Theragnostics Inc. 529 Main Street, Suite 1107 Boston, MA 02129

IMPORTANT PRESCRIBING INFORMATION

January 29, 2018

Subject: Temporary importation of Kit for the Preparation of Technetium Tc99m Succimer Injection to address drug shortage issues

Dear Healthcare Professional,

Due to the current critical shortage of DMSA Kit for the Preparation of Technetium Tc99m Succimer, Theragnostics Inc. (Theragnostics) is coordinating with the U.S. Food and Drug Administration (FDA) to increase the availability of the drug. Theragnostics has initiated temporary importation of DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection into the U.S. market. This product is marketed in Germany and is manufactured in Dresden, Germany by ROTOP Pharmaka GmbH for Theragnostics.

At this time, no other entity except ROTOP Pharmaka GmbH, Germany through its US Agent, Theragnostics, and Theragnostics' distributor, Medi-Physics Inc., dba GE Healthcare, is authorized by the FDA to import or distribute the DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection in the U.S. FDA has not approved ROTOP Pharmaka GmbH's Kit for DMSA Preparation of Technetium Tc99m Succimer Injection product in the U.S.

Effective immediately, and during this temporary period, Theragnostics will offer the following presentation of ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection:

Product	Strength	Size	Marketing
			Authorization #
ROTOP DMSA (Kit for	One vial contains 1.74 mg	5 vials in a	3003663.00.00
the Preparation of powder with the active		carton	Germany
Technetium Tc99m	substance: 1.0 mg succimer		(NDC 71647-001-01)
Succimer Injection)			

The vial and carton labels will display the text, translated to English, as approved via the Marketing Authorization of EEA in Germany. At the end of this letter you will find a product comparison table with the prescribing information in English, as well as images of the labels for your reference.

There are some differences in the labeling between the FDA-approved DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection (GE Healthcare) product and ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection (Theragnostics) product (please see the product comparison tables below). These differences do not alter the favorable risk/benefit of the drug:

- In alignment with current practice, the ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer label does not include a statement under the heading "Pediatric Use" that appears in the GE Healthcare label as follows: "Safety and effectiveness in pediatric patients have not been established."
- Unlike the GE Healthcare label, the ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer label contains pediatric dosing information under the heading "How to Use ROTOP DMSA". Pediatric doses can also be calculated online through the Society of Nuclear Medicine and Molecular Imaging website's Pediatric Injected Activity Tool.
- The ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer label does not state the product is sterile; however, like the GE Healthcare product, ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer is manufactured to be sterile.
- Side effects encountered with use of the ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer within the U.S. can be reported directly to Theragnostics, Inc., at 1-888-286-3848 rather than the foreign site referenced in the label for ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer.

ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection is available only by prescription in the U.S.

Please refer to the package insert for the FDA-approved DMSA Kit for the Preparation of Technetium Tc99m Succimer drug product for full prescribing information.

ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection (Theragnostics) does not contain a barcode. Institutions should manually input the product into their systems. Alternative procedures should be followed to assure that the correct drug product is being used and administered to individual patients.

If you have any questions about the information contained in this letter or the use of ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection (Theragnostics), please contact Theragnostics, Inc., Boston, Massachusetts, 1-617-286-7479, 9:00 AM to 5:00 PM Eastern time.

To place an order for ROTOP DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection (Theragnostics), please contact Medi-Physics Inc., dba GE Healthcare, 1-800-292-8514, 8:00 AM to 6:30 PM Eastern time; Email address: <u>CUSTSVSNUCLEAR@ge.com</u>.

To report adverse events or quality problems associated with the use of this product, please call Theragnostics, Inc., Boston, Massachusetts, 1-888-286-3848

CONTACT NUMBERS: Please use the following contact numbers as appropriate: Phone: 1-617-286-7479 Fax: 1-617-398-6337

Adverse reactions or quality problems experienced with the use of this product may be reported to the FDA's MedWatch Adverse Event Reporting program either online, by regular mail or by fax.

- Complete and submit the report **Online**: <u>www.fda.gov/medwatch/report.htm</u>
- **Regular Mail or Fax**: Download form <u>www.fda.gov/MedWatch/getforms.htm</u> or call 1-800-332-1088 to request a reporting form, then complete and return to the address on the pre-addressed form, or submit by fax to 1-800-FDA-0178 (1-800-332-0178)

Sincerely,

Gregory Mullen President & CEO

Attachments:

- 1. Product Comparison Table
- 2. Label Comparison Table
- 3. Vial and Carton Labels

