The role of thyroid hormones in depression

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Introduction
During the last 30 years a huge number of scientific articles have appeared on the subject of relationships between psychiatric disease and thyroid hormones. These studies have demonstrated the presence of numerous changes in the hypothalamo–pituitary–thyroid (HPT) axis, mainly in patients with depression, but also in patients with other psychiatric diseases. Simultaneously, many studies have been published on the possible therapeutic effects in depression of the hormones involved in the HPT axis.

Despite great efforts to standardize the classification of depression, this is still a less well-defined disease, possibly including several subtypes with different pathogenesis and biochemical abnormalities. Furthermore, the classification has changed over time, making it difficult to interpret previously published data. The most accepted classifications are the DSM-IV and ICD-10. These cover a spectrum from minor depression (neurotic depression) through major depression to melancholic (endogenous and psychotic) depression. Furthermore major and melancholic depressions are divided into unipolar and bipolar depressions, the latter also demonstrating episodes of mania. Rapid cycling bipolar psychosis is a subgroup of the bipolar depression, demonstrating four or more episodes in any year of observation.

The present review focuses on the concentrations of hormones of the HPT-axis seen in unipolar and bipolar depression, their relation to relapse after antidepressive treatment, and a survey of therapeutic trials with thyrotropin-releasing hormone (TRH), thyroid-stimulating hormone (TSH) and thyroid hormones, as well as the effect of the various drugs used in depression on the HPT-axis. The hypothesis is put forward that the changes seen in the HPT-axis during depression might be explained by cerebral serotonin deficiency, and that tri-iodothyronine (T₃) treatment, to some degree, can revert this deficiency.

Iodothyronines in depression
An overall picture of thyroid-related disturbances seen in depression is given in Table 1.

Thyroxine (T₄)
Serum T₄ levels, both total and non-protein-bound (free), are consistently found as normal to increased in groups of depressed patients (reviewed in 1, 2), approximately 25% of the patients having levels above the reference range. The different findings might be explained by the different severity of depression among the patients studied, since some studies have found a correlation between the severity of depression and serum T₄ levels (2). An additional explanation might be that the depressed patient is often in a state of semi-starvation, and thus may present changes in the HPT-axis similar to those seen in patients with nonthyroidal somatic illness, elevated serum T₄ and 3,3',5'-triiodothyronine (rT₃), and reduced serum T₃.

Turnover studies using radiolabeled T₄ in a small group of depressed patients have demonstrated that also the daily production rate (PR) of T₄ is significantly increased, by 30% (3). This finding contrasts with the unaltered T₄ PR found in groups of patients with different nonthyroidal illnesses (4). These studies used a noncompartmental kinetic approach for the evaluation of PRs. This method correctly estimates the PR of T₄. Increased PR of T₄ in depressed patients thus suggests that the thyroid gland is stimulated abnormally in the depressed patient.

The concentration of free T₄ in cerebrospinal fluid (CSF) seems relatively increased during depression, since recovery is followed by a reduction (5). The ratio between CSF and serum free T₄ levels has been found to be 0.6, both before and after recovery of depression due to electro-convulsive treatment (ECT) (5). This suggests that CSF T₄ concentrations follow systemic levels, and that transport of T₄ into CSF is restricted. An alternative explanation is that T₄ is taken up directly into brain tissues, and partly undergoes deiodination, thus yielding less T₄ into the CSF.

T₃
Serum T₃ levels in depressed patients are often found normal, but several studies have found reduced levels, typically in more severely depressed patients (1, 2). Serum T₃ levels are influenced by numerous factors which all may be present in the depressed patient: starvation, concomitant somatic illness and medication, and changes (increase) in cortisol levels. When present these factors all tend to decrease serum T₃ levels (4, 6). This makes interpretation of serum T₃ concentrations difficult. Serum free T₃ levels have been found both
normal and reduced (2). However, methods for the measurement of free T₃ in serum often give spurious, mainly reduced levels, in diseased patients (7). Ultrafiltration seems the method of choice (7, 8), and using this technique we have found unaltered free T₃ levels in depression (9). The daily PR of T₃ in unmedicated, moderately depressed patients has been studied using tracer turnover techniques, and T₃ PR was found normal (3). This was quite different from patients with various nonthyroidal illnesses, in whom a 40% reduction in T₃ PR has been demonstrated (4). The combination of an increased T₄ PR and an unaltered T₃ PR in depression suggests a reduced deiodination of T₄ into T₃, as also seen in nonthyroidal illness, but with more substrate (T₄) availability than in somatic nonthyroidal illnesses. However, correct interpretation of noncompartmental tracer studies is based on the assumption that the tracer injected is distributed freely into all tissue compartments (4). If anything, some degree of underestimation of T₃ PR might take place when performing T₃ kinetics in humans (4), but conflicting results are available on this issue (10).

The reduced conversion of T₄ into T₃ seen during depression (3) might be due to reduced deiodination enzyme activity. However, in which compartment of the human body this takes place is at present unknown. This could in theory be the brain, but unfortunately we are not aware of any data on intracerebral T₃ content or CSF levels of T₃ in depression.

