Challenges and strategies on radioiodine treatment for differentiated thyroid carcinoma

Abstract
Objective: Radioiodine ($^{131}$I) is considered an effective and low-risk therapeutic radionuclide for differentiated thyroid carcinoma (DTC); however, dilemmas exist in the optimization of indications, pre-treatment thyroid stimulating hormone (TSH) stimulation, dose decision, as well as in the treatment of $^{131}$I-refractory disease. Refined strategies on $^{131}$I treatment for DTC based on late evidence and novel insights are greatly needed. Conclusion: The indications of $^{131}$I ablation continue to be refined with a better understanding of the risks and benefits. For pre-treatment TSH stimulation, recombinant human thyrotropin presents a better choice as it improves the quality of life, but is indicated only for ablation of the thyroid remnant and follow-up. Decreased doses of $^{131}$I seem to be more appropriate in patients without gross residual disease or metastases, but maximal doses are suggested in patients with advanced disease. Imaging procedures contributing to decision-making for patients with advanced DTC also continue to be modified. As for the $^{131}$I-refractory disease, there is a trend to increase $^{131}$I uptake and retention by using additional therapeutic agents like kinase inhibitors with encouraging results.

Introduction

For about 65 years, radioiodine ($^{131}$I) treatment is the standard treatment for differentiated thyroid carcinoma (DTC) combined with surgery and thyroid hormone replacement/inhibition [1]. However, challenges still exist in patient selection, pre-treatment thyroid stimulating hormone (TSH) stimulation, dose decision, as well as in the treatment for $^{131}$I-refractory DTC (RR-DTC) [2]. In this review, these dilemmas are presented and corresponding strategies are provided taking into account the best available medical evidence and novel insights into the management of DTC. Significant studies necessary to further address these issues are recommended accordingly.

Initial patient selection for radioiodine remnant ablation (RRA)

The aim of $^{131}$I administration post-surgery in DTC patients falls into three categories depending on risk stratification: remnant ablation, adjuvant therapy, and $^{131}$I therapy [3]. Figure 1 displays risk stratifications according to the American Thyroid Association (ATA) and the European Thyroid Association (ETA) guidelines. The goal of remnant ablation is to eliminate any normal thyroid tissue left after surgery, thereby simplifying disease surveillance by examinations, such as thyroglobulin (Tg) measurements and whole body $^{131}$I scans ($^{131}$I WBS) [3]. Adjuvant therapy theoretically destroys suspected microscopic neoplastic cells in the hope of improving disease-free survival [3]. $^{131}$I therapy is the administration of $^{131}$I in an attempt to destroy gross extrathyroidal extension and recurrent or metastatic disease. In this setting, patient selection is fairly simple [4].

Most DTC patients in whom $^{131}$I treatment is considered are in the remnant ablation setting, deserving optimization of indications. It is worth mentioning that in clinical practice remnant ablation, adjuvant therapy and $^{131}$I therapy can be simultaneously achie-
Radioiodine remnant ablation (RRA) is definitely recommended after total thyroidectomy for patients with large (>4cm) primary tumors or with gross extrathyroidal extension in order to improve disease-specific and disease-free survival [3]. Guidelines of ATA, ETA, and the National Comprehensive Cancer Network define the population of patients who do not require RRA, mainly those with small, (<1cm for papillary and <2cm for follicular or cell thyroid cancer), intrathyroidal tumors without other adverse features such as nodal or vascular invasion, high postsurgical thyroglobulin (Tg) levels or aggressive histology [3, 5, 6] (Table 1). However, there is still considerable controversy about the appropriate role of RRA in patients with primary tumors between 1 and 4cm in maximum diameter.

Rationales for RRA are as follows: 131I can destroy any possible occult residual microscopic thyroid carcinoma and thus reduce future disease recurrence. RRA facilitates long-term disease surveillance and detection of possible persistent or recurrent disease by the measurement of Tg concentrations as stimulated by thyroid-stimulating hormone (TSH) or by 131I WBS after RRA, or during the follow-up period [7].

