- 1 **NEUTROSPEC**TM
- 2 Kit for the Preparation of Technetium (99m Tc) fanolesomab

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- 4 Diagnostic Radiopharmaceutical
- 5 For intravenous use only
- 6 Rx ONLY
- 7 CONTAINS SODIUM HYDROSULFITE

8 **DESCRIPTION**

- 9 NeutroSpecTM [Kit for the Preparation of Technetium (99m Tc) fanolesomab] is a
- 10 radiodiagnostic agent consisting of a murine IgM monoclonal antibody, formulated to be
- 11 labeled with technetium Tc 99m. Each NeutroSpecTM kit contains all the excipients
- 12 needed to reconstitute and to radiolabel this imaging agent with sodium pertechnetate Tc
- 13 99m Injection, USP. The murine monoclonal antibody fanolesomab is produced in
- suspension culture of hybridoma cells. NeutroSpecTM [Technetium (99m Tc)
- 15 fanolesomab] is an *in vivo* diagnostic radiopharmaceutical that can be visualized by
- 16 nuclear medicine instrumentation.
- 17 Each NeutroSpecTM kit contains a single use vial of fanolesomab as a sterile, non-
- pyrogenic, lyophilized mixture of 0.25 mg fanolesomab; 12.5 mg maltose monohydrate;
- 19 0.522 mg sodium potassium tartrate tetrahydrate, USP; 0.221 mg succinic acid; 54 mcg
- stannous tartrate (minimum stannous 7 mcg; maximum total stannous and stannic tin 24
- 21 mcg); 28 mcg glycine, USP; and 9.3 mcg disodium edetate dihydrate, USP. The
- 22 lyophilized powder contains no preservatives and has no US standard of potency.
- When sterile, pyrogen-free sodium pertechnetate Tc 99m Injection, USP in isotonic
- saline (no preservatives) is added to the single use fanolesomab vial, a Tc 99m complex
- of fanolesomab is formed with an approximate pH of 6.2.

26 Physical Characteristics of Technetium Tc 99m

- 27 Technetium 99m decays by isomeric transition with a physical half-life of 6.02 hours.
- The photon that is useful for imaging studies is listed in **Table 1**.

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Table 1. Principal radiation emission data for technetium Tc 99m

Radiation	Mean Percent per Disintegration	Mean Energy (keV)
Gamma-2	89.07	140.5

31 External Radiation

- 32 The specific gamma-ray constant for technetium Tc 99m is 5.4 μC·kg⁻¹·MBq⁻¹·h⁻¹
- 33 (0.78 R/mCi·h) at 1 cm. The first half-value thickness of lead for Tc 99m is 0.017 cm. A
- range of values for the relative attenuation of the radiation emitted by this radionuclide
- 35 that results from the interposition of various thicknesses of lead is shown in **Table 2**. For
- example, the use of a 0.25 cm thickness of lead will decrease the external radiation
- exposure by a factor of 1,000.

Table 2. Radiation attenuation by lead shielding

Lead Shield Thickness (cm)	Coefficient of Attenuation
0.017	0.5
0.08	0.1
0.16	0.01
0.25	0.001
0.33	0.0001

To correct for physical decay of this radionuclide, the fractions that remain at selected time intervals after the time of calibration are shown in **Table 3**.

Table 3. Physical decay chart—technetium Tc 99m half-life 6.02 hours

Hours	Fraction Remaining	Hours	Fraction Remaining
0*	1.00	7	0.45
1	0.89	8	0.40
2	0.79	9	0.36
3	0.71	10	0.32
4	0.63	11	0.28
5	0.56	12	0.25
6	0.50	18	0.13

*Calibration Time (time of preparation)

CLINICAL PHARMACOLOGY

Pharmacodynamics

- 47 Fanolesomab is directed against the carbohydrate moiety 3-fucosyl-*N*-acetyllactosamine
- 48 that defines the cluster of differentiation 15 (CD15) antigen. NeutroSpec[™] [Technetium
- 49 (99m Tc) fanolesomab] radiolabels human white blood cells and myeloid precursors.
- The CD15 antigen is expressed on the surface of polymorphonuclear neutrophils (PMNs),
- eosinophils and monocytes. Monocytes and eosinophils constitute approximately 5% of
- 52 circulating leukocytes; therefore, most of the circulating blood cellular activity resides on
- 53 PMNs. In blood cell fractions isolated from healthy volunteers who had received
- NeutroSpecTM, radioactivity was associated with PMNs (25%) or plasma (72%) when
- measured one hour after injection. The binding of fanolesomab to its antigenic sites on
- human PMNs has an apparent $K_d = 1.6 \times 10^{-11} \,\mathrm{M}$.
- 57 Cross-reactivity studies indicate the presence of CD15 antigenic sites on many human
- 58 tissues.

