bone implant comprising:

a substantially hollow cylinder having bone-engaging protrusions, and

a bone-piercing tip comprising a distal portion having a pointed tip, a medial portion having a suture aperture, and a proximal portion, wherein the pointed tip has a maximum diameter at its proximal end, the medial portion has a constant diameter along its length that is smaller than the distal portion maximum diameter, and the proximal portion has a constant diameter along its length that is smaller than the medial portion diameter;

wherein a portion of the inner tube extends distally through the substantially hollow cylinder and contacts the bone-piercing tip,

wherein the proximal portion of the bone-piercing tip is configured to be positioned within a distal portion of the substantially hollow cylinder and not extend to a proximal end of the substantially hollow cylinder and is removably coupled to the inserter,

wherein the length of the bone-piercing tip is shorter than the length of the substantially hollow cylinder,

wherein the substantially hollow cylinder has an inner diameter that is smaller than the bone-piercing tip distal portion maximum diameter, and

wherein a distal end of the outer sleeve is flush with an open, proximal end of the substantially hollow cylinder.
D. The Prior Art

Petitioner relies on the following prior art references:

<table>
<thead>
<tr>
<th>Reference</th>
<th>Date</th>
<th>Exhibit No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>U.S. Patent No. 5,527,342 issued to Pietrzak et al. (“Pietrzak”)</td>
<td>June 18, 1996</td>
<td>1007</td>
</tr>
<tr>
<td>U.S. Patent No. 5,141,520 issued to Goble et al. (“Goble”)</td>
<td>Aug. 25, 1992</td>
<td>1008</td>
</tr>
<tr>
<td>U.S. Patent No. 5,741,282 issued to Anspach, III et al. (“Anspach”)</td>
<td>Apr. 21, 1998</td>
<td>1021</td>
</tr>
</tbody>
</table>
E. Asserted Grounds of Unpatentability

Petitioner challenges claims 1–18 based on the following grounds (Pet. 19):

<table>
<thead>
<tr>
<th>Ground No.</th>
<th>Reference(s)</th>
<th>Basis</th>
<th>Challenged Claims</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Meridew</td>
<td>§ 102</td>
<td>1–10, 13–15, and 17</td>
</tr>
<tr>
<td>2</td>
<td>Meridew and Fucci or Schmieding</td>
<td>§ 103</td>
<td>11–12</td>
</tr>
<tr>
<td>3</td>
<td>Meridew and Anspach</td>
<td>§ 103</td>
<td>16</td>
</tr>
<tr>
<td>4</td>
<td>Meridew, Pietrzak or Goble, Fucci or Schmieding, and Anspach</td>
<td>§ 103</td>
<td>1–18</td>
</tr>
<tr>
<td>5</td>
<td>Burkhart and Pietrzak or Goble</td>
<td>§ 103</td>
<td>1–2, 5–10, 13–15, and 17–18</td>
</tr>
<tr>
<td>6</td>
<td>Burkhart, Pietrzak or Goble, and Fucci or Schmieding</td>
<td>§ 103</td>
<td>3–4 and 11–12</td>
</tr>
<tr>
<td>7</td>
<td>Burkhart, Pietrzak or Goble, and Anspach</td>
<td>§ 103</td>
<td>16</td>
</tr>
</tbody>
</table>

II. ANALYSIS

A. Claim Construction

In an inter partes review, a claim in an unexpired patent shall be given its broadest reasonable construction in light of the specification of the patent in which it appears. 37 C.F.R. § 42.100(b); Cuozzo Speed Techs., LLC v. Lee, 136 S. Ct. 2131, 2144–46 (2016) (upholding the use of the broadest
reasonable interpretation standard). Consistent with the broadest reasonable construction, claim terms are presumed to have their ordinary and customary meaning as understood by a person of ordinary skill in the art in the context of the entire patent disclosure. *In re Translogic Tech., Inc.*, 504 F.3d 1249, 1257 (Fed. Cir. 2007).

Here, neither party contends that any claim term should be afforded a meaning other than its ordinary and customary one. Petitioner proposes express construction of such meaning for certain claim terms and phrases. *See* Pet. 13–19. For purposes of this Decision, we need only address explicitly the construction of three claim phrases/terms: (1) “constant diameter along its length” (claims 1, 8, and 13); (2) “bone-piercing tip . . . comprising a pointed tip” (claims 1, 8, and 13); and (3) “inner tube” (claims 8 and 13).

