News for hypothyroidism
News for hypothyroidism

II Forum of Endocrinology
27-29 September 2013, Gubbio
We would like to thank Dr. Loredana Bianchi for the editorial support during the preparation of the manuscript.
SESSION 1
THE THYROID, THE HORMONES, AND THE METABOLISM:
NEW ELEMENTS FOR THE CLINIC

13 THE PLASMATIC CONTROL OF T3 AND T4 LEVELS IN HUMANS
Domenico Salvatore

18 GENETIC DEIODINASE POLYMORPHISMS
AND CLINICAL IMPLICATIONS
Robin Peeters

23 THE INDIVIDUALLY TAILORED DOSE OF LT4:
A SIMPLE ANSWER TO MANY THERAPEUTIC PROBLEMS
Camilla Virili, Cecilia Verga Falzacappa, Marco Centanni

30 WHAT ATA-ETA GUIDELINES INDICATE ON T3/T4 TREATMENT
Luca Persani

SESSION 2
NEW HORIZONS FOR HYPOTHYROIDISM

37 THYROID HORMONE IN NORMAL AND STEM CELL
Michelina Plateroti
WHAT IS THE RATIONALE FOR A COMBINED THERAPY WITH HORMONES T3 AND T4?
Salvatore Benvenga

CLINICAL STUDIES AND BENEFITS OF THE THERAPY WITH T3
Francesco Saverio Celi

A ROLE FOR T3 IN THE “LOW T3 SYNDROME”?
Antonio Mancini, Chantal Di Segni, Sebastiano Raimondo, Giuseppe Maria Corbo, Alfredo Pontecorvi

T3 SULFATE AS A POTENTIAL PHARMACOLOGICAL AGENT IN THE TREATMENT OF HYPOTHYROIDISM
Ferruccio Santini

FUTURE TREATMENT FOR HYPOTHYROID PATIENTS
Antonio Bianco, Sabina Casula

CONCLUSIONS
The aim of the II Forum of Endocrinology organized by IBSA Foundation held in Gubbio on 27-29 September 2013 was to analyze, in the context of the therapy for hypothyroidism, the data coming from basic and clinical research with an eye to the future. During the forum, which was attended by leading experts in the field, several topics have been treated and lively debated:

- the inadequacy of TSH as the sole marker of thyroid function and the need to identify other tissue markers. Currently, TSH remains the marker used in the daily clinical practice, proving not very sensitive in a small niche of patients;
- the difficulty of matching the “well being” with the feeling of “being well” of the patient, not being possible to be guided only by patient preference in the adoption of a specific therapeutic strategy;
- another aspect of the debate has been the need to follow a diagnostic iter to highlight a possible malabsorption, in case the therapeutic target is not reached after administration of the theoretical dose of levothyroxine;
- the potentiality of research in the study of polymorphisms of genes involved in the metabolism of thyroid hormones and the possible correlation between exon sequences and TSH and FT4 values in hundreds of patients, in the clinical practice;
- the correlation between peripheral metabolism of the thyroid hormone and basal metabolism. Having more information in this topic it could help, among other things, to understand, for example, the reason why there are different metabolic responses to the same stimulus, as may be the cold, in both sexes.

We are confident that the forum offered interesting insights and can be a useful study to achieve the goal desired by all: a timely and correct diagnosis and an always more personalized and effective therapy in the future of the hypothyroid patient.
The Forum held by the IBSA Foundation in Gubbio has drawn together a number of internationally acknowledged experts to discuss current therapeutic approaches to hypothyroidism. The treatment of this condition affecting about 10% of the population living in Western countries is still lively debated by clinicians and basic researchers who strive to identify the ideal treatment for this complex condition.

It has been unmistakably shown that substitution therapy based on levothyroxine alone – the main hormone secreted by the thyroid gland that has been regarded as the cornerstone of the therapy for years – is not able to ensure well-being in a subgroup of patients. Moreover, the administration of thyroxine alone causes an excessive rise in plasma concentration of T4 in a group of patients with a limited ability to convert T4 to T3. In the light of these results, T3 might be used in combination therapy with T4. There is a long way to go, though, as various steps and goals need to be achieved before such a potent hormone can be safely used in clinical practice. How is T3 to be used? When? And for which kind of patients?

Answering these questions was one of several challenges faced by the Forum, which aimed to identify the best treatment for each patient affected by hypothyroidism, a common pathology characterized by simple etiology and diagnosis, as well as by a complex therapeutic treatment.
SESSION 1

The homeostasis of thyroid hormones in plasma and in tissues is a rather complex subject. Indeed, we know that the thyroid hormone is not able to enter into target cells in a passive manner but needs transporters. A cell that does not express the carrier is not permissive to the action of thyroid hormone. Entered into the cell, the hormone becomes the target of a family of enzymes, the selenium-deiodinase, that, in some way, can completely change its biological activity.

The deiodinase are enzymes that convert thyroxine, the main thyroid product, in a molecule metabolically active, T3, or in an inactive product, the rT3 (Figure 1). The fate of T4, therefore, varies depending on the pathway of activation or degradation in which is directed, so not necessarily the pre-hormone exerts its hormone activity, but it can also be turned off and, as such, be unable to reach the nuclear targets.

The structure of deiodinase is complex. These enzymes have a membrane domain whose function is not yet fully known, and it is probably necessary so that the molecules dimerize and become enzymatically active. The most surprising thing, though, is the presence in these molecules of an atom of selenium, a trace element that, in order to be incorporated into the protein, requires a considerable amount of time and energy by the cell. However, this process is essential because the selenium is localized in the catalytic site of the enzyme, site that T4 must reach in order to be metabolized. Selenium is preserved phylogenetically in all species and is common to all deiodinases. Also, in laboratory there was a significant reduction in the deiodination activity following the replacement of this oligo element with a sulfur atom in the catalytic site.

Now imagine that there is a cell deiodinase-free: in this cell T3, produced in small part by the thyroid, enters and performs its function without any control at intracellular level. If, instead, in the cell there is predominance of one enzyme on the other, we obtain different effects. In particular, when the deiodinase type 2 (D2) predominates
there is an increased production of T3 with consequent increase of receptor saturation. The opposite situation occurs if deiodinase type 3 predominates (D3), which inactivates the thyroid hormone. All this suggests that the target cell is able to control the response to thyroid hormone, or rather, its “thyroid status” regardless of what is the plasmatic level of the hormone. The plasma concentration is controlled by a powerful homeostatic system, the hypothalamus-pituitary axis, and exerts a steady-state action, namely, balancing action, compared to what, instead, happens inside single cells that are able to modulate the signaling.

The perhaps most striking example of deiodinase power is observed in brown adipose tissue responsible for thermogenesis. The production of heat due to adrenergic stimulus can only occur if the D2 is up-regulated, with increase of T3 concentration that saturates almost completely the receptor and determines the synthesis of uncoupling protein and, therefore, the production of heat. This shows how the control system operated by the deiodinase is essential for the functioning of the same tissue.

The hormones secreted by the thyroid T4 and T3 are present with a ratio of 15:1 within the colloid, while, after intra-thyroidal conversion, the ratio becomes 13:1 in the circulating plasma. But how is it regulated the synthesis of T3 in the plasma?
Understanding is crucial when you consider that in patients with thyroidectomy the T4 administered is converted in T3 ensuring the proportion of hormone needed to tissues. The enzymes responsible for the control of the synthesis of T3 in the periphery are substantially D1 and D2, with some differences. In order to determine the role of D1 compared to D2 it was used propyl-thiouracil (PTU), which selectively blocks the D1. It was found that in patients treated with PTU in euthyroid condition the T3 concentration is only reduced by 20-30%, compared to a reduction of about 50% under conditions of hyperthyroidism. This indicates that D1 predominates in hyperthyroidism, and this is the reason why the PTU is very effective in reducing the levels of thyroid hormone in hyperthyroid patients.

To recap: about 20% of the hormone synthesis of T3 is guaranteed by the thyroid, but there is also an extra-thyroid production of T3 regulated by the activity of D1 and D2. These two activating enzymes, however, have a different pattern of expression in the various tissues of the body, as well as a different degree of affinity for the T4 (● Table 1).

Another important aspect concerns the homeostatic role of deiodinase enzymes, well described from the regulatory mechanism of D2: in hypothyroidism there is an increased activity of this enzyme, with consequent increase of the conversion of T4 to T3; an opposite regulation occurs in conditions of hyperthyroidism.

It emerges, at this point, a new important question: the exogenous T4 is itself able to ensure proper concentration of T3 both at plasma and tissue level? At this regard, a study by the Catania group [2] on more than 1,800 patients without thyroid has demonstrated how, in these subjects, T3 levels were on average lower compared with control patients. The research shows that in patients treated with T4 the T3 levels, even if within the normal range, are slightly lower than the norm. Furthermore, it

● **Table 1.** The two activation pathways of T4 in T3

<table>
<thead>
<tr>
<th></th>
<th>D2</th>
<th>D1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Km(T4)</td>
<td>10⁻⁹M</td>
<td>10⁻⁴M</td>
</tr>
<tr>
<td>Tissues</td>
<td>Muscle, skin</td>
<td>Liver, kidney</td>
</tr>
<tr>
<td>Cellular location</td>
<td>End. Retic.</td>
<td>Plasma membrane</td>
</tr>
<tr>
<td>Enzyme half-life</td>
<td>1 h</td>
<td>12 h</td>
</tr>
<tr>
<td>PTU effect</td>
<td>0</td>
<td>Inhibit</td>
</tr>
<tr>
<td>Hypothyroid, iodine deficiency</td>
<td>↑</td>
<td>↓</td>
</tr>
<tr>
<td>Hyperthyroid</td>
<td>↓</td>
<td>↑</td>
</tr>
</tbody>
</table>

Source: data from Bianco et al., 2001 [1].
was found that in a small percentage of patients (approximately 15%) T3 does not normalize perfectly, even though TSH values are normal. In these cases, in order to normalize the T3 levels, it is necessary to increase the T4 dosage so much to reduce the TSH to values not more physiological. But what is the role of deiodinase enzymes in this context? To respond it is crucial to distinguish the plasma control of the hormone levels from the tissue one. In this regard an important information is provided to us from an animal model deprived of D2, in which, despite normal levels of T3 in plasma, the concentration of T3 in some tissues such as the brain, is reduced. The same occurs in mice in which both deiodinase activators are genetically deleted: the plasma T3 is normal, but TSH and T4 levels are high. In this case, the thyroid provides all the T3 necessary ensure a normal plasma concentration of T3.

From the described animal models it can be deduced that in patients treated with T4, the plasma values of T3 in the normal range does not necessarily reflect a euthyroid condition at tissue level.

Although there are still some aspects to be clarified, it can be concluded that, probably, through treatment with T4 alone you can get the normalization of T3 concentration in plasma but not in tissue.

In humans, the thyroid hormone is essential for proper development of the organism in intrauterine and neonatal life as well as in the adulthood one. A demonstration of the importance of deiodinase enzymes for life comes again from the animal world. Metamorphosis is an event regulated by thyroid hormone: in the absence of deiodination activity, this process is altered. Moreover, the thyroid hormone is essential for the proper maturation of the cochlea, a structure of the inner ear [3]: in the first days of life there is a peak of D2 which leads to an increase in T3 concentration of almost 100 times. This is necessary so that the ear can develop in a proper way, so much that mice deprived of the D2 activity are deaf.

