Optimising conditions for radiolabelling of DOTA-peptides with ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu at high specific activities

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Abstract. DOTA-conjugated peptides, such as [DOTA⁰, Tyr³]octreotide (DOTATOC) and [DOTA⁰,Tyr³]octreotate (DOTA-tate), can be labelled with radionuclides such as ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu. These radiolabelled somatostatin analogues are used for peptide receptor radionuclide therapy (PRRT). Radioligands for PRRT require high specific activities. However, although these radionuclides are produced without addition of carrier, contaminants are introduced during production and as decay products. In this study, parameters influencing the kinetics of labelling of DOTA-peptides were investigated and conditions were optimised to obtain the highest achievable specific activity. The effects of contaminants were systematically investigated, concentration dependently, in a test model mimicking conditions for labelling with minimal molar excess of DOTA-peptides over radionuclide. Kinetics of labelling of DOTA-peptides were optimal at pH 4–4.5; pH <4 strongly slowed down the kinetics. Above pH 5, reaction kinetics varied owing to the formation of radionuclide hydroxides. Labelling with ⁹⁰Y and ¹⁷⁷Lu was completed after 20 min at 80°C, while labelling with ¹¹¹In was completed after 30 min at 100°C. The effects of contaminants were systematically categorised, e.g. Cd²⁺ is the target and decay product of ¹¹¹In, and it was found to be a strong competitor with ¹¹¹In for incorporation in DOTA. In contrast, Zr⁴⁺ and Hf4+, decay products of 90Y and 177Lu, respectively, did not interfere with the incorporation of these radionuclides. The following conclusions are drawn: (a) DOTApeptides can be radiolabelled at high specific activity; (b) reaction kinetics differ for each radionuclide; and (c) reactions can be hampered by contaminants, such as target material and decay products.

Keywords: DOTA - Radiolabelling - Specific activity

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Introduction

DOTA-conjugated peptides, such as the stable somatostatin analogues [DOTA⁰,Tyr³]octreotate (DOTA-tate) and [DOTA⁰,Tyr³]octreotide (DOTATOC), can be readily labelled with radionuclides such as ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu. In order for these radiolabelled peptides to be successfully used in peptide receptor radionuclide therapy (PRRT) [1, 2, 3, 4, 5, 6], high specific activities (SAs) are required. A number of biological factors dictate the need for a high SA. First, for in vivo use the amount of (radio)ligand that can be administered is limited by affinity and the amount of receptors. Above the optimal dose a further increase in ligand will increase the competition between unlabelled and labelled ligand for the same receptor and thus lower the uptake of radiolabel into receptor-positive tissue [4]. Second, for peptides that display pharmacological (side)effects, such as DOTA-substance P or DOTA-bombesin, only very small quantities of peptides may be tolerated. For the latter peptide, the amount that can be administered intravenously is limited to 0.1 nmol per minute [7]. Therefore, a high SA will reduce the total peptide amount to be administered. Third, endocytotic mechanisms that affect the cellular internalisation of peptides may become desensitised at high peptide concentrations [8], resulting in lower uptake of radiolabel into target tissue. Additionally, in vitro investigations aimed at measuring receptor-binding affinities require low concentrations of these radioligands (e.g. $10^{-10} M$) in order to measure receptor-ligand interactions accurately. Unfortunately, the need for high SA is often compromised by conflicting radiochemical parameters that determine reaction rates and yields, i.e. the rate of formation of the metal-DOTA complexes increases with pH [9], but the solubility of In³⁺, Y³⁺ and Lu³⁺ decreases with pH owing to formation

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Table 1. Production methods, targ	get materials, decay products.	, physical constants and m	naximal SA for the radionuclides considered

