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3 Committee for Medicinal Products for Human Use (CHMP)

4 **Guideline on core SmPC and Package Leaflet for**
5 **(⁹⁹Mo/^{99m}Tc) generator**

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Comments should be provided using this [template](#). The completed comments form should be sent to radiopharmaceuticalsDG@ema.europa.eu

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Keywords	<i>Radiopharmaceuticals, radionuclide, kit for radiopharmaceutical preparation, core SmPC, core Package Leaflet, (⁹⁹Mo/^{99m}Tc) generator</i>
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17 **Executive summary**

18 This guideline describes the information to be included in the Summary of Products Characteristics
19 (SmPC) and Package Leaflet for (⁹⁹Mo/^{99m}Tc) generator.

20 **1. Introduction (background)**

21 The purpose of this core SmPC and Package Leaflet is to provide applicants and regulators with
22 harmonised guidance on the information to be included in the Summary of product characteristics
23 (SmPC) for (⁹⁹Mo/^{99m}Tc) generator ¹. This guideline should be read in conjunction with the core SmPC
24 and Package Leaflet for Radiopharmaceuticals, the QRD product information templates and the
25 guideline on Summary of Product Characteristics.

26 This Core SmPC has been prepared on the basis of national SmPCs taking into account the approved
27 Core Safety Profile (CSP) from the recently finished PSUR worksharing procedure and recently finalised
28 European procedures. However, any new application for a (⁹⁹Mo/^{99m}Tc) generator should be submitted
29 with all the needed and adequate data in order to be valid.

30 The indications concerning labelling of red blood cells are not part of this generator's Core SmPC as
31 they are authorised with the respective kit for pre-treatment with a reducing agent. The indication
32 brain scintigraphy which was included in the previous core SmPC is considered no more clinically
33 relevant and was therefore deleted.

34 The activities to be administered to children and to adolescents may be calculated according to the
35 EANM Dosage Card [Lassmann M et al. Eur J Nucl Med Mol Imaging (2008) 35:1667].

36 Note: Section 4.8 Undesirable effects was completely revised to meet the current medical knowledge.
37 The content of the CSP resulting from the PSUR worksharing was included completely. Section 4.8. of
38 the CSP is probably based on the original SmPC version from 1992 and has not been updated since.
39 Therefore, a comprehensive search of the marketing authorisation holder's (MAH) adverse reactions
40 database was performed covering the complete lifetime of the medicinal product, and all reported side
41 effects were medically assessed for inclusion into the SmPC. The frequency was set to unknown for all
42 symptoms, since all information was solely derived from the spontaneous reporting system. Compared
43 to the CSP a table was introduced and includes a MedDRA tabulation of adverse reactions to adapt the
44 structure to the current requirements for SmPCs. The symptom "coma" as listed in the CSP was
45 included into the SmPC even though no spontaneous reports of coma have been received by the MAH
46 to date.

47 **2. Scope**

48 This core SmPC and Package Leaflet covers (⁹⁹Mo/^{99m}Tc) generator.

49 **3. Legal basis**

50 This guideline has to be read in conjunction with Article 11 of Directive 2001/83 as amended, and the
51 introduction and general principles (4) and part I of the Annex I to Directive 2001/83 as amended.

52 **4. Core SmPC and Package Leaflet for (⁹⁹Mo/^{99m}Tc) 53 generator**

54 ¹ Concept paper on the harmonisation and update of the clinical aspects in the authorised conditions of use for
radiopharmaceuticals and other diagnostic medicinal products (EMA/CHMP/EWP/12052/2008)

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CORE SmPC and Package Leaflet for (⁹⁹Mo/^{99m}Tc) generator

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ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS

161 <▼ This medicinal product is subject to additional monitoring. This will allow quick identification of
 162 new safety information. Healthcare professionals are asked to report any suspected adverse reactions. See
 163 section 4.8 for how to report adverse reactions.>
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165 **1. 1. NAME OF THE MEDICINAL PRODUCT**

166 {(Invented) name strength} GBq radionuclide generator
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 170 **2. QUALITATIVE AND QUANTITATIVE COMPOSITION**

171 Sodium pertechnetate (^{99m}Tc) injection is produced by means of a (⁹⁹Mo/^{99m}Tc) generator. Technetium
 172 (^{99m}Tc) decays with the emission of gamma radiation with a mean energy of 140 keV and a half-life of
 173 6.02 hours to technetium (⁹⁹Tc) which, in view of its long half-life of 2.13 x 10⁵ years, can be regarded as
 174 quasi stable.
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176 The radionuclide generator containing the parent isotope ⁹⁹Mo, adsorbed on a chromatographic column
 177 delivers sodium pertechnetate (^{99m}Tc) injection in sterile solution.
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179 The ⁹⁹Mo on the column is in equilibrium with the formed daughter isotope ^{99m}Tc. The generators are
 180 supplied with the following ⁹⁹Mo activity amounts at activity reference time which deliver the following
 181 technetium (^{99m}Tc) amounts, assuming a 100% theoretical yield and 24 hours time from previous elution
 182 and taking into account that branching ratio of ⁹⁹Mo is about 87%: *[Product specific]*
 183
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^{99m} Tc activity (Maximal theoretical eluable activity at calibration date, {XX} CET)	<i>i.e.</i> 2.0																			GBq	
⁹⁹ Mo activity (at calibration date, {XX} CET)	<i>i.e.</i> 2.5																				GBq

185 The technetium (^{99m}Tc) amounts available by a single elution depend on the real elution yield of generator
 186 itself declared by manufacturer and approved by NCA.
 187

188 Excipient(s) with known effect:

189 Each mL of sodium pertechnetate (^{99m}Tc) solution contains {XX} mg of sodium.

190 For the full list of excipients, see section 6.1.
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 194 **3. PHARMACEUTICAL FORM**

195 Radionuclide generator.
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197 *[Appearance product specific]*
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 201 **4. CLINICAL PARTICULARS**

