Iodine, Thyroid Autoimmunity and Cancer

Emilio Fiore    Francesco Latrofa    Paolo Vitti
Department of Experimental and Clinical Medicine, Endocrinology Unit 1, University Hospital of Pisa, Pisa, Italy

Key Words
Iodine · Hashimoto’s thyroiditis · Thyroid autoimmunity · Papillary thyroid cancer

Abstract
This review focuses on two different topics: (a) iodine and autoimmune thyroid disease (AITD) and (b) AITD and papillary thyroid carcinoma (PTC). Iodine intake modifies the expression of thyroid diseases and has been associated with induction of AITD. Thyroglobulin (Tg) is an important target in iodine-induced autoimmune response due to post-translational modifications of iodinated Tg, as suggested in animal models. We have shown that the unmasking of a cryptic epitope on Tg contributes to iodine-induced thyroid autoimmunity in humans. The relationship between AITD and PTC has been suggested in many studies. The presence of two different mechanisms has been hypothesized, one typical of AITD and the other of an immune reaction to PTC. We have shown that in AITD, the pattern of Tg recognition by anti-Tg antibodies (TgAb) is ‘restricted’ to the immunodominant regions of Tg, while in patients with non-AITD, such as nodular goiter and PTC devoid of thyroid lymphocytic infiltration at histology, TgAb show a less restricted epitopic pattern and bind also to other regions of Tg. Thyroid function may also affect the frequency of PTC, the risk of cancer increasing with serum TSH levels. We have shown that this mechanism, rather than thyroiditis per se, plays a major role in the association of PTC with Hashimoto’s thyroiditis, as a consequence of the autoimmune process leading to a progressive increase of serum TSH in these patients.

Introduction
This review will focus on two different, related topics: (a) iodine and thyroid autoimmunity and (b) thyroid autoimmunity and papillary thyroid carcinoma (PTC). We will review the latest data on these issues and describe our more recent contributions to these fields.

Iodine and Thyroid Disease

Iodine Nutrition and the Spectrum of Thyroid Diseases
The iodine dietary intake is insufficient in over 2 billion of people and, therefore, deficiency of iodine is one of the main issues of public health worldwide [1]. The term iodine deficiency disorders (IDD) refers to the several consequences that iodine deficiency imposes on individuals, including goiter, which is the most common clinical manifestation, and cerebral impairment ranging from mild cognitive defects to cretinism. The frequency
and severity of IDD are proportional to the magnitude of iodine deficiency [2] and can be prevented by improving iodine nutrition through salt iodization [3].

Only few countries (Switzerland, some Scandinavian countries, Australia, the USA and Canada) were completely iodine sufficient before 1990. Since then, iodized salt (IS) programs have been implemented in many countries worldwide, and two thirds of the world’s population (71%) are nowadays estimated to be covered by IS [2].

Iodine supplementation affects the epidemiology and eventually the natural history of thyroid diseases. In old Chinese medical writings of 3600 B.C., a decrease in goiter size was reported upon ingestion of seaweed and overcooked sea sponge [4]. A large number of studies have shown the effect of iodine supplementation on the epidemiology of goiter [5–7]. In two surveys conducted in Pescopagano [8, 9], an Italian village with a previous moderate iodine deficiency, before and after the introduction of a iodine prophylaxis program on a voluntary basis, a significant reduction of the prevalence of goiter was observed, mainly due to the reduction of diffuse goiter in 2010 compared to the previous survey of 1995 (10.3 vs. 34.0%). The overall prevalence of nodular goiter was not significantly different in the two surveys, but when age was taken into account, its frequency was significantly lower in 2010 than in 1995 in subjects aged 26–35 years (3.8 vs. 11.3%), while no difference was observed in subjects >35 years old. The absence of a significant difference in the frequency of nodular goiter in subjects older than 35 years may be explained by the fact that in the majority of the subjects of this class of age the process of goiter development was initiated already before the increase of iodine intake.