	⁵⁷ Co	⁶⁷ Ga	⁹⁰ Y	¹¹¹ In	¹⁷⁷ Lu
Production method	Reactor	Cyclotron	Generator	Cyclotron	Reactor
_	(n, pn)	(p, 2n)	00-	(p, 2n)	(n)
Target	⁵⁸ Ni	⁶⁸ Zn	⁹⁰ Sr	¹¹² Cd	¹⁷⁶ Lu
Decay product	⁵⁷ Fe	⁶⁷ Zn	⁹⁰ Zr	¹¹¹ Cd	¹⁷⁷ Hf
Physical constants					
$t_{1/2}$ (days)	270.9	3.26	2.67	2.83	6.71
pmoles per mCi	2080	25	20.5	21.5	51.4
Maximal SA (mCi per nmol)					
Theory ^a	0.5	40	49	46	19
In practice ^b	n.d.	10	11	22	3

Labelling conditions with 57 Co and 67 Ga were not optimised, but taken to be identical to those of 111 In: 30 min at 100°C

SA, Specific activity, expressed as mCi per nmol; n.d., not determined

^a Since 1 nmol DOTA can incorporate 1 nmol (radio)nuclide, this number indicates the maximal theoretical SA of the radiolabelled DOTA-peptides

of hydroxides [10]. Although recently labelling of DOTA analogues with ⁹⁰Y and ¹⁷⁷Lu was reported at pH 7–8 [11, 12], we encountered the above-mentioned solubility problems when high concentrations of these radionuclides were used at pH 7–8. Therefore we decided to perform this study at pH 5 or lower.

The studies presented here were undertaken to determine the optimal conditions for radiolabelling DOTApeptides, using the radionuclides ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu and DOTATOC and DOTA-tate as model reactants. We investigated parameters that influence reaction kinetics and radiochemical yields in order to define conditions that result in maximal achievable SAs. This report also summarises the effects of ever-present contaminants, such as nuclides formed by decay of the radionuclides.

Materials and methods

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Ligands and radionuclides. DOTA-peptides were dissolved in 0.01–0.05 *M* acetic acid in Milli-Q water. As the buffer, 25 m*M* sodium ascorbate (quencher) in 50 m*M* sodium acetate was used. ⁹⁰Y was from Pacific Northwest National Laboratory (Richland, Wash., USA) and from MDS Nordion (Fleurus, Belgium). ¹⁷⁷Lu was from Missouri University Reactor Research (MURR, St. Louis, Mo., USA) and from NRG (Petten, The Netherlands). ⁵⁷CoCl₂ in 0.1 *M* HCl was from MDS Nordion (Vancouver, BC, Canada). ¹¹¹InCl₃ was from Mallinckrodt Medical (Petten, The Netherlands). All these radionuclides were delivered in 0.01–0.2 *N* HCl. ⁶⁷GaCl₃ in 0.7–0.8 *M* HCl was from MDS Nordion (Vancouver, BC, Canada) in 0.05 *M* HCl.

Reaction conditions: the effects of pH and temperature. The experiments were performed in small volumes (typically 40–75 µl)

^b Highest value achieved; this implies a ratio of DOTA over radionuclide [practice (b) over theory(a)] of 4, 4 1/2 and 2 1/4 for ⁶⁷Ga, ⁹⁰Y and ¹¹¹In, respectively

using double-sealed plastic reaction tubes (PCR thermocycler tubes, max volume 125 μ l, MoBiTec, ITK Diagnostics, Uithoorn, The Netherlands). Heating was performed in a temperature-controlled heating block (Grant, Fisher Scientific, Zoeterwoude, The Netherlands). The pH of the reaction mixture was measured after the reaction. All experiments were carried at least in duplicate, using at least three different production batches of the radionuclides from the suppliers mentioned above. All chemicals were purchased from Aldrich Chemicals (Zwijndrecht, The Netherlands), and were of the highest analytical grade available. The radiochemical purity of the radiolabelled DOTATOC and DOTA-tate was studied at room temperature in the presence of 4 mM DTPA pH 5. High-performance liquid chromatography and instant thin-layer chromatography were performed as described previously (see [13] and [14] respectively)..

