Thyroglobulin as a Biomarker of Iodine Deficiency: A Review

Zheng Feei Ma and Sheila A. Skeaff

Background: Thyroglobulin, produced exclusively by the thyroid gland, has been proposed to be a more sensitive biomarker of iodine status than thyrotropin or the thyroid hormones triiodothyronine and thyroxine. However, evidence on the usefulness of thyroglobulin (Tg) to assess iodine status has not been extensively reviewed, particularly in pregnant women and adults.

Summary: An electronic literature search was conducted using the Cochrane CENTRAL, Web of Science, PubMed, and Medline to locate relevant studies on Tg as a biomarker of iodine status. Since urinary iodine concentration (UIC) is the recommended method to assess iodine status in populations, only studies that clearly reported both Tg and UIC were included. For the purpose of this review, a median Tg <13 μ g/L and a median UIC ≥100 μ g/L (UIC ≥150 μ g/L for pregnant women) were used to indicate adequate iodine status. We excluded studies conducted in subjects with either known thyroid disease or those with thyroglobulin antibodies. The search strategy and selection criteria yielded 34 articles of which nine were intervention studies. The majority of studies (six of eight) reported that iodine-deficient pregnant women had a median Tg ≥13 μ g/L. However, large observational studies of pregnant women, including women with adequate and inadequate iodine status, as well as well-designed intervention trials that include both Tg and UIC, are needed. In adults, the results were equivocal because iodine-deficient adults were reported to have median Tg values of either <13 or ≥13 μ g/L. Only studies in school-aged children showed that iodine-sufficient children typically had a median Tg <13 μ g/L. Some of the inconsistent results may be partially explained by the use of different methodological assays and failure to assess assay accuracy using a certified reference material.

Conclusions: These data suggest that Tg does hold promise as a biomarker of iodine deficiency. However, it is associated with limitations. A median Tg cutoff of $13 \mu g/L$ warrants further investigation, particularly in adults or pregnant women, as there is a lack of both observational and intervention studies in these groups.

Introduction

I ODINE IS NEEDED BY THE THYROID GLAND to produce thyroid hormones required for normal growth and development (1). Insufficient iodine intake causes iodine deficiency, which affects millions of people worldwide (2). Iodine deficiency is most commonly assessed by measuring urinary iodine concentration (UIC) because approximately 90% of dietary iodine is excreted in the urine (3). Due to large intraand interindividual variation, UIC cannot be used to assess iodine status in individuals and is only appropriate for groups (4). A median UIC <100 μ g/L in children and nonpregnant adults indicates iodine deficiency (5). Since UIC only assesses recent iodine intake (i.e., days) (5), a low UIC in a single urine sample does not necessarily indicate iodine deficiency in that individual (4). In addition to UIC, other measures of iodine status include thyroid volume, thyrotropin (TSH), triiodothyronine (T3), and thyroxine (T4); each of these indices has limitations. Thyroid volume reduces gradually (i.e., months to years) in previously iodine-deficient subjects (6). TSH, T3, and T4 concentrations typically fall within the normal range in mildly iodine-deficient populations of school-aged children and adults (7,8) such as those who live in developed countries such as the United States, the United Kingdom, Australia, and New Zealand. Another biomarker of iodine status sensitive to an intermediate change (i.e., weeks to months) in iodine intake would be useful.

Thyroglobulin (Tg) plays an important role in the synthesis of thyroid hormones T3 and T4 (9). It is a glycoprotein comprising two 330 kDa protein chains synthesized in the thyrocyte (10). After synthesis, Tg is transported and stored in the follicular colloid of the thyrocyte (11). In the follicular lumen, the tyrosine residues of Tg undergo iodination to

Department of Human Nutrition, University of Otago, Dunedin, New Zealand.

produce mono- (MIT) and di-iodotyrosines (DIT) catalyzed by thyroid peroxidase (12) and hydrogen peroxide (13). Subsequent coupling of these iodotyrosines produces T3 and T4 (14,15). Tg is pinocytosed into the thyroid cell (16) and undergoes proteolysis by lysosomes to release T3 and T4 (17), which are then secreted into the bloodstream (18).

When iodine intake is insufficient, low circulating levels of T4 stimulate the release of thyrotropin-releasing hormone from the pituitary gland, which subsequently increases the production of TSH. In addition to increasing the synthesis and proteolysis of Tg, TSH also stimulates the growth and division of the follicular cells, which causes the thyroid gland to enlarge (i.e., goiter) (19). In iodine deficiency, an increased amount of Tg is released into the blood (20), which is positively correlated with thyroid volume (21). For example, healthy adults have a mean Tg concentration ranging from 5 to $14 \,\mu g/L$ (22–27). In contrast, adults with endemic goiter have a mean Tg ranging from 94 to $208 \,\mu g/L$ (28–30). Recently, experts attending a National Institutes of Health workshop (31) recommended that Tg be used in the evaluation of iodine status.

The most common use of Tg is to monitor the treatment of patients with differentiated thyroid cancer (DTC) (32). Several review articles have focused on Tg monitoring in patients with DTC (11,33,34) or the performance of different assays used for monitoring DTC (35,36). The evidence on the usefulness of Tg in patients with DTC is well established. However, data on the effectiveness of Tg to assess iodine status in healthy populations is scarce. This review will report on: first, the analytical issues of Tg methods; second, observational studies measuring Tg to assess iodine status in healthy populations of pregnant women, newborns, children, and adults; and third, intervention studies investigating the effect of iodine supplementation on Tg in populations of pregnant women, newborns, children, and adults. This information will be used to determine if Tg can be used as a biomarker to assess iodine status.

Search Strategy

We conducted an electronic literature search using the Cochrane CENTRAL, Web of Science, PubMed, and Medline (OvidSP) to locate relevant studies published in English between January 1960 and October 2013 using Tg as a biomarker of iodine status. We used the following combined keywords: serum thyroglobulin, thyroglobulin, blood, children, infants, adults, pregnant women, pregnancy, maternal iodine status, iodine status, iodine deficiency, iodine insufficiency, iodine sufficiency, and iodine supplementation. We also located additional studies from references in the retrieved articles. Since UIC is the recommended biomarker of iodine status in populations (5), only studies that clearly report both Tg and UIC were included. We excluded studies conducted in subjects with either known thyroid disease or those with thyroglobulin antibodies (TgAb) because such subjects can have falsely low or high Tg that are not caused by insufficient iodine intake. The search resulted in 34 articles (i.e., 38 studies) being selected (Table 1). Of these, nine were randomized controlled trials, two were nonrandomized controlled trials, three were cohort observational studies, 23 were cross-sectional studies (10 multicenter), and one was a monitoring report of iodization programs that included a measurement before the introduction of iodized salt to a measurement after the introduction of iodized salt. In order to investigate the consistency of the relationship between iodine status as determined by UIC and Tg more clearly, for those studies that reported this information for more than one group, we considered each of these groups separately (i.e., one study of pregnant women and their newborns (37); one study of pregnant women and adults (38); one study of children and adults (30); three studies of children living in different regions (39) or countries (40,41); and seven studies of adults living in different regions (21,25,42–44) or countries (45,46)).

Discussion

Methods to measure Tg concentration

Tg can be measured using either immunometric assay (IMA) or radioimmunoassay (RIA) (35). Of the 34 articles measuring Tg (Table 1), the predominant Tg assay used was RIA (27%), followed by various IMAs, including immunoluminometric assay (22%), immunochemiluminescence assay (21%), immunoradiometric assay (12%), fluoroimmunoassay (10%), enzyme-linked immunosorbent assay (3%), electrochemiluminescence immunoassay (3%), and not reported (3%). Only one article (21) measured Tg using two different types of assays. A dried blood spot method using fluoroimmunoassay (FIA) (40) has been developed by Zimmermann et al. to assess Tg in children (5). Though Tg obtained from a dried blood spot was well correlated with serum samples (r=0.98, p<0.0001) in healthy children (n=29) (47), this relationship has yet to be validated in populations of adults including pregnant women. Furthermore, the dried blood spot method has not been reproduced in other laboratories.

Studies of Tg using RIA were first published in the 1960s. Several of these early studies (48-50) reported that Tg was undetectable in some healthy participants. For example, a small study conducted by Hjort et al. (48) used a RIA with a limit of detection (LoD) of 50 μ g/L and found that Tg was undetected in all 12 healthy subjects, indicating that these subjects would likely have had Tg concentrations $\leq 50 \,\mu$ g/L. In contrast, Torrigiani et al. (49) detected Tg in 60-70% of healthy subjects (n = 111) when they used a RIA with a LoD of $10 \,\mu g/L$; van Harle *et al.* (50) detected Tg in 74% of healthy subjects (n=95) using a RIA with a LoD of 1.6 μ g/L. Therefore, early RIAs had a relatively poorer functional sensitivity compared with first-generation Tg assays $(0.5-1.0 \,\mu\text{g/L})$ developed in the 1980s (51,52) and second-generation Tg assays $(\leq 0.1 \,\mu g/L)$ in use since the early 2000s (53,54); studies using first-generation Tg assays (21,55) did not report undetectable Tg in any healthy subjects.