Rationales against routinely administering RRA include: a) low risk of thyroid cancer associated mortality or disease recurrence [8], b) absence of consistent clear high quality evidence showing that such treatment reduces disease associated mortality or risk of recurrence [9], c) potential for short term and long term treatment associated side effects, d) availability of disease surveillance testing procedures that can be used irrespective of RRA status (such as diagnostic neck ultrasound with or without fine needle aspiration cytology, and unstimulated Tg measurements) [10]. Several following factors may contribute to decision making but their roles are not quite clear.

### Table 1. Comparison of indications on radioiodine remnant ablation for differentiated thyroid carcinoma among different guidelines

<table>
<thead>
<tr>
<th>Organization</th>
<th>Year</th>
<th>Indication</th>
</tr>
</thead>
<tbody>
<tr>
<td>American Thyroid Association Reference [3]</td>
<td>2015</td>
<td>Not recommended when diameter &lt;1cm (unifocal or multifocal, in absence of other high-risk features)</td>
</tr>
<tr>
<td>National Comprehensive Cancer Network Reference [5]</td>
<td>2014</td>
<td>Not recommended for small, &lt;1cm for papillary and &lt;2cm for follicular or Hürthle cell intrathyroidal tumors without evidence of nodal involvement or vascular invasion and with a low postsurgical thyroglobulin level</td>
</tr>
<tr>
<td>European Thyroid Association Reference [6]</td>
<td>2006</td>
<td>Not indicated when diameter is &lt;1cm and there was complete surgery, favourable histology, and no nodal or distant metastases or extrathyroidal tumor extension</td>
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**Serum Tg**

An area of controversy is about patients with elevated Tg without findings of macroscopic disease on the diagnostic 131I WBS. The discordance of Tg testing and imaging findings may be due to either false positive Tg or false negative 131I WBS. The former is usually caused by persistence of the thyroid remnant or by anti-Tg antibodies (TgAb) as tested by immunoradiometric assay. The later may be due to small volume of the lesion or to non iodine avid disease when 131I WBS is used [11].
According to ATA guidelines, for patients with relatively low (<5ng/mL) and stable serum Tg, conservative follow-up is generally recommended, but in the setting of higher or rapidly rising Tg, empiric treatment may be be indicated [3]. On the other hand, even some patients with high risk clinicopathologic features have undetectable Tg after thyroidectomy, so the role of pretreatment Tg on deciding whether to proceed with treatment is insufficient and more studies are needed to guide decision making in this area.

**Neck nodal involvement**

Another area of controversy relates to the importance of neck nodal involvement. In a single center retrospective study, investigators examined data from a subgroup of 421 patients, and reported an improvement of lymph node failure-free survival in patients with node-positive papillary thyroid cancer (PTC) treated with $^{131}$I. It is noteworthy that the greatest treatment benefits are observed in patients with lateral neck node metastases (N1b disease), as well as with lymph nodes >1cm in diameter [12]. Evidence from multivariable and propensity analyses from the National Thyroid Cancer Treatment Cooperative Study Group (NTCTCSG) showed no benefit of the adjuvant $^{131}$I treatment in Stage I subjects, including patients younger than 45 years with central neck node disease (N1a disease), but showed a beneficial effect in stage III patients, including patients younger than 45 years with N1b disease and patients with nodal metastases aged $\geq$45 years [13]. Thus, younger patients with N1a disease, especially when the size of node metastases is small, may not benefit from adjuvant therapy, though more convincing evidence is expected to withhold $^{131}$I treatment in such patients. Moreover, the 2015 ATA guidelines have classified patients with few small central neck nodal diseases as a low-risk group and do not recommend routine use of $^{131}$I in these patients [3]. However there are studies that demonstrate the effectiveness of RRA, even in low/intermediate risk patients with minimal node involvement [14].

**Molecular character of DTC**

Strong association has been found between the recurrence of PTC and genetic alterations [15-17]. Interestingly, coexisting BRAF V600E and TERT promoter C228T mutations were more often associated with high-risk clinicopathologic characteristics of PTC [16]. In view of the high negative predictive value of coexisting mutations, patients who harbor neither mutation and are conventionally low risk would be spared from RRA. The prognostic ability of BRAF and of other genetic biomarkers needs further investigation in order to make specific therapeutic suggestions.

**Pre-treatment TSH stimulation**

Thyroid hormone withdrawal (THW) is common practice for pre-treatment TSH stimulation in many countries. However, hypothyroidism may occur as an undesirable situation [18].