Maximal achievable SA, and effects of contaminants. In order to determine the maximal achievable SA (\geq 98% incorporation of ⁶⁷Ga, ⁹⁰Y, ¹¹¹In or ¹⁷⁷Lu), radiolabelling was performed using a constant amount of the radionuclides and increasing amounts of ligand. To investigate the effects of the contaminants present, experiments were performed adding known amounts of contaminants to the reaction vial at the start of the radiolabelling. For all radionuclides considered, the target material and the decay products are summarised in Table 1, and all were tested concentration dependently. The starting conditions in the absence of added contaminants were chosen to be critical, using a low mol/mol ratio of DOTA over radionuclide.

Results

Reaction kinetics

Reaction kinetics with ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu were found to be optimal at pH 4–4.5, with a steep decrease at lower

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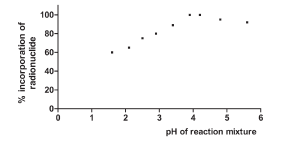


Fig. 1. Formation of 177 Lu-DOTATOC as a function of pH after 20 min at 80°C, as measured by the % incorporation of the radionuclide. Similar results were found with 90 Y and with DOTA-tate as ligand

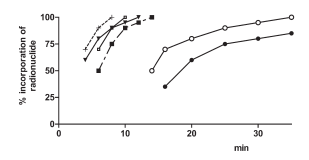


Fig. 2. Formation of radiolabelled DOTATOC at pH 4 as a function of time of incubation, as measured by the % incorporation of the radionuclide. Similar results were found with DOTA-tate as ligand. +, ¹⁷⁷Lu at 80°C; ♥, ¹⁷⁷Lu at 100°C; ■, ⁹⁰Y at 80°C; □, ⁹⁰Y at 100°C; ●, ¹¹¹In at 80°C; ○, ¹¹¹In at 100°C

pH and a slow decrease at higher pH (Fig. 1). In addition, % incorporation of ¹¹¹In and ¹⁷⁷Lu at pH \geq 5 became non-reproducible: after centrifugation of these reaction vials, precipitation was found. The reaction kinetics were also found to be time- and temperature-dependent: reactions were complete with ⁹⁰Y and ¹⁷⁷Lu after 20 min at 80°C, and with ¹¹¹In after 30 min at 100°C (Fig. 2). Table 1 shows the highest achieved SAs of ⁶⁷Ga, ⁹⁰Y and ¹¹¹In, implying a mol/mol ratio of DOTA over radionuclide of 4, 4 1/2 and 2 1/4, respectively. For ¹⁷⁷Lu the mol/mol ratio was even 1.2 (see also Discussion). We were still able to label at high SAs 2 weeks after the production of ¹⁷⁷Lu (data not shown).

Effects of contaminants of maximal achievable SA

To validate the test model the effect of the addition of unlabelled Y, In and Lu to the corresponding radionuclide was investigated. As expected, the dilution of the SA of the radionuclide decreased its incorporation in the DOTA-chelator in a concentration-dependent manner (Fig. 3, Table 2). The addition of nuclides such as Hf, Zr and Sr had no effect on the % incorporation of the radionuclides, indicating that these nuclides are not competitors under these reaction conditions. In contrast, the ef-

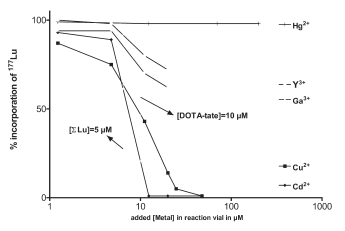


Fig. 3. Effects of contaminants on the incorporation of ¹⁷⁷Lu in DOTA-tate by the controlled addition of non-radioactive nuclides

Table 2. Effects of different concentrations of metal ions, as contaminants in the reaction vial, on incorporation of radionuclides in DPTA-ligand