Tg has been reported to be method dependent (56–58), and the interassay variation can vary between 43% and 65% in healthy subjects (35,57,59). To overcome interassay variation and allow for comparisons between studies, a certified Tg reference material (i.e., CRM-457) has been produced as a quality-control material for assay standardisation (60). Some but not all types of Tg assays have been standardized against CRM-457 in-house by the manufacturers (61). However, Tg CRM-457 only reduces interassay variation by 14–27% (59). It is suggested that this is because current Tg assays are unable to identify the heterogeneity of Tg epitopes (52,62). Of

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TABLE 1. TYPES OF TG AND TGAB ASSAYS IN STUDIES ASSESSING UIC AND TG IN VARIOUS POPULATION GROUPS

Studies	Year	Tg assay	TgAb assay
Pregnant women			
Observational			
Pedersen <i>et al.</i> $(38)^{a}$	1988	RIA	RIA
Laurberg et al. (76)	1994	Yes (NR)	NA
Eltom $et al.$ (78)	2000	RIA	NA
Costeira et al. (79)	2010	RIA	RIA
Brucker-Davis et al. (74)	2012	IRMA	NA
Raverot <i>et al.</i> (75)	2012	IRMA	NA
Andersen <i>et al.</i> $(37)^{b}$	2012	ILMA	RIPA
Brough <i>et al.</i> (77)	2013	ICMA	AA
	2015	ICIVIA	ΛΛ
Intervention Liesenkötter et al. (80)	1996	RIA	NA
Nøhr et al. (81)	2000	ILMA	RIPA
Santiago <i>et al.</i> (82)	2013	ICMA	NA
Newborns			
<i>Observational</i>	2012		DTD (
Andersen <i>et al.</i> $(37)^{b}$	2013	ILMA	RIPA
Intervention in pregnancy			
Pedersen et al. (85)	1993	ICMA	Tg recovery
Glinoer et al. (86)	1995	RIA	RĬA
Children			
Observational			
Simsek <i>et al.</i> (39)	2003	ICMA	NA
Zimmermann <i>et al.</i> $(40)^{c}$	2005	FIA	RIA
Bayram <i>et al.</i> $(30)^d$	2009	RIA	RIA
Shooff at $al (72)$	2009	RIA	NA
Skeaff <i>et al.</i> (73)			
Skeaff and Lonsdale-Cooper (55)	2013	ECLIA	NA
Zimmermann <i>et al.</i> (41)	2013	FIA	RIA
Intervention			
Benmiloud et al. (89)	1994	ILMA	NA
Zimmermann et al. (47)	2003	FIA	RIA
Zimmermann <i>et al.</i> $(40)^{c}$	2006	FIA	NA
Gordon et al. (64)	2009	RIA	RIA
Adults			
Observational			
Fenzi et al. (25)	1985	IRMA	HA
Gutekunst et al. (45)	1986	ILMA	NA
Pedersen et al. (38) ^a	1988	RIA	RIA
Hintze <i>et al.</i> (90)	1991	ELISA	RIA
Laurberg <i>et al.</i> (46)	1998	ILMA	RIPA
	2001		
Knudsen <i>et al.</i> (44)	2001	ILMA RIA	RIA NA
Thomson <i>et al.</i> (91)			
Rasmussen <i>et al.</i> (43)	2002	ILMA	RIA
Teng et al. (42)	2006	ICMA	ICMA
Bayram <i>et al.</i> $(30)^d$	2009	RIA	RIA
Vejbjerg et al. (21)	2009	ILMA & FIA	RIA & FIA
Cahoon et al. (72)	2013	ICMA	RIA & ICMA
Intervention			
Thomson <i>et al.</i> (63)	2009	ICMA	ICMA
Soriguer <i>et al.</i> (92)	2011	IRMA	RIA

^aIncluded pregnant women and adults. ^bIncluded pregnant women and newborns.

^cCounted as one article. ^dIncluded children and adults.

AA, agglutination assay; ECLIA, electrochemiluminescence immunoassay; ELISA, enzyme-linked immunosorbent assay; FIA, fluoroimmunoassay; HA, hemaglutination assay; ICMA, immunochemiluminescence assay; ILMA, immunoluminometric assay; RIA, immunoradiometric assay; NA, not assessed; NR, not reported; RIA, radioimmunoassay; RIPA, radioimmunoprecipitation assay; Tg, thyroglobulin; TgAb, thyroglobulin antibodies; UIC, urinary iodine concentration.

Another issue with regard to the measurement of Tg is the presence of TgAb. When a RIA is used, a subject positive for TgAb will most likely have a higher Tg value, while IMA tends to lower Tg in TgAb-positive subjects (33,65). Thus, subjects who have a positive test for TgAb should be excluded from the results if Tg is used as a biomarker of iodine status in a population. In adults, studies (66-69) have found that 3–13% of adults have TgAb. However, in children, the prevalence of TgAb is lower (70,71), and Zimmermann *et al.* (47) suggest that screening for TgAb in this age group is not necessary. Twenty-two of 34 studies measured TgAb prior to Tg measurement (Table 1). Of these, the predominant TgAb assay used was a RIA (58%), followed by the radioimmunoprecipitation assay (14%), immunochemiluminescence assay (12%), Tg recovery (5%), hemagglutination assay (5%), agglutination assay (5%), and FIA (2%). Only two articles (21,72) measured TgAb using two different types of TgAb assays.

In a large multicenter study of healthy children aged 5–14 years, Zimmermann et al. reported a reference range for Tg of $4-40 \,\mu\text{g/L}$ as determined by FIA (40). This is similar to reference ranges reported for adults of $3-40 \,\mu\text{g/L}$ using both RIA and IMA methods (65,68). We did not identify any consistent effects of age or sex on Tg. Only one study (73) reported that Tg decreased with advancing age. In 1994, the WHO/ICCIDD/UNICEF suggested that a median Tg concentration $<10 \,\mu g/L$ indicates adequate iodine status in populations of school-age children. However, in 2007, the WHO/ICCIDD/UNICEF, although acknowledging that Tg could be used an indicator of iodine status, did not provide a cutoff for Tg. More recently, Zimmermann et al. (41) conducted a large multicenter study of children (n = 2512) from 12 countries with varying iodine status, and suggest that a median Tg concentration $<13 \,\mu$ g/L and/or <3% of Tg values $>40 \,\mu g/L$ be used as a biomarker of adequate iodine status in children and, with caution, in adults. To date, the cutoff of $13 \,\mu\text{g/L}$ and/or <3% of Tg values >40 $\mu\text{g/L}$ has not been examined in younger children or pregnant women. Because no studies have reported the percentage of Tg values $>40 \,\mu g/L$ in populations, for the purpose of this review, a median Tg <13 μ g/L and a median UIC \geq 100 μ g/L (UIC \geq 150 µg/L for pregnant women) were used to indicate adequate iodine status.

Pregnant women

Eight observational studies measuring Tg in iodinedeficient pregnant women aged between 15 and 46 years were identified (Table 2). Six of eight studies (37,38,74–77) reported that iodine-deficient pregnant women (either first, second, or third trimester, or at delivery) had a median Tg $\geq 13 \ \mu g/L$ (range 16–67 $\ \mu g/L$). Two of the eight studies (78,79) assessed Tg concentration in iodine-deficient women throughout their pregnancy (i.e., in each trimester); in one study (79), a median Tg <13 $\ \mu g/L$ was observed in all three trimesters, and in one study (78), a Tg $\geq 13 \ \mu g/L$ was reported in the first and third trimesters, but it was <13 $\ \mu g/L$ in the second trimester. Although six of eight studies (37,38, 74,75,77,79) collected information on the use of iodine supplements in pregnancy, of these, only one study (37) reported that the Tg concentration of women taking iodine supplements was significantly lower compared with women who did not take supplements (i.e., difference of $\sim 15 \,\mu g/L$). We are unaware of any published studies of pregnant women with adequate iodine status that include measures of both UIC and Tg.

