

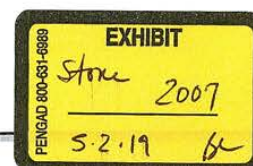
**Date:** April 16, 2019

**Case:** Certain Strontium-Rubidium Radioisotope Infusion Systems, and  
Components Thereof Including Generators

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UNITED STATES OF AMERICA  
BEFORE THE  
INTERNATIONAL TRADE COMMISSION

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IN THE MATTER OF: : Investigation Number  
CERTAIN STRONTIUM-RUBIDIUM : 337-TA-1110  
RADIOISOTOPE INFUSION SYSTEMS AND :  
COMPONENTS THEREOF INCLUDING :  
GENERATORS :

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HEARING - VOLUME IV

April 16, 2019  
Courtroom C  
U.S. International Trade  
Commission  
500 E Street, S.W.  
Washington, D.C.

The Hearing commenced, pursuant to notice of the Judge, at  
9:03 a.m., before the Honorable CLARK S. CHENEY,  
Administrative Law Judge for the United States  
International Trade Commission.

1 reporter. But I would like to move -- jointly move the  
2 admission of the exhibits that we are sending to the court  
3 reporter that were discussed -- discussed on the record  
4 from April 15th, 2019.

5 JUDGE CHENEY: I presume counsel for complainant  
6 and staff understand the list of exhibits that Mr. Hails  
7 has represented and is there any objection to the admission  
8 of those exhibits?

9 MR. DAVIS: No, Your Honor.

10 MR. KOO: No, Your Honor.

11 JUDGE CHENEY: The exhibits identified by  
12 Mr. Hails will be admitted to the record.

13 MR. HOFFMAN: Your Honor, Mr. Hails is sitting  
14 in front of me. I'm Mr. Hoffman.

15 JUDGE CHENEY: I'm sorry. Mr. Hoffman.

16 Any other housekeeping matters before we resume  
17 with Dr. Stone? Okay. Hearing none, we'll ask Dr. Stone  
18 to return to the stand. As you're coming to the stand,  
19 Dr. Stone, I'll remind you you're still under the same  
20 obligation to tell the truth under penalty of perjury.  
21 Whereupon,

22 DR. ROBERT STONE,  
23 was called as a witness by counsel for Respondents, and  
24 having been previously duly sworn, was examined and  
25 testified as follows:

1 THE WITNESS: Thank you, Your Honor.

2 JUDGE CHENEY: Mr. Hails, who has spent all  
3 evening talking slowly and clearly in the mirror so that  
4 today we'll just have a nice relaxed transcript.

5 MR. HAILS: We'll see. Yes, sir.

6 DIRECT EXAMINATION

7 BY MR. HAILS:

8 Q Dr. Stone, before we broke we were talking about  
9 shielding design. Do you recall that?

10 A Yes.

11 Q Let's pull up RX-357, page 10, if you can. Do  
12 you recognize this, Dr. Stone?

13 A Yes, I do.

14 Q Do you see any shielding compartments  
15 illustrated in this photograph?

16 A Yes, I do.

17 Q First of all, what does this picture illustrate?

18 A This is a picture of the interior of the cabinet  
19 of the CardioGen 82 or the Model 510 as it's called.

20 Q The Model 510. Okay. Do you see any shielding  
21 compartments in this design?

22 A I do.

23 Q Would you point them out for us?

24 A In the back is a shielding compartment. I  
25 believe that is for the waste container and here we have

1 the shielding compartment for the generator.

2 Q And Ricky, can we blow up that sticker in the  
3 middle center on that first.

4 All right. Do you see the sticker for the  
5 generator shield?

6 A I do.

7 Q Okay. You can back out of that. All right. So  
8 which way do the openings face on the shielding?

9 A They face vertically upward.

10 Q Let's switch over to RX-103 at page 4. Can we  
11 rotate that so the letters all line up. Thank you.

12 Do you recognize this as the Tate application  
13 that you've been discussing?

14 A I do.

15 Q Okay. Do you see any shielding containers --  
16 sorry, shielding compartments in this -- illustrated in  
17 this figure?

18 A Yes, I do.

19 Q Can you point them out for us.

20 A Yes. Here we have, I believe it's a shielding  
21 compartment for the source. Here we have the shielding  
22 compartment for the onboard dose calibrator, and I believe  
23 this is a shielding compartment for the waste bottle.

24 Q And just for the record, Dr. Stone pointed to  
25 element 111 in this diagram, 121 in this diagram, and 127

1 in this diagram.

2 Dr. Stone, do you recognize this as a top view  
3 of the Tate cart?

4 A Yes, I do.

5 Q And which way do the openings of these shielding  
6 compartments face?

7 A They face vertically upward.

8 Q Can you explain why do people build shielding  
9 compartments with openings that face vertically upward?

10 A Yes. As one is approaching a device, the  
11 shielding can be complete and enclosed without access to  
12 the device if the shielding compartments face vertically  
13 upward. So the shielding is providing adequate protection  
14 for the thorax of a person unless -- and the only way they  
15 could get exposed to the thorax is if they leaned over that  
16 source of radioactive material. So they don't have to  
17 expose themselves to radiation as they approach the devices  
18 because it's fully enclosed with openings facing vertically  
19 upward and radiation only going upward even if the lid were  
20 open.

21 Q Can we return to the PowerPoint.

22 Okay. Before we broke, we were talking about  
23 this element shown here on slide 128 at the bottom, the  
24 first door element. Do you see that?

25 A I do.

1 Q Okay. And this claim element says that the  
2 first door is configured to provide access to the shielding  
3 compartment and to close over that first opening.

4 Do you see that?

5 A I do.

6 Q Did you see any teachings of this subject matter  
7 in the prior art?

8 A Yes. In Klein, the generator was placed in the  
9 cart and surrounded by lead rings to provide maximum  
10 radiation shielding. The opening faced vertically upward  
11 in that configuration. Similarly in Tate, the radioactive  
12 source was inserted into a well and had an opening which  
13 faced vertically upward.

14 Q Can you show us the door in the Tate design?

15 A In Tate, we have a door which closes over the  
16 source of radioisotope.

17 Q For the record, we're looking at figure 4A from  
18 Tate. And did you point to element 684 in that drawing?

19 A I did.

20 Q Are doors conventional structures for shielding  
21 design in the mid 2000?

22 A Doors are conventional structures for shielding  
23 in my entire history of looking at radioisotopes and  
24 radioactive sources. They are very common.

25 Q If someone applied a door to a shielding

1 structure, for example, in the Klein design, would it be  
2 accessible via the opening through the exterior shell of  
3 the compartment?

4 A Yes, it would.

5 Q Move forward, please.

6 JUDGE CHENEY: Mr. Hails, you tend to be doing  
7 this with microphone. I need you to be doing this with  
8 your microphone. There you go.

9 BY MR. HAILS:

10 Q Okay. We are at slide 129 in the presentation.  
11 Talking about claim 1. And claim 1 refers to a second  
12 shielding compartment. Do you see that in the slide?

13 A I do.

14 Q On the third line it talks about that the second  
15 shielding compartment is for the waste bottle.

16 Do you see that in claim 1?

17 A Yes.

18 Q Does the prior art claim teach a second  
19 shielding compartment for a waste bottle?

20 A Yes, it does.

21 Q Please explain.

22 A Klein had a waste bottle on the top shelf that  
23 was in a shielded container and it had an opening facing  
24 vertically upward.

25 Q And that opening, does Klein teach that that



1 opening is one through which a waste bottle can be inserted  
2 into or removed from the shielding compartment?

3 A Yes, he does.

4 Q All right. Let's move forward, please. I  
5 should note for the record that slide 129 has an excerpt  
6 from figure 2-3 of the Klein thesis.

7 The second door -- I'm sorry, the next element  
8 is a second door that is configured to provide access to  
9 the second shielding compartment and to close over the  
10 second opening. Did you see teachings of this subject  
11 matter in the prior art?

12 A Yes.

13 Q Please explain.

14 A As Klein describes, a waste container was  
15 mounted on the top shelf inside a lead container with a  
16 lid. That lid door sliding whatever, those are very common  
17 in the art.

18 Q Okay. The claim talks about that the second  
19 door is accessible by the opening through the top surface  
20 of the exterior shell of the cart. Did you see that  
21 subject matter talked about in the prior art?

22 A Yes, I did.

23 Q Please explain.

24 A I'm sorry. We previously showed that there was  
25 a door here that opened over this area and that's

1 accessible via the top shelf of the -- the top surface of  
2 the cart.

3 Q Okay. All right. So the slide 129 also lists a  
4 wherein clause from claim 1 saying that the first opening,  
5 which is the one corresponding to the generator shield, is  
6 located at a lower elevation than the second opening, which  
7 is the one corresponding to the waste bottle.

8 Will you please remind the Court how does the  
9 prior art teach that subject matter?

10 A The prior art teaches that subject in that,  
11 first of all, it would be obvious to relocate the generator  
12 and the shielding to a lower elevation for ergonomic  
13 purposes as we've discussed as taught by Chaffin, which  
14 would be well-known to a person who is designing medical  
15 devices and designing any industrial device.

16 Q And Chaffin, for the record, is RX-96. Let's  
17 move forward, please.

18 We are at slide 130 and the claim is talking  
19 about a radioactivity detector. Does the prior art teach a  
20 radioactivity detector?

21 A Yes, it does.

22 Q Please explain.

23 A We have a radioactivity connector at the  
24 activity counter in the Klein thesis as well as in other  
25 sources.

1           Q       For the record, slide 130 shows an excerpt from  
2 Klein figure 2-2 on page 29. Is the radioactivity detector  
3 that's taught by the prior art, is it positioned to measure  
4 radioactivity of a rubidium radioactive eluant that flows  
5 through an eluant tubing line?

6           A       Yes, it does. The tubing is positioned to go  
7 under the radioactivity detector and it counts it while the  
8 tubing -- the fluid is flowing through.

9           Q       Is the eluate tubing line in fluid communication  
10 with an outlet tubing port of the strontium-rubidium  
11 radioisotope generator?

12          A       Yes, it is. It's shown here this is the outlet  
13 and it's in fluid communication.

14          Q       Let's move forward, please. We are on slide 131  
15 of the presentation.

16                   Claim 1 talks about a shielded well on board the  
17 cart configured to receive an eluate reservoir. Please  
18 explain how does the prior art teach this subject matter?

19          A       Well, the Klein thesis has indeed a shielded  
20 well in the dose calibrator and it has the eluate reservoir  
21 in the shielded well while a dose is pumped into it. He  
22 doesn't have it on board the cart, but as we stated  
23 previously, it is obvious to relocate the dose calibrator.  
24 It was done by Tate. It's done by Medrad. There is  
25 nothing inventive in that step.

1           Q       Again for the record, slide 131 shows a split  
2 screen, I guess, with both figure 2-3 and figure 2-2 from  
3 the Klein thesis. Next part of this shielded well element  
4 says that the eluate reservoir is configured to receive a  
5 test sample. Do you see that?

6           A       Yes.

7           Q       Does the prior art teach that subject matter?

8           A       Yes, it does. The computer configures the flow  
9 path. So it goes out to the patient line, which itself is  
10 now inserted into the eluate reservoir so that the eluate  
11 reservoir can receive a test sample.

12          Q       Let's move forward, please. Claim 1 also refers  
13 to the computer of the system and lists a variety of  
14 features. One is to provide a stop button on the  
15 touchscreen display. Did you see a stop button taught by  
16 the prior art?

17          A       As we've seen before, the stop button is there  
18 for all elutions.

19          Q       For the record, slide 132 shows a screenshot  
20 taken from page 64 of the Klein thesis.

21                   Does the prior art teach that that stop button  
22 aborts a function of the infusion system in response to a  
23 user input that activates the stop button?

24          A       That's correct.

25          Q       Let's move forward, please. Claim 1 says that

1 the computer is configured to pump saline from a saline  
2 reservoir. Does the prior art teach this subject matter?

3 A Yes, it does.

4 Q Please explain.

5 A A computer controls a peristaltic pump, which  
6 takes saline from the saline reservoir.

7 Q All right. Please continue.

8 A It then pumps it through the generator valve  
9 into the inlet port of the generator and the eluate is  
10 pumped out of the outlet point -- sorry, through the outlet  
11 tubing port of the generator.

12 Q Okay. So this -- for the record, slide 133  
13 again shows an excerpt from figure 2-2 of the Klein thesis  
14 and Dr. Stone was referring to the peristaltic pump that's  
15 illustrated in that diagram.

16 Okay. The claim says that the saline reservoir  
17 has to be positioned outside the interior space of the  
18 cabinet structure. Does the prior art teach that subject  
19 matter?

20 A Yes, it does.

21 Q Please show us.

22 A The peristaltic pump -- sorry, the saline  
23 reservoir is here outside the cart.

24 Q You're pointing to our own screen. Why don't we  
25 do it on the screen up here.

1           A       Thank you. I'm sorry. Here we have the saline  
2 reservoir outside the cart.

3           Q       And for the record, Dr. Stone was pointing to  
4 figure 2-3 from the Klein cart and the annotation that we  
5 provided on slide 133 for the saline bag.

6                    Okay. Does the pump -- sorry, does the computer  
7 pump saline into the strontium-rubidium radioisotope  
8 generator through an inlet tubing port?

9           A       Yes, it does. Here is the inlet tubing port we  
10 discussed previously.

11          Q       And does that process generate the rubidium  
12 radioactive eluate that is discharged through the outlet  
13 tubing port?

14          A       Yes, it does.

15          Q       Please go back. Go back, please, one more.  
16 Just show us the outlet tubing port on slide 133.

17          A       I'm sorry. Here we have the outlet tubing port.

18          Q       Very good. Let's move forward, please.

19                   All right. We are on slide 134 and we have an  
20 excerpt from figure 2-2 from the Klein thesis. The claim  
21 says that the computer of the infusion system fills the  
22 eluate reservoir in the shielded well on board the cart  
23 with a test sample of the rubidium radioactive eluate.

24                   Please remind the Court, how does the prior art  
25 teach this subject matter?

1           A        Again, this is a block diagram from the Klein  
2       thesis. We have the eluate reservoir in the shielded well  
3       with pumping and filling of that eluate reservoir is  
4       controlled by the computer configuring the valves and the  
5       pump in order to do that. And it doesn't to on board the  
6       cart, but as we say, it's obvious to relocate the cart for  
7       reasons we've discussed with regard to the weight of the  
8       cart and the movement.

9           Q        Okay. And on this drawing here, figure 2-2 from  
10      the Klein thesis, you're pointing to the dose calibrator  
11      color coded in blue; is that correct?

12          A        That is correct.

13          Q        All right. Do you see that box with the little  
14      snowman-looking icon inside?

15          A        Yes.

16          Q        What is that?

17          A        The box with the little snowman is the shielded  
18      well with the eluate reservoir inside.

19          Q        Move forward, please. Claim 1 says that the  
20      computer is configured to determine a strontium  
21      breakthrough test result on the test sample. Please remind  
22      the Court, how does that process occur in the prior art?

23          A        Well, we've discussed the Klein thesis that the  
24      strontium breakthrough test is termed by first measuring  
25      the activity when the sample is first placed in the well.

1 The computer then waits 30 minutes and takes another  
2 reading and computes the strontium breakthrough from the  
3 formula shown below.

4 Q And does that process operate on a test sample  
5 that is filled into an eluate reservoir in a shielded well?

6 A Yes, it does.

7 Q We've discussed here on board the cart, off-cart  
8 analysis already, correct?

9 A That's correct. All the system components were  
10 there in Klein. We are just talking about the  
11 rearrangement for a commercial product.

12 Q All right. Let's go through the while clause in  
13 this element. Does that process occur while the eluate  
14 reservoir remains in the shielded well?

15 A Yes, it does.

16 Q Let's move forward, please. Claim 1 further  
17 states that the computer is configured to not allow a  
18 patient infusion if the strontium breakthrough test result  
19 is greater than or equal to an allowed limit.

20 Please remind the Court, how does the prior art  
21 teach the subject matter?

22 A The Klein thesis states specifically that the  
23 computer disables patient elution -- sorry, that the  
24 computer does not enable patient elutions unless a  
25 calibration run and a successful breakthrough measurement



1 are completed. That's done daily. The amount is  
2 determined by the Health Canada guidelines for the  
3 strontium breakthrough level that's part of the daily  
4 protocol and the system software ensures that the protocol  
5 is followed; that is, each run is enabled only after the  
6 prerequisites have been completed successfully.

7 Q For the record, we are on slide 136 and the  
8 excerpts shown are from pages 39, 43 and 54 of the Klein  
9 thesis. Let's move forward, please.

10 Okay. Claim 1 talks about a shielding for a  
11 generator. Claim 2 says that the infusion system further  
12 comprises the strontium-rubidium radioisotope generator in  
13 the first two compartments. Did you see a  
14 strontium-rubidium radioisotope generator taught by the  
15 prior art?

16 A Yes.

17 Q Can you point it out to us?

18 A Here we have the strontium-rubidium generator in  
19 the interior space of the cabinet shown how we had proposed  
20 that it was obvious to move it, but it was in the interior  
21 previously.

22 Q All right. And so for the record, Dr. Stone is  
23 pointing, again, I think it's to figure 2-3 of the Klein  
24 thesis. So let's move forward, please. We are still on  
25 slide 37.

1           Claim 2 says that there is an eluate reservoir  
2 located in the shielded well on board the cart and in fluid  
3 communication with the eluate tubing line. Did you see  
4 that subject matter taught by the prior art?

5           A       Yes.

6           Q       Please explain.

7           A       We've talked about the eluate reservoir in the  
8 shielded well previously. The on board the cart we've also  
9 talked about that it's obvious to relocate that as it's  
10 done in the prior art and it's in fluid communication with  
11 the eluate tubing line as shown in the hot pink tracing  
12 that we have here.

13          Q       For the record, we are on slide 137 of this  
14 presentation and the hot pink he was referring to was taken  
15 from slide -- I'm sorry, from figure 2-2 of the Klein  
16 thesis.

17                 All right. Let's move forward. So now we are  
18 at claim 3. We are on slide 138. Claim 3 says that the  
19 cabinet structure has a lowermost portion and the platform  
20 has a lower surface. Did you see that subject matter  
21 taught by the prior art?

22          A       Yes.

23          Q       Walk us through it, please.

24          A       Well, we've talked about how it would be obvious  
25 to relocate the generator and the shielding compartment to

1 a lower elevation. Ergonomics is taught by Chaffin. If I  
2 look at this proposed concept here that would happen and  
3 compare it with a standard laboratory countertop, which is  
4 normally found at a height of 30 to 36 inches, so if I  
5 compare that to the opening that we have here in the first  
6 generator, that, by that comparison, my eye is easily  
7 determined to be nominally between one and two feet.

8 Q Let me stop you right there. Right now we are  
9 talking about part one of this claim and it says that the  
10 cabinet structure has a lowermost portion.

11 Let's talk about the cabinet structure just to  
12 make sure the record is clear. Do you see he a cabinet  
13 structure in this picture here on -- I think this is figure  
14 2-3?

15 A Yes, we have a cabinet structure as shown here.

16 Q Does it have a lowermost portion?

17 A It does.

18 Q Did you see a platform in this figure 2-3?

19 A There is a platform here at the base of the  
20 cart.

21 Q And did that platform have a lower surface?

22 A Yes, it does.

23 Q All right. So now let's talk about the next  
24 elements. You were talking about the shielding and let's  
25 walk through the claim. The claim talks about the first

1 opening, which refers back to the shielding compartment of  
2 the generator and that must be at a first elevation.

3 So with the modifications that you've identified  
4 here using the Chaffin teachings, where is the -- what is  
5 the elevation of the first opening?

6 A Here is the first opening. That's where it's  
7 located.

8 Q Okay. So let's go to the next element. And  
9 that refers to a second opening, which refers back to the  
10 shielding compartment for the waste bottle. Applying these  
11 teachings, do you see that the second opening for the waste  
12 bottle would be at a second elevation?

13 A Yes. In the original location that's shown in  
14 Klein's thesis, we had the waste container located there at  
15 that second elevation.

16 Q All right. And then the next -- sorry, the next  
17 element says that the first elevation, which again refers  
18 to the shielding compartment, is between approximately one  
19 foot and approximately two feet with respect to the  
20 lowermost portion of the cabinet structure.

21 Okay. Please walk us through your analysis of  
22 this element.

23 A All right. Thank you. Again, using the  
24 standard height of a laboratory countertop positioned  
25 nominally between 30 and 36 inches for ergonomics for a

1 workspace, using that as a reference of nominally 36  
2 inches, this first opening is halfway through. That would  
3 be between one and two feet.

4 Q And do you recall looking at the shielding  
5 generator this morning for the CardioGen 510?

6 A I do.

7 Q Would you agree with me that the shielding  
8 compartment of that structure also was at the bottom of  
9 that cart?

10 A Yes, it is.

11 Q Let's move forward please to the next element.  
12 So the next element says that the second elevation is  
13 between approximately two feet and approximately three feet  
14 with respect to the lower surface of the platform. Please  
15 explain your analysis of this element.

16 A The Klein thesis had the shielded waste  
17 container with its upwardly facing -- sorry, vertically  
18 upward facing opening was on that shelf. And as you can  
19 see in comparison with the countertop, that's between two  
20 and three feet with respect to the lower surface of the  
21 platform.

22 Q Let's move forward, please. We are at slide 139  
23 and we are talking about claim 4 of the '869 patent.

24 Claim 4 says that the infusion system has a  
25 wherein clause, wherein the first shielding compartment

1 comprises two tubing passageways formed in a perimeter  
2 surface of the first opening.

3 Let's talk about tubing passageways first. In  
4 the Klein thesis, how many tubing passageways go in and out  
5 of the generator?

6 A Well, he has two tubing passageways because we  
7 have to have saline going in. I'm sorry. The system has  
8 saline going in and has the strontium-rubidium eluate  
9 coming out.

10 Q Did you see teachings of a shielding compartment  
11 with tubing passageways formed in a perimeter surface of an  
12 opening of the shielding compartment?

13 A We saw that in the -- the tubing passageways  
14 were formed in the perimeter surface on the Tate system.

15 Q We have an excerpt here from figure 6-E of Tate  
16 on page 119. Show me the tubing passageway.

17 A We have a tubing passageway here formed in the  
18 perimeter of the shielded well.

19 Q And the shielded well, is that element 111 in  
20 this drawing?

21 A Yes.

22 Q Okay. There is only one in the Tate system.  
23 But there are two tubing -- that's tubing in the Klein  
24 system. Why don't you explain why is that -- why is it  
25 obvious to put in two?

1           A       Well, in the Tate system, using FDG, one merely  
2 needs a single tube in order to withdraw eluate -- or in  
3 order to withdraw the radioisotope source -- sample from  
4 the source. In the strontium-rubidium isotope generator,  
5 you need two tubes; one to go in with the saline and one to  
6 bring the eluate out.

7           Q       Do you believe it's obvious to apply the  
8 teachings of Tate and to add a second tubing passageway,  
9 one for the other tube?