Attachment 1: Product Comparison Table

Compa	arison Table 1: Theragnostics vs. GE H	Iealthcare Reference Product
Characteristics	Reference product: MPI DMSA KIDNEY REAGENT (Kit for the Preparation of Technetium Tc99m Succimer Injection)	Theragnostics' product: Kit for the Preparation of Technetium Tc99m Succimer Injection
Conditions of use	DMSA is indicated for the use as an aid in the scintigraphic evaluation of renal parenchymal disorders.	Theragnostics' Kit is indicated for the use as an aid in the scintigraphic evaluation of renal parenchymal disorders.
Active ingredient	meso-2,3-dimercaptosuccinic acid	meso-2,3-dimercaptosuccinic acid
Inactive	stannous chloride dihydrate	stannous chloride dihydrate
ingredients	ascorbic acid	ascorbic acid
inositol		
	sodium hydroxide	sodium hydroxide
	hydrochloric acid	hydrochloric acid
	nitrogen	nitrogen
Route of Administration	Intravenous	Intravenous
Dosage form	Injection	Injection
Strength	N/A	N/A
Description	Each vial contains a sterile, pyrogen- free freeze-dried mixture of 1.0 mg dimercaptosuccinic acid, 0.42 mg stannous chloride dihydrate [0.38 mg (minimum) stannous chloride dihydrate (SnCl ₂ •2H ₂ O) and 0.46 mg (maximum) total tin expressed as stannous chloride dihydrate (SnCl ₂ •2H ₂ O)], 0.70 mg ascorbic acid, and 50.0 mg inositol. After freeze-drying, vials are sealed under a nitrogen atmosphere with a rubber closure. Sodium hydroxide and hydrochloric acid have been used for pH adjustment. When sterile, oxidant- free, pyrogen-free sodium pertechnetate Tc ⁹⁹ m injection in isotonic saline is combined with the vial contents, following the instructions provided with the kit, a complex is formed. After 10 minutes' incubation the reconstituted solution is ready for intravenous injection.	One vial contains 1.74 mg powder with the active substance, 1.0 mg succimer. The excipients are: stannous chloride dihydrate, ascorbic acid, sodium hydroxide, hydrochloric acid 36% and nitrogen.

Attachment 2: Labeling Comparison Table

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
GE REFERENCE I RODUCT INSERT	DIFFERENCES	This Package Leaflet and Summary of Product
	DMSA English translation note	Characteristics was translated by the manufacturer based on the original German document (Vs. 4), authorized by the German Federal Institute for Drugs and Medicinal Services in November 2014.
		Package Leaflet and Summary of Product Characteristics
DMSA Kit for the Preparation of Technetium Tc99m Succimer Injection	Product name specific for market	ROTOP - DMSA, 1.0 mg Kit for radiopharmaceutical preparation
DIAGNOSTIC - FOR INTRAVENOUS USE	Insert layout	Succimer
	specific to manufacturer; GE layout adjusted to "line" up to sections with ROTOP insert for ease of review German product specific instructions	 Read all of this leaflet carefully before you start using this medicine. Keep this leaflet. You may need to read it again. If you have any further questions, ask your doctor or pharmacist. This medicine has been prescribed for you. Do not pass it on to others. It may harm them, even if their symptoms are the same as yours. If any of the side effects gets serious, or if you notice any side effects not listed in this leaflet, please tell your doctor or pharmacist.
DESCRIPTION Each vial contains a sterile, pyrogen-free freeze- dried mixture of 1.0 mg dimercaptosuccinic acid, 0.42 mg stannous chloride dihydrate [0.38 mg (minimum) stannous chloride dihydrate (SnCl ₂ •2H ₂ O) and 0.46 mg (maximum) total tin expressed as stannous chloride dihydrate (SnCl ₂ •2H ₂ O)], 0.70 mg ascorbic acid, and 50.0 mg inositol. After freeze-drying, vials are sealed under a nitrogen atmosphere with a rubber closure. Sodium hydroxide and hydrochloric acid have been used for pH adjustment. When sterile, oxidant-free, pyrogen-free sodium pertechnetate Tc ⁹⁹ m injection in isotonic saline is combined with the vial	Insert layout and details specific to manufacturer	 In this leaflet: 1. What ROTOP – DMSA is and what it is used for 2. Before you use ROTOP - DMSA 3. How to use ROTOP - DMSA 3. How to use ROTOP - DMSA 4. Possible side effects 5. How to store ROTOP - DMSA 6. Further information 1. WHAT ROTOP – DMSA IS AND WHAT IT IS USED FOR ROTOP - DMSA is a radiodiagnostic pharmaceutical. The kit contains the non-radioactive powder for reconstitution of the [^{99m}Tc]technetium succimer injection solution ([^{99m}Tc]-DMSA). The sodium [^{99m}T]pertechnetat which is needed for the preparation is not part of this kit. After labelling with sodium [^{99m}Tc]technetium pertechnetat solution, ROTOP - DMSA is used for static renal scintigraphy when adequate diagnostics are not possible using other diagnostic procedures (such as ultrasound):