Three intervention studies investigating the effect of iodine supplementation on Tg in iodine-deficient pregnant women were identified (Table 3). One of the studies (80) assessed Tg concentration in the first trimester before supplementation and then again at two weeks postpartum; one study (81) assessed Tg in the first and third trimesters; and one study (82) assessed Tg in all three trimesters and again 12-24 weeks postpartum. Tg concentrations in women in the first trimester (i.e., at baseline before supplementation) ranged from 13 to 25 μ g/L, and postpartum, in women that had received any type of additional iodine (i.e., supplements or iodized salt), Tg ranged from 8 to 18 μ g/L. Of the two studies with postpartum data (80,82), only one study (80) reported that women taking iodine supplementation in pregnancy had a postpartum median Tg <13 μ g/L. However, the interpretation of these findings is confounded by differences in study designs, including a lack of a placebo group, relatively small sample sizes (n=66-131), varying levels and types of supplemental iodine (iodized salt or supplements containing $150-300 \,\mu\text{g}$ iodine per day), duration of follow-up (2-24 weeks postpartum), and use of different Tg assays.

In summary, it appears that the majority studies typically report that iodine-deficient pregnant women have a median Tg $\geq 13 \mu g/L$. Furthermore, iodine supplementation does not consistently decrease Tg below this cutoff either during pregnancy or postpartum, although this may reflect inadequate supplementation, as UIC did not reach recommended cutoffs. More large observational studies of pregnant women, including women with adequate and inadequate iodine status, as well as good intervention trials that include both Tg and UIC, are required before conclusions can be drawn about the usefulness of Tg as a biomarker of iodine status in pregnancy. Another consideration is whether Tg in pregnancy needs to be trimester specific, as is suggested for thyroid hormones such as TSH (83) and T4 (84).

Newborns

Three studies that measured Tg in cord blood from newborns were identified (Tables 4 and 5). Two of the three studies (85,86) were supplementation trials of mothers during pregnancy. The Tg concentration of newborns born to mothers receiving a placebo or who did not take supplements in pregnancy ranged from 62 to $113 \mu g/L$, while in the newborns of mothers who took iodine supplements, Tg ranged from $31-65 \mu g/L$. The usefulness of measuring Tg in newborn cord blood is questionable. A more commonly used and relatively accessible biomarker to assess iodine status in newborns is neonatal TSH collected by heel prick two to three days after birth (5).

Children

Six observational studies measuring Tg in children aged between 5 and 14 years were identified (Table 4). Four studies (30,39,41,73) found that iodine-deficient children had a median Tg \geq 13 µg/L (range 13–59 µg/L), while two studies

	14	BLE 2. UBSERVA1	TIONAL STUDIES ME	ASURING 1G	IN KELATION TO	1 ABLE 2. UBSERVATIONAL STUDIES MEASURING 1G IN KELATION TO IODINE STATUS IN PREGNANT WOMEN	REGNANT WOMEN	
Authors	Age (years) ^a ; n ⁵ ; country	Trimesters	UIC^{c} ($\mu g/L$)	/T)	T_{ξ}	Tg^{c} ($\mu g/L$)	Findings	Comments
Pedersen <i>et al.</i> (38)	21–38; <i>n</i> =20; Denmark	3rd	52 ^d			67	Suggested high Tg might be due to an increase in iodine intake in	Women did not take iodine supplements.
Laurberg et al. (76)	NR; <i>n</i> = 20; Denmark, Sweden, and Iceland	At term	Denmark Sweden Iceland	39 78 1118	Denmark Sweden Iceland	29.7 15.9 15.9	Pregnancy Women living in Denmark had a significantly higher median Tg than those living in Sweden and Iceland $(n < 0.05)$	No data on supplement use
Eltom <i>et al.</i> (78)	20–40; <i>n</i> =48; Sweden and Sudan	1st, 2nd, and 3rd	1. Swedish 8 ⁸ Sudanese 33	<i>Trimesters</i> 1st 2nd 3rd 89 89 76 38 25 38	Swedish Sudanese	<i>Trimesters</i> 1st 2nd 3rd 15.5 10.5 18.0 27.5 25.0 30.0	Sudanese women $p < 0.00$ significantly higher median Tg than the Swedish women in the 1st ($p < 0.001$), and 3rd trimesters ($n < 0.01$)	Women were followed throughout pregnancy; no data on supplement use
Costeira <i>et al.</i> (79,93) ^e	29.9; <i>n</i> = 118; Portugal	1st, 2nd and 3 rd trimester, and 1 year PP	<i>Trimester</i> 1st 2nd 3rd 1 vear PP	65 57 70 40	<i>Trimester</i> 1st 2nd 3rd 1 vear PP	11.0 111.0 112.7 9.7	Tg in 1st and 2nd trimester increased from 11 to 13 µg/L in 3rd trimester	Women were followed throughout pregnancy; they did not take iodine sumhements
Brucker-Davis et al. (74)	18–40; $n = 110$; France		116	2	t t mad t	17.4	Tg was not correlated with UIC	Women did not take iodine supplements
Raverot et al. (75)	15.3-45.7; <i>n</i> =228; France	lst, 2 nd , and 3rd	Trimester 1st 3rd	69 91 91	<i>Trimester</i> 1st 2nd 3rd	16.6 15.7 16.2	Tg in the 1st, 2^{nd} , or 3rd trimesters were not significantly different $(p > 0.05)$	Women did not take iodine supplements
Andersen <i>et al.</i> (37)	27.3; <i>n</i> = 140; Denmark	At term	No supplements Supplements All	NR NR 41 ¹	No supplements Supplements All		A	
Brough <i>et al.</i> (77)	31; $n = 70$; New Zealand	3rd trimester or breast- feeding for >3 weeks	3rd trimester PP	85 74	3rd trimester PP	15.9 13.9	Tg was not correlated with UIC in women in the 3rd trimester and at PP	70% pregnant women and 36% breastfeeding women used iodine supplements ranging from 100 to 150 µg I/day
^a Range used u	Range used unless mean reported.							

bouly subjects with no known thyroid disease or negative for TgAb were included. ^bOnly subjects with no known thyroid disease or negative for TgAb were included. ^cMedian used unless mean or geometric mean reported. ^dUIC reported as µg/g creatinine. ^eTg was reported in Costeira *et al.* (79); UIC and the data on supplement use were reported in Costeira *et al.* (93). These two studies were counted as one study (79). ^fGeometric mean. I, iodine; PP, postpartum.

TABLE 3. INTERVENTION STUDIES INVESTIGATING THE EFFECT OF IODINE SUPPLEMENTATION ON TG IN PREGNANT WOMEN

Findings Comments	Women taking Unequal group iodine supple- ments did not in iodine have a signifi- cantly lower group while Tg in the 1st 70 women in trimester and no iodine group); 0.5 months PP doiner proviouely	Women taking All women were supplements thyroid had a signifi- peroxidase cantly lower antibody-positive: Tg than wom- PP data for Tg en taking pla- was available ceb in the 3rd from a figure trimester only $(n=0.011)$	There was no significant women were difference in not iodine T within supplemented groups $(p = 0.13)$ or be- in the study; tween groups 32% of pregnant (p = 0.21) at least 1 year before their pregnancy
ł	Wom iod hav Car trifi 0.5	Wom sur car car car car car car trict	$\begin{array}{c} 3-6^{\text{g}} \\ months PP \\ 16.7 \\ 14.9 \\ 18.1 \\ 18.1 \\ 0.1 \\ 18.1 \\ 0.1 \\ 18.1 \\ 0.1 \\ 1$
$Tg~(\mu g/L)^{ m d}$	Ist trimester ^s 0.5 months PP ⁸ 16.6 13.5 16.5 8.3	Trimesters 3rd 19.4 14.1	Trimesters ⁸ 3rd 2nd 2nd 22.7 22.1 21.6 24.8 8.4 13.9 8.4 13.9
	Ist trimester ^s 16.5 16.5	1st 17.1 18.0 ^{f.g}	1st 21.3 25.4 13.0
$UIC (\mu g/L)^{d}$	0.5 months Pp ^e 50 ^f 105		3-6 3-6 144 NA 166 NA 212 NA
UIC (<i>Ist trimester^e 0.5 1</i> <i>Pl</i> 55 50 ⁶ 49 105	Trimesters 1st 3rd 51 53 51 ^{g.h} 105	Trimesters ⁸ 1st 2nd 145 130 117 177 137 222
Group ^c	0 µg 300 µg	0μg 150μg	IS 200 μg 300 μg
Study design ^b	Women ($n = 108$) randomized to receive either iodine supple- ments ($300 \ \mu g$ I/day) or no iodine until 0.5 months PP	Women ($n = 66$) randomized to receive a vita- min and min- eral tablet with 0 (placebo) or 150 µg I daily until 9 months	Women $(n = 131)$ randomized to receive either IS, 200 µg, or 300 µg I daily until 3–6 months PP
Age (years) ^a ; country	21–40; Germany	18–35; Denmark	31; Spain
Authors	Liesenkötter 21–40; et al. (80) Germany	Nøhr 18–35; et al. (81) Denmark	Santiago <i>et al.</i> (82)

^aRange used unless mean reported. ^bOnly subjects with no known thyroid disease or negative for thyroglobulin antibody were included. ^c Actual quantity of iodine from supplement; Liesenkötter *et al.* (80) and Santiago *et al.* (82) used iodine supplements in the form of KI. ^dMedian used unless mean reported. ^eUIC reported as µg/g creatinine. ^fEstimated value from a figure.