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 203 **4.1 Therapeutic indications**

204 This medicinal product is for diagnostic use only.
 205

206 The eluate from the radionuclide generator (sodium pertechnetate (^{99m}Tc) injection) is indicated for:
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- labelling of various kits for radiopharmaceutical preparation developed and approved for radiolabelling with such solution
- Thyroid scintigraphy: direct imaging and measurement of thyroid uptake to give information on the size, position, nodularity and function of the gland in case of thyroid disease.
- Salivary gland scintigraphy: diagnosis of chronic sialadenitis (e.g. (Sjögren's Syndrom) as well as assessment of salivary gland function and duct patency in salivary glands disorders and monitoring of the response to therapeutic interventions (in particular radio iodine therapy).
- Location of ectopic gastric mucosa (Meckel's diverticulum).
- Lacrimal duct scintigraphy: to assess functional disorders of lacrimation and monitoring of the response to therapeutic interventions

4.2 Posology and method of administration

Posology

If sodium pertechnetate (^{99m}Tc) is administered intravenously, activities may vary widely according to the clinical information required and the equipment employed. The injection of activities greater than local DRLs (Diagnostic Reference Levels) should be justified. for certain indications. Recommended activities are as follows:

Adults (70kg) and elderly population

- Thyroid scintigraphy: 20-80 MBq
- Salivary gland scintigraphy: 30 to 150 MBq for static images up to 370 MBq for dynamic images
- Meckel's diverticulum scintigraphy: 300-400 MBq
- Lacrimal duct scintigraphy: 2-4 MBq per drop per eye

Renal impairment

Careful consideration of the activity to be administered is required since an increased radiation exposure is possible in these patients.

Paediatric population

The use in children and adolescents has to be considered carefully, based upon clinical needs and assessing the risk/benefit ratio in this patient group.

The activity to be administered to children and adolescents must be adapted <and may be calculated according to the recommendations of the European Association of Nuclear Medicine (EANM) paediatric dosage card>; the activity administered to children and to adolescents may be calculated by multiplying a baseline activity (for calculation purposes) by the weight-dependent correction factor given in the table below (see Table 1).

$$A[\text{MBq}]_{\text{Administered}} = \text{Baseline Activity} \times \text{Multiple}$$

Thyroid scintigraphy: Activity administered [MBq] = 5.6 MBq x correction factor (Table 1). A minimal activity of 10 MBq is necessary for obtaining images of sufficient quality.

Identification/location of ectopic gastric mucosa: Activity administered [MBq] = 10.5 MBq x correction factor (Table 1). A minimal activity of 20 MBq is necessary in order to obtain images of sufficient quality.

Table 1: Weight-dependent correction factors in the paediatric population (for thyroid scintigraphy and identification/location of ectopic gastric mucosa) according to the EANM-May 2008 guidelines

Weight [kg]	Multiple	Weight [kg]	Multiple	Weight [kg]	Multiple
3	1	22	5.29	42	9.14
4	1.14	24	5.71	44	9.57

6	1.71	26	6.14	46	10.00
8	2.14	28	6.43	48	10.29
10	2.71	30	6.86	50	10.71
12	3.14	32	7.29	52-54	11.29
14	3.57	34	7.72	56-58	12.00
16	4.00	36	8.00	60-62	12.71
18	4.43	38	8.43	64-66	13.43
20	4.86	40	8.86	68	14.00

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Salivary gland scintigraphy: The Paediatric Task Group of EANM (1990) recommends that the activity to be administered to a child should be calculated from the body weight according to the table below (see Table 2) with a minimum dose of 10 MBq in order to obtain images of sufficient quality.

Table 2: Weight-dependent correction factor in the paediatric population (for salivary gland scintigraphy) according to EANM 1990 recommendations

Weight [kg]	Factor	Weight [kg]	Factor	Weight [kg]	Factor
3	0.1	22	0.50	42	0.78
4	0.14	24	0.53	44	0.80
6	0.19	26	0.56	46	0.82
8	0.23	28	0.58	48	0.85
10	0.27	30	0.62	50	0.88
12	0.32	32	0.65	52-54	0.90
14	0.36	34	0.68	56-58	0.92
16	0.40	36	0.71	60-62	0.96
18	0.44	38	0.73	64-66	0.98
20	0.46	40	0.76	68	0.99

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Lacrimal duct scintigraphy: Recommended activities apply as well for adults as for children.

Method of administration

For multidose use.

For intravenous or ocular use.

For instructions on extemporaneous preparation of the medicinal product before administration, see section 12.

For patient preparation, see section 4.4.

In thyroid scintigraphy, salivary gland scintigraphy and identification/location of ectopic gastric mucosa, the sodium pertechnetate (^{99m}Tc) solution is administered by intravenous injection.

In lacrimal duct scintigraphy, drops are instilled in each eye (ocular use).

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Image acquisition

Thyroid scintigraphy: 20 minutes after intravenous injection.

Salivary gland scintigraphy: immediately after intravenous injection and at regular intervals for 15 minutes.

Identification/location of ectopic gastric mucosa: immediately after intravenous injection and at regular intervals for 30 minutes.

Lacrimal duct scintigraphy: dynamic acquisition within 2 minutes after instillation, followed by static images acquired at regular intervals within 20 minutes.

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4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

4.4 Special warnings and precautions for use

Potential for hypersensitivity or anaphylactic reactions

If hypersensitivity or anaphylactic reactions occur, the administration of the medicinal product must be discontinued immediately and intravenous treatment initiated, if necessary. To enable immediate action in emergencies, the necessary medicinal products and equipment such as endotracheal tube and ventilator must be immediately available.

Individual benefit/risk justification

For each patient, the radiation exposure must be justifiable by the likely benefit. The activity administered should in every case be as low as reasonably achievable to obtain the required <diagnostic information> <therapeutic effect>.