10          A       One would have tubing passageways for each of  
11 the tubings. In order to do that, that would be obvious.  
12 It would be the standard thing to do.

13          Q       On this slide 139, you color coded a tube 210 in  
14 pink; is that correct?

15          A       Yes.

16          Q       Okay. The opening in the perimeter surface of  
17 this shielded well, does it pinch that tube?

18          A       No. It certainly would not be obvious to have  
19 anything that would pinch the tube if I need to have flow  
20 through that tube.

21          Q       Does that opening crush the tube?

22          A       No, it does not.

23          Q       And you have this element 684 with the red halo  
24 around it on slide 139. Do you see that?

25          A       I do.

1 Q And remind the Court, what is that?

2 A That is the source of rubidium-82 in a shielded  
3 container.

4 Q I'm sorry. So 684 --

5 A Actually, not rubidium-82. Of FDG.

6 Q We have a shielded well, right, at 111? Please  
7 look at the screen. And what is this guy?

8 A I'm sorry. That is a door over the -- over the  
9 vial of FDG.

10 Q All right. For the record, I was pointing to  
11 684 as "this guy." So that door is closed in the shielded  
12 well. Would you agree?

13 A That's correct.

14 Q All right. Does Tate disclose that that door  
15 crushes or pinches that tube when that door is closed?

16 A No.

17 Q Let's move forward, please. All right. We are  
18 now on slide 140 and we are talking about claim 5 of the  
19 '869 patent.

20 Claim 5 says the opening through the exterior  
21 shell is configured to provide -- that is configured to  
22 provide access to the strontium-rubidium radioisotope  
23 generator within the interior space of the cabinet  
24 structure is through the front side of the exterior shell.

25 Did you see teachings of that subject matter in



1 the prior art?

2 A I do.

3 Q Show me the front side.

4 A Here is the front side.

5 Q And do you see an opening in the front side?

6 A There is indeed an opening in the front side.

7 Q And would that front side provide access to the  
8 generator?

9 A Yes, it does.

10 Q That was slide 140. Let's move to the next  
11 slide, please. So now we are at slide 141 still talking  
12 about the '869 patent. We are now at claim 8.

13 Claim 8 says wherein the infusion system is  
14 configured to determine the strontium breakthrough test  
15 result on the test sample at least once a day. Please  
16 explain how does this prior art teach this subject matter?

17 A Klein clearly describes that his system ensures  
18 compliance with the daily protocol prescribed in the  
19 previous chapter, a flush followed by a calibration run and  
20 successful breakthrough measurement must be completed in  
21 order to enable patient elutions for the remainder of the  
22 day. So at midnight, that daily protocol completed is  
23 cleared and the daily protocol chart has to be completed in  
24 order to enable patient elutions.

25 Q All right. The excerpt on this slide 141 is

1 taken from page 139 of the Klein thesis. We also have an  
2 excerpt showing figure 2-1 from the Klein thesis. Where in  
3 this daily protocol are the patient elution runs?

4 A Patient elution runs are here after the  
5 completion of the calibration run.

6 Q I was going to ask where in the daily protocol  
7 is the calibration run?

8 A It's after the daily flush.

9 Q And you said that the calibration run must  
10 achieve a successful breakthrough measurement in order to  
11 enable the patient elutions; is that correct?

12 A That's correct.

13 Q Let's move forward, please. So we are at slide  
14 142. And we'll talk about claim 14 of the '869 patent.  
15 Claim 14 talks about various functions of the computer of  
16 the infusion system. The first one is to track a volume of  
17 the saline remaining in the saline reservoir.

18 Please explain how does the prior art teach this  
19 subject matter?

20 A Well, saline is a consumable for the system.  
21 The international standard for user interfaces for medical  
22 devices calls for the user to be aware of the use of the  
23 correct consumable, the remaining amount of them, whether  
24 accessories might be used with the medical device, how to  
25 assemble them, how to check their correct functioning.

1 Saline is indeed a consumable. The user is made  
2 aware of that through the user interface as described and  
3 taught by this standard.

4 Q Okay. For the record, on this slide 141 we  
5 haven't excerpt from RX-114 at page 63. All right. And do  
6 you see -- did you see a description of tracking saline in  
7 the Medrad documentation?

8 A Yes, we did.

9 Q We'll get to that in a moment. Did you see a  
10 description of tracking saline in the Tate patent  
11 application?

12 A Yes.

13 Q All right. Let's move forward. Claim 14 also  
14 talks about that the computer is configured to alert the  
15 user via the touchscreen display when the volume of the  
16 saline remaining in the saline reservoir is below a  
17 predetermined volume threshold. Why don't you walk us  
18 through the subject matter.

19 A Yes. As we stated previously, I believe that's  
20 covered by the international standard calling for the user  
21 to be made aware of the correct usage and how to assemble  
22 them, how to check their correct functioning. And it's  
23 also provided to Medrad where they provide an alert when  
24 the remaining saline is low.

25 Q We'll get to that in a moment. Let's move

1 forward, please. Can we move forward.

2 A Yes.

3 Q Okay. So now we are on slide 144. We were  
4 talking about claim 24. Claim 24 first recites a hanger  
5 configured to hold the saline reservoir at an elevation  
6 above the top surface of the exterior shell.

7 Does the prior art teach this subject matter?

8 A Well, the first portion of this element, the  
9 hanger that hangs the saline bag is taught both in Klein  
10 and in the Tate patent application; however, it's clearly  
11 disclosed in the CardioGen Model 510.

12 Q In your experience, is it common or is it an  
13 uncommon configuration to hang saline at high elevations on  
14 medical devices?

15 A It's hung at a high elevation. It makes it  
16 clearly visible from across the room.

17 Q Let's move forward, please. The next element is  
18 a handle that's configured for the user to grasp in order  
19 to move the infusion system. Did you see teachings of  
20 handles in the prior art?

21 A Yes. Handles for moving carts are quite a  
22 standard configuration. We have handles on the Tate  
23 cart -- I'm sorry, on the Klein cart, on the Tate cart, as  
24 well as on the CardioGen Model 510. And we also had it on  
25 Medrad.

1           Q       All right. So for the record, we are on slide  
2 145 and Dr. Stone was pointing out handles on page 34 of  
3 the Klein reference, page 2 of the Tate reference, page 13  
4 of the CardioGen reference. Actually, let me read the  
5 exhibits in. 106 page 34, 103 page 2, RX-207 page 13.

6                   Let's move forward. Next thing -- now we are at  
7 slide 146. The next thing you've got to have is four  
8 wheels mounted to an underside of the platform. Please  
9 explain, how does the prior art teach this subject?

10           A       I believe this is a very standard and obvious  
11 feature. Klein had four wheels. The Tate disclosure had  
12 four wheels. The CardioGen 510 had four wheels.

13           Q       All right. And again, for the record, we are  
14 pointing to pictures taken from page 34 of Klein RX-106,  
15 page 2 of Tate RX-103, page 13 of the CardioGen manual,  
16 RX-207.

17           A       And just to be clear it was also on the Medrad  
18 though I neglected it.

19           Q       All right. Let's move forward, please. Slide  
20 147. Now we have to have a power inlet port for connecting  
21 the infusion system to a power source. Please explain, how  
22 does the prior art teach this subject matter?

23           A       All the devices that we've talked about were  
24 powered by AC mains. Specifically, Klein states that the  
25 system is plugged into a wall socket at all times and only

1 needs several minutes of battery power while being moved.

2 Q For the record, this is slide 48 discussing  
3 claim 24 and the excerpt from which Dr. Stone read was page  
4 46 of RX-106.

5 All right. And moving on. We have a printer  
6 that's configured to print a document concerning a patient  
7 infusion or a quality control test result generated by the  
8 infusion system.

9 Please explain. Does the prior art teach this  
10 subject matter?

11 A Yes, it does.

12 Q Can you walk us through it?

13 A Klein certainly has a printer as he displays  
14 here in the picture from his cart. Printer and power  
15 isolation transformers were mounted on the lower tray.  
16 When one is producing a medical device, it would be obvious  
17 to utilize that printer to print. What is done by Tate and  
18 Medrad, they disclosed printers for printing infusion data.

19 Q Okay. Just for the record, you were referring  
20 to an excerpt from Klein at page 35. You were referring  
21 to -- what are the portions of Tate that you relied on?

22 A We relied on the Tate. The RX-103 page 93.

23 Q Okay. And you said also that you saw evidence  
24 of that in Medrad; is that correct?

25 A Yes. That's RX-0200C page 91.

1 Q Let's move forward, please. Okay. We are at  
2 slide 149 and we are talking about tubing passageways  
3 again. The first shielding compartment comprising two  
4 tubing passageways formed in a perimeter surface of the  
5 first opening. Have we discussed this already earlier in  
6 your testimony?

7 A Yes, we have.

8 Q All right. You have an excerpt taken from Tate  
9 on page 119 figure 6E. Are you relying on the same  
10 disclosure in Tate as before in that prior claim element?

11 A Yes.

12 Q All right. You believe -- so just point out,  
13 please, where is the opening formed in the printer surface  
14 of the shielding compartment?

15 A We have the opening formed in the perimeter of  
16 the surface.

17 Q For the record, that's the opening through which  
18 the tube 210 proceeds. The next portion of this claim is  
19 that each of the two tubing passageways has a depth  
20 configured to prevent pinching or crushing of a  
21 corresponding tubing line routed there through when the  
22 first door is closed thereover. Is the tubing passageway,  
23 is it crushed or pinched in the Tate disclosure?

24 A No.

25 Q Is it crushed or pinched when that cap is closed

1 over the shielding well?

2 A No.

3 Q All right. Let's move to the next one. You  
4 also have to have a first door that's mounted by a hinge.  
5 How is this subject matter taught by the prior art?

6 A Well, we are showing the first door here and  
7 while it's still being mounted on a post, mounting a door  
8 on a hinge, a post, on rails, there is nothing inventive  
9 about how a door is mounted.

10 Q Have you seen in your experience shielding  
11 compartments with doors mounted by hinges before?

12 A Yes, indeed.

13 Q Let's move forward, please. Okay. Now we are  
14 on slide 150. We are still talking about claim 24. It  
15 says that access to an operation of the computer is  
16 regulated through a user login credential.

17 Did you see this subject matter taught by the  
18 prior art?

19 A Yes.

20 Q Please explain.

21 A In the Klein thesis, he states that the  
22 generator information screen is displayed first, which  
23 gives information as to the state of the generator activity  
24 and history. Following is a prompt for a user ID code,  
25 which is useful for avoiding tampering by unauthorized



1 personnel as well as enabling test runs, which were  
2 commonly used during a development cycle, but have no  
3 clinical application.

4 Q All right. And for the record, this excerpt is  
5 taken from page 63 of RX-106. Let's move forward, please.  
6 We are at slide 151 still talking about claim 24. It says  
7 the strontium breakthrough test result is for at least one  
8 of strontium-82 and strontium-85. How does the prior art  
9 teach this subject matter?

10 A Klein discloses the formulas which are used by  
11 the computer and he states the breakthrough of each  
12 isotope, strontium-82 and strontium-85, is calculated as a  
13 relative activity ratio of strontium activity to 82  
14 rubidium activity delivered as demonstrated below.

15 Q For the record, this excerpt is taken from page  
16 61 of Klein. Let's move forward, please. Slide 152. It  
17 says the exterior shell further includes a saline tubing  
18 opening configured for a saline tubing line to pass from  
19 the reservoir outside the exterior shell to the interior  
20 space of the cabinet structure.

21 Why don't you walk us through this subject  
22 matter.

23 A We have a saline bag on the exterior of the  
24 shell and it has a tubing line that passes through to the  
25 interior of the shell.

1 Q For the record, slide 152 has an excerpt taken  
2 from figure 2-4 of the Klein thesis. Dr. Stone was  
3 pointing to the tubing line that we've color coded in pink  
4 that terminates in the pointer for the generator.

5 A And to clarify, I said to the interior shell.  
6 It's to the interior space.

7 Q Thank you. Let's move forward, please. Slide  
8 153. The computer is configured to determine the strontium  
9 breakthrough test result on the test sample at least once  
10 per day. I think we've discussed this, but please just  
11 refresh the Court, how does the prior art show this?

12 A This is a reminder at midnight the system is  
13 reset such that a daily protocol must be completed prior to  
14 enabling -- enabling patient run elutions for the remainder  
15 of the day.

16 Q And for the record, this slide has an excerpt  
17 from page 39 of the Klein thesis and also an excerpt  
18 showing figure 2-1. Let's move forward, please.

19 All right. So staying with claim 24, we are at  
20 slide 154, the computer is configured to pump saline  
21 through the strontium-rubidium radioisotope generator at a  
22 rate that's less than approximately 70 milliliters per  
23 minute. Please explain how the prior art teaches this  
24 subject matter.

25 A Well, Klein clearly states in his thesis that

1 the pump operates at less than 70 milliliters per minute.  
2 For example, during a flush run, flushing of all the lines  
3 in the system as well as 50 milliliters of flush of the  
4 generator at 15 milliliters per minute. That's certainly  
5 less than 70 milliliters per minute.

6 Q And for the record, that excerpt is taken from  
7 page 53 of the Klein thesis. All righty. Let's keep  
8 going. Slide 155. The computer is configured to track a  
9 volume of the rubidium radioactive eluate discharged from  
10 the generator to the waste bottle.

11 Does the computer track a volume of eluate  
12 discharge from the generator to the waste bottle in the  
13 prior art?

14 A Yes, it does.

15 Q Please explain.

16 A During a flush run, he flushes all of the lines  
17 in the system as well as 50 milliliters of flush of the  
18 generator at 15 milliliters per minute. That flush of the  
19 generator goes to the waste bottles.

20 Q So what's the quantity that is tracked by the  
21 computer?

22 A 50 milliliters.

23 Q 5-0; is that correct?

24 A 5-0.

25 Q And that excerpt is taken from page 53?

1           A        I'm sorry. I did not hear that.

2           Q        Just for the record, that excerpt is taken from  
3 page 53. The claim also talks about the computer being  
4 configured to control the touch screen display to display a  
5 user screen guiding the user to empty the waste bottle.

6                    How does the prior art teach this subject  
7 matter?

8           A        Again, Klein provides guidance to replace some  
9 of the consumables to the generator, but he also provides  
10 direct guidance to present reminders to empty the waste  
11 bottle. If the level switch is tripped, the current  
12 elution continues to completion, but a new run is not  
13 permitted. If a new run is attempted without emptying the  
14 waste container, an error is produced and the elution does  
15 not proceed until the waste container is emptied and the  
16 elution is restarted.

17          Q        And that excerpt is taken from page 45 of the  
18 Klein thesis. Let's move forward, please. Okay. Did you  
19 also see teachings of this subject matter in other  
20 references?

21          A        Yes. As we read before, the user is -- has to  
22 be aware of the current use of the correct consumable, the  
23 remaining amount, whether accessories might be used with  
24 the device, how to assemble them and how to check the  
25 correct functioning.

1 Q And what about Medrad? Why do you have Medrad  
2 listed here?

3 A Medrad specifically has a reminder to empty the  
4 waste bag.

5 Q Let's move forward, please. The computer is  
6 also configured to track a volume of the saline remaining  
7 in the saline reservoir and to alert the user via the  
8 touchscreen display when the volume of the saline remaining  
9 in that reservoir is below a predetermined volume  
10 threshold. Again, walk us through your analysis of this  
11 subject matter, please.

12 A Again, the same international standard calls for  
13 the remaining amount of consumables to be made known and  
14 how the user is to use them. And Medrad tracks the volume  
15 of saline remaining in the saline reservoir and provides  
16 alerts to replace it.

17 Q And we don't have it here, but did you also see  
18 teachings of that subject matter in the Tate reference?

19 A Yes.

20 Q Let's move forward, please. All right. So we  
21 are done with claim 24 and now we are on to claim 27. We  
22 are on slide 158 of your presentation. This claim calls  
23 for a dose calibrator located in the shielded well on board  
24 the cart and in communication with the computer. Where is  
25 the dose calibrator in the Klein system?

1           A        In the Klein system, the dose calibrator was off  
2 the cart, but it was in communication with the computer.

3           Q        Does it have a dose calibrator located in the  
4 shielded well?

5           A        Yes, it is.

6           Q        The next part of this claim says wherein the  
7 dose calibrator is configured to determine the strontium  
8 breakthrough test result. Why don't you explain how does  
9 the dose calibrator participate in strontium breakthrough  
10 test results?

11          A        The dose calibrator acts, as we've described  
12 previously, a sample of eluate is pumped into the  
13 reservoir. The radiation levels are taken by the computer.  
14 The computer waits, takes another radiation level and  
15 computes the strontium breakthrough level.

16          Q        All right. For the record, Dr. Stone was  
17 pointing earlier to the dose calibrator shown on figure 2-2  
18 on page 29 of the Klein thesis.

19                 Let's move forward, please. Okay. We are at  
20 slide 159. We are still talking about claim 27. It says  
21 wherein the opening through the exterior shell configured  
22 to provide access to the strontium-rubidium radioisotope  
23 generator within the interior of the cabinet structure is  
24 through the front side of the exterior shell. Did you see  
25 this subject matter taught by the prior art?

1           A       Yes.

2           Q       Show me the front side opening, please.

3           A       We have the front side opening of the cart in  
4 the Klein thesis.

5           Q       And does it provide access to the generator?

6           A       Yes, it does.

7           Q       All right. And is that generator inside the  
8 interior space of the cabinet structure?

9           A       Yes, it is.

10          Q       All right. For the record, we were pointing to  
11 the picture from page 34 of the Klein thesis. Let's move  
12 forward, please.

13                   So now we are at slide 160. We have an excerpt  
14 again, the block diagram taken from figure 2-2 of the Klein  
15 thesis on page 29. And we are talking about claim 8 of the  
16 '869 patent. All right. It says the infusion system  
17 comprises the generator with the inlet tubing port  
18 configured to receive saline. Did you see this subject  
19 matter taught by the prior art?

20          A       Yes.

21          Q       Show me.

22          A       Here we have the generator configured to  
23 receive --

24          Q       Does it receive saline?

25          A       Yes. It receives saline.

1           Q       All right. So that's the -- thank you. Just  
2 for the record, that's the hot pink coded stuff here on  
3 slide 160 that connects the saline IV through the pump to  
4 the generator.

5                    The second part of this element says the  
6 generator also has the outlet tubing port configured to  
7 discharge the rubidium radioactive eluate. Why don't you  
8 point that out to the Court.

9           A       Here we have a discharge -- sorry, an outlet  
10 tubing port that discharges the rubidium radioactive  
11 eluate.

12           Q       And for the record, that's the pink color-coded  
13 extension from the generator to the input of the activity  
14 detector shown here on slide 160.

15                    Let's move forward please. Claim 28 also says  
16 that the system comprises the eluate reservoir located in  
17 the shielded well on board the cart and in fluid  
18 communication with the eluate tubing line. Will you point  
19 out the eluate reservoir for us in this drawing?

20           A       Once again, the eluate reservoir is here in blue  
21 in the shielded well off cart, but we've discussed how it's  
22 obvious to relocate the dose calibrator on the cart.

23           Q       So for the record, Dr. Stone is pointing to the  
24 blue color-coded assembly that we have shown here on slide  
25 161 and specifically to the inside of the box, let's say,



1 in the center top. We talked about on board the cart, off  
2 the cart. Why don't you explain, how is this eluate  
3 reservoir in fluid communication with the eluate tubing  
4 line?

5 A From the outlet port of the generator through  
6 the patient valve on to the eluate reservoir we have fluid  
7 communication.

8 Q And so for the record, that's the tubing  
9 extensions that we color coded between the generator that  
10 was shown in red and the dose calibrator that's coded in  
11 blue. The extension is hot -- coded in hot pink.

12 Okay. Let's move forward. We are at slide 162.  
13 We are still talking about claim 28. The next element says  
14 there is a waste tubing line in fluid communication with  
15 the eluate tubing line and the waste bottle. Why don't you  
16 point out where is the waste bottle in Klein?

17 A The waste bottle is here shown in green.

18 Q All right. And where is the waste tubing line  
19 that's in fluid communication with the eluate tubing line?

20 A We have the waste tubing line in fluid  
21 communication with the eluate tubing line.

22 Q Okay. And again, then for the record, that's  
23 the extension that is coded in pink from the generator,  
24 through the activity detector, through the patient valve  
25 and to the green coded waste container. Let's move

1 forward, please.

2 Claim 28 says you've got to have a valve  
3 configured to control fluid flow between the eluate tubing  
4 line and the waste bottle via the waste tubing line.  
5 Please show us how does the prior art teach this valve?

6 A Here we have the patient valve, which controls  
7 the fluid flow between the eluate tubing line and the waste  
8 bottle being in the waste line.

9 Q For the record, we are on slide 163. Dr. Stone  
10 was pointing to the patient valve assembly that we've color  
11 coded in orange. Let's move forward, please. Can we move  
12 forward.

13 Now we are at slide 164. And we've moved to  
14 claim 29 of the '869 patent. This claim says that the  
15 infusion system has a computer that's configured to measure  
16 an activity of the test sample filled into the eluate  
17 reservoir in the shielded well on board the cart. Which  
18 component in the Klein system measures activity of the test  
19 sample in the eluate reservoir?

20 A The computer measures the activity.

21 Q Okay. Where is the activity?

22 A The activity is in the sample vial that is in  
23 the dose calibrator.

24 Q Okay. Does the computer measure this activity  
25 while the eluate reservoir remains in the shielded well on

1 board the cart?

2 A Remains in the shielded well. He doesn't have  
3 it on board the cart. We've talked about how it's obvious  
4 to relocate it for ergonomic and utility reasons.

5 Q The wherein clause of this first element says  
6 that the activity is measured with the dose calibrator in  
7 the shielded well on board the cart. How does the prior  
8 art teach this subject matter?

9 A It's measured in the shielded well in the dose  
10 calibrator. It's not on board the cart, but it's obvious  
11 to relocate that on board the cart.

12 Q Let's move forward, please. Okay. This new  
13 element says that the computer calibrates the infusion  
14 system based on the activity measured by the dose  
15 calibrator. Will you walk us through the calibration  
16 operation?

17 A The Klein thesis he describes the calibration  
18 run as flushing of the generator at 15 milliliters per  
19 minute over 60 seconds into the dose calibrator.  
20 Obviously, that's into the sample vial in the dose  
21 calibrator.

22 Q And what is being calibrated?

23 A What is being calibrated in this case is the  
24 activity detector that's on board the cart.

25 Q Okay. And -- go ahead. I'm sorry.

1           A       The integral activity is recorded from the dose  
2   calibrator and it's used to calibrate the activity counter  
3   and verify that the calibration constant is within  
4   tolerance from previous records.

5           Q       And do you recall how we color coded the  
6   activity counter in prior drawings?

7           A       I believe that was in purple.

8           Q       All right. Let's move forward. Okay. So now  
9   we are at slide 166 and on claim 30 of the '869 patent.  
10   Have we seen this subject matter of claim 30 earlier in  
11   your testimony?