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Pharmacokinetics

- 60 In a study of 10 healthy volunteers, following intravenous injection of NeutroSpecTM,
- blood concentrations of radioactivity decreased rapidly with an initial half-life of 0.3
- 62 hours and a second phase half-life of approximately eight hours. Whole-body
- scintigraphy at two hours post-injection indicated that the liver had the highest

- radioactivity uptake and retention (50% of the injected dose), followed by the kidney,
- spleen and red marrow. Over the 26–33 hours after injection, 38% of the injected dose of
- radioactivity was recovered in urine.

CLINICAL STUDIES

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- A multicenter, single-arm study evaluated 200 patients (5 to 86 years of age) with
- 69 equivocal signs and symptoms of appendicitis defined as absence of one or more of the
- 70 following: periumbilical pain migrating to right lower quadrant (RLQ), gradual onset of
- pain, increasing intensity of pain over time, pain aggravated by movement and coughing,
- McBurney's point tenderness, referred tenderness to RLQ with palpation in other
- quadrants, abdominal muscular spasm with RLQ tenderness, temperature > 101° F, white
- blood cell count > 10,500/mm³. Readers blinded to clinical information (except for age,
- 75 gender and body habitus) assessed the diagnosis of appendicitis by NeutroSpecTM
- 76 imaging. The diagnosis by the blinded readers was compared with a final clinical
- diagnosis based upon a surgical pathology report (in cases that proceeded to
- appendectomy) or upon two weeks of follow-up (in cases without surgical intervention).
- 79 The study investigators had access to other diagnostic modalities (e.g., CT scan and
- 80 ultrasound) and were instructed not to rely on NeutroSpecTM imaging for their diagnosis
- of appendicitis. Appendicitis prevalence in this study was 30%. The image evaluation
- was limited to the assessment of the planar images performed in specified projections at
- 83 defined time points and single photon emission tomography was not used to assess
- 84 performance in this study.
 - The performance rates for the diagnosis of appendicitis by the blinded readers and by the clinical investigators are shown in **Table 4.**

Table 4. Diagnostic performance of NeutroSpecTM

		Performance Rates (n=200)	
Evaluation	Blinded Readers	Study Investigators	
	percentages (95%CI)	percentages(95%CI)	
Sensitivity	75 (62, 85)	91 (80, 97)	
Specificity	93 (87, 97)	86 (79, 91)	
Accuracy	87 (82, 92)	87 (81, 91)	
Positive Predictive Value	82 (69, 91)	74 (62, 84)	
Negative Predictive Value	90 (84, 94)	96 (90, 99)	

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- 90 In a supportive single-arm, two-center study of the detection of appendicitis in 56 patients
- of whom 50% had a final diagnosis of appendicitis, the diagnostic performance of
- 92 NeutroSpecTM was similar to the performance observed in the larger study.

93 Other intra-abdominal conditions

- Among 30 study patients with other types of intra-abdominal infection (surgical and non-
- 95 surgical), 13 scintigrams were read as positive for appendicitis.

96 INDICATIONS AND USAGE

- 97 NeutroSpecTM [Technetium (99m Tc) fanolesomab] is indicated for scintigraphic imaging
- 98 of patients with equivocal signs and symptoms of appendicitis who are five years of age
- 99 or older.

100 CONTRAINDICATIONS

- NeutroSpecTM should not be administered to patients who are hypersensitive to any
- murine proteins or other component of the product.