After the introduction of iodine prophylaxis, an increased incidence of hyperthyroidism was observed [10]. This is due to the development of iodine-induced thyrotoxicosis in subjects with preexisting autonomous multinodular goiter. This condition is transient and correlated to the level of iodine deficiency and the amount of iodine administered. In 1980, when in Switzerland the salt iodine content was increased from 7.5 to 15 mg/kg, doubling iodine intake, the incidence of toxic nodular goiter increased by 12% in the first 2 years, but then declined to only 25% of the initial incidence [11]. In 2013, the WHO indicated that 10 countries have iodine intakes above 300 μg/l, the threshold of ‘excessive’ iodine intake [1]. These data emphasize the importance of regular monitoring of iodine status to detect both low and excessive intakes of iodine. In the survey conducted in Pescopagano, after the introduction of an iodine prophylaxis program on a voluntary basis [9], thyroid autonomy was not observed in younger subjects and was associated with a significant reduction in the frequency of hyperthyroidism due to toxic nodular goiter in older age. We did not observe the increased frequency of hyperthyroidism after the beginning of iodine prophylaxis that has been reported in other studies [12, 13], conceivably because of the lower level of iodine supplementation in this population. Taken as a whole, these data indicate that iodine prophylaxis is associated with a reduced frequency of thyroid autonomy in younger subjects and of toxic nodular goiter in older patients. This phenomenon may have a relevant impact on the health of this population, thyrotoxicosis being associated with arrhythmias and mortality in older people.

Iodine Nutrition and Thyroid Autoimmunity

A potential side effect of iodine supplementation is the induction or the worsening of autoimmune thyroiditis [14, 15]. Cross-sectional studies of populations with different iodine intakes in Great Britain [16], in Denmark and in Iceland [17] showed that the frequency of thyroid autoantibodies (TAb) and hypothyroidism is higher in iodine-replete than in iodine-deficient populations. Boukis et al. [18] showed that in goitrous patients from a mildly iodine-deficient area in Greece, TAb became positive in 42.8% of the subjects 3 and 6 months after treatment with iodized oil. Recently, Pedersen et al. [19] found an increased prevalence of TAb after the beginning of a cautious iodization program, supporting the view that even a small increase in iodine supplementation may be associated with an increased thyroid autoimmunity. An increased frequency of hypothyroidism has also been reported after the careful introduction of IS in Denmark [14]. Similar findings were reported in the Pescopagano survey, where an increased frequency of TAb and of Hashimoto’s thyroiditis (HT) was observed after the beginning of iodine prophylaxis [20]. This study showed that iodine intake modulates the pattern of thyroid diseases, even in case of slight differences in intake and doses below 150 μg daily recommended for preventing IDD. The role of iodine in inducing thyroid autoimmunity is also strongly supported by animal models. Excessive iodine intake can precipitate spontaneous thyroiditis in genetically predisposed animals by increasing the immunogenicity of thyroglobulin (Tg) [21]. This may be explained by the fact that Tg is the only self-antigen that undergoes post-translational modification as a consequence of the environmental supply of iodine, with the exposure of previously hidden epitopes. In experimental animals, enhanced iodination of Tg facilitates processing and presentation of a cryptic pathogenic peptide [22–25]. In agreement with this concept is the observation that io-

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dine may play a role in central tolerance, and T cells undergoing thymic selection likely recognize only noniodinated Tg epitopes [21]. Furthermore, in experimental models of autoimmune thyroiditis, it has been shown that breaking B-cell self-tolerance occurs first for Tg and subsequently for thyroperoxidase [26].

It is not known how Tg modifications induced by posttranslational iodination are involved in the development of spontaneous thyroid autoimmunity in humans. We have approached this matter in the population of Pesco-pagano. In a second survey conducted in this village in 2010 [20], 70% of subjects declared to routinely use IS (IS users) and 30% never to use IS (IS non-users). Accordingly, median urinary iodine excretion was slightly higher in IS users than in non-users (Mann-Whitney U test: p < 0.0001).

Table 1. Prevalence of TAb according to use of IS

<table>
<thead>
<tr>
<th></th>
<th>Users (n = 906)</th>
<th>Non-users (n = 389)</th>
<th>p</th>
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<tbody>
<tr>
<td>TgAb</td>
<td>171 (18.9%)</td>
<td>53 (13.6%)</td>
<td>0.02</td>
</tr>
<tr>
<td>TPOAb</td>
<td>153 (16.9%)</td>
<td>51 (13.1%)</td>
<td>n.s.</td>
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n.s. = Not significant.

Thyroid Autoimmunity and PTC

HT and PTC

The relationship between thyroid autoimmunity and PTC has been suggested in many studies, but its meaning is still debated. The association between these two disorders is appealing, and it has been hypothesized that in HT, the inflammatory response may create a favorable setting for malignant transformation as well established for other cancers [35].