What happens, though, if the balance guaranteed by deiodinase enzymes is lost? That’s what occurs in consumptive hypothyroidism, a frequent paraneoplastic syndrome in children, characterized by a pathological increased expression of D3. An article appeared in the *New England Journal of Medicine* [4] described the case of a child suffering by hepatic hemangioma who presented a severe hypothyroidism. The hemangioma is a highly vascularized tumor that may have a highly variable clinical presentation: typically begins with a phase of expansive growth for a period of about a year, to which follows a phase of regression. The growth phase is the most dangerous from a metabolic point of view, since cancer cells express very high amount of D3, resulting in massive degradation of T3 and T4 normally produced by the thyroid. The activation of the hypothalamic-pituitary-thyroid feedback in these children is not sufficient to buffer the hepatic hormone degradation and to ensure euthyroidism, and, therefore, occurs a particularly severe hypothyroidism, which in some cases, can lead to death. Several studies have shown that consumptive hypothyroidism can also affect adulthood and may be associated also with other types of cancer.
In conclusion, we can say that the deiodinase type 1 is involved in the plasmatic homeostasis of T₃ in the same way as the deiodinase type 2. The latter is also crucial for the intracellular homeostasis of T₃ and for the initiation of the feedback operated by T₄ on TSH secretion. The deiodinase type 3 plays a key role during intrauterine life and in different pathophysiological mechanisms, such as pregnancy.

Finally, deiodinase enzymes as homeostatic circuit presiding the control of plasma T₃, meet the functional requirements of the different tissues and are able, to some extent, to protect tissues from modest changes in plasmatic T₃.

References


Genetic plays an essential role in control of thyroid hormone’s set-point. The importance of hereditariness in the regulation of TSH and FT$_4$ is well described in the study of Hansen et al. [1], in which there is a significant correlation between TSH and FT$_4$ values in dizygotic twins, and, much more, in monozygotic twins. Complicated calculations have showed that in the last ones about 60% of total TSH variability is related to genetic factors.

In another work [2], a group of healthy voluntaries was monitored during the year through monthly measurements of TSH and FT$_4$: for every subject hormones levels fluctuated around a particular set-point, that was different from set-point of the other subjects. The normal range of TSH is calculated on the basis of variability of values in general population, but as we can see in • Figure 1, this may not be optimal for each individual. So, it may be that a TSH value in the normal range can be still too high for a particular subject. The same holds true for FT$_4$.

The • Figure 2 [3] shows the relation between TSH and FT$_4$: as we can see, small changes in FT$_4$ levels correspond to logarithmic alterations of TSH: high values of FT$_4$ are associated to low values of TSH and vice versa. In general, there is a good correlation between FT$_4$ and TSH, but a lot of values are scattered. For example, if we consider the two subjects marked in • Figure 2, we can see that both of them have had a similar decrease in FT$_4$, leading to a similar increase in TSH; but, while the one with FT$_4$ under the normal range is now considered hypothyroid, for the other one the plasmatic concentration of the hormone is still normal, and he or she is therefore considered subclinically hypothyroid. This demonstrates the importance of individual variation of hormonal set-point.

Another study [4] shows the possible contribution of genetic factors in determining the clinical set-point: in patients with syndrome of resistance to thyroid hormones cau-
sed by mutations in the THR-β gene, the reduced pituitary’s sensitivity to the negative feedback of thyroid hormones causes high levels of FT$_4$ and FT$_3$; there is still a log-linear relationship between TSH and FT$_4$, but the set-point has shifted.

About 10 years ago, we analyzed polymorphisms of different genes involved in thyroid pathway [5], including genes for deiodinases, and we showed the existence of two particular polymorphisms in a region of the gene for type 1 deiodinase. We know that this enzyme is responsible for plasmatic FT$_3$ production and for clearance of rT$_3$ (reverse T$_3$). So we speculated that a defective deiodinase would, somehow or other, alter hormone’s homeostasis. In 156 healthy voluntaries, asymptomatic carriers of the polymorphism had lower levels of T$_3$, while rT$_3$ was high.

Moreover, because of the enormous increase in technological possibilities associated to genetic studies in the last 10 years, we have switched from analyzing single gene variants to whole genomes, through the so called GWAS (Genoma Wide Association Studies). In these kind of studies we analyze about 2.5 million polymorphisms in the genome at the same time.

In one of these works [6] we identified a lot of functional variants associated to TSH and FT$_4$ and we saw that the most significant variant for FT$_4$ was the variant in type 1 deiodinase, that we had already analyzed 10 years before (● Figure 3). By using these large consortia we have identified 23 independent polymorphisms associated to TSH, and these contributed not only to TSH variation within the normal range, but also to TSH values outside the reference range. However, these findings explained only 5.64% of the trait variation in TSH, and 2.30% of total variation in FT$_4$.

Of course, the most important question concerns consequences of these findings in clinical practice. There are three reasons why it is important to study this kind of polymorphisms. First of all, we are able to obtain novel insights into mechanisms in-
involved in the development of the diseases of interest, also for identification of new drug targets. A study from Butler et al. (2010) [7] analyzed functional effects of a particular polymorphism in the type 2 deiodinase (Thr92Ala), showing that subjects who with the rare variant of this polymorphism had an altered sensitivity of the pituitary in the regulation of hypothalamic-pituitary-thyroid axis. When these subjects were injected with TRH they showed a different response in increase of T3 in comparison with healthy subjects, also suggesting that this polymorphism is indeed relevant for the control of the hypothalamic-pituitary-thyroid axis and for the regulation of thyroid hormone’s set-point.

These studies of polymorphisms associated with the HPT-axis set-point may also help for diagnosis: if we are able to predict disease risk, consequently it may help us to personalize medicine, and to decide whether or not we should treat patients. But, actually, we can explain only a little percentage of TSH variations so far, and so we can’t yet develop reliable tests for that particular patients. In other words, we are not already able to predict individual risk of disease.

At last, these studies could help us to predict the response to treatment. Panicker et al. (2009) [8] analyzed relationship between different polymorphisms, including polymorphisms of type 1, 2 and 3 deiodinase, and well-being in hypothyroid patients on thyroxine therapy. In particular we know that type 2 deiodinase is the major responsible for local production of T3, so it represents the most important resource of active

---

**Figure 2.** Log-linear relationship between TSH and FT4

![Log-linear relationship between TSH and FT4](source: Spencer et al., 1990 [3], adapted.)
hormone in the brain. So, if we speculate that hypothyroid patients depend almost completely on local conversion of T₄ to T₃ by DIO₂, less functional variants of this enzyme would be associated to a reduced pool of T₃ into the brain, and, consequently, to decreased well-being of patient. The authors demonstrated that this variant was indeed associated with well-being in patient on LT₄ therapy. Moreover, the authors demonstrated that patients with this DIO₂ polymorphism may have more benefit from T₄+T₃ combination therapy.

However, there are two important remarks here: first of all, a lot of tests were performed, and if we apply strict multiple testing corrections, the results are not considered significant. This remark is also made by the authors in this paper. Second, it’s a small study, so we need confirmation from independent studies showing the same results.

In conclusion, although the current evidence suggests that variants in deiodinase genes may explain why some patients continue to have complaints under LT₄ therapy, we have not yet sufficient knowledge to predict response to treatment for the individual patient.
References


Replacement therapy with sodium levothyroxine involves the use of a hormone with a narrow therapeutic index, usually framed in a treatment of long/very long period. In the past, on empirical grounds, there has been a substantial anarchy of doses and routes of administration, with significant side effects due to a chronic hyper-or hypo-treatment.

On the basis of these considerations it is evident the importance of using an individualized dose of thyroxine: first because the minimum effective dose of T\textsubscript{4} should be administered, but also to avoid continuous oscillations of the dosage and consequent loss of compliance by the patient. Moreover, customizing the dose allows suspecting possible malabsorption, identifying gastrointestinal disorders. Further advantage of this therapeutic modality regards the ability to obtain substantial savings in terms of pharmaco-economics, avoiding lengthy and costly monitoring. There are however several variables to consider to find an optimal and predictable daily dose of thyroxine (\textbullet Figure 1).

The hormonal need changes in the absence of the thyroid gland and several studies have shown that the dose of thyroxine must be incremented after thyroidectomy. Crucial element when the thyroid gland is still in situ is a correct assumption of the therapy, therefore, an effective compliance by the patient. The dose of LT\textsubscript{4} should be calculated according to the BMI (Body Mass Index) or the amount of lean body mass of the subject [1]. The administration also can occur at any time of the day, denying the belief that levothyroxine must necessarily be taken in the morning [2, 3]. The opportunity to change and to “tailor” the time of assumption guarantees better compliance by the patient and the possibility of a greater intestinal absorption of the hormone. Studies from Salvatore Benvenga have determined that the initial rate of absorption of thyroxine occurs in the first 60-90 minutes after assumption. The in-
The interval of time necessary to avoid interference with any type of foods and beverages, including coffee, must therefore be of at least 1 hour [4]. To properly assess the effect of therapy with thyroxine it is also crucial having a good marker of adequacy. The reliability of TSH as a marker of the thyroid homeostasis is widely accepted, although there are doubts both quantitative, relative to the reference range [5], and qualitative, about the influence that the tissue-specific metabolism of the hormone exerts on the effectiveness of oral thyroxine [6]. To compromise the reliability of TSH, however, contribute other factors, often not considered (Figure 2), which interfere with the dosage of this marker [7]. In particular, some substances contained in cosmetics that falsely lower TSH, affecting the understanding of the real thyroid situation. Therefore, the need to re-evaluate the effectiveness of this marker as isolated predictor of the adequacy of thyroxine therapy, associating it always to the dosage of iodothyronines.

To obtain individually tailored dose, it is also necessary to exclude the factors that can cause an increase in the daily intake of the hormone: a) changes in anthropometric characteristics of the patient (weight, gestational status); b) different biochemical properties of the pharmaceutical preparation of LT₄ (bioequivalence); c) nutritional interference [8]; d) concomitant medications (poly-treatment).

An increased need for levothyroxine dose than the theoretical dose, which persists despite the exclusion of all these conditions, suggests the presence of a gastrointestinal malabsorption of the hormone. In general, it was found that can exist an increased need for T₄ in the course of gastric diseases (active infections and chronic gastritis from *Helicobacter pylori*, atrophic chronic gastritis and gastrointestinal resections) or intestinal diseases (coeliac disease, lactose intolerance, etc.) [8-10].

Among the putative mechanisms of increased requirements which is observed in the course of gastric pathologies we remember essentially the possible consequences of increased gastric pH: on one hand, the changes in the ionization state of the thyroxine molecule that may impact on the ability to cross the membrane of enterocytes,
on the other hand the variability of the degree of dissolution of the pharmaceutical preparation. In this respect, a study of 2009 from Pabla et al. compares the ability of dissolution of pharmaceutical preparations of thyroxine in tablets and in softgel to increasing pH values in the medium [11]. Among the putative mechanisms of malabsorption of thyroxine in intestinal diseases are mainly counted the following: a) the reduction of the absorptive surface for shortening or destruction of intestinal villi or surgical removal; b) alterations of intestinal motility; c) permanence of undigested material capable of adsorbing thyroxine within the intestinal lumen; d) qualitative and quantitative variations of mucus that lines the monolayer of intestinal columnar cells; e) parasites [12].

An essential requirement for any study aimed to identify malabsorption is the presence of a reference standard. Taking into account all the variables mentioned above, we formed a group of patients who had certainly respected all the criteria described in order to establish a targeted therapy and, on this basis, we derived the median dose of levothyroxine that allowed to reach target values of TSH: 1.31 µg/kg/day vs. 1.09 µg/kg/day in replacement mode and 1.56 µg/kg/day vs. 1.34 µg/kg/day in semi-suppressive mode respectively, in adults and elderly. By administering these ideals doses in our sample of patients during almost twenty years we have observed that about 20% of them did not reach the therapeutic target while 80% responded biochemically as expected. After the exclusion of patients with poor compliance, drug interference, or incomplete screening, the remaining patients, defined poor responders, accounted for about 12% of the sample. In such patients, the need for thyroxine showed to be increased of approximately 30% both in adults and in elderly (>60 years) (● Figure 3).

