The diagnostic value of $^{124}$I-PET in patients with differentiated thyroid cancer

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Abstract

Background The purpose of this prospective study was to evaluate the clinical diagnostic value of iodine-124 ($^{124}$I)-positron emission tomography (PET) in patients with advanced differentiated thyroid carcinoma (DTC) and to compare the $^{124}$I-PET imaging results with the $^{131}$I whole-body scan (WBS).

Materials and methods Twenty patients with histologically proven advanced DTC (including T4, extra-nodal tumour growth, or distant metastases) underwent diagnostic $^{131}$I-WBS, $^{124}$I-PET scan, and post-treatment $^{131}$I-WBS 4 months after ablation. The findings on the $^{124}$I-PET were compared with the findings on the diagnostic and post-therapeutic $^{131}$I-WBS and were also correlated with radiologic and/or cytological investigations.

Results $^{124}$I-PET vs diagnostic $^{131}$I-WBS. Eleven patients showed uptake on the $^{124}$I-PET. Only 3 of these 11 patients also showed uptake on the diagnostic $^{131}$I scan, but the uptake was more clearly visible and the abnormalities were more extensive on the $^{124}$I-PET.

$^{124}$I-PET vs post-treatment $^{131}$I-WBS. Eleven patients showed uptake on the $^{124}$I-PET, which was also visible on the post-treatment scan in nine patients; in the other two patients, no uptake was observed on the post-treatment scan and no anatomical localisation could be confirmed. Two patients showed only uptake on the post-treatment scan without uptake on the $^{124}$I-PET: in one, the uptake was confirmed by MRI, and in the other, no anatomical localisation was found. In seven patients, no uptake was observed on both the scans.

Conclusion $^{124}$I-PET proved to be a superior diagnostic tool as compared to low-dose diagnostic $^{131}$I scans and adequately predicted findings on subsequent high-dose post-treatment $^{131}$I scans.

Keywords Iodine-124 · Positron emission tomography · Differentiated thyroid cancer · Diagnostic value

Introduction

Papillary and follicular thyroid cancer are the most frequent histological types of thyroid cancer (85–90%) [1, 2]. Total
thyroidectomy with or without lymph node dissection is the initial therapy, followed by radioactive iodine therapy.

In the routine follow-up of low-risk patients, diagnostic $^{131}$I whole-body scanning (WBS) is not recommended in the recently published guidelines [3] (http://www.british-thyroid-association.org/draft_thyca_23.12.06.pdf; http://www.oncoline.nl/uploaded/docs/Schildkliercarcinoom/schildklier%20ebro/richtlijn%20schildklier.pdf), but is still considered to be valuable in the follow-up of the high-risk patients. Measurement of the serum level of thyroglobulin (Tg), under recombinant human thyroid-stimulating hormone (rhTSH) stimulation or suppression, and ultrasonography have gained a more central role in monitoring for recurrent thyroid cancer [3, 4–7] (http://www.british-thyroid-association.org/draft_thyca_23.12.06.pdf). In patients with increasing or elevated Tg, a blind treatment with high-dose $^{131}$I can be applied, followed by a post-treatment $^{131}$I scan which also serves as a diagnostic tool [21]. However, with this strategy, unnecessary high radiation exposure and high TSH level must be taken into account especially in patients who subsequently have no $^{131}$I uptake on their post-treatment scan. Improvement of diagnostic imaging for the detection of recurrent or metastatic disease and a better (anatomical) localisation, e.g. using advanced imaging technique such as $^{124}$I-positron emission tomography (PET)/CT would allow more selective application of $^{131}$I therapy and might avoid unnecessary high-dose treatments.

Several iodine isotopes such as $^{123}$I, $^{125}$I, $^{131}$I play an important role in nuclear medicine, both for diagnostic purposes and for therapy. Iodine-124 is a positron emitting isotope and therefore suitable PET imaging. Its half-life is 4.2 days with a very complex decay scheme leading to extra non- or partially annihilation radiation coincidence detection [8, 9]. Approximately 23% of the disintegrations results in positron emissions.

