Improved GMP-compliant multi-dose production and quality control of 6-[18F]fluoro-L-DOPA

Luurtsema, Geert; Boersma, Hendrikus; Schepers, Marianne; de Vries, Michel; Maas, Bram; Zijlma, Rolf; de Vries, Erik; Elsinga, Philippus

Published in:
EJNMMI Radiopharmacy and Chemistry

DOI:
10.1186/s41181-016-0009-1

IMPORTANT NOTE: You are advised to consult the publisher's version (publisher's PDF) if you wish to cite from it. Please check the document version below.

Document Version
Publisher's PDF, also known as Version of record

Publication date:
2016

Link to publication in University of Groningen/UMCG research database

Citation for published version (APA):
Luurtsema, G., Boersma, H., Schepers, M., de Vries, M., Maas, B., Zijlma, R., de Vries, E., & Elsinga, P. (2016). Improved GMP-compliant multi-dose production and quality control of 6-[18F]fluoro-L-DOPA. EJNMMI Radiopharmacy and Chemistry, 1(7), Article 7. https://doi.org/10.1186/s41181-016-0009-1

Copyright
Other than for strictly personal use, it is not permitted to download or to forward/distribute the text or part of it without the consent of the author(s) and/or copyright holder(s), unless the work is under an open content license (like Creative Commons).

The publication may also be distributed here under the terms of Article 25fa of the Dutch Copyright Act, indicated by the “Taverne” license. More information can be found on the University of Groningen website: https://www.rug.nl/library/open-access/self-archiving-pure/taverne-amendment.

Take-down policy
If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

Downloaded from the University of Groningen/UMCG research database (Pure): http://www.rug.nl/research/portal. For technical reasons the number of authors shown on this cover page is limited to 10 maximum.

Download date: 03-12-2024
Improved GMP-compliant multi-dose production and quality control of 6-[^18]F]fluoro-L-DOPA

G. Luurtsema*, H. H. Boersma, M. Schepers, A. M. T. de Vries, B. Maas, R. Zijlma, E. F. J. de Vries and P. H. Elsinga

* Correspondence: g.luurtsema@umcg.nl
Department of Nuclear Medicine and Molecular Imaging, University of Groningen, University Medical Center Groningen, Hanzeplein 1, PO Box 30.001 9700 RB Groningen, The Netherlands

Abstract

Background: 6-[^18]F]Fluoro-L-3,4-dihydroxyphenylalanine (FDOPA) is a frequently used radiopharmaceutical for detecting neuroendocrine and brain tumors and for the differential diagnosis of Parkinson’s disease. To meet the demand for FDOPA, a high-yield GMP-compliant production method is required. Therefore, this study aimed to improve the FDOPA production and quality control procedures to enable distribution of the radiopharmaceutical over distances. FDOPA was prepared by electrophilic fluorination of the trimethylstannyl precursor with [^18]F]F₂, produced from [^18]O₂ via the double-shoot approach, leading to FDOPA with higher specific activity as compared to FDOPA which was synthesized, using [^18]F]F₂ produced from ^20Ne, leading to FDOPA with a lower specific activity. The quality control of the product was performed using a validated UPLC system and compared with quality control with a conventional HPLC system. Impurities were identified using UPLC-MS.

Results: The [^18]O₂ double-shoot radionuclide production method yielded significantly more [^18]F]F₂ with less carrier F₂ than the conventional method starting from ^20Ne. After adjustment of radiolabeling parameters substantially higher amounts of FDOPA with higher specific activity could be obtained. Quality control by UPLC was much faster and detected more side-products than HPLC. UPLC-MS showed that the most important side-product was FDOPA-quinone, rather than 6-hydroxydopa as suggested by the European Pharmacopoeia.

Conclusion: The production and quality control of FDOPA were significantly improved by introducing the [^18]O₂ double-shoot radionuclide production method, and product analysis by UPLC, respectively. As a result, FDOPA is now routinely available for clinical practice and for distribution over distances.