This group of patients with increased requirement of T4 was subjected to a well-defined diagnostic pathway (● Figure 4), which aims to investigate the cause of ma-

---

**Figure 2.** Factors interfering with the interpretation of serum TSH

- TSH-secreting pituitary adenomas
- Asymptomatic hyperthyrotropinemia
- Drugs (antidopaminergic, anti-serotonergic)
- Interference on the dosage (macro-thyrotropin, heterophile antibodies, rheumatoid factor)
- First trimester of pregnancy
- Marginal Malnutrition
- Central Hypothyroidism
- Drugs (neuroleptics, glucocorticoids)
- Nutritional supplements, cosmetics and hair colorants
- Not thyroidal illnesses

Source: data extrapolated from Centanni et al., 2007 [7].
absorption of the drug. On this basis we have identified, in most cases, the presence of gastric diseases: atrophic gastritis, Helicobacter pylori infection, chronic gastritis Hp-related. In the remaining patients the malabsorption was secondary to intestinal diseases, in particular to lactose intolerance and to coeliac disease\textsuperscript{13}. Unlike intestinal malabsorption, reversible in many cases with specific diets, the gastric malabsorption are difficult to reverse because dependent on outcomes of chronic inflammation. There are, however, in the market new drug formulations that may be useful to resolve these forms of malabsorption. Among these must be counted the liquid preparations (drops or mono-dose) and those in softgel. In this regard, our preliminary data of a study with the preparation of T\textsubscript{4} in softgel on patients with gastric malabsorption, histologically defined, have suggested the possibility of reducing the dose of thyroxine using this new pharmaceutical preparation.

It can therefore be concluded that the dose of thyroxine should be individualized to reach the therapeutic target at the minimum necessary dose and that the T\textsubscript{4} therapeutic efficacy can be affected by numerous biases. The adoption of individually tailored dose as therapeutic standard allows determining the actual needs of T\textsubscript{4} and identifying, therefore, patients with T\textsubscript{4} malabsorption. Thyroxin malabsorption oc-

**Figure 3.** Thyroxine need in patients with malabsorption

![Graph showing thyroxine need in patients with malabsorption](image-url)

Source: Centanni et al., 2007\textsuperscript{7}.
**Figure 4.** Diagnostic iter for the principal gastrointestinal diseases

Source: data extrapolated by Centanni et al., 2007 [13].
curs in a patient treated out of 6 and the cause recognizes a gastric pathogenesis in two thirds of patients. Knowledge of the thyroxine pharmacokinetics enables the development of more absorbable preparations and potentially able to optimize the thyroxine therapy, adapting it to the characteristics and the needs of the individual patient.

References


Replacement therapy with levothyroxine is, at the present, the preferential therapeutic approach for hypothyroidism. The guidelines of the ATA-AACE (American Thyroid Association-American Association of Clinical Endocrinologists) [1] and those of ETA (European Thyroid Association) [2] strongly recommend monotherapy with T4, since there is not yet enough evidence in favor of combined treatment T3+T4.

It is known, however, that 5-10% of patients treated with levothyroxine, while achieving euthyroidism in plasma, continues to complain of symptoms of hormonal deficiency. The explanations for this phenomenon may be different. For example, the awareness of being affected by a chronic disease could in some way, affect the feeling of well-being reported by the patient; in addition, several conditions related to thyroid disease or condition of the replacement, may compromise the achievement of a state of well-being. We know, in this regard, that often the autoimmune thyroid disease is associated with other diseases (diabetes mellitus, Addison’s disease, early menopause, etc.) in the so-called polyglandular autoimmune syndromes (PGAS). In these cases it is necessary to take into account that the patients complaints attributed to hypothyroidism may actually have a different origin from the thyroid.

At the base of discomfort reported by the patient, theoretically euthyroid, there could be an inadequacy of the therapy itself: several studies reviewed in the guidelines have shown that monotherapy with LT4 may be insufficient to ensure a proper plasma relationship of FT4/FT3 and, in contrast, a combination therapy T3+T4 has proved capable, in some cases, to lower this ratio to more physiological values (Table 1). An inadequate response to treatment with levothyroxine may also depend on individual variables, such as the existence of deiodinase polymorphisms [2].

If, then, for that famous 5-10% of patients in therapy, symptomatic despite normal hormone levels in plasma, we exclude interfering factors previously described, there may be a rationale for the use of combination therapy and get more clinical benefits.
In this regard, the study of Panicker et al. [3] analyzed the correlation between diodi-
nase polymorphisms and feeling of well-being reported by the patient, demonstrating
that a particular polymorphism for D2, present in approximately 16% of subjects tre-
ated with levothyroxine, was associated with a worse feeling of sickness and a better
response to combined treatment.

The ETA guidelines [2] emphasize the concept, stating that these data on polymor-
phisms involved in the thyroid hormone pathway, especially for genes of transporters
and desiodinase, could play a role in order to justify the use combined T3+T4 the-
rapy. Several crossover studies show that, generally, patients tend to prefer combina-
tion therapy to monotherapy with LT4 [2] (• Table 2). The work of Appelhof et al.
[4] (• Figure 1) confirms this trend: the subgroups of patients undergoing therapy
T3+T4 in various combinations defined themselves more satisfied compared to pa-
tients with levothyroxine monotherapy. Various aspects make, however, questionable
this study: short duration, as well as a condition of thyrotoxicosis associated with the
combination therapy, both at laboratory (suppressed TSH) and clinical level, with a
significant increase in heart rate and decrease of body weight. It is therefore necessary
to clarify whether the feeling of satisfaction reported by patients is, in fact, the result
of an overtreatment and not more of a replacement therapy aimed at restoring the
euthyroid state.

The limits of the comparison studies between the two treatment modalities are
different: patient samples of small size, lack of homogeneity of the hypothyroid state
of the populations in exam (post-thyroidectomy, primary hypothyroidism...); use of
various associations of T3 and T4.

ETA guidelines suggest that the combination therapy may constitute an experi-
mental approach in hypothyroid patients that, despite the normalization of bioche-
Both the A

Table 2. Preference of patients receiving replacement therapy with T4+T3 or T4 derived from controlled crossover or parallel studies

<table>
<thead>
<tr>
<th>Author</th>
<th>n</th>
<th>Preference T4 monotherapy</th>
<th>Preference none</th>
<th>Preference T4+T3 therapy</th>
<th>p value</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Crossover studies</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Walsh et al. J Clin Endocrinol Metab 2003;88:4543-50</td>
<td>100</td>
<td>46</td>
<td>18</td>
<td>36</td>
<td>0.32</td>
</tr>
<tr>
<td>Bunevicius et al. Endocrine 2002;18:129-33</td>
<td>10</td>
<td>2</td>
<td>2</td>
<td>6</td>
<td>-</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td>228 (100%)</td>
<td>61 (27%)</td>
<td>58 (25%)</td>
<td>109 (48%)</td>
<td></td>
</tr>
<tr>
<td><strong>Parallel study</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Appelhof et al. J Clin Endocrinol Metab 2005;90:2666-74</td>
<td>140</td>
<td>14/48 (29%)</td>
<td>43/92 (47%)</td>
<td>0.024</td>
<td></td>
</tr>
</tbody>
</table>

* Including 6 patients who preferred the combination of 87.5 μg L-T4 + 7.5 μg L-T3 for a final 8-week addon regimen that was not randomized.

Source: Wlersinga et al., 2012 [2].

mical parameters, continue to complain about the clinical disorders, and for which have been excluded all interfering factors on the well-being above described. Therapy should be discontinued in absence of clinical improvement after three months of treatment [2].

Both the ATA and ETA do not recommend this type of therapy in pregnant patients and in patients at risk of cardiac arrhythmias.

At this point emerges the need to calculate the proper dosage of the two hormones for combination therapy. The ETA guidelines [2] suggest a relationship L\textsubscript{T}4/LT\textsubscript{3} between 13:1 and 20:1 according to body weight. Since the currently available formulations contain a L\textsubscript{T}4/LT\textsubscript{3} ratio of less than 13:1, it is preferable to use separate tablets of the two hormones. The dose of T\textsubscript{4} should be administered only once per day, while the T\textsubscript{3} should be divided into a lower dose taken in the morning together with T\textsubscript{4}, and a higher dose administered in the evening on an empty stomach 1-2 hours after
dinner, before bedtime. This mode of administration is based on the results of a study appeared on JCEM in 2008 [5]: it is known that TSH has a circadian rhythm with an acrophase with between 2 and 3 a.m., while the FT4 will remain virtually stable during the day. This study has shown that even the FT3 has an acrophase that occurs about half an hour after the peak of TSH.

The ETA still recommends to monitor the combination therapy with thyroid function test on blood samples taken before administration of the drugs, with the aim of restoring normal plasma values of TSH, FT3 and FT4, but also an adequate ratio T4/T3. If there is need of dose adjustment of the two hormones, it is preferable to start from T3, under the supervision of an experienced endocrinologist.

The ETA suggestions for future research include prospective studies in patients initiated on monotherapy with T4, comparing the characteristics at baseline between those who declare themselves satisfied with the treatment or not; trial on large numbers of patients to identify the most appropriate dose to reproduce the normal FT4/FT3 ratio in the blood; randomized clinical trials comparing the two therapeutic modes in hypothyroid subjects recognized to be carriers of polymorphisms of deiodinase and transporters; studies with slow-release T3 preparations (e.g. T3 sulfate), prospective studies to evaluate the long-term effects and the safety of combined therapy [2].

**Figure 1.** Percentage distribution of the subjective preferences of the patients in the different regimens used in the in parallel study of Appelhof et al.

Source: Appelhof et al. 2005 [4].
References


SESSION 2

NEW HORIZONS FOR HYPOTHYROIDISM
This paper provides an overview on current knowledge regarding the stem cell, with particular reference to the potential future therapeutic applications; at the same time it will describe the ways in which thyroid hormones influence the so-called “stem cell”.

Two properties define the stem cell: the ability to self-renew, i.e. to generate a cell identical to itself and the ability to generate differentiated cells. There are two types of stem cells. Embryonic cells can be recovered in large quantities from the blastocyst and once cultured, they maintain the ability to self-renewovate. They are defined pluripotent, as they are capable of giving rise, if properly stimulated, to any type of differentiated cell. Cells are easy to handle in vitro, but their use requires a number of bioethical issues. The situation is different for somatic stem cells who, instead, are present in tissues in relatively small quantities and are characterized by multipotency, i.e. they have a limited differentiation capacity, relative to the tissue of origin (\textit{Table 1}).

Precondition for the stem cell to be that is the presence of the so-called “niche”, a collection of cells and proteins that surround the stem cell itself and form a conducive environment so that it may divide symmetrically or asymmetrically, depending on the case and in function of the stimuli to which is subjected.

The current interest in stem cells comes in part from the desire to give answers to questions related to their physiology, within fundamental research. On the other hand, stem cells represent an attractive target in the scenario of regenerative medicine because they have the potential to differentiate into any tissue of the body.

In this regard, it is interesting the work done in the laboratory of Professor Costagliola in Brussels [1], who attempted to recreate the thyroid follicles from undifferentiated stem cellular. For the development of the thyroid, it is essential the co-expression of two transcription factors: NKX 2.1 and Pax8. If in vitro stem cells are properly stimulated so that express both the above factors, the differentiation in thyroid folli-
cicular cells occurs. If this in vitro system is explanted under the kidney capsule of mice, there is an increase in the levels of circulating thyroid hormones, with the possibility of re-establishment a condition of euthyroid in animals previously subjected to experimental hypothyroidism (Figure 1).

However, it is also interesting to understand whether and how thyroid hormones influence the stem cell. Different tissues respond to the thyroid stimulus by modulating the activity of the stem cells. For example, at the level of the central nervous system, the subventricular zone contains stem cells of astrocytic type, from which originate proliferating progenitors that migrate and differentiate into neurons of the olfactory bulb. In this context, it is crucial for the maintenance of stem cell the expression of the transcription factor Sox-2. The thyroid hormones (as well as the nuclear receptor TRα1) inhibit this process and the stem cell differentiates into neural progenitors [2].

Another example emerges from our work on the intestinal crypt [3], on whose bottom different types of stem cells have been described. Our previous studies had demonstrated that the nuclear receptor TRα1 has a maximum expression exactly towards the bottom of the crypts [4].

