We investigated factors underlying the varying effects of a high dietary iodide intake on serum T4 levels in a wide spectrum of mouse strains, including thyroiditis-susceptible NOD.H2h4, NOD.H2k, and NOD mice, as well as other strains (BALB/c, C57BL/6, NOD.Lc7, and B10.A4R) not previously investigated. Mice were maintained for up to 8 months on control or iodide-supplemented water (NaI 0.05%). On iodized water, serum T4 was reduced in BALB/c (males and females) in association with colloid goiters but was not significantly changed in mice that developed thyroiditis, namely NOD.H2h4 (males and females) or male NOD.H2k mice. Neither goiters nor decreased T4 developed in C57BL/6, NOD, NOD.Lc7, or B10.A4R female mice. In further studies, we focused on males in the BALB/c and NOD.H2h4 strains that demonstrated a large divergence in the T4 response to excess iodide. Excess iodide ingestion increased serum TSH levels to the same extent in both strains, yet thyroidal sodium iodide symporter (NIS) messenger RNA (mRNA) levels (quantitative polymerase chain reaction) revealed greatly divergent responses. NOD.H2h4 mice that remained euthyroid displayed a physiological NIS iodine autoregulatory response, whereas NIS mRNA was inappropriately elevated in BALB/c mice that became hypothyroid. Thus, autoimmune thyroiditis-prone NOD.H2h4 mice adapted normally to a high iodide intake, presumably by escape from the Wolff-Chaikoff block. In contrast, BALB/c mice that did not spontaneously develop thyroiditis failed to escape from this block and became hypothyroid. These data in mice may provide insight into the mechanism by which iodide-induced hypothyroidism occurs in some humans without an underlying thyroid disorder.

Abbreviations: cAMP, cyclic adenosine monophosphate; Duox1, dual oxidase 1; Duox2, dual oxidase 2; GAPDH, glutaraldehyde-3-phosphate dehydrogenase; mRNA, messenger RNA; NIS, sodium iodide symporter; PCR, polymerase chain reaction; Pds, pendrin; RT, real-time; SD, standard deviation; T4, thyroxine; Tg, thyroglobulin; TPO, thyroid peroxidase; TSHR, TSH receptor.
In mice, increased iodide ingestion causes divergent outcomes in two strains not spontaneously susceptible to thyroid autoimmunity (Table 1). SJL mice develop colloid goiter and hypothyroidism, whereas CBA mice remain euthyroid with normal thyroid histology [6]. Hypothyroidism and TSH-induced goiter in the SJL mice are caused by their inability to downregulate thyroid sodium iodide symporter (NIS) expression [6] and consequent failure to escape from the Wolff-Chaikoff block [7]. Very limited, or contradictory, information on thyroid function is available for strains that spontaneously develop thyroid autoimmunity, namely NOD.H2h4 and NOD.H2k mice [8, 9], as well as the parent NOD strain that primarily develops type I diabetes and, to a lesser extent, thyroid autoimmunity [10–12]. For example, excess iodide administration to diabetes-prone NOD mice modestly decreased serum T4 levels [13], whereas in the NOD.H2h4 strain, iodide had either no effect [10] or consistently reduced serum T4 levels [14]. Unlike in the SJL and CBA strains, there are no reports on potential mechanisms underlying iodide-induced thyroid dysfunction in these thyroiditis-susceptible mouse strains.

Because of this strain-dependent variability, we investigated the serum T4 and thyroid histologic responses to excess dietary iodide in a wide spectrum of mouse strains, including thyroiditis-susceptible NOD.H2h4, NOD.H2k, and NOD mice, as well as other strains (BALB/c, C57BL/6, NOD.Lc7, and B10.A4R) for which the effect of increased dietary iodide on thyroid function has not been investigated. To examine the expression of genes related to thyroid function, namely thyroglobulin (Tg), thyroid peroxidase (TPO), the TSH receptor (TSHR), NIS, Duox1, Duox2, and pendrin (Pds), we focused on two mouse strains that do (NOD.H2h4), or do not (BALB/c), spontaneously develop thyroid autoimmunity. We confirmed that excess iodide ingestion had no effect on serum T4 levels in NOD.H2h4 mice [10–12, 15–17]. Further, hypothyroidism and goiter did develop in nonautoimmune-prone BALB/c mice because of an increase (rather than the anticipated decrease) in NIS on exposure to excess iodide.