1. “constant diameter along its length”

   Each of claims 1, 8, and 13 requires a medial portion and a proximal portion associated with a bone-piercing tip. Each of those portions is required to have “a constant diameter along its length.” According to Petitioner, that requirement should be broadly construed as encompassing a constant diameter along only “a portion of” the lengths of the proximal and medial portions. Petitioner, however, provides no meaningful explanation as to why the broadest reasonable interpretation of the pertinent phrase should, in effect, be read in a manner that inserts the phrase “a portion of” in front of the word “length.” It is apparent from the claim language itself that each of the medial and proximal portions must have a length, and that such length has a constant diameter along its extent.
Thus, although we agree with Petitioner (Pet. 15) that the “along its length” language does not limit the claims to details of the embodiment shown at Figure 11 of the ’663 patent, that does not justify a construction that permits a constant diameter of the medial and proximal portions that is something regarded as less than along their entire length.

2. “pointed tip”

Each of claims 1, 8, and 13 requires a “bone-piercing tip” having a distal portion with a “pointed tip.” According to Petitioner, the term “pointed” means “having a sharpened or tapered tip or end,” and that “[a] tip can be ‘tapered’ and still have a blunted or softened terminal end.” Pet. 16. Thus, Petitioner is of the view that a tip remains “pointed” even if it has a “blunted or softened terminal end.” That construction is unreasonable.

Petitioner’s contorted position that construction of the term “tapered,” which is not a claim term, infuses a meaning of “pointed tip” as encompassing a tip that is blunted, and thus clearly not pointed, cannot be correct. A blunted tip is essentially the opposite of a pointed tip. Petitioner’s position is contrary to the ordinary meaning of “pointed” and inconsistent with the specification of the ’663 patent, which discloses pointed tips lacking a blunted or softened terminal end in the embodiments covered by the claims of the ’663 patent. See, e.g., Ex. 1001, Figs. 10A, 10B, 11, 13. We, thus, do not adopt Petitioner’s construction of the term ‘pointed tip,’ but determine that, for purposes of this Decision, we need not make express the construction of that term.

3. “inner tube”

Petitioner provides no express construction of the claim term “inner tube.” Pointing to a dictionary definition (Ex. 2004), Patent Owner contends
that a “tube” must be a “hollow” structure. Prelim. Resp. 27. Patent Owner also notes that the specification of the ’226, in discussing inner tubes, describes structures that are hollow. *Id.* at 27–28. We agree with Patent Owner that in the context of the ’663 patent, a tube is something that is hollow. Thus, we construe “inner tube” as an inner structure that is hollow.

4. Remaining claim terms

We have given all claim terms, including those discussed above, their ordinary and customary meaning. We determine that it is not necessary to address or make express that meaning for any other claim term of the ’663 patent at this time. *See Vivid Techs., Inc. v. Am. Sci. & Eng’g, Inc.*, 200 F.3d 795, 803 (Fed. Cir. 1999) (“[O]nly those terms need be construed that are in controversy, and only to the extent necessary to resolve the controversy.”).

B. Grounds 1–3 – Anticipation and Obviousness Based on Meridew

1. Overview of Meridew

Meridew discloses an “Expanding Suture Anchor Having an Actuator Pin.” Ex. 1005, Title. Meridew summarizes its invention as “[a] method and apparatus for reattaching soft tissue to a pre-selected boney structure using an expanding suture anchor.” *Id.* at 1:32–35. Meridew’s Figures 1 and 2 are reproduced below:
Figure 1 depicts expanding suture anchor 10 having actuator pin 12. *Id.* at 2:13–14; 55–56. Figure 2 illustrates an exploded view of suture anchor 10. *Id.* at 2:15. Suture anchor 10 includes insert 18 molded to actuator pin 12 and partially disposed in sleeve 20. *Id.* at 2:61–62. Insert 18 includes suture receiving portion 22, with tip 32 and cylindrical body 34, and end section 24 displaced from portion 22 by breakaway section 26. *Id.* at 2:64–66; 3:12–13.
Meridew’s Figures 4A–4C are reproduced below:

Figures 4A–4C depict “environmental view[s]” of the suture anchor in use. *Id.* at 2:21–25. As shown Figures 4A–4C, suture anchor 10 is inserted into “pre-drilled hole 14” in boney structure 16. *Id.* at 4:52–56. Squeezing trigger 84 of actuation gun 56 applies retractive force F1 to actuator pin 12. *Id.* at 4:60–62. Meridew further explains the following:

The application of the retractive force F1 causes the insert 18 to displace rearwardly with respect to the sleeve 20 as illustrated in FIG. 4B. This rearward displacement causes the tapered portion 46 to apply a force F2 to the tapered interior bearing surface 80 of the expanding members 72 of the sleeve 20. As the insert 18 continues to move rearward, the tapered portion 46 applies an
increasingly greater force $F_2$ to the tapered interior bearing surface 80 of the expanding members 72 until the expanding members 72 are engaged with the boney structure 16.