For subsequent studies we took as a reference the phenomenon of metamorphosis in amphibian, in which occurs a remodeling of the intestinal epithelium after a significant increase in the plasmatic concentration of thyroid hormones. Even in mice during the weaning phase, between the second and third week of life, there is an increase in the release of hormones, necessary for the intestine adaptation to a different diet rather than milk [5]. In particular, the intestinal adult epithelium, starting from a simple tubular form, acquires a more complex structure, even with the appearance of the crypts that contain progenitor and stem cells. Then, on the basis of these observations, through the use of knock-out mice for genes TRα and TRβ, we concluded that the receptor TRα1 controls the development, but also the continuous remodeling of the intestinal adult epithelium, modulating the activity of the cells of the intestinal crypts (stem and progenitors).

This is achieved through the direct or indirect control by the receptor of the expression of a whole series of genes functionally related to each other, belonging to diffe-

| **Table 1. Main differences between embryo and adult stem cells** |
|---------------------------------|-----------------|---------------------------------|
| **Adult tissues**               | **ES**          |
| Multipotency                    | Pluripotency    |
| Small number                    | High number     |
| Hard to maintain and differentiate in vitro | Easy to maintain and differentiate in vitro |
| Limited/no problems of bioethics constraints | Bioethics problems |
rent pathways of signaling of the cell (Wnt, Notch, c-Fos) involved in the regulation of proliferation and differentiation of the intestinal crypts cells (● Figure 2).

Subsequently, through other studies we have made it clear that the action of thyroid hormone involves in particular the stem cells of the crypts and not just the progenitors, and that TRα1 receptor expression is also essential for the regeneration of the epithelium subsequent to intestinal damage; in knock-out mice for the gene TRα1, in fact, the intestinal epithelium was incapable of dealing with the damage caused by massive doses of radiation \[6\].

The receptor TRα1 also has an important action in vitro on the growth of organoids cell, inducing a progressive increase in complexity through a direct monitoring of the activity of stem cells. In fact, in absence of receptor activity there is a considerable delay in this development process \[7\].

Our future goals are aimed at a better understanding of the described phenomena, possibly recreating the same conditions in animals of different genotype. It will also be interesting to evaluate the effects, on the contrary, of an over-repression of TRα1 receptor on the same processes.

In conclusion, there are several demonstrations, both in vivo and in vitro, of the important role of thyroid hormone and TRα1 receptor in controlling the activity of
proliferation and differentiation of stem cells. On the other hand, several studies open new scenarios on the possible use of these cells in regenerative medicine, leading the way towards new therapeutic approaches, for example in the field of neurodegenerative diseases.

References


The issue of combination therapy T₃+T₄ is quite controversial, as shown by recent reviews [1-3]. The important work of Gullo et al. [4] shows that the only therapy with levothyroxine is not able to guarantee euthyroidism in all patients without thyroid: in fact, approximately 15% of these patients supplemented with LT₄ alone (but about 17% in the age group over 70 years) had a FT₃ lower than in euthyroid subjects (used as normal controls). In addition, values of FT₃/FT₄ ratio relatively lower of the control group were present in approximately 30% of patients without thyroid treated with single LT₄.

In a study conducted in Leiden, the Netherlands, [5] were enrolled about 600 patients 85 years old, 85% of which were euthyroid. Patients were followed for 4 years, monitoring the values of FT₃, FT₄ and TSH. Already in the process of recruitment was observed a direct and significant correlation between serum levels of type FT₃ and well-being reported from the patient. In particular, to values relatively higher of hormone corresponded a better quality of life, lower incidence of depression and a better global cognitive ability. Similar results were observed during follow-up. In addition, during follow-up, the correlation with mortality could be assessed. It was seen that the FT₃ serum levels were inversely correlated with mortality: the lower was the serum FT₃, the higher the mortality from all causes. Even the FT₃/FT₄ relationship proved to be predictive of a better outcome, being it 0.25 in the fourth year survivors vs. 0.23 in died patients. An exactly opposite correlation was observed with FT₄ levels at enrollment and during follow-up.

Although monotherapy with LT₄ is considered the recommended one for hypothyroidism [3], about 10% of hypothyroid patients continue to complain of residual hypothyroidism disorders despite adequate dosage of LT₄ [1]. The important study of Panicker et al. [6] reveal that only the group of patients with a particular poly-
morphism of deiodinase type 2 (i.e. carriers of the polymorphism genotype CC of rs225014) benefited from combined therapy LT3+LT4. These carriers, already at the start (pre-treatment) had the worst values of psychological well-being than the carriers of the other two polymorphic variants (genotype TT and genotype TC).

In the study by De Jong et al. [7] data emerge that should make us reflect: in the period between 2005 and 2011 there has been an increase of percentage of cases in thyroid replacement therapy among the Dutch population. In particular, there has been a 53% increase of patients in monotherapy with levothyroxine, but surprising data are related to the 67% increase of the cases treated with T3+T4 combination therapy and 36% with T3 monotherapy. Another trend that emerges from this study regards the increase of the prescriptions, for all the above types of treatment, made by general practitioners, compared to a relative decrease of prescriptions by specialists.

A situation predisposing to the fact that different hypothyroid patients do not respond to the same dose of LT4, becoming euthyroid, and that residuals hypothyroid disorders may persist (variable from patient to patient) is that, given the systemic action of thyroid hormones, the clinical picture of hypothyroidism is also systemic. Furthermore, the thyroid hormonal signal is extremely complex and not comparable with other hormonal signals. Indeed, in addition to the fact that there are three deiodinase enzymes in charge of activation/inactivation of the thyroid hormone signal, there are numerous plasmatic transport proteins for thyroid hormones, there are many transporters on the plasma membrane to facilitate the entry/exit of thyroid hormones and numerous are also the isoforms of nuclear receptors of thyroid hormones (and different peripheral tissues have a peculiar distribution of these isoforms). And yet, over the main action (the one mediated by nuclear receptors, and known as genomic action), there is one that is not mediated by nuclear receptors (the so-called non-genomic action), with signaling pathways common between T4 and T3 and other routes specific for one or the other hormone. The field of the combined therapy T3+T4 is rather controversial and reflects the complexity of the thyroid hormone pathway in the body. The situation is made more complicated by the existence of a wide range of nuclear receptors for thyroid hormones which have different degrees of expression in different tissues as well as a variable affinity for the hormones themselves. This explains why a certain amount of T3 in the different cells of the organism does not involve the same degree of hyperthyroidism.

To illustrate the concept of non-correspondence between plasma levels and tissue concentrations of thyroid hormone comes in handy a study on polymorphisms of transporters [8], which shows that, in spite of the reduction in intracellular hormone uptake favored by some polymorphic variants compared to the wild type transporter, there was no significant change in circulating hormone concentrations. The work of Liao et al. [9], carried out on different types of knock-out, even triple, for the main carrier of T3, the MCT-8, and for deiodinase (D1, D2), has emphasized that in three types of knock-out mice (MCT-8, and MCT8/D2 MCT8/D1/D2) there was an increase in circulating
T3. However, in the same three genotypes of knock-out mice, the tissue concentration of T3 was systematically increased in the liver, but systematically reduced in the cerebral cortex. In practice, as demonstrated also by the expression of some liver genes and some brain genes, there was a condition of hyperthyroidism in the liver, but, in contrast, a condition of hypothyroidism in the cerebral cortex. Thus, the biochemical hyperthyroidism (increased circulating concentrations of FT3 and/or FT4) does not necessarily mean universal tissue hyperthyroidism. Even the Italian study of Sampaolo et al. [10] shows how poorly conclusive is the concentration of circulating thyroid hormones: the plasma levels assessed in a group of patients suffering from Alzheimer’s disease were similar to those observed in the controls, on the contrary, the concentration of T3 in the liquor was about three times lower than in healthy subjects.

In this respect, precursors have been the fundamental studies of the Madrid group Escobar-Morreale et al. on mice made hypothyroid and then variously supplemented in order to make them euthyroid [11-13]. While the increase of the plasmatic T3 concentration (following the increase in the dose of T3 infused) results in an equivalent increase in the intra-tissue concentration of T3 for various tissues (even for the ovaries, the adrenal cortex and the lungs, the intra-tissue increase is higher than the plasmatic one), for other tissues (pituitary, cerebellum and, especially, for cerebral cortex and brown adipose tissue), the increase of intra-tissue concentration of T3 is much lower than the expected one based on the increase of plasmatic concentration. In the cerebral cortex, the infusion of T3 (and not the one of T4) is the one that does increase the T3 tissue concentration in a dose-dependent manner. And again, the same posology combination of T4+T3 does not result in concentrations of T4 and T3 which are exactly identical in the various tissues; and, in effect, a tissue concentration of both T3 and T4 fully corresponding to the normal range of a given tissue is achieved with a posology combination of T4+T3 that is different from the one which provides a tissue concentration of both T4 and T3 fully corresponding to the normal range of another tissue. The final practical conclusion of these studies of Escobar-Morreale et al. is that the normalization of the intracellular content of both T4 and, especially, T3, in all tissues of hypothyroid rats, requires the combined administration of T4 and T3.

Returning to the human species, regarding the various parameters of neuropsychological wellness, investigated by the works that have addressed the possible beneficial effect of combination therapy compared with monotherapy with T4, it seems that there is a certain benefit of the combination T4+T3 in depression.

In psychiatric literature, in fact, there is evidence of how the administration of T3, as monotherapy or as an additive to antidepressant therapy or as an accelerator in initial association with drugs, is effective in improving the outcome of patients being treated with antidepressants. In most cases, T3 was administered at a dose between 5 and 50 g/day. Other studies have evidence of the greater effectiveness of T3 compared to T4 for this purpose [14], as well as of a better response in females than males [15, 16].

However, according to U.S. guidelines, there is insufficient evidence to support the
use of thyroid hormones in the treatment of depression in euthyroid patients [3].

This mention of the “antidepressant” use of T3 leads to recall a particular mode of action of thyroid hormones, a mode which is seldom or never mentioned in the literature and textbooks/treatises of endocrinology. The studies of Dratman and Gordon [17] show, in a very convincing way, how thyroid hormones work as neurotransmitters. In particular, the structure of the central nervous system richer deiodinase type 2 (D2) is the locus coeruleus, localized in the sub-ependymal area. This site has a high concentration of tyrosine-hydroxylase, an enzyme that converts tyrosine into norepinephrine. Incidentally, it is not superfluous to recall that even T3 and T4 originate from tyrosine; however, unlike the norepinephrine, are iodine molecule. It is important to emphasize the role played as powerful activator of D2 by norepinephrine and, on the other hand, the ability of thyroid hormones to stimulate strongly the expression of the adrenergic receptors in the brain. So, although the reasons are still unknown, nature has made sure that the metabolic pathways of T3 and norepinephrine are closely related to each other, that the two neurotransmitters travel along the same axons and interface post-synaptically to transmit signals.

An interesting aspect that emerges from the literature on the neuro-transmissal role of the thyroid hormones T3 and T4 is that they have even different effects. In the classical field of genomic action of thyroid hormones is well known, however, that T3 and T4 act in the same direction, although the T3 is more potent than T4. Caria et al. [18] evaluated the effects of the two hormones separately on the frequency of neuronal discharge induced by norepinephrine in cultured hippocampus. When the noradrenergic stimulation was preceded by the infusion of T4, it was noted a depressive effect on the frequency of neuronal discharge, on the contrary, when the noradrenergic stimulation was preceded by the infusion of T3, we obtained a significant increase of neuronal discharge (Figure 1). Other works have also demonstrated the different effect of the two hormones on glutamatergic [19] and GABAergic neurotransmission, both phasic and tonic [20]. Taking into consideration the above background (thyroid hormones as neurotransmitters), found that there was no electrophysiological studies on the motor cerebral cortex in hypothyroid patients (and we just said that the motor cortex is one of the main targets of noradrenergic neurons located in the locus coeruleus) and using the non-invasive transcranial magnetic stimulation we wanted to test parameters of human motor cortex excitability in adult patients with acquired hypothyroidism before and after replacement therapy with LT4 [21]. Trans-cranial magnetic stimulation is a non-invasive, painless, safe and accurate technique to induce electrical potentials in brain through electromagnetic induction. The electrical impulse generated in brain (in our case specifically, the motor cortex) is registered with motor-evoked potential in a distal skeletal contra-lateral muscle (in our study the left motor cortex was stimulated and it was recorded in muscle called the first dorsal interosseous of the right index finger).