Renal impairment

Careful consideration of the benefit risk ratio in these patients is required since an increased radiation exposure is possible.

Paediatric population

For information on the use in paediatric population, see section 4.2 <or 5.1>.

Careful consideration of the indication is required since the effective dose per MBq is higher than in adults (see section 11).

Thyroid blocking is of special importance in the paediatric patient population except for thyroid scintigraphy.

Patient preparation

Pre-treatment of patients with thyroid-blocking medicinal products may be necessary for certain indications.

The patient should be well hydrated before the start of the examination and urged to void as often as possible during the first hours after the examination in order to reduce radiation.

To avoid false positives or to minimise irradiation by reduction of pertechnetate accumulation in the thyroid and salivary glands, a thyroid blocking agent should be given prior to lacrimal duct scintigraphy or Meckel's diverticulum scintigraphy. Conversely a thyroid blocking agent must NOT be used before thyroid, parathyroid or salivary glands scintigraphy.

Before the application of sodium (^{99m}Tc)pertechnetate solution for scintigraphy of Meckel's diverticulum the patient should keep an empty stomach for 3 to 4 hours to reduce intestinal peristalsis.

After in vivo labelling of erythrocytes using stannous ions for reduction sodium pertechnetate (^{99m}Tc) is primarily built into erythrocytes, therefore Meckel's scintigraphy should be performed before or some days after in vivo labelling of erythrocytes.

After the procedure

Close contact with infants and pregnant women should be restricted during 12 hours.

Specific warnings

<Sodium pertechnetate (^{99m}Tc) solution for injection contains {XX} mg/mL of sodium.

Depending on the time when the injection is administered, the content of sodium given to the patient may in some cases be greater than 1 mmol (23 mg). This should be taken into account in patient on low sodium diet.>

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348 In salivary gland scintigraphy a lower specificity of the method should be expected compared to MR
349 sialography.

350
351 When sodium pertechnetate (^{99m}Tc) solution is used for labelling of a kit, the determination of the overall
352 sodium content must take into account the sodium derived from the eluate and the kit. Please refer to the
353 package leaflet of the kit.

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355 For precautions with respect to environmental hazard, see section 6.6.

356 357 **4.5 Interaction with other medicinal products and other forms of interaction**

358
359 Atropine, isoprenaline and analgesics may cause a delay of gastric emptying and thereby cause a
360 redistribution of (^{99m}Tc) pertechnetate in abdominal imaging.

361
362 Administration of laxatives should be withheld since they irritate the gastrointestinal tract. Contrast-
363 enhanced studies (e.g. barium) and upper gastro-intestinal examination should be avoided within 48h prior
364 to administration of pertechnetate (^{99m}Tc) for Meckel's diverticulum scintigraphy.

365
366 Many pharmacological medicinal products are known to modify the thyroid uptake.

- 367 • antithyroid medicinal products (e.g. carbimazole or other imidazole derivatives such as
368 propylthiouracil), salicylates, steroids, sodium nitroprusside, sodium sulfobromophthalein, perchlorate
369 should be withheld for 1 week prior thyroid scintigraphy ;
- 370 • phenylbutazone and expectorants should be withheld for 2 weeks ;
- 371 • natural or synthetic thyroid preparations (e.g. sodium thyroxine, sodium liothyronine, thyroid extract)
372 should be withheld for 2-3 weeks
- 373 • amiodarone, benzodiazepines, lithium should be withheld for 4 weeks
- 374 • intravenous contrast agents should not have been administered within 1-2 months.

375 376 **4.6 Fertility, pregnancy and lactation**

377 378 Women of childbearing potential

379 When an administration of radiopharmaceuticals to a woman of childbearing potential is intended, it is
380 important to determine whether or not she is pregnant. Any woman who has missed a period should be
381 assumed to be pregnant until proven otherwise. If in doubt about her potential pregnancy (if the woman
382 has missed a period, if the period is very irregular, etc.), alternative techniques not using ionising radiation
383 (if there are any) should be offered to the patient.

384 385 Pregnancy

386 Administration of pertechnetate (^{99m}Tc) to a woman who is known to be pregnant should be justified by
387 medical need and a positive individual benefit risk assessment for the mother and the foetus. Alternative
388 non-irradiating diagnostic modalities should be taken into account.
389 ^{99m}Tc (as free pertechnetate) has been shown to cross the placental barrier.

390 391 Breastfeeding

392 Before administering radiopharmaceuticals to a mother who is breastfeeding, consideration should be
393 given to the possibility of delaying the administration of radionuclide until the mother has ceased
394 breastfeeding and to what is the most appropriate choice of radiopharmaceuticals, bearing in mind the
395 secretion of activity in breast milk. If the administration is considered necessary, breastfeeding should be
396 interrupted for 12 hours post administration and the expressed feeds discarded.
397 Close contact with infants should be restricted during this period.

398 399 **4.7 Effects on ability to drive and use machines**

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401 No studies on the effects on the ability to drive and use machines have been performed.

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4.8 Undesirable effects

Summary of the safety profile

Information on adverse reactions is available from spontaneous reporting. The reported reaction types are anaphylactoid reactions, vegetative reactions, as well as different kinds of injection site reactions. Sodium pertechnetate (^{99m}Tc) from the {(Invented) name} radionuclide generator is used for radioactive labelling of a variety of compounds. These medicinal products generally have a higher potential for adverse reactions than ^{99m}Tc , and therefore the reported adverse reactions are rather related to the labelled compounds than to ^{99m}Tc . The possible types of adverse reactions following intravenous administration of a ^{99m}Tc -labelled pharmaceutical preparation will be dependent on the specific compound being used. Such information can be found in the SmPC of the kit used for radiopharmaceutical preparation.

Tabulated list of adverse reactions

The frequencies of undesirable effects are defined as follows:

Very common ($\geq 1/10$), common ($\geq 1/100$ to $< 1/10$), uncommon ($\geq 1/1,000$ to $< 1/100$), rare ($\geq 1/10,000$ to $< 1/1,000$), very rare ($< 1/10,000$) and not known (cannot be estimated from the available data).