Keywords: PET, Radionuclide production, Radiochemistry, Automation, Quality control

Background

Positron emission tomography (PET) with the radiopharmaceutical 6-[^18]F]fluoro-L-3,4-dihydroxyphenylalanine (FDOPA) is frequently used for measuring dopamine metabolism in vivo. Clinical applications of FDOPA PET are the detection and staging of neuroendocrine and brain tumors, and the differential diagnosis between Parkinson’s disease and other degenerative disorders of the central nervous system (Jager et al. 2008; Santhanam and Taïeb 2014). Since clinical indications for FDOPA PET imaging
are increasingly being established, there will be a high demand for this radiopharmaceutical. Because of the growing clinical interest, there is a need for a high-yield GMP-compliant production method for FDOPA that enables the distribution of this radiopharmaceutical to other hospitals.

So far, the most common synthesis route to produce FDOPA is via an electrophilic substitution reaction with $[^{18}\text{F}]$fluorine gas ($[^{18}\text{F}]F_2$). This approach has already been used for more than one decade (De Vries et al. 1999). A disadvantage of this synthesis route is the complex production of the $[^{18}\text{F}]F_2$ labeling reagent, giving low yields and requires the use of carrier fluorine gas. As a consequence of the use of $[^{18}\text{F}]F_2$, the radiochemical yield of the radiopharmaceutical is maximally 50 % and only a product with a low specific activity can be produced. The consequence of using FDOPA with low specific activity is that it could lead to adverse reactions such as flushes (Koopmans et al. 2006). To increase the specific activity and the relatively low radiochemical yield of FDOPA, two general approaches have been pursued: first, to develop a radiolabeling method for FDOPA using nucleophilic $[^{18}\text{F}]$fluoride (Kuik et al. 2014), and second, to improve the $[^{18}\text{F}]F_2$ radionuclide production (Bishop et al. 1996; Bergman and Solin 1997) and the electrophilic labeling method. Considerable efforts were invested in the development of new synthesis approaches based on non-carrier-added $[^{18}\text{F}]$fluoride as the starting material (Shen et al. 2009). These approaches generally required complex multistep synthetic strategies, resulting in low overall radiochemical yields. At this moment, new strategies are under development to obtain higher radiochemical yields (Lemaire et al. 2015; Zlatopolvskyi et al. 2015).

In this article, we report on our efforts to optimize the FDOPA production using an improved method for the production of $[^{18}\text{F}]F_2$ for the electrophilic synthesis of FDOPA. The installation of a Cyclone 18 twin cyclotron (IBA, Belgium) recently allowed us to change our $[^{18}\text{F}]F_2$ production method from the $^{20}\text{Ne}(d,\alpha)^{18}\text{F}$ single-shoot method (De Vries et al. 1999) to the $^{18}\text{O}(p,n)^{18}\text{F}$ double-shoot method (Bishop et al. 1996). As the Cyclone 18 twin cyclotron has a newly designed small volume target, the double-shoot production method needs less carrier $F_2$ gas, which increases the specific activity of $[^{18}\text{F}]F_2$ in comparison to the single-shoot method using an MC17 Scanditronix cyclotron.

Finally, the overall aim of this study is to improve the radionuclide production and the electrophilic labeling method for FDOPA via $[^{18}\text{O}]_2$ double shoot approach using Cyclone 18 twin cyclotron. This improved method, leads to FDOPA with higher specific activity (FDOPA-H). The percentage radiochemical yield (%) and the amount of $[^{18}\text{F}]$FDOPA radioactivity (practical yield), the radiochemical purity, specific activity is compared with conventional approach using MC17 Scanditronix cyclotron, which leads to FDOPA with a lower specific activity (FDOPA-L). Furthermore, an improved, more sensitive and accurate QC analysis for FDOPA is developed and validated using UPLC.

