

# United States Court of Appeals for the Federal Circuit

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**BECTON, DICKINSON AND COMPANY,**  
*Appellant*

**v.**

**BAXTER CORPORATION ENGLEWOOD,**  
*Appellee*

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2020-1937

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Appeal from the United States Patent and Trademark  
Office, Patent Trial and Appeal Board in No. IPR2019-  
00119.

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Decided: May 28, 2021

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THOMAS SAUNDERS, Wilmer Cutler Pickering Hale and  
Dorr LLP, Washington, DC, argued for appellant. Also rep-  
resented by OMAR KHAN, New York, NY; KATHERINE P.  
KIECKHAFFER, Boston, MA.

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IL, argued for appellee. Also represented by KATHERINE  
ALLOR, GEORGE C. SUMMERFIELD, JR.; MICHAEL IRA COHEN,  
Baxter International Inc., Deerfield, IL.

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Before PROST\*, CLEVINGER, and DYK, *Circuit Judges*.

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\* Sharon Prost vacated the position of Chief Judge on  
May 21, 2021.

DYK, *Circuit Judge*.

Becton, Dickinson and Company (“Becton”) appeals a decision from the Patent Trial and Appeal Board (“Board”), determining that certain claims of U.S. Patent No. 8,554,579 (“the ’579 patent”) were not invalid as obvious. We reverse.

#### BACKGROUND

Baxter Corporation Englewood (“Baxter”) is the owner of the ’579 patent, which is directed to “[s]ystems for preparing patient-specific doses and a method for telepharmacy in which data captured while following [a protocol associated with each received drug order and specifying a set of steps to fill the drug order] are provided to a remote site for review and approval by a pharmacist.” ’579 patent, Abstract.

Becton petitioned for inter partes review of claims 1–13 and 22 of the ’579 patent. Claims 2–7 and 22 depend, directly or indirectly, from independent claim 1. Claims 9–13 depend, directly or indirectly, from independent claim 8. The parties agree that claims 1 and 8 of the ’579 patent are illustrative.

There are two contested limitations on appeal. The first is the “verification” limitation, which appears in claim 8 but not claim 1. The second is the “highlighting” limitation, which appears in both claims 1 and 8. The relevant portion of claim 8, containing both limitations, states:

8. A system for preparing and managing patient-specific dose orders that have been entered into a first system, comprising:

...

- a dose preparation station for preparing a plurality of doses based on received dose orders, the dose preparation station being in bi-directional communication with the order processing server and

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having an interface for providing an operator with a protocol associated with each received drug order and specifying a set of drug preparation steps to fill the drug order, the dose preparation station including an interactive screen that includes prompts that can be highlighted by an operator to receive additional information relative to one particular step and includes areas for entering an input;

. . . and wherein each of the steps must be verified as being properly completed before the operator can continue with the other steps of drug preparation process, the captured image displaying a result of a discrete isolated event performed in accordance with one drug preparation step, wherein verifying the steps includes reviewing all of the discrete images in the data record . . . .

*Id.* col. 32 l. 52–col. 33 l. 30 (highlighting and verification limitations emphasized). Claims 1 and 8 are set forth in full in an Attachment to this opinion.

In asserting that the challenged claims were invalid, Becton relied primarily on three prior art references: U.S. Patent No. 8,374,887 (“Alexander”), U.S. Patent No. 6,581,798 (“Liff”), and U.S. Patent Publication No. 2005/0080651 (“Morrison”).

The Board found that Becton had established that one of ordinary skill in the art would have been motivated to combine Alexander and Liff, as well as Alexander, Liff, and Morrison. The Board also determined that Baxter’s “evidence of secondary considerations [was] weak.” J.A. 34.

However, the Board determined that Alexander did not teach or render obvious the verification limitation and that combinations of Alexander, Liff, and Morrison did not teach or render obvious the highlighting limitation. The

Board concluded that, as a result, none of the challenged claims (1–13, 22) was shown to be unpatentable.<sup>1</sup>

Becton appeals. We have jurisdiction under 28 U.S.C. § 1295(a)(4)(A) and 35 U.S.C. § 141(c).