Immune system disorders

Frequency unknown*: Anaphylactoid reactions (e.g. dyspnoea, coma, urticaria, erythema, rash, pruritus, oedema at various location e.g. face oedema)

Nervous system disorders

Frequency unknown*: Vasovagal reactions (e.g. syncope, tachycardia, bradycardia, dizziness, headache, vision blurred, flushing)

Gastrointestinal disorders

Frequency unknown*: Vomiting, nausea, diarrhoea

General disorders and administration site conditions

Frequency unknown*: Injection site reactions due to extravasation (e.g. cellulitis, pain, erythema, swelling)

* Adverse reactions derived from spontaneous reporting

Exposure to ionising radiation is linked with cancer induction and a potential for development of hereditary defects. As the effective dose is 5.2 mSv when the maximal recommended activity of 400 MBq is administered these adverse reactions are expected to occur with a low probability.

Description of selected adverse reactions

Anaphylactic reactions (e.g. dyspnoea, coma, urticaria, erythema, rash, pruritus, oedema at various locations [e.g. face oedema])

Anaphylactic reactions have been reported following intravenous injection of sodium perchtechnetate (^{99m}Tc) and include various skin or respiratory symptoms like skin irritations, oedema, or dyspnoea.

Vegetative reactions (nervous system and gastrointestinal disorders)

Single cases of severe vegetative reactions have been reported, however, most of the reported vegetative reactions include gastrointestinal reactions like nausea or vomiting. Other reports include vasovagal reactions like headache or dizziness. Vegetative reactions are rather considered to be related to the examination setting than to technetium (^{99m}Tc), especially in anxious patients.

General disorders and administration site conditions

Other reports describe local injection site reactions. Such reactions are related to extravasation of the radioactive material during the injection, and the reported reactions rank from local swelling up to cellulitis. Depending on the administered radioactivity and the labeled compound, extended extravasation may necessitate surgical treatment.

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Reporting of suspected adverse reactions

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in [Appendix V*](#).

*[*For the printed material, please refer to the guidance of the annotated QRD template.]*

4.9 Overdose

In the event of administration of a radiation overdose with sodium pertechnetate (^{99m}Tc), the absorbed dose should be reduced where possible by increasing the elimination of the radionuclide from the body by defaecation, forced diuresis and frequent bladder voiding.

The uptake in the thyroid, salivary glands and the gastric mucosa can be significantly reduced when sodium perchlorate is given immediately after an accidentally high dose of sodium pertechnetate (^{99m}Tc) was administered.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: Diagnostic radiopharmaceuticals, various thyroid diagnostic radiopharmaceuticals, ATC code: V09FX01.

No pharmacological activity has been observed in the range of doses administered for diagnostic purposes.

<Paediatric population>

<The European Medicines Agency has waived the obligation to submit the results of studies with <{(Invented) Name}> [or for generics: <the reference medicinal product containing {name of the active substance(s)}>] in all subsets of the paediatric population in {condition as per Paediatric Investigation Plan (PIP) decision, in the granted indication} (see section 4.2 for information on paediatric use).>

<The European Medicines Agency has deferred the obligation to submit the results of studies with <{(Invented) Name}> [or for generics: <the reference medicinal product containing {name of the active substance(s)}>] in one or more subsets of the paediatric population in {condition, as per Paediatric Investigation Plan (PIP) decision in the granted indication} (see section 4.2 for information on paediatric use).>

<This medicinal product has been authorised under a so-called ‘conditional approval’ scheme. This means that further evidence on this medicinal product is awaited. The European Medicines Agency will review new information on the product every year and this SmPC will be updated as necessary.>

<This medicinal product has been authorised under ‘exceptional circumstances’. This means that due to <the rarity of the disease> <for scientific reasons> <for ethical reasons> it has not been possible to obtain complete information on this medicinal product. The European Medicines Agency will review any new information which may become available every year and this SmPC will be updated as necessary.>

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509 **5.2 Pharmacokinetic properties**

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511 Distribution

512 The pertechnetate ion has similar biological distribution to iodide and perchlorate ions, concentrating
513 temporarily in salivary glands, choroid plexus, stomach (gastric mucosa) and in the thyroid gland, from
514 which it is eliminated, unchanged. The pertechnetate ion also tends to concentrate in areas with increased
515 vascularisation or with abnormal vascular permeability, particularly when pre-treatment with blocking
516 agents inhibits uptake in glandular structures. With intact blood brain barrier, sodium pertechnetate (^{99m}Tc)
517 does not penetrate into the brain tissue.

518

519 Organ uptake

520 In the blood 70-80% of the intravenously injected sodium pertechnetate (^{99m}Tc) is bound to proteins,
521 primarily in an unspecific way to albumin. The unbound fraction (20-30%) accumulates temporarily in
522 thyroid and salivary glands, stomach and nasal mucous membranes as well as in the plexus chorioideus.

523

524 Sodium pertechnetate (^{99m}Tc) in contrast to iodine, nevertheless, is neither used for the thyroid hormone
525 synthesis (organification), nor absorbed in the small intestine. In the thyroid the maximum accumulation,
526 depending on functional status and iodine saturation (in euthyroidism approx. 0.3-3%, in hyperthyroidism
527 and iodine depletion up to 25%) is reached about 20 min after injection and then decreases quickly. This
528 also applies for the stomach mucous membrane parietal cells and the salivary glands acinar cells.

529

530 In contrast to the thyroid which releases sodium pertechnetate (^{99m}Tc) in the bloodstream the salivary
531 glands and the stomach secrete sodium pertechnetate (^{99m}Tc) in the saliva and gastric juice, respectively.
532 The accumulation by the salivary gland lies in the magnitude of 0.5% of the applied activity with the
533 maximum reached after about 20 minutes. One hour after injection, the concentration in the saliva is about
534 10-30 fold higher than in the plasma. The excretion can be accelerated by lemon juice or by stimulation of
535 the parasympathetic nerve system, the absorption is reduced by perchlorate.