103 WARNINGS

104 **Hypersensitivity Reactions**

- Allergic reactions, including anaphylaxis, can occur in patients who receive murine
- antibodies such as fanolesomab.
- 107 CenolateTM Ascorbic Acid, USP injection (diluent) contains sodium hydrosulfite, a sulfite
- that may cause allergic reactions, including anaphylaxis. Serious hypersensitivity
- reactions were not observed in the 523 patients who received NeutroSpec[™] in the clinical
- studies. Emergency resuscitation personnel and equipment for the treatment of
- 111 hypersensitivity reactions should be immediately available during administration of this
- agent.

113 **PRECAUTIONS**

114 **Repeat Administration**

- NeutroSpecTM has not been studied in repeat administration to patients. Murine
- monoclonal antibodies are frequently immunogenic. The development of human anti-
- mouse antibodies (HAMA) can alter the pharmacokinetics, biodistribution, safety, and
- imaging performance properties of the administered agent.

119 Use in Patients with Neutropenia

- 120 The biodistribution and the imaging performance of NeutroSpecTM in neutropenic patients
- have not been studied. NeutroSpecTM induces transient neutropenia and a downward shift
- in white blood cell counts. (See ADVERSE REACTIONS Laboratory Values). The
- safety and effectiveness of NeutroSpecTM in patients with neutropenia have not been
- established.

125 General Use and Handling

- NeutroSpec[™] [Technetium (99m Tc) fanolesomab], like other radioactive medical
- products, must be handled with care and appropriate safety measures should be used to
- minimize radiation exposure to clinical personnel. Care should also be taken to minimize
- radiation exposure to the patient consistent with proper patient management.
- Radiopharmaceuticals should be used by or under the control of personnel who are
- qualified by specific training and experience in the safe use and handling of
- radionuclides, and whose experience and training have been approved by the appropriate
- governmental agency authorized to license the use of radionuclides.

134 Information for patients

- Murine monoclonal antibodies such as fanolesomab are foreign proteins and their
- administration can induce hypersensitivity reactions. Patients should be informed that the
- use of this product could affect their future use of other murine based products, and
- should be advised to discuss prior use of murine antibody based products with their
- health care provider.
- To minimize the radiation-absorbed dose to the bladder, adequate hydration should be
- encouraged to permit frequent voiding during the first few hours after injection. To help
- protect themselves and others in their environment, patients should take the following
- precautions for 12 hours after injection. Whenever possible, a toilet should be used,
- rather than a urinal and the toilet should be flushed several times after each use. Spilled
- urine should be cleaned up completely. After each voiding or fecal elimination, patients
- should thoroughly wash their hands. If blood, urine or feces soil clothing, the clothing
- should be washed separately.

148 Carcinogenesis, Mutagenesis, Impairment of Fertility

- 149 Studies have not been conducted to evaluate carcinogenic potential, mutagenic potential,
- or effects on fertility.

151 **Pregnancy**

- 152 Pregnancy Category C. Animal reproductive studies have not been conducted with
- NeutroSpecTM. It is also not known whether NeutroSpecTM can cause fetal harm when
- administered to a pregnant woman or can affect reproductive capacity. NeutroSpecTM
- should not be used during pregnancy unless the potential benefit to the patient justifies
- the potential risk to the fetus.

157 **Nursing Mothers**

- 158 It is not known whether this drug is excreted in human milk. Because many drugs are
- excreted in human milk, caution should be exercised when NeutroSpecTM is administered
- to a nursing woman. Whenever possible, infant formula should be substituted for breast
- milk until the radioactivity has cleared from the body of the nursing woman.

162 **Pediatric Use**

- In clinical studies of NeutroSpecTM, 29 (5%) patients were 5–11 years old and 32 (6%)
- were 12–16 years old. No overall differences in safety or effectiveness were observed
- between these patients and patients in other age brackets, however, this number of
- patients is too few to exclude differences.

167 Geriatric Use

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- In clinical studies of NeutroSpecTM, 64 (12%) patients were 65 years or older. No overall
- differences in safety or effectiveness were observed between these patients and younger
- patients, but this number of patients is too few to exclude differences.