The possible association between HT and PTC was first proposed more than 50 years ago by Dailey et al. [36] who found a high frequency of thyroid carcinomas in patients with HT who underwent thyroidectomy. Hirabayashi and Lindsay [37] examined 9,287 thyroid specimens and found that papillary carcinoma was present in 22.5% of the glands with HT and only in 2.4% of those with no pathological signs of HT. Okayasu et al. [38] and Okayasu [39] found that the prevalence of lymphocytic infiltrates, suggestive of autoimmune thyroiditis, was significantly higher in patients from IS users and IS non-users. We have previously characterized 4 human recombinant monoclonal TgAb-Fab (A, B, C and D) [27, 28]. In agreement with the well-known restriction of TgAb epitope pattern [29, 30], the 4 TgAb-Fab recognize 4 different but partially overlapping epitopes, A epitope lying at one pole, B epitope at the other pole and epitopes C and D being on the fringe of the immunodominant region of Tg [28]. By inhibiting TgAb binding to Tg with human recombinant monoclonal TgAb-Fab (fig. 2), we showed that region A was the major component of the immunodominant region on Tg recognized by TgAb from autoimmune thyroid disease (AITD) and non-AITD patients, while epitopic region B was recognized to the greatest extent by HT patients [28, 31, 32].

When we compared sera from IS users and IS non-users, we observed a similar pattern of inhibition of Tg binding by TgAb-Fab A, C and D, while inhibition by TgAb-Fab B was higher in IS users than in IS non-users (27.5 vs. 3.0%; p = 0.04). These results suggest that iodine contributes to autoimmunity to Tg by unmasking a cryptic epitope on Tg (fig. 3), showing for the first time in humans the results reported in animal models. In addition, the observation that neither age nor genetic background influence TgAb epitope pattern in HT [33], and that the rise in TgAb levels observed after 131I treatment for Graves’ hyperthyroidism [34] is not associated with an epitope spreading of TgAb, underlines that in AITD, TgAb epitope pattern, once established, remains restricted.

Fig. 1. Box-whisker plot of urinary iodine excretion (UIE) measured in subjects not using (white box) or using (gray box) IS. Results are reported as median values (black lines), interquartile (25th to 75th percentiles) ranges (boxes) and 10th to 90th percentiles (whiskers). UIE was significantly higher in IS users than in non-users (Mann-Whitney U test: p < 0.0001).
with PTC than in those with adenomatous goiter or follicular adenoma. Ott et al. [40, 41] reported a high incidence of thyroid carcinoma (32%) in 146 patients with HT and solitary cold nodules. However, other studies yielded conflicting results, reporting a low incidence of thyroid carcinoma in patients with HT [42–44]. Boi et al. [45] found a significant association between HT and PTC in a cytological series and Jankovic et al. [46] reported a significant relationship between HT and PTC in histological, but less consistently in cytological series. Indeed, the average prevalence of PTC in patients with HT was 27.56 and 1.20% in histological versus cytological series, respectively. The authors underline that a selection bias may account for the higher prevalence of PTC in HT in histological series, because in the vast majority of these patients, the indication for surgery is the presence of nodules suspicious for malignancy, while it is more commonly the size of goiter or the presence of compressive symptoms in patients with nodular goiter. These findings were confirmed by Castagna et al. [47] who reported that the prevalence of malignant cytology in patients with nodular HT diagnosed on clinical grounds was similar (4.5%) to that found in nodular goiter with or without TAb and in nodular Graves’ disease (GD). When the same analysis was performed in the subgroup of patients submitted to thyroid surgery, the prevalence of PTC was significantly higher in patients with nodular HT (67.8%) compared with the other groups (36.9, 37.2 and 40%, respectively).

The heterogeneity of criteria used to define HT may hamper the drawing of conclusions on this issue. By strict criteria, HT is a histological diagnosis characterized by a widespread lymphocytic infiltration of the thyroid. On the other hand, PTC is often associated with a significant lymphocytic infiltration [38] that may represent a response to the tumor. On clinical grounds, the diagnosis of HT is based on the presence of serum TAb and of spontaneous hypothyroidism that may be present at the initial evaluation or may develop during follow-up [15, 16, 48]. However, TAb may also be detected in euthyroid subjects in the absence of a diffuse lymphocytic infiltration of the thyroid, especially in nodular goiter. Thyroid ultrasound has been proposed as a tool for the clinical diagnosis of
HT, even if this point is not universally accepted. The typical hypoechoic pattern at ultrasound is related to the structural changes caused by the lymphocytic infiltration of the thyroid gland and has been associated with the onset of hypothyroidism [49, 50].