#### DISCUSSION

A patent may not be obtained . . . if the differences between the subject matter sought to be patented and the prior art are such that the subject matter as a whole would have been obvious at the time the invention was made to a person having ordinary skill in the art to which said subject matter pertains.

35 U.S.C. § 103(a).<sup>2</sup> In reviewing the Board’s determination on the question of obviousness, “[w]e review the Board’s legal conclusions de novo and its factual findings for substantial evidence.” *MCM Portfolio LLC v. Hewlett-Packard Co.*, 812 F.3d 1284, 1293 (Fed. Cir. 2015).

#### I

We first address the verification limitation, “wherein each of the steps must be verified as being properly completed before the operator can continue with the other steps of drug preparation process.” ’579 patent, col. 33 ll. 18–21. The Board construed the limitation under the broadest

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<sup>1</sup> The Board also found that the challenged claims were not shown to be unpatentable on a separate third ground asserted by Becton, which Becton does not appeal.

<sup>2</sup> Congress amended § 103 when it enacted the Leahy-Smith America Invents Act (“AIA”). Pub. L. No. 112-29, § 3(c), 125 Stat. 284, 287 (2011). However, because the challenged claims of the ’579 patent have an effective filing date before March 16, 2013, the pre-AIA version of § 103 applies. *See id.* § 3(n)(1), 125 Stat. at 293.

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reasonable interpretation standard.<sup>3</sup> Under the Board's construction, which neither party appeals, the verification limitation requires that "the system will not allow the operator to proceed to the next step until the prior step has been verified." J.A. 17. The Board further determined that the plain language of the verification limitation does not require "automatic system function" to "trigger verification." *Id.*<sup>4</sup>

The Board determined that Alexander does not teach or render obvious the verification limitation. Specifically, the Board found persuasive Baxter's argument that Alexander "only discusses that 'a remote pharmacist may verify each step'; not that the remote pharmacist must verify each and every step before the operator is allowed to proceed." *See id.* at 36–37 (quoting *id.* at 5284) (citation and emphasis omitted). We conclude that the Board's determination is not supported by substantial evidence.

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<sup>3</sup> Because the filing date of the petition for inter partes review, October 29, 2018, was before November 13, 2018, the broadest reasonable interpretation standard applies. *See* Changes to the Claim Construction Standard for Interpreting Claims in Trial Proceedings Before the Patent Trial and Appeal Board, 83 Fed. Reg. 51,340 (Oct. 11, 2018) (codified at 37 C.F.R. § 42.100); *see also Personalized Media Commc'ns, LLC v. Apple Inc.*, 952 F.3d 1336, 1340 n.2 (Fed. Cir. 2020).

<sup>4</sup> Baxter refers to the verification limitation as a "hard stop," Appellee's Br. 25, as opposed to a "soft stop," which, according to Baxter, "provides information to the clinician about a potential drug safety or efficacy problem and may offer alternative suggestions for the clinician to consider." J.A. 5283 (citation omitted). Under the Board's construction, "'hard stop' is merely a short hand for [the verification limitation] and does not impute any additional meaning to the claim term." *Id.* at 15 n.8.

The passage from Alexander on which the Board relied states:

[I]n some embodiments, a remote pharmacist may supervise pharmacy work as it is being performed. For example, in one embodiment, a remote pharmacist may verify each step as it is performed and may provide an indication to a non-pharmacist performing the pharmacy that the step was performed correctly. In such an example, the remote pharmacist may provide verification feedback via the same collaboration software, or via another method, such as by telephone.

Alexander, col. 9 ll. 47–54 (emphasis added).

In the context of Alexander, “may” does not mean “occasionally,” but rather that one “may” choose to systematically check each step. This is quite clear from the context of Alexander.