12          A       Indeed we saw this in claim 3, and my analysis  
13   of these elements remain the same as it was in claim 3.

14          Q       All right. So can we just walk through them.  
15   The cabinet structure element from claim 30, is it the same  
16   as claim 3?

17          A       Yes.

18          Q       Let's move forward, please. The first opening,  
19   are they recited the same in claim 30 and in claim 3?

20          A       Yes.

21          Q       Move forward, please. The second opening, are  
22   they recited the same in claim 30 and claim 3?

23          A       Yes.

24          Q       Next element, first elevation with one foot and  
25   two feet. Is the recitation the same between claim 30 and

1 claim 3?

2 A My analysis remains the same.

3 Q All right. And let's move forward. In the last  
4 element, is that the same between these two claims?

5 A Yes.

6 Q All right. So that was slide 166. Let's move  
7 forward, please. All right. So now we are at the '870  
8 patent at slide 167. And if you would advance, now we are  
9 at slide 168.

10 We've got to start all over again. We've got  
11 claim 1, which is a method of using an infusion system on  
12 board a cart to deliver a rubidium radioactive eluate.  
13 Does Klein disclose an infusion system on board a cart to  
14 deliver a rubidium radioactive eluate?

15 A Yes, he does.

16 Q All right. The first element says you've got to  
17 install a saline reservoir on the infusion system. Does  
18 the prior art teach this subject matter?

19 A Yes.

20 Q Can you point out the saline reservoir?

21 A Here is the saline reservoir.

22 Q For the record, Dr. Stone was pointing to the  
23 illustration taken from page 34 of the Klein thesis, the  
24 markings we have on slide 168 for the saline bag. The  
25 wherein clause of this element says that the infusion

1 system comprises a platform and an exterior shell extending  
2 upwardly above the platform. Did you see this subject  
3 matter taught by the prior art?

4 A Yes, I did.

5 Q All right. Why don't you advance. I think it  
6 would be easier. Why don't you point out where is the  
7 platform and where is the exterior shell?

8 A As previously discussed, here is the platform.  
9 Here is an exterior shell.

10 Q So for the record, Dr. Stone was pointing to the  
11 annotations provided in yellow on the illustration taken  
12 from page 34. Why don't you advance.

13 Claim 1 says wherein the platform and the  
14 exterior shell collectively define an interior space of the  
15 cabinet structure. Did you see an interior space disclosed  
16 by the prior art?

17 A As we see here labelled an interior space in the  
18 Klein cart.

19 Q And again, for the record, Dr. Klein -- wow --  
20 Dr. Stone was pointing to the yellow annotations for the  
21 interior space that are labelled here on slide 168.

22 All right. Let's keep going, please. Claim 1  
23 says you've got to place the saline reservoir in fluid  
24 communication through a saline tubing line with an inlet  
25 tubing port of a strontium-rubidium radioisotope generator.

1 Does the prior art teach this subject matter?

2 A Yes. We still have the saline reservoir placed  
3 in communication with an inlet tubing line the inlet port  
4 of the strontium-rubidium generator.

5 Q So we are on slide 169. And for the record,  
6 Dr. Stone was pointing to figure 2-2, the block diagram,  
7 pointing out the saline IV in brown, the generator in red  
8 and the tubing lines in pink that extend between them.

9 All right. We are up to the generator. That  
10 generator, the claim says, must be located in a first  
11 shielding compartment in the interior space of the cabinet  
12 structure. How does the prior art teach that subject  
13 matter?

14 A We've already discussed the shielded rings that  
15 Klein utilized and it's in the interior space of the  
16 cabinet. When I say utilized, in order to place the  
17 shielded generator.

18 Q All right. Is the generator inside the cart?

19 A Yes, it is.

20 Q Move forward, please. We are at slide 170.

21 There is a wherein clause. The strontium-rubidium  
22 radioisotope generator further comprises an outlet tubing  
23 port configured to discharge the rubidium radioactive  
24 eluate. Do you see an outlet tubing port?

25 A I do. Here it is located and it discharges the

1 rubidium radioactive eluate.

2 Q For the record, we are on slide 170. Dr. Stone  
3 pointed out the pink extension, let's say, on the far  
4 right-hand side of the excerpt from figure 2-2 of the Klein  
5 thesis on page 29. Let's move forward, please.

6 Claim 1 says the first shielding compartment has  
7 to have a first opening facing vertically upwardly. Do you  
8 see that subject matter taught by the prior art?

9 A Yes.

10 Q What's the structure?

11 A The structure was with Klein, the generator was  
12 placed in the cart surrounded by lead rings to provide  
13 maximum radiation shielding that still faces vertically  
14 upward. The shielded wells were also taught by Tate and  
15 Medrad with openings that faced vertically up.

16 Q So that's slide 171. Let's keep going. Woops.  
17 Let's not do that. And we move to the next slide. We are  
18 at slide 172. Talking about claim 1 of the '870 patent.  
19 The claim says inserting a waste bottle into a second  
20 shielding compartment on board the cart. Show us how the  
21 prior art teaches this subject matter, please.

22 A I indeed had a shielding compartment for a waste  
23 bottle that was on board the cart.

24 Q Okay. The second shielding compartment on board  
25 the cart has to have a second opening facing vertically



1 upwardly. How does the prior art teach this subject  
2 matter?

3 A Klein had a lead container for the waste bottle  
4 with a lid that faced vertically upward.

5 Q Okay. And then the last piece of this claim  
6 element says that the -- I think the waste bottle opening  
7 must be at a higher elevation than the first opening, which  
8 is the one corresponding to the generator. And remind the  
9 Court, how is this subject matter taught by the prior art?

10 A It would be obvious for ergonomic reasons to  
11 place the generator at the lower level of the cart.

12 Q Okay. Let's move forward, please. We are at  
13 slide 173. We have excerpts taken from both pages 34 and  
14 the block diagram on page 29 of the Klein thesis. We are  
15 talking about claim 1 and the claim element refers to  
16 placing the waste bottle in fluid communication with the  
17 outlet tubing port of the strontium-rubidium radioisotope  
18 generator through an eluate tubing line. Please walk us  
19 through the subject matter.

20 A We have the outlet tubing port in hot pink here  
21 from the strontium generator through the patient valve  
22 directed to the waste container in fluid communication.

23 Q And for the record, we are on slide 173.  
24 Dr. Stone was pointing to the pink coded tubing lines on  
25 the slide and also to the waste container coded in pink --

1     sorry, in green. Let's move forward, please.

2                     Still on slide 173. Wherein a computer on board  
3     the cart is configured to control the fluid communication  
4     between the waste bottle and the outlet tubing port. First  
5     of all, why don't you call out the computer for us.

6             A        The computer is shown here in light green.

7             Q        All right. So that's the green-coded computer  
8     on figure 2-2. Please explain how does that computer  
9     control the fluid communication between the waste bottle  
10    and the outlet tubing port of the generator?

11            A        Klein's thesis discloses an interface board that  
12    is utilized to control the patient valve, which in turn  
13    controls the communication between the generator and the  
14    waste bottle.

15            Q        All right. And for the record, Dr. Stone was  
16    pointing to the patient valve that is coded in orange on  
17    slide 173. All right. Let's move forward, please. A  
18    wherein clause. Wherein the computer has a touchscreen  
19    display. Does Klein disclose a computer with a touchscreen  
20    display?

21            A        He does over here.

22            Q        That's -- for the record, Dr. Stone is pointing  
23    to the photograph on slide 173 taken from page 34. Is that  
24    computer mounted on a vertical post with a top end  
25    extending above the cabinet structure?

1           A       It is. Here is the top of the cabinet structure  
2 and here is the vertical post.

3           Q       All right. And again, for the record, we are  
4 pointing to the excerpt taken from page 34 of RX-106.  
5 Let's move forward please.

6                    Claim 1 says we are inserting an eluate  
7 reservoir in a shielded well on board the cart. Please  
8 explain, how does the prior art teach that subject matter?

9           A       Klein has the eluate reservoir in the shielded  
10 well of the dose calibrator and we've talked about how that  
11 would be obvious to relocate the dose calibrator and the  
12 eluate reservoir on board the cart as disclosed and taught  
13 by Tate and Medrad.

14          Q       So that's slide 174. Let's keep going. We are  
15 still on slide 174. The next element is placing the eluate  
16 reservoir in fluid communication with the eluate tubing  
17 line. Please explain, how does that eluate reservoir get  
18 put in communication with the eluate tubing line?

19          A       In this case, the computer in green once again  
20 controls the patient valve to direct the fluid to the  
21 patient line which now is connected to the eluate reservoir  
22 and as we've said, that would be obvious to place it on  
23 board the cart.

24          Q       And just to knock this out, the computer is  
25 configured to control the fluid communication between the

1 eluate reservoir and the eluate tubing line. Remind us how  
2 does the computer control this?

3 A The computer controls it with software through  
4 this computer control -- through this interface port to the  
5 valve.

6 Q All right. And for the record, Dr. Stone is  
7 pointing to the patient valve color coded in orange on  
8 slide 174.

9 Okay. Let's move forward. We are on to slide  
10 175. Claim 1 says you're pumping a -- sorry, you're  
11 pumping a sample of the rubidium radioactive eluate into  
12 the eluate reservoir in the shielded well on board the  
13 cart. Why don't you show us the pumping.

14 A So here we have the computer controlling the  
15 peristaltic pump which in turn pumps saline into the  
16 generator. The generator has now the rubidium radioactive  
17 eluate coming out going through the patient valve into the  
18 reservoir, and we've discussed the on board the cart fairly  
19 extensively.

20 Q Okay. For the record, we are -- Dr. Stone is  
21 referring to color codings on slide 175 extending from the  
22 pump through to the dose calibrator. Let's move forward,  
23 please.

24 Claim 1 says there is measuring a radioactivity  
25 of the sample of the rubidium radioactive eluate flowing

1 through the eluate tubing line with a radioactivity  
2 detector on board the cart. Did you see the subject matter  
3 in the prior art?

4 A Yes.

5 Q Would you point out the radioactivity detector?

6 A Here we have the radioactivity detector, which  
7 is measuring the activity while the eluate is flowing  
8 through tubing on its way to the reservoir.

9 Q For the record, we are on slide 176 and  
10 Dr. Stone has called out the activity counter color coded  
11 in purple and also the tubing line color coded in pink that  
12 extends from the generator to the dose calibrator. All  
13 right. Is that radioactivity detector on board the cart?

14 A Yes, it is.

15 Q All right. And does that radioactivity detector  
16 do its job while the sample of the rubidium radioactive  
17 eluate is flowing through the eluate tubing line?

18 A Yes, it does.

19 Q Let's move forward, please. All right. We are  
20 on slide 177 still talking about claim 1 of the '870  
21 patent. This claim element says that we are measuring a  
22 calibration radioactivity of the sample pumped into the  
23 eluate reservoir, and let's stop there.

24 How does this -- how does the prior art teach  
25 measuring a calibration radioactivity of the sample?

1           A       Klein teaches measuring the radioactivity of  
2   that sample when it is first pumped into the eluate  
3   reservoir.

4           Q       All right.  So that sample is pumped into the  
5   eluate reservoir; is that right?

6           A       That's correct.

7           Q       Is it in the shielded well?

8           A       It is.

9           Q       All right.  It's not on board the cart, but  
10   we've discussed that, right?

11          A       No.

12          Q       Does that measuring occur while the eluate  
13   reservoir remains in the shielded well?

14          A       Yes.  That's the way the system works.

15          Q       All right.  Let's move forward, please.  All  
16   right.  We are on slide 178.  We've got a comparison step.  
17   This one is kind of wordy.  All right.  Let's look four  
18   lines from the bottom.  Do you see this?

19          A       Yes.

20          Q       All right.  I'm going to try to walk you through  
21   you this claim element.  First, we are comparing, one, the  
22   radioactivity of the sample of the rubidium radioactive  
23   eluate flowing through the eluate tubing line measured by  
24   the radioactivity detector on board the cart while the  
25   sample of the rubidium radioactive eluate is flowing

1 through the eluate tubing line. That's one thing that we  
2 are -- we've taken.

3 And then second, the claim says we are comparing  
4 that with the calibration radioactivity of the sample  
5 pumped into the eluate reservoir in the shielded well on  
6 board the cart. Why don't you explain this to the Court.  
7 How does the prior art teach this subject matter?

8 A Well, Klein discloses that flushing of the  
9 generator into the dose calibrator and the integral  
10 activity recorded from the dose calibrator while that is  
11 done as the integral activity while it's flowing through  
12 this line is used to calibrate the activity counter --  
13 sorry, the integral activity and then the dose calibrator's  
14 reading here is used to calibrate that activity counter and  
15 verify the calibration constant is within tolerance from  
16 previous readings.

17 Q All right. For the record, this slide 178  
18 refers to an activity counter color coded in purple and a  
19 dose calibrator color coded in blue taken from this block  
20 diagram on page 29 of the Klein thesis. So am I correct  
21 that the activity counter measures one radioactivity?

22 A Yes.

23 Q And the dose calibrator measures a second  
24 radioactivity?

25 A That's correct.

1 Q And are they for the same sample of rubidium  
2 eluate?

3 A Yes, they are.

4 Q All right. And who does the comparison of those  
5 readings taken from those two components?

6 A The computer.

7 Q All right. Let's move forward, please. We are  
8 at slide 179 still talking about claim 1, and this element  
9 refers to determining a strontium breakthrough test result.  
10 You've got to determine the strontium breakthrough test  
11 result on the sample pumped into the eluate reservoir in  
12 the shielded well on board the cart.

13 Again, walk us through, how does the prior art  
14 show this subject matter?

15 A Well, again, Klein has the sample pumped into  
16 the reservoir and radioactivity readings taken at the  
17 appropriate times does the computation. It doesn't do it  
18 on board the cart, but we've shown that it's obvious to  
19 place it on board the cart.

20 Q All right. This claim element says that that  
21 process occurs while the eluate reservoir remains in the  
22 shielded well on board the cart. We've talked about the on  
23 board the cart piece. But the rest of it, does this  
24 process occur while the eluate reservoir remains in the  
25 shielded well?



1           A        Radioactivity readings are taken while the  
2 eluate reservoir is in the shielded well of the dose  
3 calibrator.

4           Q        All right. Claim 1 has -- this element has a  
5 wherein clause. The computer of the infusion system is  
6 further configured to not allow a patient infusion if the  
7 strontium breakthrough test result is greater than or equal  
8 to an allowed limit. Again, please walk us through this  
9 subject matter.

10          A        Enablement of patient elutions is reset to not  
11 being enabled each night at midnight and a daily protocol  
12 must be completed in order to -- with a successful  
13 breakthrough run measurement as it states here -- must be  
14 completed in order to enable patient elutions for the  
15 remainder of the day. This is done by the system. The  
16 amount is utilized compared to strict Health Canada  
17 guidelines and the data protocol computes that strontium  
18 breakthrough in accordance with the formulas as we've  
19 discussed previously.

20          Q        Okay. Thank you. So for the record, we are on  
21 page 179 of the presentation. And the excerpts from which  
22 Dr. Stone was referring are pages 39, 43 and, in this case,  
23 28 of the Klein thesis. Let's move forward, please.

24                 All right. We are at slide 180. And now we get  
25 to talk about claim 2 of the '870 patent. You have to

1 place the eluate tubing line in fluid communication with a  
2 patient. Does that happen in the prior art?

3 A Yes, it does.

4 Q Where does the patient go?

5 A The patient line is here at the imaging system.  
6 The patient line is placed in eluate -- I'm sorry, the  
7 eluate tubing line is placed in fluid communication with  
8 the patient by the patient valve of the system and that's  
9 done -- controlled by the computer through the interface  
10 board.

11 Q For the record, we have an excerpt here on slide  
12 180 from page 29 of the Klein thesis. When Dr. Stone was  
13 referring to the patient, he was referring to the thing  
14 color coded in yellow. What did you call it? It's not a  
15 person. What is that?

16 A I'm sorry.

17 Q What is the thing color coded in yellow?

18 A In yellow, that's the imaging system with the  
19 patient lying on the tray there.

20 Q And Dr. Stone also referred to the tubing that  
21 connects the generator that's coded in pink to that imaging  
22 system. All right. And then I'm not sure the record is  
23 clear, so let's go back through this piece. There's a  
24 wherein clause here in claim 2 that says the computer is  
25 further configured to control the fluid communication

1 between the eluate tubing line and the patient. Again,  
2 walk us through how -- what are we seeing here on slide 180  
3 showing?

4 A The first portion here is the eluate tubing line  
5 and the valve -- the patient valve controlled by the  
6 computer controls the flow to the patient.

7 Q All right. So for the record, Dr. Stone is  
8 pointing to the patient valve that's color coded in orange  
9 on slide 180.

10 A And I should have said the fluid communication  
11 rather than the flow.

12 Q Let's keep going. All right. We are at slide  
13 181 still talking about claim 2. The method includes  
14 pumping a dose of the rubidium radioactive eluate to the  
15 patient. Does that occur in the prior art?

16 A Yes, it does.

17 Q Tell us.

18 A We'll start with the computer which controls the  
19 peristaltic pump taking saline from the bag in hot pink  
20 line that goes on through the generator valve to the inlet  
21 port of the generator eluating now rubidium chloride  
22 through the outlet port of the generator through the -- or  
23 sorry, through the activity detector through the patient  
24 valve to the patient and that's the pumping action all  
25 controlled by the computer.

1 Q All right. And for the record, slide 181 has  
2 the same excerpt taken from page 29 of the client thesis,  
3 and Dr. Stone was walking us through those components.  
4 Let's go forward.

5 All right. Claim 2 says there is flushing of  
6 the rubidium radioactive eluate remaining in at least a  
7 portion of the eluate tubing line into the patient. So how  
8 does Klein teach flushing eluate that remains in a tubing  
9 line into the patient?

10 A As shown here in the diagram, when the  
11 appropriate dose has been measured here at the activity  
12 counter, this valve is -- sorry, this valve shifts now to  
13 pump saline from this location on and flushes any remaining  
14 radioactive eluate into the patient.

15 Q Okay. All right. So the system decides that  
16 the patient has had enough. Is that really where we are  
17 at?

18 A That's where we are at.

19 Q All right. And when that decision is made,  
20 where is the remaining radioactivity that's relevant under  
21 this claim?

22 A The radioactivity here, which hasn't been  
23 measured yet, but measured activity is from this point on  
24 to the patient.

25 Q All right. For the record, Dr. Stone is

1 pointing to the activity counter on the slide 182, which  
2 probably is the only component not color coded for us.

3 Okay. So that's where the radioactivity occurs.  
4 How does the system push that radioactivity into the  
5 patient?

6 A By switching the inlet -- sorry, the saline  
7 that's being pumped from the generator input inlet through  
8 the bypass line and forcing any fluid remaining from this  
9 point on on into the patient.

10 Q All right. So let's walk through the rest of  
11 this claim to make sure that we've got it captured. That  
12 flushing must occur by pumping saline from the saline  
13 reservoir to the eluate tubing line through a bypass line.  
14 Show us the bypass line, please.

15 A Bypass line is shown here.

16 Q All right. So for the record, Dr. Stone is  
17 pointing to a pink color coded line that extends through  
18 the orange generator valve. It is the top of the two lines  
19 extending into the generator valve let's say.

20 All right. That bypass line has to bypass the  
21 strontium-rubidium radioisotope generator. Does that occur  
22 in the prior art?

23 A It does. Here is the generator. There is no  
24 longer flow in this line. Instead, it bypasses the  
25 generator and goes to the junction point just before the

1 activity counter.

2 Q And then the claim also says that the computer  
3 is configured to control fluid communication via the bypass  
4 line. Just remind the Court, how does the computer control  
5 all this?

6 A The computer controls that by controlling this  
7 valve through the interface board that's shown in orange.

8 Q All right. Thank you. So let's move forward,  
9 please. So now we are on slide 183. We get to talk about  
10 claim 8 of the '870 patent. Claim 8 says that the computer  
11 of the infusion system is configured to present on the  
12 touchscreen display a screen for starting the patient  
13 infusion by touching a button on the touchscreen display.

14 Does the prior art teach this subject matter?

15 A Yes. Klein displays a screen with a button to  
16 start the patient infusion. They are asked if they want to  
17 start the constant activity elution here in this sample  
18 screen and the patient must push on the button outlined in  
19 yellow here the yes button in order to start that constant  
20 activity elution.

21 Q All right. So for the record, there is a  
22 screenshot illustrated here on slide 183 that's taken from  
23 page 64 of the Klein thesis. And Murphy's Law, there's  
24 probably like eight of those screenshots, but this one is  
25 the one that's entitled "Start constant activity elution"

1 question mark on the green gray screen.

2 Let's move forward, please. All right. So  
3 claim 8 also says that the computer is configured to  
4 present on the touchscreen display a screen reminding the  
5 user to insert the eluate reservoir in the shielded well on  
6 board the cart. Please explain your analysis of this  
7 element.

8 A Well, here we have one of the differences  
9 between a prototype and a device that's configured for  
10 operations. The system provides reminders in Klein to  
11 operating personnel, but not for an eluate reservoir. But  
12 we have pointed out that the -- that he does have warnings  
13 and he has -- and we have other evidence that shows that  
14 that would be obvious to do.

15 Q Let's go on to slide 184. Why don't you  
16 advance, please, and now we get to go to slide 185. You  
17 have an excerpt from this international standard. Is that  
18 the subject matter that you're referring to?

19 A Yes. This is where the international standard  
20 teaches that the user has to be aware of the correct  
21 consumable, how to assemble them and how to check the  
22 correct functioning. That vial is a consumable and its use  
23 must be made known to the user.

24 Q I don't think we've shown the Court this picture  
25 shown here on the lower left-hand side of slide 185. What

1 is that?

2 A That's a diagram from the Klein thesis that  
3 shows the vial in the dose calibrator, which is itself a  
4 shielded well, but it's also had additional shielding  
5 surrounding it, though he's not showing the additional  
6 shielding. He's showing that ion chamber that is a  
7 shielded well for measurements.

8 Q Okay. So we've been using the word "eluate  
9 reservoir." What's the eluate reservoir in Klein's system?

10 A Klein calls it a vial.

11 Q Let's move forward, please. We are at slide  
12 186. Claim 8 further refers to a computer that is  
13 configured to present on the touchscreen display a screen  
14 indicating that the patient infusion is in process. Does  
15 the prior art teach this subject matter?

16 A It does.

17 Q Why don't you walk us through it, please.

18 A So Klein's thesis states that the realtime  
19 graphics display must include a system diagram with updated  
20 information about the state of the system. This includes  
21 the current activity rate reading, the flow rate, the valve  
22 status, expected accumulated activity at the patient  
23 outlet. In addition, progress bars must be included for  
24 each stage of the elution so as to facilitate monitoring of  
25 the system. And an emergency stop button must be enabled



1 throughout the elution and take immediate effect to bring  
2 the system to the safe mode.

3 Q Okay. So do you think this disclosure teaches a  
4 display screen that indicates a patient infusion is in  
5 process?