536

537 Elimination

538 Half life in plasma is approximately 3 hours. Sodium pertechnetate (^{99m}Tc) is not metabolised in the
539 organism. One fraction is eliminated very quickly renally, the rest more slowly via faeces, salivary and
540 tear liquid. Excretion during the first 24 hours following administration is mainly urinary (approximately
541 25%) with faecal excretion occurring over the next 48 hours. Approximately 50% of the administered
542 activity is excreted within the first 50 hours. When selective uptake of pertechnetate (^{99m}Tc) in glandular
543 structures is inhibited by the preadministration of blocking agents, excretion follows the same pathways
544 but there is a higher renal clearance.

545

546 The above data are not valid when sodium pertechnetate (^{99m}Tc) is used for labelling of another
547 radiopharmaceutical.

548

549 **5.3 Preclinical safety data**

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551 There is no information on acute, subacute and chronic toxicity from single or repeated dose
552 administration. The quantity of sodium pertechnetate (^{99m}Tc) administered during clinical diagnostic
553 procedures is very small and, apart from allergic reactions, no other adverse reactions have been reported.
554 This medicinal product is not intended for regular or continuous administration.

555 Mutagenicity studies and long-term carcinogenicity studies have not been carried out.

556

557 Reproductive toxicity

558 Placental transfer of ^{99m}Tc from intravenously administered sodium pertechnetate (^{99m}Tc) has been studied
559 in mice. The pregnant uterus was found to contain as much as 60% of the injected ^{99m}Tc when
560 administered without perchlorate pre-administration. Studies performed on pregnant mice during
561 gestation, gestation and lactation, and lactation alone showed changes in progeny which included weight
562 reduction, hairlessness and sterility.

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6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

[Product specific, including accessories which belong to the radionuclide generator as the solution for elution]

6.2 Incompatibilities

This medicinal product must not be mixed with other medicinal products except for those mentioned in section 12.

6.3 Shelf life

[Product specific, including shelf – life of the accessories as solution for elution and elution vials]

Generator: {XX} days from manufacturing date.

The calibration date and the expiry date are stated on the label.

Sodium pertechnetate (^{99m}Tc) eluate: After elution, use within {XX} hours. This medicinal product does not require any special storage conditions.

Elution vials: {XX} months.

6.4 Special precautions for storage

[Product specific]

Eluate: For storage conditions after elution of the medicinal product, see section 6.3

Storage of radiopharmaceuticals should be in accordance with national regulation on radioactive materials.

6.5 Nature and contents of container

[Product specific, including containers of the accessories, package sizes]

6.6 Special precautions for disposal and other handling

General warnings

Radiopharmaceuticals should be received, used and administered only by authorised persons in designated clinical settings. Their receipt, storage, use, transfer and disposal are subject to the regulations and/or appropriate licences of the competent official organisation.

Radiopharmaceuticals should be prepared in a manner which satisfies both radiation safety and pharmaceutical quality requirements. Appropriate aseptic precautions should be taken.

If at any time the integrity of the generator or the vial with the eluted solution is compromised, it should not be used.

Administration procedures should be carried out in a way to minimise risk of contamination of the medicinal product and irradiation of the operators. Adequate shielding is mandatory.

The administration of radiopharmaceuticals creates risks for other persons from external radiation or contamination from spills of urine, vomiting, etc. Radiation protection precautions in accordance with national regulations must therefore be taken.

The residual activity of the generator must be estimated before disposal.

617 Any unused medicinal product or waste material should be disposed of in accordance with local
618 requirements.
619

620

621 7. MARKETING AUTHORISATION HOLDER

622

623 {Name and address}

624 <{tel}>

625 <{fax}>

626 <{e-mail}>

627

628

629 8. MARKETING AUTHORISATION NUMBER

630

631

632 9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

633

634 <Date of first authorisation: {DD month YYYY}>

635 <Date of latest renewal: {DD month YYYY}>

636

637

638 10. DATE OF REVISION OF THE TEXT

639

640 <{MM/YYYY}>

641 <{DD/MM/YYYY}>

642 <{DD month YYYY}>

643

644

645 11. DOSIMETRY

646

647 The data listed below are from ICRP 80:

648

Organ	Absorbed dose per unit activity administered (mGy/MBq)				
	Adults	15 years	10 years	5 years	1 year
Adrenals	0.0037	0.0047	0.0072	0.011	0.019
Bladder	0.018	0.023	0.030	0.033	0.060
Bone surfaces	0.0054	0.0066	0.0097	0.014	0.026
Brain	0.0020	0.0025	0.0041	0.0066	0.012
Breast	0.0018	0.0023	0.0034	0.0056	0.011
Gall bladder	0.0074	0.0099	0.016	0.023	0.035
Gastrointestinal tract					
Stomach	0.026	0.034	0.048	0.078	0.16
Small intestine	0.016	0.020	0.031	0.047	0.082
Colon	0.042	0.054	0.088	0.14	0.27
(Upper large intestine	0.057	0.073	0.12	0.20	0.38)
(Lower large intestine	0.021	0.028	0.045	0.072	0.13)
Heart	0.0031	0.0040	0.0061	0.0092	0.017
Kidneys	0.0050	0.0060	0.0087	0.013	0.021
Liver	0.0038	0.0048	0.0081	0.013	0.022
Lungs	0.0026	0.0034	0.0051	0.0079	0.014

