Endemic Goiter - an update

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INTRODUCTION

“Goiter” means thyroid enlargement, and “endemic” means frequent in a certain locality. The problem is from what volume and up a thyroid is considered to be large enough as to be classified as goiter and also from what prevalence and up an area is considered to be endemic. It is obvious that by playing with these definitions one may increase or decrease the prevalence of goiter perceived in an area and also change the characterization of an area from endemic to non-endemic or vice versa.

In the past, goiter was defined as a thyroid gland enlarged 4 to 5 times above normal, and more practically as a thyroid with lobes greater than the terminal phalanges of the thumb of the person examined. Nowadays, the size of the thyroid gland is assessed by ultrasonography. By these means, Delange et al suggested variable limits for children and adolescents at different ages e.g. an upper limit of 16ml for boys and girls aged 15 years and 5ml for boys and girl aged 6 years. These data have been obtained in areas with a urinary iodine excretion of at least 100 μg/l. These limits have also been recommended by WHO and ICCIDD. For adults an upper limit of 25 ml for men and 18 ml for women has been suggested with smaller values for children. There is now a tendency to further decrease the upper normal limits. This would result in many persons with a small subclinical and non-palpable enlargement being classified as goitrous. The author doubts whether these subclinical enlargements should raise a concern.

Regarding the word endemic, this has been defined as a prevalence rate of more than 10%1,5. There is now a tendency to decrease this figure from 10% to 5%. Therefore, more areas will be classified as having a problem of endemic goiter.

To conclude, the tendency to decrease the upper normal limit of the thyroid volume together with the altered definition of endemicity from 10% to 5% will result in classifying several regions as endemic, whereas, in fact, the problem is not truly significant.

In this paper the problem of endemic goiter is reviewed with special emphasis on the studies of endemic goiter in Greece performed by our own team.

ETIOLOGY AND PATHOGENESIS

The main cause of endemic goiter is the lack of iodine. There is no renal homeostatic mechanism to keep the plasma inorganic iodine (PII) within the normal range. Thus, the PII fluctuates more or less directly with the iodine intake. The lower the iodine content in the diet, the lower the PII.

Adaptation to the iodine intake is achieved by a thyroidal mechanism. When the PII falls, the Thyroid Clearance of iodide (Th. Cl.) increases, i.e. the thyroid clears a higher volume of plasma from its iodide.
content. In this way, the absolute amount of iodide taken up by the thyroid (AIU = Absolute Iodine Up-
take) stays to a certain degree constant. That is, in the equation

\[ AIU = Th. \text{ Cl.} \times PII \]

the Th. Cl. changes inversely to the PII so as to keep the AIU normal. If the PII drops, owing to iodine-
deficiency, Th. Cl. increases and this increase in function is associated with an increase in the volume of the thyroid gland\(^7,8\). The thyroid normally needs about 2.5 \(\mu\)g/hr for thyroid hormone synthesis. When the PII level is 2.5 \(\mu\)g/l, the thyroid has to clear one l of plasma to obtain these 2.5 \(\mu\)g of iodine. If the PII is 1 \(\mu\)g/ l, the thyroid must clear 2.5 l of plasma. If the PII falls to below 0.8 \(\mu\)g/l and the thyroid has to clear more than three l of plasma per hour, thyroid enlargement (goiter) may develop.

The situation is complicated by other factors. Exogenous goitrogens\(^9-18\), such as cassava\(^9,16\), may lead to goiter, either by potentiating a mild iodine deficiency or autonomously. Vegetables from the genus Brassica may release thiocyanate, a well-known goitrogen\(^11-13,15-18\). Several other goitrogens have been described in various plants as well as in the drinking water. Iodine in large concentrations is in itself a goitrogen. “Iodide goiter” has been described in Japan in a coastal area where the inhabitants consumed large quantities of sea-plants with a high iodine content\(^19\), and recently in China due to a high iodine concentration in the drinking water\(^20\). The authors of the above studies suggest that iodine should not exceed 300 \(\mu\)g/l in the drinking water or 800 \(\mu\)g/l in the urine of the inhabitants.

In iodine-deficient populations not everyone is goitrous\(^5,21\). This phenomenon may be due to genetic factors. We have shown that there is a higher concordance rate for goiter in monozygotic than in dizygotic twins\(^22\). Such a difference in the thyroid size may be due to differences in the efficiency for utilising iodine, for instance iodide binding\(^23\).