We carefully selected, from the of neurological and neuropsychiatric point of view, 11 patients with primary overt hypothyroidism, which were evaluated before replace-
Figure 1. Non-genomic action of thyroid hormones: effects on excitability of hippocampal neurons

Source: Carla et al., 2009 [18], adapted.

ment with levothyroxine monotherapy and at the end of the 3rd and 6th month post-therapy. As controls, were recruited as many healthy subjects, matched for various features. In the patients group, all reached the euthyroid condition (normalization of TSH and FT4) already at the end of the 3rd month of therapy. The results, summarized in **Figure 2**, for three illustrative parameters, show normalization of parameters both at the end of the 3rd month and at the end of the 6th month of therapy with levothyroxine.

However, it must be said that, in the evaluation phase of the work, it was asked to exclude from the case studies a patient who, despite having reached the biochemical euthyroid, still had abnormal values (as per overt hypothyroidism) of the various parameters of trans-cranial magnetic stimulation. In retrospect, this subject must be interpreted as a patient in which, despite the normalization of circulating thyroid hormones, the biological action of the hormones on the nervous districts underlying those parameters was still insufficient. Likely in this patient those parameters would have changed in euthyroid sense if the T4 monotherapy had been switched to
Figure 2. Parameters illustrative of excitability of the motor cortex (assessed in the trans-cranial magnetic stimulation) in 10 patients with primary hypothyroidism (before, 3 and 6 months after replacement therapy with LT4) and 10 euthyroid controls.

The symbols in red (▲ for the 3rd month ● 6th month) refer to the eleventh patient that it was asked to exclude from the case study because it was considered outliers (see text). In fact, this patient had parameters of motor excitability, both at the end of the 3rd month and the end of the 6th month, which were still values as per hypothyroidism condition, despite he had reached the biochemical euthyroid condition already at the 3rd month.

Source: Rizzo et al., 2008 [21], adapted.

T4+T3 combination therapy. In fact, some of the parameters of trans-cranial magnetic stimulation underly the activity of glutamatergic and GABAergic interneurons.

We conclude by stressing the complexity of the treatment modalities of hypothyroidism, which descends from the complexity of the thyroid hormonal signal at all levels (synthesis, plasmatic transport, cell transport, metabolism, cell and nuclear target) and from the systematicity of the thyroid hormone action (therefore systematic-
ity of the clinical manifestations of hypothyroidism). It is therefore readily apparent that the universe of hypothyroid patients cannot be released by 100% of hypothyroidism disorders only with the use of a single hormone (indeed, pro hormone, namely the T₄), and not even with administration of the same in a single daily dose. It arises, therefore, the need to be able to identify those patients that may take greater benefit from the combination therapy of T₄+T₃.

References


Tissue-specific patterns of changes in 3,5,3'-triiodo-L-thyronine concentrations in thyroidectomized rats infused with increasing doses of the hormone. Which are the regulatory mechanisms? Biochimie 1999 May;81(5):453-62.


Thyroid hormones as neurotransmitters. Thyroid 1996 Dec;6(6):639-47.


In physiological conditions, the total amount of T3 in the body is given by the algebraic sum of the hormone produced and secreted by the thyroid, the T3 coming from intra-thyroidal and peripheral conversion of T4, and, finally, by the catabolism and excretion of the active hormone. In conditions of replacement therapy, namely in the absence of an endogenous T3 production, the available quantity of the hormone is only guaranteed by the mechanism of peripheral conversion of T4.

The current standard of care of hypothyroidism is the monotherapy with levothyroxine, of which we know the pharmacokinetic mechanisms, methods of administration and therapeutic targets.

With regard to liothyronine (the synthetic formulation of T3), the pharmacokinetic and pharmacodynamic mechanisms are less clear because the data currently available have been obtained through studies with obsolete methods and non-physiological situations.

In order to characterize and understand properly the pharmacokinetics of therapy with T3 and its metabolic effects, in our studies we have tried to eliminate some bias which compromised the results of previous works enrolling, in the first place, well-standardized groups of patients.

In a double-blind study (2011) [1, 2], we enrolled 14 hypothyroid volunteers, 12 of whom had undergone a total thyroidectomy, one had undergone radioiodine therapy for Basedow’ disease, and one had a diagnosis of Hashimoto’s thyroiditis. Patients with residual glandular >1 cm³ were subjected to study with scintigraphic uptake of radioiodine, to rule out a residual endogenous hormone production.

A prerequisite to be able to participate in the study was that each patient reaches the euthyroid state at pituitary level, by treatment with levothyroxine or with liothyronine, lasting at least one month. At the end of the adjustment period of therapy the
dose of levothyroxine was equal to 1.6 g/kg vs. a dose approximately three times less of liothyronine. Both drugs were taken with three daily doses.

Between the two groups of patients TSH values at baseline were fully overlapping, as well as the response to TRH test [3]. Even during 24 hours TSH did not show significant differences. Our patients were thus perfectly identical and so we had achieved our target, i.e. the euthyroid condition at pituitary level. Moreover, we observed a significant difference in the levels of plasmatic T3 between patients treated with levothyroxine and those treated liothyronine. In fact, while in patients treated with liothyronine we found average levels of the hormone within the normal range in patients treated only with levothyroxine the T3 was maintained on average at the lower limits of the standard (● Table 1). Our interest however, rather than to the hormonal levels in plasma, was focused on the study of the metabolic effects of hormone action in the two cases under consideration. In fact, we found significant differences between the two groups of patients: in patients treated with liothyronine there had been a significant reduction in the amount of total cholesterol and LDL, and, in a marginal way, even in HDL (● Table 2). We have observed, moreover, an increase in serum levels of SHBG and a reduction in the average weight of about 2 kg, in absence of dietary modifications. No difference between the two groups was observed within the cardiovascular parameters (systolic blood pressure, diastolic blood pressure, heart rate and echo-cardiographic parameters), the basal and glucose metabolism.

In another study that is not yet complete and from which we present preliminary data we enrolled patients in hypothyroidism after suspension of replacement therapy with levothyroxine, waiting to be subjected to nuclear medicine procedures post-thyroidectomy for thyroid cancer. In the 30 days prior to the suspension of treatment was performed a shift from levothyroxine to liothyronine in suppressive mode and in

● Table 1. Values of thyroid hormones and TSH in terms of drug-equivalence between levothyroxine (LT4) and liothyronine (LT3)

<table>
<thead>
<tr>
<th></th>
<th>LT4 (mU/L)</th>
<th>LT3 (mU/L)</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>TSH ± SD 0.4-4.0</td>
<td>1.3 ± 0.79</td>
<td>1.4 ± 0.79</td>
<td>0.674</td>
</tr>
<tr>
<td>T3 (ng/dL) ± SD 90-215</td>
<td>92.9 ± 19.0</td>
<td>172.0 ± 88.2</td>
<td>0.003</td>
</tr>
<tr>
<td>fT4 (ng/dL) ± SD 0.8-1.5</td>
<td>1.57 ± 0.3</td>
<td>&lt; 0.3 &lt; 0.0001</td>
<td>&lt; 0.0001</td>
</tr>
</tbody>
</table>

Source: Celi et al., 2011 [1] adapted.
divided doses. The therapeutic target, of course, was represented by the suppression of TSH. We therefore analyzed the pharmacokinetics of the last dose with sequential doses of T3 and TSH. The latter, immediately before discontinuation of therapy, not surprisingly resulted suppressed, while the average T3 at baseline was located at the higher limits of the normal range compared to the mean values measured during treatment with levothyroxine. A preliminary data analysis indicates that the apparent half-life of the drug appeared around 27 hours, with a half-life of distribution of 3-4 hours and half-life of elimination of approximately two days. The administration fractionated in three daily doses, also assured normal plasmatic T3 levels in a constant and prolonged manner; in contrast, giving the entire dose in a single time or in two times would have favored the onset of peaks and important oscillations of the hormone concentration in circulation.

These data indicate that the treatment with liothyronine in the correct dosage per weight and per initial dose of levothyroxine in multiple administrations, it ensures an adequate TSH suppression with T3 levels within normal limits. Discontinuation of the therapy ensures TSH levels adequate to radio-metabolic treatment for a period of about two weeks, with minimal symptoms of hypothyroid. The study showed, in addition, that the pharmacokinetics of liothyronine comprises two components: distribution and elimination. An administration in single or double dose of the drug would result in prolonged periods of over-and under-dosing. These results are empirical data on which to set up long-term controlled studies regarding combined levothyroxine-liothyronine therapy, which identify clinically valid endpoint in order to evaluate the metabolic effects of this treatment modality.

Table 2. Lipidic metabolism under conditions of drug-equivalence of levothyroxine (LT4) and liothyronine (LT3)

<table>
<thead>
<tr>
<th></th>
<th>LT4</th>
<th>LT3</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Total cholesterol</td>
<td>195.9</td>
<td>173.9</td>
<td>0.002</td>
</tr>
<tr>
<td>(mg/dL) ± SD</td>
<td>25.9</td>
<td>27.7</td>
<td></td>
</tr>
<tr>
<td>HDL (mg/dL) ± SD</td>
<td>63.0</td>
<td>57.5</td>
<td>0.067</td>
</tr>
<tr>
<td>LDL (mg/dL) ± SD</td>
<td>122.6</td>
<td>106.2</td>
<td>0.002</td>
</tr>
<tr>
<td>Triglycerides (mg/dL) ± SD</td>
<td>78.2</td>
<td>78.8</td>
<td>0.937</td>
</tr>
<tr>
<td></td>
<td>30.8</td>
<td>28.6</td>
<td></td>
</tr>
</tbody>
</table>

Source: Celi et al., 2011 [1] adapted.
References


In literature, the topic “low T3 syndrome” is controversial for at least three reasons: the complexity of the pathophysiology, and, consequently, of the treatment; the criticality of the clinical conditions of the patients that involves the intervention of numerous confounding factors (comorbidities, medications, etc.), the discordance of treatment studies currently in the literature.

For nearly forty years we know that, in course of fasting or of acute or chronic diseases, may occur alterations of the thyroid homeostasis, a condition that in literature does not present an univocal name: we speak, in most cases of “low T3 syndrome” or “euthyroid sick syndrome”, but the most suitable definition is probably that of “Not thyroidal Illness syndrome” (NTIS), as to emphasize that the patient is normal from the point of view of thyroid function and that it is a secondary response of the gland to a condition of systemic disease.

The reduction in the plasma concentration of T3, in fact, is only one of the phenomena that occur in the context of a clinical systemic picture much more complex. Specifically, the deficit of the active hormone, resulting from the inhibition of deiodination of T4, is a very early event in the involvement of the hypothalamic-pituitary-thyroid axis in response to situations of acute or chronic stress. With the worsening or the prolongation of the underlying disease, in fact, there is a progressive extension of the functional block to the whole axis, which results, in most serious cases, in a reduction even of the circulating levels of T4 associated with values of TSH inappropriately normal. This last data has been interpreted, by some, as an index of hypothalamic alteration, hypothesis supported by the concomitant alteration of other hormones such as GH or cortisol. Furthermore, a normal thyroid function is restored not only in response to TRH infusion, but also by co-administration of other hypothalamic peptides such as, for example, the secretagogues GH.
Other authors [1] have attempted to justify the normality of TSH in these patients as a result of normal deiodination activities at central level but not at peripheral level. All the mechanisms downstream of hormone secretion, implicated in the transmission of the thyroid signal were analyzed in order to better understand the physiopathology of the low T₃ syndrome. In literature there are many conflicting studies on plasmatic transport proteins, on receptors of thyroid hormones, on intracellular transporters, which do not allow drawing a univocal picture of this clinical condition.