**Methods**

**Materials for synthesis and analysis**

For preparation of FDOPA the 6-trimethylstannyl-L-DOPA precursor was purchased from ABX (Germany). Dry chloroform >99 % for spectroscopy stabilized with amylene (stored on molecular sieves) and ammonium dihydrogen phosphate p.a. were obtained from Acros (Belgium) and 47 % hydrobromic acid, 25 % ammonia and diammonium
hydrogenphosphate p.a. from Merck (Germany). A sterile solution for injection of ascorbic acid (100 mg/ml) was purchased from Centrapharm (Netherlands). A sterile 0.1 M sodium acetate solution, pH 4.7, was prepared in the hospital pharmacy. Reference compounds for calibration curves of FDOPA, L-DOPA and 6-hydroxy-DL-DOPA were purchased from respectively ABX (Germany) and Sigma Aldrich (Netherlands). HPLC supra gradient acetonitrile, used as solvent for LC-MS analysis, was obtained from Biosolve (Netherlands), ~98 % formic acid was bought from Fluka (Netherlands) and LC-MS grade water (>18.2 MΩ) was purchased from the Department of Clinical pharmacy & Pharmacology, UMCG. The reference compound leucine encephalin for calibration of the MS detector was purchased from Merck (Germany). For calibration of the MS, a solution was prepared of 200 μL 10 % formic acid, 100 μL 0.1 M sodium hydroxide (Merck, Germany) and 20 ml 80 % acetonitrile in water.

The production of [18F]F₂ using deuterons and proton particles

Deuteron particles were accelerated using a MC17 cyclotron. Carrier-added [18F]F₂ was produced via the 20Ne(d,α)18F nuclear reaction. A 350-ml nickel target was filled with 0.25 % F₂ (150 μmol) in neon and irradiated with 30 μA 8.5 MeV deuterons for 2 h. After irradiating the F₂ target, [18F]F₂ gas was transported via ~20 m 1/8 in. stainless steel tubing from Scanditronix MC-17 cyclotron to production laboratory. Passivation of transport lines was required.

Proton particles were accelerated using Cyclone 18 twin cyclotron. Production of [18F]F₂ via the 18O(p,n)18F nuclear reaction was performed. A 35-ml aluminum target was filled with 18O-enriched oxygen gas ([18O]₂, > 97 %) at a pressure of 20 bar and irradiated with 35 μA 18 MeV protons for 1 h. The produced radioactivity is recovered from the target wall via a double shoot method. Therefore, the target was emptied to recover the [18O]₂ gas using a cryogenic system and a vacuum was created inside the target to remove traces of [18O]oxygen. The target was filled with 24 bar 0.5 % F₂ (75 μmol) in neon and irradiated for another 10 min. Passivation of transport lines was not necessary.

Transport of [18F]F₂ gas from the Cyclone 18 twin cyclotron to the GMP laboratory is through 25 m long 1/8 in. stainless steel tubing with a flow of 250 ml.min⁻¹.

Synthesis of FDOPA

The synthesis procedure using [18F]F₂ produced via the 20Ne(d,α)18F nuclear reaction was based on previous described synthesis (De Vries et al. 1999), but with some improvements and modifications. The modifications were; the activity was trapped into the reactor vial in the synthesis module (Raytest, Synchrom FDOPA F2, see schematic overview in Fig. 1) which was filled with 55–65 mg 6-trimethylstannyl-L-DOPA precursor in 3 ml of chloroform and cooled at a temperature of -20 °C. After trapping the [18F]F₂ gas in the reaction mixture, the chloroform was evaporated at 70 °C. Thereafter, the hydrolysis using 2 ml 47 % hydrobromic acid was performed in 5 min at 130 °C. After hydrolysis, the reactor was cooled to 40 °C and 1.3 ml 25 % aqueous ammonia was added, followed by 1.3 ml buffer solution consisting of 1 M ammonium dihydrogen phosphate and 1 M diammonium hydrogenphosphate to neutralize the solution prior HPLC purification. A schematic overview of the synthesis is given in Fig. 1 and a schematic overview of the synthesis is illustrated in Fig. 2.
The product was eluted with a flow of 3 ml/min from two consecutive (connected in series) preparative HPLC columns (Hamilton PRP1 polymeric column in series (250 × 10 mm, 10 μm), Switzerland) with sterile 0.1 M sodium acetate, pH 4.7 and collected in a sterile vial with 100 μL ascorbic acid solution (100 mg/ml) to prevent oxidation. Aseptic filtration of the collected product was done via Millex LG filter and divided in three vials, patient batch, QC sample and a retain sample. Filtration and dispensing was performed in a hot cell with class A classification. The total synthesis time was 50–60 min.