The adaptation of man to iodine deficiency does not involve only an increase in the thyroidal iodide clearance. In iodine deficient areas, the ratio of \(T_3\) to \(T_4\) increases not only in the thyroid but also in the plasma\(^24\). \(T_3\) contains less iodine than \(T_4\) and is also metabolically more potent, hence this represents an additional mechanism to compensate for iodine deficiency. Which is the optimum iodine intake? Considering iodine kinetics, Wayne et al\(^6\) concluded that although some persons could adapt with 70\(\mu\)g iodine/ day, others would need 120 \(\mu\)g, depending on the renal iodide clearance, while 160 \(\mu\)g iodine/day constitute a safe level. We have found\(^25\) that serum TSH is lower when the urinary iodine is 150-200 \(\mu\)g /g Cr, or equivalent to an intake of about 200 \(\mu\)g /day if the fecal excretion is also taken into account.

Most authors and authorities broadly agree with these figures and the general conclusion may be that although euthyroidism may be maintained with as little as 50 \(\mu\)g/day, usually at the expense of goiter formation the optimum intake is 150 or 200 \(\mu\)g/day. Pregnant women have higher requirements since in pregnancy the renal clearance of iodide increases to about twice the normal rate, thus reducing the level of PII\(^26,27\).

Another trace element interplaying with iodine and influencing thyroid function is selenium\(^28\). Type I iodo-thyronine deiodinase, which plays a crucial role in the action of thyroid hormones, contains selenium\(^29\). Currently the synergistic effect of selenium deficiency in the development of endemic cretinism is well recognized\(^30-32\). Surprisingly perhaps, selenium supplements decrease the levels of anti-TPO autoantibodies in autoimmune thyroiditis\(^33\).

**Epidemiology**

Since iodine deficiency is the main cause of endemic goiter, the epidemiology of endemic goiter largely depends on the iodine intake of the population.

Primitive societies depend on locally produced food. The iodine content of this food in turn, depends on the iodine content of the soil and the water. Thus, if the soil and the water do not contain enough iodine, nor does the locally produced food, endemic iodine deficiency goiter appears. A notable exception must be made for populations consuming sea-food, which is rich in iodine. Iodine-poor soils are usually found in mountainous areas remote from the sea, with considerable soil erosion, mainly due to intense previous glaciation\(^34\).

In ancient times goiter was very common and was depicted in many ancient statues, including those of the Buddha, the famous Queen Cleopatra of Egypt, etc. As recently as some decades ago endemic goiter was very prevalent\(^35,36\), affecting for instance, the northwest and south-east USA, several areas of Central and
South America and notably the Andes and southern Brazil, several central European countries and notably the Alps and the Pindos mountain range in Greece, several areas of Turkey, several areas of Africa, of which Congo has been extensively studied by Belgian teams, many areas in Asia, including the Himalayas and South-East Asia, New Guinea, New Zealand, etc. In these areas a variable proportion of the population was affected by endemic goiter. In general, the prevalence of goiter (by palpation) peaked during puberty, and then decreased in the males.

In addition to exogenous goitrogens and genetic factors, the socio-economic conditions play a role in goiter formation. Najjar and Woodruff reported that rural populations and lower socio-economic classes are especially affected. Our findings and those of Brahmlatt et al. that endemic goiter is associated with evidence of generalised malnutrition support the findings of Najjar and Woodruff. It should be noted that the more expensive animal sources of food contain more iodine than the cheaper vegetable food items. Hence, poverty must be regarded as a cause of endemic goiter, as discussed later in the chapter on silent iodine prophylaxis.

The epidemiology of endemic goiter has now radically changed thanks to the supply of iodine, either deliberately as described below or through the so-called silent iodine prophylaxis. In Greece, iodine intake is now generally normal and several recent unpublished studies have confirmed the virtual eradication of endemic goiter from Greece. In most villages, the only goitrous children were those who had recently emigrated from less developed countries. Today, the predominant type of non-toxic goiter in Greece is autoimmune thyroiditis. Tsatsoulis et al. also found increased autoimmunity and urinary iodine excretion in a previous iodine-deficient area of northwestern Greece. A similar improvement in iodine intake has been reported from Brazil, the Netherlands and Bosnia-Herzegovina. The entire western hemisphere now nears iodine sufficiency.

Nevertheless, the problem still does exist. Several endemic areas persist, especially in rural areas and in developing countries. Since the situation varies from year to year, in order to draw up a map of the global incidence of endemic goiter, a worldwide collaboration of health authorities and research workers is essential. According to WHO, UNICEF and IC-CIDD, in 1998 over one third of the total world population lived in iodine deficient areas. According to Delange, in 1990 out of 5438 million people round the world, 1572 were at risk of iodine deficiency, and 655 millions (12% of the total world population) were actually goitrous, 2% were cretins and even more had some degree of mental retardation. It is not clear how many of these goitrous persons actually had a clinically significant goiter rather than a small thyroid enlargement.