We addressed the issue of the relationship between thyroid autoimmunity and PTC in a large series of patients submitted to fine-needle aspiration of thyroid nodules and to thyroid surgery [51]. We showed that the prevalence of TAb was similar in patients with benign nodules as compared to patients with malignant nodules at cytology (38.7 vs. 35.6%, respectively). However, in women with multinodular goiter, TAb were more common in presence of benign nodules than in PTC (40.2 vs. 32.5%, respectively; p = 0.02), while in men with single/isolated thyroid nodules, TAb were more common in PTC than in nodular goiter (31.2 vs. 20.4%, respectively; p = 0.02). At variance with these clinical observations, a moderate to severe lymphocytic infiltration at histology was more common in PTC than in benign nodular thyroid disease, both in patients with multinodular goiter (37.8 vs. 9.2%, respectively; p < 0.01) and in those with single thyroid nodules (58.7 vs. 30.1%, respectively; p < 0.01). Interestingly, in subjects with moderate/severe thyroid lymphocytic infiltration, the frequency of positive serum TAb was significantly higher in patients with benign diseases than in those with PTC (78.2 vs. 49.7%, respectively; p = 0.0002), indicating that several patients with PTC had undetectable serum TAb despite the moderate/severe thyroid lymphocytic infiltration observed at histology. Furthermore, in a group of patients treated with total thyroidectomy for PTC [52], thyroiditis at histology was detected in 40.7% of patients, while a clinical diagnosis of HT or GD was performed in half of these patients (20.3%), supporting the observation that in PTC, lymphocytic thyroiditis on histology is more common than clinically diagnosedAITD. Based on these data, (1) we confirmed that PTC was not predicted by positive serum TAb; (2) we concluded that PTC was associated with TAb only in the subgroups of patients with a lower prevalence of thyroid autoimmunity (i.e. men vs. women); (3) we showed that a moderate to severe lymphocytic thyroiditis was much more common in PTC than in nodular goiter and (4) we demonstrated that the correlation between histological thyroiditis and serum TAb was lower in PTC than in benign nodules.

**Thyroid-Stimulating Hormone, Thyroid Autoimmunity and PTC**

Thyroid function may also affect the frequency of clinically detectable PTC [53]. In the last few years, it has been reported that in patients with nodular thyroid diseases, the risk of thyroid malignancy increases with increasing concentrations of serum thyroid-stimulating hormone (TSH), and even within normal ranges, higher TSH values are associated with a higher frequency and more advanced stage of thyroid cancer [54–61]. In HT patients, the progressive reduction of thyroid function as a consequence of the autoimmune process leads to the progressive increase in serum TSH. Therefore, it is possible to hypothesize that increased TSH levels may also play a role in the development of PTC in patients with nodular HT. To verify this hypothesis [62], we have evaluated the relationship between TSH and PTC in 9,824 untreated patients with nodular HT (n = 893) or nodular goiter (n = 8,931). Patients were diagnosed as affected by nodular HT if they had a high titer of TAb, i.e. had both TgAb and TPOAb higher than 100 IU/ml, or a low level of TAb, but had hypothyroidism or a clear hypoechoic ‘thyroiditis’ pattern at thyroid ultrasound. Patients with nodular goiter had no hypoechoic pattern at thyroid ultrasound, were euthyroid and had either undetectable serum TAb (TAb– nodular goiter; n = 6,571) or low-level serum TAb (TAb+ nodular goiter; n = 2,360). The frequency of PTC in patients with nodular HT (9.4%) was significantly higher compared to patients with TAb– nodular goiter (6.4%) and TAb+ nodular goiter (6.5%). Altogether, these data suggest that the high frequency of PTC in patients with nodular HT is mainly due to the higher TSH levels compared to patients with nonnodular goiter. The results of the statistical analysis were confirmed in a second group of patients with nodular HT (n = 638) or nodular goiter (n = 3,276) under treatment with L-thyroxine (LT4) [62]. This therapy was inhomogeneous in the patients included in the study, the purpose of LT4 treatment being to reduce TSH levels to lower values of a normal range in patients with nodular goiter and to correct hypothyroidism, bringing serum TSH within the range of normal values in more than half of the patients with nodules in HT. As a result of this uneven treatment, higher levels of TSH and a higher frequency of PTC were observed in nodular HT with respect to nodular goiter in the whole series of LT4-treated patients. However, when only patients with serum TSH levels below the median value (0.90 mIU/l) were taken into account, no significant difference in PTC frequency was found, suggesting the hypothesis that TSH is the main factor associated with the risk of PTC independently of the diagnosis of nodular HT or goiter (fig. 4).