*Id.* at 5:8–17. Once expanding member 72 fully engages with boney structure 16, subsequent application of retractive force $F_1$ causes breakaway section 26 to fracture as shown in Figure 4C. *Id.* at 5:18–23.

2. Discussion

For Ground 1, Petitioner argues that Meridew anticipates claims 1–10, 13–15, and 17. Pet. 19–36. For Grounds 2 and 3, Petitioner asserts that claims 11–12 and 16 are obvious based on Meridew in combination with additional prior art. *Id.* at 36–41. Grounds 1–3 all rest on Petitioner’s proposed interpretation of “pointed tip” as encompassing a tip that is blunted. As discussed above in connection with our claim construction, we decline to adopt Petitioner’s proposed interpretation. According to Petitioner, Meridew’s tip 32, shown for instance in Figures 1, 2, and 4A–4C reproduced above, is a pointed tip. *See,* e.g., Pet. 19–21. We agree with Patent Owner, however, that it is clear from those figures that tip 32 includes a flattened end portion. *See* Prelim. Resp. 22. That flattened end portion is not a “pointed tip.” We are cognizant that Meridew generally contemplates a possibility of some undescribed “impacting tip” that may be employed with a suture anchor. *See* Ex. 1005, 2:47–49. Meridew, however, provides no detail of the structure of an “impacting tip,” or further mention of such a tip. Based on this record, we find that Meridew does not expressly or inherently disclose the “pointed tip” required by all of the claims of the ’663 patent.

1 Force $F_2$ appears to be labeled “F3” in Figure 4B.
In addition, Petitioner does not assert, as part of Grounds 1–3, any modification of Meridew that would result in the claimed “pointed tip.” Rather, Petitioner bases Grounds 1–3 on the flawed position that Meridew’s tip 32 with a flattened end is, in its own right, a pointed tip.

For at least the reasons noted above, Petitioner has not established a reasonable likelihood of success in connection with Grounds 1–3.

C. Ground 4 – Obviousness Based on Meridew

Petitioner proposes an alternative ground to claims 1–18 based on Meridew. Although Petitioner initially styles Ground 4 applied to claims 1–18 as being collectively over “Meridew in view of Pietrzak or Goble, Fucci or Schmieding, and Anspach” (Pet. 19), Petitioner refines the statement of the ground in justifying its merits. In particular, Petitioner applies the references as follows: (1) claims 1–10, 13–15, 17, and 18 are rendered unpatentable over Meridew in view of Pietrzak or Goble (id. at 41–47); (2) claims 11 and 12 are rendered unpatentable over Meridew, Goble or Pietrzak, and Fucci or Schmieding (id. at 46); and (3) claim 16 is rendered unpatentable over Meridew, Goble or Pietrzak, and Anspach (id. at 46).

Accordingly, we consider the claims in the above-noted groupings.

1. Claims 1–10, 13–15, 17, and 18

For these claims, Petitioner asserts that even if Meridew fails to disclose a “pointed tip,” Pietrzak and Goble both disclose such a tip and it would have been obvious to modify Meridew to incorporate the pointed tip of either Pietrzak or Goble.

a. Overview of Pietrzak

Pietrzak discloses a “Method and Apparatus for Securing Soft Tissues, Tendons and Ligaments to Bone.” Ex. 1007, Title. Pietrzak’s
Figure 2 is reproduced below:

Figure 2 illustrates a side elevational view of “the apparatus for securing soft tissues, tendons and ligaments to bone” according to an embodiment of the invention. *Id.* at 3:1–4. Pietrzak describes suture anchor 10 with spear member 12 having cone-shaped head portion 14 that comes to “point 24 which is suitable for piercing and being driven into bone.” *Id.* at 5:16–22. Pietrzak sets forth that an object of the invention “is to provide a method and apparatus to secure a suture to bone without the need to pre-drill the bone nor require the use of a drill to install the anchor.” *Id.* at 2:45–49.