One of the release criteria is the integrity of the used filter. To reduce the finger dose of the technicians we implemented an online pressure hold test directly in the module.

The synthesis procedure using $^{18}$F$\text{F}_{2}$ produced via the $^{18}\text{O}(p,n)^{18}\text{F}$ nuclear reaction was performed like described above, but with a slight modification. Due to low amount of carrier F$_2$ gas, the amount of stannyl precursor was decreased to 30 mg (48.5 μmol).

**Quality control of FDOPA and identification of side products**

For the determination of the specific activity, the radiochemical and chemical purity of FDOPA an QC analysis was performed on conventional HPLC system. To improve the analysis with respect to retention times and resolution we implemented an Acquity Ultra Performance Liquid Chromatography (UPLC®) with UV and an online radioactivity detector from Berthold (Flowstar LB513) in our laboratory. For a direct comparison on retention times of L-DOPA and FDOPA and the resolution of the separation, same
samples were injected on the two analytical methods. The conventional HPLC method was performed on Waters system (pump model 515 and tunable UV detector model 486), using a Hamilton PRP-1 150 × 4,1 mm column, with a eluent of sodium citrate 8,71 g and sodium dodecylsulfon 0,8 g in 900 ml H2O pH 2,9/100 ml acetonitrile and a flow rate of 1 ml/min.

A UPLC method was executed, using a Waters ACQUITY UPLC® HSS T3 1.8 μm, 3.0 × 50 mm analytical column with an eluent of 0.05 M phosphate buffer pH 2.5 with an isocratic flow of 0.8 ml/min. For both methods a wavelength of 254 nm was used for UV-detection. As specified in the European pharmacopeia 8, FDOPA, L-DOPA, 6-hydroxy-DOPA were used as reference standards. An additional test on the UPLC was performed to provide information about the stability of FDOPA during the time of transport (max 8 h). Over a period of 8 h at room temperature, samples were taken at different time points for analysis of the radiochemical purity using UPLC.

Identification of specific compounds like; FDOPA, L-DOPA, 6-hydroxy-DOPA was performed on a Waters (Milford, MA, USA) UPLC H-class system coupled to a Waters Xevo® G2 QToF mass spectrometer (UPLC-MS). Samples (1 μL) were injected onto a Waters 5.0 × 2.1 mm I.D., 1.7 μm Ethylene Bridged Hybrid (BEH) C18 column and eluted at a flow rate of 0.6 ml/min with a 1.5-min linear gradient of 10 mM ammonium bicarbonate pH 9.4 and acetonitrile, starting at 2 % and ending at 80 % acetonitrile.

Negative electrospray ionization in resolution mode was used. Both MS and MS^E scans were performed simultaneously in a mass range from 50 to 1200 Da. MS^E is a method of data acquisition that records exact mass precursor and fragment ion information from every detectable component in a sample. The following settings were used: capillary voltage 0.5 kV, sampling cone 65 V, extraction cone 4.0 V, source temperature 150 °C and desolvation temperature 500 °C. The collision energy was set to 6 V during the MS acquisition and it was ramped from 20 to 45 during the MS^E acquisition. Leucine-Enkephalin was used as a lock mass (m/z 554.2615 Da → negative mode) with a concentration of 2 ng/μL, at a flow rate of 20 μL/min. The capillary voltage was set to 2.5 kV.

Results

[^18F]F₂ production

The radioactivity amount of [^18F]F₂ using irradiation with deuterons ranged between 4 and 8 GBq (70-100 MBq. μAh⁻¹) and using proton irradiation, the [^18F]F₂ production ranged between 25 and 35 GBq (800-1100 MBq. μAh⁻¹). Both activities were measured under the same conditions in the reaction vial of the module, 10 min after EOB. The total time, including irradiations, flushing, pre- and double shoot beam, was 150 min using deuteron and 90 min using the double shoot proton procedure.