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
OF INSERT Contents, following the instructions provided with the kit, a complex is formed. After 10 minutes incubation the reconstituted solution is ready for intravenous injection. Chemical Name: meso-2,3-dimercaptosuccinic acid SH SH HOOC - C - C - COOH H H HOOC - C - C - COOH H H The succimer component of DMSA consists of more than 90% meso isomer and less than 10% d,1 isomer. PHYSICAL CHARACTERISTICS Technetium Tc99m decays by isomeric transition with a physical half-life of 6.02 hours ¹ . The principal photon that is useful for detection and imaging studies is listed in Table 1. Table 1. Principal Radiation Energy n Disintegratio n Disintegratio Energy n Disintegratio Energy n Mean % / Mean n Disintegratio Energy n <t< td=""><td>Insert layout and details specific to manufacturer</td><td> to identify focal renal parenchymal changes (e.g. in the case of renal infarction) to identify norm variants such as atypical double kidney, small kidney, dysplastic kidney, horseshoe kidney, as well as to identify ectopic kidneys to confirm absence of renal function in multicystic kidneys. </td></t<>	Insert layout and details specific to manufacturer	 to identify focal renal parenchymal changes (e.g. in the case of renal infarction) to identify norm variants such as atypical double kidney, small kidney, dysplastic kidney, horseshoe kidney, as well as to identify ectopic kidneys to confirm absence of renal function in multicystic kidneys.
 INDICATIONS AND USAGE DMSA is to be used as an aid in the scintigraphic evaluation of renal parenchymal disorders. PRECAUTIONS General As in the use of any radioactive material, care should be taken to minimize radiation exposure to the patient consistent with proper patient management and to ensure minimum radiation exposure to occupational workers. DMSA should be used between 10 minutes and 4 hours following reconstitution (see "Preparation" section). Any unused portion should be discarded after that time.		 2. BEFORE YOU USE ROTOP - DMSA Take special care with ROTOP – DMSA ROTOP - DMSA is not suitable for determining global renal function from the DMSA accumulation. In the case of proximal tubulopathies [^{99m}Tc]DMSA does not lead to a sufficient diagnostic renal accumulation. The patient must be well hydrated before and after administration. In order to keep radiation exposure to a minimum, patients must be encouraged to empty their bladders as often as possible during the first hours after the examination. For each patient it should be carefully considered whether the expected diagnostic benefits outweigh the risk linked to radiation exposure. In order to keep

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
Some patients with advanced renal failure may		the radiation dose as low as possible, the
exhibit poor renal intake of Tc99m DMSA. It has		administered activity may not be higher than that
been reported that satisfactory images may be		required for eliciting the diagnostic information.
obtained in some of these patients by delaying		
imaging for up to 24 hours.		Radiopharmaceuticals may be received, used and
		administered only by authorised persons in areas
The contents of the kit vials are intended only for		specially designated for this purpose. The
use in the preparation of DMSA Injection and are		manipulation and use of these products is subject to
not to be directly administered to the patient.		the regulations of the local supervisory authority
		and/or requires appropriate permission.
The contents of the kit vials are not radioactive.		Contraindications
However, after Tc99m is added, adequate shielding		ROTOP-DMSA should not be used in case of
of the final preparation must be maintained.		hypersensitivity to the active substance or to any of
Dediantemperations about the word only her		the excipients listed in section 6.
Radiopharmaceuticals should be used only by		
physicians who are qualified by training and experience in the safe use and handling of		Using other medicines
radionuclides and whose experience and training		
have been approved by the appropriate government		Chemotherapeutic agents such as methotrexate,
agency authorized to license the use of		cyclophosphamide and vincristine can alter the
radionuclides.	Insert layout and	biodistribution of [99mTc]DMSA.
	details specific to	
Carcinogenesis, Mutagenesis, Impairment of	manufacturer	Shifting the acid/base balance, e.g. through
Fertility		ammonium chloride or sodium hydrogen carbonate,
No long term animal studies have been performed		effects in vivo a change in the valence of the
to evaluate carcinogenic potential, mutagenic		[99mTc]DMSA complex and in turn a lower
potential, or whether technetium Tc99m succimer		accumulation in the renal cortex with simultaneous
injection affects fertility in males or females.		strong accumulation in the liver and rapid urine
		excretion. Mannitol leads to dehydration and in turn
		to a reduction in the extraction of [^{99m} Tc]DMSA.
Pregnancy Category C		In the case of renal artery stenosis, ACE inhibitors
Animal reproduction studies have not been		can lead to a reversible insufficiency of the tubular
conducted with technetium Tc99m succimer		function and in turn to a reduced accumulation of
injection. It is also not known whether technetium		[^{99m} Tc]DMSA as a result of the reduction in filtration
Tc99m succimer injection can cause fetal harm		pressure in the affected kidney.
when administered to a pregnant woman or can		
affect reproduction capacity. Technetium Tc99m		If high doses of other chelating agents are injected at
succimer injection should be administered to a		the same time, the stability of the [99mTc]DMSA
pregnant woman only if clearly needed.		DMSA may be influenced, thus effecting a change in
Ideally, examinations using radiopharmaceuticals,		kinetics.
especially those elective in nature, of a woman of		
child bearing capability should be performed during		Pregnancy and lacation
the first few (approximately 10) days following the		
onset of menses.		<u>Pregnancy</u> : No data on the clinical use of
		[^{99m} Tc]DMSA with pregnant women is available. If
Nursing Mothers		it is necessary to administer a radiopharmaceutical
Technetium Tc99m is excreted in human milk		product to a woman of child-bearing age, she must
during lactation; therefore, formula feedings should		have a pregnancy test first.
be substituted for breast feedings.		If a woman has missed a newigit it would be a set
Ŭ		If a woman has missed a period, it must be assumed that she is program. In some of doubt, indication
Pediatric Use		that she is pregnant. In case of doubt, radiation
Safety and effectiveness in pedriatric patients have		exposure must be reduced to the minimum amount required to acquire the needed clinical information.
not been established.		In this case, alternative investigative methods must
		be considered that do not use ionising radiation.
Geriatric Use		se considered that do not use follising fadiation.