b. **Overview of Goble**

Goble discloses a “Harpoon Suture Anchor.” Ex. 1008, Title. Goble’s Figure 3 is reproduced below:

Figure 3 shows “an enlarged profile sectional view of a first embodiment of a harpoon suture anchor of the present invention.” *Id.* at 3:49–51. Goble
describes suture anchor mount 28 and suture anchor 21 that incorporates “a pointed forward end 35 . . . for penetrating a bone 33 surface.” *Id.* at 5:15–27 (bone surface 33 not shown in Figure 3). Goble describes that an “object of the present invention is to provide a suture anchor that can be installed in a bone that does not require a prior site preparation.” *Id.* at 2:20–23.

c. *Discussion*

In connection with the above-noted ground, Petitioner specifies where all the limitations of claims 1–10, 13–15, 17, and 18 are found in Meridew with a proposed modification of tip 32 so as to account for the feature of a bone-piercing tip having a pointed tip. *See* Pet. 41–47; *see also* *id.* at 21–36. Specifically, Petitioner contends that it would have been obvious to implement the pointed tip portions of the suture anchors of either Pietrzak or Goble into the suture anchor of Meridew. Pet. 43–46.

Although Meridew’s tip 32 does not form a pointed tip, the tips disclosed by Pietrzak and Goble do form pointed tips that fall within the scope of “pointed tip” in the claims of the ’663 patent.

Petitioner sets forth several reasons why it would have been obvious to modify the blunted end with the pointed ends of either Pietrzak or Goble. *Id.* 43–46. Based on the current record and for purposes of this decision, we adopt those reasons as our own.

As Petitioner points out, Meridew generally recognizes the possible use of an “impacting tip.” *Id.* at 43; Ex. 1005, 2:47–49. Further, it is apparent from each of Pietrzak and Goble that the tip of a suture anchor may be configured with a pointed tip operable to impact and pierce bone. *See* Ex. 1007, 5:1–23; Ex. 1008, 5:15–27. Pietrzak and Goble also make explicit that such a pointed tip, and its associated bone-piercing ability, provides a
benefit in eliminating the need to provide prior site preparation, such as the need to pre-drill a hole in the bone. See Ex. 1007, 2:45–49 (seeking “to secure suture to bone without the need to pre-drill the bone”); Ex. 1008, 2:20–23 (setting goal of providing suture anchor that can be installed without “a prior site preparation”). As discussed above, Meridew contemplates a pre-drilled hole prior to the use of the anchor. Ex. 1005, 4:52–56. Accordingly, by providing a tip that removes the need for a pre-drilled hole, both Pietrzak and Goble provide an alternative, pointed tip design with an advantage over the tip used in Meridew. On the record before us, there is adequate reason presented in the Petition that a skilled artisan would have modified Meridew’s tips to incorporate a tip understood as one that is bone-piercing with a pointed tip. Indeed, a skilled artisan seeking to eliminate the necessity of pre-drilling a hole in the bone has good reason to pursue the known, viable options that are conveyed by the prior art to facilitate such pursuit, i.e., implementing the pointed tips of either Pietrzak or Goble. See KSR Int’l Co. v. Teleflex Inc., 550 U.S. 398, 421 (2007) (“When there is a design need or market pressure to solve a problem and there are a finite number of identified, predictable solutions, a person of ordinary skill has good reason to pursue the known options within his or her technical grasp. If this leads to the anticipated success, it is likely the product not of innovation but of ordinary skill and common sense.”).

Patent Owner argues that Meridew fails to disclose the limitation of claims 1, 8, and 13 directed to a “proximal portion having a constant diameter along its length that is smaller than the medial portion diameter.” Prelim. Resp. 30–34. Petitioner identifies the portions of Meridew’s suture anchor that Petitioner believes corresponds to the claimed proximal, medial,
and distal portions. See, e.g., Pet. 20, 22–23. In considering those identified portions as shown, in Meridew’s Figure 2, we are satisfied, based on the current record, that Petitioner has made an adequate showing that Meridew discloses the claimed “proximal portion.”