^gMean. ^hEstimated value from a table. IS, iodized salt; KI, potassium iodide.

	TABLE 4. OB	3SERVATIONAL STUDIES MEASU	JRING]	G IN RELATION TO IODINE	STATUS IN	OBSERVATIONAL STUDIES MEASURING TG IN RELATION TO IODINE STATUS IN NEWBORNS, CHILDREN, AND ADULTS	JLTS
Authors	Age (years) ^a ; n ^b ; country	UIC^{c} ($\mu g/L$)		Tg^{c} $(\mu g/L)$		Findings	Comments
Newborns Andersen et al. (37)	Newborns; n = 140; Denmark	No supplements Supplements All	44 ^d	No supplements Supplements All	61.6^{d} 31.1^{d} 50.0^{d}	Infants of mothers taking iodine supplements of 150 μ g I/day had a significantly lower Tg than infants born to mothers not taking supplements ($n < 0.001$)	A lag period (i.e., 5 days) between the collection of urine from infant and cord blood samples
Children Simsek et al. (39) $8-10$; $n=727$; Turkey	8–10; <i>n</i> =727; Turkey	<i>Urban</i> ^e Düzce Bolu <i>Rural</i> Yiğica Mudurnu Akçakoca Gerede	96 108 13 46 71 75	<i>Urban</i> ^e Düzce Bolu <i>Rural</i> Yiğilca Mudurnu Akçakoca Gerede	10.9 8.4 59.1 14.2 114.2 12.8	Severely iodine deficient children had a significantly lower Tg than children with mild to moderate iodine deficiency and iodine suffi- cient children ($p < 0.001$); Tg was negatively correlated with urinary iodine ($r = -0.27$, p < 0.001).	
Zimmermann et al. (40)	5-14; $n=710$; 5 countries	<i>Countries:</i> Bahrain Peru South Africa China Switzerland All	177 161 266 234 130 198	<i>Countries:</i> Bahrain Peru South Africa China Switzerland All	19.3 11.6 13.3 11.2 14.5	Purpose of this study was to determine the Tg reference interval of children (i.e., 4–40 μg/L). Of the five coun- tries, children from three countries had high Tg despite being iodine sufficient	
Bayram <i>et al.</i> (30)	10–14; <i>n</i> = 109; Turkey	51°		49.9°		Nongoifcout children had a significantly lower Tg than goitrous children ($p < 0.001$); Tg was negatively correlated with urinary iodine ($r=-0.611$, $p < 0.05$)	
Skeaff et al. (73)	5–14; <i>n</i> = 1153; New Zealand	Age (years) 5-7 8-10 11-14 All	63 67 68 68	Age (years) 5-7 8-10 11-14 All	16.2 12.5 11.1 12.9	Indine sufficient children had a significantly lower Tg than indine deficient children (p < 0.001)	
Skeaff and Lonsdale- Cooper (55)	8-10; n=147; New Zealand	113		10.8		UIC indicated adequate iodine status but Tg indicated mild iodine deficiency, suggesting that the children still had thyroid enlargement	Correction factor of 0.588 used to adjust for intraassay variation in Tg values between ECLIA and RIA method
							(continued)

	Age (years) ^a ; n ^b ;			IABLE 4. (CONTINUED)			
Authors	country	UIC^{c} ($\mu g/L$)		Tg^{c} ($\mu g/L$)		Findings	Comments
Zimmermann et al. (41)	6–12; <i>n</i> =2512; 12 countries	<i>Countries</i> : Morocco Tajikistan Switzerland Philippines Bahrain Peru Croatia Croatia Indonesia Paraguay South Africa All	$\begin{array}{c} 16\\ 15\\ 15\\ 15\\ 15\\ 10\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23\\ 23$	<i>Countries^d</i> : Morocco Tajikistan Switzerland Philippines Bahrain Peru Croatia China Indonesia Paraguay South Africa All	25.5 10.9 11.5 11.5 11.5 9.8 13.6 13.6 13.6 13.3	Tg followed a U-shaped curve from severely deficient to excessive iodine intake as assessed by UIC	
Adults Fenzi et al. (25)	38.8; <i>n</i> = 840; Italy	Endemic goiter Iodine sufficient	44 ^{e,f} 88 ^{e,f}	Endemic goiter Iodine sufficient	49.9° 9.5°	Adults living in an iodine suffi- cient area had a significantly lower Tg than adults living in an endemic goiter area (p < 0.01); Tg was negatively correlated with urinary iodine (r = -0.185, p = 0.001)	
Gutekunst <i>et al.</i> (45)	≥ 17 ; $n = 1291$; Germany and Sweden	Germany Sweden	63^{g} 141 ^g	Germany Sweden	43.0 21.2	German adults had a signifi- cantly lower Tg than Swedish adults ($p < 0.0001$)	
Pedersen <i>et al.</i> (38)	22–37; $n = 20$; Denmark	42 ^g		32.5			Six adults were positive for TgAb but were not included in Tg results
Hintze <i>et al.</i> (90) $60-97$; $n = 286$; Germany	60-97; n = 286; Germany	64 ^g		8.9		Adults with no goiter had a significantly lower median Tg than adults with goiter $(p < 0.001)$	
Laurberg et al. (46)	66-70; n = 523; Denmark and Iceland	Denmark Iceland	38 150	Denmark Iceland	15.5 9.5	In Denmark (Jutland), 14.2% adults living had Tg >50 μg /L; while in Iceland, 3.4% adults had Tg >50 μg /L	Differences in Tg between adults living Denmark and Iceland not reported
Knudsen <i>et al.</i> (44)	18–65; <i>n</i> = 3759; Denmark	Copenhagen Aalborg	68 53	Copenhagen Aalborg	11.3 15.2	in had than	Adults included those who were taking iodine sup- plements.
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Authors	Age (years) ^a ; n ^b ; country		UIC^{c} ($\mu g/L$)		T_{δ}	Tg^{c} $(\mu g/L)$		Findings	Comments
Thomson <i>et al.</i> (91)	18-49; n=233; New Zealand		54			5.1		Tg was negatively correlated with UIC $(r = -0.210, p = 0.003)$	
Rasmussen <i>et al.</i> (43)	18–65; <i>n</i> = 4649; Copenhagen Denmark Aalborg	Copenhagen Aalborg		68 53	Copenhagen Aalborg		11.5 15.4	Adults living in Copenhagen had Adults included pregnant a significantly lower Tg than women (1.3%) and adults living in Aalborg lactating women (1.8%) (p < 0.001); Tg was negatively correlated with iodine intake	Adults included pregnant women (1.3%) and lactating women (1.8%)
Teng et al. (42)	14-95; n = 3761; China	Panshan Zhangwu Huanghua	Baseline 103 375 615	5 years 97 350 635	Panshan Zhangwu Huanghua	Baseline 5 years 7.7 11.7 6.0 9.1 6.4 10.2	5 years 11.7 9.1 10.2	Adults living in Zhangwu city had a significantly lower Tg than adults living in Huanghua and Pangshan cities at baseline $(p < 0.001)$ and 5 years $(p < 0.001)$	
Bayram <i>et al.</i> (30)	28.7; <i>n</i> = 109; Turkey		31 ^e			68.9 ^e		Adults with no goiter had a significantly lower Tg than adults with goiter $(p < 0.001)$	
Vejbjerg <i>et al.</i> (21)	18–65; <i>n</i> = 4649; Denmark	Copenhagen Aalborg	IS Before 61 45	After 99 86	Copenhagen Aalborg	IS Before 11.5 15.4	After 9.1 9.3	Adults living in Copenhagen and Aalborg had a significantly lower Tg after introduction of mandatory iodization of salt (p < 0.001)	This study included two cross-sectional samples
Cahoon <i>et al</i> . (72)	Cahoon <i>et al.</i> (72) $10-33$; $n = 7890$; NR Belarus	NR			UIC (μg/L) 0-20 20-50 50-100 100-2120		$12.1 \\ 9.7 \\ 8.0 \\ 6.6$	Adults with UIC 100–2120 μ g/L had a significantly lower median Tg than subjects with UIC of 50–100, 20–50, and 0–20 μ g/L (p <0.001)	
^a Range used unless mean reported.	ss mean reported.		E						

TABLE 4. (CONTINUED)

^bOnly subjects with no known thyroid disease or negative for TgAb. ^cMedian used unless mean or geometric mean reported. ^dGeometric mean. ^eMean. ^fUIC reported as urinary iodine excretion (μg /day).