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
Clinical studies of DMSA did not include sufficient		Radiopharmaceutical examinations of pregnant
numbers of subjects age 65 and over to determine		women also expose the foetus to radiation. For this
whether they respond differently from younger patients. Other reported clinical experience has not		reason, [^{99m} Tc]DMSA may only be used if there is a vital indication and if the expected benefit
identified differences in responses between the		outweighs the risk to mother and child.
elderly and younger patients. In general, dose		
selection for an elderly patient should be cautious		Lactation: Before administering [99mTc]DMSA to a
usually starting at the low end of the dosing range,		breast-feeding mother, it must be considered
reflecting the greater frequency of decreased		whether the investigation could also be delayed
hepatic, renal or cardiac function, and of concomitant disease or other drug therapy.		until the mother has ceased breast-feeding and as to whether using a radiopharmaceutical is the most
concomitant disease of other drug therapy.		appropriate examination method, bearing in mind
This drug is known to be substantially excreted by		the secretion of activity into breast milk. If
the kidney, and the risk of toxic reactions to this		administering [^{99m} Tc]DMSA is deemed necessary,
drug may be greater in patients with impaired renal		breast-feeding must be interrupted for at least 12
function. Because elderly patients are more likely		hours, and the expressed breast milk discarded.
to have decreased renal function, care should be	Insert layout and	
taken in dose selection, and it may be useful to monitor renal function.	details specific to manufacturer	Driving and using machines
monitor renar function.	manufacturer	Effects on the ability to drive or use machines have
		not been described.
		Precautions for avoiding hazards for the environment
		Radiopharmaceuticals must be prepared and used by
		the user under precautions for the protection from
		ionizing radiation and taking pharmaceutical quality
		standards into account. In accordance with the
		guidelines for Good Pharmaceutical Manufacturing Practice, work must be done under aseptic
		conditions.
		Patients treated with radiopharmaceuticals pose a
		risk for other persons based on external radiation
DOGACE AND ADMINISTRATION		exposure or contamination due to spilling urine,
DOSAGE AND ADMINISTRATION The suggested dose range for slow I.V.		vomiting, etc. For this reason, the precautionary measures provided by the national radiation
administration to be employed in the average		protection regulations must be observed.
patient (70 kg) for renal parenchymal imaging is		Contamination brought about by radioactivity that
74-222 MBq, 2-6 mCi technetium Tc99m succimer		has been excreted by the patient must be avoided.
injection.		
The product must be used between 10 minutes to 4		3. HOW TO USE ROTOP - DMSA
hours following preparation (see "Preparation"		Single intravenous use after preparation with
section). Acceptable renal images may be obtained		sodium [^{99m} Tc]pertechnetate solution.
beginning 1 to 2 hours post injection. Any unused		Adults are given 0.3 to 1.0 mg succimer and
portion should be discarded after that time.		activities of 70 MBq.
The patient dose should be measured by a suitable		Scintigraphic examinations should not be carried
radioactivity calibration system immediately prior		Scintigraphic examinations should not be carried out until at least 1 hour after application; waiting 3
to administration.		hours is preferable. In the case of very poor renal
		function, waiting periods of up to 6 hours should be
Do not use after the expiration date stated on the		observed. The patient must be well hydrated.
label. The components of the kit are supplied sterile		
and pyrogen-free. Aseptic procedures normally		 ٥