Liothyronine (L-T3) substitution is one of the alternatives intended to avoid hypothyroidism due to THW, but a randomized controlled study showed that L-T3 substitution did not prevent hypothyroidism, and delayed TSH elevation [19,20]. Recombinant human thyrotropin (rhTSH), approved by Food and Drug Administration in 2007 for use in pre-ablation settings, induced high TSH levels without clinical hypothyroidism. Whole-body radiation exposure was thus decreased due to more rapid clearance of $^{131}$I [21]. Multiple randomized controlled trials (RCTs) showed that exogenous rhTSH offered a much better short term quality of life in low risk DTC patients while the efficacy of RRA treatment remained sufficient [22]. Therefore, rhTSH is an acceptable alternative to THW in these patients for remnant ablation.

Studies comparing the use of rhTSH to THW are not sufficient to decide which of these procedures is better in higher-risk DTC patients. Exogenous stimulation was suggested by some investigators [23, 24] while Hugo et al. (2012) [23] showed in 586 patients that the 9 years clinical outcome of both procedures was almost the same with similar recurrence rates 1.5% for rhTSH vs. 1.2% for THW. Studies on metastatic DTC showed that the radiation absorbed dose by the tumors was 4-10 times lower under rhTSH as compared to THW, which is a determinant factor for treatment response [25, 26].

Since rhTSH administered by the standard method of two rhTSH injections is beyond the reach of many patients, some researchers suggested a combined protocol i.e. a shorter withdrawal period and a single dose of rhTSH [27]. Vrachimis et al. (2012) reported that the overall efficacy of the combined endogenous and exogenous TSH stimulation was not significantly different between the above two procedures [27], and thus the combined protocol may be an attractive alternative for RRA in DTC.

**Dosage of $^{131}$I and complication management**

The dose of $^{131}$I given to DTC patients ranges depending on the aim of $^{131}$I administration: remnant ablation, adjuvant therapy, or $^{131}$I therapy.

**Remnant ablation**

RRA is the only purpose for administering $^{131}$I in DTC patients without macroscopic or microscopic tumor focus post surgery. The main consideration is minimizing radiation toxicity while maintaining a reasonable uptake in the thyroid remnant. In order to reduce the dose of $^{131}$I in low risk patients in whom RRA is indicated, it is suggested to give much less $^{131}$I than previously thought (i.e. 1.1GBq rather than 3.7GBq) [28]. Equivalent ablation success rates using 1.1GBq or 3.7GBq was also confirmed by two separate, large, randomized studies. However, it should be noted that persistent disease was found in 3.7% of the patients studied and a second dose was needed in 9.5% of the patients, indicating that a more concise evaluation of the disease was needed before deciding about the amount of the administered dose [29, 30].
Adjuvant therapy
In high risk for recurrence patients, $^{131}$I is administered with the actual aim of both remnant ablation and adjuvant therapy although the optimum $^{131}$I dose is not exactly known. There is little evidence to suggest that increased dose activity of $^{131}$I is necessarily associated with improvement of clinical outcomes. A recent multivariable analysis using data from 1,171 DTC patients without distant metastases suggested that there was no significant difference in the risk of disease recurrence when using more or less than 2.78Bq of $^{131}$I post operatively (higher dose activity hazard ratio 1.57, 95% CI 0.61 to 3.98, $P=0.341$, with a mean follow-up period of 60 months) [31]. Thus, a lower dose with a potential of fewer adverse effects seems more appropriate.

In a recent study, among 319 subjects with an intrathyroid tumor and no metastases (T1-T2, N0, M0), BRAF V600E-mutated PTC was found to require a higher number of $^{131}$I courses to obtain disease-free status [32]. A possible explanation was that in the mutated cases, tumoral foci may be more frequently present in the postsurgical remnant because the BRAF V600E mutation correlates with multifocality. Similarly, DTC patients with TERT promoter mutated tumors were submitted to more $^{131}$I treatments with higher cumulative activities [33]. Therefore, it is conceivable to include a search of the BRAF mutation and TERT pro-moter mutations in the workup of DTC patients without other risk features, in order to distinguish those requiring more aggressive treatments from those that can be treated with a more conservative approach. RCT are also needed to explore the impact of other genetic factors on the efficacy of $^{131}$I in order to establish their role in guiding dose selection of post operative adjuvant therapy.