LTS	Comments	A lag period (i.e., 5 days) between the collection of urine from infants and cord blood samples	Cord blood used	The study only reported Tg values for baseline and 5 months		Main outcome was cognition
16 THE EFFECT OF IODINE SUPPLEMENTATION ON TG IN NEWBORNS, CHILDREN, AND ADULTS	Findings	Inf	(p = 0.002) Infants of mother supplemented with 131 μ g I and 131 μ g 1+L-T4 had a significantly lower Tg than infants of mothers taking placebo $(p < 0.001)$	All groups of children had a decrease in Tg after 5 months	Tg after using IS for 12 months (p < 0.001) Children had a significantly lower Tg after using IS for 10 months	($p < 0.001$) Children taking iodine tablets had a significantly lower Tg after 7 months ($p < 0.001$)
NEWBORN		67 38	113 ^d 65 ^d 52 ^d		12 4.4 10 8.0	
on Tg in Ì	Tg (μg/L) ^c			5 months 31.0 22.0 24.0 14.0 12.0 Months	Months 5 6.2 Months 5 13.0	7 months 11.6 8.5
EMENTATION	$T_{\mathcal{E}}$	0 µg 200 µg	0μg 131μg 131μg+L-T4	Baseline 98.5 175.0 77.0 62.0 Raveline	24.5 24.5 Baseline 49.0	Baseline 15.2 16.5
ve Suppl				15	12 165 165 100 102	57
OF IODIN	c)د	27 64	43 ^d 77 ^d 80 ^d	5 months 41 99 109 132 1185 Months	5 181 <i>Months</i> 5 74	7 months 81 145
he Effect	UIC (μg/L) ^c			Baseline 27 25 25 25 25 27 27 Baseline	Baseline 12	Baseline 62 66
VESTIGATING T		0μg 200μg	0μg 131μg 131μg+L-T4	<i>Orally</i> 120 mg 240 mg 480 mg <i>Injection</i> 480 mg 480 mg		0 μg 150 μg
TABLE 5. INTERVENTION STUDIES INVESTIGATIN	Study design ^b	Data obtained from infants ($n = 54$) born to mothers who were randomized to receive either 200 µg I/day or no iodine supplement	Data obtained from infants ($n = 180$) born to mothers who were randomized to receive either 131 μ g I/day, 131 μ g + 100 μ g L-T4, or placebo	Children $(n = 169)$ randomized to receive either a single dose of iodized poppy seed oil orally (120, 240, 480, or 960 mg 1) or intramuscular injection of 480 mg 1 for 150–395 days	Children $(n = 86)$ received IS for 12 months Children $(n = 86)$ received IS for 10 months	Children $(n = 184)$ randomized to receive either placebo or $150 \mu g$ I tablet daily for 28 weeks
TABLE 5. Ir	Age (years) ^a ; country	Newborns; Denmark	Newborns; Belgium	6–11; Algeria 6–15-	5-14; Morocco Morocco	10–13; New Zealand
	Authors	Newborns Pedersen et al. (85)	Glinoer et al. (86)	Children Benmiloud et al. (89) Zimmermann	Zimmermann 5–14; <i>et al.</i> (40) Morocco	Gordon et al. (64)

(continued)

						(777)			
Authors	Age (years) ^a ; country	Study design ^b		UIC (μg/L) ^c	0		Tg (μg/L) ^c	Findings	Comments
Adults Thomson et al. (63)	60–80; New Zealand	Adults $(n = 100)$ randomized to receive	0 μg	Baseline ^e 3 months ^e 49 44	3 months ^e 44	Baseline ^e 14.4	3 months ^e 14.0	Both groups taking iodine supplements	
		either placebo, 100 μ g Se, 80 μ g I, or 100 μ g Se+80 μ g I daily for 12 weeks	Se 80 μg Se + 80 μg	45 33 63	44 71 84	15.4 21.2 17.3	15.2 15.4 14.7	had a significantly lower Tg after 3 months compared to baseline $(p < 0.01)$	
Soriguer et al. (92)	34.9; Spain	A cross-over study of adults $(n = 30)$ randomized to 100,	100 μg 200 μg	Baseline ^{a,f} 192 140	$\begin{array}{rcl} Baseline^{a,f} & 2 \ months^{a,f} \\ 192 & 233 \\ 140 & 230 \end{array}$	Baseline ^a 3.9 7.6	$2 months^{d}$ 0.8 8.8	There was no difference A cross-over in Tg between the study with groups after 1 month	A cross-over study with 1 month
		200, or 300 µg I/day	300 μg	201	377	6.8	4.0	supplementation	washout; adults recruited were regular users of IS
(

TABLE 5. (CONTINUED)

^aRange used unless mean reported. ^bOnly subjects with no known thyroid disease or negative for TgAb. ^cMedian used unless mean or geometric mean reported. ^dMean. ^eGeometric mean. ^fUIC reported as urinary iodine excretion (μg/day). Se, selenium.

(40,41) reported that iodine-sufficient children also had a median Tg $\geq 13 \,\mu$ g/L (range 13–19 μ g/L). Four of six studies (39–41,55) reported that children with adequate iodine status had a median Tg <13 μ g/L. However, one study (39) reported that iodine-deficient children had a median Tg <13 μ g/L. The study by Zimmermann et al. (41) included 2512 children from 12 countries with severe iodine deficiency (i.e., median UIC $<20 \,\mu g/L$), mild iodine deficiency (i.e., median UIC 50–99 $\mu g/L$ L), adequate iodine status (i.e., median UIC 100–299 μ g/L), and iodine excess (i.e., median UIC \geq 300 µg/L). It showed that median Tg appeared to follow a U-shaped curve with the nadir at an UIC of 100–300 μ g/L. When iodine intake is very high, excess iodide transiently inhibits the activity of thyroid peroxidase and proteolysis of Tg, which subsequently reduces the synthesis and secretion of thyroid hormones (i.e., the Wolff-Chaikoff effect) (87). However, when prolonged excess iodine intake occurs, Tg could increase because the thyroid gland fails to escape from the Wolff-Chaikoff effect (88). Nonetheless, close examination of data reported by Zimmermann et al. (41) suggests that the relationship between UIC and Tg is highly variable. It is not known, however, how much of this variability is associated with UIC and/or Tg because a single UIC can be confounded by the hydration status, dietary intake, and diurnal variation (4).

Four intervention studies investigating the effect of iodine supplementation on Tg in iodine-deficient children aged 5–14 years for a duration of 5–12 months were identified (Table 5). Three of the four studies (40,47,64) were more than six months long, and reported that median Tg decreased significantly and fell below 13 μ g/L when UIC increased from < 100 to \geq 100 μ g/L. The remaining study (89) included five treatment groups but not a control group, was only five months long, and had fewer children in each group. In the three groups where the children became iodine sufficient, Tg was <13 μ g/L in only one group.

The majority of observational and intervention studies in school children appear to support the $13 \mu g/L$ cutoff proposed by Zimmermann *et al.* (41) to assess iodine status in this age group. However, the relationship between UIC and Tg is not always consistent, suggesting that Tg alone should not be used to assess iodine status in this group.

Adults

Twelve observational studies measuring Tg in adults aged between 18 and 97 years were identified (Table 4). Seven studies (21,25,30,38,43,45,46) showed that iodine deficient adults had a median Tg \geq 13 µg/L (range 16–69 µg/L), while only one study (45) reported that iodine-sufficient adults had a median Tg \geq 13 µg/L. However, 8 of 12 studies (21,25,42– 44,72,90,91) reported that adults who were categorized as iodine deficient had a median Tg <13 µg/L. One of these studies (42) included adults with iodine excess (i.e., median UIC \geq 300 µg/L) who, in contrast to the findings of Zimmermann *et al.* (41) in schoolchildren, had a median Tg <13 µg/L.

Two intervention studies investigating the effect of iodine supplementation on Tg in adults were identified (Table 5). One study included iodine-sufficient middle-aged adults supplemented with additional iodine for 8–12 weeks (92); at baseline, the median Tg was <13 μ g/L, which decreased, but not significantly, after supplementation. Another study of

older adults (60–80 years) (63) who were moderately iodine deficient reported a median Tg $\geq 13 \,\mu g/L$ at baseline. Although iodine status improved, the subjects remained mildly iodine deficient, which likely explains that, although Tg concentration significantly decreased after supplementation, it remained $\geq 13 \,\mu g/L$.

Based on these observational studies, it is difficult to conclude that the Tg cutoff of $13 \mu g/L$ suggested by Zimmermann *et al.* (41) for children can be used as a biomarker of iodine status in adults. Furthermore, there are no randomized placebo-controlled trials in adults that have shown an improvement in iodine status (indicated by an increase in baseline UIC from <100 to $\geq 100 \mu g/L$) results in a concomitant fall in Tg concentration from ≥ 13 to <13 $\mu g/L$.