FDOPA synthesis

Results of the synthesis of FDOPA with [^18F]F₂ from both production methods are summarized in Table 1. The radiochemical yield (%) for the production of FDOPA-H was higher in comparison to the FDOPA-L-production. The radiochemical yield (%) was calculated from trapped activity in the reactor. The average overall amount of [^18F]FDOPA and the average specific activity were measured at the end of synthesis (EOS). The average specific activity of FDOPA-H was > 15 times higher compared to FDOPA-L.
FDOPA quality control and identification of impurities

Concentrations of FDOPA and L-DOPA meet the pharmacopeia specifications (FDOPA 15 mg/dose and L-DOPA 1 mg/dose, respectively) and were used to calculate the SA which is a pre-release criterion (Table 2).

The total analysis time using UPLC was 3 min, whereas execution of the HPLC method took 15 min. The retention times of FDOPA and L-DOPA measured with UPLC were 1.5 and 1.1 min, respectively. The retention time on HPLC for FDOPA was 7 min and 5 min for L-DOPA. The calculated resolution factor (Rf) was 3.0 and 1.4 for UPLC and HPLC, respectively.

The radiochemical purity, the specific activity and the concentration FDOPA and L-DOPA were determined using the UPLC method and all met the pharmacopeia requirements. FDOPA was at least stable (>95 % RCP) during 8 h after synthesis. An example of QC chromatogram performed with UPLC and HPLC coupled with UV and radioactivity detector is given in Fig. 3.

Additionally, both FDOPA and L-DOPA were identified using exact mass (n = 6 for FDOPA-H, n = 6 FDOPA-L). However, identification of 6-hydroxy-DOPA in the drug product was not possible. In contrast, a molecule with almost the same mass as 6-hydroxy-DOPA (212.036 Da), but with a different retention time as the reference standard was found. Calculation of the elemental composition from the exact mass and fragment ion information using MS² scans (Fig. 4b) proved that this product is 2-amino-3-(6-fluoro-3,4-dioxocyclohexa-1,5-dienyl)propanoic acid (FDOPA-quinone), which can be formed by oxidation of FDOPA. FDOPA–quinone was only detected in lower specific activity FDOPA samples (Fig. 4a). We did not find changes in FDOPA-quinone concentration over time. Because no reference compound FDOPA-quinone was

### Table 1

| Method (18F₂) production | Number of productions (n) | Radiochemical yield (%) | Specific activity (GBq/mmol) | Radio-chemical Purity (%) | Amount of FDOPA activity (MBq) |
|-------------------------|---------------------------|-------------------------|-----------------------------|--------------------------|-----------------------------|
| FDOPA-Lower             | ²⁰Ne(d,α)¹⁸F single-shoot  | 68                      | 15 ± 5                      | 8.5 ± 3.3                | 97 ± 3                      | 526 ± 192                  |
| FDOPA-Higher            | ¹⁸O(p,n)¹⁸F double-shoot   | 42                      | 23 ± 4                      | 121 ± 27                 | 97 ± 3                      | 4521 ± 967                |

### Table 2

Concentrations and identification of FDOPA and L-DOPA (g/l) and side products in FDOPA-L and FDOPA-H batches analyzed with UPLC-MS (both n = 6)

| FDOPA-L | MC17- cyclotron | FDOPA 214.052 Da (g/L) | L-DOPA 196.061 Da (g/L) | 6-hydroxy-DL-DOPA 212.056 Da | FDOPA-quinone 212.036 Da |
|---------|----------------|------------------------|-------------------------|------------------------------|------------------------|
| X ± SD  | 1.21 ± 0.15    | 0.04 ± 0.04            | -                       | +                           |

| FDOPA-H | Cyclone 18 twin cyclotron | FDOPA 214.052 Da (g/L) | L-DOPA 196.061 Da (g/L) | 6-hydroxy-DL-DOPA 212.056 Da | FDOPA-quinone 212.036 Da |
|---------|--------------------------|------------------------|-------------------------|------------------------------|------------------------|
| X ± SD  | 0.53 ± 0.10              | 0.007 ± 0.003          | -                       | -                           |

+ = side product detected and – means not detectable
available, it was not possible to quantify FDOPA-quinone concentrations in the samples (see Fig. 4a and b for the mass spectra).

