The use of ^{99m}Tc as a tracer in the determination of ⁹⁹Tc by ICPmass spectrometry

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The aim of this paper is to carry out a deep study on the possibilities of using ⁹⁹mTc from ⁹⁹Mo–⁹⁹mTc clinical isotope generators to trace the radiochemical separations for ⁹⁹Tc measurements by ICP-MS. By appropriately defining a minimum detectable mass concentration (MDMC) for the mass spectrometer it is possible to conclude that such use is acceptable for a wide range of activities of the tracer. The MDMC increases from 0.41 to 0.51 ppt when the amount of ⁹⁹mTc added to the samples ranges from 0 to 2 MBq. The purification of the final sample from Ru must be the main objective of the ⁹⁹Tc measurement method, since its presence noticeably enhances the ⁹⁹Tc MDMC.

1 Introduction

⁹⁹Tc is a pure beta-emitter, $E_{\text{max}} = 294$ keV, with a long half-life of 2.11×10^5 y. It can be considered as an extinct radionuclide although there is some experimental documentation on its natural production during the spontaneous fission of ²³⁸U in uranium ores.¹

Nevertheless, ⁹⁹Tc has been reintroduced into nature during the nuclear era, since it is also produced during the fission of ²³⁵U and ²³⁹Pu at a rate of an average of 6%, very similar to that of ¹³⁷Cs. It is estimated that some 15 PBq have been produced throughout the nuclear fuel cycle.² About 10% has been released into the environment which masks any ⁹⁹Tc trace of natural origin. Nuclear military tests have released an additional quantity of 140 TBq mainly into the atmosphere. The environmental relevance of ⁹⁹Tc is well recognised, as its

The environmental relevance of ⁵⁵Tc is well recognised, as its long half-life assures its presence in nature for a long time. This fact makes ⁹⁹Tc a significant radionuclide for future nuclear waste management, as well as an important contributor to the committed dose of radiation to the population. In addition, ⁹⁹Tc is considered a good potential tracer of some environmental processes such as marine water movements.

Despite this fact, there is no ample Tc environmental data archive. This lack of documentation is not surprising considering the difficulty in determining its environmental concentrations. Sub-ppt ⁹⁹Tc levels in non-perturbed sites pose serious limitations on the use of traditional radiometric methods for its determination $(1 \text{ mBq g}^{-1} \text{ }^{99}\text{Tc} = 1.6 \text{ pg g}^{-1})$. It is true, however, that some of these methods, developed during the early eighties, were able to produce a relevant ⁹⁹Tc data file.^{2–4} Nevertheless, some environmental issues still remain unexplored for ⁹⁹Tc and only the recent use of mass spectrometry has enabled the scientific community to meet the experimental challenge of ⁹⁹Tc determination at very low levels.

ICP-MS,⁵ RIMS⁶ and even accelerator mass spectrometry (AMS)⁷ have been used to determine ⁹⁹Tc in a variety of environmental samples. Comparisons between radiometric and non-radiometric methods have also been carried out.⁸ In all cases, source preparation requires more or less intensive radiochemistry to avoid isobaric interferences, or to condition the sample in the ion source. Consequently, it is necessary to

use a tracer to calculate the yield obtained after sample preparation. 95m Tc, 97m Tc or even 99m Tc are good candidates as tracers in

^{95m}Tc, ^{97m}Tc or even ^{99m}Tc are good candidates as tracers in Tc chemical extractions and, in fact, all of them have already been used in connection with radiometric methods. ^{99m}Tc could be a good option as it is very easy to obtain. However, it does not seem the best choice when using ICP-MS, at least from a formal point of view, because ⁹⁹Tc is its daughter product. In principle, this could lead to interferences in the sample measurements. ^{95m}Tc ($T_{1/2}=61$ days) is the most suitable radiochemical tracer for mass spectrometric measurements, due to its convenient long half-life and the absence of possible mass interferences. Nevertheless, some authors claim that commercially available solutions of ^{95m}Tc contain considerable amounts of ⁹⁹Tc.⁹ The same scientists propose the production of laboratory-made solutions of this radionuclide using the reaction ⁹⁵Nb(α ,2n)^{95m}Tc. However not all the laboratories have access to such facilities and easier alternatives such as that of ^{99m}Tc should be explored.

In this paper, we will study in depth the advantages and disadvantages of using ^{99m}Tc as a tracer for sample preparation in ICP-MS. Several ^{99m}Tc solutions obtained from isotope generators in general use in nuclear medicine are studied by using ICP-MS. The objective is to determine to what extent it is possible to use ^{99m}Tc as a tracer for sample preparation, and what kind of limitations are introduced on the sensitivity of the method.