ADVERSE REACTIONS

- 172 The data described below reflect exposure to NeutroSpecTM in 523 patients and normal
- volunteers receiving a mean antibody dose of 121 mcg (33–250 mcg) and a mean

- 174 radioactive dose of 15 mCi (1-33 mCi). The median patient age was 35 years (5-91
- years); 53% of patients were women and 61% of patients were Caucasians.
- 176 Two patients enrolled in studies of post surgical infection or abscess had serious adverse
- events associated with fatality (hypotension and worsening of sepsis). Underlying
- medical conditions may have also contributed to the fatality and the relationship of the
- fatality to NeutroSpecTM cannot be determined.
- Overall, 49 adverse events occurred in 37 (7%) of the 523 patients exposed to
- NeutroSpecTM. Four of these events were classified as severe (hypotension, worsening of
- sepsis, chest pressure and decreased SaO₂ pain). The most frequently reported adverse
- events were flushing (n=10, 2%) and dyspnea (n=5, 1%). Other less common adverse
- events (< 1%) included syncope, dizziness, hypotension, chest pressure, paresthesia,
- nausea, injection site burning/erythema, pain, and headache.
- Because clinical trials are conducted under widely varying controlled conditions, adverse
- reaction rates observed in clinical trials of a drug cannot be directly compared with rates
- in the clinical trials of another drug, and may not reflect the rates observed in practice.
- The adverse reaction information from clinical trials does, however, provide a basis for
- identifying the adverse events that appear to be related to drug use and for approximating
- 191 rates.

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Laboratory Test Values

- 193 NeutroSpecTM induced transient decreases in neutrophil counts in a study of 10 healthy
- volunteers. Neutrophil counts began to decrease within 3 to 5 minutes post-injection and
- returned to pre-injection values within four hours. Downward shifts in neutrophil counts
- have been observed in 18% of patients (28/151). Three of 284 patients were observed to
- develop transient elevations of AST and ALT after NeutroSpec[™] administration.

198 **Immunogenicity**

- 199 The incidence of antibody development in patients receiving NeutroSpecTM has not been
- adequately determined because the assay was not directly quantitative and its ability to
- detect low titers could not be assured. Human anti-mouse antibody (HAMA) response
- 202 following a single NeutroSpecTM administration was evaluated in a total of 54 patients 3-
- 203 16 weeks post injection. None of the patients had a positive HAMA response. In 30
- 204 healthy volunteers who were exposed to two administrations of fanolesomab separated by
- 205 two weeks, fanolesomab induced HAMA response in five volunteers.
- Immunogenicity data are highly dependent on the sensitivity and specificity of the assay.
- Additionally, the observed incidence of antibody positivity in an assay may be influenced
- by several factors, including sample handling, timing of sample collection, concomitant
- 209 medications, and underlying disease. For these reasons, comparison of the incidence of
- antibodies to NeutroSpec[™] with the incidence of antibodies to other products may be
- 211 misleading.

212 **OVERDOSAGE**

213 There is no experience with overdosage in clinical trials.

DOSAGE AND ADMINISTRATION

215	Adults
216	To prepare NeutroSpec TM the reaction vial containing fanolesomab is reconstituted with
217	sodium pertechnetate Tc 99m Injection, USP solution prior to use. (See
218	INSTRUCTIONS FOR PREPARATION).
219	Fanolesomab is not intended for direct administration to the patient without reconstitution
220	and labeling with sodium pertechnetate Tc 99m Injection, USP. NeutroSpec TM
221	[Technetium (99m Tc) fanolesomab] is intended for a single intravenous (IV)
222	administration through an intravenous access that has been demonstrated to be patent,
223	e.g., butterfly, running IV line, or equivalent injection system to assure that no dose
224	infiltration occurs. Following administration, flush the injection line with an appropriate
225	volume of saline to assure administration of the total dose.
226	For imaging, 75 to 125 mcg of fanolesomab is labeled with 10 to 20 mCi (370 to 740
227	MBq) and administered as a single dose of NeutroSpec TM .
228	Planar imaging should be performed using a large field of view camera fitted with a low-
229	energy, parallel-hole, high-resolution collimator. The camera should be positioned so
230	that the lower edge of the liver is at the upper end of the field of view at the midline of
231	the patient.
232	Dynamic image acquisition over the lower abdomen should begin at the time of injection
233	and consist of 10 sequential four-minute images. Following dynamic image acquisition,
234	the patient should ambulate for approximately 10 to 15 minutes and void. Static planar
235	images should then be collected, including supine anterior, posterior, 10–25 degree RAO
236	and LAO views of the lower abdomen, followed by a standing anterior image of the
237	lower abdomen. After the camera has been positioned (as described above), it is
238	recommended that a total of one million counts be collected for the anterior supine
239	image. All remaining images should be collected for the same duration of time required
240	for the anterior supine image.
241	Children (Five years and older)
242	NeutroSpec [™] is administered in a single dose of 0.21 mCi/kg to a maximum of 20 mCi.
243	Recommended imaging times and procedures are the same as for adults.
244	
245	Dose adjustment has not been established in patients with renal insufficiency, in geriatric
246	patients or in pediatric patients under five years of age.
247	Image Interpretation
248	The biodistribution of the NeutroSpec TM radiopharmaceutical is imaged in the blood pool,
249	reticuloendothelial system (liver, spleen, bone marrow), and urinary excretion organs
250	(kidneys and urinary tract). Imaging of the uterus has been noted, consistent with blood
251	pool activity of NeutroSpec TM .
252	In the 200-patient clinical trial (see CLINICAL STUDIES), based on the average of the
253	three blinded reader interpretations, 75% of the 59 true positive cases of appendicitis
254	were identified (range 66-81%).