6 A Yes.

7 Q All right. And then the wherein clause of this  
8 element says that the screen indicating that the patient  
9 infusion is in process displays a stop button to abort the  
10 patient infusion. How does the prior art teach this  
11 subject matter?

12 A Indeed, Klein shows a stop button for one of the  
13 steps that's taking place as shown here on the diagram in  
14 yellow.

15 Q All right. For the record, Dr. Stone is  
16 pointing to the screenshot excerpted from page 64, the stop  
17 button that is coded in yellow on slide 186 of the  
18 presentation. All right. Let's move forward, please.

19 Claim 8 says that the computer of the infusion  
20 system is configured to present on the touchscreen display  
21 the strontium breakthrough test result. Do you see this  
22 subject matter taught by the prior art?

23 A Yes.

24 Q Why don't you explain it to us.

25 A First of all, he says that a record is kept of

1 all the completed elutions for analysis and filing. But he  
2 also states at the end of an elution, reports must be  
3 generated based on the type of elution and its mode of  
4 completion. In addition, a separate window must list a  
5 comprehensive display of all statistics in addition to  
6 activity curves relating to the activity rate and the  
7 integrated activity at the patient outlet.

8 Q Okay.

9 A I believe it's obvious that strontium  
10 breakthrough test results are a statistic that is related  
11 to the elution.

12 Q So you think a person of skill would think that  
13 strontium breakthrough test results are a relevant  
14 statistic to the calibration run?

15 A Yes.

16 Q All right. Let's move forward. All right. So  
17 we are on slide 188 and we are talking about claim 9 and  
18 there are four elements listed here. The first one says  
19 that there is logging into the computer by entering a user  
20 login credential on the touchscreen display.

21 Please explain how the prior art teaches this  
22 subject matter.

23 A As we've seen before, that the generator  
24 information screen is displayed first and following there's  
25 a prompt for number one, a user ID code, which is useful

1 for avoiding tampering by unauthorized personnel as well as  
2 enabling test runs.

3 Q All right. So that excerpt is taken from page  
4 63 of RX-106. The next element of claim 9 is entering a  
5 patient ID on the touchscreen display. How does prior art  
6 teach this subject matter?

7 A Again, from the Klein thesis, depending on the  
8 chosen run type, the user is prompted for additional  
9 information such as the patient ID number, number two.

10 Q All right. And that excerpt is from page 65 of  
11 the Klein thesis. Sub-element 3 of claim 9 is entering a  
12 patient dose on the touchscreen display. Is a patient dose  
13 taught by the prior art?

14 A Continuing from that same location in Klein, it  
15 also displays that a person can enter the dose activity,  
16 number three.

17 Q All right. And the fourth element here in claim  
18 9 is entering a flow rate on the touchscreen display. Does  
19 the prior art teach this subject matter?

20 A Again, continuing from the same location, that  
21 flow rate and priming of the patient line, etc., number  
22 four as in the Klein thesis.

23 Q All right. So again, for the record, Dr. Stone  
24 is referring to an excerpt taken from page 64 of the Klein  
25 thesis.

1 All right. Let's move forward, please. We are  
2 at slide 192 and we are talking about claim 10 of the '870  
3 patent. Claim 10 refers to tracking a volume of saline  
4 remaining in the saline reservoir. Have we seen this  
5 element before?

6 A We have. The '869 patent claim 24 and my  
7 analysis remains the same.

8 Q All right. Let's cycle forward. Claim 10  
9 refers to providing an alert via the touchscreen display  
10 when the volume of saline remaining in the saline reservoir  
11 is below a predetermined volume threshold.

12 And actually let's move forward, please. Claim  
13 10 also refers to presenting on the touchscreen display a  
14 screen reminding the user to empty the waste bottle. These  
15 three elements that are shown here on slide 192 for claim  
16 10, where have we seen these elements described?

17 A Those are in the '869 patent claim 4 and my  
18 analysis remains identical.

19 Q All right. Let's move forward, please. We are  
20 at slide 193 and we're talking about claim 11. It says  
21 that the method further comprises initiating a generator  
22 column wash through the touchscreen display. Please  
23 explain what does -- how does Klein describe a generator  
24 column wash?

25 A Klein refers to that as a flush run that's

1 initiated from the user interface, and we see on the screen  
2 a sample screen from the Klein thesis that shows how a  
3 flush run is initiated.

4 Q All right. For the record, Dr. Stone is  
5 referring to page 64, another one of these screenshots  
6 taken from the client thesis. All right. So the flush  
7 run, is it initiated from the user interface?

8 A Yes.

9 Q All right. Let's move forward. Claim 11 says  
10 that a predetermined amount of saline is pumped through the  
11 strontium-rubidium radioisotope generator and directed to  
12 the waste bottle during the generator column wash. How  
13 does Klein describe operation of the generator column wash?

14 A From the Klein thesis, we have that the flush  
15 run flushes all the lines in the system as well as a 50  
16 milliliter flush of the generator at 15 milliliters per  
17 minute.

18 Q All right. And this excerpt that you're  
19 referring to is taken from page 53 of the Klein thesis; is  
20 that correct?

21 A That's correct.

22 Q Let's move forward, please. All right. We are  
23 at slide 194 and we are still talking about claim 11 from  
24 the '870 patent. And this new element refers to initiating  
25 a purging process through the touchscreen display to purge

1 a patient tubing line of air. How does Klein describe  
2 purging patient tubing lines of air?

3 A Well, once again, the same citation from Klein  
4 that the flush run flushing all of the lines in the system  
5 as well as 50 milliliter flush of the generator at 15  
6 milliliters per minute, which ensures flushing of air  
7 bubbles in the saline and strontium breakthrough from the  
8 generator.

9 Q And the claim also refers to the patient tubing  
10 line being in fluid communication with the eluate tubing  
11 line. Did you see that subject matter taught by the prior  
12 art?

13 A Yes.

14 Q All right. Let's move forward. We are at claim  
15 12 of the '870 patent on slide 196 of the presentation.  
16 The first element says that the saline tubing line and the  
17 eluate tubing line are routed through tubing passageways.

18 Let's move forward, please. Claim 12 also says  
19 that the tubing passageways have depths configured to  
20 prevent pinching or crushing of the tubing lines routed  
21 there through. Have we seen this subject matter before in  
22 your analysis?

23 A Yes. We saw those elements in claim 4 of the  
24 '869 patent and my analysis remains the same.

25 Q Let's move forward, please. We are at slide 198

1 and we are talking about claim 13 of the '870 patent which  
2 refers to a handle and also to four wheels. Have we seen  
3 this subject matter in your analysis already?

4 A My analysis of handles and wheels remains the  
5 same as in the '869 patent claim 4.

6 Q Claim?

7 A 24.

8 Q Claim 24. Thank you. Let's move forward,  
9 please. We are at slide 200. Talking about claim 16 of  
10 the '870 patent. The infusion system further comprises a  
11 dose calibrator in the shielded well on board the cart and  
12 in communication with the computer to determine the  
13 strontium breakthrough test result. Have we seen this  
14 subject matter before in your analysis?

15 A Yes, we have.

16 Q Where?

17 A It's in the '869 patent claim 27. My analysis  
18 remains the same.

19 Q Your analysis at claim 16 is the same as claim  
20 27; is that correct?

21 A That's correct.

22 Q Let's move forward, please. All right. Now we  
23 are at slide 202. We are talking about claim 17 of the  
24 '870 patent and you have a comparison here between claim 17  
25 and claim 3 of the '869 patent. Have we seen the elements

1 of claim 17 already in your analysis?

2 A We have. These were in the '869 patent claim 3.

3 Q All right. So let's just move forward. Would  
4 you cycle through this to show the Court how they  
5 correspond.

6 All right. And how does your analysis of claim  
7 17 of the '870 patent correspond to your analysis of claim  
8 3 from the '869 patent?

9 A My analysis of the '869 patent claim 3 remains  
10 the same and it's the same analysis for claim 17.

11 Q Let's move forward, please. All right. We are  
12 at slide 203 talking about claim 27 of the '870 patent.  
13 I'm not sure we've seen this before. This one says the  
14 computer of the infusion system is configured to track time  
15 passed from the completion of pumping the sample of  
16 rubidium radioactive eluate into the eluate reservoir to  
17 determining the strontium breakthrough test result.

18 Please explain, how does the prior art teach  
19 this subject matter?

20 A In Klein, we see here that the activity of the  
21 dose calibrator is registered 30 minutes after the end of  
22 the elution to compute the strontium-82 and strontium-85  
23 breakthrough activity.

24 Q All right. So this excerpt is taken from page  
25 28 of the RX-106. How much time does the computer track?



1           A           The computer tracks 30 minutes. The computer  
2 does the measurements. It does the tracking of time.

3           Q           All right. Let's move forward, please. All  
4 right. So we are at slide 204 and we get to shift to the  
5 '826 patent. All right. And we are at slide 205. Thank  
6 you.

7                       All right. So let's talk about claim 1. Claim  
8 1 talks about a method of building an infusion system to  
9 deliver a rubidium radioactive eluate. Does Klein teach  
10 how to build an infusion system to develop rubidium  
11 radioactive eluate?

12          A           He does.

13          Q           All right. So let's move into the elements.  
14 The first thing you've got to do is install a first  
15 shielding compartment, a second shielding compartment, and  
16 a shielded well on a platform of a cart. Walk us through  
17 where are the first, second shielding compartments and  
18 where is the shielded well that's relevant to this element?

19          A           Klein has first shield -- Klein has a first  
20 shielding compartment, a second shielding compartment. We  
21 talked about the obviousness of placing this shielded well  
22 for the dose calibrator on board the cart.

23          Q           And for the record, Dr. Stone is pointing to,  
24 again, this photograph taken from page 34 of the Klein  
25 thesis and to the red, green and blue annotation provided

1 on slide 205.

2 Okay. So claim 1 has a couple of wherein  
3 clauses. The first wherein clause is that the first  
4 shielding compartment has a first opening facing vertically  
5 upwardly. Have we encountered this before?

6 A We have. And we've explained that Klein has his  
7 shielding compartments with openings facing vertically up.

8 Q And we've also discussed that with respect to  
9 CardioGen and Tate; is that correct?

10 A That is correct.

11 Q The next part of the wherein clause is that the  
12 first opening is configured for a strontium-rubidium  
13 radioisotope generator to be inserted into and removed from  
14 the first shielding compartment. How does the prior art  
15 teach this subject matter?

16 A As we spoke previously, the generator was placed  
17 in the cart surrounded by lead rings to provide maximum  
18 radiation shielding. They had an opening facing vertically  
19 upward through which the strontium-rubidium generator could  
20 be inserted into and removed from that compartment.

21 Q All right. And the other prior art references  
22 that we've discussed, the Tate, the Medrad and CardioGen,  
23 did they have radiopharmaceuticals placed?

24 A They had radiopharmaceuticals. The source was  
25 placed in shielded wells with vertically opening -- with

1 vertical openings facing upward.

2 Q All right. Let's move forward, please. We are  
3 at slide 206. We are talking about the second shielding  
4 compartment which is color coded and was color coded in  
5 green in the prior slide. This one says also that that  
6 shielding compartment must have a second opening facing  
7 vertically upwardly. Have we discussed this in your  
8 analysis already?

9 A We have.

10 Q All right. Again, explain how does the prior  
11 art show this subject matter?

12 A We had a waste container mounted on the top  
13 shelf inside a lead container with a lid.

14 Q All right. Claim 1 also refers to the second  
15 opening being configured for a waste bottle this time to be  
16 inserted into and removed from the second shielding  
17 compartment. How does the prior art teach this?

18 A Well, again, we spoke of the waste container  
19 shielding compartment that Klein has.

20 Q And then, I guess, the last part of this wherein  
21 clause is that the first opening, that's the one for the  
22 generator shield, is located at a lower elevation than the  
23 second opening, which is the one for the waste bottle  
24 shield. Why is this subject matter obvious over the prior  
25 art?

1           A       As we've spoken before, it would be obvious to  
2 take the heavier objects that have to be handled by the  
3 user and place them at the lower elevation than that second  
4 opening that we dealt with.

5           Q       Let's move forward, please. Now we are at slide  
6 207 still talking about claim 1. This one says that the  
7 shielded well is configured to receive an eluate reservoir  
8 that is configured to receive a sample of the rubidium  
9 radioactive eluate. Again, walk us through your analysis  
10 of this piece.

11          A       Again, we have the calibrator chamber that's a  
12 shielded chamber located in a shielded well and inside is  
13 the vial, the eluate reservoir that is configured to  
14 receive a sample of the radioactive rubidium.

15          Q       So for the record, Dr. Stone is pointing to the  
16 excerpt at page 57 that we have here on slide 207 and to  
17 the vial coded in blue and to -- would you call it a  
18 doughnut assembly, the doughnut cylinder illustrated as the  
19 calibrator chamber. All right. Let's move forward,  
20 please.

21                   All right. We are on slide 208. We are talking  
22 about claim 1. And claim 1 starts -- this excerpt starts  
23 off with configuring a computer with a touchscreen display  
24 for the infusion system to do a bunch of stuff. Again,  
25 remind the Court, does the prior art teach a computer with

1 a touchscreen display?

2 A Yes. Klein's thesis described a computer with a  
3 touchscreen display.

4 Q All right. The first thing that the computer  
5 must configure the system to do is to fill the eluate  
6 reservoir in the shielded well on board the cart with the  
7 sample of rubidium radioactive eluate. Remind the Court,  
8 how does this happen?

9 A We've discussed this previously. The computer  
10 controls -- the computer controls the peristaltic pump  
11 which pumps saline through the generator to the eluate  
12 reservoir. The only difference here is that it's not on  
13 board the cart. And we talked about how it's obvious to  
14 relocate the dose calibrator and the eluate reservoir on  
15 board the cart that is taught by Tate and Medrad.

16 Q So for the benefit of the record, Dr. Stone is  
17 referring to the color-coded elements here in this excerpt  
18 from page 29 of the Klein thesis.

19 All right. You've got to fill the eluate  
20 reservoir by pumping saline from the saline reservoir into  
21 the strontium-rubidium radioisotope generator via a saline  
22 tubing line. So just to be absolutely clear, where is the  
23 saline tubing line?

24 A We have saline tubing lines going from here  
25 through the generator.

1           Q       Dr. Stone is referring to the pink color-coded  
2 lines extending from the saline IV to the generator.

3           A       That's correct. And that's controlled by the  
4 computer.

5           Q       It's got to thereby generate the rubidium  
6 radioactive eluate that is discharged through an eluate  
7 tubing line. So where is the rubidium radioactive eluate  
8 discharged from?

9           A       It's discharged from the discharge port through  
10 the eluate tubing line.

11          Q       All right. So for the record, Dr. Stone is  
12 pointing to the red generator and to the pink line that  
13 exits that generator and goes into the activity counter.

14                   All right. Let's move forward, please.  
15 Configuring the computer for the infusion system to  
16 determine a strontium breakthrough test result on the  
17 sample of the rubidium radioactive eluate built into the  
18 eluate reservoir in the shielded well on board the cart.  
19 Remind us, how did that occur in the prior art?

20          A       The only missing element here is on board the  
21 cart. We have -- in Tate, we have readings from the dose  
22 calibrator taken by the computer, which also computes the  
23 time and ultimately calculates the strontium breakthrough  
24 test result while the sample remains in the eluate  
25 reservoir in the shielded well.

1           Q       All right. So you stole a little bit of my  
2       thunder. But I want to make sure we've knocked out this  
3       while clause at the bottom. So it must occur while the  
4       eluate reservoir remains in the shielded well on board the  
5       cart. And again, tell me the while.

6           A       The while is the sample readings are taken by a  
7       dose calibrator while it's in the shielded well.

8           Q       Okay. And again, for the benefit of the record,  
9       we are on slide 209 and Dr. Stone is testifying about the  
10      blue color coded dose calibrator and the green color coded  
11      computer and the communication between them.

12                   Let's move forward, please. All right. So now  
13      we are at slide 210. Claim 1 says you've got to configure  
14      the computer for the infusion system to not allow a patient  
15      infusion if the strontium breakthrough test result is  
16      greater than or equal to an allowed limit. Again, remind  
17      us how does this occur?

18           A       The system ensures compliance with the daily  
19      protocol, which includes a calibration run and successful  
20      breakthrough in order to enable patient elution runs. The  
21      amount is based on the strontium breakthrough activity  
22      limited to the Health Canada guidelines, and only after a  
23      calibration run with low strontium breakthrough has been  
24      successfully completed can patient elutions be carried out.

25           Q       All right. Let's move forward, please. That

1 was slide 210. And now we are on slide 211 and we have  
2 claim 2 of the '826 patent. So claim 2 says we are  
3 configuring the computer to measure a radioactivity of the  
4 sample of the rubidium radioactive eluate while the sample  
5 is flowing through the eluate tubing line to the eluate  
6 tubing -- to the eluate reservoir. Excuse me. How does  
7 the computer measure this radioactivity?

8 A The computer measures it by being in  
9 communication with the activity counter which measures the  
10 activity while the eluate is flowing through the tubing  
11 line to the eluate reservoir.

12 Q All right. So we are on slide 211. And for the  
13 record, Dr. Stone is referring to the purple coded activity  
14 counter and the pink coded tubing lines.

15 Let's move forward, please. We are at slide  
16 212. This refers to configuring the computer to measure a  
17 calibration radioactivity of the sample while the sample  
18 remains in the eluate reservoir in the shielded well on  
19 board the cart. How does the computer measure a calibrator  
20 radioactivity?

21 A When the sample is in the eluate reservoir where  
22 it's first pumped in, the computer measures the activity,  
23 receives that activity from the dose calibrator. The only  
24 thing missing is on board the cart and we've shown how  
25 that's obvious to place that on board the cart.



1           Q       Let's move forward, please. All right. Claim 2  
2 further refers to configuring the computer to compare the  
3 radioactivity of the sample measured while flowing through  
4 the eluate tubing line and compare that value with the  
5 calibration radioactivity of the sample measured in the  
6 eluate reservoir in the shielded well on board the cart.

7                   So please explain how does this comparison occur  
8 in the prior art?

9           A       As we've discussed before, the integral activity  
10 is used to calibrate the activity counter and verify the  
11 calibration constant is within tolerance from the previous  
12 records. Those activities are taken while the sample is  
13 flowing through the dose -- the activity counter and the  
14 activity that's read when the sample vial is filled.

15           Q       So that's slide 213 and the excerpt that you're  
16 referring to is taken from page 53 of RX-106. Let's move  
17 forward, please.

18                   JUDGE CHENEY: That's probably a good place to  
19 take our morning break. We'll take 15 minutes. We are off  
20 the record.

21                   (Recess.)

22                   JUDGE CHENEY: Let's go back on the record. We  
23 are on the record in the 1110 investigation. Before the  
24 break we were in the direct examination of Dr. Stone the  
25 respondents' expert on the issues of patent validity.

1 Please continue when you're ready Mr. Hails.

2 MR. HAILS: Thank you, Your Honor.

3 BY MR. HAILS:

4 Q Before we broke we were on slide 215 of your  
5 presentation. This shows claim three of the '826 patent  
6 and the claim element refers to installing a dose  
7 calibrator in the shielded well on board the cart with the  
8 dose calibrator in communication with the computer to  
9 measure the strontium breakthrough test result. Have we  
10 seen this part of this claim earlier in your analysis?

11 A Yes. We saw that in the '869 patent claim 27.

12 Q And has your analysis of claim three of the '862  
13 patent correspond with claim 27 of the '869 patent?

14 A It remains the same.

15 Q There is a second piece of this claim that says  
16 the dose calibrator is not only to measure the  
17 strontium-rubidium test result. It is also to measure the  
18 calibration radioactivity of the sample pumped into the  
19 eluate reservoir. Please explain how does this process  
20 occur in the prior art?

21 A As we've disclosed previously, the calibration  
22 radioactivity at the sample pump into the eluate reservoir  
23 is measure by the dose calibrator as soon as the saline has  
24 been pumped into the dose reservoir and is used for that  
25 calibration factor.

1 Q And, again, what is the system calibrating?

2 A It's calibrating the activity counter on board  
3 the cart.

4 Q Let the record reflect Dr. Stone is pointing,  
5 has pointed to the blue dose calibrator color coded on  
6 slide 215 and the communication connectivity between that  
7 element and the computer shown in green. All right. Let's  
8 move forward please all right. We just talked about a dose  
9 calibration radioactivity, so why don't we move on from  
10 slide 216. All right. Now we are at slide 218. Talking  
11 about claim five of the '826 patent. This claim says  
12 configuring the computer to allow a user to initiate a  
13 purging process through the touch screen display to purge a  
14 patient tubing line of air and that the tubing line is in  
15 fluid communication with the eluate tubing line. Have we  
16 seen this subject matter before in your analysis?

17 A Yes. We described our coverage of that in claim  
18 11 of the '870 patent. My analysis remains the same.

19 Q So your analysis of claim five is the same as  
20 your analysis of claim 11 of the '870, is that correct?

21 A That's correct.

22 Q Let's move forward, please. All right. Now we  
23 are on slide 225 and we are talking about claim nine of the  
24 '826 patent and you have a comparison here between claim  
25 nine and claim eight of the '870 patent. This is the one

1 that talks about starting the patient infusion by touching  
2 a button on the touch screen display --

3 And why don't you cycle through these elements.

4 How do the --

5 Keep going, please. And one more.

6 So how do the elements of claim nine from the  
7 '826 patent compare to the elements of claim eight from the  
8 '870 patent?

9 A Those are the same elements and my analysis of  
10 claim eight, or sorry, of claim nine of the '826 is  
11 identical to my analysis of claim nine of the '870.

12 Q You said claim nine of the '870 but just for the  
13 record...

14 A Sorry. Claim eight of the '870.

15 Q That's all right. This is a toughy for us both.  
16 Let's keep going. We are on slide 227 talking about claim  
17 10 from the '826 patent. Have we seen the subject matter  
18 of claim 10 already in your analysis?

19 A We have. Cycling through each of these  
20 elements, these elements are identical to the elements of  
21 claim nine of the '870 patent so my analysis of claim 10 of  
22 the '826 patent is identical to my analysis of claim nine  
23 of the '870 patent.

24 Q Thank you. That was slide 227 of your  
25 presentation.

1           Can we move forward, please. All right. We are  
2 at claim 11 of the '826 patent on slide 228.

3           This says configuring the computer to track time  
4 passed from completion of pumping the sample of the  
5 rubidium radioactive eluate into the eluate reservoir to  
6 measuring the strontium breakthrough test result. Remind  
7 the court, how does the computer track time between these  
8 events?

9           A       Indeed, as Klein states, the activity in the  
10 dose calibrator is registered 30 minutes after the end of  
11 the elution to compute the breakthrough of strontium-82 and  
12 strontium-85 activity. Those measurements are done by the  
13 computer and the tracking of the time is done by the  
14 computer.

15          Q       Okay. And so this excerpt on slide 228 is taken  
16 from page 28 of RX-106. Again, how much time is tracked by  
17 the computer?