2 The spectrometer

The ICP-MS system used is a Hewlett-Packard 4500 (Yokogawa Analytical Systems, Tokyo, Japan). Tuning of the system is accomplished by minimising the so called instrumental limit of detection (LOD) defined as:

$$\text{LOD} = \frac{3\sigma_{\text{B}}}{S - B}c\tag{1}$$

where $\sigma_{\rm B}$ is the standard deviation of the counts obtained in the blank during a given integration time, 10 s in our case, *c* is the mass concentration of the standard used for tuning, *S* is the

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number of counts under the mass peak of interest and B the background counts of the process obtained during blank measurement runs.

Our system can perform an automatic tuning for the complete mass range (7-254 u). Using the optimum set of operational parameters, the LOD for ⁹⁹Tc reaches a value of 46 ppt. A complete account of the tuning process for ⁹⁹Tc can be found in the thesis of J. L. Mas.¹⁰ The following is a short description.

A complete series of optimisation experiments with a solution of ⁹⁹Tc containing 65.4 ± 1.1 ppm diluted in deionized water (DAMRI, Gif-sur-Yvette Cedex, France) to obtain 13.1 ± 0.3 ppb were carried out. The first stage of the experiments consisted of the application of the variable control method to check the dependence of the S/B on several operational parameters: rf power, carrier gas flow, sampling depth and ion lens voltages. By maximising the S/B a set of operational parameters was found which gave an instrumental LOD of 0.2 ppt for ⁹⁹Tc.

To check the suitability of the achieved working conditions, a direct search method was used to optimise the S/B by applying Spendley's Simplex¹¹ to five lenses of the ion optics system: the extraction lens 1, the Einzel plate 2, the bias omega lens, the omega minus lens and the quadrupole focus lens. During that optimisation run, the set of operational parameters and mechanical variables previously found, were kept constant.

The Simplex was carried out by adapting the algorithm from the Numerical Recipes in Pascal to our working conditions. As can be seen in Fig. 1, there is a fast convergence towards a S/B peak. With the obtained conditions an operational LOD of 0.8 ppt was found. It is clearly lower than that found by automatic tuning and is equivalent to the value obtained during the tuning of the mechanical variables. The final set of parameters is listed in Table 1. The LOD is an operational limit of detection which serves to optimise the system for ⁹⁹Tc counting, in our case. A more realistic and fundamental approach to the problem of detection limits and minimum detectable concentrations is given in Section 5.

3 Samples: preparation and measurement

^{99m}Tc samples were obtained from a ⁹⁹Mo–^{99m}Tc clinical isotope generator, Amertec (Amersham, Buckinghamshire, UK). As usual ^{99m}Tc was eluted from an alumina chromatographic column into a physiological serum solution in 0.9% NaCl.

Consecutive elutions were carried out after each 20-24 h from the same isotope generator. Typically, the eluate volume was 5 ml, with the exception of one sample in which the volume was 12 ml (see next Section).

In Fig. 2 the theoretically expected time evolution of the



Fig. 1 S/B evolution with the 5-D Spenley's Simplex to tune the ion optics.

1370 J. Anal. At. Spectrom., 2000, 15, 1369–1373

Table 1 Operating conditions for ICP-MS

Rf power	1240 W
Coolant gas flow	$151 \mathrm{min}^{-1}$
Carrier	$1.17 \mathrm{l}\mathrm{min}^{-1}$
Nebulizer	Babington type
Sampling depth	6 mm
Extraction lens 1	-150 V
Einzel 2 lens	5.5 V
Bias omega lens	-51 V
Omega minus lens	-3 V
Quadrupole focus	6 V



Fig. 2 Theoretical predictions of normalized (to 99 Mo) activities on the column. The continuous line represents the 99 Mo activity time evolution. Dotted lines give the 99m Tc and 99 Tc activities time evolution.