In any case, it is the author’s impression that endemic iodine deficiency goiter, although far from being eradicated, is at least not so frequent or serious as in the past in the developed countries and is also gradually becoming a less serious problem in the developing countries. This, of course should not diminish our efforts to completely eradicate iodine deficiency worldwide.

HEALTH CONSEQUENCES OF ENDEMIC GOITER

The first and most obvious consequence of iodine deficiency goiter is the goiter itself. If large enough, it constitutes more than a cosmetic problem and may also cause pressure symptoms.

Furthermore, iodine deficiency may be accompanied by hypothyroidism. In severe iodine deficiency, the thyroid gland cannot compensate by the means of the mechanisms previously described. The formation of thyroid hormones is reduced and TSH levels rise. A tragic result of this may be endemic cretinism, which is more likely to occur if selenium deficiency is superimposed upon iodine deficiency, as previously discussed. In severely iodine-deficient areas, the thyroid hormones produced by pregnant women are not enough to ensure the normal development of the fetal brain. Later in gestation, the hormones, produced by the fetus are also not sufficient. The result is endemic cretinism, accompanied by neurological symptoms owing to the lack of thyroid hormones during fetal life and postnatally, if hypothyroidism persists after birth. Cretinism does not follow the rule “whole or none”. In addition to overt cretinism, there are other milder degrees of mental retardation and hearing impairment. A more extensive review of endemic cretinism is outside the scope of this paper other than the mention that endemic cretinism has been noted mainly in the Himalayas, the Andes and Congo and elsewhere, and was also present in central Europe sev-
eral centuries ago.

As stated above, iodine requirements are increased during pregnancy. In iodine deficient areas with endemic goiter, pregnancy is accompanied by increased abortions, neonatal deaths and defective progeny, with various degrees of mental retardation extending even to overt cretinism.

Another problem of endemic goiter is hyperthyroidism. In long-standing goiters autonomous nodules develop. This may lead to hyperthyroidism, especially if the iodine intake is increased, as discussed later in the context of iodine prophylaxis. This also happens spontaneously. Barker and Phillips found in 12 British cities that the incidence of hyperthyroidism was greater in areas where iodine-deficiency had been present in the past, and this high incidence was due to toxic nodular goiter. This is also in agreement with more recent papers, according to which in iodine-deficient Denmark there were more cases of Graves’ disease in the young and of hypothyroidism in the elderly. The higher prevalence of Graves’ disease and hypothyroidism in Iceland must be attributed to the increased prevalence of thyroid autoimmunity due to a high iodine intake.

Finally, carcinogenesis must be discussed. An increase in thyroid cancers in the endemic areas has been reported in the past. Now, however, the contrary has been found through fine sections of the glands: in iodine-replete areas, not only the proportion of papillary carcinomas but also the total prevalence of thyroid cancers increases. The practical conclusion may be stated as follows: in iodine deficiency, although the total number of thyroid carcinomas may be lower, the mortality from thyroid cancer is increased owing to the increased frequency of aggressive follicular and anaplastic thyroid carcinomas. On the other hand, in iodine-replete areas, although the total frequency of thyroid cancers is increased, mortality is decreased since the overwhelming majority of thyroid tumours in iodine sufficient areas are subclinical papillary micro-carcinomas.

THE TREATMENT OF ENDEMIC GOITER

The obvious advice, i.e. to supply iodine in iodine-deficient persons, may be miraculous for prevention but is only moderately effective in established goiters. According to the author’s experience, only diffuse goiters in young persons have a chance of shrinking, usually only to a limited extent. In long-standing nodular goiters, iodine is contraindicated since it may precipitate hyperthyroidism.

In these cases thyroxine administration is more effective, especially in diffuse goiters. If autonomous nodules are present, exogenous thyroxine is added to the hormones secreted by the nodules and a hyperthyroid condition state may result.

Surgery is usually required for large goiters with pressure symptoms and/or autonomous nodules. This procedure requires an experienced surgeon and a well organized hospital, and none of these is readily available in the developing countries where endemic goiter is usually present.

Instead of surgery, many now give $^{131}$I in large amounts, but again modern facilities are essential. The injection of ethanol in prominent nodules is simpler and may be used without elaborate equipment.

THE PREVENTION OF ENDEMIC GOITER

Since treatment of endemic goiter is neither easy nor very effective, prevention must be our main concern. The theoretical basis is very simple: for iodine deficiency supply iodine, and when goitrogens are present eliminate them. This is more easily said than done. Goitrogens are especially difficult to avoid. They may be contained in the staple foods on which a population survives, for instance cassava (manioc), millet, etc. It is difficult to advise their avoidance if there are not healthier alternatives available. It may also be difficult to modify the water supply in order to avoid water-borne goitrogens.