18          A       30 minutes is tracked by the computer.

19          Q       Let's move forward, please.

20                 Claim 11 also says you got to track a volume of  
21 saline remaining in the saline reservoir --

22                 And will you advance, please.

23                 You also have to provide an alert via the touch  
24 screen display when the volume of saline in the saline  
25 reservoir is below a predetermined volume threshold. Have

1 we seen this subject matter before in your analysis?

2 A We have in the '869 patent at claim 14. Those  
3 elements are identical. And so my analysis of claim 11 of  
4 the '826 is identical to my analysis of those elements in  
5 claim 14 of the '869 patent.

6 Q Or at least with respect to these two newly  
7 added elements, correct?

8 A With those elements, yes.

9 Q Let's move forward.

10 All right. So that was slide 232 and now we are  
11 at slide 233 talking about the final two elements of claim  
12 11 tracking a volume of the rubidium radioactive eluate  
13 discharge from the generator to the waste bottle. Thank  
14 you. And presenting the touch screen, on the touch screen  
15 display a screen reminding the user to empty the waste  
16 bottle. Have we seen this subject matter before in your  
17 analysis?

18 A Yes. My analysis of those elements in claim 24  
19 of the '869 patent are identical to these elements in claim  
20 11 of the '826 patent.

21 Q All right. So let's move forward, please.

22 All right. So now we are at slide 237. We are  
23 talking about claim 12 from the '826 patent. Let's cycle  
24 through the elements of claim 12. Have we seen the  
25 elements of claim 12 already in your analysis?

1           A       Yes.  Those elements were present in claim 11 of  
2 the '870 patent.

3           Q       Okay.  And how does your analysis of claim 12  
4 for the '826 compare to your analysis from claim 11 of the  
5 '870?

6           A       My analysis of claim 12 of the '826 remain  
7 identical to the analysis of those elements in claim 11 of  
8 the '870 patent.

9           Q       So that's slide 237.  Let's move forward,  
10 please.

11                    Claim 13 of the '826.  I think we are talking  
12 about tubing passageways again and perimeter surfaces of  
13 openings and pinching and crushing of tubing lines.  Have  
14 we seen this subject matter of claim 13 before in your  
15 analysis?

16           A       Yes.  We saw those elements in claim 4 of the  
17 '869 patent.  My analysis of these elements in claim 13 of  
18 the '826 remains identical to my analysis of those elements  
19 in claim four of the '869 patent.

20           Q       Okay.  So that's slide 240.  Why don't we move  
21 forward.  So now we are at slide 241 and we are going to  
22 talk about claim 14 of the '826.  The first element is an  
23 exterior shell extending upwardly above the platform  
24 wherein the platform and the exterior shell collectively  
25 define an interior space of a cabinet structure.  Why don't

1 you just walk us through this. Have we seen this subject  
2 matter before?

3 A We have indeed. Here we have the Klein -- here  
4 we have the Klein thesis with a platform and exterior shell  
5 on the platform.

6 Q And an anterior space, do we have an anterior  
7 space?

8 A We have an anterior space.

9 Q All right.

10 A Similarly we have those elements in the Tate  
11 disclosure, as well as in the CardioGen Model 510.

12 Q Let the record reflect that with respect to his  
13 analysis of Klein he was pointing to the structures  
14 illustrated in this picture taken from page 34 of the Klein  
15 reference. All right. Let's move forward.

16 Handles and wheels. Have we seen this subject  
17 matter before in your analysis?

18 A We covered this subject matter in the '870  
19 patent claim 13 and my analysis of claim 14 of the '826  
20 where those elements remains identical.

21 Q All right. So that's slide 244. Let's move  
22 forward, please.

23 So now we are at slide 246. Talking about claim  
24 17 of the '826 patent which refers to a dose calibrator in  
25 the shielded well on board the cart. That's in



1 communication with the computer to measure the strontium  
2 breakthrough test result. Have we seen this subject matter  
3 before in your analysis?

4 A We have indeed. We've analyzed that as claim 27  
5 of the '869 patent and that element, my analysis remains  
6 the same for that element of claim 17 of the '826 patent.

7 Q All right. You said of claim -- which -- your  
8 analysis of claim 17 is identical to your analysis of which  
9 claim for the '869 patent?

10 A Claim 27.

11 Q All right. Let's move forward, please.

12 That was slide 246 and now we are up to 248.  
13 Claim 18. Have we seen the subject matter of claim 18  
14 before in your analysis?

15 A Yes. Each of these elements we saw previously  
16 in claim three of the '869 patent. Those elements in claim  
17 18 are analyzed in my analysis remains identical to my  
18 analysis of the claim three of the '869 patent.

19 Q So just to be clear, your analysis of claim 18  
20 for the '826 is identical to the analysis of claim three of  
21 the '869. Is that what you meant?

22 A That is correct.

23 Q Let's move forward, please.

24 All right. Now we are at slide 249. This claim  
25 is claim 19 from the '826 patent. It requires configuring

1 the computer to do a bunch of things. The first thing is  
2 to control a fluid communication between the  
3 strontium-rubidium radioisotope generator and the saline  
4 reservoir. Please walk us through how does the prior art  
5 teach this subject matter?

6 A The computer on the Klein thesis controls the  
7 peristaltic pump which controls fluid communication between  
8 the saline reservoir. It also controls the generator valve  
9 to control fluid communication to the inlet of the  
10 strontium-rubidium generator.

11 Q All right. For the record, Dr. Stone is  
12 pointing to this excerpt taken from page 29 of Klein  
13 referring to the blue color-coded peristaltic pump, the  
14 orange color-coded generator valve and the connections  
15 between the saline IV and the strontium-rubidium  
16 radioisotope in red. Let's move forward.

17 Claim 19 says you got to configure the computer  
18 to control a fluid communication between the eluate tubing  
19 line and the eluate reservoir. How does the computer do  
20 this?

21 A Well, the hot pink eluate tubing line is  
22 configured to the fluid communication with the eluate  
23 reservoir by the patient valve which is controlled by the  
24 computer.

25 Q And just for the record, Dr. Stone is pointing

1 to the picture taken from page 29, the orange color-coded  
2 patient valve. All right. Let's move forward, please.

3 The next piece of claim 29 is the computer is  
4 configured to control a fluid communication between the  
5 eluate tubing line and the waste bottle. Let's talk about  
6 how this occurs.

7 A The eluate tubing line shown in pink is in fluid  
8 communication with the waste bottle by the positioning of  
9 the patient valve controlled by the computer.

10 Q All right. Let's move forward, please.

11 All right. There is two elements in claim 29  
12 that are compared to '870, placing the eluate tubing line  
13 in fluid communication with a patient, pumping a dose of  
14 the rubidium radioactive eluate to the patient. Have we  
15 seen this subject matter before?

16 A We have. In the '870 patent claim two we saw  
17 those two elements and my analysis for claim 19 of -- those  
18 two elements of claim 19 of the '826 is identical to my  
19 analysis of the -- those two elements in the '870 patent,  
20 claim two.

21 Q All right. And then let's move forward, please.

22 Claim 19 also says you got to flush the rubidium  
23 radioactive eluate remaining in a portion of the eluate  
24 tubing line to the patient using this bypass line. Have we  
25 seen this subject matter before?

1           A        Yes. We saw that element in claim two of the  
2 '870 patent and so my analysis of this element for claim 19  
3 of the '826 is identical to my analysis for claim two of  
4 that element. That element of claim two in the '870  
5 patent.

6           Q        Let's move forward, please.

7                    All right. We are at slide 258 and we are  
8 talking about claim 28 from the '826 patent. Talks about  
9 initiating a column wash through the touch screen display.  
10 Also talks about a predetermined amount of saline being  
11 pumped through the generator and directed to the waste  
12 bottle during the generator column wash. Have we seen this  
13 subject matter before?

14          A        We have. We covered those two elements in the  
15 '870 patent claim 11 so my analysis of these two elements  
16 in claim 28 of the '826 is identical to my analysis of  
17 those elements in claim 11 of the '870 patent.

18          Q        All right. Let's move forward, please.

19                    All right, Your Honor, I'd like to run through  
20 the Medrad documents, please.

21                    JUDGE CHENEY: Okay. So there has been some  
22 discussion about what is public about the Medrad prior art,  
23 including dates of sale and things like that. What exactly  
24 are you going to be getting into now that has a third-party  
25 assertion of confidentiality?

1 MR. HAILS: So to my understanding, all of these  
2 documents have third-party claims of confidentiality and I  
3 recognize they are untested. We are going to go through  
4 the sales documents and documents showing dates of  
5 installation and location of installation and we are going  
6 to go through user manuals to essentially confirm the  
7 analysis that Dr. Stone has provided earlier, those yellow  
8 redaction bubbles.

9 JUDGE CHENEY: Are you going to call any  
10 witnesses about these documents?

11 MR. HAILS: We have a declaration from a Medrad,  
12 I guess a Bayer person, attesting to these documents that's  
13 included.

█ [REDACTED]

█ [REDACTED]

█ [REDACTED]

█ [REDACTED]

█ [REDACTED]

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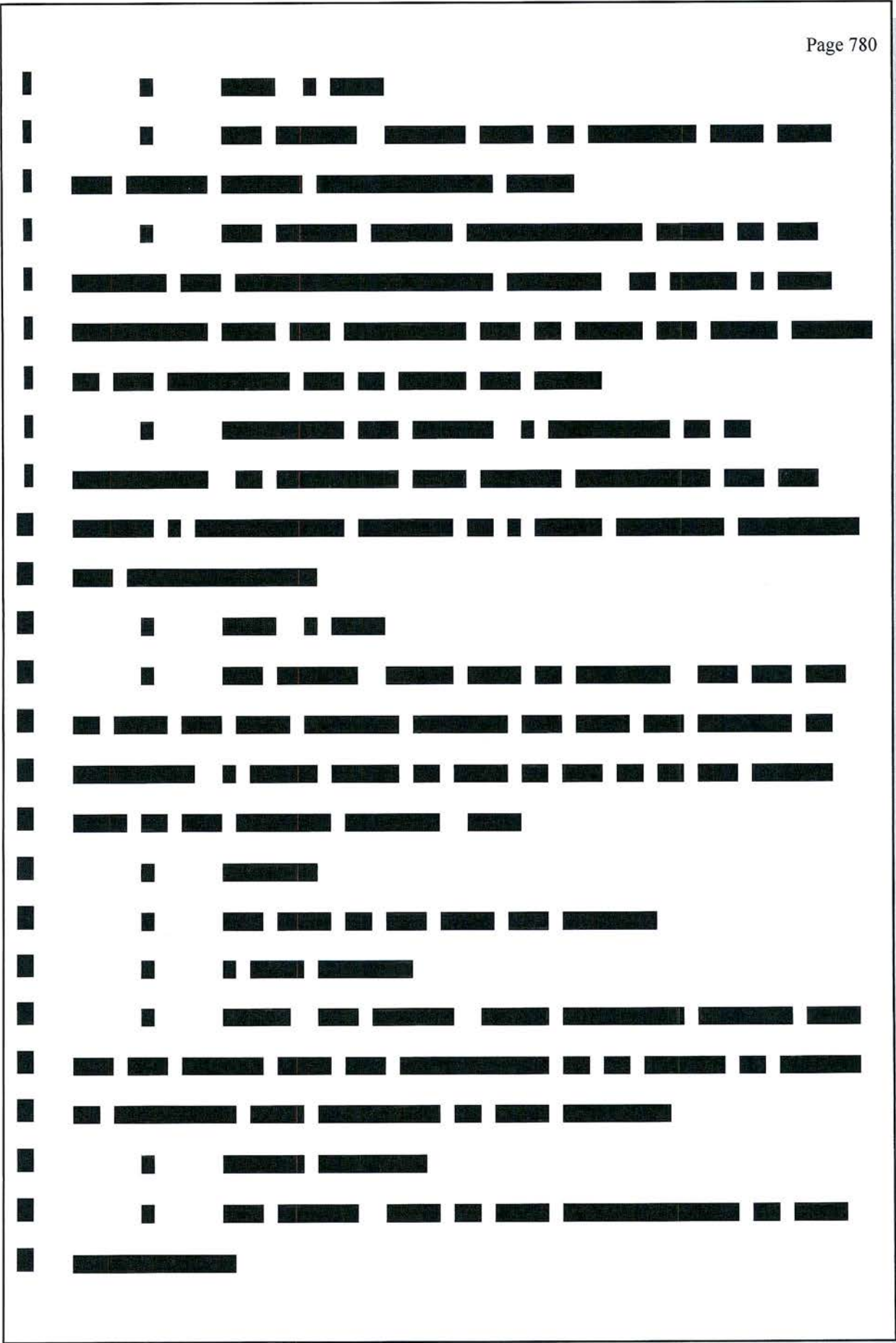
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[REDACTED] Your Honor, we can go back  
6 to the public record. Thank you.

7 JUDGE CHENEY: We are back on the public record.

8 (End of confidential session.)

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1 OPEN SESSION CONTINUED

2 BY MR. HAILS:

3 Q Okay. Now let's talk about your anticipation  
4 opinions. Can we move forward, please. And let's start  
5 with slide 69. So earlier, I had asked you if you believed  
6 that the June 2009 date that you had used for priority for  
7 your obvious analysis, whether that was the proper priority  
8 date. Do you remember that?

9 A I do.

10 Q Okay. And what was your answer?

11 A I don't believe that that's the proper priority  
12 date.

13 Q Okay. And what priority date do you believe is  
14 proper under the circumstances of these patents?

15 A I believe no earlier than March of 2017 is the  
16 proper priority date to use for these patents.

17 Q Okay. Let's say it was March 2017. What does  
18 that mean for the status of the RUBY product, particularly  
19 the user manual as against these claims?

20 A It means that the RUBY product was priority, is  
21 prior art for those patents and invalidates them.

22 Q Okay. Why do you think that the priority date  
23 does not extend backwards to June 2009?

24 A Because there is no evidence that those  
25 inventors had possession of the claims that are in those

1 patents.

2 Q All right. Let's move forward, please.

3 We are jumping to slide 35 of your presentation.

4 And here you have another timeline. Will you walk us

5 through this material.

6 A All right. First of all, in 2008 we have four

7 patents that were filed by Bracco and its

8 contractor/employees. And in 2009 we have the June of 2009

9 we have the PCT application which enters the United States

10 which for the first time claims an on board -- sorry, it

11 doesn't claim, it discloses an on-board dose calibrator and

12 then we have the U.S. patent for that in 2010.

13 Q Okay. And so those are not the asserted

14 patents, is that correct?

15 A Those are not the asserted patents.

16 Q All right. So when were the asserted patents

17 filed?

18 A In 2016 after Bracco obtained the user manual

19 for the FDA approved RUBY-FILL product, the patents were --

20 the '869 was filed and then in 2017 the '826 and the '870

21 patents.

22 JUDGE CHENEY: I'm sorry to interrupt.

23 Dr. Stone, is your microphone on?

24 THE WITNESS: I'm sorry. Yes, Your Honor.

25 JUDGE CHENEY: You can move it like this.

1                   THE WITNESS: I have some water here that I  
2 spilled, Your Honor. I'm good. Thank you. Paper towels  
3 would be nice.

4                   JUDGE CHENEY: Will someone please assist  
5 Dr. Stone. There is no need to suffer in silence. We have  
6 so many resources available to us.

7                   THE WITNESS: Thank you. I'm prepared, Your  
8 Honor.

9                   JUDGE CHENEY: Okay. Thanks.

10 BY MR. HAILS:

11           Q       All right. So let's -- let's talk about  
12 foundation. Do you have a binder that contains the  
13 materials that form the foundation of your opinions on this  
14 priority issue?

15           A       Yes.

16           Q       All right. Can you pull it out for me. All  
17 right. Just administrative note, rather than provide the  
18 entireties of prosecution histories we have excerpts for  
19 certain exhibits but let me run through this list and see  
20 in my list is the same as yours.

21                   Ready?

22           A       Yes.

23           Q       First of all, JX-1, 2 and 3, those are the  
24 asserted patents?

25           A       Yes.



1 Q JX-4, pages 18 through 25?  
2 A Yes.  
3 Q JX-5 and JX-6?  
4 A Yes.  
5 Q All right. JX-26, pages 1535 through 1543?  
6 A Yes.  
7 Q JX-63, pages 2105 through 2110.  
8 A Yes.  
9 Q All right. And then a couple excerpts from  
10 JX-64, page 2347. Also page 2, 351 through 2538.  
11 A Yes.  
12 Q All right. Two excerpts from JX-65, the first  
13 one from 1498 through 1525.  
14 A Yes.  
15 Q The second one from pages 1738 through 1747.  
16 A Yes.  
17 Q I think this is three excerpts from CX-169.  
18 Pages 21 and 24.  
19 A Yes.  
20 Q 2940?  
21 A Yes.  
22 Q 2948 through 2958?  
23 A Yes.  
24 Q All right. And RX-373?  
25 A Yes.

1 Q All right. So those documents form the  
2 foundation of your opinions, is that correct?

3 A Yes. They do.

4 Q All right. All right. So we've talked about  
5 the filing of the asserted patents. The asserted patents  
6 were filed in 2016 and 2017 respectively, is that right?

7 A That is correct.

8 Q All right. And what was the trigger event for  
9 the filing of these applications?

10 A The triggering event was when Bracco obtained  
11 the user manual for the RUBY-FILL Version 3.

12 Q All right. Let's move forward, please.

13 All right. And that is what you've shown here.  
14 Let's move forward, please. Okay. So let's start back in  
15 2008. Why don't you walk us through what's the same and  
16 what's different about these 2008 filings.

17 A What's the same about these 2008 filings is the  
18 disclosure. What is different are two factors. There are  
19 different inventors for different claim elements that are  
20 asserted. The tubing circuits we have one set of  
21 inventors. The shielding assembly is still another set of  
22 inventors, at least, we add two more inventors there. The  
23 cabinet structure still a slightly different set of  
24 inventors and then finally we have computer controls which  
25 has a very different set of inventors.

1 Q Okay. So all of these patents were filed as  
2 applications on the same day, is that correct?

3 A That's my understanding.

4 Q Okay. And did you say that they all had the  
5 same disclosure?

6 A Yes.

7 Q Okay. But you said that the claims were  
8 different. Again, just walk us through how are the claims  
9 different among these four cases?

10 A The '534 relates to tubing circuits. The '674  
11 relates to shielding assemblies. The '352 claims relate to  
12 cabinet structure and the '053 patent relates to computer  
13 controls of the process.

14 Q Okay. And the different patents have different  
15 sets of inventors, is that correct?

16 A That is correct.

17 Q All right. Let's move forward, please.

18 Okay. So now we are on slide 52 and you have  
19 the PCT application shown which was filed in 2009. Why  
20 don't you explain to the Court what is the claim and who  
21 are the named inventors on this?

22 A Well, this disclosure for the first time adds  
23 computer controls. It's very similar disclosure with a  
24 couple of lines added with regard to computer control and  
25 it has the same inventors, most of the same inventors that

1 were on the '053 but it adds a couple.

2 Q All right. Let's move forward, please.

3 So now we are at slide 59. Here you have  
4 comparison of the inventorship, is that correct?

5 A That's correct.

6 Q Why don't you just summarize for the Court when  
7 you're talking about claims to computer controls, what kind  
8 of subject matter do you see in these claims?

9 A Well, they claimed pumping an eluate through a  
10 generator could generate an eluate. They claimed providing  
11 indication that the elution is completed, providing  
12 indication of time lapse since completion, since the  
13 elution completion was completed. But there are no claim  
14 elements that are drawn to shielding structures or cabinet  
15 structures in either of those two patents.

16 Q All right. So just to make sure the record is  
17 clear, the '053 element doesn't have any claim elements  
18 drawn to shielding structures?

19 A No.

20 Q And does the '053 have any claims directed to  
21 the elements of the cabinet structure?

22 A No.

23 Q The PCT application, when it was filed did it  
24 have any claim elements drawn to shielding structures?

25 A No.

1 Q And did the PCT application when it was filed  
2 have any claims directed to elements of a cabinet  
3 structure?

4 A No. It did not.

5 Q All right. Let's move forward.

6 All right. Slide 53. You have the asserted  
7 claims and we've been through them. What kinds of claim  
8 elements do we see in the asserted claims?

9 A We see not only computer controls but now we see  
10 also claim elements directed towards shielding assemblies  
11 and to cart configurations.

12 Q All right. Let's move forward, please.

13 What do we see by way of inventorship for the  
14 asserted patents?

15 A We see inventors claimed who were the inventors  
16 for the PCT filing.

17 Q All right. Let's move forward, please.

18 So now we are at slide 54. And you have the  
19 '674 patent shown here on the left. Why don't you give us  
20 a sense of what kinds of claim elements do we see for the  
21 shielding assembly claims from the '674 patent?

22 A Well, a couple of key elements are that they  
23 talk about openings for a generator compartment and a waste  
24 bottle compartment that are being oriented upward, openings  
25 for a second waste bottle and a compartment at a second

1 elevation being greater than the first elevation of the  
2 opening.

3 Q Okay. Let's move forward.

4 Okay. So the '674 patent when it was filed in  
5 2008 and it had these claim elements, did it identify any  
6 of the people who are identified as named inventors on the  
7 asserted claims as inventors for the '674 application?

8 A Now, the inventors of those elements in the  
9 asserted claims were -- the only ones that had those  
10 elements previously were from the '674 patent and that was  
11 Charles Quirico, Ernest Balestracii, Daniel Dorst, Eric  
12 Krause, Vishal Lokhande, Jacob Childs, Peter Madson, Daniel  
13 Clements. It does not include the -- none of those were  
14 included in the '674 patent that are on the later patent,  
15 Stephen Hidem, Aaron Fontaine, Janet Gelbach, Patrick  
16 McDonald, Kathryn Hunter, Rolf Swenson, or Jules Szoda.

17 Q Let's move forward, please.

18 So now we are at slide 57. Do we see claim  
19 elements that were filed in the patent in 2008 in the  
20 asserted claims that were filed in the 2016 -2017 time  
21 frame?

22 A Yes. We do. These new patents have some of the  
23 same claim elements. They have a shielding compartment  
24 that's opening facing vertically upwardly. Again, second  
25 shielding compartment with an opening facing vertically

1 upward and different locations. The first opening is  
2 located at a lower elevation than the second opening.

3 Q Okay. Let's move forward.

4 Again, any overlap between the inventorship  
5 between those sets of patent filings?

6 A Between those claim elements, no.

7 Q Okay. Let's move forward, please.

8 So now we are at slide 62 of your analysis. In  
9 this investigation was Bracco requested to provide  
10 information on contribution of the various inventors of  
11 these patent filings?

12 A Yes.

13 Q And what information do they provide on the  
14 inventive contributions of the people who are identified as  
15 inventors on the asserted patents?