⁹⁹Mo activity contained in the isotope generator is plotted. The predicted behaviour of the ^{99m}Tc activity present in the generator is given, which includes the periodic cut-offs produced in the growth of the activity due to the elutions. All the results were obtained by applying the well known Bateman equations:

$$a_1 = a_{1,0} e^{-\lambda_1 t} \tag{2}$$

$$a_2 = \frac{\lambda_2 a_{1,0}}{\lambda_2 - \lambda_1} (e^{-\lambda_1 t} - e^{-\lambda_{12} t})$$
(3)

 $a_3 = \lambda_2 \lambda_3 a_{1,0}$

$$\times \left[\frac{e^{-\lambda_1 t}}{(\lambda_2 - \lambda_1)(\lambda_3 - \lambda_1)} + \frac{e^{-\lambda_2 t}}{(\lambda_1 - \lambda_2)(\lambda_3 - \lambda_2)} + \frac{e^{-\lambda_3 t}}{(\lambda_1 - \lambda_3)(\lambda_2 - \lambda_3)}\right] (4)$$

 a_1 and $a_{1,0}$ being the specific activities at time t and t=0 for the i^{th} radionuclide, λi (h⁻¹) its decay constant and t the time. The subindices 1, 2, 3 correspond to ⁹⁹Mo, ^{99m}Tc and ⁹⁹Tc, respectively. ^{99m}Tc in the eluate was left to decay for at least two weeks. In this way the health risks associated with the handling of such high activity samples were avoided. The ^{99m}Tc activity was previously measured by NaI(Tl) γ -spectrometry (EG&G ORTEC, Nuclear Ibérica S.A., Madrid, Spain) and the eluate activity concentration at the time of elution was determined.

Once the ^{99m}Tc had totally decayed, the ⁹⁹Tc mass and activity concentrations were determined using our ICP-MS system with optimum counting conditions. The solutions were diluted nine-fold in 5% HNO₃ before measurement by ICP-MS in order to avoid matrix effects from the massive presence of NaCl in the eluate, which could block the extraction cones. Deionized water (>18 M Ω cm) and suprapure grade HNO₃ (Merck, Darmstadt, Germany) were used for the dilution.

Table 2 ⁹⁹Tc and ^{99m}Tc activity concentrations in the eluates obtained at different elution times from the first elution. Relative uncertainties of ^{99m}Tc activity concentrations range from 1-3%

Time/hours	⁹⁹ Tc activity concentration/Bq g ⁻¹	^{99m} Tc activity concentration/MBq g ⁻¹
0	9.2 ± 0.3	1422.6
21.0	7.3 ± 0.3	596.4
46.5	14.4 ± 0.2	977.5
69.0	9.7 ± 0.2	781.2
141.0	19.5 ± 0.6	394.9
167.2	4.87 ± 0.11	285.6
190.2	3.1 ± 0.1	214.6
213.9	2.4 ± 0.1	168.7
237.7	2.0 ± 0.1	128.1
309.9	3.96 ± 0.14	65.8
331.4	0.75 ± 0.02	45.8
357.7	0.68 ± 0.04	35.4
380.9	0.49 ± 0.02	27.5

4 Results

In Table 2 the ^{99m}Tc and ⁹⁹Tc activity concentrations in the different eluates are presented. Time origin was taken at the first elution instant. Thus, the meaning of time in Table 2 is "time elapsed from the first elution". The results are plotted in Fig. 3. ^{99m}Tc activity follows an exponential decay law with the half-life of ⁹⁹Mo, as is expected. With the exception of the two first points, the rest perfectly fit to an exponential decay law $(r^2 = 0.9996)$, with $a_0 = (1.58 \pm 0.02) \times 10^9 \text{ Bq g}^{-1}$ and $T_{1/2} = 67.4 \pm 1.4$ h. Sample 2 does not follow general behaviour, because the elution volume was 12 ml (see previous Section) instead of 5 ml, the physiological serum volume used for the rest of samples. The consequence was a higher dilution factor for the obtained activity and, therefore, a smaller activity concentration than expected. It is evident that a simple volume normalisation would permit the inclusion of such data into the general exponential curve.

After the first elution, the correlation among the ^{99m}Tc and ⁹⁹Tc activity concentrations in the eluate is very clear, with the exceptions of samples 5 and 10. It can be seen in Table 2 that, in both cases, the time elapsed from the previous elution was larger than normal. In fact they were collected after a week-end waiting time period. The consequence is an increase of the ⁹⁹Tc activity in the generator which is recovered in the following elution. This can be predicted by the Bateman equations, as seen in Fig. 4, where we depict the experimental time evolution of the ⁹⁹Tc activity concentration together with the predictions of eqn. 3. To carry out the calculations, the ⁹⁹Mo initial activity was assumed to be that of ^{99m}Tc in the first eluate, instead of the ⁹⁹Mo activity which appears in the isotope generator description. This last activity is normally given with a very high



Fig. 3 Time evolution of activities on the chromatographic column. Lines are depicted to guide the eye.