Muscles	0.0032	0.0040	0.0060	0.0090	0.016
Oesophagus	0.0024	0.0032	0.0047	0.0075	0.014
Ovaries	0.010	0.013	0.018	0.026	0.045
Pancreas	0.0056	0.0073	0.011	0.016	0.027
Red marrow	0.0036	0.0045	0.0066	0.0090	0.015
Salivary glands	0.0093	0.012	0.017	0.024	0.039
Skin	0.0018	0.0022	0.0035	0.0056	0.010
Spleen	0.0043	0.0054	0.0081	0.012	0.021
Testes	0.0028	0.0037	0.0058	0.0087	0.016
Thymus	0.0024	0.0032	0.0047	0.0075	0.014
Thyroid	0.022	0.036	0.055	0.12	0.22
Uterus	0.0081	0.010	0.015	0.022	0.037
Remaining organs	0.0035	0.0043	0.0064	0.0096	0.017
Effective dose (mSv/MBq)	0.013	0.017	0.026	0.042	0.079

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The effective dose resulting from the intravenous administration of 400 MBq of sodium pertechnetate (^{99m}Tc) to an adult weighing 70 kg is about 5.2 mSv.

The radiation dose absorbed by the lens of the eye following administration of sodium pertechnetate (^{99m}Tc) for lacrimal duct scintigraphy is estimated to be 0.038 mGy/MBq. This results in an effective dose equivalent of less than 0.01 mSv for an administered activity of 4 MBq.

The specified radiation exposure is only applicable if all organs accumulating sodium (^{99m}Tc) pertechnetate will function normally. Hyper/hypofunction (e.g. of the thyroid, gastric mucosa or kidney) and extended processes with impairment to the blood-brain-barrier or renal elimination disorders, may result in changes to the radiation exposure, locally even in strong increases of it.

External radiation exposure

	^{99}Mo - ^{99m}Tc dose rate on the surface of generator (mGy/h/GBq)	^{99}Mo - ^{99m}Tc dose rate at 1 m distance from the generator (mGy/h/GBq)
Shielding with ... mm lead

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Measurements on the location and during work are critical and should be practised for more precise and instructive determination of overall radiation dose to the staff.

12. INSTRUCTIONS FOR PREPARATION OF RADIOPHARMACEUTICALS

[Product specific: e. g. handling of the radionuclide generator, elution frequency, configuration of the radionuclide generator, use and specifics of the accessories, description of the correlation technetium (^{99m}Tc) elution yield, mandatory quality testing by the user, e. g. tests on molybdenum (^{99}Mo) break through.]

Elution of the generator must be performed in premises complying with the national regulations concerning the safety of use of radioactive products.

The solution eluted is a clear and colourless sodium pertechnetate (^{99m}Tc) solution, with a pH between {XXX} and {XXXX} and a radiochemical purity equal to or greater than {XX}.

681 When sodium pertechnetate (^{99m}Tc) solution is used for kit labelling, please refer to the package leaflet of
682 the concerned kit.

683

684 Quality control

685 Clarity of the solution, pH, radioactivity and the molybdenum (^{99}Mo) break-through must be checked
686 before administration.

687

688 The test for molybdenum (^{99}Mo) break-through can be performed either according to Ph. Eur. or to any
689 other validated methods able to determine a molybdenum (^{99}Mo) content below 0.1 per cent of total
690 radioactivity at the date and hour of administration.

691

692 <The first eluate obtained from this generator can be normally used, unless otherwise specified. The eluate
693 can be used for kit labelling even eluted after 24 hours from the last elution, except if the use of fresh
694 eluate is specified in the relevant kit SmPC.>

695

696 Detailed information on this medicinal product is available on the website of the European Medicines
697 Agency <http://www.ema.europa.eu>

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B. PACKAGE LEAFLET

720 **Package leaflet: Information for the patient**

721
722 **{(Invented) name strength} GBq radionuclide generator**
723 Sodium pertechnetate (^{99m}Tc) solution
724

725 < ▼ This medicine is subject to additional monitoring. This will allow quick identification of new safety
726 information. You can help by reporting any side effects you may get. See the end of section 4 for how to
727 report side effects.>

728
729 **Read all of this leaflet carefully before you are given this medicine because it contains important**
730 **information for you.**

- 731
732 - Keep this leaflet. You may need to read it again.
733 - If you have any further questions, ask your nuclear medicine doctor who will supervise the procedure.
734 - If you get any side effects talk to your nuclear medicine doctor. This includes any possible side effects
735 not listed in this leaflet. See Section 4.
736

737 **What is in this leaflet:**

- 738
739 1. What X is and what it is used for
740 2. What you need to know before the sodium pertechnetate (^{99m}Tc) solution obtained with X is used.
741 3. How sodium pertechnetate (^{99m}Tc) solution obtained with X is used
742 4. Possible side effects
743 5. How X is stored
744 6. Contents of the pack and other information
745
746

747 **1. What X is and what it is used for**

748
749 This medicine is a radiopharmaceutical product for diagnostic use only.

750
751 X is a technetium (^{99m}Tc) generator, which means it is a device used to obtain a solution for injection of
752 sodium pertechnetate (^{99m}Tc). When this radioactive solution is injected, it temporarily collects in certain
753 areas of the body. The low quantity of radioactivity injected can be detected outside of the body by special
754 cameras. The nuclear medicine doctor will then take an image (scan) of the concerned organ which can
755 give him valuable information about the structure and the function of this organ.
756

757 After injection the sodium pertechnetate (^{99m}Tc) solution is used to obtain images of various body parts
758 such as the:

- 759 - thyroid gland
760 - salivary glands
761 - appearance of stomach tissue in an abnormal location (Meckel diverticulum)
762 - tear ducts of the eyes
763

764 The sodium pertechnetate (^{99m}Tc) solution can also be used in combination with another product to
765 prepare another radiopharmaceutical medicine. In this case, please refer to the corresponding package
766 leaflet.
767

768 The nuclear medicine doctor will explain to you what type of examination will be performed with this
769 product.
770

771 The use of sodium pertechnetate (^{99m}Tc) solution does involve exposure to small amounts of radioactivity.
772 Your doctor and the nuclear medicine doctor have considered that the clinical benefit that you will obtain
773 from the procedure with the radiopharmaceutical outweighs the risk due to radiation.
774

775 **2. What you need to know before the sodium pertechnetate (^{99m}Tc) solution obtained with X is used**

776

777 **The sodium pertechnetate (^{99m}Tc) solution obtained with X must not be used**

778 - if you are allergic to sodium pertechnetate (^{99m}Tc) or any of the other ingredients of this
779 medicine (listed in section 6).