While the radioisotopes $^{123}$I and especially $^{131}$I are used on a wide scale in diagnosis and treatment of all thyroid disorders, $^{124}$I has received little attention. This isotope would allow thyroid cancer imaging using the high-resolution PET technique [8, 10]. Iodine-124 has so far mainly been used for dosimetry or thyroid volume measurements [10–16]. However, the recent development of combined PET/CT scanners may increase clinical application in thyroid cancer patients, as detailed anatomical information is combined with the location of iodine positive tissue [17]. Iodine-124 has recently been applied for staging of differentiated thyroid cancer, as reported in a few case reports and small series, with promising results [17–19]. However, the diagnostic value of $^{124}$I-PET imaging as compared to (low- and high-dose) $^{131}$I scintigraphy has to be further investigated.

Iodine-124 PET may be able to detect recurrent or residual disease in differentiated thyroid carcinoma (DTC) with a higher sensitivity than the conventional (diagnostic) $^{131}$I scans because of the higher spatial resolution. In this way, $^{124}$I-PET imaging could possibly be a useful diagnostic tool during the follow-up of DTC patients. Therefore, we have performed a pilot study in the early treatment phase of thyroid cancer to get an impression of the diagnostic potential of $^{124}$I-PET for detection and staging of advanced DTC. We compared the $^{124}$I-PET imaging results with the diagnostic and post-treatment $^{131}$I-WBS.

**Materials and methods**

Twenty patients with histologically proven advanced DTC, including extrathyroidal tumour growth (T4), extra-nodal tumour growth or distant metastasis (M1), were examined in this study. These patients had undergone (near) total thyroidectomy. Four to 6 weeks after surgery, diagnostic $^{131}$I-WBS after 37 MBq of $^{131}$I was obtained, followed by an ablative dose of 5,550 MBq $^{131}$I. Post-treatment WBS was obtained after 10 days. At the time of ablation, serum Tg and Tg antibodies (TgAb) levels were also determined.

All 20 studied patients underwent diagnostic $^{131}$I-WBS, $^{124}$I-PET scan and post-treatment $^{131}$I-WBS 4 months after ablation therapy (Fig. 1). The Tgoff levels (under TSH stimulation after thyroid hormone withdrawal or after rhTSH injection) were also measured just before the administration of the diagnostic low-dose $^{131}$I. The Medical Ethics Committee of University Medical Center Groningen approved the study protocol, and all patients gave written informed consent.

Tg was measured by immunoradiometric assay (Brahms, Henningsdorf, Germany), with a functional sensitivity (i.e. the Tg concentration with a vital capacity of 20%) of 0.3 ng/ml as determined by the laboratory. Thyroglobulin antibodies (TgAb) were detected by both immunoradiometric assay (Brahms, Henningsdorf, Germany) with a cutoff of 60 U/ml and by immunoluminometric assay (Abbott, Hoofddorp, Netherlands) on the Architect i 2000 platform (Abbott) with a cutoff of 4.1 U/ml. Both cutoff concentrations were determined by Brahms and Abbott, respectively. In 19 patients, the Tg level was determined after thyroid hormone withdrawal and in one patient after rhTSH due to poor physical condition (Table 1).

**Tracer**

$^{124}$I-sodium iodide solution was obtained from Ritvec Isotope Products, St. Petersburg, Russia and was imported by I.D.B. Holland BV (Baarle-Nassau, The Netherlands). Radiochemical purity using instant thin layer chromatography and radionuclide purity using germanium (HP-Ge) detector were tested before release according to standard...
Radiopharmaceutical procedures. Before use, the solution was filtered using a Millipore bacterial filter (0.22 μm) and diluted with sterile saline. A sterility test was performed on each batch, but data were only available after administration due to the length of the test procedure (7 days).