Summary and Conclusions

Tg does hold promise as a biomarker of iodine deficiency. However, it is also associated with limitations. The variety of methods used to analyze Tg makes it difficult to compare studies. It would be helpful if studies that measured Tg standardized their assays with CRM-457. Furthermore, particularly in adult populations, subjects should be screened for TgAb. Despite these problems, the studies included in this review support the use of Tg as a biomarker of iodine status in school children using the $<13 \,\mu g/L$ cutoff as suggested by Zimmermann et al. (41). However, it is not possible to draw conclusions regarding the efficacy of Tg in adults because the data are equivocal, while there are no studies of pregnant women with adequate iodine status that also include data on Tg concentration. In particular, few intervention studies have investigated the diagnostic performance of Tg assays and its clinical relevance in assessing iodine status in healthy populations. Well-designed randomized placebo-controlled trials are required to investigate further the effect of iodine supplementation on Tg in mild to moderately iodine-deficient populations, particularly in adults and pregnant women.

Author Disclosure Statement

The authors declare that they have no conflict of interest.

References

- Zimmermann MB 2009 Iodine deficiency. Endocr Rev 30:376–408.
- Pearce EN, Andersson M, Zimmermann MB 2013 Global iodine nutrition: where do we stand in 2013? Thyroid 23: 523–528.
- Nath SK, Moinier B, Thuillier F, Rongier M, Desjeux JF 1992 Urinary excretion of iodide and fluoride from supplemented food grade salt. Int J Vitam Nutr Res 62:66–72.
- König F, Andersson M, Hotz K, Aeberli I, Zimmermann MB 2011 Ten repeat collections for urinary iodine from spot samples or 24-hour samples are needed to reliably estimate individual iodine status in women. J Nutr 141:2049–2054.
- WHO/UNICEF/ICCIDD 2007 Assessment of Iodine Deficiency Disorders and Monitoring Their Elimination: A Guide for Programme Managers. Third editions. WHO, Geneva.
- Zimmermann MB, Hess SY, Adou P, Toresanni T, Wegmüller R, Hurrell RF 2003 Thyroid size and goiter prevalence after introduction of iodized salt: a 5-y prospective study in schoolchildren in Côte d'Ivoire. Am J Clin Nutr 77:663–667.

- Zimmermann MB 2008 Methods to assess iron and iodine status. Br J Nutr 99:S2–S9.
- Bourdoux PP 1993 Biochemical evaluation of iodine status. In: Delange F, Dunn JT, Glinoer D (eds) Iodine Deficiency in Europe: A Continuing Concern. Plenum Press, New York, pp 119–125.
- Lamas L, Anderson PC, Fox JW, Dunn JT 1989 Consensus sequences for early iodination and hormonogenesis in human thyroglobulin. J Biol Chem 264:13541–13545.
- van de Graaf SA, Ris-Stalpers C, Pauws E, Mendive FM, Targovnik HM, de Vijlder JJ 2001 Up to date with human thyroglobulin. J Endocrinol **170:**307–321.
- Torréns JI, Burch HB 2001 Serum thyroglobulin measurement: Utility in clinical practice. Endocrinol Metab Clin North Am **30**:429–467.
- Deme D, Gavaret JM, Pommier J, Nunez J 1976 Maximal number of hormonogenic iodotyrosine residues in thyroglobulin iodinated by thyroid peroxidase. Eur J Biochem **70**:7–13.
- Song Y, Driessens N, Costa M, De Deken X, Detours V, Corvilain B, Maenhaut C, Miot F, Van Sande J, Many M-C, Dumont JE 2007 Roles of hydrogen peroxide in thyroid physiology and disease. J Clin Endocrinol Metab **92:**3764– 3773.
- Dunn JT, Anderson PC, Fox JW, Fassler CA, Dunn AD, Hite LA, Moore RC 1987 The sites of thyroid hormone formation in rabbit thyroglobulin. J Biol Chem 262:16948– 16952.
- Lamas L, Dorris ML, Taurog A 1972 Evidence for a catalytic role for thyroid peroxidase in the conversion of diiodotyrosine to thyroxine. Endocrinology 90:1417–1426.
- Kostrouch Z, Bernier-Valentin F, Munari-Silem Y, Rajas F, Rabilloud R, Rousset B 1993 Thyroglobulin molecules internalized by thyrocytes are sorted in early endosomes and partially recycled back to the follicular lumen. Endocrinology 132:2645–2653.
- Nunez J, Pommier J 1982 Formation of thyroid hormones. Vitam Horm **39:**175–229.
- de Vijlder JJ, den Hartog MT 1998 Anionic iodotyrosine residues are required for iodothyronine synthesis. Eur J Endocrinol 138:227–231.
- Pezzino V, Vigneri R, Squatrito S, Filetti S, Camus M, Polosa P 1978 Increased serum thyroglobulin levels in patients with nontoxic goiter. J Clin Endocrinol Metab 46:653–657.
- 20. de Vijlder JJ, Ris-Stalpers C, Vulsma T 1999 On the origin of circulating thyroglobulin. Eur J Endocrinol **140**:7–8.
- 21. Vejbjerg P, Knudsen N, Perrild H, Laurberg P, Carlé A, Pedersen IB, Rasmussen LB, Ovesen L, Jørgensen T 2009 Thyroglobulin as a marker of iodine nutrition status in the general population. Eur J Endocrinol 161:475–481.
- Ikekubo K, Jutton J, Schneider AB 1980 Radioimmunoassay of human thyroglobulin with use of "thyroglobulinfree" plasma prepared by ultracentrifugation as diluent. Clin Chem 26:1566–1568.
- Nakamura S, Sakata S, Minamori Y, Komaki T, Kojima N, Kamikubo K, Yasuda K, Miura K 1984 Serum thyroglobulin (Tg) concentration in healthy subjects: absence of ageand sex-related differences. Endocrinol Jpn 31:93–98.
- Pacini F, Pinchera A, Giani C, Grasso L, Doveri F, Baschieri L 1980 Serum thyroglobulin in thyroid carcinoma and other thyroid disorders. J Endocrinol Invest 3:283–292.
- Fenzi GF, Ceccarelli C, Macchia E, Monzani F, Bartalena L, Giani C, Ceccarelli P, Lippi F, Baschieri L, Pinchera A 1985 Reciprocal changes of serum thyroglobulin and TSH

in residents of a moderate endemic goitre area. Clin Endocrinol (Oxf) **23:**115–122.

- Mitchell ML, Klein RZ, Sargent JD, Meter RA, Haddow JE, Waisbren SE, Faix JD 2003 Iodine sufficiency and measurements of thyroid function in maternal hypothyroidism. Clin Endocrinol (Oxf) 58:612–616.
- Izumi M, Larsen PR 1978 Correlation of sequential changes in serum thyroglobulin, triiodothyronine, and thyroxine in patients with Graves' disease and subacute thyroiditis. Metabolism 27:449–460.
- Van Herle AJ, Hershman JM, Hornabrook RW, Chopra IJ 1976 Serum thyroglobulin in inhabitants of an endemic goiter region of New Guinea. J Clin Endocrinol Metab 43:512–516.
- 29. Bayram F, Borazan B, Torun E, Tanrıverdi F, Güven M, Erdoğan N, Muhtaroğlu S, Ünlühızarcı K, Tutuş A, Keleştimur F 2003 The prevalence of endemic goiter and iodine deficiency and evaluation of thyroid function in an area of central Anatolia. Turk J Endocrinol Metab 7:37–43.
- 30. Bayram F, Beyazyildiz A, Gökçe C, Budak N, Erdoğan N, Kurtoğlu S, Kula M, Ünlühızarcı K, Keleştimur F 2009 The prevalence of iodine deficiency, serum thyroglobulin, antithyroglobulin and thyroid peroxidase antibody levels in the urban areas of Kayseri, Central Anatolia. Exp Clin Endocrinol Diabetes 117:64–68.
- 31. Swanson CA, Zimmermann MB, Skeaff S, Pearce EN, Dwyer JT, Trumbo PR, Zehaluk C, Andrews KW, Carriquiry A, Caldwell KL, Egan SK, Long SE, Bailey RL, Sullivan KM, Holden JM, Betz JM, Phinney KW, Brooks SP, Johnson CL, Haggans CJ 2012 Summary of an NIH workshop to identify research needs to improve the monitoring of iodine status in the United States and to inform the DRI. J Nutr 142:1175S–1185S.
- 32. Black EG, Sheppard MC, Hoffenberg R 1987 Serial serum thyroglobulin measurements in the management of differentiated thyroid carcinoma. Clin Endocrinol (Oxf) 27:115–120.
- Spencer CA, Wang CC 1995 Thyroglobulin measurement: techniques, clinical benefits, and pitfalls. Endocrinol Metab Clin North Am 24:841–863.
- 34. Clark P, Franklyn J 2012 Can we interpret serum thyroglobulin results? Ann Clin Biochem **49:**313–322.
- 35. Spencer CA, Takeuchi M, Kazarosyan M 1996 Current status and performance goals for serum thyroglobulin assays. Clin Chem **42:**164–173.
- 36. Spencer CA, LoPresti JS 2008 Technology insight: measuring thyroglobulin and thyroglobulin autoantibody in patients with differentiated thyroid cancer. Nat Clin Pract End Met **4**:223–233.
- 37. Andersen SL, Nøhr SB, Wu CS, Olsen J, Pedersen KM, Laurberg P 2013 Thyroglobulin in smoking mothers and their newborns at delivery suggests autoregulation of placental iodide transport overcoming thiocyanate inhibition. Eur J Endocrinol 168:723–731.
- Pedersen KM, Börlum KG, Knudsen PR, Hansen ES, Johannesen PL, Laurberg P 1988 Urinary iodine excretion is low and serum thyroglobulin high in pregnant women in parts of Denmark. Acta Obstet Gynecol Scand 67:413–416.
- Simsek E, Safak A, Yavuz O, Aras S, Dogan S, Kocabay K 2003 Sensitivity of iodine deficiency indicators and iodine status in Turkey. J Pediatr Endocrinol Metab 16:197–202.
- 40. Zimmermann MB, de Benoist B, Corigliano S, Jooste PL, Molinari L, Moosa K, Pretell EA, Al-Dallal ZS, Wei Y, Zu-Pei C, Torresani T 2006 Assessment of iodine status using dried blood spot thyroglobulin: Development of reference