Alexander is directed to “[a] system and method for remotely supervising and verifying pharmacy functions performed by a non-pharmacist at an institutional pharmacy.” *Id.* Abstract. Alexander discloses that “software may be installed at both an institutional pharmacy site and at a remote pharmacist site allowing a pharmacist to view in real-time, or near real-time, images of the pharmacy work being performed.” *Id.* col. 9 ll. 31–34. “Captured images and corresponding documentation may be transmitted from institutional pharmacy to the remotely located pharmacist, either directly or via a web site accessible to both.” *Id.* Abstract. The purpose is to allow the pharmacist to “authorize” the work. *See id.*

In this process, the Alexander specification provides that the non-pharmacist is not authorized to proceed absent verification by the pharmacist. The abstract states that “[r]eceiving the pharmacist’s verification may authorize the non-pharmacist to further process the work.” *Id.*

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Likewise, column two of the specification provides that “a pharmacist supervises and verifies the work, and subsequently authorizes non-pharmacist personnel to further process the work.” *Id.* col. 2 ll. 51–53. Plainly, Alexander discloses systematic step-by-step review and authorization by the pharmacist.

It is also clear that, without the pharmacist’s verification of “each step as it is performed,” *id.* col. 9 ll. 49–50, the non-pharmacist is not “authorize[d]” to “further process the work,” *id.* col. 2 ll. 52–53; *see also id.* Abstract. There is no significant difference between that teaching of Alexander and the ’579 patent’s verification requirement, which the Board construed as requiring that “the system will not allow the operator to proceed to the next step until the prior step has been verified.” J.A. 17.

Baxter attempts to sustain the Board on two grounds not adopted by the Board. Baxter first contends that “Alexander does not disclose a system that would stop the operator from proceeding if a prior step was unverified, and that such an improperly prepared dose ‘could go out to the patient and cause harm,’” quoting from the deposition testimony of Becton’s expert witness, Dr. Marc Young. Appellee’s Br. 29 (quoting J.A. 3376). Requiring authorization before proceeding necessarily stops the work if the authorization is not forthcoming. The remote operator cannot further process the work without authorization. Counsel for Baxter conceded at oral argument that, in Alexander’s system, a non-pharmacist who, without authorization to proceed, did not stop processing work would likely be disciplined. Oral Arg. 25:55–26:40, [http://oralarguments.cafc.uscourts.gov/default.aspx?fl=20-1937\\_04082021.mp3](http://oralarguments.cafc.uscourts.gov/default.aspx?fl=20-1937_04082021.mp3).

Baxter also contends that Alexander’s disclosure is insufficient because the verification limitation requires a “mechanical . . . prohibition” on continuing the work absent verification. *Id.* at 23:37–24:01. The Board’s construction

requires that “the system will not allow the operator to proceed to the next step until the prior step has been verified,” J.A. 17 (emphasis added), and Baxter contends that the use of the word “system” implies a mechanical stop.

Nothing in the construction requires a mechanical stop as opposed to requiring authorization from a pharmacist to continue. Nor does the specification of the ’579 patent indicate that the “stop” cannot be in the form of an instruction from a pharmacist. See, e.g., ’579 patent, col. 15 ll. 39–45 (“If during any step, a verification error arises and there is a question as to whether the step was properly performed, the dose order processing is prevented from continuing to the next step until the step is verified as being properly performed or until the dose order is flagged as being not completed due to an error.” (emphasis added)); *id.* col. 18 ll. 25–27, 56–58 (similar).

Finally, Baxter presents the ’579 patent as an improvement to the “pull-back’ method” of pharmacist verification in sterile compounding. See Appellee’s Br. 4. “Often a pharmacy technician (a non-pharmacist) performs the sterile compounding under a pharmacist’s supervision, with the pharmacist responsible for final verification of the prepared dose.” *Id.*

According to the “pull-back” method, after combining ingredients using one or more syringes, the technician pulls each syringe back to the position it was in when it was full of the added component, but since the ingredients have already been combined, the syringe would be filled with air. The pulled-back syringe(s) along with other dose preparation materials would be in a basket, which the pharmacist would then use to reconstruct the process and verify that the technician had properly prepared the dose.

*Id.* (citations omitted).

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According to Baxter, the pull-back method “left a lot of room for error in dose preparation of sterile compounds.” *Id.* Baxter contends that the ’579 patent improved the prior art by disclosing a “a system for dose preparation flow and verification of preparation steps, whereby the system prevents the dose preparer from proceeding to the next preparation step if the previous step has not been verified.” *Id.* at 5.