O	+	++
Ag+	Ga ³⁺ Y ³⁺	Cd ²⁺
Hf ⁴⁺	Y ³⁺	Cd ²⁺ Co ²⁺ Cu ²⁺
Hg ²⁺		Cu ²⁺
Hg ²⁺ Sr ²⁺		In ³⁺
Zr ⁴⁺		Fe ²⁺
		Lu ³⁺
		Ni ²⁺
		Zn ²⁺

0, ≤10% at 10 µ*M*; +, ≥10% at 1–10 µ*M*; ++, ≥10% at 1 µ*M*

fect of addition of nuclides such as Fe and Cd clearly showed that they are strong competitors for the incorporation of radionuclide in the DOTA-chelator, as shown in Fig. 3 and Table 2. Table 1 also contains data on labelling experiments with ⁵⁷Co and ⁶⁷Ga using the same conditions as those for ¹¹¹In, 30 min at 100°C. With the Mallinckrodt-produced ⁶⁷Ga, pH was difficult to control since the radionuclide is delivered in 0.7–0.8 *M* HCl. More importantly, the [Zn] (Zn is the target and decay product of ⁶⁷Ga, see Table 1) was measured by ICP (data not shown) and found to exceed [Ga] frequently by >100-fold, with a dramatic effect on achieving high SAs.

Discussion

We recently reported that Cd is a strong competitor for ¹¹¹In incorporation in the DOTA-chelator, and since ¹¹¹Cd is formed from decaying ¹¹¹In, the highest SA is achieved immediately after the production of ¹¹¹In [15]. Analogously, achieving a high SA with ⁶⁷Ga will be very difficult owing to the [^{67/68}Zn], present from target (⁶⁸Zn)

and formed during decay. Even if the $[^{68}Zn]$ is low or zero at the end of production of ^{67}Ga , after one half-life of ^{67}Ga the $[^{67}Ga]=[^{67}Zn]$.

This study provides insights into the effects of contaminants in daily radiolabelling practice when high SAs are required. It also improves the interpretation of the concentrations of contaminants as mentioned in the data sheet provided by the manufacturer of the radionuclides. For instance, the data sheet of 90 Y (MDS Nordion) states that the maximal concentration of Zn in 90 Y will not exceed the level of 30 µg per Ci of 90 Y. If this is so, however, it implies that the mol/mol ratio for Zn will be more than 20 times that for Y; achieving a high SA would then not be possible.

In Table 1 the highest achieved SAs of ⁶⁷Ga, ⁹⁰Y, ¹¹¹In and ¹⁷⁷Lu are presented, and imply a mol/mol ratio of DOTA over radionuclide of 4, 4 1/2, 2 1/4 and 6, respectively. This reflects the reaction conditions, including pH, temperature and the use of very pure reactants. From Table 2 it can be seen that Hf is not a competitor for ¹⁷⁷Lu under our reaction conditions. This implies that the ingrowth of Hf has no consequences for the maximal achievable SA. The rate of incorporation is >99%, even at a mol/mol ratio (DOTA-peptide over ¹⁷⁶⁺¹⁷⁷Lu) of 1.2. In addition, a high SA can still be achieved 2 weeks after the production of ¹⁷⁷Lu, confirming that Hf is not a competitor for Lu in the incorporation in DOTA. Although stability constants of DOTA with many nuclides are available in the NIST database [16], data on reaction kinetics are scarce. However, from the data presented here it can be concluded that the reaction kinetics of Lu, Y and In with DOTA are in the order: Lu>Y>>In.

In conclusion, DOTA-peptides can be radiolabelled at high SA. Reaction kinetics differ for each radionuclide: with ⁹⁰Y and ¹⁷⁷Lu, conditions were optimal at pH 4–4.5 and the reactions were complete after 20 min at 80°C, while labelling with ¹¹¹In was completed after 30 min at 100°C. Reactions can be hampered by contaminants, such as target material and decay products.

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