16 A They had no information what these individuals  
17 may have contributed to the claimed invention.

18 Q Okay. So did Bracco have any information on  
19 what Mr. Hidem provided?

20 A No.

21 Q Did they have any information on what  
22 Mr. Fontaine provided?

23 A No.

24 Q Mr. McDonald?

25 A No.

1 Q Ms. Hunter?

2 A No.

3 Q Any information on what Mr. Swenson may have  
4 provided?

5 A No.

6 Q What about Dr. Zodda?

7 A None.

8 Q But you have an exception here for Janet  
9 Gelbach. What information did Bracco provide on her  
10 contributions?

11 A Mr. LaVanway thought she might have proposed the  
12 idea of an on-board dose calibrator.

13 Q What is Mr. LaVanway?

14 A Mr. LaVanway is the patent attorney who wrote  
15 these claims and filed the patent.

16 Q Let's move forward, please.

17 Okay. So now we are on slide 63. Was Bracco  
18 asked to provide information on inventive contributions  
19 that may have been made by the named inventors of the '674  
20 patent?

21 A Yes. They were.

22 Q And what did they identify as the contributions  
23 made by these individuals?

24 A They had no information on what they may have  
25 contributed to the claimed invention.



1 Q So does Bracco have any information on what  
2 Mr. Quirico may have done, Mr. Balestracii may have done,  
3 Mr. Dorst may have done, what Mr. Krause may have done,  
4 what Mr. Lokhande may have done, what Mr. Childs may have  
5 done, what Mr. Madson may have done or what Mr. Clements  
6 may have done?

7 A No.

8 Q Okay. Earlier you said that Bracco had gotten a  
9 copy of JDI's product literature and drafted the asserted  
10 claims. Are you familiar with that?

11 A Yes.

12 Q Okay. Was Bracco asked if they targeted JDI  
13 specifically with those asserted claims?

14 A Yes.

15 Q And what did they say?

16 A Yes.

17 Q And who was the person who was testifying on  
18 such issues?

19 A Mr. LaVanway.

20 Q Who was the person who drafted the asserted  
21 claims for filing at the Patent Office?

22 A Mr. LaVanway.

23 Q Okay. And who actually filed them at the Patent  
24 Office?

25 A It's my understanding it was Mr. LaVanway.

1 Q And who testified he had no information  
2 regarding any inventive contributions made by the named  
3 inventors on the asserted patents?

4 A Mr. LaVanway.

5 Q Let's move forward, please.

6 We are at slide 68. So let's say you file a  
7 patent application and you claim priority back to a prior  
8 application and you have no idea what the named inventors  
9 on your new filing may have contributed to the subject  
10 matter that you've claimed. Do you have an understanding  
11 of what that affect is for priority date?

12 A It means you really can't claim that as a  
13 priority date.

14 Q Okay. And so is priority date, are priority  
15 date claims effective in that circumstance?

16 A No. They are not.

17 Q Okay. And if a priority date made by the  
18 asserted patents in this case were ineffective then what is  
19 the priority date that should be assigned to those patents?

20 A Priority date is when the claimed inventors  
21 actually sign a declaration that they were in possession of  
22 the invention.

23 Q Okay. The date that you've ascribed, does that  
24 predate or does that postdate Bracco's acquisition of the  
25 RUBY-FILL product materials that they removed?

1           A       It postdates it.

2           Q       And you're aware of the analysis that's been  
3 performed in this investigation already, the final  
4 determination that the asserted claims read on the RUBY  
5 products?

6           A       Yes.

7           Q       Okay. So are you aware that it's been decided  
8 already that there is element-for-element correspondence  
9 between the asserted claims and the RUBY product?

10          A       I have.

11          Q       And if RUBY is prior art to the asserted claims,  
12 what does that mean?

13          A       It invalidates these patents.

14          Q       Thank you. Let me --

15                   MR. HAILS: Your Honor, thank you. We pass the  
16 witness.

17                   JUDGE CHENEY: Is there any cross-examination of  
18 Dr. Stone?

19                   MR. DAVIS: Yes, Your Honor. If we could take a  
20 minute to pass out the binders.

21                   JUDGE CHENEY: While the binders are being  
22 passed out I'm concerned about completing this hearing on  
23 time. So it seems like time estimates are getting blown  
24 through.

25                   MR. WALKER: Given where we are, I think I'd

1 probably go ahead and provide -- I think we are going to go  
2 ahead and probably drop Dr. Clark. Mr. Clark. Forgive me.  
3 So I just want to let the other side know that, given where  
4 we are right now.

5 JUDGE CHENEY: Okay.

6 MR. DAVIS: Your Honor, we believe we can still  
7 meet our time constraints.

8 JUDGE CHENEY: Okay. Please proceed when you're  
9 ready, Mr. Davis.

10 CROSS-EXAMINATION

11 BY MR. DAVIS:

12 Q Thank you, Your Honor. Good morning still. I'm  
13 Mark Davis. I'm one of the attorneys who represents the  
14 complainant Bracco.

15 A Good morning.

16 Q Now, in opining that the RUBY-FILL Version 3  
17 anticipates the claims of the asserted patents, you rely on  
18 a priority date of no earlier than September 30th, 2016,  
19 correct?

20 A That is correct.

21 Q Okay. But you can see that the specification of  
22 the parent '031 application filed on June 11th, 2009,  
23 discloses, among other things, a dose calibrator, an eluate  
24 reservoir and shielded well, right?

25 A I concede that.

1 Q Okay. And the '031 application is a parent to  
2 the asserted patent, correct?

3 A I'm sorry. You'll have to show me which one the  
4 '031 is. I don't keep them all memorized.

5 Q Okay. So why don't we pull up JX-1 and the  
6 related U.S. application data, page 2 of the patent. So do  
7 you see here where it mentions the continuation of  
8 application 12/808,467 filed as application number  
9 PCT/US2009/047031 on June 11th, 2009, now patent number  
10 9,607,722?

11 A Thank you. I recognize that now. Now what was  
12 your question.

13 Q Yes. So that's the parent application to the  
14 asserted patents, correct? That's a parent to not only  
15 JX-1, but JX-2 and JX-3?

16 A The '031 and the '722. Yes.

17 Q Okay. And the disclosure of the parent '031  
18 application, that disclosures are repeated in the  
19 specification of the asserted patents, correct?

20 A Yes.

21 Q Okay. And you did not offer an opinion that any  
22 of the asserted claims failed to satisfy the written  
23 description requirement, correct?

24 A I did not offer such an opinion. That's  
25 correct.

1 Q Okay. And you didn't offer an opinion that any  
2 of the asserted claims are not enabled by the disclosure of  
3 the '031 application?

4 A I did not discuss any of them.

5 Q And now in your initial report, you have a  
6 section regarding the applicable law regarding written  
7 description and priority dates, correct?

8 A I believe that's correct.

9 Q So that's page 21 of your initial report. And  
10 at paragraph 51, you state I understand that all patent  
11 applications must contain a written description of the  
12 invention claimed by the application. The written  
13 description requirement has two primary elements, the  
14 specification must not only describe the subject matter  
15 claimed by the patent, it must, it also must describe the  
16 claimed subject matter in a manner that demonstrates the  
17 attorneys had possession of the claimed subject matter.  
18 Possession of the invention is demonstrated when the  
19 subject matter of the invention is described in a manner  
20 that conveys to the reader that the inventors recognized  
21 the claimed subject matter of their invention. So that's  
22 what you were instructed with regard to your written  
23 description, correct?

24 A I see that.

25 Q Okay. Now let's go to paragraph 52. Now, in

1 523 of your report, you state that I understand that, to  
2 benefit from priority of an earlier-filed application, a  
3 patent must claim priority to the earlier-claimed  
4 application and that earlier-filed application must satisfy  
5 the written description requirement. Correct?

6 A I see that.

7 Q Okay. Now, you agree, we just showed that the  
8 patents in suit all claim priority back to the '031  
9 application, correct?

10 A I see that.

11 Q All right. And in your deposition, you admitted  
12 that if priority of the claims is based on what is  
13 disclosed in the specification then the priority date is no  
14 earlier than 2009, as opposed to 2016, correct?

15 A I didn't say it was as opposed to. I believe I  
16 said that it was no earlier than 2009.

17 Q Okay. So if you base priority on what's  
18 disclosed in the specification then you have a 2009  
19 priority date?

20 A If I'm basing my analysis on obviousness, I used  
21 2009.

22 Q Okay. Well let's look at your deposition. This  
23 is page 187, line 19 to 188, line 1. All right. And you  
24 say based on inventorship and when it was filed, it can be  
25 no earlier than 2007. Based on the disclosure when the

1 material was first added, it could be no earlier than 2009.

2 Correct?

3 A That's what I said.

4 Q Okay. So the written description requirement  
5 and the priority -- I'm sorry. Let me restate that  
6 question. So you indicated that your understanding of  
7 priority law was that you needed to claim priority and you  
8 needed to disclose the subject matter in the specification,  
9 and you admit that the subject matter claimed in the  
10 asserted patents is disclosed in the specification,  
11 correct?

12 A Again, with regard to obviousness, I utilized  
13 the date of June 2009. That's not the only thing with  
14 regard to priority that I did an analysis on. I also did  
15 an analysis based on inventorship.

16 Q Okay. So my question was, was the subject  
17 matter of the claim disclosed in the specification?

18 A As we've acknowledged the subject matter, the  
19 idea of the -- the idea was disclosed but not claimed  
20 previously.

21 Q Okay. Let's turn to inventorship. You read the  
22 transcript of Janet Gelbach's deposition, correct?

23 A I did.

24 Q And she testified that she has no reason to  
25 believe the incorrect inventors are listed on the asserted



1 patents, correct?

2 A She may very well have.

3 Q Okay. Well, let's pull it up. It's JX-176 at  
4 page 85. Now, Exhibits 4, 5, 6 and 7 we are talking about  
5 the inventorship of various patents. Do you have any  
6 reason to believe that the incorrect inventors are listed  
7 on any of those?

8 A No.

9 Q So she didn't think there was a problem with  
10 inventorship for the asserted patents, correct?

11 A She did not.

12 Q And you read her testimony that she was one of  
13 the people who conceived of putting the dose calibrator on  
14 board the cart, correct?

15 A I believe she said that it was a joint decision,  
16 but she knew of a dose calibrator being on a cart at the  
17 time of this disclosure.

18 Q Okay. And she said that she was part of the  
19 team that contributed that idea and that contribution was  
20 back at the time when she was in Bracco working on the new  
21 design in this 2006 to 2009 time period, right?

22 A She indeed brought that idea to the design team  
23 but at the time they didn't claim it as a patent and I  
24 don't think they recognized it as such. I don't think they  
25 -- it was already in commerce at that time.

1           Q       Okay. Well, let's look at your deposition.  
2       140, line 23 to 141, 25. I think admitted a little bit  
3       more forcefully that Janet Gelbach contributed to the  
4       on-board dose calibrator. Do you see where the question  
5       is? So it seems as though we've established your opinion  
6       that Ms. Gelbach contributed to the on-board dose  
7       calibrator. She was named on the 2008 application, and  
8       then also all the way through to the asserted patents which  
9       claim the on-board dose calibrator, right? Answer. I  
10      believe that is correct that she is the only one. Was that  
11      your testimony?

12           A       That was my testimony.

13           Q       Do you still believe it to be true?

14           A       She brought the idea. It was not recognized as  
15      an invention until after Jubilant had their product on the  
16      market. It was an idea. It was a part of a product, not a  
17      claimed invention.

18           Q       Right. And it was disclosed in the  
19      specification in 2009, correct?

20           A       The idea was indeed disclosed in the  
21      specification.

22           Q       And again, she got this idea while working at  
23      Bracco before she went to JDI?

24           A       She got this idea, as she stated, looking at and  
25      having seen an on-board dose calibrator in commerce, that's

1 correct, that was while she was working at JDI -- I'm  
2 sorry, Bracco.

3 Q Did you read her testimony also that the only  
4 reason that the design team not to -- decided not to  
5 incorporate the on-board dose calibrator was for  
6 cost-saving purposes?

7 A I saw that.

8 Q Now, in forming your opinion regarding  
9 inventorship, you did not make a determination of what each  
10 inventor contributed, correct?

11 A No. I did not.

12 Q All right. And beyond the fact that Janet  
13 Gelbach contributed the idea of the on-board dose  
14 calibrator, you had no information that links various claim  
15 elements to various inventors, correct?

16 A No. The only thing we could look at was what  
17 had been disclosed and who were the claimed inventors of  
18 those elements in the previous patents and those were not  
19 the claimed inventors of the patents that are in suit now.

20 Q And just to be clear, in your deposition you  
21 stated that your opinion was not that the named inventors  
22 should not have been named, but rather you thought that  
23 additional people should have been added as inventors,  
24 correct?

25 A I think the correct inventors are not the ones

1 that were named.

2 Q Okay. But I just want to clarify for the record  
3 who you think is wrongly named. At your deposition you  
4 said you didn't think anybody listed on the patent should  
5 not have been listed on the patent, correct?

6 A That's correct.

7 Q All right. So we are talking about people named  
8 on other patents you think should have been added as  
9 additional inventors?

10 A We are talking about people who actually claimed  
11 that they invented certain claim elements that were left  
12 off the patent.

13 Q And you haven't identified any motivation by  
14 Bracco for purposely leaving any inventors off, correct?

15 A I would have to speculate with regard to any  
16 motivations.

17 Q Right. And indeed on the named -- on the  
18 asserted patents, there are named inventors who worked for  
19 third-party vendors such as North Pole Engineering,  
20 correct?

21 A That's correct.

22 Q Now, to your knowledge, did JDI depose any of  
23 the people that you thought should have been named as  
24 inventors to find out what they contributed?

25 A I have no knowledge of that.

1           Q       Okay. Let's turn to prosecution laches. In  
2 your presentation as RDX-2, page 35, you failed to address  
3 multiple patent applications that relate to the patents in  
4 suit that were being prosecuted in the relevant time  
5 period, correct, and this continues on throughout your  
6 slides?

7           A       I did.

8           Q       Right. And -- and at your deposition you  
9 indicated that you purposely left off some of those  
10 applications, correct?

11          A       I don't believe I said I purposely left them  
12 off.

13          Q       Well, you knowingly left them off, correct?

14          A       I don't think I tracked that through. I don't  
15 think it was relevant to the analysis we were doing.

16          Q       Okay. So you left them off despite knowing the  
17 applications because you thought those other applications  
18 being prosecuted were not relevant to the issue of  
19 prosecution laches?

20          A       I don't think they were relevant to these  
21 particular patents.

22          Q       Okay. So it's not that you opine that Bracco  
23 wasn't actively prosecuting patent applications in this  
24 patent family during the relevant time frame, correct?

25          A       That would be likely a valid statement.

1 Q Okay. So your opinion is based purely on the  
2 raw amount of time that passed between the priority filings  
3 and the time that claims were written that correspond to  
4 what are in the asserted patents?

5 A It's fundamentally based on the amount of time  
6 that went by from disclosure of an idea which seems to be  
7 germane to this topic before one files it and after one  
8 sees another product on the market so converting an idea  
9 into a claimed patent. I believe that's my analysis with  
10 regard to prosecution laches.

11 Q Okay. Let's talk about obviousness. So you  
12 don't have any experience working with elution infusion  
13 systems, correct?

14 A I do not work with it. I'm not a medical  
15 practitioner.

16 Q Okay. And your obviousness analysis is largely  
17 based on four references, correct?

18 A That's correct.

19 Q And the sole primary reference upon which you  
20 rely is the Klein thesis?

21 A I believe that the Klein thesis is a primary  
22 reference. I don't use that term as sole because there  
23 were other things that were brought in that are key to a  
24 couple of the components there.

25 Q Okay. But you didn't use any of the other art

1 that you relied on as the primary reference that would be  
2 modified by other references, correct?

3 A If you want to call that the sole reference or  
4 the primary reference that's your calling. I looked at all  
5 of those as being applicable in doing my analysis.

6 Q Okay. And the analysis you did was how would  
7 the Klein thesis be modified based on these other  
8 references, correct?

9 A I believe that most all the terms are addressed  
10 by Klein. Other terms are addressed by the other  
11 references that I utilized.

12 Q Okay. So for example, Tate. You didn't say it  
13 was obvious to modify Tate to match the claimed -- the  
14 claims in this case. You said it was, it would be obvious  
15 to modify Klein using Tate?

16 A That's a fair representation.

[REDACTED]

1 Q Okay. And so was Tate and the CardioGen,  
2 correct?

3 A Yes.

4 Q And you clarified at your deposition that you  
5 weren't relying on the Version 1 or the Version 0 or the  
6 Version 2 or the RUBY-FILL as prior art. You were relying  
7 on the Klein thesis, right?

8 A That is correct.

9 Q Now, so with Klein, CardioGen and Tate all of  
10 record before the Patent Office, the Patent Office still  
11 allowed the claims at issue in this case?

12 A I believe the Patent Office was looking mostly  
13 at whether they were anticipated and I saw him referring  
14 very often to deKemp and I saw very few references to  
15 Klein.

16 Q Okay. But they were all of record before the  
17 Patent Office?

18 A They were.

19 Q Okay. And yet the Patent Office still allowed  
20 the claims.

21 A They did.

22 Q Okay. Now, Tate is an FDG system, right?

23 A It is.

24 [REDACTED]

25 A Yes.





1 not a variable source of background radiation, correct,  
2 when it's sitting on the cart?

3 A When it's sitting on the cart, that's correct.

4 Q Okay. And so the amount of background radiation  
5 that the dose calibrator is exposed to as a result is also  
6 not variable.

7 A That is a true statement.

8 [REDACTED]

9 [REDACTED]

10 A No. Those systems do not include the necessary  
11 plumbing for a rubidium generator.

12 Q And for the Tate and Medrad saline isn't used  
13 for the same purpose as it is in Klein, correct?

14 A That's a partially correct statement but it's  
15 not totally true.

16 Q Okay. So I mean you're certainly not using  
17 saline with a generator, correct, to create the radio --  
18 radiopharmaceutical on board the cart?

19 A Saline is not used to generate a rubidium  
20 elution. It is instead used to push the sample of the  
21 radioactive material into the patient just as it is in the  
22 RUBY system.

23 Q Right, but in the claimed invention you use the  
24 saline and it actually travels through the generator to get  
25 the rubidium?

1           A        It does indeed.

2           Q        Okay.  And that's not what's happening in Tate  
3 or Medrad?

4           A        No.  That particular feature doesn't happen.

5           Q        Okay.  So let's go through and see if we can get  
6 agreement on what the Klein thesis doesn't disclose.  So  
7 the Klein thesis does not disclose a first opening in the  
8 first shielding compartment being at a lower elevation than  
9 the second opening of the second shielding compartment,  
10 correct?

11          A        The Klein thesis does not disclose that.

12          Q        Okay.  And the Klein thesis also does not  
13 disclose a shielded well configured to fill the eluate  
14 reservoir in the shielded well on board the cart?

15          A        No.  As we stated, the on board the cart feature  
16 is not there.

17          Q        Okay.  Klein thesis does not disclose a computer  
18 configured to determine a strontium radioactive eluate  
19 filled on board the cart while the eluate reservoir remains  
20 in the shielded well on board the cart, correct?

21          A        The on board the cart is missing from the Klein  
22 thesis.

23          Q        Okay.  And the Klein thesis also does not  
24 disclose a computer configured to measure a calibration  
25 radioactivity of the sample while the sample remains in the

1 eluate reservoir in the shielded well on board the cart?

2 A As we stated, the Klein thesis does not include  
3 the on board element.

4 Q Okay. And so, for example, the dose calibrator  
5 is not in a shielded cart on board the cart in Klein?

6 A The system has all those components but it does  
7 not do it on board the cart.

8 Q Okay. And Klein thesis does not disclose  
9 configuring a computer to present on a touch screen display  
10 a screen reminding the user to insert the eluate reservoir  
11 in the shielded well on board the cart?

12 A The Klein thesis prototype does not indeed have  
13 that particular reminder disclosed.

14 Q Okay. And the Klein thesis does not disclose a  
15 specific elevations of the openings of the first and second  
16 shielding compartments, correct?

17 A No. Those elevations -- as I've stated, those  
18 are obvious configuration choices that one would utilize in  
19 changing a prototype into a product.

20 Q Well, I think your testimony was that they are  
21 just irrelevant. Is that what you stated on direct?

22 A I believe these are standard things that a  
23 person would do when they are going from a prototype to a  
24 product so the relative heights I believe aren't relevant.  
25 That's correct.

1 Q Okay. And if they are irrelevant there is no  
2 motivation to change them?

3 A I don't think there is anything inventive about  
4 putting components at different heights.

5 Q Okay. But I mean, if people don't care about  
6 it, there is no motivation to change those features, right?  
7 Especially on a system as complicated as the RUBY?

8 A We spoke about putting heavy items where they  
9 would not have to be lifted great distances for ergonomic  
10 purposes which is what one would do going from a laboratory  
11 prototype to a production device that would be used in an  
12 occupational use.

13 Q Okay. Oh, I'm sorry. I thought you were done.  
14 Klein thesis does not disclose tracking the  
15 volume of saline remaining in the saline reservoir,  
16 correct?

17 A The Klein thesis does not track what's remaining  
18 in the reservoir. That is correct.

19 Q Okay. And the Klein thesis does not disclose  
20 providing an alert on the touch screen display when the  
21 volume of saline remaining in the reservoir is below a  
22 predetermined volume?

23 A That's correct. That laboratory prototype did  
24 not disclose that.

25 Q All right. Similarly, Klein thesis did not

1 disclose a first door accessible via the opening through  
2 the exterior shell, the first door being configured to  
3 provide access to the first shielding compartment and to  
4 close over the first opening, correct?

5 A I'm sorry. If the first shielding compartment  
6 -- again.

7 Q Sure. I'll just repeat it. So Klein thesis  
8 does not disclose a first door accessible via the opening  
9 through the exterior shell, the first door being configured  
10 to provide access to the first shielding compartment and to  
11 close over the first opening.

12 A The Klein thesis had the first shielding  
13 compartment, which I believe was for the strontium-rubidium  
14 generator, is that correct? So I can answer this  
15 correctly.

16 Q So why don't we bring up your deposition at page  
17 260, lines 6 through 21. So do you see the question?  
18 Okay. And just going by the claim element to make sure  
19 that we understand what your opinions are. All right. I'd  
20 like to talk to you about the '869 patent for a moment,  
21 specifically element 1.3, which is on page 413 that's of  
22 your report, correct?

23 A I see that.

24 Q In the images here from the Klein thesis you've  
25 not labelled the first door accessible via the opening

1 through the exterior shell, the first door being configured  
2 to provide access to the first shielding compartment and to  
3 close over the first opening. Right? And you answer, I  
4 have not?

5 A That's what I said.

6 Q And the follow-up question was, well, is it  
7 disclosed by Klein? And you said, no. Correct?

8 A That's what I said at this location. That's  
9 correct.

10 Q Okay. And you still maintain that testimony,  
11 correct? That was true then? It's true now?

12 A To the best of my memory.

13 Q Okay. And the Klein thesis does not disclose  
14 that the infusion system is configured for the saline  
15 tubing line and the eluate tubing line to be routed through  
16 two tubing passageways formed in a perimeter surface of the  
17 first opening wherein each of the two tubing passageways  
18 has a depth configured to prevent pinching or crushing of  
19 the corresponding tubing line, correct?