The embodiment of Alexander, in which “a remote pharmacist may verify each step as it is performed and may provide an indication to a non-pharmacist performing the pharmacy that the step was performed correctly,” Alexander, col. 9 ll. 49–52, is not the pull-back method (or an electronic version thereof). The Board’s suggestion to the contrary, *see* J.A. 36–37, is not supported by substantial evidence.

We conclude that the Board’s determination that Alexander does not teach the verification limitation is not supported by substantial evidence.

## II

We next address the highlighting limitation, which requires “an interactive screen that includes prompts that can be highlighted by an operator to receive additional information relative to one particular step.” ’579 patent, col. 31 ll. 55–57, col. 33 ll. 4–6. Figure 10 of the ’579 patent, “an exemplary display of a product preparation screen and procedure,” *id.* col. 5 ll. 47–48, shows

an interactive screen in that the user can simply highlight different areas either to receive more information or to enter information. For example, there is a Detail button 622 near the drug identification and if additional information is needed concerning this particular drug order, the user can simply highlight this particular button (as by “clicking” the box).

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*Id.* col. 15 l. 58–col. 16 l. 3.

The screenshot displays the IntelliFlow software interface. At the top, it shows 'IntelliFlow LAF Hood1' and user information: 'Use: demo', 'Login Time: 8/7/2008 5:10 PM', and buttons for 'Exit' and 'ForHealth'. The interface is organized into a sidebar on the left, a main central workspace, and a task/action panel on the right.

**Left Sidebar:** Contains buttons for 'ALL [245]', 'STAT / Dose Spool [99]', 'First Dose [10]', 'BQ [15]', 'WAIT / HOLD [3]', 'Make Now [11]', 'Unknown Orders [8]', 'PHT Doses [245]', 'Padman Doses [2]', 'High-Risk Doses [34]', and 'Override [251]'.

**Central Workspace:**

- Top status bar: 'Cefazolin 100mg in 1mL', 'Pending(1)', 'Completed(0)', 'Total(1)'. Buttons: 'Cancel Pending Doses', 'Task [3]'.
- 'Selected Products Information' section for 'Cefazolin 1g (Apoth)'. It shows '100 mg / 100mg/mL = 1 mL', 'Recon: 9.6 mL', 'Current: 10mL', 'Qty: 10'. Buttons: 'Detail', 'Cancel'.
- A message box states: 'The product list is complete'.
- 'Activity log' section:
  - 1 Started Dose preparation for DoseIDs: 16 Dose Description: Cefazolin
  - 2 Selected a new Product 'Cefazolin 1g (Apoth)' (555). Volume=10.00

**Right Task/Action Panel:**

- 'Cefazolin 1g (Apoth)' section: 'Prepare vial with reconstitution', 'Print and apply Product label' (with 'Print Label' button), 'Scan product to verify'.
- 'Enter Product Lot Number. 620' (with 'Keyboard' button).
- 'Enter Product Expiration Date. 8/7/2008' (with dropdown arrow).
- 'Scan diluent for reconstitution' (with 'Done' button).
- 'Acquire diluent for reconstitution'.
- 'Capture image of diluent'.
- 'Capture image of completed vial'.
- 'Scan a Dose to begin preparation'.
- Bottom buttons: 'Settings', 'Run Reports', 'Change Password', 'Scan Barcode', 'Support', 'Help'.

**FIG. 10**

*Id.* fig. 10.

The Board determined “that in implementing an electronic system for preparing medications, one of ordinary skill in the art would have considered it obvious to provide a set of drug preparation steps on a computer.” J.A. 41. Given that determination, the only missing element of this limitation is the ability to highlight prompts to receive more information concerning drug preparation steps.

The Liff reference teaches highlighting in the pharmacy context. *See id.* at 1496–97 (declaration of Dr. Marc Young in support of petition for inter partes review). Liff is directed to “[a]n automated drug dispensing system [that] includes a cabinet adapted to store a variety of

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prepackaged pharmaceuticals in a plurality of bins for filling patient prescriptions.” Liff, Abstract. Liff teaches a user interface for a workstation, *see id.* col. 4 ll. 5–9, figs. 14A–14V, which provides the operator with options, such as “entering a new prescription” and “refilling a prescription.” *Id.* col. 17 ll. 28–31 (referring to fig. 14A).