However, there is a biologic conundrum about the role of adjuvant therapy. Beta particles omitted by $^{131}$I have a range in tissues of the order of millimeters, and thousands of beta particles must traverse a cell to provide a high likelihood of lethal DNA damage. In microscopic foci of the disease, the radiation dose may be insufficient to kill tumor cells because a significant fraction will be deposited outside the cells. Therefore, a relatively low but effective remnant ablation dose may be less successful in achieving the adjuvant goal of killing residual microscopic cells of the disease [34]. On the other hand, there are studies suggesting reduced local recurrence with postoperative RRA, a benefit beyond remnant ablation as an adjuvant effect. To explain this effect, research in patients with a higher risk for local recurrence is needed, randomizing patients to low-dose $^{131}$I for RRA only versus higher doses for treating potentially small residual cancer deposits [34].

$^{131}$I therapy in locoregional or metastatic disease
There are usually three ways to choose the therapeutic dose of $^{131}$I in locoregional or metastatic thyroid cancer disease: a) To administer an empirically fixed dose, b) To administer a dose determined by the upper permissible limit of radiation to be absorbed in blood and the whole body and c) To administer a dose calculated according to tumor and/or lesions dosimetry. Published data suggest that although comparison of the outcome after administering the above doses of $^{131}$I is difficult, dosimetric approaches obviously have some advantages over empirical approaches [35].

Empirical doses are generally safe for the typical patient but are not based on $^{131}$I kinetics of a particular patient. Factors related to tumor burden, cardiac and renal function, and age can dramatically affect the maximal safe dose [36].

The dosimetric approach determines the maximum dose that can be safely given to prevent severe radiation toxicity. A $^{131}$I dose of 7.4MBq would exceed the maximum tolerated radiation absorbed dose, defined as 200cGy to the blood, in 8%-15% of patients younger than 70 years and in 22%-38% of patients aged 70 years and older [37]. Since whole-body and blood dosimetry determines the maximum $^{131}$I dose that can be safely given, they can maximize the probability of controlling the tumor and additionally prevent severe radiation toxicity.

In the treatment of DTC by $^{131}$I, several parameters vary for every patient and may be more important than the administered $^{131}$I activity. These parameters are: a) The tissue's avidity to $^{131}$I, b) The residence time of $^{131}$I per volume of blood or plasma and c) The effective half-life of $^{131}$I in the target volume. Lesional dosimetric approaches can overcome the uncertainty of individual bio kinetics, and thus are theoretically superior to empiric activity approaches. Lesional dosimetry, based on the absorbed dose of the target tissue, can prevent both suboptimal administration, which entails further $^{131}$I treatment, and an excessive administration of radioactivity, which may increase radiation toxicity. One retrospective study concluded that patients with loco-regional disease were more likely to respond to dosimetric treatment than to empiric treatment [35].

Numerous investigators have used $^{124}$I in PET or in PET/CT imaging to facilitate lesional dosimetry [38-40]. However, absorbed doses in a tumor may be inhomogeneous, which may lead to undertreatment of lesions not included in the dosimetric evaluation [40]. Future prospective studies are needed to determine formalisms, to weigh the biologically effective dose or the equivalent of a uniform $^{131}$I dose, study the three-dimensional radiobiological dosimetry and provide more reliable recommendations for lesional dosimetry in DTC.

Complications management
A minority of patients may have early-and late-onset complications from $^{131}$I treatment, such as taste dysfunction salivary dysfunction, xerophthalmia, obstruction to the nasolacrimal duct and pulmonary fibrosis [41-43]. For acute transient loss of taste or change in taste and sialadenitis, current measures to prevent damage to the salivary glands include hydration, sour candies, amifostine, and cholinergic agents [44]. However, observed effects of salagogues on radiation exposed salivary glands differ in various studies, so the exact role, so details of use of salagogues to prevent salivary gland damage remains uncertain [45, 46]. For chronic salivary gland complications, such as dry mouth, cholinergic agents may increase salivary flow [44]. As for nasolacrimal duct obstruction, definite preventive measures have not been reported. Perhaps, the application of artificial tears may serve to prevent ophthalmic sequelae in the same way that sour candies have been used to potentially alleviate sialoadenitis; endoscopic dacrycystori-
nontomy is an effective cure for this complication [42]. Finally, in patients with diffuse pulmonary metastases, pneumonitis or pulmonary fibrosis are associated with extremely high individual doses and repeated administration of $^{131}I$. Reduction in dose and courses of $^{131}I$ administration may minimize these complications [43].