1. Methods

A. Mice

The following mouse strains were investigated:

a. Male and female BALB/c, male and female NOD.H2h4 (NOD.Cg.H2h4/DiLTacUmmJ), and female B10.A4R [B10.A.H2h4/(4R) SgDvEgJ] mice, originally from The Jackson Laboratory (Bar Harbor, ME), were bred at Cedars-Sinai Medical Center (Los Angeles, CA).

| Strain     | I-A   | M or F | Lymphocytic Infiltration | Goiter (Colloid) | T4        | Reference              |
|------------|-------|--------|--------------------------|-------------------|-----------|------------------------|
| BALB/c     | I-Ad  | M, F   | No                       | Yes               | Decreased | Present study          |
| NOD.H2h4   | I-Ak  | M, F   | Moderate                 | No                | No change | [10–12, 15–17]        |
|            |       |        |                          |                   |           | [18], present study    |
| NOD        | I-Ag7 | M      | Small                    | No                | Not done  | [12, 61, 62]          |
|            |       |        |                          |                   |           |                       |
| NOD.Lc7    |       |        | Small                    | No                | No change | [18], present study    |
| NOD.H2k    |       |        | Small                    | No                | No change | [18], present study    |
| B10.A.4R   | I-Ak  | M, F   | Moderate                 | No                | Decreased (M) | Present study      |
| C57BL/6    | I-Ab  | F      | No                       | Yes               | Decreased | Present study          |
| CBA        | I-Ak  | F      | None                     | No                | None      | [6, 11]               |
| SJL        | I-As  | F      | Small                    | Yes               | Decreased | [6]                   |

Sex indicated by male or female.
Abbreviations: F, female; I-A, mouse major histocompatibility class II antigens; M, male.
b. Female C57BL/6 mice were purchased from The Jackson Laboratory and housed at Cedars-Sinai Medical Center.

c. Sera and thyroid tissue were available from previous studies of thyroid autoimmunity in male mice of the following strains: NOD.H2h4, NOD.H2k, NOD, B10.A4R, and NOD.Lc7 mice [18]. NOD.Lc7 mice are congenic for a segment on the distal end of mouse Chr 7, which coincides with the B10.A4R interval retained in the generation of NOD.H2h4 mice [18]. NOD mice had been tested for urinary glucose using Diastix Reagent Strips for Urinalysis (Glucose Bayer Health Care LLC, Mishawaka, IN) to exclude data from mice that developed diabetes.

All mouse studies were performed in accordance with the guidelines and approval of the Institutional Animal Care and Use Committee at Cedars-Sinai Medical Center and performed with the highest standards of care.

B. Exposure to Iodide-Supplemented Water

From the age of 8 weeks (2 months), when both males and females are sexually mature, mice were maintained on regular water or water supplemented with 0.05% sodium iodide (iodized water) [19] for 16 weeks in all mouse strains and for 32 weeks in NOD.H2h4, NOD.H2k, and BALB/c. Blood was drawn before treatment and after 8 or 16 weeks on regular water or NaI. At the 16-week time point, some mice (including all NOD and NOD.Lc7) were euthanized to harvest blood and thyroid tissue.