Among the mechanisms involved in the pathogenesis of the syndrome, it has recently been revealed a significant increase of Nuclear Factor-kB (NFkβ) in response to the massive release of cytokines such as Interleukin-6 (IL-6) and Tumor Necrosis Factor-α (TNF-α) [2]. Various studies have shown that the NFkβ, mitigating the expression of deiodinase type 1, is able to reduce the circulating levels of T₃ in both rats and humans.

To complicate the picture of the syndrome, which makes more difficult to attempt understanding the pathogenic mechanisms, is the complexity of the condition of the patient with “low T₃ syndrome”, which comes from the underlying pathology, acute or chronic, from the state of malnutrition that often accompanies it, from the wide range of drugs that can interfere with the endocrine axis at various levels. Often, these patients take glucocorticoids, β-blockers or anti-arrhythmics, all drugs that block deiodinase enzymes. Other drugs, such as heparin, alter the hormone binding to plasma proteins and still others, such as catecholamines and dopamine, interfere with the release of TSH at pituitary level.

The clinical conditions that frequently accompany the low T₃ syndrome are undoubtedly heart and lung diseases. Followed by major surgery interventions, sepsis, trauma. At the moment, we try desperately to understand if thyroid changes are only a mechanism of adaptation to the underlying disease or, conversely, a pathogenic factor also implicated in the progression and aggravation of the disease. It is important to give an answer to this question, especially in view of the therapeutic approach: if it were a mere form of adaptation does not require any intervention, otherwise, it would be necessary a substitution treatment. In this context, it is important to identify valid indexes of tissue hypothyroidism, able to reflect the thyroid situation at peripheral level, and that allow clarifying dubious situations from the clinical point of view. Coenzyme Q₁₀, an important component of the mitochondrial respiratory chain and a powerful antioxidant, would seem suitable for this purpose [3, 4]: since the eighties we know, in fact, that the levels of this molecule correlate inversely with circulating values of T₃ and T₄, in a highly significant manner, to the point to “pick up” even subclinical conditions.

Another important biochemical marker of real hypothyroidism at tissue level is represented by oxidative stress, in particular, the hormone deficiency is associated with an increase in oxidizing and a parallel reduction of antioxidants; the replacement therapy reverses this condition. In the clinical practice, 20-30% of patients
with dilated cardiomyopathy have a low T₃ syndrome, with a higher incidence in decompensated forms. There is a significant correlation between low plasma levels of T₃ and mortality from heart disease, and it has also been shown an increase in morbidity and mortality due to heart conditions in the form of subclinical hypothyroidism.

The opportunity to use T₃ in cardiac pathology is supported by biological and clinical evidence, we know that thyroid hormones modulate myocardial contractility, increasing, thus, the performance of both systolic and diastolic; inducing coronary vasodilation also they promote myocardial perfusion.

In a pioneering study of Hamilton et al. [5], patients with dilated cardiomyopathy unresponsive to other inotropic agents were treated with intravenous infusion of T₃, with amazing effects: an increase in cardiac output, reduction of vascular resistance, reduction of blood pressure and pulmonary capillary wedge pressure (● Figure 1).

In another study of the Pisan physiology school [6], the infusion of T₃ in patients with heart disease resulted in a significant improvement in both systolic and diastolic parameters (● Table 1).

Despite this evidence, however, it is impossible to find a unique perspective with which to look at this complex syndrome in the literature.

De Groot, an eminent expert in low T₃ syndrome, confirms the beneficial effects of T₃ administration from the cardiovascular point of view, in different categories of patients, especially cardiological patients, without however in fact, significant changes in the prognosis [7].

On the other hand, alarming studies demonstrate even an increase in mortality in patients with renal insufficiency; in addition, the fear of cardiologists and endocrinologists about the fact that the administration of T₃, modifying the tissue request of oxygen, may, in patients at risk, precipitate an ischemic event limits its use in clinical practice.

In the field of pneumology, are present in the literature numerous discordant studies regarding the association between low T₃ syndrome and COPD (chronic obstructive pulmonary disease). According to some authors, in course of this systemic disease would not occur any change in the thyroid function, while others have shown that there is a reduction in circulating hormone proportional to the hypoxia, associated or not to TSH alteration. In a isolated study, it was even observed an increase of T₃ in these patients [8].

In our study on 32 patients with COPD [9] (● Table 2) we put in evidence, in fact, alterations in circulating hormone levels compared to healthy controls: the average TSH, even within the normal range was maintained at high limits, the average T₃ was lower than normal, while the T₄ stood at the lower limits of the reference range. In addition, we have demonstrated in these subjects a significant reduction in antioxidant defenses. Surprisingly, there was a very significant correlation between low levels of circulating T₃, then hypothyroidism, and increased oxidative stress, a
condition that it was possible to normalize with replacement therapy. The low values of T₃, however, were not uniformly present in the entire sample under examination: about 1/3 of the patients showed, in fact, a syndrome in Low T₃, but 20 had still normal hormonal values. We, therefore, separated the results from the total, comparing the two subsets of patients with healthy controls, and we have confirmed, in patients with low T₃, the data previously described. It is also interesting to note that the hormonal deficiency did not correlate significantly with the alteration of any respiratory parameter.

Another preliminary work of our group on patients in pulmonary sub-intensive therapy for acute respiratory failure (unpublished data) (● Figure 2), showed an inverse correlation, which tended to be significant, between values of FT₃ and pCO₂,
even if the sample under examination was rather small. Hypoxia, in these patients, was responsible for the reduction of plasma levels of the hormone, while hypercapnia proved then actually associated with a poorer prognosis.

Preliminary data also demonstrate that if these patients are treated with T3 (20 µg/day orally in two doses) show an improvement in gas exchange in a rather short period of time (two weeks), compared to untreated patients. These data, although preliminary, suggest the need to evaluate T3 levels in pulmonary critical patient, especially if hypercapnic.

To conclude, there is still no consensus on whether to treat patients with low T3 syndrome, but we have enough data to say that the treatment is necessary in cases where it can be proved the actual presence of tissue hypothyroidism.

Table 1. Cardiovascular parameters at baseline and after 3 days of treatment with infusion of synthetic LT3 or placebo

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Patients treated with LT3 Before LT3</th>
<th>Patients treated with LT3 After LT3</th>
<th>P value before LT3 vs. after LT3 Basal</th>
<th>P value before LT3 vs. basal</th>
</tr>
</thead>
<tbody>
<tr>
<td>LV EDV (ml/m² bs)</td>
<td>133 (114-158)</td>
<td>142 (132-161)</td>
<td>0.02</td>
<td>130 (117-153)</td>
</tr>
<tr>
<td>LV ESV (ml/m² bs)</td>
<td>103 (84-127)</td>
<td>108 (89-124)</td>
<td>ns</td>
<td>91 (86-115)</td>
</tr>
<tr>
<td>LV SV (ml/m² bs)</td>
<td>35 (28-39)</td>
<td>40 (34-44)</td>
<td>0.01</td>
<td>36 (29-48)</td>
</tr>
<tr>
<td>CO (liter/min)</td>
<td>4.1 (3.3-5.4)</td>
<td>4.8 (3.4-5.4)</td>
<td>ns</td>
<td>4.7 (4.0-5.3)</td>
</tr>
<tr>
<td>CI (liter/m² bs x min)</td>
<td>2.2 (1.7-2.8)</td>
<td>2.5 (1.9-2.7)</td>
<td>ns</td>
<td>2.5 (2.1-2.9)</td>
</tr>
<tr>
<td>LV EF (%)</td>
<td>25 (18-32)</td>
<td>28 (22-32)</td>
<td>ns</td>
<td>27 (23-41)</td>
</tr>
<tr>
<td>SVR (dyne/sec x cm)</td>
<td>2.07 (1.92-3.13)</td>
<td>2.10 (1.87-2.48)</td>
<td>ns</td>
<td>2.03 (1.86-2.36)</td>
</tr>
<tr>
<td>Elastance</td>
<td>1.36 (0.93-1.63)</td>
<td>1.27 (0.91-1.36)</td>
<td>ns</td>
<td>1.32 (0.97-2.14)</td>
</tr>
<tr>
<td>External cardiac work (ml x mm Hg x bpm)</td>
<td>201,226 (161,084-3,002,307)</td>
<td>226,519 (169,276-266,388)</td>
<td>ns</td>
<td>253,950 (190,929-306,180)</td>
</tr>
<tr>
<td>Internal cardiac work (ml x bpm x mm Hg/2)</td>
<td>401,849 (348,910-534,505)</td>
<td>396,885 (343,080-473,613)</td>
<td>ns</td>
<td>360,260 (314,153-440,763)</td>
</tr>
<tr>
<td>Total cardiac work</td>
<td>626,859 (492,291-787,522)</td>
<td>592,085 (540,060-756,684)</td>
<td>ns</td>
<td>599,945 (538,645-748,639)</td>
</tr>
</tbody>
</table>

Data are expressed as median (25º and 75º percentiles); bpm, beats per minute; CI, cardiac index. Parameters statistically significant are highlighted.

Source: Pingitore et al., 2008 [6].
Table 2. Mean values (± SD) of the parameters of thyroid function, values of coenzyme Q10 corrected for the concentrations of cholesterol, and total antioxidant capacity (expressed as LAG) in subjects with chronic obstructive pulmonary disease (COPD) and normal controls

<table>
<thead>
<tr>
<th></th>
<th>Normal subjects (n = 45)</th>
<th>COPD patients (n = 32)</th>
<th>F</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>TSH μUI/ml</td>
<td>1.15 ± 1.12</td>
<td>1.92 ± 1.63</td>
<td>4.93</td>
<td>0.03</td>
</tr>
<tr>
<td>FT3 pg/ml</td>
<td>3.55 ± 1.11</td>
<td>2.82 ± 0.76</td>
<td>9.51</td>
<td>0.003</td>
</tr>
<tr>
<td>FT4 pg/ml</td>
<td>14.6 ± 7.46</td>
<td>11.4 ± 4.27</td>
<td>4.48</td>
<td>0.04</td>
</tr>
<tr>
<td>CoQ10 μg/ml</td>
<td>0.75 ± 0.24</td>
<td>0.85 ± 0.27</td>
<td>2.55</td>
<td>ns</td>
</tr>
<tr>
<td>CoQ10/chol nmol/mmol</td>
<td>213.45 ± 67.73</td>
<td>205.59 ± 81.03</td>
<td>0.12</td>
<td>ns</td>
</tr>
<tr>
<td>LAG sec</td>
<td>74.44 ± 15.7</td>
<td>61.3 ± 13.16</td>
<td>5.79</td>
<td>0.002</td>
</tr>
</tbody>
</table>

F, value with the test of variance; P, level of significance
Differences statistically significant are highlighted.

Source: Mancini et al., 2012 [9].

Figure 2. Changes in PaCO2 in patients with respiratory failure treated (left) or not treated with T3

Source: original elaboration, unpublished data.
References


The sulfo-conjugation path of thyroid hormones has been known for many years. Already in 1957 Roche et al. [1] had identified sulfo-conjugates hormones in the blood and in the bile. These observations led to assume, rightly, that sulfo-conjugation was a way to facilitate the biliary excretion of T3. Today we have, without doubt, a wider knowledge about the biological significance of this metabolic pathway.

The sulfo-conjugation of T3 is the addition of a sulfate group at the level of a hydroxyl of the outer ring of the hormone molecule (● Figure 1).

This process is mediated by enzymes localized at the level of various tissues such as liver, placenta, kidney, central nervous system, and cause a significant impact on the fate of the molecule. The sulfo-conjugation of T3 transforms the hormone from lipophilic molecule to molecule soluble in water. Furthermore, the T3 sulfate is inert from a functional point of view as unable to bind the nuclear receptor, it has a high affinity for the plasmatic carrier proteins and is easily hydrolyzed in acid environment (● Table 1).