As Dr. Young testified in his declaration, Liff “teaches that the user can highlight various inputs and information displayed on the screen, as illustrated in Figure 14F.” J.A. 1496–97. More specifically, the Board found that Liff taught “highlight[ing] patient characteristics when dispensing a prepackaged medication,” *id.* at 43, and Baxter does not contend that this aspect of the Board’s decision was in error.

Figure 14F of Liff is below:

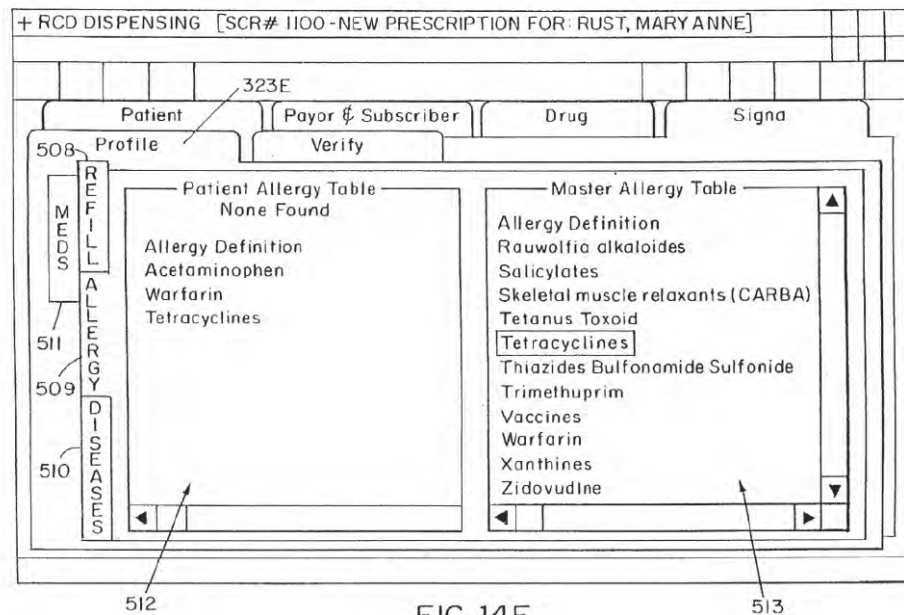


FIG. 14F

Liff, fig. 14F.

Becton does not argue that Liff “directly discloses highlighting to receive additional language about a drug

preparation step.” Appellant’s Br. 4. Becton instead argues that “Liff discloses basic computer functionality—i.e., using prompts that can be highlighted by the operator to receive additional information—that would render the highlighting limitation obvious when applied in combination with other references,” primarily Alexander. *Id.* at 4 (emphasis omitted); *see also id.* at 24–25.

Becton relies on the following testimony of Dr. Young:

A person of ordinary skill in the art would have understood that additional information could be displayed on the tabs taught by Liff and that additional tabs, with additional information, could also be displayed in the user interface, depending on the design needs and expected use of the software. For example, a person of ordinary skill in the art would have found it obvious to include in the user interface taught by Liff a tab for the prescription order and information regarding the prescription order that the operator was fulfilling. Such information could have included the text of the order itself, information relating to who or how the order should be prepared, or where the order should be dispensed.

J.A. 1497. The testimony of Baxter’s expert, Dr. Jeffrey Brittain, was not to the contrary.

The Board found that “this present[ed] a close case.” *Id.* at 43. As noted above, the Board agreed that, in light of Alexander and Liff, “one of ordinary skill in the art would have considered it obvious to provide a set of drug preparation steps on a computer.” *Id.* at 41. Nevertheless, the Board determined that

Dr. Young fail[ed] to explain why Liff’s teaching to highlight patient characteristics when dispensing a prepackaged medication would lead one of ordinary skill to highlight prompts in a drug

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formulation context to receive additional information relative to one particular step in that process, or even what additional information might be relevant.

*Id.* at 43. The Board found that Becton's arguments with respect to Morrison did not address the deficiency in its position based on Alexander and Liff.