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
employed in making additions and withdrawals from sterile, pyrogen-free containers should be used during addition of sodium pertechnetate Tc99m injection solutions and during the withdrawal of doses for patient administration.		<u>Children</u> The recommendation of the Paediatric Task Group of the European Association of Nuclear Medicine (EANM) of 1990 lists the paediatric dose scaled to body weight as a fraction of the adult dose:
Parenteral drug products should be inspected visually for particulate matter and discoloration		3 kg = 0.1 22 kg = 0.50 42 kg = 0.78
prior to administration.		4 kg = 0.14 24 kg = 0.53 44 kg = 0.80
		6 kg = 0.19 26 kg = 0.56 46 kg = 0.82
		8 kg = 0.23 28 kg = 0.58 48 kg = 0.85
		10 kg = 0.27 30 kg = 0.62 50 kg = 0.88
		12 kg = 0.32 32 kg = 0.65 52 - 54 kg = 0.90
	T	14 kg = 0.36 $34 kg = 0.68$ $56 - 58 kg = 0.92$
	Insert layout and details specific to	16 kg = 0.40 36 kg = 0.71 60 - 62 kg = 0.96
	manufacturer	18 kg = 0.44 38 kg = 0.73 64 - 66 kg = 0.98
		20 kg = 0.46 40 kg = 0.76 68 kg = 0.99
WARNINGS None. ADVERSE REACTIONS Rare instances of syncope, fever, nausea and maculopapular skin rash have been reported. CONTRAINDICATIONS None known.		Activity of less than 20 % (15 MBq) of the adult dose generally does not allow a satisfactory assessment to be derived from the examination.If you use more ROTOP – DMSA than you shouldDue to the low amounts of substances used, overdosage in the pharmacological sense is not expected. Exposure to radiation resulting from an overdosage of radioactivity can be reduced by forced diuresis.4. POSSIBLE SIDE EFFECTSAs all medicinal products, ROTOP - DMSA can cause side effects, although not everybody gets them.For assessing the side effects the frequency is classified as follows:Veryobserved in more than 1 patients in 10Commonobserved in less than 1 patient in 100, but more than 1 patient in 1,000Rareobserved in less than 1 patient in 1,000, but more than 1

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
		patient in 10,000
		Very rare observed in less than 1 patient
		in 10,000 or not known
		In very rare cases (< 0.01 %) after intravenous
		injection of the ready-to-use solution,
		hypersensitivity reactions have occurred such as
		locally confined or general rashes, itching, drop in
		blood pressure, headache, dizziness, nausea and vomiting. Reactions can occur up to 24 hours after
		the injection.
		Although such reactions are very rare and usually
		very minor, appropriate instruments and
		medications for immediate treatment of allergic reactions (adrenaline, corticosteroids and
		antihistamines) should be within reach for possible
		emergency treatment at all times.
		Since the administered amounts of active substances
	Insert layout and	are very low, the risks of use are mainly related to radiation exposure. Ionising radiation can cause
	details specific to	cancer and genetic mutations.
	manufacturer	Since most radiopharmaceutical examinations are
		conducted with low effective radiation doses of less
		than 20 mSv, the probability of such effects occurring is expected to be low.
		occurring is expected to be low.
		The effective radiation dose is 0.62 mSv when the
		maximum recommended activity of this medicinal
		product is applied.
		Reporting of side effects
		If you notice any side effects please contact your
		nuclear physician responsible for supervising the
		administration. This also applies to any side effects
		not listed in this leaflet.
		You can also report any side effects directly to:
		Bundesinstitut für Arzneimittel und
		Medizinprodukte, Abt. Pharmakovigilanz, Kurt-
HOW SUPPLIED		Georg-Kiesinger Allee 3, D-53175 Bonn, website: <u>http://www.bfarm.de</u> .
Kit Contents		<u>http://www.totatalade</u> .
5 Vials containing a freeze-dried mixture of 1.0 mg		By reporting side effects you can help provide more
dimercaptosuccinic acid, 0.42 mg stannous chloride dihydrate [0.38 mg (minimum) stannous chloride		information on the safety of this medicine.
dihydrate [0.38 mg (mmmull) standous chloride dihydrate (SnCl ₂ •2H ₂ O) and 0.46 mg (maximum)		5. HOW TO STORE ROTOP - DMSA
total tin expressed as stannous chloride dihydrate		. HOW TO STORE ROTOL - DIISA
(SnCl ₂ •2H ₂ O)], 0.70 mg ascorbic acid, and 50.0 mg		Keep out of the reach and sight of children.
inositol.		Do not use this medicinal product after the expiry
5 Labels 1 Package Insert		date stated on the label.
		Storage conditions
NDC 017156-525-01		

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
Storage Store the kit at 2°-8°C (36°-46°F) and protect from light.		Store refrigerated (2 to 8 °C) in the original package. Radiopharmaceuticals must be stored in accordance with the regulations for radioactive protection and in particular be kept from unauthorised access. Shelf life after opening and reconstitution The product labelled with [^{99m} Tc]technetium can be
		 injected within 4 hours after reconstitution and has to be stored at room temperature (15–25 °C) during this time. 6. FURTHER INFORMATION
		What ROTOP – DMSA contains
		One vial contains 1.74 mg powder with the active substance:
		1.0 mg succimer
	T . 1 . 1	The other ingredients are:
This reagent kit is approved for use by persons	Insert layout and details specific to	Stannous chloride dihydrate
licensed by the Illinois Emergency Management	manufacturer	Ascorbic acid
Agency pursuant to 32 Ill. Code Adm. Section, Section 330.260(a) and 335.4010 or under		Sodium hydroxide Hydrochloric acid 36%
equivalent licenses of the U.S. Nuclear Regulatory		Nitrogen
Commission, or an Agreement State.		
Manufactured for:		What ROTOP – DMSA looks like and contents of the pack:
GE Healthcare Medi-Physics, Inc. 3350 North Ridge Avenue		The package consists of a carton with 5 vials.
Arlington Heights, IL 60004 1-800-633-4123 (Toll Free)		Marketing Authorisation Holder and Manufacturer
By:		ROTOP Pharmaka GmbH,
GE Healthcare Ltd.		Bautzner Landstr. 400,
Little Chalfont, HP7 9NA, UK		01328 Dresden,
GE and the GE Monogram are trademarks of		Germany Tel: 0049 + (0) 351 – 26 310 210
General Electric Company.		Fax: $0049 + (0) 351 - 26 310 210$
		e-mail: service@rotop-pharmaka.de
43-4349H L/2331/04		This medicinal product is sufficient in the
L/2331/04		This medicinal product is authorised in the Member States of the EEA under the following
Revised February 2006		names:
		Germany: ROTOP - DMSA
CLINICAL PHARMACOLOGY		This leaflet was last approved in May 2017.
After intravenous administration, technetium		
Tc99m succimer injection is distributed in the		
plasma, apparently bound to plasma proteins.		