Fig. 4 Theoretical and experimental values of ⁹⁹Tc specific activity.

uncertainty. The assumption is perfectly reasonable since both ⁹⁹Mo and ^{99m}Tc are in secular equilibrium before the first elution.

The time used in eqns. 2–4 is the time elapsed between elutions. For every calculation, $a_{1,0}$ is supposed to be that of ⁹⁹Mo activity in the previous elution. In addition $a_{2,0}$ and $a_{3,0}$ are 0. The agreement between the experimental values and the theoretical predictions is clear although the experimental values systematically exceed the theoretical ones. That excess can be explained on the basis of the elution recovery yield for ⁹⁹Tc. Theoretically it is supposed that 100% of the ⁹⁹Tc present in the column is recovered, whereas it is more realistic to suppose that some ⁹⁹Tc remains in the column after each elution. The amount recovered in the following elution is contributing to an increase in the ⁹⁹Tc activity concentration.

The ⁹⁹Tc accumulation effect is perfectly coherent with the predictions of eqn. 4. It is observed that the maximum ^{99m}Tc activity in the column is obtained at t=23.5 h. After such a time the negative term in eqn. 4 dominates and the activity of the daughter product, ⁹⁹Tc, increases. This can also be seen when studying the time evolution of the ⁹⁹Tc/^{99m}Tc activity ratio in the eluates (plotted in Fig 5, together with the experimental data).

All these results are very relevant if ^{99m}Tc from clinical isotope generators is going to be used for tracing radiochemical extractions for ICP-MS measurements. It is clear that previous to the elution of ^{99m}Tc which will be taken for tracing, the accumulated ⁹⁹Tc has to be removed from the generator. Therefore, a prior elution step is compulsory.

Therefore, a prior elution step is compulsory. A given amount of 99 Tc is added to the sample during the spiking process; theoretically some 5 pg of 99 Tc is added to the sample per MBq of 99m Tc. Or in real terms, according to our experimental results, some 10 pg of 99 Tc per MBq of 99m Tc are added to trace the separation, *i.e.*, twice the theoretical expected result.

This has to be taken into account in the calculation of the ⁹⁹Tc concentration in the real sample, and, of course, it affects the LOD of the procedure as can be seen in the following Section.

5 Limits of detection

It is apparent that the LOD is sample-dependent and has to be reformulated. According to Currie¹² the detection limit, LOD, can be written as follows:

$$LOD = 2.54(1 + \sqrt{1 + 2.48\sigma_0^2})$$
 (5)

at a confidence level of 0.05 for the I- and II-species errors with σ_0 being the standard deviation of a null response.



Fig. 5 Time evolution of the activity ratios.

Several contributions have to be taken into account in order to evaluate σ_0 ; firstly, the background *B* due to aleatory counts, photons, overlaps from ¹⁰⁰Mo on mass 99 (due to abundance sensitivity), ⁹⁸MoH generation, electric noise, *etc.*, obtained in a blank procedure sample in the absence, of course, of the tracer. In our working conditions the net counting due to abundance sensitivity could be 1–2 orders of magnitude higher than the contribution from hydride generation.¹⁰ On the other hand, the isotopic abundance of ⁹⁸Mo is more than twice that of ¹⁰⁰Mo natural abundance. Thus, special care must be taken to purify the sample from Mo.⁸

A second contribution comes from the ⁹⁹Ru present in the sample, the reagents or internal standard, which cannot be totally removed with the chemical procedure. This contribution is sample-dependent and can be estimated by checking the presence of other Ru isotopes, *i.e.*, ¹⁰¹Ru or ¹⁰²Ru, in the mass spectrum. The most traditional choice is ¹⁰¹Ru, as we used in this work, although ¹⁰²Ru could be alternatively used provided there is no Pd in the final sample. The well known atomic abundance ratio between ⁹⁹Ru and ¹⁰¹Ru mass region, *S*_{Ru-101}. Finally, the ⁹⁹Tc added to the sample as a consequence of the use of ^{99m}Tc as tracer, *S*_{Tc-99m}, is sample-dependent and will also vary according to the chemical recovery of the separation procedure, *R*_Q.

Therefore, the gross background response B_t at a given integration time becomes:

$$B_{t} = B + \alpha S_{\text{Ru}-101} + S_{\text{Tc}-99\text{m}} = B + \frac{12.6}{17.8} S_{\text{Ru}-101} + \frac{R_{\text{Q}} T_{1/2}^{99\text{m}} a_{\text{t}} m_{\text{t}} M}{N_{\text{A}} m_{\text{s}} \ln 2} b$$
(6)

where $T_{1/2}^{99m}$ is the half-life of 99m Tc, a_t its specific activity added to the sample to measure R_Q , M its molar weight (99 grams), N_A the Avogadro's number, m_t the mass of the tracer added to the sample, m_s the sample mass, and b the slope of the calibration curve of the ICP-MS system.