C. T4, TSH, and Thyroid Histology

Total serum thyroxine (T4) was measured in undiluted mouse serum (25 μL) using a radio-immunoassay (Diagnostic Products Corporation, Los Angeles, CA). T4 values were computed from kit standards and expressed as μg/dL. Serum TSH was measured (50 μL) in a radio-immunoassay by Dr. Samuel Refetoff (University of Chicago; fee for service). T4 values for male and female NOD.H2h4 mice after 16 weeks on NaI were included in a previous study [20]. Thyroid glands were preserved as previously described [18], and paraffin-embedded and serial sections were stained with hematoxylin and eosin (Research Animal Diagnostic Laboratory, University of Missouri, Columbia, MO). Thyroid histology was examined in Los Angeles. The extent of thyroid lymphocytic infiltration was previously reported for male NOD.H2h4, NOD.H2k, NOD, NOD.Lc7, and B10.A4R mice [18]. Histological data have not previously been reported for BALB/c, C57BL/6, or female B10.A4R mice.

D. Intrathyroidal mRNA Expression Measured by RT-PCR

Thyroids from mice exposed for 16 weeks to regular water (NOD.H2h4, n = 2; BALB/c, n = 4) or NaI water (NOD.H2h4, n = 4; BALB/c, n = 4) were stored in RNAlater (LifeTechnologies, Carlsbad, CA). Quantitative real-time (RT) polymerase chain reaction (PCR) was performed essentially as previously described [21]. Tissue was homogenized with QIAshredder columns (QIAGEN, Valencia, CA). Total RNA was prepared using RNeasy Plus Mini kit and treated with TURBO DNase (LifeTechnologies, Carlsbad, CA) to remove genomic DNA. Reverse transcription was performed with the AffinityScript QPCR cDNA Synthesis Kit (Agilent Technologies, Cedar Creek, TX) using oligo(dT) and random primers. Quantitative RT-PCR was performed using the FastStart SYBR Green Master Mix (Roche, Basel, Switzerland) with 2.5% of complementary DNA (20 μL final volume). Reactions were run on an iCycler Thermal Cycler with an iQ5 Real Time PCR Detection System module (Bio-Rad Laboratories, Hercules, CA). An initial denaturation step at 95°C (10 minutes) was followed by denaturation at 95°C (30 seconds) and annealing and extension at 55°C (30 seconds) for 40 cycles. Relative gene expression levels were calculated using the comparative Ct method (Delta delta Ct), according to the Pfaffl model [22] using Bio_Rad iQ5 2.0 software. Samples were tested in triplicate; parallel controls lacked reverse transcription. The genes studied included: NIS,
Pds, dual oxidase 1 (Duox1) and Duox2, and the housekeeping genes β-actin and glutaraldehyde-3-phosphate dehydrogenase (GAPDH). Primers were obtained from QIAGEN, except for Pds primers, which were obtained from Bio-Rad. Data were normalized to mouse β-actin and GAPDH; results are shown as mean ± standard deviation (SD).

**E. Statistical Analyses**

Significant differences between responses in different groups were determined by Mann-Whitney rank sum test or, when normally distributed, by Student t test. Multiple comparisons were made using analysis of variance. Tests were performed using SigmaStat (Jandel Scientific Software, San Rafael, CA).

**2. Results**

**A. Serum T4 in Mice on Iodized vs Regular Water**

We initially compared the response to iodide supplementation of water in two mouse strains that do (NOD.H2h4), or do not (BALB/c), spontaneously develop thyroiditis when maintained on regular water. In the former, excess iodide ingestion exacerbates the thyroiditis. After 16 and 32 weeks on iodized drinking water, serum T4 levels were significantly reduced in BALB/c mice of both sexes (Fig. 1A). However, excess iodide ingestion in thyroid autoimmunity-prone NOD.H2h4 mice (male and female) did not alter serum T4 levels (Fig. 1B).