Iodine-131 treatment for thyroid malignancy is also associated with a small risk of secondary malignancies in bones, breast, colorectal, kidneys, salivary glands, and leukemia; which seems to be associated with cumulative $^{131}I$ dose activity [47-49]. Methods of reducing abdominal pelvic viscera radiation dose, such as laxatives which decrease radiation exposure of the bowel and vigorous oral hydration which reduces exposure of the bladder and gonads, may help reduce the risk of secondary malignancies [47].

Imaging procedures contributing to decision-making in patients with advanced DTC

Tg testing and neck ultrasonography are routinely used for follow-up after initial treatment of DTC. However, once persistent/recurrent disease is detected by these tests, functional imaging or hybrid imaging should be proceeded in order to decide on subsequent treatment including $^{131}I$ therapy. For instance, for $^{131}I$ therapy, both the size of the lesions and their avidity for $^{131}I$ are important factors during decision-making. For example: persistent/recurrent disease with positive or negative $^{131}I$ WBS findings may need completely different treatment schedules (Figures 2 and 3). Large lesions, even if they are $^{131}I$-avid, are not suitable for $^{131}I$ therapy due to poor treatment efficacy. We should mention that $^{131}I$ WBS provides whole body information, which also contributes to the plan of $^{131}I$ therapy or other treatments. Radiiodine-whole body scan ($^{131}I$ WBS) used for follow-up after initial treatment detects $^{131}I$-avid diseases which may require subsequent $^{131}I$ therapy. However, several problems exist in the use of $^{131}I$ WBS: equivocal findings are frequent in the absence of anatomical imaging, for example, it is difficult to differentiate between uptake due to lesions in the lung and the ribs. Furthermore, even a routine dose of $^{131}I$ in diagnostic $^{131}I$WBS may cause stunning [50].

Incremental diagnostic value of $^{131}I$ PET/CT over $^{131}I$ WBS was examined in a retrospective study, in which $^{131}I$ PET/CT was reported to improve detection and localization of $^{131}I$ accumulation in lymph node metastases and distant metastases, and help change therapeutic planning in 3 (2.0%) of 147 patients [51]. An even greater additional value of $^{131}I$ PET/CT was reported by a study of ours, in which therapeutic strategy was changed in 8 of 17 patients whose planar images showed doubtful lesions. With additional information of the lesions derived from $^{131}I$ SPET/CT, six of the 8 patients were transferred to surgery, one patient was transferred to interventional treatment, and the other referred to external-beam radiotherapy [52]. It is worth mentioning that for patients with bone and brain metastases that cannot be adequately controlled with additional surgery for patients with $^{131}I$ refractory disease, the external beam radiotherapy should be considered [53]. This treatment helps to control pain, preserve or improve neurologic function, optimize local tumor control, and improve the quality of life [54]. In other studies carried out by colleagues of ours, hybrid imaging modalities were found to be more suitable for detecting more mediastinal lymph node metastases and for supporting treatment planning in DTC patients [55]. The $^{131}I$ PET/CT imaging is also of incremental value in detecting rare metastases from DTC [56].

The $^{124}I$ PET image allows foci of highly specific $^{124}I$ uptake to be localized with a low radiation dose, which is specifically important in pretherapeutic diagnostics. The sensitivity of $^{124}I$-PET for the detection of residual thyroid tissue and/or metastatic DTC was reported to be higher than that of a usual diagnostic $^{131}I$ WBS [57]. However, $^{124}I$ has not yet been widely available for routine clinical use and $^{124}I$ PET/CT has not yet been compared with $^{131}I$ SPET/CT in large

Figure 2. A 35 years old female in our department with papillary thyroid cancer and $^{131}I$-avid lymph nodes metastases after surgery who had complete response after $^{131}I$ treatment. Anterior: A) and posterior: B) therapeutic WBS 5 days after the administration of 555MBq $^{131}I$ showed two “hot spots” in the neck with a stimulated serum thyroglobulin level of 8.65ng/mL and normal TgAb. SPET/CT revealed thyroid remnant (not shown) and lymph nodes metastases. C) CT image; D) SPET image; E) SPET/CT fusion image. Six months later, a diagnostic $^{131}I$ scan showed no visible remaining disease accompanied by an undetectable stimulated serum Tg level of $<0.04$ng/mL and a normal TgAb.