780

781 **Warning and precautions**

782 Inform your nuclear medicine doctor in the following cases:

- 783 - if you suffer from allergies, as a few cases of allergic reactions have been observed after
- 784 administration of sodium pertechnetate (^{99m}Tc) solution
- 785 - if you suffer from kidney disease
- 786 - if you are pregnant or believe you may be pregnant
- 787 - if you are breast-feeding

788

789 You nuclear medicine doctor will inform you if you need to take any special precautions after using this
790 medicine. Talk to your nuclear medicine doctor if you have any questions.

791

792 Before administration of sodium pertechnetate (^{99m}Tc) solution you should:

- 793 - drink plenty of water before the start of the examination in order to urinate as often as possible during
794 the first hours after the study.
- 795 - You should be fasting for 3 to 4 hours before Meckel's diverticulum scintigraphy to keep the small
796 bowel peristalsis low.

797

798 **Children and adolescents**

799 Please talk to your nuclear medicine doctor if you are under 18 year old.

800

801 **Other medicines and sodium pertechnetate (^{99m}Tc) solution**

802 Tell your nuclear medicine doctor if you are taking, have recently taken or might take any other
803 medicines since they may interfere with the interpretation of the images; and especially the following
804 medicines:

- 805 - **atropine**, used for example:
 - 806 - to reduce stomach, bowel or gall bladder spasms
 - 807 - to reduce pancreas secretions
 - 808 - in ophthalmology
 - 809 - before administering an anaesthesia
 - 810 - to treat reduced heart beat or
 - 811 - as an antidote
- 812 - **isoprenaline**, a medicine to treat reduced heart beat
- 813 - **pain killers**
- 814 - **laxatives** (they should not be taken during this procedure since they irritate the gastrointestinal tract)
- 815 - if you had **contrast-enhanced studies** (e.g. with the contrast agent barium) or **upper gastro-**
816 **intestinal examination** (as these should be avoided within 48h prior to Meckel's diverticulum
817 scintigraphy)
- 818 - **antithyroid medicines** (e.g. carbimazole or other imidazole derivatives such as propylthiouracil),
819 **salicylates, steroids, sodium nitroprusside, sodium sulfobromophthalein, perchlorate** (as they
820 should not be taken for 1 week prior to scintigraphy)
- 821 - **phenylbutazone** to treat fever, pain and inflammation in the body (as it should not be taken for 2
822 weeks prior to scintigraphy)
- 823 - **expectorants** (as they should not be taken for 2 weeks prior to scintigraphy)
- 824 - **natural or synthetic thyroid preparations** (e.g. sodium thyroxine, sodium liothyronine, thyroid
825 extract) (as they should not be taken for 2-3 weeks prior to scintigraphy)
- 826 - **amiodarone** an antiarrhythmic agent (as it should not be taken for 4 weeks prior to scintigraphy)

- 827 - **benzodiazepines** used for example for sedation, or as anti-anxiety or anti-convulsion or muscle
828 relaxant medication or **lithium** used as a mood stabiliser in manic-depressive illness (as both should
829 not be taken for 4 weeks prior to scintigraphy)
830 - **intravenous contrast agents** for radiologic examinations of the body (as they should not have been
831 administered for 1-2 months prior to scintigraphy)
832

833 Please ask your nuclear medicine doctor before taking any medicines.
834

835 **Pregnancy and breast-feeding**

836 If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your
837 nuclear medicine doctor for advice before you are given this medicine.
838

839 You must inform the nuclear medicine doctor before the administration of sodium pertechnetate (^{99m}Tc)
840 solution if there is a possibility you might be pregnant, if you have missed your period or if you are breast-
841 feeding. When in doubt, it is important to consult your nuclear medicine doctor who will supervise the
842 procedure.
843

844 If you are pregnant, your nuclear medicine doctor will only administer this medicine during pregnancy if a
845 benefit is expected which would far outweigh the risks.

846 If you are breast-feeding, please tell your nuclear medicine doctor, as he/she will advise you to stop doing
847 so until the radioactivity has left your body. This takes about 12 hours. The expressed milk should be
848 discarded. Resuming breast-feeding should be in agreement with the specialist in Nuclear Medicine who
849 will supervise the procedure.
850

851 **Driving and using machines**

852 Sodium pertechnetate (^{99m}Tc) solution has no influence on the ability to drive and use machines.
853

854 **Sodium pertechnetate solution contains sodium**

855 Sodium pertechnetate solution contains {XX} mg/mL of sodium. Depending on the volume injected, the
856 limit of 1 mmol (23 mg) of sodium per dose administered may be exceeded. This must be taken into
857 account if you are on a low-salt diet.
858

860 **3. How the sodium pertechnetate (^{99m}Tc) solution obtained with X is used**

861 There are strict laws on the use, handling and disposal of radiopharmaceutical products. X will only be
862 used in special controlled areas. This product will only be handled and given to you by people who are
863 trained and qualified to use it safely. These persons will take special care for the safe use of this product
864 and will keep you informed of their actions.
865
866

867 The nuclear medicine doctor supervising the procedure will decide on the quantity of sodium
868 pertechnetate (^{99m}Tc) solution to be used in your case. It will be the smallest quantity necessary to get the
869 desired information.
870

871 The quantity usually recommended to be administered for an adult ranges depending on the test to be
872 performed, and ranges between 2 and 400 MBq (megabecquerel, the unit used to express radioactivity).
873

874 **Use in children and adolescents**

875 In children and adolescents, the quantity to be administered will be adapted to the child's weight.
876

877 **Administration of sodium pertechnetate (^{99m}Tc) solution and conduct of the procedure**

878 Depending on the purpose of the examination, the product will be administered by injection into an arm
879 vein or may be instilled into the eyes in the form of drops.
880

881 One administration is sufficient to conduct the test that your doctor needs.

882 **Duration of the procedure**

883 Your nuclear medicine doctor will inform you about the usual duration of the procedure.
884 Scans can be performed at any time, between the time of injection and for up to 24 hours after the
885 administration, depending on the type of examination.