With respect to claims 8–18, Patent Owner contends that Meridew does not disclose an inserter that includes an “inner tube” as required by those claims. Prelim. Resp. 23–28. Petitioner contends that Meridew’s end section 24, which includes tapered portion 46 and annular body 48, discloses the claimed inner tube. Pet. 27. Meridew’s Figures 2 and 4C (reproduced above) shows the end section and associated tapered portion and annular body. Meridew describes tapered portion 46 and annular body 48 as forming central bore 58 that receives actuator pin 12 therein. Ex. 1005, 3:56–58, Fig. 2.

Patent Owner first argues that these components do not form the claimed “inner tube” because they are “actually part of the suture anchor and therefore cannot be part of the claimed inserter.” Prelim. Resp. 23. According to Patent Owner, end section 24, as well as the end section’s tapered portion 46 and annular body 48, are “all part of insert 18, which is part of suture anchor 10” and, therefore, “cannot also be part of the claimed inserter.” Id. at 26. We disagree. Although end section 24 is initially associated with insert 18, once the insert is implanted within a bone, the end section is compelled, via retractive force F1 provided by actuator gun 56, to separate from insert 18 via breakaway section 26 and is removed from the bone via operation of actuator gun 56. Ex. 1005, 3:24–36. At this time, we are persuaded that because end section 24 is removed along with the inserter device formed by actuator gun 56, and does not remain implanted within
bone, the end section is regarded reasonably as a part of the inserter rather than part of the suture anchor as alleged by Patent Owner. Indeed, the primary function of end section 24 appears to be facilitating insertion of insert 18 within a bone.

Patent Owner’s second contention is that Meridew does not disclose the claimed “inner tube” because the structure relied on by Petitioner is not hollow and therefore not a “tube” at all. Prelim. Resp. 27–28. On the contrary, Meridew’s tapered portion and annular body together form central bore 58, and, thus, constitute a structure that is hollow.

Lastly, with respect to claims 13–18, Patent Owner contends that Meridew fails to teach a bone implant that is “coupled to a distal end of the inserter” as recited in claim 15. Prelim. Resp. 28–30. The premise of that contention is that components initially attached to insert 18, such as end section 24, but that ultimately are removed by operation of actuator 56, nevertheless, must be considered part of an implant rather than an inserter. Id. at 29. For purposes of this Decision, we do not agree with Patent Owner, and conclude, at this time, that the portions of Meridew’s insert 18 removed along with actuator 56 are viewed reasonably as part of an inserter. In that case, Meridew discloses a bone implant coupled to a distal end of the inserter, i.e., the distal end of section 24. See Ex. 1005, Fig. 2.

For the reasons given above, we are satisfied, on the present record, that Petitioner has shown that the combination of Meridew with either Pietrzak or Goble discloses all of the limitations of claims 1–10, 13–15, 17, and 18. We have considered Patent Owner’s additional arguments against the combination of Meridew with either Pietrzak or Goble and are not persuaded that institution should be denied for the reasons stated by Patent
Owner. See Pet. 46–51. We are satisfied, on the present record, that Petitioner has shown a reasonable likelihood of success in its challenge that claims 1–10, 13–15, 17, and 18 are unpatentable over Meridew and Pietrzak and unpatentable over Meridew and Goble.

2. Claims 11 and 12

Claims 11 depends from claim 10, which depends from claim 8. Claim 12 depends from claim 11. Claim 11 adds the limitation of a suture that “extends through an axial bore in the inner tube and an axial bore in the handle of the inserter.” Claim 12 further adds that “the suture exits the inserter through a hole in the proximal end of the handle.”

Petitioner contends that although such features are absent from Meridew, claims 11 and 12 would have been obvious when Meridew’s teachings are considered along with those of either Fucci or Schmieding. Pet. 46 (incorporating analysis at Pet. 36–39).

a. Overview of Fucci

Fucci discloses a “Suture Anchor Driver with Suture Retainer.” Ex. 1019, Title. Fucci’s Figures 1 and 7 are reproduced below:

![FIG. 1](image)
Figures 1 and 7 depict a suture anchor driver according to Fucci’s invention. As shown in Figure 1, driver 10 includes drive shaft 12 extending to distal end 16, and handle portion 18 at proximal end 20. \textit{Id.} at 4:26–30. Throughbore 22 extends from distal end 16 to proximal end 20. \textit{Id.} at 4:30–31. As illustrated in Figure 7, suture ends 38a and 38b of suture 38 (not numbered) “are passed through bore 22 of the cannulated driver from the distal tip 16 and out of the proximal end 20.” \textit{Id.} at 5:27–30. Fucci describes that driver 10 is intended for use with implants such as anchor 30 or anchor 30a (not shown in Figures 1 and 7 above), and that the driver “captures and secures either implant to itself by means of tension translated to the implant via suture 38 that is passed through the eyelet of the implant.” \textit{Id.} at 4:47–51.