In addition to the two foregoing mouse strains, we studied the serum T4 responses to excess iodide ingestion in five other strains with variable susceptibility to autoimmune thyroiditis, and for whom such data have not previously been reported (Fig. 2). Like NOD.H2h4 (Fig. 1), NOD.H2k and NOD.Lc7 are derived from the original NOD strain. In a long-term study with NOD mice, development of overt diabetes requiring insulin therapy is a confounding factor in assessing thyroid function, and mice becoming hyperglycemic were excluded from analysis. Further, because diabetes develops earlier in females, we studied NOD males and, for comparison, males in the other NOD-derived strains (NOD.H2k and NOD.Lc7). Excess iodide ingestion only decreased serum T4 levels in the original NOD founders (Fig. 2C), not in the two NOD variants (Fig. 2A and B).

Thyroiditis-prone NOD.H2h4 mice (the focus of studies described later) are derived by crossing regular NOD with B10.A4R mice. In the latter strain, 16-week exposure to excess dietary iodide significantly decreased serum T4 levels in males (Fig. 2D), but the suggestive decline in females did not attain statistical significance (Fig. 2E). Finally, we studied C57BL/6 females because they differ markedly from BALB/c females in failing to develop hyperthyroidism in an induced model of Graves disease [23]. Despite this difference, ingestion of a high iodide diet by C57BL/6 mice, like BALB/c mice (Fig. 1A), clearly decreased serum T4 levels, a phenomenon even observable after 8 weeks, the earliest time point studied (Fig. 2F).

**B. Thyroid Histology in Relation to Iodide Intake**

The histological changes induced by a high iodide diet were strikingly different in BALB/c and NOD.H2h4 mice of both sexes. BALB/c mice developed large colloid goiter in association with the decline in serum T4. In contrast, thyroid of NOD.H2h4 mice (whose T4 levels did not decrease) were not enlarged but had moderate thyroid lymphocytic infiltrates. Representative examples after 32 weeks on control or iodized water are shown for female BALB/c (Fig. 3A and 3B) and female NOD.H2h4 (Fig. 3C and 3D) mice. Moderate thyroiditis was previously reported for male NOD.H2h4 and NOD.H2k mice [18] and NOD.H2h4 females [20] after iodide exposure for 16 weeks. Thyroid tissue from male NOD mice on control vs iodized water revealed small lymphocytic infiltration (Fig. 3E and 3F) as reported for NOD.Lc7 [18]. After exposure to iodized water, C57BL/6 mice developed colloid goiter (although less striking than in the BALB/c strain), and thyroid histology was unchanged in B10.A4R mice (data not shown).
C. Serum TSH in Mice on Iodized Water

We measured TSH levels in BALB/c and NOD.H2h4 mice, the two strains showing the greatest divergence in response to iodized water, namely hypothyroid goiters in BALB/c and euthyroidism in association with moderate lymphocytic infiltration in NOD.H2h4 mice. Sera were from mice after 16 weeks on iodide, the first time point at which T4 levels were significantly reduced in the BALB/c strain. Because of limited serum available from venepuncture (serum used for multiple assays), and with TSH requiring a large volume, data from only male mice could be subject to statistical analysis. Despite the difference between the BALB/c and NOD.H2h4 mice in their T4 responses to excess dietary iodide (Fig. 1), the extent of the iodide-induced increases in serum TSH were comparable in males of these two strains (Fig. 4). A similar tendency was observed in female mice (data not shown), but the numbers were too small for statistical analysis.

D. Expression of Genes Related to Thyroid Function

To examine factors underlying the marked difference between BALB/c and NOD.H2h4 mice in their response to excess dietary iodide intake, we assessed the messenger RNA (mRNA) expression level of thyroid genes related to regulation of thyroid hormone synthesis by quantitative
PCR. Although we did not quantitate protein expression levels for these thyroid genes, protein concentrations correlate with their respective mRNA when measured under the same conditions [26]. Consistent with the comparable elevation of serum TSH in both mouse strains, expression of Tg, TPO, and the TSHR mRNA coding for the major thyroid-specific genes were all significantly increased on a high iodide intake vs control chow with one exception, namely TPO in NOD.H2h4 mice (Fig. 5A). The expression levels of Duox1 and Pds were reduced in both BALB/c and NOD.H2h4 mice on a high iodide diet (Fig. 5B). Duox2 expression was unchanged.