- Among those with a blinded diagnosis of appendicitis, 76% displayed uptake of
- radiotracer activity in the appendix within 30 minutes following injection and 98% did so
- by 60 minutes following injection.
- 258 In the trial the acquisition of image collection was performed for a 90 minute period. The
- 259 image finding of a persistent or intensifying uptake in the right lower quadrant (appendix
- zone) that is seen before the completion of the entire imaging sequence may be
- considered a positive study, and imaging may be terminated at this time. In the case of a
- 262 negative image finding at 30 and 60 minutes, collection to 90 minutes is recommended
- 263 prior to termination of the study.
- A diagnostic abnormality is characterized by the presence of an irregular, asymmetric
- 265 uptake of radiotracer localized in the right lower quadrant of the abdomen. The abnormal
- localization of radiotracer remains constant or increases in intensity in follow up imaging.

RADIATION DOSIMETRY

- Based on human data, the absorbed radiation dose to an average human adult (70 kg)
- 269 from an intravenous injection of NeutroSpecTM is listed in **Table 5**. The values were
- 270 calculated using the Standard Medical Internal Radiation Dosimetry (MIRD) method.
- The values are listed as rad/mCi and mGy/MBq and assume urinary bladder emptying at
- 4.8 hours. Radiation absorbed dose estimates for children are given in **Table 6.**

Table 5. Absorbed radiation dose in adults (NeutroSpecTM)

Target Organ	rad/mCi	mGy/MBq
Spleen	0.23	0.062
Kidneys	0.19	0.051
Liver	0.18	0.048
Urinary Bladder Wall	0.12	0.032
Heart	0.061	0.017
Gallbladder	0.056	0.015
Upper Large Intestine Wall	0.051	0.014
Adrenal Glands	0.044	0.012
Lungs	0.043	0.012
Thyroid Gland	0.042	0.011
Red Marrow	0.038	0.010
Lower Large Intestine Wall	0.034	0.0091
Bone Surface	0.031	0.0083
Brain	0.0052	0.0014
Testes / Ovaries	0.0039 / 0.019	0.0010 / 0.0052
Total Body	0.019	0.0050

Dose calculations were performed using the standard MIRD method (MIRD Pamphlet No. 1 rev., Soc. Nucl. Med., 1976). Effective dose equivalent was calculated in accordance with ICRP 53 (Ann. ICRP 18, 1-4, 1988) and gave a value of 0.018 mSv/MBq (0.068 rem/mCi).

Table 6. Estimated absorbed radiation dose for a five-year old child

Target Organ	rad/mCi	mGy/MBq
Spleen	0.70	0.19
Kidneys	0.43	0.11
Liver	0.41	0.11
Urinary Bladder Wall	0.27	0.072
Upper Large Intestine Wall	0.21	0.056
Thyroid Gland	0.19	0.052
Lower Large Intestine Wall	0.16	0.042
Heart	0.15	0.041
Gallbladder	0.13	0.036
Red Marrow	0.11	0.030
Lungs	0.11	0.028
Adrenal Glands	0.095	0.026
Bone Surface	0.085	0.023
Testes / Ovaries	0.019 / 0.059	0.0052 / 0.016
Brain	0.0075	0.0020
Total Body	0.049	0.013

Dose calculations were performed using the standard MIRD method based upon biodistribution studies conducted in adults. Effective dose equivalent was calculated in accordance with ICRP 53 and gave a value of 0.047 mSv/MBq (0.17 rem/mCi).