GE REFERENCE P	DODUCT INSEDT	DIFFERENCES	ROTOP-DMSA INSERT
There is negligible activity		DIFFERENCES	The following information is intended for medical
The activity is cleared from			or healthcare professionals only:
time of about 60 minutes a			or nearthcare professionals only.
renal cortex. Approximate			PHARMACOLOGICAL PROPERTIES
excreted in the urine within			THARMACOLOGICAL TROTERTIES
about 20% of the dose is c			Pharmacodynamic properties
kidney.	oncentrated in caen		Pharmacotherapeutic group: Diagnostic
Kidiley.			radiopharmaceutical for renal diagnostics (ATC:
EXTERNAL RADIATIO)N		V09CA02). Based on current research, for the low
The specific gamma ray co			amounts of substances used for imaging techniques
Tc99m is 0.78 R/hr-mCi a			no clinically relevant pharmacodynamic effects of
value layer is 0.017 cm of			[^{99m} Tc]DMSA are expected.
			[IC]DMSA are expected.
of the radiation exposure f of this radionuclide, the us			Dhormooolinatic properties
			Pharmacokinetic properties
of Pb will attenuate the rad	nation enlitted by a factor		After introvenous injection within 5 minutes over
of about 1,000.			After intravenous injection, within 5 minutes over 70% of the 199 mToIDMSA is bound to the 2
Table 1 D - 2-4	Attonuotion ha		70% of the [99m Tc]DMSA is bound to thea-2
Table 2. Radiation Lood Shielding	Attenuation by		microglobulin fraction in blood plas ma. Binding to
Lead Shielding			erythrocytes may be disregarded. One hour post
Shield Thickness	Coefficient of		injection, 25% of the radiopharmaceutical is already located in the renal cortex and only 30% remains in
(Pb) cm	Attenuation	Insert layout and	
0.02	0.5	details specific to	the plasma. Approx. 10% appears in the urine.
0.08	0.1	manufacturer	In boolthy persons, the plasma algorance of
0.16	0.01	manufacturer	In healthy persons, the plasma clearance of [^{99m} Tc]DMSA amounts to approx. 10 ml/min.
0.25	0.001		(scaled to 1.73 sqm body surface). After approx. 3
0.33	0.0001		hours, the maximum renal accumulation is reached.
To correct for physical dec			In healthy persons, at this point approx. 50% of the
the fractions that remain at	t selected intervals after		radiopharmaceutical is located in the renal cortex,
the time of calibration are	shown in Table 3.		approx. 20% remains in the plasma and just under
			10% in the liver and muscles. Within 24 hours,
Table 3. Physical De	cay Chart: Tc99m,		approx. 30% is excreted with the urine.
half-life 6.02 hours			[^{99m} Tc]DMSA accumulates in the pars recta and
Hour Fraction	Hour Fraction		convoluta of the proximal renal tubules – most
s Remaining	s Remaining		likely due to peritubular reabsorption. On an
0* 1.000	7 0.447		intracellular level, the majority of the [^{99m} Tc]DMSA
1 0.891	8 0.398		
2 0.794	9 0.355		is bound to a soluble protein in the cytosol. This mechanism, which has not yet been explained in
3 0.708	10 0.316		detail, is disrupted in the case of proximal
4 0.631	11 0.282		tubulopathies (such as nephritides or the Fanconi
5 0.562	12 0.251		syndrome), which can be recognised by the
6 0.501			increased plasma clearance of [^{99m} Tc]DMSA and
* Calibration Time	1		low renal accumulation.
			low renar accumulation.
			Toxicological properties
DISPOSAL			Toxicological properties
Any unused portion of the	Tc99m-labeled kit must		Due to the low amounts of DMSA and stannous
be stored and disposed of i			chloride contained in the kit, toxic effects brought
conditions of NRC radioad			
pursuant to 10 CFR Parts 2			about by the substances are not expected if used
conditions pursuant to Agr			according to directions. Data on investigations on
or other regulatory agency			reproduction toxicity as well as on mutagenicity and
use of radionuclides.	autionized to neense the		cancerogenity are not available.
use of futionactions.			Special propositions for dispersal and further
1		1	
			Special precautions for disposal and further
			directions for handling