The major, and highly surprising, difference between the two mouse strains was in the expression of NIS, the iodide transporter normally downregulated in the escape from the Wolff-Chaikoff block (Fig. 5A). Increased iodide intake reduced NIS expression in NOD.H2h4 mice that remained euthyroid. In contrast, NIS expression was increased in the BALB/c mice that became hypothyroid. Of note, this difference in NIS expression between BALB/c and NOD.H2h4 was unrelated to TSH stimulation, with TSH levels being increased by a similar extent in both strains (Fig. 4).

3. Discussion

Deficiency of dietary iodide, an essential element for thyroid hormone synthesis, leads to hypothyroidism with severe metabolic consequences. Paradoxically, it has been recognized for
more than 60 years [27] that ingestion of excess iodide, for which there are numerous sources [5], can, in some individuals, also cause hypothyroidism. Most susceptible to "iodide myxedema" are euthyroid individuals with pre-existing, subclinical thyroid disorders, particularly Hashimoto thyroiditis [28, 29] and Graves disease following radioiodine therapy or subtotal thyroidectomy [30] and after partial thyroidectomy for benign thyroid nodules [31]. Hemithyroidectomy of thyroid autoimmune-prone BbBioBreeding Wistar (BB/W) rats also predisposes to iodide-induced hypothyroidism [32]. On the other hand, iodide-induced hypothyroidism may be idiosyncratic, occurring in normal individuals [33], or may not develop in some patients with Hashimoto thyroiditis with features identical to affected individuals [28]. The underlying mechanism for iodide-induced hypothyroidism in humans is generally accepted to be failure to escape from the Wolff-Chaikoff block [2, 3] because of the inability to reduce iodide transport into the thyroid [4], although it should be appreciated that this

\[ \text{Figure 3. BALB/c mice develop colloid goiters and NOD.H2 \textsuperscript{a,b} mice develop moderate thyroiditis after exposure to iodized drinking water for 32 weeks. Some NOD mice develop small lymphocytic infiltrates on iodized water (16 weeks). Thyroid histology in representative (A, B) female BALB/c and (C, D) NOD.H2 \textsuperscript{a,b} mice and (E, F) male NOD mice on regular water (left panels) and iodized water (right panels). Magnification: ×10. Number of thyroids examined in each group: A, 5; B, 5; C, 5; D, 5; E, 9; F, 12.} \]
phenomenon can only be directly observed \textit{ex vivo} in experimental animals after removal of their thyroids.

The current study, taken together with previous reports, provides insight into the mechanism of “iodide myxedema,” and it is useful to categorize the contributing factors, or lack thereof, as follows:

1. \textit{Thyroid lymphocytic infiltration} is not a factor in iodide-induced hypothyroidism in rodents. Consistent with previous reports [10, 16], excess iodide ingestion increases the extent of thyroiditis (lymphocytic infiltration) in thyroid autoimmunity-prone mouse strains, particularly NOD.H2h4 and NOD.H2k, without suppressing serum T4 levels. Conversely, increased dietary iodide did suppress serum T4 levels in BALB/c and NOD.H2h4 mice on iodide supplementation are not significantly (ns) different, but are higher than in recombinant inbred males (analysis of variance, \( P < 0.05 \)). “Pool” comprises small aliquots of sera from five male BALB/c mice to provide sufficient volume (50 uL) for assay. Note this value is shown for interest and is not used for statistical analysis. T4 values for male NOD.H2h4 mice after 16 weeks on NaI were included in a previous study [20].