PET scanning

Diagnostic 124I-PET imaging was performed 24 h after intravenous administration of 74 MBq of 124I [17]. A Siemens PET camera (Exact HR+, Knoxville, TN, USA) was used for imaging. The patient was positioned in the scanner, and a standard clinical whole-body PET study was performed in 2D ETTE mode over seven to eight positions.

Table 1 Patient characteristics and imaging results

| Pt, sex | TNM   | Tgoff (ng/ml) | TSH (mU/l) | 124I-PET | Pt 131I-WBS | Dx 131I-WBS | Validation |
|---------|-------|---------------|------------|----------|-------------|-------------|------------|
| 1 59M   | pT1N0M1 | 48            | 50         | CV, pelvis +++ | CV, pelvis +++ | CV, pelvis + | MR cv: p, MRI pelvis:p |
| 2 67F   | fT4N1M1 | 201           | 48         | Neck, thyroid bed +++ | Neck, thyroid bed +++ | Neck ++ | CT lung: no abnormal lesions, MRI neck: p |
| 3 85M   | fT4N0M1 | 2.6a          | 0.01a      | Skull, CV, pelvis +++ | CV+++ | CV + | Skull: p, MR cv: p |
| 4 48F   | pT4N1M0 | 49            | 77         | Neck ++ | Neck ++ | – | MRI:neck: p |
| 5 74F   | pT4N1M0 | <0.30         | 44         | Neck ++ | Neck ++ | – | MRI:neck: p |
| 6 74M   | fT2N0M1 | 691           | >200       | Neck/SC +++ | Neck/SC, pelvis +++ | – | MRI pelvis: p |
| 7 73F   | fT4N1M0 | 105           | 56         | Neck/thyroid bed ++ | Neck/thyroid bed ++ | – | US neck/FNAC: p |
| 8 18F   | fT4N1M1 | <0.30         | 51         | Pelvis ++ | Pelvis + | – | MRI pelvis:p |
| 9 39M   | pT4N1M0 | 85            | 75         | Neck +++ | Neck +++ | – | MRI neck: p |
| 10 73F  | pT4N1M0 | 3             | 43         | Neck + | – | – | US neck and FNAC: inconclusive, FDG PET: n |
| 11 59M  | fT3N0M1 | 1680          | 73         | Femur prosthetic ++ | – | – | MRI pelvis/femur: reactive tissue around the femur prosthesis; FDG PET: lesions in costae and pelvis |
| 12 73M  | fT3N0M1 | <0.30         | 33         | – | skull ++ | – | MRI skull: p, FDG PET: n |
| 13 67M  | pT4N0M0 | <0.30         | 142        | – | pelvis + | – | X-pelvis: n, bone scan: n |
| 14 42F  | fT4N1M0 | 15            | 81         | – | – | – | US neck and FNAC: n, FDG PET: n |
| 15 49F  | pT4N0M0 | 18            | 66         | – | – | – | US neck and FNAC: n, FDG PET: n |
| 16 59M  | pT4NxM0 | 2.8           | 76         | – | – | – | MRI neck, US neck and FNAC: n, FDG PET: n |
| 17 68M  | fT2N0M1 | 14            | 44         | – | – | – | US neck/FNAC: n, FDG PET and CT: lung lesions |
| 18 79F  | fT4NxM0 | 1.6           | 53         | – | – | – | MRI neck: p, US neck: lymph node not accessible for FNA, FDG PET: n |
| 19 71F  | fT4N0M0 | 0.54          | 36         | – | – | – | – |
| 20 37M  | pT1N1bM0 | <0.30         | 95         | – | – | – | – |

p Papillary, f follicular, Pt 131I-WBS post-treatment 131I-WBS, Dx 131I-WBS diagnostic 131I-WBS, CV cervical vertebrae, SC sternal clavicular, – not visible, + just visible, ++ visible, +++ clearly visible, n negative, p positive, FNAC fine needle aspiration cytology, FDG-PET fluorodeoxyglucose positron emission tomography

*a rhTSH stimulated

*b In these two patients, no additional radiologic imaging of the neck (no. 3) and pelvis (no. 6) was performed to confirm the findings on the PET and 131I-WBS due to poor physical condition and due to the lack curative therapeutic options
(from the upper thigh up until the top of the skull) of 5-min emission and 3-min transmission, standard energy window setting of 350–650 keV. Images were reconstructed with attenuation-weighted OSEM, two iterations and eight subsets. The total time needed for the scan was approximately 60 min.