To understand the metabolic consequences of sulfo-conjugation, we can take as a reference a work that dates back to the eighties [2]. We know that in the liver deiodinase type 1 converts T4 to T3 and that the latter, on the contrary, constitutes a poor substrate for the enzyme itself. In the study by Otten, incubating a sample of hepatocytes with T3 and adding increasing amounts of sulfate, occurred an acceleration of the degradation of the hormone with the appearance of free iodine. The explanation of this phenomenon was obtained by adding propyl-thiouracil (PTU) to the culture liquid of the same cells: the PTU is an inhibitor of D1 and its presence inhibited the formation of free iodine while it was associated to the appearance of T3 sulfate. The authors could thus conclude that the sulfate promotes the production of T3 sulfate and that the latter, unlike the active hormone, is rapidly deiodazed from liver D1.
Based on these studies, we now know that the process of sulfo-conjugation is not only a mechanism to facilitate the biliary excretion of T3 but is a way that accelerates its degradation.

In contrast, the sulfo-conjugation guarantees protection from the action of deiodinase type 3 (D3), the enzyme that degrades the active T3 and that is found in many tissues such as placenta and skin [3].

This has implications in practical terms, if we measure the serum levels of T3 sulfate we notice significant differences in various clinical conditions. In particular, low amounts of the hormone can be detected in euthyroid subject, while circulating levels are elevated in hyperthyroid patient (for increase of the precursor, that is T3), but also in hypothyroid patient, in non-thyroidal illnesses, in the fetus and newborn, all conditions in which it is inhibited the D1 [4, 5].

To confirm these observations, when to hyperthyroid patients was administered sodium ipodate, a contrast agent that inhibits the D1, it was observed a reduction of plasmatic levels of T3 accompanied by an increase of T3 sulfate and rT3.

The presence of very high concentrations of the metabolite in the fetal and neonatal life has led to the hypothesis that the T3 sulfate, after desulphation, can represent to some extent a source of active T3. In this regard, it has been demonstrated the presence in the liver of an enzyme system capable of desulfating T3 sulfate and, therefore, to reactivate it [6, 7].

Furthermore, by administering T3 sulfate intra-peritoneally in hypothyroid rats it was possible to restore a smooth curve of growth and normalize other parameters re-
lated to the thyroid status of the animals [8]. In particular, following a single injection of T3 sulfate it was obtained a normalization of serum levels of T3, which remained constant for about 8 hours.

The results of preliminary studies in humans [9] indicate that the T3 sulfate can be absorbed after oral administration in hypothyroid patients, in addition, after administration of T3 sulfate it is observed a proportional increase in T3 serum concentrations which remains stable for at least 48 hours, unlike what is observed after administration of the active hormone. After T3 administration it occurs in fact a rapid absorption peak followed by an equally fast descent that ends in a matter of hours. In theory, therefore, the T3 sulfate may represent a slow release formulation able to improve the satisfaction of those hypothyroid patients who continue to complain about hormonal deficit disorders, despite the normalization of serum parameters obtained with the administration L-thyroxine alone.

---

**Table 1. Properties of T3 sulfate**

<table>
<thead>
<tr>
<th>Property</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrophilic molecule</td>
</tr>
<tr>
<td>Negligible affinity for the nuclear receptor of T3</td>
</tr>
<tr>
<td>High affinity for plasmatic carrier proteins</td>
</tr>
<tr>
<td>Susceptible to acid hydrolysis</td>
</tr>
<tr>
<td>Rapid deiodination by deiodinase type 1 (D1)</td>
</tr>
<tr>
<td>Resistant to deiodination by deiodinase type 3 (D3)</td>
</tr>
</tbody>
</table>

---

**References**


The hypothalamus-pituitary-thyroid axis (HPT) preserves thyroid hormone’s homeostasis producing T₃ and T₄ and controlling concentrations of the two hormones through a sensitive feedback mechanism. Hypothalamic TRH neurons and TSH producing cells can sense at the same time both T₄ and T₃, because of the presence of type 2 deiodinase that convert T₄ to T₃. So, the HPT senses at the same time T₃ produced from local conversion of T₄ in hypothalamus and in pituitary, and the T₃ in the circulation (● Figure 1).

We know that most of the biological effect of thyroid hormones, such as cognition, metabolism, growth and development, are mediated by the interaction of T₃ with nuclear receptors.

Every day the organism produces about 30 µg of T₃, but only 5 µg comes from the thyroid; the rest comes from peripheral conversion from T₄ by type 1 and type 2 deiodinases (D₁ e D₂). These enzymes show different expression patterns in different tissues. For example, D₁ is expressed above all in kidneys and liver, while D₂ is expressed in brain, skin, skeletal muscle and brown adipose tissue. T₄ that comes into the tissue is transformed to T₃ in a compartment that is near the cell nucleus, stays in the cell for a few hours (“equilibration time” with the plasma is about 3-4 hours for D₂ and approximately 30 minutes for D₁), and exits to the plasma. So, every day, about 20 µg of T₃ are produced from conversion of T₄ by D₂, while D₁, that has low affinity for T₄, produces about 5 µg/day of T₃. Ultimately T₃, which is a good substrate for type 3 deiodinase, is then inactivated to T₂ (● Figure 1).

Studies by Andersen measuring TSH, FT₃ and FT₄ over a long period of time indicated that serum levels of T₃ are stable [1].

To test how well the organism defends serum T₃, a few decades ago Larsen evaluated rats kept on low iodine diet. While their serum levels of T₄ dropped and serum TSH
increased, serum T3 remained stable. This has been observed also in humans living in areas of low iodine diet. Overall these observations suggest that the HPT defends serum T3, preserving it even during iodine deficiency [2].

But what happens when we inactivate D2, considering that the most T3 is produced by this enzyme? The study of Christoffolete et al. shows that serum T3 is normal in the D2 knock-out mouse mostly by a compensatory increased production of T3 from the thyroid gland [3].

Moreover, when we evaluated the serum T3 of another mouse model that lacked D2 just in the pituitary cells producing TSH (we confirmed that D2 activity was lost only in these cells and not in other regions of the brain), it was normal despite alterations in TSH and FT4 levels.

We can therefore conclude that the HPT is wired to maintain a normal serum T3, and that in this mechanism the thyroid gland plays a major role [4].

What happens in primary hypothyroidism? Current guidelines suggest that patients with primary hypothyroidism should be treated with levothyroxine (T4) monotherapy. Such treatment usually leads to resolution of the symptoms and normalization of TSH and T4 levels and, in most cases, T3 levels as well, which is derived exclusively
from peripheral activation of T4 mediated by deiodinases. However, about 10-15% of hypothyroid patients on levothyroxine monotherapy, despite being biochemically euthyroid based on a normal TSH, remain symptomatic. One of the hypotheses therefore is that the small amount of T3 that the thyroid gland produces has some key role in achieving that sense of wellbeing that some hypothyroid patients on levothyroxine monotherapy seem to struggle to obtain.

A study of Gullo et al. showed that euthyroidism cannot be restored in all post-surgical hypothyroid patients through monotherapy with levothyroxine (Figure 2). Stratifying these data by serum TSH, they compared euthyroid and hypothyroid patients on monotherapy that had the same level of TSH and observed that for every level of TSH, the athyrotic patients had a relatively lower serum T3 and a higher serum T4. Consequently, post-surgical hypothyroid patients maintain a normal serum TSH because of an excess replacement of T4 that compensates for a relative deficiency of serum T3. Therefore, serum TSH, in these cases, is not a reliable indicator of euthyroid state since even if TSH is normal, T3 may be low. This could affect

* Figure 2. Levothyroxine monotherapy cannot guarantee euthyroidism in all athyreotic patients

![Diagram showing the relationship between TRH, TSH, T4, T3, and L-thyroxine](Image)

**CAN SERUM T3 BE MAINTAINED IN L-T4-TREATED HYPOTHYROID PATIENTS?**

*Source: Gullo et al., 2011 [5], adapted.*
especially those organs/tissues that do not have D2 and that therefore depend solely on serum T3 [5].

Another bias for the comprehension of thyroid status regards the techniques of measuring T3. Jonklaas et al. compared serum T3 levels of patients after thyroidectomy as measured by immunoassay or by mass spectrometry. Immunoassays overestimate serum T3 values in comparison with mass spectrometry when values are about 100 ng/dl or lower, concluding that immunoassays underestimate the number of patients with low serum T3 [6].

In the study of Geffner et al., PTU, an inhibitor of type 1 deiodinase was given to hypothyroid patients on monotherapy with levothyroxine. They observed that while FT4 did not change, a relatively small decrease in serum T3 caused serum TSH to double. We could then speculate that the fall in serum T3, albeit small, results in pituitary hypothyroidism. Therefore, we should expect that other tissues that depend on serum T3 can become hypothyroid as well [7].

We know that there is an equilibrium between tissue T3 and plasma T3 and that not all cells depend exclusively on plasma T3. In the brain T3 plays a major role in cognitive function. D2 is responsible for more than half of the T3 present in the murine brain. Accordingly, D2KO animals have half as much brain T3 content as their wild type siblings greatly supporting the idea that any interference in the D2 pathway could affect brain function and/or result in intellectual or cognitive symptoms [8].

Is there a clinical syndrome that involves D2 deficiency? If so, could this explain the efficacy of combined therapy for some patients?

The group of Celi discovered a particular polymorphism (Thr92Ala DIO2) that was associated to a specific metabolic phenotype: increased BMI and glucose intolerance. D2 KO mice are obese and have intolerance to glucose as well. Later, many other studies confirmed this association and tried to explain how this polymorphism could produce a similar phenotype, since the association is not sufficient to confirm that the polymorphism is really influencing D2 activity [9].

Studies by Panicker et al. showed that hypothyroid patients carrying the polymorphism in the DIO2 gene had a better neurocognitive outcome with combined therapy (T3 and T4) compared to standard monotherapy. From these data we could speculate that in these patients D2 doesn’t produce sufficient quantities of T3. However this hypothesis needs to be confirmed and does not necessarily preclude the possibility that there is another rational behind [10].

When we give combination therapy, what happen to T3? To answer that question we performed a systematic review and meta-analysis in 10 randomized controlled trials, 7 of which were crossover. In particular we evaluated and compared the post treatment levels of TSH, free T4, and total T3 (unpublished data). Despite similar TSH level, our analysis revealed a significant difference in the serum level of free T4 and total T3 in patients on monotherapy compared with patients on combined therapy. In particular patients on levothyroxine monotherapy had a higher plasmatic level of
free T₄ suggesting that this may be needed in order to maintain a normal TSH level. Additionally, patients on combined therapy had a significantly higher level of total T₃ (● Table 1, ● Figure 3).

This increased in serum T₃ can be useful for tissue in which D₂ is defective, such as the brain. In this meta-analysis, whether this difference in T₃ level really reflects on a better clinical outcome remains still unclear at the moment. More studies are needed to define the benefits and the complications of maintaining a supra-physiologic level of T₃ over a long period of time.

In conclusion, the HPT is wired to maintain serum T₃, and the thyroid gland does play a major role in this process. We need more studies to define whether serum T₃ can be maintain in the absence of fully functional thyroid gland and how much of the residual thyroid gland activity in hypothyroid patients is enough to defend normal serum T₃.

Also we need more studies to define whether normalization of serum T₃ in hypothyroid patients is clinically relevant.