The Board's determination is not supported by substantial evidence. That Liff's teaching was "to highlight patient characteristics when dispensing a prepackaged medication," *id.*, does not suggest that a person of ordinary skill would not have used highlighting (accomplished in the '579 patent by "clicking" [a] box" labeled "[d]etail," '579 patent, col. 15 l. 64–col. 16 l. 3) with respect to other information in the pharmacy field. Dr. Young, without contradiction, testified to the opposite, stating that "a person of ordinary skill in the art would have found it obvious to include in the user interface taught by Liff a tab for the prescription order and information regarding the prescription order that the operator was fulfilling." J.A. 1497. As the Supreme Court has made clear, "[t]he combination of familiar elements according to known methods is likely to be obvious when it does no more than yield predictable results." *KSR Int'l Co. v. Teleflex Inc.*, 550 U.S. 398, 416 (2007).

The Board erred in looking to Liff as the only source a person of ordinary skill would consider for what "additional information might be relevant." J.A. 43. "A person of ordinary skill is also a person of ordinary creativity, not an automaton." *KSR*, 550 U.S. at 421. Dr. Young testified that "[a] person of ordinary skill in the art would have understood that additional information could be displayed on the tabs taught by Liff" and that "such information could have included the text of the order itself, information relating to who or how the order should be prepared, or where the order should be dispensed." J.A. 1497. Dr. Young further

testified that “[a] medication dose order for compounding a pharmaceutical would have been accompanied by directions for how the dose should be prepared, including step-by-step directions for preparing the dose.” *Id.* Baxter points to no contrary testimony.

We conclude that the highlighting limitation would have been obvious to one of ordinary skill in the art in view of Alexander and Liff. The Board’s determination that the highlighting limitation is not obvious over Alexander and Liff is not supported by substantial evidence. We need not reach Becton’s arguments regarding Morrison.

### III

As an alternative ground to affirm the Board’s determination of non-obviousness, Baxter argues that the Board erred in determining that Alexander is prior art under 35 U.S.C. § 102(e)(2) (pre-AIA).<sup>5</sup> This section provides that “[a] person shall be entitled to a patent unless . . . the invention was described in . . . a patent granted on an application for patent by another filed in the United States before the invention by the applicant for patent.” 35 U.S.C. § 102(e)(2).<sup>6</sup> It is undisputed that the filing date of the

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<sup>5</sup> Congress amended § 102 when it enacted the AIA. Pub. L. No. 112-29, § 3(b)(1), 125 Stat. at 285–87. However, because the application that led to the ’579 patent was filed before March 16, 2013, the pre-AIA version of § 102 applies. *See id.* § 3(n)(1), 125 Stat. at 293.

<sup>6</sup> The Board noted that “Alexander is not prior art under 35 U.S.C § 102(e)(1).” J.A. 23. That section provides that “an application for patent, published under [35 U.S.C. § 122(b)], by another filed in the United States before the invention by the applicant for patent,” is prior art. 35 U.S.C § 102(e)(1) (pre-AIA). The Board found that “the applicant [for Alexander] expressly requested that the application that matured into Alexander ‘not be published under

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application for Alexander is February 11, 2005, which is before the earliest filing date of the application for the '579 patent, October 13, 2008; that the Alexander claims were granted; and that the application for Alexander was filed by another.

Baxter contends that Alexander nonetheless is not prior art because all claims in Alexander (granted on February 12, 2013) were cancelled on February 15, 2018, following inter partes review. Baxter argues that “because the Alexander ‘grant’ had been revoked, it can no longer qualify as a patent ‘granted’ as required for prior art status under Section 102(e)(2).” Appellee’s Br. 35.<sup>7</sup>

The text of the statute requires only that the patent be “granted,” meaning the “grant[]” has occurred. 35 U.S.C. § 102(e)(2) (pre-AIA). The statute does not require that the patent be currently valid.

#### IV

Finally, we address “secondary considerations” of non-obviousness. *See Graham v. John Deere Co. of Kansas City*, 383 U.S. 1, 17 (1966). The Board found that Baxter’s evidence of secondary considerations was “weak.” J.A. 33–

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35 U.S.C 122(b)’ and was, therefore, never published under that section.” J.A. 23 (citation omitted). On appeal, Becton does not argue that Alexander is prior art under grounds other than 35 U.S.C. § 102(e)(2).