**Discussion**

This study demonstrates that optimized radionuclide production, via a proton double-shoot method on $^{18}$O$_2$ leads to significant higher amounts of FDOPA with higher specific activity.

Although the ratio $\mu$mol precursor versus carrier F$_2$ was equal for both syntheses, the radiochemical yield with the improved FDOPA-H method was higher compared to the conventional FDOPA-L. This phenomenon could be due to the higher purity of $[^{18}\text{F}]$F$_2$. Intensive cleaning of the target using flushing and vacuum was always done after the first shoot and therefore the $[^{18}\text{F}]$F$_2$ produced via the double-shoot method may contain less chemical impurities and therefore its reaction with the $[^{18}\text{F}]$FDOPA precursor appears to be more efficient. Taken together, it is evident that the new method leads to significantly higher radiochemical yields and together with higher radionuclide production yield of $[^{18}\text{F}]$F$_2$ with less carrier F$_2$ consequently more amount of FDOPA with higher specific activity was achieved.
As a result of the higher specific activity, lower amounts of carrier FDOPA need to be administered to the patient, which reduces the risk on side effects, such as carcinoid crisis (Koopmans et al. 2006).

In general, it can be concluded that, although the complex radionuclide production of $^{18}$F$^2$ gas, the radiochemistry via electrophilic substitution is quite straightforward resulting in short reaction times. Disadvantages of using $^{18}$F$^2$ gas, as labeling reagent,
is the theoretical maximum radiochemical yield of 50% and the corrosive character of [\(^{18}\text{F}\)]F\(_2\) gas which can lead to malfunctioning of valves and equipment.

During this study, 6-hydroxy-DOPA could not be detected in the produced FDOPA batches, irrespective of its specific activity. Another impurity, the so called FDOPA-quinone, could be detected, but only in FDOPA batches with lower specific activity. The identification of this impurity was based on the exact mass, as determined by UPLC-MS, in combination with the proposed fragmentation pattern for this product. The difference in the exact mass of 6-hydroxy-DOPA and FDOPA-quinone is only 0.02 Da, but could be readily detected by UPLC-MS. Together with the observed fragments, FDOPA-quinone could be reliably identified with this method. Why FDOPA-quinone was only identified in the FDOPA-L batch is not completely clear. The major difference in both production methods are the used concentrations of precursor, radionuclide production of [\(^{18}\text{F}\)]F\(_2\) and the differences in SA of FDOPA.

Performing state-of-the-art quality control analysis using UPLC in combination with radioactivity detector and MS-TOF leads to accurate identification of the impurities. Using UPLC-MS, impurities, like the FDOPA-quinone, in low molecular concentrations can be identified. This can provide more accurate information which is needed for the release of radiopharmaceuticals. In this study, we have shown that 6-hydroxy-DOPA is not formed as an impurity in FDOPA batches. There is no logical chemical pathway that could lead to the formation of this impurity, as hydrolysis of the trimethylstannyl precursor should lead to the formation of L-DOPA (and trimethyltin hydroxide). Considering the similarity in molecular weight, it seems plausible that FDOPA-quinone has been falsely identified as 6-hydroxy-DOPA in the past. Therefore, an adjustment of the FDOPA monograph in the European Pharmacopeia should be considered.

It is expected that the population of patients with endocrine tumors will grow in the coming years and the application of FDOPA will grow for differential diagnosis between Parkinson's disease and other degenerative disorders. In order to meet with the increasing demand for FDOPA PET, we improved the [\(^{18}\text{F}\)]F\(_2\)-based electrophilic production method of FDOPA to make it suitable for production of FDOPA for multiple patients in our institute and distribution over distance. The fact that the described FDOPA production is GMP compliant, our institute was granted an exemption to have a marketing authorization by the government to distribute FDOPA to other hospitals without a dossier for marketing authorization. This exemption is applicable for the Netherlands, because FDOPA is a licensed radiopharmaceutical, but not available for use, since the current manufacturing authorization holder does not produce FDOPA. With the described production method, production of FDOPA batches for distribution to other hospitals became feasible.