Figure 3. A 32 years old male in our department with papillary thyroid cancer and non $^{131}I$-avid lymph nodes metastases after initial surgery who had a complete response after reoperation. Anterior: A) and posterior: B) therapeutic WBS 5 days after the administration of 555MBq $^{131}I$ showed a “hot spot” in the neck with a stimulated serum Tg of 15.93ng/mL and normal TgAb level. SPET/CT fusion image revealed only remnants in the left thyroid lobe: C) and the pyramidal lobe: D). However, calcified lymph nodes (arrow) were detected by ultrasonography: E) and plane CT: F) and G) indicating possible metastases. Pathologic examination of resected lymph nodes verified lymph node metastases from papillary thyroid cancer. Two months later, another ultrasonography: H) and enhanced CT: I) and J) were negative with an undetectable stimulated serum Tg level <0.04ng/mL and a normal TgAb.
society of patients with DTC. Moreover, problems or pitfalls exist such as the “shine-through” effect, which hamper the interpretation of $^{131}$I PET/CT imaging.

The ability of molecular imaging modalities mentioned above to detect persistent or recurrent DTC disease relies on the avidity of different radiotracers to lesions. As for non $^{131}$I avid DTC lesions, the fluorine-18-fluorodeoxyglucose ($^{18}$F-FDG) PET imaging is a better diagnostic modality (than $^{131}$I WBS) [58]. The incremental value of $^{18}$F-FDG PET/CT over $^{131}$I WBS in patients with extensive or metastatic high-risk DTC has been confirmed in a study by Rosenbaum-Krumme et al. (2012) [59]. Furthermore, high $^{18}$F-FDG uptake on PET imaging in metastatic DTC is a reliable predictive factor for RR-DTC, which we will discuss in detail in the following part of this review [60].

As to $^{18}$F-FDG PET/MRI, its diagnostic potential was evaluated very recently in comparison to PET/CT in patients with DTC suspected or known to have dedifferentiation, and it was found that PET/MRI was equal to contrast-enhanced neck PET/CT for the assessment of cervical lesions but was inferior to low-dose PET/CT for the assessment of pulmonary lesions [61].

Means of increasing $^{131}$I uptake and retention in $^{131}$I-refractory differentiated thyroid carcinoma

In patients with advanced disease RR-DTC is defined by the following outcomes: a) The presence of at least one tumor focus without any uptake of $^{131}$I, b) Structural progression of the disease shortly after a $^{131}$I treatment course (12 or 16 month later), c) No signs of remission after the administration of a cumulative activity of 22GBq or more. Although RR-DTC is rare, with an estimated incidence of four cases per million population or 5% of patients with clinical cancer, it is usually aggressive, with a median survival after distant metastases, ranging from 3 to 6 years [62]. The poor survival and the absence of effective treatment make the clinical management of RR-DTC extremely difficult [63].

Given the positive outcome from $^{131}$I treatment in patients with $^{131}$I-avid disease, enthusiastic physicians suggested to adopt additional therapeutic agents such as retinoids, epigenetic drugs and, kinase inhibitors and lithium in order to increase $^{131}$I uptake and retention in DTC tumor cells in $^{131}$I-refractory disease. It is also important to notice that RR-DTC can be asymptomatic and stable for a period of time and in such cases the benefits of novel therapies (kinase inhibitors therapy etc.) might be largely outweighed by drug toxicities [64,65].

The main selection criterion in clinical trials in patients with RR-DTC is documented progression of target lesions within less than 1.5 years after $^{131}$I therapeutic dose according to Response Evaluation Criteria In Solid Tumors. Patients with a large tumor burden might require systemic treatment, including cytotoxic chemotherapy or targeted therapy, before assessment of progression. The decision to initiate treatment may be based on high $^{18}$F-FDG uptake of the lesions on PET scan, on histological characteristics of the tumor and on Tg doubling time less than one year. These are signs predicting RR-DTC during the follow-up period [66].