**rT₃**

Changes in serum total as well as free rT₃ levels in depression seem to follow those seen for serum total and free T₄, respectively (2, 5, 11).

CSF levels of rT₃ have been studied in different types of depression, and have been found highest in the endogenous type (12). Similar to changes in serum, CSF free rT₃ concentrations seem to follow CSF free T₄ levels (5), but the ratio of CSF to serum free rT₃ has been found to be approximately 26, which is quite different from that of T₄ (0.6) (5). This ratio for rT₃ did not change after recovery from the depression, and this suggests that intracerebral concentration of rT₃ is high in humans, and that rT₃ in brain is mainly derived from local production from T₄. This enzymatic production seems not changed in depression, and this argues against rT₃ as a pathogenic factor in depression.

### TSH in depression

Previously, changes in circulating levels of TSH in depressed patients were evaluated using the TSH response to intravenous TRH, due to lack of sufficiently sensitive TSH assays. The response in TSH has usually been evaluated as the peak value minus basal TSH (\( \Delta_{\text{TSH}} \)). Normal \( \Delta_{\text{TSH}} \) to 200 μg TRH i.v. is typically 2–16 mU/l. Patients with depression, as a group, have reduced \( \Delta_{\text{TSH}} \) and approximately 25% have \( \Delta_{\text{TSH}} \) below 2 mU/l in endogenous (melancholic) depression (1). Using newer, and more sensitive TSH methods, basal TSH values correlate closely to peak TSH as well as \( \Delta_{\text{TSH}} \) (13–15). According to this, depressed patients seem to have some degree of reduced basal serum TSH, but within the normal range (13, 15). Basal as well as \( \Delta_{\text{TSH}} \) using a sensitive TSH assay have been studied in various types of depression, and those patients with endogenous depression had the lowest levels of both basal TSH and \( \Delta_{\text{TSH}} \) (13). \( \Delta_{\text{TSH}} \) being a mean of 3.6 mU/l.

The diurnal variation of serum TSH is, in healthy man, associated with a surge in TSH around midnight. In untreated depression this diurnal variation seems attenuated (16). After complete recovery from depression the diurnal TSH variation seemed to be re-established, but
Thyroid hormones in depression

In one study the interrelation between serum TSH concentrations and the daily production of T4 and T3 in endogenous depression was evaluated (3). Six depressed patients were compared with seven subjects with L-T4-treated hypothyroidism. These two groups had quite similar T4 and T3 PRs, both having an equal T4 PR/T3 PR ratio. Serum TSH level using a sensitive assay was a median 0.11 mU/l among L-T4-treated hypothyroid subjects, but 0.90 mU/l in depressed patients, significantly higher than in the L-T4-treated hypothyroid group (3). This suggests an inappropriate secretion of TSH (in relation to the elevated thyroidal production of T4) in patients with endogenous depression, compatible with some degree of central over-stimulation of the thyroid in the untreated depressed state.

The apparent paradox that serum TSH levels are found slightly decreased among depressed patients, and our statement of inappropriately elevated serum TSH levels in comparison to daily production of T4 and T3 (3), can be compared with the situation of chronic TRH stimulation (by oral administration) in man. In this experimental setting, TRH induced an initial increase in serum TSH, whereas later, at the time when serum T4 and T3 began to increase, serum TSH levels were normalized and were unable to respond to repeated TRH stimulation (18, 19). Thus chronic stimulation with TRH in man can provoke a pattern in serum levels of TSH, T4 and T3 similar to those seen in the depressed patient.

Chronic stimulation of the thyrotrhops in the pituitary during depression might be caused by TRH, since TRH levels have been found elevated in CSF in two studies (in total 31 patients) (20, 21), although normal levels were found in a third study including 17 patients (22). Serum TSH levels are also influenced by somatostatin, which inhibits the TSH release from the pituitary (23). Three studies have found that the CSF concentration of somatostatin is reduced during depression (24–26). This might contribute to an increase in serum TSH levels.

Two subgroups of the depression syndrome seem to present with a different pattern in the HPT-axis. One group consists of patients with bipolar depression (27, 28), and another group of patients is characterized by a depression resistant to treatment with tricyclic antidepressants (TCA) (29, 30). Both groups have a tendency towards slightly elevated basal serum TSH levels (approximately 20% have levels above the upper normal reference range), or exaggerated TSH response to TRH stimulation. These changes seem independent of previous or ongoing lithium treatment, and probably reflect some degree of thyroid insufficiency. Immunological mechanisms seem to play a role, since patients with bipolar depression have increased frequency of sera positive for antithyroid peroxidase (anti-TPO) antibodies (28). Whether this is an epiphenomenon to bipolar depression, or might be considered as one of several pathogenic factors, is at present unknown. Interestingly, postpartum changes in mood seem to be associated with the presence of anti-TPO antibodies and thyroid dysfunction (31). Thyroid hormone substitution might therefore be of potential benefit in alleviating depressive symptoms in the pueroerpal period. A preliminary study, in which L-T4 was administered immediately after delivery, in fact resulted in reduced depression score among the L-T4-treated women compared with placebo treatment (32).