In the initial four patients, the $^{124}$I-PET images were repeated 96 h after $^{124}$I administration (thus, 72 h after therapeutic dose of $^{131}$I) using the same acquisition settings as after 24 h. Narrowing of the energy window (425–650 or 460–562 keV, 3D mode) for $^{124}$I-PET imaging during high-dose $^{131}$I therapy to reduce the effects of $\gamma$-rays (364, 637 keV) of $^{131}$I and $^{124}$I (602 keV) and to improve image quality, as described in the phantom study (and clinical application in one thyroid cancer patient) by Lubberink et al. [20], was not possible in our institution due to technical reasons, e.g. camera characteristics. The 96-h $^{124}$I-PET images were very noisy and blurred and were excluded from the study. This poor image quality could partially be explained by the poor statistics due to the very low radioactivity left in the body after 96 h and partially by technical reasons as described above.

Data analysis

The $^{131}$I-WBS and $^{124}$I-PET scans were visually interpreted by two independent, experienced nuclear medicine physicians (HTP, PLJ). The findings on the $^{124}$I-PET scans were compared with the findings on the diagnostic and post-therapeutic $^{131}$I-WBS. Correlation with radiologic imaging (US, CT, MRI) and/or cytological (fine needle aspiration cytology, FNAC) investigation was done to confirm the findings or in case of discordant findings on the $^{124}$I-PET scan and $^{131}$I-WBS. If no abnormal uptake was seen on the PET scan and $^{131}$I-WBS, additional radiologic imaging (MRI, US), with or without FNAC, and fluorodeoxyglucose (FDG) PET were also performed to detect local or metastatic disease and/or used as a follow-up diagnostic tool.

Results

From December 2005 until April 2007, 20 consecutive patients with advanced DTC were included in this prospective study. The group consisted of ten women and ten men, median age of 67 years (range 18–85 years). Individual patient characteristics and the findings on the $^{124}$I-PET scan and $^{131}$I-WBS are summarised in Table 1.

Physiological uptake of $^{124}$I was observed in the salivary glands, esophagus, gastrointestinal tract and bladder as it is also normally seen on the $^{131}$I scans.

$^{124}$I-PET vs diagnostic $^{131}$I-WBS

Eleven patients (no. 1–11) showed uptake on the $^{124}$I-PET scan. In only three of them (no. 1–3) was the uptake also observed on the diagnostic $^{131}$I scan, but the uptake was clearer and the abnormalities were more extensive on the $^{124}$I-PET scan (Fig. 2). In nine patients (no. 12–20), no uptake was observed on both the scans. Results of additional radiologic imaging (MRI and/or US–FNAC) are also listed in Table 1.

It has also been noticed that the Tg off level (after endogenous TSH stimulation) was not detectable (<0.30 ng/ml, TSH >30 mU/L) without the presence of Tg antibodies in five (Table 1, no. 5, 8, 12, 13, 20) of the 20

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Fig. 2 This patient (no. 1) showed a clearly visible lesion in the cervical vertebrae on the $^{124}$I-PET (a, arrow), comparable with the lesion visible on the post-treatment $^{131}$I-WBS (c, arrow). This lesion was vaguely visible on the diagnostic $^{131}$I-WBS (b, arrow). Physiologic uptake in the esophagus, gastrointestinal tract and bladder is observed on the $^{124}$I-PET
patients. However, two of these five patients showed lesions on the ²¹²⁴I-PET (and post-treatment ¹³¹I scan) confirmed by MRI. Diagnostic ¹³¹I-WBS was negative in all five.