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT					
The unlabeled residual materials may be discarded in ordinary trash, provided that the vials and syringes read background with an appropriate low- range survey meter. It is suggested that all identification labels be destroyed before discarding.	abeled residual materials may be discarded ary trash, provided that the vials and s read background with an appropriate low- urvey meter. It is suggested that all		The empty package is considered to be regular waste if the permitted level for [99m Tc]technetium is not exceeded (≤ 0.5 Bq/g or 0.5 Bq/cm ²). Particulars indicating radioactivity must be removed prior to disposing of the non-radioactive waste and must be destroyed separately. Radioactive waste must be disposed of as provided by law.				
RADIATION DOSIMETRY		MARKET 3003663.0	TING AU 0.00	THORI	ISATIO		IBER
The estimated absorbed radiation doses ^{2,3} to an average adult (70 kg) are shown in Table 4.		DATE OF RENEWA 24/11/2005	L OF TI				N
Table 4. Absorbed Radiation Dose Tissue mGy / rads / 222 MBq 6 mCi Bladder Wall 4.2 0.42		DOSIME7					
Kidneys (total) 37.8 3.78 Renal Cortices 51.0 5.10 Liver 1.9 0.19 Bone Marrow 1.3 0.13	Insert layout and details specific to manufacturer	Radiation According following 1	ICRP	publicati			1) the
Ovaries 0.8 0.08 Testes 0.4 0.04		Absorbe	ed dose pe	r unit of (mGy/M		adminis	tered
Total Body 0.9 0.09		Organ	Adult s	15 years	10 years	5 years	1 year
² Method of Calculation: A schema for Absorbed- Dose Calculations for Biologically Distributed Radionuclides, Supplement No. 1, MIRD Pamphlet		Adrenals Bladders wall	0.012 0.018	0.016	0.024	0.035	0.060
No. 1, J. Nucl. Med., p. 7, 1968. ³ Biological Data: Arnold, R.W; Subramanian, G.;		Bone surface	0.0050	0.006	0.009	0.014	0.026
McAfee, J.G.; Blair, R.J.; Thomas, F.D.;		Brain	0.0012	0.001 5	0.002 5	0.004 0	0.007
Comparison of Tc99m complexes for renal imaging, J. Nucl. Med., 16, pp. 357-367, 1975.		Breast	0.0013	0.001 8	0.002 8	0.004 5	0.008
		Gall bladder	0.0083	0.010	0.014	0.022	0.031
		Stomach wall	0.0052	0.006	0.010	0.014	0.020
		Colon	0.0050	0.006	0.010	0.014	0.024
		Intestine	0.0043	0.005 5	0.008	0.012	0.020
		Upper large intestine	0.0050	0.006 4	0.095	0.014	0.023
		Lower large intestine	0.0035	0.004 3	0.006 5	0.009 6	0.016
		Heart	0.0030	0.003 8	0.005 8	0.008 6	0.014
		Kidneys Liver	0.18 0.0095	0.22 0.012	0.30 0.018	0.43 0.025	0.76 0.041
		Lungs	0.0025	0.003	0.005	0.008	0.015

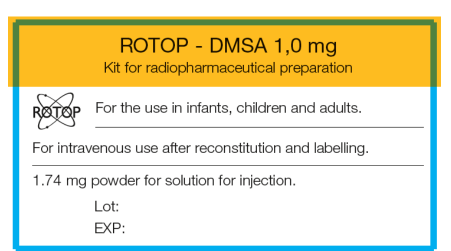
GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT					
				5 0.003	2 0.005	0.007	
		Muscles	0.0029	6	2	7	0.014
		Oesopha gus	0.0017	0.002	0.003	0.005 4	0.009 4
		Ovaries	0.0035	0.004 7	0.007	0.011	0.019
		Pancreas	0.0090	0.011	0.016	0.023	0.037
		Red marrow	0.0039	0.004	0.006	0.009 0	0.014
		Skin	0.0015	0.001	0.002	0.004	0.008 5
		Spleen	0.013	0.017	0.026	0.038	0.061
		Testes	0.0018	0.002	0.003 7	0.005 3	0.010
		Thymus	0.0017	0.002	0.003	0.005 4	0.009 4
		Thyroid	0.0015	0.001 9	0.003	0.005 2	0.009 4
		Uterus	0.0045	0.005 6	0.008	0.011	0.019
		Remaini ng organ	0.0029	0.003	0.005	0.007 7	0.014
		Effective Dose per unit of activity administ ered (mSv/M Bq)	0.0088	0.011	0.015	0.021	0.037
		In an adult (70 kg), after intravenous injection of MBq (maximum dose) [^{99m} Tc]DMSA, the effect dose is approx. 0.62 mSv. The absorbed dose in target organ kidney is approx. 12.6 mGy and in critical organ bladder wall 1.26 mGy. Radiophysical Properties					
 Preparation The following directions must be carefully followed for optimum preparation of technetium Tc99m succimer injection: Note: Use aseptic procedures throughout and take precautions to minimize radiation exposure by the use of suitable shielding. Waterproof gloves should be worn during the preparation procedure. Place one of the vials in a suitable shielding container and swab the closure with a bacteriostatic swab. Using a 10 mL sterile syringe, inject an appropriate amount (see notes 1 and 2) of the eluate from a Tc99m generator into the shielded vial. Before removing the syringe from the vial withdraw an equivalent volume 		[99mTc]ted [⁹⁹ Mo/ ^{99m} T gamma rad a half-life of turn decays to a long considered INSTRUC RADIOPH Instruction [^{99m} Tc]tech prepared u	c] sterile iation wi of 6.02 ho s to stable half-life to be sta TTIONS HARMA n for lab anetium s nder steri	generat th an ene urs to ^{[99} e [⁹⁹ Ru]r of 214,(ble. FOR PH CEUTIC elling uccimer le condi	rgy of 1 Tc]tech utheniur 00 year CALS injectio tions wit	decays 1 40/142 h netium, n; Howe rs, ⁹⁹ Tc ATION n solution	ceV with which in ever, due itself is OF
		[^{99m} Tc]pert	ecimetate	mjectio	Soluti	on (Euro 	-

	GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DM	MSA INSERT
	of nitrogen from the space above the solution		Pharmacopoeia quality 4.	
	to normalize the pressure in the vial.		directly before use. Oxyg	enation must be avoided.
3.	Carefully invert the vial a few times until the			
	powder is completely dissolved.		Place the vial with powde	
4.	Assay the total activity, complete the label			ce and disinfect the stopper
~	provided and attach to the vial.		(allow disinfectant to dry	
5.	Incubate the vial for at least 10 minutes at		smallest possible cannula	
6	room temperature. Use the preparation between 10 minutes and 4		sodium [^{99m} Tc]technetium	a pertechnetate solution aq to the vial. Use the same
6.	hours following reconstitution.		syringe to withdraw the a	
	nours following reconstitution.		from the vial for pressure	
Not	e:			compensation
	Not more than 1.48 GBq, 40 mCi technetium-		Lightly shake the vial in o	order to completely
	99m in a volume of 1-6 mL should be added to		dissolve the powder. The	
	the vial.		moistened as well. After	10 minutes reaction time,
2.	Before reconstitution, the eluate may be			ity. If needed, the finished
	adjusted to the correct radioactive			diluted with sterile isotonic
	concentration by dilution with preservative-		sodium chloride to a total	volume
2	free, non-bacteriostatic saline for injection.		of up to 10 mL.	
3.	The use of technetium-99m solution complying with the specifications prescribed by the USP		Quality Control	
	Monograph on Sodium Pertechnetate (99mTc)		Quanty Control	
	injection will yield a preparation of an		Prior to use in the natient	, the radiochemical purity
	appropriate quality.			succimer injection solution
4.	It is recommended that with proper shielding	Insert layout and details specific to	must be tested using the r	
	and equipment, the final formulation be tested			
	for radiochemical purity. If radiochemical	manufacturer	Preparation:	
	purity is not adequate, discard the finished		-	
	drug.		Type of test:	Thin layer
			chromatography Plates used:	Silica gel on a glass fibre
			plate, heated for 10 min.	Silica ger oli a glass fibre
			place, neared for to min.	at 110 °C prior to testing
			Starting point:	1.5 cm from lower end of
			the plate	
			Migration distance:	10 to 15 cm (in approx.
			15 minutes)	
			Execution:	
			TT	
				pette to extract a volume of
			approx. 5 µl and apply it Chromatography begins i	
1			solution of methylethylke	
			migration distance of 10	
			to air-dry, and use a detec	
			distribution of radioactivi	ity.
			Evaluation:	
Rx	ONLY		The [99mTc] technetium succimer complex remains	
1			the starting point while [99	
			migrates near the solvent	nont.
			Target value: \geq 95.0 % [⁹⁹	^{9m} Tc]technetium succimer
			$\leq 2.0\%$ [^{99m}	Tc]pertechnetate
I		1	_ 2.070 [por teennetute

GE REFERENCE PRODUCT INSERT	DIFFERENCES	ROTOP-DMSA INSERT
		CLASSIFICATION FOR SUPPLY
		Pharmacy-only medicine

Attachment 3: Product Labels

Vial



Carton

	ROTOP – DMSA 1.0 mg Kit for radiopharmaceutical preparation Succimer			
For the use in infants, children and adults.				
5 vials Content/vial: 1.74 mg powder for solution for injection active substance: 1.0 mg succimer excipients: stannous chloride dihydrate, ascorbic acid, sodium hydroxide, hydrochloric acid, nitrogen				
For intravenous use after reconstitution and labelling. Store in the original package in or to protect from light. Store in a refrigerator at 2 – 8 °C. Keep out of the sight and reach or children.				
MA Number: 3	003663.00.00 pharmacy only medicine			
ROTOP Pharmaka GmbH, Bautzner Landstraße 400, 01328 Dresden, Germany				