20 A No. We were not relying on the Klein thesis for  
21 anticipation. We were relying on how it would be modified  
22 for obviousness; and as I stated in my prior testimony, if  
23 one is going to route those tubing lines through the  
24 perimeter it would be obvious to anyone skilled in the art  
25 not to configure those so that they would not be pinched or

1 crushed by a door. That is the only thing that makes  
2 sense.

3 Q At this point I'm just trying to make sure we  
4 have agreement on what Klein discloses and doesn't  
5 disclose. We'll get to combinations a little bit later.  
6 So do you agree that Klein didn't disclose that claim  
7 element, correct?

8 A Klein did not disclose that claim element.

9 Q All right. And you didn't identify anything in  
10 the Klein thesis indicating that the system disclosed  
11 therein should be modified in any way, correct?

12 A I'm sorry.

13 Q Yeah. So in Klein itself you didn't point to a  
14 passage in Klein that said, you know, you may want to  
15 consider modifying this feature that I disclosed?

16 A No. I don't recall, although he does talk about  
17 improvements that could be made in his system before he  
18 closes things out.

19 Q Okay. But you haven't identified anything in  
20 your direct with regard to a motivation from the Klein  
21 thesis itself to make any of the modifications that you  
22 propose?

23 A I did not identify anything directly in the  
24 Klein thesis for those motivations. That's correct.

25 Q Okay. And so could you turn to page 417 of your



1 initial report. Or you can just read it on the screen if  
2 that's easier. I just want to ask you about element 1.5.  
3 So see here, a second door accessible via the opening  
4 through the top surface of the exterior shell, the second  
5 door being configured to provide access to the second  
6 shielding compartment and to close over the second opening.

7 A I see that.

8 Q Okay. Now, let's turn now to RDX-2.196 and talk  
9 about what you considered to be the door in Tate. So do  
10 you identify on the record what you consider to be the door  
11 shown here?

12 A I believe we've shown a door in Tate here. Or  
13 down below. Can we back up.

14 Q Yes. Why don't we go to the one down below the  
15 figure 4A.

16 A We talked about this door right here.

17 Q So this is the red highlight is what you  
18 considered to be the door disclosed by Tate?

19 A That is the door disclosed by Tate.

20 Q All right. Now, Tate doesn't actually call it a  
21 door, does it?

22 A I think he calls it a lid. Lids, door. Access  
23 hatches. These are all things you would find together, I  
24 believe, in any thesaurus.

25 Q Tate calls it a vial access system, correct?

1           A       Vial access lid, I believe.

2           Q       So why don't we pull up Tate RX-103, page 97,  
3 paragraph 139. You have the vial access system, right? So  
4 you've got it's a system. It's got the vertical support  
5 arm that goes to the housing. You've got a cap member and  
6 a handle member and they are all connected to an upper end  
7 of the vertical support arm, correct?

8           A       Yes. I believe he calls it the cap member.

9           Q       Okay. And then what you do is you grab the  
10 handle, you pull it up and you twist, correct?

11          A       That's one of the things that you do. Yes.

12          Q       All right. Let's go back to the RDX-2 at 196.  
13 All right. Now, in Tate, this is designed to be used on  
14 top of the cart, correct?

15          A       It's designed to be accessed via the top of the  
16 cart. That's correct.

17          Q       Okay. And that access system is mounted via  
18 that sliding arm, correct?

19          A       That's correct.

20          Q       Now, the claimed door that you're equating the  
21 system in Tate to, that's the door for access to the  
22 generator, correct, and that's why you colored it red?

23          A       That's correct.

24          Q       Okay. Now, let's look at RDX-2 at 166. So if  
25 we can blow up the RX-106 at 34 portion. So here in your

1 modification of Klein the generator is on the bottom,  
2 correct?

3 A That's correct.

4 Q And you've got the dose calibrator and the lead  
5 shielding right above it?

6 A That's how it's shown configured. Yes.

7 Q Okay. So you couldn't actually use what you  
8 describe as the door in Tate to obtain access to the  
9 generator in this configuration, could you?

10 A No.

11 Q All right. So one wouldn't be motivated to use  
12 the door in Tate to access the generator if the door in  
13 Tate wouldn't work in the design?

14 A That particular configuration would not work.  
15 It would be obvious to a person of skill in the art to  
16 configure his door such that one could have access to the  
17 generator as described in Klein.

18 Q Okay. But so the door in Tate as it's shown is  
19 not used?

20 A I disagree. He has a door. He has it where it  
21 provides access. The manner in which it provides access,  
22 how it's attached, those are obvious design choices that a  
23 person would make as one configures a commercial system.

24 Q So Tate, the door in Tate as shown -- well, the  
25 vial access system -- when you -- to pull out the rod you

1 would just, your hand would just run into the dose  
2 calibrator and lead shielding above before you could gain  
3 access to the generator, correct?

4 A I'm sorry?

5 Q Sure. If you were using the vial access system  
6 actually disclosed in Tate that you showed with the -- that  
7 uses the slidable bar that you pull up on, if you were to  
8 try to use that as the door to the generator, your hand  
9 would just run into the dose calibrator and shielding or  
10 whatever supporting that before you actually gained access  
11 to the generator?

12 A I believe you were the one that said it was the  
13 dose access system. I said it was the cap that was the  
14 door and the attachment to the door I don't believe is an  
15 inventive process.

16 JUDGE CHENEY: And with that, we will take our  
17 lunch break. We'll see you in one hour.

18 MR. DAVIS: Thank you, Your Honor.

19

20 (Whereupon, at 12:33 p.m., the hearing in the  
21 above-entitled matter was recessed, to reconvene at 1:33 p.m.)

22

23

24

25

1 AFTERNOON SESSION

2 (1:33 p.m.)

3 JUDGE CHENEY: We are back on the record in the  
4 1110 investigation. Before our lunch break we were  
5 listening to the cross-examination by complainants of  
6 Dr. Stone, who has been called by respondents as an expert  
7 on issues relating to patent validity. Mr. Davis, you may  
8 resume.

9 MR. DAVIS: Thank you, Your Honor.

10 BY MR. DAVIS:

11 Q Dr. Stone, could you turn to RX-106 and page 34,  
12 the Klein thesis, and that photo. 34 of RX-106. That's 34  
13 of the document. Sorry. There we go. So could we blow up  
14 the top photo. All right. Now, could you explain to me  
15 again what, in your opinion, is the front side of the  
16 exterior shell?

17 A Certainly. Front slide of the exterior shell is  
18 determined by these four corners on the front of the device  
19 of the cabinet.

20 Q Okay. So in your opinion, the front side of the  
21 exterior shell includes this opening, correct?

22 A That's correct.

23 Q All right. What's your understanding of the  
24 word shell?

25 A A shell is something that tends to surround

1 something. It has a cover, for example, a turtle has a  
2 shell across its back and openings for its legs. A shell  
3 is an enclosure.

4 Q Okay. So it's something that encloses or  
5 protects. Is that fair?

6 A That's a fair assumption. Yes.

7 Q All right. Now, is there anything in the front  
8 side enclosing or protecting the various filament that's  
9 shown here?

10 A It doesn't say the front side has to do with the  
11 completion of the enclosing and protecting, just that the  
12 cabinet does that.

13 Q So but it is a four-sided exterior shell,  
14 correct?

15 A It is a four-sided cart.

16 Q It's a four-sided shell.

17 A It's a four-sided cart enclosed on three sides,  
18 top and bottom.

19 Q Okay. So, so your opinion that this meets the  
20 shell is based on your understanding that the claims don't  
21 call for a four-sided exterior shell?

22 A No. In fact the claims do call for two side  
23 walls. It does not call for a front wall.

24 Q So let's -- all right. Could we go to JX-2 at  
25 page 45, column 27, lines 47 to 50. So here we have a

1 limitation that the exterior shell further includes an  
2 opening, correct?

3 A That's correct.

4 Q Now, so is it your opinion that the front of the  
5 cart constitutes both the opening and part of the shell?

6 A The opening that it calls for here is for a  
7 saline tubing line.

8 Q Let's go to the picture, page 34 of 106 again.  
9 RX-106, page 34. All right. So where is the saline --  
10 where is the opening in the exterior shell for the saline  
11 tube?

12 A The opening for that, as we disclosed earlier,  
13 is through the top surface here.

14 Q All right. So Ms. Gelbach considered FDG to be  
15 a whole different product that does not do the same type of  
16 study as a system like a CardioGen, correct?

17 A I believe she may have stated that. Yes.

18 Q Okay. And you have opined that it would have  
19 been obvious for a person of ordinary skill to take the  
20 on-board dose calibrator of Tate and incorporate it on  
21 board into the Klein thesis, correct?

22 A I have.

23 Q All right. And you agree that the on-board dose  
24 calibrator of Tate measures radioactivity that will be  
25 delivered to the patient, right?

1           A       That is correct.

2           Q       All right.  And the Klein thesis that already  
3 disclosed a detector separate from the dose calibrator that  
4 measures the radioactivity that's delivered to the patient,  
5 right?

6           A       Yes.  After it's been calibrated by dose  
7 calibrator.

8           Q       Okay.  So the Klein thesis doesn't need Tate's  
9 dose calibrator to measure the radioactivity delivered to  
10 the patient?

11          A       No.  Only to calibrate the detector that is  
12 measuring that.

13          Q       Now, Klein conducts breakthrough testing,  
14 correct?

15          A       He does.

16          Q       And it does that with the off-board dose  
17 calibrator, correct?

18          A       He does.

19          Q       And you've pointed to nothing in the Klein  
20 thesis that indicates that Klein thought that the  
21 breakthrough testing with an off-board dose calibrator  
22 should be changed, correct?

23          A       Klein includes an off-board calibrator as part  
24 of his system.  He doesn't say whether -- he doesn't teach  
25 any changing in the Klein thesis.



1 Q All right. Let's go to 247 of RDX-2. All  
2 right. So this shows your proposed placement of the dose  
3 calibrator, the generator and the shielded waste container,  
4 correct?

5 A That's correct.

6 Q All right. Now --

7 A Excuse me. That's what a person of the art  
8 might do. It's not necessarily I'm proposing that. But it  
9 would be obvious do that.

10 Q Okay. So this is, but this is the configuration  
11 that you testified regarding?

12 A That's correct.

13 Q All right. Now, first of all, the front opening  
14 is blacked out but there is pre-existing equipment on the  
15 shells not shown in this picture, correct?

16 A Yes. He used pre-existing off-the-shelf  
17 equipment to put together his laboratory prototype.

18 Q And to put the shielded waste container there  
19 and the shielded generator and the dose calibrator in  
20 shielding you'd have to move all that equipment that was  
21 previously there. You'd have to find a new home for it?

22 A As I'm configuring a product I would probably  
23 find new homes for that equipment. That's correct.

24 Q Okay. And you opine that one would have been  
25 motivated to put the generator down low for ergonomic

1 purposes, correct?

2 A I do.

3 Q All right. Now, the premise of your ergonomic  
4 analysis is that with a lower generator you're closer to  
5 the floor, correct?

6 A You're closer to the level of a cart that would  
7 be rolled up having the generator on it to transfer it into  
8 the device and you're also dropping the center of gravity  
9 making the cart more stable.

10 Q So I believe you're -- now, you did not actually  
11 do any analysis of how users of the system actually handle  
12 the generator, correct?

13 A I did not.

14 Q All right. And so if, for example, somebody was  
15 motivated not to bend over to pick something up from down  
16 low, but rather to keep it up high because it's heavy, the  
17 ergonomics would be to keep the generator up high so that  
18 you could move it to the shelf more easily, correct?

19 A If the device were there on the shelf. However,  
20 I note that the CardioGen-82, the so-called Model 510,  
21 already had the generator down low.

22 Q Okay. But I'm just asking about ergonomics.

23 JUDGE CHENEY: Mr. Davis, when you're away from  
24 the mic then the court reporter doesn't hear you through  
25 her headset.

1 MR. DAVIS: I apologize, Your Honor.

2 JUDGE CHENEY: Please stick close to the mic.

3 BY MR. DAVIS:

4 Q But you didn't do an ergonomic analysis of how  
5 people actually handle the generator?

6 A I did not.

7 Q And the University of Ottawa -- I'm sorry.

8 A I am aware of how heavy objects are transported  
9 and the types of carts that they are. It's very typical  
10 for that to be a low, near-floor cart so a person of skill  
11 in the art would know that.

12 Q You understand that as an expert you're supposed  
13 to disclose your opinions in your report, correct?

14 A To the best of my knowledge. Yes.

15 Q Okay. So I just want to ask you about the  
16 opinions you've already disclosed in your report and  
17 testified today. I'm not, I'm not asking you to form any  
18 new opinions.

19 A I'm sorry. You asked me about my analysis as to  
20 whether I had done that. I formed my opinion based on what  
21 I as a person of skill in the art knew already.

22 Q Are you an expert in ergonomics?

23 A I utilize ergonomics and have to review those  
24 when I'm doing a product definition.

25 Q So now, the University of Ottawa's request for

1 information included in RX-144 touts the ergonomic design  
2 of the existing design for the Version 1, correct?

3 A Yes.

4 Q All right. So that's the last bullet point on  
5 page 10?

6 A Yes.

7 Q All right. So, in this version the generator  
8 was up high, correct?

9 A What they are discussing is they are talking  
10 about how the device can easily be used without powering  
11 down minimizing the amount of motion that has to take  
12 place, the amount of time. They are not discussing here  
13 ergonomics with regard to the weight and the orientation of  
14 the materials.

15 Q They are talking about an overall ergonomic  
16 design, correct?

17 A No. They are talking about adding convenience  
18 through ergonomic design. They are not talking about  
19 necessarily what would be in a finished product.

20 Q So they are talking about this design being  
21 ergonomic, correct?

22 A They are talking about they have added  
23 convenience through some ergonomics.

24 Q All right. And that was with regard to the  
25 existing configuration?

1           A           With the existing configuration which is a  
2 prototype ready to be productized.

3           Q           All right. And there is also restrictions on  
4 the placement of the dose calibrator on the cart due to  
5 possible interference between the dose calibrator and the  
6 generator, correct?

7           A           That's correct. That's a matter of how the  
8 shielding is set up and where the components are place.

9           Q           So one of ordinary skill wouldn't want to place  
10 the dose calibrator right above the generator, you'd want  
11 some distance?

12          A           Depending on the shielding, distance would be a  
13 factor that they would consider.

14          Q           Okay. But if you kept it close you'd have to  
15 add additional shielding which would add to the weight of  
16 the cart?

17          A           Actually, I believe they actually measured this.  
18 It's even reported, I believe, in one of our documents that  
19 we referenced that they measured it and didn't have a  
20 problem with the generator. They had a problem with  
21 something else.

22          Q           But in your deposition, you stated that there  
23 are restrictions on the placement of the dose calibrator on  
24 the cart due to possible interference between the dose  
25 calibrator and the generator, correct?

1           A        There could be possible interferences. That's  
2 correct.

3           Q        Right. One of ordinary skill would know that  
4 and one of the ways to address that would be to separate  
5 the two?

6           A        That's correct.

7           Q        You're also aware that when the University of  
8 Ottawa met with JDI to discuss a possible modifications to  
9 the Version 2, they suggested that the tubing be changed so  
10 that the system could be used with a variety of existing  
11 dose calibrators, right?

12          A        That was a suggestion. That's correct.

13          Q        Right. So that was in the 2008 time frame?

14          A        I believe 2007, 2008 time frame.

15          Q        And according to Mr. Donnelly, as of 2004,  
16 the -- the Version 1 had already been used to perform  
17 procedures on 667 patients, right?

18          A        That's correct.

19          Q        Right. So that number would be even larger by  
20 the 2008 meeting?

21          A        That's the assumption I would make. Yes.

22          Q        Okay. So even after, and the first use by  
23 University of Ottawa of a Rubidium Elution System was 1997,  
24 right? The Version 0?

25          A        The Version 0 -- by the University of Ottawa,

1 1997. I believe that's correct.

2 Q Right. So you remember --

3 A I'd have to refresh my memory on the document.  
4 I don't recall that.

5 Q Okay. So even after 11 years approximately from  
6 '97 to 2008, and somewhere, you know, somewhere north of  
7 700 procedures, Dr. Klein and the University of Ottawa  
8 still thought the dose generator should remain off the  
9 cart, correct?

10 A I think you misstated that. Do you want to read  
11 back your own question?

12 Q I'll just restate the question. So, so the  
13 first -- Ottawa starts using the rubidium system '97 with  
14 Version 0. They are talking to JDI in 2008 time frame. We  
15 are talking about 11 years and we are talking about, you  
16 know, nearly 700 procedures just using the Version 1 alone  
17 by 2004. Even after all that experience and all that time,  
18 University of Ottawa was still thinking and Dr. Klein was  
19 still thinking when they are discussing options with JDI,  
20 that the dose calibrator should remain off the cart and  
21 that way the cart could be used with various existing dose  
22 calibrators that the facilities already had?

23 A I don't think they necessarily think it should,  
24 but it did.

25 Q But that was the, that was the design they were

1 proposing to JDI. They are saying, you know, a couple of  
2 things we need to update. One of them is to change the  
3 tubing so we can have it interchangeable with several  
4 different dose calibrators rather than just the one that we  
5 were using at Ottawa?

6 A I don't think the tubing is related to the dose  
7 calibrator, is it?

8 Q Well, they were talking about changing the  
9 tubing in order to make the cart compatible with multiple  
10 dose calibrators. Why don't we bring that up. So day two,  
11 page 340 and 41, Mr. Donnelly's testimony was that the  
12 technology transfer happened and you'll need to reprove --  
13 one of the things was to change the tubing so that the cart  
14 would work with multiple types of dose calibrators because  
15 there were multiple types of dose calibrators on the  
16 market. Right. And Mr. Donnelly said correct. So --

17 A Okay. Sorry. Go ahead.

18 Q So as of this time period, Ottawa and Dr. Klein  
19 were suggesting to JDI that one of the changes they should  
20 make going forward is to make the cart compatible with  
21 multiple dose calibrators because various facilities had  
22 different dose calibrators?

23 A This seems to be discussing tubing and the  
24 tubing goes and is connected to the vial. The dose  
25 calibrator being off the cart. The only thing I can think



1 of that that might have to do with the length of the  
2 tubing. It's not discussed so it would be very unclear as  
3 to why that would be significant.

4 Q So you need to connect the dose calibrator to  
5 the cart with a tube, correct?

6 A No. You need to connect the vial that's in the  
7 dose calibrator to the cart with the tube.

8 Q So you need -- so you had various vials used  
9 with the dose calibrators?

10 A That's a possibility. I don't know the answer  
11 to that.

12 Q All right. Now, please turn to CX-413C --  
13 excuse me. Let me just check. Your Honor, I'm sorry. We  
14 need to go on the confidential record. [REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

[REDACTED]

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[Redacted text block]

[REDACTED]

9 MR. DAVIS: Your Honor, I can switch to a  
10 different line of questioning to give him time to review.

11 JUDGE CHENEY: Okay.

12 BY MR. DAVIS:

13 Q So before we address 413, could I ask you to  
14 turn to JX-2 and specifically page 43 and I want to pull up  
15 claim one and the claim relating to an exterior shell which  
16 is the first main clause after the preamble. I'm going to  
17 blow that up. Okay. So do you see the claim language  
18 there where it talks about a shell that extends upwardly  
19 above the platform and has a front side, a rear side, and  
20 two side walls connecting the front side to the rear side?

21 A Yes.

22 Q Okay. So let's go back to RX-106, page 34, but  
23 let's keep this as well. Sorry. So let's blow up that top  
24 photo and blow up that one paragraph. Okay. So we've got  
25 a shell and it extends upwardly above the platform and it's

1 got a front side, a rear side, and two side walls that  
2 connect the front side to the rear side. You know, we've  
3 established that, so where is the front side of the shell  
4 in the Klein, in the device shown in the Klein thesis?

5 A The front side of the shell is this portion  
6 right here with a large opening in the center of it.

7 Q Okay. So can you identify anything that's not  
8 the opening that's the front side that would constitute the  
9 shell?

10 A Certainly at least these folded-down edges are  
11 part of the front side.

12 Q Okay. Those edges don't enclose the components  
13 that are shown in the photo, correct?

14 A I'm not sure what you're trying to get at here.  
15 These are a front side. These are part of the front side.  
16 They are certainly not of the side walls. They are not at  
17 the backside. They are not at the top side. They are part  
18 of the front side and they have helped form the enclosure  
19 of the entire system, interior of the shell.

20 Q Okay. So it's your position that the, those  
21 narrow sides in the front that that encloses the, the  
22 equipment that's shown in the photograph?

23 A I see nothing there that requires a front wall.  
24 It says a front side. Sides don't necessarily make a total  
25 enclosure.

1 Q Okay. But you would agree that the claim  
2 language is an exterior shell that has a front side?

3 A It has a front side. It doesn't say it forms a  
4 complete enclosure.

5 Q All right. So okay. Now, let's look at the  
6 next element. So you've got an opening through the  
7 exterior shell configured to provide access to the  
8 generator. What part is the opening to provide access and  
9 what part is the shell in the front?

10 A There is an opening right here. There is a  
11 shell through, in the front that extends around from the  
12 sides. I'm not sure where you're trying to go.

13 Q Do you recall Mr. Donnelly's testimony earlier,  
14 that given the variety of things that can go wrong when you  
15 implement design changes to a device like the RUBY-FILL  
16 that you typically don't change a feature that's -- that --  
17 let me repeat the question for you. You heard Mr. Donnelly  
18 testify that given the variety of things that can go wrong  
19 when you implement design changes to a device like the  
20 RUBY-FILL, you typically don't change a feature that is  
21 known to work?

22 A I think that's an incomplete opinion. You  
23 wouldn't change a feature that's known to work where your  
24 change could affect how it works. You might change a  
25 feature if it would not affect how it works.

1           Q       Okay. Well, let's look at his testimony at page  
2 349 of the transcript, lines 7 through 19. To the  
3 question, well, when you're designing, if you know  
4 something works you're loathe to change it to something  
5 else because you're going to have to go back and verify  
6 that the -- that the new design works and there may be  
7 unforeseen consequences like the tubing or the bending of  
8 the cart so you try to avoid changing something that you  
9 know works?

10                    Answer, we would -- I think it's common in the  
11 industry that you don't make changes if not necessary, but  
12 of course there are changes that you have to make if there  
13 are expected to be any issue with the design that you have  
14 so we always make a change if we need to make a change. If  
15 we don't need to make a change we would not make it.

16           A       I see that he said that.

17           Q       Do you agree with that principle?

18           A       In general.

19           Q       Okay. And at your deposition you admitted that  
20 you did not identify in your report any market forces that  
21 would prompt one of ordinary skill to make the various  
22 changes to Klein that you opine would have been obvious,  
23 correct?