281 INSTRUCTIONS FOR THE PREPARATION OF NEUTROSPEC™

USE ASEPTIC TECHNIQUE THROUGHOUT

- 283 The user should wear waterproof gloves during the entire procedure and while
- withdrawing the patient dose from the NeutroSpecTM vial.
- 285 Transfer Sodium Pertechnetate Tc 99m Injection, USP with an adequately shielded,
- sterile syringe.

- Adequate shielding should be maintained at all times until the preparation is administered
- 288 to the patient, disposed of in an approved manner, or allowed to decay to background
- levels. A shielded, sterile syringe should be used to withdraw and inject the labeled
- 290 preparation.
- Before reconstituting a vial, it should be inspected for cracks and any indication that the
- integrity of the vacuum seal has been lost. The material should not be used if integrity of
- 293 the vacuum seal has been lost. After reconstitution, examine the vial contents for
- 294 particulates and discoloration prior to injection. The material should not be used if
- 295 particulates or discoloration are observed.
- 296 The dose should be injected via an indwelling catheter, butterfly, or equivalent injection
- 297 system to assure that no dose infiltration occurs. Following administration, flush the
- injection line with an appropriate volume of saline to assure administration of the total
- 299 dose.

300 Labeling and Preparation of NeutroSpecTM 301 Required Materials, Not Supplied within the NeutroSpecTM kit: 1. 302 a. Sodium Pertechnetate Tc-99m, USP, oxidant-free 303 b. ITLC-SG Strips, Heat Treated 304 c. Methyl Ethyl Ketone (MEK) 305 d. Developing Chambers - 50 mL disposable tubes 306 Pipettors and tips 307 **Forceps** 308 **Gamma Counter** 309 h. Dose Calibrator 310 Sodium Chloride for Injection, USP 311 Alcohol (or Germicidal) 312 k. Lead Shield 313 1 mL Sterile Syringes 314 m. Water Bath stabilized at 37±1° C 315 316 Remove a fanolesomab reaction vial from refrigerated storage (2 to 8° C) and 2. 317 allow it to come to room temperature (usually 5 to 10 minutes). NOTE: Keep 318 Cenolate ampule refrigerated and protected from light until needed (Step 5). 319 320 3. Swab the rubber stopper of the fanolesomab reaction vial with an appropriate 321 antiseptic and allow the stopper to dry. 322 323 4. Aseptically add 20 to 40 mCi (740 to 1480 MBq) Sodium Pertechnetate Tc 99m 324 Injection, USP in 0.20 to 0.35 mL generator eluate. Gently swirl (**Do not shake**) the vial until the lyophilized product is completely dissolved, ensuring the vial is 325 326 not inverted. 327 328 **Cautionary Notes:** 329 Use only eluate from a technetium Tc 99m generator that was 330 previously eluted within the last 24 hours. 331 Technetium 99m eluate which is more than 8 hours old from the time 332 of elution should NOT be used. The amount of Sodium Pertechnetate Tc 99m Injection, USP used to 333 reconstitute the reaction vial should be determined based on the 334 335 desired radioactive dose and the estimated time of use. If Sodium Pertechnetate Tc 99m Injection, USP must be diluted prior 336 337 to kit reconstitution, only sterile sodium chloride for injection, USP, 338 (without preservatives) should be used. 339 340 5. Incubate at 37° C for 30 minutes. (Shorter incubation times may result in

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inadequate labeling.)