Since iodine deficiency is the main cause of endemic goiter, its eradication is the basis for prevention of endemic goiter. Iodine may be supplied in various ways. Iodine tablets or other forms of iodine to be taken on a daily or weekly basis are not very practical. The main ways to administer iodine to a population are: a) iodized salt, b) iodized oil, c) water iodination, d) others.

Iodized salt (10-100 parts of potassium iodide or iodate per million) is perhaps the most practical measure for developed countries with a modern salt industry.
Injection of iodized oil supplies enough iodine for one or more years\textsuperscript{68-70}. Iodized oil can be taken also per os. Water iodination can be used in communities with a central water supply. Furthermore, adding iodine to bread or to anything else widely consumed may be effective. All these procedures are so well known that we need not discuss them further.

In addition to this deliberate supply of iodine, there is also the so-called “silent iodine prophylaxis”, i.e. a spontaneous increase in iodine intake due to various factors. These factors, also previously discussed, include a) economic growth, allowing people to buy more expensive but also iodine-rich food, b) better communications, with the result that previously isolated areas can now also consume food produced elsewhere, and c) the industrialization of food production, which results in an increased iodine content of food, as reviewed by Koutras et al\textsuperscript{34}.

Although in Greece iodized salt was introduced following the author’s efforts, silent iodine prophylaxis is probably the main reason for the elimination of iodine deficiency\textsuperscript{42}. Nowadays, the predominant form of non-toxic goiter in Greece is autoimmune thyroiditis\textsuperscript{43}. Iodine deficiency goiter has been virtually eradicated. In some previously endemic areas which we have recently re-examined, the iodine intake, as judged by the urinary iodine excretion, has been found adequate.

In Athens, the average urinary iodine was in 1964 $45 \pm 5 \mu g/d$\textsuperscript{71}, increased to $94 \mu g/gCr$ in 1980\textsuperscript{72}, to $208 \pm 156 \mu g/gCr$ in 1992\textsuperscript{42} and was in 1999 $204.7 \pm 21.6 \mu g/gCr$ in goitrous persons with autoimmune goiter compared to $204.7 \pm 21.6$ in persons with non-autoimmune goiter\textsuperscript{43}. In many villages known for their endemic goiter, we have observed the following situation: children were more or less non-goitrous, thanks to better iodine nutrition, while their parents in many cases had an obvious and significant goiter as a result of a previous iodine deficiency.

The results of iodine prophylaxis must be monitored. Clinical examination of the thyroid is valuable but since the results appear later (established goiters usually do not regress), it is better to monitor the iodine excretion in the population\textsuperscript{73}.

**SIDE EFFECTS OF IODINE**

Iodine, whether given for prophylaxis or for treatment, has several untoward effects\textsuperscript{73}. Hyperthyroidism is not only the most important of these but perhaps also the one best studied, described as long as 180 years ago by Coindet\textsuperscript{74}. Iodine-induced hyperthyroidism is usually associated with autonomous nodular goiters, as shown in Tasmania\textsuperscript{75-77}, in Britain by Barker et al\textsuperscript{58,59} in Zaire\textsuperscript{78} and elsewhere. This subject has been reviewed by Stanbury et al\textsuperscript{79}.

Iodine-induced autoimmunity is also a problem, though less important. Our group reported the emergence of antithyroid auto-antibodies after administration of iodized oil i.m.\textsuperscript{80,81} or KI orally\textsuperscript{82}. Our results have been challenged by some authors, but fully confirmed by Kahaly et al\textsuperscript{83,84} who were the first to find lymphocytic infiltration after iodine administration. An increased prevalence of thyroid autoimmune disorders in countries with a high iodine intake has also been detected. Harach and Williams\textsuperscript{65} found in Argentina, in surgical specimens from females, lymphocytic infiltration in 8% before and 25% after iodine prophylaxis. Furthermore, iodine-induced autoimmunity is well known in animals. This subject has been reviewed by several authors, including ourselves\textsuperscript{81}. In any case, iodine-induced autoimmune is not such a problem as to deter the authorities from iodine supplementation.

Thyroid cancer was also mentioned previously. It seems that after iodine supplementation the total prevalence of thyroid cancer increases\textsuperscript{65}, but since this concern an increase in subclinical papillary thyroid cancers and is accompanied by a decrease in lethal anaplastic and follicular cancers, the over-all mortality from thyroid cancer decreases.

All in all, it seems that iodine-induced hyperthyroidism is the only serious consequence of iodine administration\textsuperscript{3,79}. This, however, is only a transient phenomenon: once iodine deficiency has been eradicated there are practically no more autonomous thyroid nodules, and therefore no iodine-induced hyperthyroidism. Iodine-induced autoimmunity may result in more cases of Graves’ and Hashimoto’s diseases, but these entities are easily controlled.

Conclusion: if iodine deficiency is present, do eradicate it!
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