24           A       I may have said that in my deposition.

25           Q       Okay. And the -- you still agree with that?

1           A       The changes to Klein that would have been  
2 obvious.

3           Q       I'm sorry. Let me restate the question. You  
4 have not identified any market forces, have you, that would  
5 prompt one of ordinary skill in the art to make the various  
6 changes in Klein that you opined would be obvious, correct?

7           A       I believe I've identified what Miss Gelbach said  
8 that market forces were saying that it would only make  
9 sense to put the dose calibrator on board the cart.

10          Q       So let's look at your deposition, page 340 --  
11 354, lines 11 to 23. So -- I'm sorry. I'll get -- there  
12 appears to be an error. The weight of the shielding can  
13 also complicate placement of the dose calibrator on the  
14 cart, correct?

15          A       I would say it would complicate the design of  
16 the frame of the cart.

17          Q       Right. And as late as 2015, JDI was still  
18 having issues with the cart bending and pieces not fitting  
19 because of the weight putting the dose calibrator on board  
20 the cart, correct?

21          A       I think what they actually had was a problem  
22 with the enclosure not closing correctly, having a little  
23 bit of warp to it so when you refer to bending there was a  
24 displacement that occurred that somehow they were trying to  
25 track down exactly what prevented the enclosure of the



1 cart, the plastic panels from fitting correctly.

2 Q Right. Because of the weight of the dose  
3 calibrator?

4 A That was theorized as the cause. That's  
5 correct.

6 Q Okay. So multiple years into the design they  
7 were still addressing issues that they thought related to  
8 putting the dose calibrator on board?

9 A They discovered a minor issue and decided, and  
10 determined how to correct it. The minor issue was they  
11 hadn't quite handled the weight plus the strength of the  
12 materials or how they were arranged so they did a finite  
13 element, suggested doing a finite element analysis to  
14 determine if that were the cause.

15 Q All right. And you also rely on what you  
16 describe as the Medrad system, correct?

17 A Yes. We do.

18 Q All right. And the exhibit you use for your  
19 analysis of that Medrad system is RX-200C.

20 Don't put it up.

21 Correct?

22 A You're asking me if I recall the exhibit number  
23 yet you don't want to put it up?

24 Q Yes. Let's look at RDX-2.100. Right.

25 So I'm sorry. What?

█ [REDACTED]

█ [REDACTED]

3 JUDGE CHENEY: Okay. We are on the Medrad  
4 confidential record. That means if you're not authorized  
5 on either side to view Medrad confidential information, you  
6 need to leave the hearing room now.

7 (Confidential session follows.)

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1

CONFIDENTIAL SESSION

2

BY MR. DAVIS:

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Q The courtroom is cleared, Your Honor.

4

[REDACTED]



[REDACTED]

[Redacted text block containing multiple lines of blacked-out content]



[REDACTED]

4 JUDGE CHENEY: Back on the public record.

5 (End of confidential session.)

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1 OPEN SESSION CONTINUED

2 BY MR. DAVIS:

3 Q Is there any indication in the Klein thesis that  
4 there was a problem with the user not keeping an eye on the  
5 volume of saline which is in plain view of the user?

6 A No. There is no such indication in the Klein  
7 thesis. That's not a product that's out for ordinary  
8 users. We've talked about that being motivated instead by  
9 the usability guide when a product is out for use.

10 Q So the only, the only prompts or alerts that you  
11 point to in Klein are for the generator and waste bottles  
12 which are in the shielding and out of your sight, correct?

13 A Those are the only prompts or alerts. There are  
14 other warnings that he discusses.

15 Q Okay. Now, there were multiple versions of the  
16 RUBY-FILL system, correct?

17 A That's my understanding.

18 Q Right. The -- so let's go to RDX-2C14. All  
19 right. So here is your slide that talks about the  
20 evolution of the elution system. We've got Version 0 in  
21 '97, Version 1 in 2004, Version 2 in 2010 and Version 3 in  
22 2015, correct?

23 A That's correct.

24 Q All right. Now, neither the Version 0 -- well,  
25 Version 0, 1 and 2 never were approved for commercial sale

1 in the U.S., were they?

2 A In the United States. That's correct.

3 Q Okay. And in 2007, Ottawa licensed to JDI the  
4 technology regarding the rubidium PET imaging technology  
5 that they had developed, correct?

6 A That's correct.

7 Q All right. And JDI and Ottawa worked together  
8 on Version 2 and 3, correct?

9 A That's my understanding.

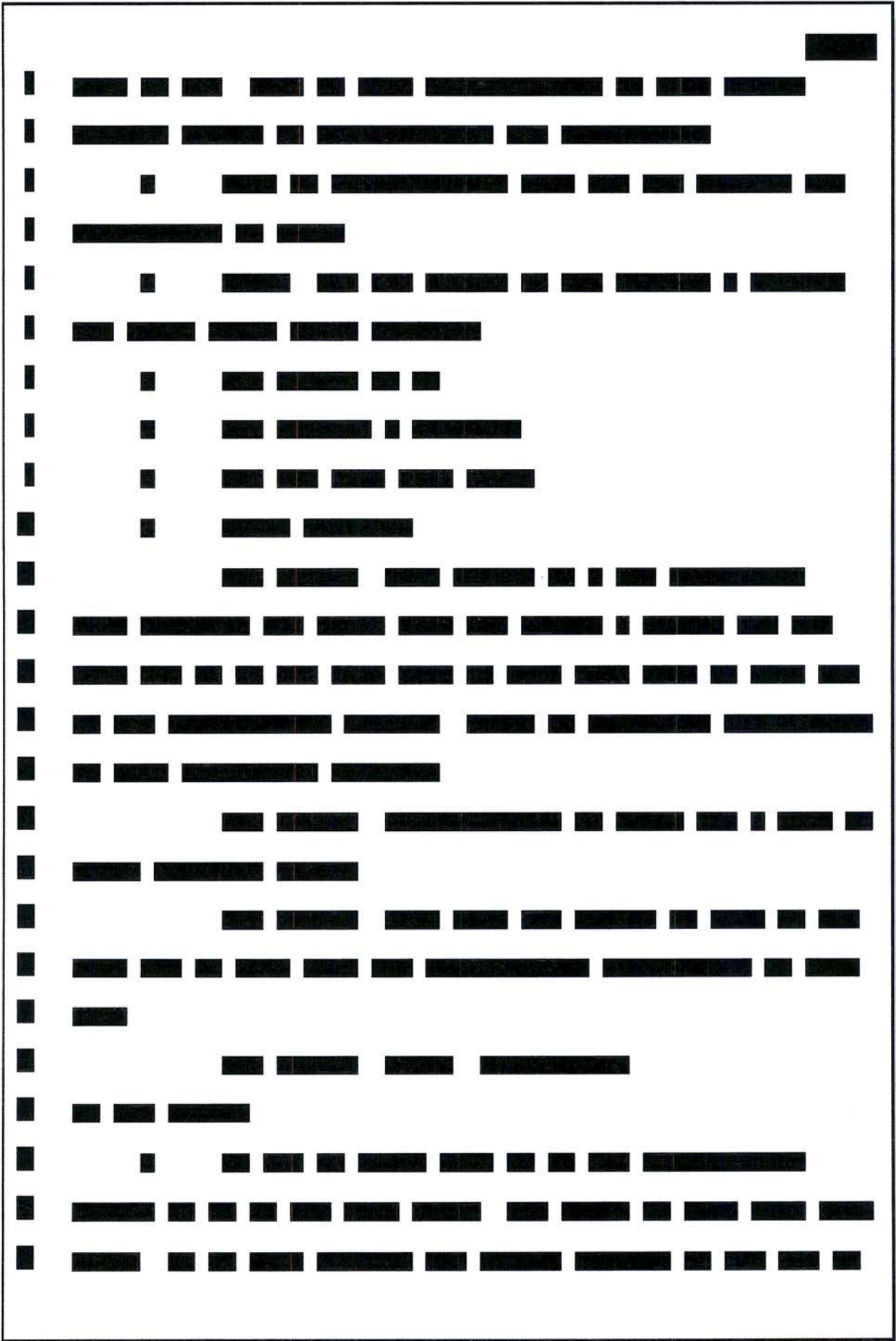
10 Q And so, but prior to working on Version 3, so  
11 Ottawa Heart and JDI had worked on a rubidium PET system  
12 either together or on their own for approximately 13 years  
13 without creating a system that was approved for commercial  
14 sale in the U.S.?

15 A They began working together, it's my  
16 understanding, in 2007. And they ended up with the device  
17 release in 2016. That would be nine years.

18 Q So I was asking either together or alone. So  
19 either Ottawa on its own or Ottawa and JDI were working  
20 over this time period?

21 A Ottawa was not developing a product. Ottawa was  
22 developing a technology, a prototype that would be licensed  
23 for development into a product.

24 Q Okay. Now, and they started trying to get a  
25 commercial partner to develop that at least as early as



1 MR. DAVIS: Your Honor, I pass the witness.

2 JUDGE CHENEY: Okay. Is there cross-examination  
3 by the Commission investigative staff.

4 MR. KOO: Yes, Your Honor. If you would just  
5 give me one minute to find a page number.

6 EXAMINATION BY ITC STAFF

7 BY MR. KOO:

8 Q Good afternoon, Dr. Stone.

9 A Good afternoon.

10 Q I just wanted to follow up on something, to  
11 things that Mr. Davis discussed with you earlier today. If  
12 we could turn to, I believe it's RX-106. And it's the page  
13 that has figures 2-3. So you don't have to expand it or  
14 anything. But looking at that top figure, Mr. Davis asked  
15 you about the front view of that cart that has the opening  
16 so that we can see the computer and the printer and the  
17 other components inside.

18 A Yes.

19 Q Do you recall that discussion? Okay. And I  
20 think you were trying to make the point that because it's  
21 an opening, it's not a side. Would you agree with that  
22 characterization?

23 A I believe that characterized what he was trying  
24 to make. Yes.

25 Q All right. I think what puzzled me up to this

1 point is you'd agree since there is a shielded waste  
2 container, a shielded generator inside the cart, you'd  
3 agree with me that at least there are some, some amount of  
4 tubing that goes inside the cart. Would you agree with me?

5 A Yes.

6 Q Okay. And the materials that are inside the  
7 tubing will carry at least at some point some radioactive  
8 materials?

9 A That's correct.

10 Q As a person of ordinary skill in the art, does  
11 it make sense to you that this cart would only be shielded  
12 on three sides?

13 A Absolutely not. If there is radioactive  
14 material in open tubing, one would supply shielding in  
15 order to reduce the exposures outside the cart.

16 Q Okay. And would you agree that that is what  
17 they have done on the exposed tubing on the top of the cart  
18 with the heavy-duty plastic shield that's shown, I believe,  
19 in figure 2-4?

20 A Yes. I believe it's referred to as high density  
21 polycarbonate shielding that's used to provide shielding  
22 from the beta radiation.

23 Q Okay. Would it surprise you if they had a door  
24 on a hinge that closed that opening on that cart there?

25 A Not in the least.

1 Q There has also been a lot of discussion over the  
2 past two days, I believe, about the features that Ms. Janet  
3 Gelbach had invented or is alleged to have invented as one  
4 of the named inventors in the three asserted patents. Do  
5 you recall that?

6 A Yes.

7 Q And I believe you testified that you agreed that  
8 Miss Gelbach is alleged to have invented the aspect of  
9 bringing the dose calibrator on board the cart?

10 A I believe I used the terminology she brought  
11 that idea to the team.

12 Q Okay. You stated that you read her deposition  
13 transcript?

14 A Yes.

15 Q Okay. Do you recall if, if -- do you recall if  
16 she stated how she came up with that idea?

17 A Yes. She talked to the sites that were using  
18 these devices and she said they were asking for it and said  
19 it only made sense to put the dose calibrator on the cart.

20 Q If we could turn to JX-176C at page 101. Let's  
21 start at page 100 at line 16 and go through 101, line 11.

22 Is this the -- I'll let you read this to  
23 yourself, Dr. Stone, but is this the portion of the  
24 transcript that you're referring to?

25 A Yes.

1 MR. KOO: Okay. I don't think I have anything  
2 further at this time. Thank you.

3 JUDGE CHENEY: Okay. Dr. Stone, I have just a  
4 couple of questions for you. If we could go back to what  
5 Mr. Koo had on the screen, RX-106, page 34. If someone  
6 among the trial wizards could help me with that.

7 Dr. Stone, have you seen this device represented  
8 in this image in real life?

9 THE WITNESS: I'm not certain that I saw the  
10 very one that was in the Klein thesis. I have seen  
11 representations of the Version 1 which includes all of  
12 those components arranged in exactly the same fashion.

13 JUDGE CHENEY: Do you see any Version 1 device  
14 in this hearing room?

15 THE WITNESS: Yes, Your Honor.

16 JUDGE CHENEY: Will you identify it for me.

17 THE WITNESS: It's right there, this first shiny  
18 cart.

19 JUDGE CHENEY: Will someone please identify for  
20 the record the exhibit number that Dr. Stone has  
21 identified.

22 MR. WALKER: Yes. It's RDX-12.

23 JUDGE CHENEY: Okay. Do you see any doors on  
24 RDX-12?

25 THE WITNESS: Yes, Your Honor. It tried to make

1    itself shown as I moved it around.  There is a, there are  
2    hinges there for the door.  It's not clear from the  
3    photograph whether that door was on the cart that was  
4    actually used by Mr. Klein as he did his work.  But it's  
5    certainly obvious to put one on it.

6                JUDGE CHENEY:  So looking at the paper  
7    documentation of Klein, you don't find express disclosure  
8    of the door that we see in the courtroom on RDX-12, is that  
9    right?

10               THE WITNESS:  That's correct, Your Honor.

11               JUDGE CHENEY:  Can we go back to the  
12   demonstrative Exhibit RX -- RDX-2, which I believe are  
13   slides illustrating your testimony.  I'd like to look at  
14   slide 52.  Do you recall giving testimony illustrated by  
15   this slide, Dr. Stone?

16               THE WITNESS:  Yes.

17               JUDGE CHENEY:  I seem to remember you making  
18   some points about what was claimed in the PCT application  
19   identified on this slide.  Do I recall you making a point  
20   about the claims of that PCT application correctly?

21               THE WITNESS:  Yes.  Those relate to computer  
22   controls.

23               JUDGE CHENEY:  And then I also recall, if we  
24   could advance or go back one slide.  Let's try slide 53.  
25   I'm not, I'm not seeing -- did you have a slide where you



1 discussed the provisional application to which the patents  
2 in suit claim priority?

3 THE WITNESS: The PCT application?

4 JUDGE CHENEY: Was there no provisional  
5 application?

6 THE WITNESS: I don't recall, Your Honor.

7 MR. HAILS: Your Honor, there is some 2008  
8 filings, those are all non-provisional filings filed in  
9 parallel. Is that what you're asking about?

10 JUDGE CHENEY: Well, I guess what I'm hearing is  
11 a clarification about my memory that there is no  
12 provisional application to which priority is claimed, is  
13 that right, Counsel?

14 MR. HAILS: Yes. That's correct.

15 JUDGE CHENEY: Okay. Returning to slide 52  
16 where you talked about the PCT priority application. Why  
17 was it important to you to evaluate what was claimed in the  
18 PCT?

19 THE WITNESS: To determine whether there was a  
20 claim of actually putting the dose calibrator on the cart  
21 as a patentable feature as opposed to it was disclosed as  
22 an idea that was incorporated but it was not claimed until  
23 we get all the way down to 2016.

24 JUDGE CHENEY: Why is it important to you that  
25 it -- well, let me, let me back up one step and say, did

1 you find that putting the dose calibrator on the cart was  
2 disclosed in the PCT, but not claimed?

3 THE WITNESS: I did find that it was disclosed  
4 but not claimed. There was a couple of lines that say you  
5 could put the dose calibrator on the cart. Had that been  
6 significant to me I think we would have claimed an  
7 enablement issue but we did not because there was no claim  
8 that that was a patent at that time.

9 JUDGE CHENEY: So in your mind, there is a legal  
10 distinction between what was disclosed and what was claimed  
11 on the issue of priority? That's how I understood this  
12 testimony -- that's the context in which I understood this  
13 testimony about slide 52. Am I misunderstanding?

14 THE WITNESS: I believe there is an issue there,  
15 Your Honor.

16 JUDGE CHENEY: Okay. And how did you come to  
17 this legal understanding about what is disclosed versus  
18 what is claimed and its importance to priority?

19 THE WITNESS: It's a very, it's a broad issue.  
20 I discussed it with the patent attorneys. Know that very  
21 often things are disclosed in a patent that are not  
22 patentable, they are not considered patentable and in this  
23 case it looked like the, the complainants in this case  
24 didn't consider it patentable until they saw it utilized in  
25 someone else's device.

1 JUDGE CHENEY: So in some sense your opinion is  
2 based on your understanding of the inventor's subjective  
3 intent about what the invention is?

4 THE WITNESS: When they recognize that it is an  
5 invention, Your Honor.

6 JUDGE CHENEY: Okay. Can we go to slide 143.  
7 Same Exhibit, RDX-2. And Dr. Stone, do you recall giving  
8 testimony about this international standard disclosed on  
9 this slide?

10 THE WITNESS: Yes, Your Honor.

11 JUDGE CHENEY: I think you testified that the  
12 international standard requires a user to track  
13 consumables. Is that right?

14 THE WITNESS: Yes, Your Honor.

15 JUDGE CHENEY: Is there anything in the standard  
16 that requires a user to use a computer to track  
17 consumables?

18 THE WITNESS: The standard relates to a user  
19 interface, Your Honor. If we go back to the first slide.  
20 The first components there. It's referring to user  
21 interface in order to do that.

22 JUDGE CHENEY: A computer user interface.

23 THE WITNESS: Not necessarily. Whatever  
24 interface there is.

25 JUDGE CHENEY: Could it be a pad of paper with

1 some columns?

2 THE WITNESS: That's typically not considered a  
3 user interface, Your Honor.

4 JUDGE CHENEY: So what is a noncomputerized user  
5 interface?

6 THE WITNESS: I could have a digital system that  
7 actually is, would be noncomputer. It might be something  
8 called a programmable logic array. There are a number of  
9 ways of coming up with a user interface that's a display  
10 that the user, or controls that the user interacts with.

11 JUDGE CHENEY: Dr. Stone, did you offer any  
12 opinion about the field of endeavor of the patented  
13 invention?

14 THE WITNESS: I'm trying to recall from my  
15 report, Your Honor. I believe I would have stated that we  
16 are dealing with a device to administer  
17 radiopharmaceutical, but I don't recall specifically.

18 JUDGE CHENEY: Okay. Those are all the  
19 questions I have. Are there any more questions or redirect  
20 for this witness. It looks like Mr. Hails has some  
21 redirect.

22 MR. HAILS: Yes, sir, there is.

23 REDIRECT EXAMINATION

24 BY MR. HAILS:

25 Q Can we go to slide 52 of the presentation, 223.

1 I think there has been some confusion on the priority  
2 analysis here. Do you recall your testimony talking about  
3 slide 57?

4 A Yes.

5 Q What are the features that drove your priority  
6 analysis for the asserted patents?

7 A The features are related to the openings for the  
8 generator compartment and those openings for the generator  
9 and the waste bottle being oriented upward and the section  
10 elevation for the waste bottle, for example, being at a  
11 greater elevation than the first elevation. Same thing in  
12 the new patents. They have openings that face vertically  
13 upward and they have a shielding compartment that has a  
14 second opening facing vertically in a first opening located  
15 at a lower elevation and the second opening.

16 Q So in 2008 when Bracco filed the first wave of  
17 applications who did Bracco identify as the inventors of  
18 these shielding assembly features?

19 A Mr. Quirico, Mr. Balestracii, Mr. Dorst,  
20 Mr. Krause, Mr. Lokhande, Jacob Childs, Peter Madson, and  
21 Daniel Clements.

22 Q Thank you.

23 Can we go to page 89, please. Can we put up  
24 RX-103. Can you put up paragraph 2 which should be the  
25 first major paragraph.

1                   At first there was a lot of discussion on  
2 cross-examination about the differences between FDG and  
3 rubidium. Are you familiar with that discussion?

4           A        Yes.

5           Q        Are you suggesting to turn Klein's system into  
6 an FDG system?

7           A        Would you repeat that.

8           Q        In your obvious analysis were you suggesting  
9 that you're trying to change Klein's system from a rubidium  
10 system into an FDG system?

11          A        Klein? No.

12          Q        Here, this is the background of the invention  
13 talking about what Tate is directed to. Do you see the  
14 reference to positron emission topography?

15          A        Yes.

16          Q        Are FDG and rubidium variants of positron  
17 emission topography?

18          A        Yes.

19          Q        Can we go to paragraph 7, which is the second  
20 column of the same document.

21                   What kinds of radiopharmaceuticals or  
22 radioisotopes does Tate discuss?

23          A        He discussed a large number such as fluorine 18,  
24 technetium 99, carbon 11, copper 64, gallium 64.

25          Q        You don't have to read it. Do you see rubidium

1 in that list?

2 A He has rubidium-82 in there, Your Honor, as well  
3 as others.

4 Q You had discussed about Tate's teachings for  
5 monitoring saline for locations of dose calibrators and  
6 other kinds of things. Do people run out of saline in both  
7 an FDG system and rubidium system?

8 A They do.

9 Q Do people have to change waste bottles in both  
10 systems?

11 A They do.

12 Q With respect to the dose calibrator that's  
13 described in Tate, do you recall what kind of dose  
14 calibrator Tate used?

15 A Yes. It's an ion chamber.

16 Q And do you recall what kind of design dose  
17 calibrator Klein uses?

18 A Ion chamber with a well force sample.

19 Q With FDG positron for -- in that dose calibrator  
20 in an FDG system, what kind of energy levels are being  
21 measured from the photons?

22 A 511,000 electron volts.

23 Q And what kinds of energy levels are being  
24 measured in a rubidium system using a dose calibrator?

25 A 511,000 electron volts.

1 MR. HAILS: Thank you. No further questions.

2 JUDGE CHENEY: Any other questions for this  
3 witness?

4 MR. DAVIS: No, Your Honor.

5 MR. KOO: Nothing from the staff, Your Honor.

6 JUDGE CHENEY: Thank you, Dr. Stone. You may be  
7 excused.

8 Respondents, call your next witness.

9 MR. BRANDYBERRY: Respondents call Dr. Thomas  
10 Vander Veen.

11 JUDGE CHENEY: Let's go off the record for a  
12 moment.

13 (Discussion off the record.)

14 JUDGE CHENEY: Let's go back on the record.

15 Dr. Vander Veen, please raise your right hand.

16 I will administer the oath.

17 Whereupon,

18 THOMAS V. VANDER VEEN,  
19 was called as a witness, and having been duly sworn, was  
20 examined and testified as follows:

21

22 JUDGE CHENEY: You may be seated.

23 Please proceed when you're ready,

24 Mr. Brandyberry.

25

DIRECT EXAMINATION