As thyroid hypoechogenicity at ultrasound is not a widely accepted criterion for the diagnosis of HT, we analyzed our data taking into account only TAb level and hy-
In untreated patients, the frequency of PTC was significantly higher (9.3%) in those with hypothyroidism and/or high serum levels of TAb compared to patients with both low serum levels of TAb (6.8%) and negative serum TAb (6.4%; fig. 5a). Patients with hypothyroidism and/or high serum levels of TAb also had higher TSH levels compared to patients with lower serum levels of TAb or negative serum TAb (fig. 5b). The question at this point was whether autoimmunity per se or rather autoimmunity as a cause of increased serum TSH was responsible for progression.

In untreated patients, the frequency of PTC was significantly higher (9.3%) in those with hypothyroidism and/or high serum levels of TAb compared to patients with both low serum levels of TAb (6.8%) and negative serum TAb (6.4%; fig. 5a). Patients with hypothyroidism and/or high serum levels of TAb also had higher TSH levels compared to patients with lower serum levels of TAb or negative serum TAb (fig. 5b). The question at this point was whether autoimmunity per se or rather autoimmunity as a cause of increased serum TSH was responsible for progression.

Fig. 5. a Frequency of PTC according to serum levels of TAb and hypothyroidism. Patients with high (both TgAb and TPOAb >100 U/l) serum TAb or with hypothyroidism (black column) showed a significantly higher frequency of PTC compared to subjects with lower serum TAb levels (gray column) or negative TAb (white column). b Box-whisker plot of serum TSH (μU/ml) in patients with high serum TAb levels or hypothyroidism (black box), lower serum TAb levels (gray box) or negative TAb (white box). Results are reported as median values (black lines), interquartile (25th to 75th percentiles) ranges (boxes) and 10th to 90th percentiles (whiskers). The statistical difference between groups was evaluated using the Mann-Whitney U test. Patients with high serum TAb also showed serum TSH (1.16 μU/ml, interquartile range 0.52–2.00) compared to both patients with low TAb levels (0.75 μU/ml, interquartile range 0.39–1.34; Mann-Whitney U test: p < 0.001) and patients with negative TAb (0.72 μU/ml, interquartile range 0.40–1.10; Mann-Whitney U test: p = < 0.001). n.s. = Not significant.

Fig. 4. A schematic representation of the role of TSH in the development of PTC. TSH is a factor involved in the progression of thyroid cancer, and the risk of clinically detectable thyroid malignancy increases with serum TSH concentrations; even within normal ranges, higher TSH values are associated with a higher frequency of clinically detectable cancers. Thyroid diseases that affect the function of the gland, influencing pituitary secretion of TSH, are associated with a different risk of PTC, the likelihood of thyroid malignancy being reduced when TSH is lower, as in thyroid autonomy, and increased when TSH is higher, as in HT. Moreover, in patients with nodular thyroid disease, LT4 treatment, which reduces serum TSH, is associated with a significantly lower risk of developing clinically detectable thyroid cancer.
the increased frequency of PTC. A multivariate analysis was performed to identify the variables independently associated with an increased probability of PTC. According to a forward stepwise binary logistic regression model, serum TSH was the most significant factor associated with PTC, TPOAb as well as the clinical diagnosis of HT were unrelated with PTC, whereas TgAb showed a slight association with PTC (table 2). These data may be explained by the hypothesis that patients with PTC have circulating antibodies to Tg unrelated to the classical HT. In a recent paper, Boi et al. [63] found that both TAb and serum TSH levels represented independent risk factors of thyroid carcinoma in patients with nodular goiter and circulating TAb (non-AITD), but devoid of tissue lymphocytic infiltration at histology show a pattern similar to that in non-AITD patients.

**Table 2.** Independent predictors of the diagnosis of PTC in patients with nodular goiter or HT

<table>
<thead>
<tr>
<th>Variable</th>
<th>p</th>
<th>OR</th>
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<tbody>
<tr>
<td>TSH</td>
<td>&lt;0.0001</td>
<td>66.5</td>
</tr>
<tr>
<td>TgAb</td>
<td>0.0003</td>
<td>2.0</td>
</tr>
<tr>
<td>TPOAb</td>
<td>n.s.</td>
<td>–</td>
</tr>
<tr>
<td>Clinical diagnosis</td>
<td>n.s.</td>
<td>–</td>
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OR = Cumulative odds ratio; n.s. = not significant.