<table>
<thead>
<tr>
<th>Study</th>
<th>TSH</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Monotherapy</td>
</tr>
<tr>
<td>Crossover</td>
<td>Escobar-Morreale 2005</td>
</tr>
<tr>
<td></td>
<td>Rodriguez 2005</td>
</tr>
<tr>
<td></td>
<td>Siegmund 2004</td>
</tr>
<tr>
<td></td>
<td>Walsh 2003</td>
</tr>
<tr>
<td></td>
<td>Bunevicius 2002</td>
</tr>
<tr>
<td></td>
<td>Bunevicius 2009</td>
</tr>
<tr>
<td>Subtotal</td>
<td></td>
</tr>
<tr>
<td>Test for Heterogeneity</td>
<td>Q-value = 32.336 (df =5)</td>
</tr>
<tr>
<td></td>
<td>Overall effect (Random)</td>
</tr>
<tr>
<td>Non-Crossover</td>
<td>Valizadeh M 2009</td>
</tr>
<tr>
<td></td>
<td>Sawka 2003</td>
</tr>
<tr>
<td></td>
<td>Clyde 2003</td>
</tr>
<tr>
<td>Subtotal</td>
<td></td>
</tr>
<tr>
<td>Test for Heterogeneity</td>
<td>Q-value = 2.158 (df = 2)</td>
</tr>
<tr>
<td></td>
<td>Overall effect (Fixed)</td>
</tr>
<tr>
<td>Total</td>
<td></td>
</tr>
<tr>
<td>Test for Heterogeneity</td>
<td>Q-value = 34.788 (df = 8)</td>
</tr>
<tr>
<td></td>
<td>Overall effect (Random)</td>
</tr>
</tbody>
</table>

Source: Casula et al., unpublished data.
Figure 3. Effect of the monotherapy compared with the combined therapy on total T3

<table>
<thead>
<tr>
<th>Study</th>
<th>Monotherapy</th>
<th>Combined Therapy</th>
<th>SMD* (95%CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Crossover</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Appelhof 2005</td>
<td>1709.40 (292.60)†</td>
<td>1832.60 (400.40)</td>
<td>0.352 (-0.053, 0.757)</td>
</tr>
<tr>
<td>Rodríguez 2005</td>
<td>1131.90 (294.14)</td>
<td>1603.14 (385.00)</td>
<td>1.375 (-0.782, 1.969)</td>
</tr>
<tr>
<td>Bunevicius 2002</td>
<td>3500.00 (1000.00)</td>
<td>3800.00 (1400.00)</td>
<td>0.247 (-0.633, 1.126)</td>
</tr>
<tr>
<td>Bunevicius 2009</td>
<td>1339.00 (585.20)</td>
<td>1801.80 (646.80)</td>
<td>0.750 (0.251, 1.250)</td>
</tr>
<tr>
<td><strong>Subtotal</strong></td>
<td></td>
<td></td>
<td><strong>0.696 (0.215, 1.177)</strong></td>
</tr>
<tr>
<td><strong>Test for Heterogeneity</strong></td>
<td>Q-value = 8.783 (df =3)</td>
<td>$I^2 = 65.843$</td>
<td>p = 0.032</td>
</tr>
<tr>
<td><strong>Overall effect (Random)</strong></td>
<td>Z-value = 2.835</td>
<td></td>
<td>p = 0.005</td>
</tr>
<tr>
<td>Non-Crossover</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Valizadeh M 2009</td>
<td>2032.00 (385.00)</td>
<td>2525.60 (215.60)</td>
<td>1.582 (1.002, 2.162)</td>
</tr>
<tr>
<td><strong>Overall effect</strong></td>
<td>Z-value = 5.347</td>
<td></td>
<td>p &lt; 0.001</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td></td>
<td><strong>0.101 (-0.358, 1.390)</strong></td>
</tr>
<tr>
<td><strong>Test for Heterogeneity</strong></td>
<td>Q-value = 16.833 (df = 4)</td>
<td>$I^2 = 76.237$</td>
<td>p = 0.002</td>
</tr>
<tr>
<td><strong>Overall effect (Random)</strong></td>
<td>Z-value = 3.320</td>
<td></td>
<td>p = 0.001</td>
</tr>
</tbody>
</table>

Source: Casula et al., unpublished data.
Ultimately, it will be important to confirm the clinical role of D2 pathway and specifically to prove that a deficient D2 activity could really explain some residual symptom in hypothyroid patients that have normal serum TSH on monotherapy with levothyroxine.

References


During the discussion in this second Forum of Endocrinology, organized by IBSA, emerged several topics that have been an opportunity for confrontation, but also for debate amongst the participants.

In particular, a hot topic was the one related to the inadequacy of TSH as the sole marker of thyroid function and the need for other tissue markers. The question is somehow tricky, especially considering that the thyroid tissue condition depends only in part on the plasma concentration of T₃, coming also from the expression of membrane transporters, the interaction with the selenium-deiodinases and the expression and the activity of nuclear receptors. TSH, then, for how much is a good pituitary marker, does not accurately reflect what is going on in tissues other than the pituitary.

The circulating T₃ may be helpful in some case but, also, it cannot be considered an optimal marker, for the reasons previously expressed.

On the other hand, it is not the case to generalize: TSH currently remains a “robust” marker in daily clinical practice, resulting less sensitive in a small niche of patients. And it is for the latter that is necessary to identify other markers. Identify a “optimal” tissue marker is a difficult task, especially if one considers that theoretically there should be a specific marker for each individual tissue.

About the reliability of TSH, there is currently a debate about the need and the possibility to perform the simultaneous dosing of the free hormonal fractions, or at least of one of the two hormones. While it is true that the so-called TSH SLR reflex is an expedient to limit health care expenditure, it is inadequate in different circumstances; as we have already seen the value of TSH may be distorted by many interfering agents, along with these there are other bias related to the dosage, and, therefore, to the kits used in the laboratory, as well as to the individual variations of this markers, including the Circadian ones. TSH reflex, then, it may be a good economic compro-
mise to reduce costs, reserving the dosage of at least one of the two hormones (better FT4 according to American guidelines) for the most suspicious cases (autoimmunity, familiarity, perspective of a pregnancy, etc.), as well as for the suspicious forms of central hypothyroidism.

Another aspect that emerged from the discussion is related to the ambiguity that is associated with the definition of well-being of the patient. It is, in fact, a parameter obviously subjective and we do not currently have adequate tools and markers to evaluate it. Not necessarily well-being coincides with the feeling of “being-well” of the patient and, in addition, we cannot be guided solely by patient preference in adopting a specific therapeutic strategy. In this regard, several studies of correlation between therapy and “well-being” were revisited in light of the fact that, in many cases, the well-being of patients resulted from a slight hormone overdose, and therefore from a state of mild hyperthyroidism. This can be also the reason for which, often, hyperthyroid patients report feeling worse after having restored the euthyroid state through therapy. However, if the ultimate goal of treatment was just to make the patient feel good, a slight increase in the hormone dosage could have a rational, monitoring, then, strictly the patient, also in view of the increased cardiovascular risk to which he would be inevitably exposed. So the debate is still open, but if, on one hand, what the patient refers has its relevance, on the other hand we have to take into account the long term complications that a condition not more physiological might have in different organs and tissues.

Among the conditions that interfere with the sense of well-being of the patient in therapy with levothyroxine and normal TSH levels, you can also count the vitamin D deficit, which may be accompanied by fatigue, muscle fatigue and depression. This forum also discussed the opportunity of dosing vitamin D in these subjects, and, in the case of deficit, to establish an adequate replacement therapy.

Another element of debate was that relating to the need to perform a diagnostic process to highlight a possible malabsorption in case the therapeutic target is not reached following the administration of the theoretical dose of levothyroxine. The justified objection to this practice comes from the significant physiological individual variability of drug metabolism: absorption, speed degradation, clearance etc. Furthermore, each clinical parameter, by definition, can be misleading, so it could be sufficient to repeat the dosage and possibly minimally adjust the dose, rather than trigger a whole series of investigations also expensive and invasive, to be reserved, perhaps, to cases with more justified clinical suspicion. It should be clear, however, that the screening for malabsorption, in fact, provides initial tests both little invasive and expensive, such as the urea breath test, which, considering the high incidence of Helicobacter pylori infection in the general population, provide, in any case, clinically useful information, regardless the potential malabsorption. The more invasive investigations, such as endoscopy, may be reserved to the later stages of the diagnostic process.
From the discussion emerged also another problem: if it is true that the dose in the elderly should be reduced by 20% compared to the standard, how to resolve the conflict with the WHO guidelines which, instead, suggest increasing the dose in these patients in relation to the natural reduction of drug absorption with the increasing age? In fact, this discrepancy does not seem to exist, as in the elderly the reduction of the absorption is associated with a concomitant slowing down of T4 metabolism, which makes sure that the permanence time of the hormone in circulation is sufficient to ensure circulatory homeostasis.

An additional problem of the treatment of hypothyroidism in the elderly derives from the reduced ability of the body after a certain age to adapt to hormonal deficiency and to ensure tissue homeostasis of the thyroid hormone. Numerous studies also showed a physiological increase in the levels of TSH, “normal” in individuals of advanced age. The treatment of hypothyroidism, especially in the subclinical forms, may therefore be counterproductive because of the potential negative effects that even a slight excess of the hormone may cause in the elderly.

During the forum, it was also discussed at length the potential of research in the study of various polymorphisms of genes involved in the metabolism of thyroid hormones and the possible correlation between the exon sequences and values of TSH and FT4 in hundreds of patients, with the clinical practice.

An important and striking point, then, is the association between polymorphisms of some genes and T3 in serum: studies done in a small numbers of patients appear to demonstrate the existence of a significant correlation, but further studies are needed to demonstrate the real meaning.

In clinical practice, it is not uncommon to observe patients who present randomly elevated values of TSH, totally asymptomatic, with hormone free fractions normal, in absence of a family history of thyroid disease or autoimmunity. These cases could be an expression of a polymorphism at pituitary level as to explain the isolated increase of TSH? In part, maybe it is so, but at present there are no sufficient data available to confirm this. Another issue related to this subgroup of patients concerns the therapeutic approach. In general they have to be considered subclinical hypothyroid, but there is no consensus regarding the treatment, considering that some of them do not benefit from hormone treatment. Therefore, it would be more appropriate to carefully evaluate each case and decide in relation to the individual patient whether to treat.

They emerged also a number of ideas for the future of the research on the peripheral metabolism of thyroid hormone. It would be, for example, interesting to see what would happen if the gene for the deiodinase type 2 was inactivated not only in the thyroid but also in other organs, such as the hypothalamus. The issue is very complex. It has been demonstrated a conspicuous presence of D2 in tanyocytes, specialized cells of glial origin that act as stem cells, but it has not yet been possible to create tanyocytes deprived of D2.
Another stimulating topic concerns the correlation between the peripheral metabolism of thyroid hormone and the basal metabolism. Surely the identification of the response of D2 gene to cyclic AMP is one of the first molecular link identified between thyroid metabolism and adrenergic signal, but the metabolic determinants of this crosstalk are still unknown. Having more information about it could help, among other things, to understand, for example, the reason why there are different metabolic responses to the same stimulus, as may be the cold, in both sexes.

There is also need of studies to analyze the ways in which different drugs may interfere with thyroxine therapy in hypothyroidism. Often hypothyroid patients take many therapies that we know interfere with the metabolism of levothyroxine, such as absorption, binding to carrier proteins etc. We know still very little, on the contrary, on the possibility of these drugs to interfere with the activity of deiodinase; some, such as amiodarone and β-blockers have been shown effects in that regard, but the currently available data are somehow inadequate.

With regard to the possible clinical use of T3 sulfate, it was exposed the doubt about reintroducing a substance in the body that is physiologically eliminated. Indeed, administering T3 sulfate, there is no claim to mimic a physiological condition, and its metabolic effects have to be yet fully defined.

In conclusion, there are still many questions concerning the future of the hypothyroid patient and many of the suggestions received are certainly still to be explored. The importance of therapy customization (individually tailored dose) on individual hypothyroid patients is the message that must enter into the clinical practice in the daily contact with a disease which has a strong social significance such as hypothyroidism.
The analysis of data from both basic and clinical research and the future perspectives in therapy for hypothyroidism are the topics discussed and lively debated during the II Forum of Endocrinology organized by the Fondazione IBSA.

The Forum, which was attended by prominent international experts, provides interesting insights and is an useful contribution to achieve the goal desired by all: a timely and correct diagnosis and a more personalized and effective therapy in the future of the hypothyroid patient.