In progression-free patients with few or asymptomatic metastatic foci, local treatment techniques, such as radiofrequency, external radiation treatment and surgery can be used and serum TSH should be maintained at a low or undetectable concentration [39, 67]. Figure 4 is a flowchart which outlines the current clinical management strategies for advanced DTC in cases of local recurrence/persistence and metastases.

Targeting signaling pathways

In early studies of DTC, multiple kinase signaling pathways (PI3K/Akt pathway, cAMP pathway, MAPK pathway etc.) were found activated due to genetic alterations, like RET/PTC rearrangement, mutations in RAS, BRAF genes etc. or due to other factors; which lead to low natrium iodine symporter (NIS) expression and/or to impaired targeting of NIS to plasma membrane, and ultimately lack of $^{131}$I uptake of the lesions [68, 69]. The above findings suggest that inhibiting the over-activated pathways may be an effective way to reconstitute NIS function.

Preclinical investigations and clinical trials exhibit target agents ability to induce $^{131}$I uptake. In-vitro studies [70, 71] showed that targeting of BRAF, MEK, or AKT in the MAPK and PI3K-AKT pathways could restore thyroid gene expression and $^{131}$I uptake, which was enhanced by TSH in thyroid cancer (TC) cells. These findings are supported by a later in vivo study in which murine TC cells with inducible BRAF V600E (TC) cells. These findings are supported by a later in vivo study in which murine TC cells with inducible BRAF V600E expression were re-sensitized to $^{131}$I treatment by using MEK inhibitor PD0325901 and the selective BRAF V600E /Raf inhibitor PLX4720 [72]. Similarly, in our in vitro study [73], enhancement of $^{131}$I uptake and decrease of glucose metabolism were simultaneously observed in the BHP-2-7 cells after treatment with either sorafenib or carbozantinib, and so the iodide-handling gene expression was restored and glucose transporter 1 and 3 expression was inhibited. Corresponding in vivo study is now ongoing. In a recent clini-
cal study, by combining selumetinib with $^{131}$I treatment in advanced DTC, an increase in $^{131}$I uptake was obtained in 12 of 20 evaluated patients [74]. Only 8/12 patients whose $^{131}$I PET/CT scans showed that DTC lesions were exposed to at least 20Gy were selected to continue $^{131}$I treatment and all had imaging evidence of response. More details will be revealed in an ongoing phase III study (NCT01843062) using selumetinib in combination with adjuvant radiodine in high risk DTC patients after thyroidectomy. High induction of $^{131}$I uptake (60%) was also obtained in patients with metastatic BRAF-mutant, and $^{131}$I-refractory PTC treated with dabrafenib, an inhibitor of mutated BRAF. Two of these 10 enrolled patients had partial response and 4 patients had stable disease on standard radiographic restaging at 3 months [75]. Another two clinical trials (NCT01534897, NCT02145143) aiming to increase $^{131}$I uptake and retention in RR-DTC patients using BRAF inhibitors are ongoing [76].

Although there is limited evidence of targeting overactive signaling pathways to induce $^{131}$I uptake and contribute to increased efficacy of $^{131}$I treatment in RR-DTC, combination treatment of target agents and $^{131}$I is of great promise. Larger studies are necessary for confirming the magnitude and reproducibility of re-inducing $^{131}$I uptake and longer observation time is needed to monitor the indolent DTC and to see if prolonged survival time can be obtained. In such studies, it should be clarified whether improved efficacy is mainly due to the antitumor effect of target agents or to improved $^{131}$I uptake induced by the action of target agents. Optimization of therapeutic strategies based on genetic alterations remains to be further explored.