**Fig. 6.** Schematic representation of the immunodominant region recognized by human TgAb. Only a portion of the very large Tg molecule is depicted. Four immunodominant epitopes (A, B, C and D) are recognized by recombinant human TgAb-Fab directed to a specific epitope. The binding of TgAb to these epitopes on Tg is inhibited by the relative TgAb-Fab. In patients with AITD (including both HT and GD), the pattern of Tg recognition by TgAb was ‘restricted’ to the immunodominant regions of Tg, while in patients with non-AITD (including nodular goiter and PTC), TgAb bind also to other regions of Tg. Sera from patients with PTC associated with thyroiditis show an epitope recognition pattern similar to that of AITD, while those with PTC and devoid of severe lymphocytic infiltration at histology show a pattern similar to that of non-AITD patients.

**Epitope Pattern of TgAb and PTC**

The presence of two different autoimmune mechanisms, one typical of AITD and the other of an immune reaction to PTC, was demonstrated by our study on the epitope pattern of TgAb [31], using the same method as described in the previous paragraph and in figure 2. In this study, we evaluated the inhibition exerted by monoclonal TgAb-Fab A, B, C and D on the binding to Tg of sera from patients with AITD, namely HT and GD, from patients with nodular goiter and circulating TAb (non-AITD), but devoid of tissue lymphocytic infiltration at histology, and from patients with PTC and circulating TAb subdivided in those without (PTC-no-thyroiditis) or with associated thyroiditis at histology (PTC-thyroiditis). The pattern of AITD patients differed from that of non-AITD patients, because the latter group showed a significantly lower inhibition by monoclonal TgAb-Fab A, B, C and D. Sera from patients with PTC-thyroiditis showed an epitope recognition pattern similar to that of AITD, while those with PTC-no-thyroiditis had a pattern similar to patients with nodular goiter, indicating that in these conditions, TgAb are not restricted to the epitopes identified by the 4 TgAb-Fab that are dominant in AITD and, therefore, are characterized by a less restricted and more heterogeneous epitope pattern (fig. 6).
Therefore, in PTC patients, the pattern of TgAb epitopes is quite variable, being of a non-AITD type in the absence of lymphocytic infiltration and AITD-like in its presence. Discrepancies of Tg molecules could cause the different TgAb patterns of AITD and non-AITD. Another hypothesis for this discrepancy is the existence of different mechanisms of antigen presentation or processing of Tg in AITD and non-AITD.

**Conclusion**

Iodine intake modulates the pattern of thyroid diseases, and iodine supplementation is associated with an increased frequency of thyroid autoimmunity. As far as breaking of self-tolerance is concerned, the iodine effect is correlated with the intrinsic characteristics of Tg, namely its modifications induced by post-translational iodination. In the presence of genetic susceptibility, iodine, which is essential for normal thyroid function, induces thyroid autoimmunity by unmasking a cryptic epitope on Tg.

Thyroid autoimmunity is associated with an increased risk of PTC; however, different factors should be taken into account when evaluating this association. On historical grounds, the lymphocytic infiltration of the thyroid is the typical hallmark of HT, but PTC may be associated with a significant lymphocytic infiltration in the absence of the characteristic signs of autoimmune thyroiditis, as an immune response to the tumor. On clinical grounds, the diagnosis of HT is based on the presence of TAb, but these antibodies may be detected in euthyroid subjects in the absence of a diffuse lymphocytic infiltration of the thyroid, and PTC is associated with serum TAb only in patients with a lower prevalence of thyroid autoimmunity (i.e. men vs. women). The different epitope patterns of TgAb in AITD and non-AITD patients support the hypothesis that two different autoimmune mechanisms may be involved, one typical of AITD and the other of an immune reaction to PTC. Thyroid function may also affect the frequency of PTC, the risk of cancer increasing with serum TSH levels. We have shown that this mechanism, rather than thyroiditis per se, plays a major role in the association of PTC with HT, as a consequence of the autoimmune process leading to a progressive increase in serum TSH in these patients, even if the direct involvement of immunological mechanisms cannot be excluded.

**Disclosure Statement**

The authors have no conflicts of interest to disclose.