2. \textit{NaI symporter autoregulation}. The inorganic iodide transporter at the thyrocyte basal surface, subsequently identified as NIS [39], is the focal point of organic iodine autoregulation [1, 4]. Excess dietary iodide intake reduced NIS mRNA expression in thyroiditis-prone NOD.H2h4 mice that remained euthyroid on this dietary regimen,
consistent with a normal iodine autoregulatory response. These data confirm the previous finding in Sprague-Dawley rats that iodide administration reduced both NIS mRNA and protein expression in the thyroid [7]. On the other hand, the most remarkable observation in the current study is that hypothyroidism induced by excess dietary iodide in BALB/c mice was associated with an unphysiological, marked increase in NIS mRNA expression (Fig. 5B). Consequently, it is likely that iodide-induced hypothyroidism in BALB/c, but not NOD.H2h4, mice can be ascribed to the inability of the former strain to escape from the Wolff-Chaikoff block by decreasing NIS mRNA levels. We did not study a role for Mct8 or anoctamin_1 [40] in iodine autoregulation in these animals, a possible future investigation.

Li and Carayanniotis [6] observed that iodide-induced hypothyroidism in SJL mice was associated with failure to suppress NIS mRNA (which remained unchanged), whereas in CBA mice that remained euthyroid on the same regimen, NIS mRNA levels were greatly reduced.
Although not obtained using quantitative PCR, these data taken together with our quantitative data suggest that there is a graded inherent variability among different mouse strains in iodine autoregulation of NIS expression. Thus, excess dietary iodide increased NIS mRNA levels in BALB/c mice, had no effect in SJL mice, and suppressed NIS mRNA levels in NOD.H2b4 and CBA mice. Hypothyroidism ensues when autoregulation fails in BALB/c and SJL mice.

3. TSH stimulation of NIS expression or functional activity. Iodide transporter functional activity is influenced in a yin-yang manner by opposing factors. Suppression by excess dietary iodide of NIS activity is counterbalanced by TSH stimulation of NIS mRNA expression and translation in rat thyroid cells [41] as well as in human thyroid tissue in vivo [42]. Therefore, conditions in which TSH levels are elevated could counter the physiological iodine autoregulatory suppression of NIS activity. Indeed, subclinical thyroid disorders that are susceptible to iodide myxedema such as Hashimoto thyroiditis [29] and treated Graves disease [30] may have slightly elevated TSH levels. Depression of serum T4 would further increase TSH secretion in a vicious cycle with ensuing overt hypothyroidism. Consistent with a role for TSH in failure to escape from the Wolff-Chaikoff block is the observation of an increased TSH level in SJL, but not CBA, female mice, only the former becoming hypothyroid on iodized water [6].

At face value, the current study provides evidence against the foregoing concept that a TSH-driven increase in NIS expression is an important factor determining whether excess iodide ingestion leads to hypothyroidism. Iodide administration to BALB/c and NOD.H2b4 mice led to comparable increases in serum TSH levels (Fig. 4), yet hypothyroidism only developed in the former strain (Fig. 1). However, as discussed later, the likely reason for this apparent inconsistency is that, because of the iodine autoregulation mechanism, the TSH level in serum does not reflect the potency of the TSH stimulus at the level of adenylyl cyclase activation.

4. Iodine autoregulation and TSH functional activity. A corollary of iodine autoregulation is that this phenomenon modulates TSH functional activity at the level of the thyrocyte. When the serum TSH level is maintained constant, excess iodide diminishes the potency of TSH action on a number of metabolic responses in experimental animals in vivo (TSH injection posthypophysectomy), as well as in isolated thyroids in vitro [reviewed in 1]. Conversely, iodide depletion amplifies TSH action, for example increasing its goitrogenic effect [43]. Inhibition of iodide organification by methimazole or propylthiouracil reverses this effect, as well as having effects on multiple thyrocyte functions [reviewed in 1], implicating an, as yet unidentified, autoregulatory organic iodine intermediary (termed “X-I”) [44], perhaps an iodolactone or iodoaldehyde [reviewed in 45].