**Acting on nuclear receptors**

Retinoids (retinol and its derivatives) were reported to increase both expression of thyroid specific proteins and $^{131}$I uptake in TC cells through acting on the nuclear receptors retinoic acid receptor (RAR) and retinoid X receptor (RXR) (mainly by activating RAR-α), whereas they decrease $^{131}$I uptake in normal thyrocytes [77]. However, their clinical efficacy is limited. Isotretinoin (13-cis-RA) re-induced $^{131}$I uptake in few patients with advanced DTC [78, 79]. Other studies reported inconsistent increase in $^{131}$I uptake [80, 81] using tretinoin (all-trans RA). In a recent publication, Damle et al. (2015) showed that retinoic acid increased $^{131}$I uptake in 55% of subjects but the magnitude of increase was very small [81]. Another less well-studied first generation retinoid, retinol, showed enhanced $^{131}$I uptake in DTC cell lines but had a low anti-proliferative effect [82].

Peroxisome proliferator-activated receptor (PPAR) agonists, such as rosiglitazone, bind to PPAR-γ which is a transcription factor of the superfamily of nuclear receptors showed a decreased expression in DTC. These agonists form a heterodimer with the RXR receptor at the response elements, activating the transcription of target gene PTEN, which in turn inhibits PI3K pathway [83]. Evidence of rosiglitazone’s ability to induce $^{131}$I uptake has been accumulated in the previous decade. There are case reports describing successful induction of $^{131}$I uptake after treatment with rosiglitazone in patients with non-$^{131}$I avid metastases of DTC [84, 85]. Furthermore, in trials including more patients, an increase in $^{131}$I uptake was observed in 40% and 26% of the patients after >6 weeks of pretreatment with up to 8mg of rosiglitazone [86, 87].

**Directing against epigenetic alterations**

Both methylation and deacetylation have been reported as mechanisms of dedifferentiation of cancer cells. Therefore, compounds that can reverse methylation or inhibit histone deacetylation may lead to the re-expression of silenced thyroid-specific genes.

In a study on mRNA expression of thyroid-specific genes (NIS, Tg, TPO and TSHR) and on $^{131}$I uptake in the PTC cell line B-CPAP, various gene modulators were examined, however, the DNA methyltransferase inhibitor (DMI) 5-azacytidine was the only compound which increased $^{131}$I uptake [88]. Another DMI, 5-aza-20-deoxycytidine, which was showed to increase expression of thyroid transcription factor 1 in follicular TC, PTC and anaplastic TC cell lines [89], is under evaluation in an ongoing phase II study (NCT00085293).

A new hydroxamic acid-derived histone deacetylase inhibitor panobinostat was reported to cause re-expression of NIS and to increase the cytotoxic effect of $^{131}$I treatment [90]. Romidepsin, another novel compound, was tested in a phase II trial, and was reported to restore $^{131}$I avidity in 2 of the 20 treated patients, but no objective response was showed after $^{131}$I treatment [91].

**Lithium and other agents**

Lithium causes an increased cell retention of iodide due to inhibition of its efflux [92], and short-term treatment with LiCO₃ as adjunct to $^{131}$I improves the efficacy of $^{131}$I in hyperthyroidic patients [93]. Recent presentation by Ilhan Lim on the 84th annual meeting of ATA showed that a response rate of 45% was obtained in metastatic RR-DTC after combined treatment of $^{131}$I and lithium for 6 months, and a slightly higher 10 years survival rate was achieved compared to patients with only $^{131}$I treatment (67.9% vs. 66.8%, P<0.05). Other compounds, such as genistein, alpha-liponic acid, arsenic trioxide and metformin have also been reported to increase $^{131}$I uptake to different extents in vitro and in vivo [94-98].

In conclusion, although $^{131}$I has become a prototypical theranostic agent for DTC, challenges still exist in the $^{131}$I treatment of DTC. Based on recent clinical experience, $^{131}$I can be more appropriately administered via better patient selection, further modified pre-treatment TSH stimulation and more precise dose decision. During long-term management, although serum Tg measurements and neck ultrasonography are routinely used with excellent diagnostic efficacy, functional/hybrid imaging modalities including WBS, SPET/CT or PET/CT using various radiotracers provide incremental information for decision-making of therapeutic strategies. As to management of RR-DTC, clinical data on various redifferentiation compounds failed to meet the initial high expectations, while novel target agents have displayed their ability to control the disease as well as to increase $^{131}$I uptake and retention in cases of RR-DTC, indicating that combined treatment with target agents and $^{131}$I has become one of the most important issues and deserves further investigation.
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The authors declare that they have no conflicts of interest.

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