¹²⁴I-PET vs post-treatment ¹³¹I-WBS

Nine (no. 1–9) out of the 11 patients with uptake on the ¹²⁴I-PET scan had lesions which were also visible on the post-treatment scan (Fig. 3). In two patients (no. 10, 11), no uptake was observed on the post-treatment ¹³¹I scan and no anatomical localisation could be confirmed. FDG PET showed, however, lesions in the costae and pelvis in patient no. 11 (Fig. 4).

Two patients (no. 12, 13) showed uptake on the post-treatment ¹³¹I scan, which was not visible on the ¹²⁴I-PET scan: In one, MRI confirmed the ¹³¹I uptake in the skull, and in the other, no anatomical localisation could be found for the ¹³¹I uptake in the pelvis region. In seven patients (no. 14–20), no uptake was observed on both the scans. In four of these seven patients (no. 14–17), no lesions could be found with MRI and/or US–FNAC; in one of these four (no. 17), FDG-PET showed uptake in the lungs which was confirmed by CT. One patient (no. 18) showed slightly enlarged lymph node in the superior mediastinum on the MRI which was not accessible for US-guided FNAC. In two patients (no. 19, 20), MRI/US of the neck will be obtained (due to patient delay). Results of additional radiologic imaging are listed in Table 1.

Discussion

This pilot study showed that ¹²⁴I-PET detected more abnormalities in comparison to the diagnostic ¹³¹I-WBS, but showed comparable findings with the post-treatment ¹³¹I-WBS. Eleven patients had positive ¹²⁴I-PET scanning, and only three had visible abnormalities with the low-dose ¹³¹I scan. Moreover, ¹²⁴I-PET also showed abnormalities in two of the five patients with undetectable Tg, while the low-dose diagnostic ¹³¹I scan was negative in all five. These findings suggest that ¹²⁴I-PET better predicts the outcome of high-dose ¹³¹I treatment and would be better suited as a diagnostic tool to base clinical decisions on such as additional surgery or application of high dose ¹³¹I. Therefore, ¹²⁴I-PET should be performed before considering high-dose ¹³¹I treatment. A negative ¹²⁴I-PET could mean omitting ¹³¹I treatment, and further additional imaging should be performed to detect (non-iodine avid) metastatic disease.

Two patients (no. 12, 13) showed uptake in the skull and pelvis region, respectively, on the post-treatment ¹³¹I-WBS which was not visible on the ¹²⁴I-PET. Possible explanations for these findings might be the low dose of ¹²⁴I in which (small) lesions in the skull might be missed and false-positive uptake of ¹³¹I in the bowel which could be mistaken for abnormality.

Our results are in agreement with the study by Freudenberg et al. [17]. In their study, 12 patients with advanced DTC underwent high-dose ¹³¹I-WBS, ¹²⁴I-PET, CT, combined

Fig. 3 This patient (no. 6) showed a clearly visible lesion in the left lower neck region or on the ¹²⁴I-PET (a, arrow), comparable with the lesion visible on the post-treatment ¹³¹I-WBS (c, arrow). No uptake was observed on the diagnostic ¹³¹I-WBS (b)
124I-PET/CT, FDG-PET and US post-thyroidectomy during routine clinical staging. The overall lesion detectability for high-dose 131I-WBS was comparable with the 124I-PET, 83 vs 87%, respectively. Moreover, combined 124I-PET/CT modality, which showed an overall lesion detectability of 100%, resulted in a change of staging in two patients and a change in management in one.