material and establishment of an international reference range in iodine-sufficient children. J Clin Endocrinol Metab **91:**4881–4887.

- 41. Zimmermann MB, Aeberli I, Andersson M, Assey V, Yorg JA, Jooste P, Jukić T, Kartono D, Kusić Z, Pretell E, San Luis TO, Jr., Untoro J, Timmer A 2013 Thyroglobulin is a sensitive measure of both deficient and excess iodine intakes in children and indicates no adverse effects on thyroid function in the UIC range of 100–299 μg/L: a UNICEF/ ICCIDD study group report. J Clin Endocrinol Metab **98**:1271–1280.
- 42. Teng W, Shan Z, Teng X, Guan H, Li Y, Teng D, Jin Y, Yu X, Fan C, Chong W, Yang F, Dai H, Yu Y, Li J, Chen Y, Zhao D, Shi X, Hu F, Mao J, Gu X, Yang R, Tong Y, Wang W, Gao T, Li C 2006 Effect of iodine intake on thyroid diseases in China. N Engl J Med **354**:2783–2793.
- 43. Rasmussen LB, Ovesen L, Bülow I, Jørgensen T, Knudsen N, Laurberg P, Perrild H 2002 Relations between various measures of iodine intake and thyroid volume, thyroid nodularity, and serum thyroglobulin. Am J Clin Nutr 76: 1069–1076.
- 44. Knudsen N, Bülow I, Jørgensen T, Perrild H, Ovesen L, Laurberg P 2001 Serum Tg-a sensitive marker of thyroid abnormalities and iodine deficiency in epidemiological studies. J Clin Endocrinol Metab 86:3599–3603.
- 45. Gutekunst R, Smolarek H, Hasenpusch U, Stubbe P, Friedrich HJ, Wood WG, Scriba PC 1986 Goitre epidemiology: thyroid volume, iodine excretion, thyroglobulin and thyrotropin in Germany and Sweden. Acta Endocrinol (Copenh) **112:**494–501.
- 46. Laurberg P, Pedersen KM, Hreidarsson A, Sigfusson N, Iversen E, Knudsen PR 1998 Iodine intake and the pattern of thyroid disorders: a comparative epidemiological study of thyroid abnormalities in the elderly in Iceland and in Jutland, Denmark. J Clin Endocrinol Metab 83:765–769.
- 47. Zimmermann MB, Moretti D, Chaouki N, Torresani T 2003 Development of a dried whole-blood spot thyroglobulin assay and its evaluation as an indicator of thyroid status in goitrous children receiving iodized salt. Am J Clin Nutr 77:1453–1458.
- Hjort T, Friis T, Lauridsen UB, Persson I 1974 Thyroglobulin in serum after TSH stimulation in hyperthyroidism. Acta Endocrinol (Copenh) 75:699–706.
- 49. Torrigiani G, Doniach D, Roitt IM 1969 Serum thyroglobulin levels in healthy subjects and in patients with thyroid disease. J Clin Endocrinol Metab **29:**305–314.
- Van Herle AJ, Uller RP, Matthews NI, Brown J 1973 Radioimmunoassay for measurement of thyroglobulin in human serum. J Clin Invest 52:1320–1327.
- Laurberg P, Pedersen KM 1987 A sensitive radioimmunoassay for serum thyroglobulin—including a correct screening for thyroglobulin autoantibodies. Scand J Clin Lab Invest 47:685–689.
- 52. Spencer CA, Bergoglio LM, Kazarosyan M, Fatemi S, LoPresti JS 2005 Clinical impact of thyroglobulin (Tg) and Tg autoantibody method differences on the management of patients with differentiated thyroid carcinomas. J Clin Endocrinol Metab **90**:5566–5575.
- Wunderlich G, Zöphel K, Crook L, Smith S, Smith BR, Franke WG 2001 A high-sensitivity enzyme-linked immunosorbent assay for serum thyroglobulin. Thyroid 11:819–824.
- 54. Spencer C, Fatemi S, Singer P, Nicoloff J, Lopresti J 2010 Serum basal thyroglobulin measured by a second-generation assay correlates with the recombinant human thyrotropin-

stimulated thyroglobulin response in patients treated for differentiated thyroid cancer. Thyroid **20:**587–595.

- 55. Skeaff SA, Lonsdale-Cooper E 2013 Mandatory fortification of bread with iodised salt modestly improves iodine status in schoolchildren. Br J Nutr **109:**1109–1113.
- 56. Schlumberger M, Hitzel A, Toubert ME, Corone C, Troalen F, Schlageter MH, Claustrat F, Koscielny S, Taieb D, Toubeau M, Bonichon F, Borson-Chazot F, Leenhardt L, Schvartz C, Dejax C, Brenot-Rossi I, Torlontano M, Tenenbaum F, Bardet S, Bussière F, Girard JJ, Morel O, Schneegans O, Schlienger JL, Prost A, So D, Archambeaud F, Ricard M, Benhamou E 2007 Comparison of seven serum thyroglobulin assays in the follow-up of papillary and follicular thyroid cancer patients. J Clin Endocrinol Metab **92:**2487–2495.
- Feldt-Rasmussen U, Schlumberger M 1988 European interlaboratory comparison of serum thyroglobulin measurement. J Endocrinol Invest 11:175–181.
- 58. Stanojevic M, Savin S, Cvejic D, Djukic A, Jeremic M, Zivancević Simonovic S 2009 Comparison of the influence of thyroglobulin antibodies on serum thyroglobulin values from two different immunoassays in post surgical differentiated thyroid carcinoma patients. J Clin Lab Anal 23:341–346.
- Zucchelli GC, Pilo A, Masini S, Prontera C, Ferdeghini M 1996 Large between-laboratory variability of thyroglobulin immunoassays: data collected in a collaborative study. J Clin Ligand Assay 19:234–238.
- 60. Feldt-Rasmussen U, Profilis C, Colinet E, Black E, Bornet H, Bourdoux P, Carayon P, Ericsson UB, Koutras DA, Lamas de Leon L, DeNayer P, Pacini F, Palumbo G, Santos A, Schlumberger M, Seidel C, Van Herle AJ, De Vijlder JJ 1996 Human thyroglobulin reference material (CRM 457). 1st part: assessment of homogeneity, stability and immunoreactivity. Ann Biol Clin (Paris) 54:337–342.
- 61. Spencer CA 2013 Measurement of thyroglobulin. In: Braverman LE, Cooper DS (eds) Werner and Ingbar's The Thyroid: A Fundamental and Clinical Text. Tenth edition. Lippincott Williams & Wilkins, Philadelphia, p 301.
- 62. Schulz R, Bethäuser H, Stempka L, Heilig B, Moll A, Hüfner M 1989 Evidence for immunological differences between circulating and thyroid tissue-derived thyroglobulin in men. Eur J Clin Invest 19:459–463.
- Thomson CD, Campbell JM, Miller J, Skeaff SA, Livingstone V 2009 Selenium and iodine supplementation: effect on thyroid function of older New Zealanders. Am J Clin Nutr **90**:1038–1046.
- Gordon RC, Rose MC, Skeaff SA, Gray AR, Morgan KM, Ruffman T 2009 Iodine supplementation improves cognition in mildly iodine-deficient children. Am J Clin Nutr 90:1264–1271.
- 65. Baloch Z, Carayon P, Conte-Devolx B, Demers LM, Feldt-Rasmussen U, Henry JF, LiVosli VA, Niccoli-Sire P, John R, Ruf J, Smyth PP, Spencer CA, Stockigt JR 2003 Laboratory medicine practice guidelines. Laboratory support for the diagnosis and monitoring of thyroid disease. Thyroid 13:3–126.
- Pedersen IB, Knudsen N, Jørgensen T, Perrild H, Ovesen L, Laurberg P 2003 Thyroid peroxidase and thyroglobulin autoantibodies in a large survey of populations with mild and moderate iodine deficiency. Clin Endocrinol (Oxf) 58:36–42.
- 67. Okamura K, Nakashima T, Ueda K, Inoue K, Omae T, Fujishima M 1987 Thyroid disorders in the general population of Hisayama Japan, with special reference to prevalence and sex differences. Int J Epidemiol 16:545–549.