Consequently, the net response in an experimental measurement would be:

$$S = S_{\rm g} - B - 0.708 \ S_{\rm Ru-101} - \frac{R_{\rm Q} T_{1/2}^{99m} a_{\rm t} m_{\rm t} M}{N_{\rm A} m_{\rm s} \ln 2} b \tag{7}$$

where S_g is the total number of counts under the ⁹⁹Tc mass peak. For a null response the value of σ_0 can be calculated by applying the error propagation formula:

$$\sigma_{0} = \sqrt{\begin{bmatrix} \sigma_{\rm B}^{2} + (0.708 \ \sigma_{\rm Ru-101})^{2} + \left(\frac{T_{1/2}^{99m} Ma_{t}m_{t}bR_{\rm Q}}{N_{\rm A}m_{\rm s}\ln 2}\right)^{2} \times \\ \left[\left(\frac{\sigma_{\rm RQ}}{R_{\rm Q}}\right)^{2} + \left(\frac{\sigma_{\rm at}}{a_{\rm t}}\right)^{2} + \left(\frac{\sigma_{\rm mt}}{m_{\rm t}}\right)^{2} + \left(\frac{\sigma_{\rm b}}{b}\right)^{2} + \left(\frac{\sigma_{\rm ms}}{m_{\rm s}}\right)^{2} \right]$$
(8)

1372 J. Anal. At. Spectrom., 2000, 15, 1369–1373



Fig. 6 MDMC and backgound increment dependence on tracer activity added.



Fig. 7 ⁹⁹Tc MDMC and equivalent Ru concentration dependence on the standard deviation at m/z = 102.

The full mathematical expression for the LOD can be easily written from the above equation. To quantify the capabilities of the method we define a minimum detectable mass concentration, MDMC, as

$$MDMC = \frac{LOD}{b}$$
(9)

where b = 1600 counts ppt⁻¹ and is the slope of the calibration curve of our system for the typical working conditions. The influence of the MDMC of the ^{99m}Tc activity, in Bq

The influence of the MDMC of the ^{99m}Tc activity, in Bq (added to trace the sample) is depicted in Fig. 6. An integration time of 10 s was taken, which supposes that $\sigma_{\rm B}$ =160 counts. On the other hand, a value of $\sigma_{\rm Ru-101}$ =45 counts, a normal value, was fixed for the calculations. Finally $R_{\rm Q}$ was assumed to be 100%, the worst case, although, unfortunately, from another point of view, the less frequent case. With these values a MDMC of 0.41 ppt in the absence of tracer can be calculated. Within a range of 0 to 2 MBq, the MDMC changes from 0.41 to 0.55 ppt, *i.e.*, a 34% relative variation. It seems clear that the modification of the MDMC is not very important and, within a reasonable range of tracer activity from 0 to 100 kBq, the variation is irrelevant. This means that, under our experimental conditions, the influence of the assumed Ru contamination.

The influence of σ_{Ru-101} , as seen in Fig. 7, is more important. The calculations have been performed for a constant tracer activity of 50 kBq, and the above mentioned values for the rest of variables. The plot shows that the need for refining the radiochemical methodology to purify the sample from Ru is a key point with regard to the sensitivity of the method. It is even more important than the influence of the amount of ^{99m}Tc added to the sample, provided the guidelines given in the

6 Conclusions

Measurements of the concentration of 99 Tc in eluates from 99 Mo $^{-99m}$ Tc clinical isotope generators have been carried out. Results have been compared to the predictions from Batemann equations. An algorithm has been developed for the calculation of detection limits in the determination of 99 Tc by ICP-MS. The algorithm can be easily extended to other problems in which the presence of interferences could affect the determination of the nuclide of interest.

It can be concluded that, under well defined working conditions, the use of ^{99m}Tc from clinical isotope generators to trace the radiochemical separation for the measurement of ⁹⁹Tc by ICP-MS is acceptable. Within a wide range of ^{99m}Tc activities, the added tracer does not appreciably change the minimum detectable ⁹⁹Tc concentration of the method. It is more important, however, to decontaminate the sample from Ru carefully, because its presence noticeably enhances the minimum determination level.

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