An interesting finding in our study is the divergence among molecules involved in thyroid hormone synthesis in their iodine autoregulatory responses, suggesting variability in the yin-yang balance between TSH (positive) and the putative X-I (negative) influences on molecules involved in thyroid hormone synthesis. Thus, elevated serum TSH levels in BALB/c and NOD.H2b4 mice were associated with increases in Tg, TPO, and TSHR mRNA (with the exception of unchanged TPO mRNA in NOD.H2b4 mice). These data, consistent with previous reports of TSH stimulation of these thyroid-specific proteins [for example, 46–52] suggest that Tg, TPO, and the TSHR are not involved in iodine autoregulation. That is, TSH action on these molecules is not significantly opposed by the putative X-I. On the other hand, iodide-induced suppression of Pds mRNA levels in the face of an elevated TSH level is consistent with the greater influence of X-I. Despite an increase in serum TSH, excess dietary iodide either had no effect (Duox2) or suppressed (Duox1) mRNA levels, consistent with transcriptional control independent of the TSH/X-I balance (Duox2) or a balance toward X-I (Duox1).
Most important, in terms of understanding the underlying mechanism for idiosyncratic susceptibility to iodide-induced hypothyroidism in rodents (and possibly in humans), our data suggest an imbalance in the TSH/X-I ratio between BALB/c and NOD.H2h4 mice. Despite similar elevations in serum TSH in both strains, reduced or absent inhibition of NIS by X-I in BALB/c mice [whether directly or indirectly via cyclic adenosine monophosphate (cAMP), discussed later] could lead to unrestrained TSH stimulation of NIS with consequent failure to escape from the Wolff-Chaikoff block. On the other hand, in NOD.H2h4 mice, the balance in the TSH/X-I ratio favors the latter. Consequently, escape from the Wolff-Chaikoff block and euthyroidism is maintained by a combination of NIS suppression and a compensatory increase in TSH, blunted, at least in part, by an autoregulatory reduction in TSH activation of adenylyl cyclase (discussed later).

5. Adenylyl cyclase activity. As mentioned previously, despite a constant TSH level in serum, iodine autoregulation modulates TSH functional activity. TSH action on the thyroid is largely effected by activation of adenylyl cyclase. Cyclic AMP, the product of adenylyl cyclase, is the major positive regulator for expression of NIS [41], TPO [53], and Tg [54], and also stimulates thyrocyte proliferation [55] in synergy with IGF-1 [56]. Control by cAMP of other thyroid-specific proteins is more complex. The functional activity of Pds (one of the apical transporters involved in iodide efflux from the thyrocyte) is increased by adenylyl cyclase activation [57], although its mRNA expression is not stimulated by TSH [42]. Duox2, and to a lesser extent Duox1, generate H2O2 necessary for iodide organification at the thyrocyte follicular lumen and are activated by TSH in a nontranscriptional mechanism [58]. The functional activity of only Duox1 is increased by forskolin via cAMP and protein kinase A [58].

Adenylyl cyclase is itself a pivotal target for iodine autoregulation in vivo [19] and in vitro [44, 59]. The unidentified organic iodine intermediary (X-I) directly suppresses the adenylyl cyclase catalytic unit [60]. Because of its central role in modulating the expression or function of different molecules involved in thyroid hormone synthesis, adenylyl cyclase could be the proximal target of X-I and be the mechanism by which iodine autoregulation modulates TSH activation of more downstream molecules. Alternatively, X-I could be targeting numerous other proteins, at a transcriptional or posttranscriptional level.

In conclusion, the idiosyncratic development of iodide myxedema in some humans, even in the absence of thyroid autoimmunity [33] as well as in diverse mouse strains, suggests a genetic contribution to this susceptibility. As mentioned previously, there is a graded inherent variability among different mouse strains in their susceptibility to iodine autoregulation of NIS expression. Future identification of genes contributing to this variability will be more readily accomplished in genetically identical mouse strains than in genetically diverse humans.

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