It has been questioned why 124I-PET would be a suitable diagnostic tool when blind treatment is given anyway in the clinical practice. Blind treatment with high-dose 131I in patients known with negative low-dose diagnostic 131I scanning but with elevated Tg has been used as a diagnostic, therapeutic and prognostic tool. Patients without iodine accumulation on the post-treatment WBS, which could be an indication for tumour dedifferentiation, had a worse prognosis compared to those with a positive post-treatment WBS [21]. Repeating high doses of 131I may bear the risks in terms of long-term risk of secondary malignancies. In addition, repeated long-lasting TSH stimulation might have adverse effects on tumour growth. This aspect of long-lasting TSH stimulation may be prevented when performing the 124I-PET after rhTSH stimulation before the decision of giving additional high-dose 131I. However, comparative studies are needed to evaluate the yield of the 124I-PET under endogenous TSH stimulation and after rhTSH stimulation.
Additional advantages of $^{124}$I-PET would include the better resolution of this tomographic method and the ease of combining this with CT data to increase the diagnostic value, as shown by Freudenberg et al. [17]. Moreover, $^{124}$I-PET would allow more precise dosimetric calculations [25, 26, 28, 29], although that was not a focus of this study.

Another aspect which favours performing of $^{124}$I-PET before blind high-dose $^{131}$I is the radiation dose. The effective dose of $^{124}$I is in the same order of magnitude as $^{131}$I [10, 22–24]. The effective dose of $^{124}$I-iodide is 0.095 mSv/MBq with a thyroid uptake of 0% and increases to 1.5 mSv/MBq with a thyroid uptake of 35% [23]. The total radiation exposure has been calculated at 7.0 mSv for a dose of 74 MBq $^{124}$I. The administered dose is comparable to the dose administered for routine nuclear medicine scans. However, the total radiation exposure of $^{124}$I is just a fraction compared to the total radiation exposure of the therapeutic dose of $^{131}$I (340 mSv for 5,550 MBq [23], which was also mentioned in the study by Freudenberg et al. [17].

The issue whether stunning is a real phenomenon and its clinical relevance/consequence is debatable. Stunning effect after (higher) diagnostic dose of $^{131}$I has been described and discussed in the literature. The applied diagnostic $^{131}$I dose could impair the ability of the residual thyroid carcinoma tissue to accumulate the subsequently applied high-dose $^{131}$I dose. The degree of stunning probably depends on the absorbed radiation dose and the time between the diagnostic and therapeutic $^{131}$I dose.

Stunning was not seen with diagnostic $^{131}$I doses of 185 MBq (5 mCi) or lower [30, 31], whereas stunning was frequently observed after a diagnostic dose of 370 MBq (10 mCi) $^{131}$I [32]. There is also evidence that a short time interval between the administration of the diagnostic and therapeutic $^{131}$I dose may diminish the effect of stunning [31]. Most authors who have described stunning administered the therapeutic $^{131}$I dose several days after the completion of the diagnostic $^{131}$I scan [32–35]. Stunning is not seen when the therapeutic dose is administered within several hours. Furthermore, the time interval between the administration of a high dose of $^{131}$I and the performance of a post-therapy WBS may influence the observation of stunning [30, 36]. A longer time interval allows more time for soft tissue clearance of $^{131}$I, which results in a higher sensitivity of the post-therapy WBS. No stunning was seen at the post-therapy $^{131}$I-WBS, performed 5–10 days after doses of 1,110–3,700 MBq (30–100 mCi) $^{131}$I after 74 MBq (2 mCi) and 370 MBq (mCi) diagnostic scan [37].

In our study, we used a rather a low dose of $^{124}$I (74 MBq) and a short interval (1 day) between the administration of the diagnostic dose of $^{124}$I and the administration of the therapeutic $^{131}$I dose and a long interval (10 days) between the therapeutic $^{131}$I dose and the post-therapy $^{131}$I-WBS. If stunning does exist after $^{124}$I, it is therefore unlikely to have reduced the efficacy of $^{131}$I treatment in our study.

Iodine-124 is, however, poorly available with high costs, but the advantages of $^{124}$I-PET could outweigh these disadvantages and can lead to more clinical application in the follow-up of DTC patients when $^{124}$I will become more available.

Conclusions

In this study, $^{124}$I-PET proved to be a superior diagnostic tool as compared to low-dose diagnostic $^{131}$I scans and adequately predicted findings on subsequent high-dose post-treatment $^{131}$I scans. In combination with the high resolution and the possibilities to combine with CT, this could lead to improved clinical decision making.

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