- 68. Spencer CA, Takeuchi M, Kazarosyan M, Wang CC, Guttler RB, Singer PA, Fatemi S, LoPresti JS, Nicoloff JT 1998 Serum thyroglobulin autoantibodies: Prevalence, influence on serum thyroglobulin measurement, and prognostic significance in patients with differentiated thyroid carcinoma. J Clin Endocrinol Metab 83:1121–1127.
- Hollowell JG, Staehling NW, Flanders WD, Hannon WH, Gunter EW, Spencer CA, Braverman LE 2002 Serum TSH, T(4), and thyroid antibodies in the United States population (1988 to 1994): National Health and Nutrition Examination Survey (NHANES III). J Clin Endocrinol Metab 87:489–499.
- Marwaha RK, Tandon N, Karak AK, Gupta N, Verma K, Kochupillai N 2000 Hashimoto's thyroiditis: countrywide screening of goitrous healthy young girls in postiodization phase in India. J Clin Endocrinol Metab 85:3798–3802.
- 71. Kaloumenou I, Mastorakos G, Alevizaki M, Duntas LH, Mantzou E, Ladopoulos C, Antoniou A, Chiotis D, Papassotiriou I, Chrousos GP, Dacou-Voutetakis C 2008 Thyroid autoimmunity in schoolchildren in an area with long-standing iodine sufficiency: correlation with gender, pubertal stage, and maternal thyroid autoimmunity. Thyroid 18:747–754.
- 72. Cahoon EK, Rozhko A, Hatch M, Polyanskaya O, Ostroumova E, Tang M, Nadirov E, Yauseyenka V, Savasteeva I, McConnell RJ, Pfeiffer RM, Brenner AV 2013 Factors associated with serum thyroglobulin levels in a population living in Belarus. Clin Endocrinol (Oxf) 79:120–127.
- 73. Skeaff SA, Thomson CD, Wilson N, Parnell WR 2012 A comprehensive assessment of urinary iodine concentration and thyroid hormones in New Zealand schoolchildren: a cross-sectional study. Nutr J 11:31.
- Brucker-Davis F, Ferrari P, Gal J, Berthier F, Fenichel P, Hieronimus S 2012 Iodine status has no impact on thyroid function in early healthy pregnancy. J Thyroid Res 2012: 168764.
- 75. Raverot V, Bournaud C, Sassolas G, Orgiazzi J, Claustrat F, Gaucherand P, Mellier G, Claustrat B, Borson-Chazot F, Zimmermann M 2012 Pregnant French women living in the Lyon area are iodine deficient and have elevated serum thyroglobulin concentrations. Thyroid 22:522–528.
- 76. Laurberg P, Bjarnadottir R, Pedersen KM, Børlum KG, Hreidarsson AB 1994 Pregnancy is associated with an increase in s-thyroglobulin independent of iodine deficiency. A comparative study in countries with different levels of iodine intake. J Endocrinol Invest 17:72.
- 77. Brough L, Jin Y, Shukri NH, Wharemate ZR, Weber JL, Coad J 2013 Iodine intake and status during pregnancy and lactation before and after government initiatives to improve iodine status, in Palmerston North, New Zealand: a pilot study. Matern Child Nutr [Epub ahead of print]; DOI: 10.1111/mcn.12055.
- Eltom A, Elnagar B, Elbagir M, Gebre-Medhin M 2000 Thyroglobulin in serum as an indicator of iodine status during pregnancy. Scand J Clin Lab Invest 60:1–7.
- Costeira MJ, Oliveira P, Ares S, Roque S, de Escobar GM, Palha JA 2010 Parameters of thyroid function throughout and after pregnancy in an iodine-deficient population. Thyroid **20**:995–1001.
- Liesenkötter KP, Göpel W, Bogner U, Stach B, Grüters A 1996 Earliest prevention of endemic goiter by iodine supplementation during pregnancy. Eur J Endocrinol 134:443–448.
- 81. Nøhr SB, Jørgensen A, Pedersen KM, Laurberg P 2000 Postpartum thyroid dysfunction in pregnant thyroid peroxidase antibody-positive women living in an area with mild to moderate iodine deficiency: is iodine supplementation safe? J Clin Endocrinol Metab 85:3191–3198.

- 82. Santiago P, Velasco I, Muela JA, Sánchez B, Martínez J, Rodriguez A, Berrio M, Gutierrez-Repiso C, Carreira M, Moreno A, García-Fuentes E, Soriguer F 2013 Infant neurocognitive development is independent of the use of iodised salt or iodine supplements given during pregnancy. Br J Nutr **110**:831–839.
- 83. Stagnaro-Green A, Abalovich M, Alexander E, Azizi F, Mestman J, Negro R, Nixon A, Pearce EN, Soldin OP, Sullivan S, Wiersinga W 2011 Guidelines of the American Thyroid Association for the diagnosis and management of thyroid disease during pregnancy and postpartum. Thyroid 21:1081–1125.
- 84. De Groot L, Abalovich M, Alexander EK, Amino N, Barbour L, Cobin RH, Eastman CJ, Lazarus JH, Luton D, Mandel SJ, Mestman J, Rovet J, Sullivan S 2012 Management of thyroid dysfunction during pregnancy and postpartum: an Endocrine Society clinical practice guideline. J Clin Endocrinol Metab 97:2543–2565.
- 85. Pedersen KM, Laurberg P, Iversen E, Knudsen PR, Gregersen HE, Rasmussen OS, Larsen KR, Eriksen GM, Jøhannesen PL 1993 Amelioration of some pregnancyassociated variations in thyroid function by iodine supplementation. J Clin Endocrinol Metab 77:1078–1083.
- Glinoer D, De Nayer P, Delange F, Lemone M, Toppet V, Spehl M, Grün JP, Kinthaert J, Lejeune B 1995 A randomized trial for the treatment of mild iodine deficiency during pregnancy: Maternal and neonatal effects. J Clin Endocrinol Metab 80:258–269.
- Leung AM, Braverman LE 2013 Consequences of excess iodine. Nat Rev Endocrinol. doi: 10.1038/nrendo.2013.251.
- Scientific Committee on Food 2002 Opinion of the Scientific Committee on Food on the Tolerable Upper Intake Level of Iodine. SCF/CS/NUT/UPPLEV/26 Final. Health & Consumer Protection Directorate-General, European Commission, Brussels.
- Benmiloud M, Chaouki ML, Gutekunst R, Teichert HM, Wood WG, Dunn JT 1994 Oral iodized oil for correcting iodine deficiency: optimal dosing and outcome indicator selection. J Clin Endocrinol Metab 79:20–24.
- 90. Hintze G, Windeler J, Baumert J, Stein H, Köbberling J 1991 Thyroid volume and goitre prevalence in the elderly as determined by ultrasound and their relationships to laboratory indices. Acta Endocrinol (Copenh) 124:12–18.
- Thomson CD, Woodruffe S, Colls AJ, Joseph J, Doyle TC 2001 Urinary iodine and thyroid status of New Zealand residents. Eur J Clin Nutr 55:387–392.
- 92. Soriguer F, Gutiérrez-Repiso C, Rubio-Martin E, Linares F, Cardona I, López-Ojeda J, Pacheco M, González-Romero S, Garriga MJ, Velasco I, Santiago P, García-Fuentes E 2011 Iodine intakes of 100–300 μg/d do not modify thyroid function and have modest anti-inflammatory effects. Br J Nutr **105**:1783–1790.
- 93. Costeira MJ, Oliveira P, Ares S, de Escobar GM, Palha JA 2009 Iodine status of pregnant women and their progeny in the Minho Region of Portugal. Thyroid **19**:157–163.

Address correspondence to: Sheila Skeaff, PhD Department of Human Nutrition University of Otago Dunedin 9054 New Zealand

E-mail: sheila.skeaff@otago.ac.nz