\textit{b. Overview of Schmieding}

Schmieding discloses a “Corkscrew Suture Anchor.” Ex. 1020, Title. Schmieding’s Figure 4 is reproduced below:

Figure 4 shows a side view of a suture anchor assembly according to an embodiment of Schmieding’s invention. The suture anchor assembly includes pieces of suture 32 extending within driver 34 and within a handle
portion of the assembly and wraps around cleat 36. See id. at 4:45–47.

Schmieding sets forth that “[t]ension on the suture aids in retaining the suture anchor in the distal end of the driver.” Id. at 4:47–49.

c. Discussion

Petitioner contends that in light of the teachings of either Fucci or Schmieding, a person of ordinary skill in the art would have recognized that in connection with a suture anchor assembly, such as that of Meridew, suture may be arranged within the assembly such that it extends from a suture anchor to, and through, a hole in the proximal end of a handle. Pet. 36–39. According to Petitioner, a skilled artisan would have configured the handle of Meridew’s suture anchor assembly, and associated suture, in the manner required by claims 11 and 12 “in order to hold the eyelet implant in place under tension which increases the ease of insertion, as taught by Fucci or Schmieding.” Pet. 38 (citing Ex. 1027 ¶ 133). Petitioner also asserts that the suture-holding arrangements of Fucci and Schmieding “would perform their same functions when incorporated into the Meridew anchor as they do alone.” Id. at 39 (citing Ex. 1027 ¶¶ 134–135).

Patent Owner urges that Petitioner has not advanced any “reasonable rationale” for the proposed combination of Meridew and either Fucci or Schmieding on the theory that the back portion of Meridew’s handle “houses necessary functional components,” and that incorporation of Fucci or Schmieding’s teachings would require that Meridew’s handle “have to be significantly redesigned.” Prelim. Resp. 41–45.

2 Although characterized as relating to “Ground 2,” these portions of the Petition and Dr. Jordan’s testimony here, and below, are referenced at page 46 of the Petition in connection with the ground identified as “Ground 4.”
We have considered Patent Owner’s preliminary arguments pertaining to claims 11 and 12. *Id.* Patent Owner does not dispute that the features of claims 11 and 12 are known in the art of suture anchors and that the proposed combination discloses all of the limitations of claims 11 and 12. At this initial stage, we are satisfied that Petitioner has made a sufficient showing to warrant institution of *inter partes* review of claims 11 and 12 based on Meridew and either Fucci or Schmieding. In particular, to the extent that some redesign of Meridew may be necessary in light of Fucci or Schmieding’s teachings, it is not apparent that such redesign would be beyond the ordinary skill of a skilled artisan. Based on the teachings of Fucci or Schmieding, and in crediting, at this time, the testimony of Petitioner’s declarant, Dr. Steve E. Jordan (Ex. 1027), we are persuaded that Petitioner has shown a reasonable likelihood of success in its challenge to claims 11 and 12 based on Meridew and either Fucci or Schmieding.

3. *Claim 16*

Claim 16 depends from claim 13 and adds a step of “piercing soft tissue with the bone-piercing tip.” Petitioner relies on Anspach in combination with Meridew to account for that additional step. Pet. 46 (incorporating arguments made at Pet. 40–41). Anspach discloses a “Soft Tissue Fastener Device.” Ex. 1021, Title. Anspach describes the use of its device during a surgical procedure and during such procedure, “soft tissue is pierced.” *Id.* at 2:3–6. Petitioner urges that Meridew and Anspach are generally similar devices and that, in light of the teachings of Anspach, a person of ordinary skill in the art would have appreciated that a suture anchor having a pointed, bone-piercing tip, such as that suggested by Meridew as modified by either Goble or Pietrzak, may be employed to
pierce soft tissue with predictable results. Pet. 40–41 (citing Ex. 1027 ¶¶ 138–140)).

Patent Owner disagrees with Petitioner. In particular, Patent Owner is of the view that Anspach and Meride are not similar devices, and that Anspach’s approach to piercing tissue would not apply to Meridew’s device. Prelim. Resp. 45–46. Patent Owner also contends that any combination of Anspach and Meridew would necessitate a “substantial redesign” of Meridew that undercuts Petitioner’s obviousness position. Id. at 46.