886
887 **After administration of sodium pertechnetate (^{99m}Tc) solution has been performed, you should:**

- 888 – avoid any close contact with young children and pregnant women for the 12 hours following the
- 889 injection
- 890 – urinate frequently in order to eliminate the product from your body
- 891 – After administration, you will be offered a drink and asked to urinate immediately preceding the test.

892
893 The nuclear medicine doctor will inform you if you need to take any special precautions after receiving
894 this medicine. Contact your nuclear medicine doctor if you have any questions.

895
896 **If you have been given more sodium pertechnetate (^{99m}Tc) solution obtained with X than you should**

897 An overdose is almost impossible because you will only receive a single dose of sodium pertechnetate
898 (^{99m}Tc) solution precisely controlled by the nuclear medicine doctor supervising the procedure. However,
899 in the case of an overdose, you will receive the appropriate treatment. In particular, the nuclear medicine
900 doctor in charge of the procedure may recommend that you drink plenty of fluids to remove the traces of
901 radioactivity from your body.

902
903 Should you have any further questions on the use of this product, please ask nuclear medicine doctor who
904 supervises the procedure.

905
906
907 **4. Possible side effects**

908
909 Like all medicines, this medicine can cause side effects, although not everybody gets them.

910 Frequency not known, cannot be estimated from the available data:

- 911 - allergic reactions, with symptoms such as
 - 912 - skin rash, itching
 - 913 - hives
 - 914 - swelling at various locations, e.g. of the face
 - 915 - shortage of breath
 - 916 - redness of the skin
 - 917 - coma
- 918 - circulatory reactions, with symptoms such as
 - 919 - rapid heart beat, slow heart beat
 - 920 - fainting
 - 921 - blurred vision
 - 922 - dizziness
 - 923 - headache
 - 924 - flushing
- 925 - gastrointestinal disorders, with symptoms such as
 - 926 - being sick (vomiting)
 - 927 - feeling sick (nausea)
 - 928 - diarrhoea
- 929 - injection site reactions, with symptoms such as
 - 930 - skin inflammation
 - 931 - pain
 - 932 - swelling
 - 933 - redness

936 This radiopharmaceutical will deliver low amounts of ionising radiation associated with the least risk of
937 cancer and hereditary abnormalities.

938

939 **Reporting of side effects**

940 If you get any side effects, talk to your <doctor> <or> <,> <pharmacist> <or nurse>. This includes any
941 possible side effects not listed in this leaflet. You can also report side effects directly via the national
942 reporting system listed in [Appendix V](#)*. By reporting side effects you can help provide more information
943 on the safety of this medicine.

944

945 [*For the printed material, please refer to the guidance of the annotated QRD template.*]

946

947

948 **5. How X is stored**

949

950 You will not have to store this medicine. This medicine is stored under the responsibility of the specialist
951 in appropriate premises. Storage of radiopharmaceuticals will be in accordance with national regulation on
952 radioactive materials.

953

954 The information is intended for the specialist only.

955

956 This medicine must not be used after the expiry date which is stated on the <label> <carton> <bottle>
957 <...> <after {abbreviation used for expiry date}>. <The expiry date refers to the last day of that month.>

958

959 <This medicine will not be used if it is noticed {description of the visible signs of deterioration}>.

960

961

962 **6. Contents of the pack and other information**

963

964 **What X contains**

965 The active substance is sodium pertechnetate (^{99m}Tc) solution.

966

967 The other ingredients are: *[Product specific]*.

968

969 **What X looks like and contents of the pack**

970

971 The product is sodium pertechnetate (^{99m}Tc) solution provided by a radionuclide generator.

972

973 X has to be eluted and the obtained solution may be used itself or to radiolabelled some particular kits for
974 radiopharmaceutical preparation.

975

976 Pack size

977 *[Product specific]*

978

979 **Marketing Authorisation Holder and Manufacturer**

980

981 {Name and address}

982 <{tel}>

983 <{fax}>

984 <{e-mail}>

985

986 **This leaflet was last revised in {MM/YYYY} {month YYYY}**

987

988 <This medicine has been given ‘conditional approval’. This means that there is more evidence to come
989 about this medicine.

990

991 The European Medicines Agency will review new information on the medicine every year and this leaflet
992 will be updated as necessary.>

993
994 <This medicine has been authorised under ‘exceptional circumstances’. This means that <because of the
995 rarity of this disease> <for scientific reasons> <for ethical reasons> it has been impossible to get complete
996 information on this medicine.

997 The European Medicines Agency will review any new information on the medicine every year and this
998 leaflet will be updated as necessary.>

999
1000 **<Other sources of information>**

1001
1002 Detailed information on this medicine is available on the European Medicines Agency web
1003 site: <http://www.ema.europa.eu>

1004
1005 <This leaflet is available in all EU/EEA languages on the European Medicines Agency website.>

1006
1007 -----

1008 The following information is intended for healthcare professionals only:

1009 The complete SmPC of {(Invented) name} is provided <as a separate document> <as a tear-off section at
1010 the end of the printed leaflet> in the product package, with the objective to provide healthcare
1011 professionals with other additional scientific and practical information about the administration and use of
1012 this radiopharmaceutical.

1013
1014 Please refer to the SmPC [SmPC should be included in the box].