**Conclusion**

The production and quality control of FDOPA were significantly improved by introducing the [\(^{18}\text{O}\)]\(_2\) double-shoot radionuclide production method, improved synthesis, and product analysis by UPLC, respectively. As a result, FDOPA is now routinely available for clinical practice and for distribution over distances. Adjustment of the FDOPA monograph in the European Pharmacopeia should be considered.

**Competing interests**

All authors declare that they have no competing interests.
Authors’ contributions
GL analysed the data, and drafted the manuscript. HHB helped with the design of the experiments and with the drafting of the manuscript. MS designed and performed the mass spectrometry analysis. AMTdV and BM performed the radio-synthesis and implementation of the GMP production. RZ designed and performed QC analysis. EFJdV helped with the design of the radio-synthetic protocols and interpretation of the results and critically revised the manuscript. All authors read and approved the final manuscript.

Acknowledgements
This work was financially supported by NWO, project number 91111007 (UPLC-MS). We thank Michel Ghyoot from IBA RadioPharma Solutions, Louvain-la-Neuve, Belgium for his work on the radionuclide production.

Received: 29 January 2016 Accepted: 14 March 2016
Published online: 04 April 2016

References
Bergman J, Solin O. Fluorine-18-labeled fluorine gas for synthesis of tracer molecules. Nucl Med Biol. 1997;24:677–83.
Bishop A, Satyamurthy N, Bida G, Barrio JR. Chemical reactivity of the 18F electrophilic reagents from the 18O(p, n)18F gas target systems. Nucl Med Biol. 1996;23:559–65.
De Vries EFJ, Luurtsema G, Brusselman M, Eisinga PH, Vaalburg W. Fully automated synthesis module for the high yield one-pot preparation of 6-[18 F]fluoro-L-DOPA. Appl Radiat Isot. 1999;51:389–94.
Jager PL, Chirakal R, Marriott CJ, Brouwers AH, Koopmans KP, Guleschyn KY. 6-L-18F-fluorodihydroxyphenylalanine PET in neuroendocrine tumors: basic aspects and emerging clinical applications. J Nucl Med. 2008;49:573–86.
Koopmans KP, de Vries EG, Kema IP, Eisinga PH, Neels OC, Stutter WU, et al. Staging of carcinoid tumours with 18F-DOPA PET: a prospective, diagnostic accuracy study. Lancet Oncol. 2006;7:728–34.
Kulk WI, Kema IP, Brouwers AH, Zijlm N, Neumann KD, Dierckx RAJO, et al. In Vivo Biodistribution of No-Carrier-Added 6-18F-Fluoro-3,4-Dihydroxy-L-Phenylalanine (18F-DOPA), Produced by a New Nucleophilic Substitution Approach, Compared with Carrier-Added 18F-DOPA, Prepared by Conventional Electrophilic Substitution. J Nucl Med. 2014;56:106–12.
Lemaire C, Libert L, Franci X, Genon J-L, Kuci S, Giacomelli F, et al. Automated production at the curie level of no-carrier-added 6-[18 F]fluoro-dopa and 2-[18 F]fluoro-tyrosine on a FASTlab synthesizer. J Label Compd Radiopharm. 2015;58(7):281–90.
Santhanam P, Taleb D. Role of 18F-FDOPA PET/CT imaging in endocrinology. Clin. Endocrinol. 2014;81(6):789–98.
Shen B, Ehrlöchmann W, Uebel M, Machulla HJ, Reischl G. Automated synthesis of n.c.a. [18F]FDOPA via nucleophilic aromatic substitution with [18F]fluoride. Appl Radiat Isot. 2009;67:1650–3.
Zlatopolskiy BD, Zischler J, Urusova EA, Endepols H, Kordys E, Freauendorf H, et al. A Practical One-Pot Synthesis of Positron Emission Tomography (PET) Tracers via Nickel-Mediated Radiofluorination. ChemistryOpen. 2015;4(4):457–62.