342		
343	6.	Aseptically add sufficient Cenolate [™] [Ascorbic Acid Injection, USP (500
344		mg/mL)] to make the final preparation volume up to 1 mL.
345		
346		Note: Further dilution is not recommended.
347		
348	7.	Assay the product in a suitable calibrator and record the time, date of preparation
349		and the activity of NeutroSpec [™] onto the string tag label and attach to lead
350		dispensing shield ("pig").
351		
352	8.	Each patient should receive a dose of 10-20 mCi of NeutroSpec TM (the final
353		reconstituted product).
354		
355	9.	Discard vials, needles and syringes in accordance with local, state, and federal
356		regulations governing radioactive and biohazardous waste.
357		
358	Recor	mmended Method for Radiochemical Purity Testing
359		
360	1.	After addition of Cenolate™ (Ascorbic Acid Injection, USP) aseptically withdraw
361		approximately 10 μL of the final reconstituted product for Quality Control (QC)
362		testing. Care should be taken not to introduce air into the vial. Use of a shielded 0.5
363		- 1.0 cc syringe with a 25 or 27 gauge needle is recommended.
364		
365	2.	Apply 1 - 5 µL (a drop that has not completely formed on the tip of a 25 - 27 gauge
366		needle) of NeutroSpec [™] 2 cm (origin) from the bottom of an ITLC-SG 1.5 x 10 cm
367		strip and allow the solution to absorb into the strip (approximately 5 seconds).
368		
369	3.	Immediately place the strip, origin side down, in a development chamber containing
370		4 mL methyl ethyl ketone (MEK).
371		
372	4.	Allow the strip to develop until the solvent front is within 0.5 cm of the top of the
373		strip (3 - 5 minutes).
374		
375	5.	Remove the strip using forceps and allow to dry.
376		
377	6.	Cut the strip at the 4 cm mark, place each piece in a separate counting tube and
378		measure the radioactivity associated with each piece.
379		
380	7.	Calculate the % Free Technetium Tc 99m Pertechnetate as follows:
381		
382		% Free Pertechnetate = Radioactivity in Solvent Front Piece x 100%
383		Total Radioactivity in Strip
384		
385		Note: The product should only be used if the percentage of Free Technetium
386		Tc 99m Pertechnetate is $\leq 10\%$.

387	HOW SUPPLIED		
388	NeutroSpec [™] Kit for the Preparation of Technetium (99m Tc) fanolesomab		
389			
390	The NeutroSpec TM kit contains five individual kits each containing:		
391	One	3 mL single use vial of fanolesomab as a sterile, non-	
392		pyrogenic, lyophilized mixture of 0.25 mg fanolesomab;	
393		12.5 mg maltose monohydrate; 0.522 mg sodium potassium	
394		tartrate tetrahydrate, USP; 0.221 mg succinic acid; 54 mcg	
395		stannous tartrate (minimum stannous 7 mcg; maximum	
396		total stannous and stannic tin 24 mcg); 28 mcg glycine,	
397		USP; and 9.3 mcg disodium edetate dihydrate, USP. The	
398		lyophilized powder contains no preservatives and has no	
399		US standard of potency.	
400			
401	One	2 mL ampule Cenolate [™] [Ascorbic Acid Injection, USP	
402		(500 mg/mL)]	
403	_		
404	One	NeutroSpec [™] Package Insert	
405			
406	One	String tag label for NeutroSpec TM vials (reconstituted	
407		product)	
408	STORAGE		
409		[eutroSpec TM kits at 2 to 8° C (36 to 46° F). After labeling	
410	Refrigerate the lyophilized NeutroSpec TM kits at 2 to 8° C (36 to 46° F). After labeling with Sodium Pertechnetate Tc 99m Injection, USP and addition of Cenolate TM (Ascorbic		
411		should be kept at room temperature, 15 to 25° C (46 to 77°	
412		Use appropriate radiation shielding.	
413	1) and ased within <u>barr</u> ingars.	ese appropriate radiation sinciding.	
414	This reagent kit is approved to	for distribution to persons licensed by the U.S. Nuclear	
415	Regulatory Commission to use byproduct material identified in 10 CFR 35.200 or under		
416	an equivalent license issued l		
417	•	•	
419	,	., Bedford, OH 44146	
420	U.S. Patent X,XXX,XXX		
421	US license number 1588		
	by Hospira, Chicago, IL 600	64	
425			
426	Distributed by:		
	•		
	50. E0010, 1110 0010T		

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431
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434 Cenolate is a registered trademark of Hospira.
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