It is apparent from the record at hand that surgical techniques involving the fastening or anchoring of structures to bone are understood to employ bone-piercing tips and piercing of soft tissue. Patent Owner also does not explain why even if the implementation of Anspach’s teachings to Meridew’s assembly requires some redesign, that any such redesign extends beyond an ordinary artisan’s skill. Based on the current record, we are persuaded to institute inter partes review of claim 16 based on Petitioner’s Ground 4.

D. Grounds 5–7 – Obviousness Based on Burkhart

Petitioner proposes alternative grounds to claims 1–18 based on Burkhart. Pet. 19. Burkhart discloses a “Graft Fixation Using a Plug Against Suture.” Ex. 1006, Title. Although, like Meridew, Burkhart describes a tool and practice of securing soft tissue to bone, Burkhart takes a different approach than Meridew. For instance, Burkhart’s tool employs various configurations, including an eyelet, through which suture is threaded. See, e.g., Ex. 1006, Figs. 4, 10, 11 (elements 50, 55, 250, 255).

It is manifest that those structures are not bone-piercing tips and do not have pointed tips. Burkhart, unlike Meridew, provides no contemplation
of incorporating any such pointed tip or impacting tip in lieu of those structures. Although Petitioner contends that combining Burkhart’s teachings with either Pietrzak or Goble accounts for the required tip, we are skeptical that Petitioner’s proposed “result” of that combination is one that a skilled artisan reasonably would have derived based on the teachings of the relevant prior art. See Pet. 51.

As Patent Owner notes, Burkhart contemplates manipulating the sutures after inserting the implant device into a pilot hole, with the sutures residing on the outside of the device. Prelim. Resp. 52–53; see also Ex. 1006, ¶¶ 27–28, Figs. 4–5. Only after tensioning the suture does Burkhart contemplate impacting the interference device into the pilot hole, locking the implant with the sutures in place. Ex. 1006 ¶ 28, Fig. 7. Patent Owner correctly points out that the interference device of Burkhart “becomes superfluous” by modifying Burkhart with the pointed tips of Pietrzak or Goble, given that the pointed tips form an interference fit with surrounding bone upon insertion. Prelim. Resp. 54–56. Petitioner also fails to explain how the procedure of Burkhart, initially requiring insertion of the anchor without an interference fit to allow manipulation of the sutures, can be performed using pointed tips that form an interference fit upon insertion. Based on these deficiencies, we are not persuaded that Petitioner has established a reasonable likelihood of success in prevailing on Grounds 5–7.

Further, with respect to claims 13–18 and as discussed above, those claims require an “inner tube,” and that such a tube in the context of the ’663 patent must be hollow. We agree with Patent Owner that it is not apparent that shank 19 of Burkhart, on which Petitioner relies to account for such a tube, is hollow. See Prelim. Resp. 34–35. In that respect, there is no
suggestion in Burkhart that any component is intended to extend through that shank. For instance, all of Burkhart’s disclosed embodiments show suture that is external to Burkhart’s suturing tool.

Petitioner relies on other prior art in combination with Burkhart, but does not rely on that art to address the deficiencies we note above in Burkhart. Based on the foregoing, we do not institute *inter partes* review on any of Petitioner’s Grounds 5–7 based on Burkhart.

III. CONCLUSION

Having evaluated the Petition, its underlying supporting evidence, and Patent Owner’s Preliminary Response, we determine that Petitioner has shown a reasonable likelihood of success in its challenge to claims 1–18 based on Meridew, Pietrzak or Goble, Fucci or Schmieding, and Anspach, as set forth in the Petition and discussed above.

At this stage of the proceeding, the Board has not made a final determination as to the construction of any claim term or the patentability of claims 1–18 of the ’663 patent.

IV. ORDER

It is ORDERED that, pursuant to 35 U.S.C. § 314(a), an *inter partes* review is hereby instituted to determine whether claims 1–18 are unpatentable based on Meridew, Pietrzak or Goble, Fucci or Schmieding, and Anspach;

FURTHER ORDERED that *inter partes* review is not instituted in this proceeding on any other grounds; and

FURTHER ORDERED that pursuant to 35 U.S.C. § 314(c) and 37 C.F.R. § 42.4, notice is hereby given of the institution of a trial. The trial will commence on the entry date of this decision.