<sup>7</sup> Baxter also argues that “even assuming that Alexander’s prior art status is evaluated at the time of patent filing, a person of skill in the art would not have considered Alexander to be prior art as of October 2008 [the ‘579 patent’s priority date], or even known of Alexander, because Alexander was not made public until issuance on February 12, 2013.” Appellee’s Br. 36. The Supreme Court’s decision in *Hazeltine Research, Inc. v. Brenner*, 382 U.S. 252, 254–56 (1965), forecloses this argument.

34.<sup>8</sup> Baxter does not argue that the Board’s determination in this respect was in error. “[W]eak evidence of secondary considerations . . . simply cannot overcome the strong showing of obviousness.” *ZUP, LLC v. Nash Mfg., Inc.*, 896 F.3d 1365, 1375 (Fed. Cir. 2018). Baxter does not meaningfully argue that the weak showing of secondary considerations here could overcome the showing of obviousness based on the prior art.

#### CONCLUSION

The Board’s determination that the verification and highlighting limitations are not obvious is not supported by substantial evidence. We reverse.

#### REVERSED

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<sup>8</sup> Baxter presented evidence of secondary considerations focusing primarily on the verification limitation.

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ATTACHMENT

1. A method for performing telepharmacy comprising the steps of:

receiving and processing a dose order;

preparing a dose at a medication preparation station based on the dose order including following a recipe, wherein the dose is a reconstituted drug and the recipe having one or more drug preparation steps including using a diluent for reconstitution;

displaying the recipe on an interactive screen that includes prompts that can be highlighted by an operator to receive additional information relative to one particular step and includes areas for entering an input;

capturing one or more images of a plurality of the drug preparation steps, each of the images being captured at, corresponding to, and confirming a performance of one discrete drug preparation step of the recipe, one captured image displaying a result of a discrete isolated event performed in accordance with one drug preparation step, the drug preparation steps including at least one step that is an intermediate step involving the diluent that shows the dose prior to completing the dose preparation and obtaining a completed dose that is in a state that is suitable for delivery to a patient, wherein one input comprises an input that is prompted by the performance of the drug preparation steps;

storing each image associated with the drug preparation steps of the recipe that has been collected together in a data record of a database, thereby allowing the captured image to be later retrieved for inspection;

accessing the data record including the images from a remote site using a portal in communication with the database;

inspecting the data record through the portal;

reviewing the images in the data record in order to verify that each of the captured drug preparation steps was properly completed; and

approving release of the dose to the patient if the reviewing step confirms that each of the captured drug preparation steps was properly completed.

'579 patent, col. 31 l. 47–col. 32 l. 18.

8. A system for preparing and managing patient-specific dose orders that have been entered into a first system, comprising:

an order processing server executing software on a processor thereof and connected by a network to the first system and configured to receive the patient-specific dose orders from the first system, the order processing server including a database configured to store the dose orders and images that relate to the dose orders, the order processing server being configured to generate a dose order queue listing all dose orders received by the order processing server;

a dose preparation station for preparing a plurality of doses based on received dose orders, the dose preparation station being in bi-directional communication with the order processing server and having an interface for providing an operator with a protocol associated with each received drug order and specifying a set of drug preparation steps to fill the drug order, the dose preparation station including an interactive screen that includes prompts that can be highlighted by an operator to receive additional information relative to one particular step and includes areas for entering an input;

the dose preparation station being configured to present the protocol and having one or more data input devices to capture images of a plurality of the set of drug preparation steps that are part of the protocol and are followed to fill the drug order, wherein each image

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associated with the drug preparation steps of the protocol is stored together in a data record of the database, wherein at least one captured image is captured at, corresponds to, and confirms a performance of one discrete drug preparation step in which the dose is not completely prepared and ready for delivery to the patient and wherein each of the steps must be verified as being properly completed before the operator can continue with the other steps of drug preparation process, the captured image displaying a result of a discrete isolated event performed in accordance with one drug preparation step, wherein verifying the steps includes reviewing all of the discrete images in the data record; and

a display communicatively coupled to the order processing server and positionable independently of the dose preparation station, the display outputting the dose order queue and metrics concerning activity at the dose preparation station.

*Id.* col. 32 l. 52–col. 33 l. 30.