Predictive value of changes of ΔTSH

Another interesting aspect of the TSH response to TRH is the possible predictive value of the changes before and after antidepressive treatment. It is a typical clinical finding that depressed patients, after recovery following ECT, demonstrate a high frequency of early relapse, typically within 6 months, if not treated prophylactically with antidepressive medication. We have repeatedly demonstrated that those patients with early relapse of endogenous depression have not normalized their reduced TSH response to TRH, despite clinical recovery. This has been demonstrated after treatment with ECT as well as sleep deprivation (i.e. nonpharmacological treatment modalities) (1, 33, 34). More than 80% of those patients with early relapse did not change their maximal TSH response to TRH (Δmax TSH). An unchanged Δmax TSH was defined as post-treatment Δmax TSH minus pre-treatment Δmax TSH (ΔΔmax TSH) less than 2 mU/l. By contrast more than 80% of those patients who remained clinically cured for more than 6 months had a ΔΔmax TSH exceeding 2 mU/l (1). Serum T4 as well as free T4 levels remained elevated in those patients with early relapse and continued reduced Δmax TSH (9). Thus a continued disturbance of the HPT-axis despite clinical improvement of endogenous depression suggests that the patient is not clinically cured. However, other studies in patients treated with antidepressive medication, rather than nonpharmacological treatment, have been unable to confirm these results (2).

In a small group of patients treated with amitriptyline we confirmed this lack of predictive value of the TRH test (33). Thus it is possible that the use of antidepressive medication might interfere with the HPT-axis on multiple sites, thus blurring the finding seen in the unmedicated, clinically cured patient. Alternatively, in the studies using TCA, the TRH tests have been performed with intervals too long to maintain a stable ΔTSH in patients not cured of depression.

Relationships between depression and thyroid hormones – a hypothesis

The pathogenesis of endogenous depression is not known, and is most probably multifactorial. The
The currently favored hypothesis is that a lack of serotonin in the brain has a central role (35). This hypothesis has generated the development of effective antidepressive drugs belonging to the group of serotonin re-uptake inhibitors. TRH seems under a constant inhibition by serotonin, and reduced intracerebral serotonin concentration will lead to increased TRH concentrations in brain tissue (reviewed in 23) (Fig. 1). As a consequence, TSH secretion will be stimulated. In addition CSF levels of somatostatin are reduced in depression. Since TSH secretion is under a constant inhibition of somatostatin (23), the consequence might be a further stimulated TSH secretion. Increased TSH levels result in enhanced thyroidal production of T₄ and T₃. Due to the thyroid–pituitary feedback, increased T₄ and T₃ levels tend to reduce serum TSH levels, reaching a new steady state usually within the normal range. The new TSH levels are, however, inappropriately elevated in relation to the increased T₄ production (3).

Acute as well as chronic T₃ treatment has been shown to increase the serotonin levels in the cerebral cortex of rats (36). In man, plasma serotonin levels correlate positively with T₃ concentrations (37, 38), and treatment of hyperthyroidism results in a reduction in serum serotonin levels (37). Brain serotonin levels in rats seem similarly affected, i.e. synthesis is reduced in hypothyroidism (39) and increased in hyperthyroidism (40, 41).

Thus the findings that reduced intracerebral serotonin concentrations lead to increased TRH and thereby to increased thyroid hormone levels, and that increased T₃ levels lead to an increase in brain serotonin, seem to constitute a classical feedback mechanism relevant to alleviation of depression.

Intracerebral T₃ seems mainly a result of local production by deiodination of T₄. In general, at least three deiodinating enzymes seem capable of affecting the turnover of T₄ and T₃ (reviewed in 42). The type-I deiodinase (D-I) results in both inner (5-) and outer (5₀-) ring deiodination, mainly of T₄, which thereby provides a circulating source of T₃ to the peripheral tissues. The type-II deiodinase (D-II) results in outer (5₀-) ring deiodination, mainly of T₄. This enzyme functions to regulate intracellular T₃ levels in those tissues where T₃ is most critical, such as brain and pituitary. In line with this, the enzyme activity is increased in hypothyroidism and reduced in hyperthyroidism. The type-III (D-III) enzyme results in inner (5-) ring deiodination, of both T₄ into rT₃, and T₃ into 3,3₀-T₂, and, in contrast to the D-II, but similar to D-I, its activity is increased in hyperthyroidism and decreased in hypothyroidism. Whereas all three enzymes are present in rat brain,:

![Figure 1 Decreased brain serotonin levels (serotonin (−)) activate (indicated by (+)) the HPT-axis, and increased T₃ (by endogenous as well as exogenous sources) increases brain serotonin levels (feed-back) – a hypothesis. D-II and D-III refer to iodothyronine deiodinases type II and III, respectively. 3,3₀-T₂: 3,3₀-diodothyronine.](image)
human brain tissues seem to contain only D-II and D-III enzymes (43). The properties of the D-II and D-III enzymes in rat and human tissues seem quite similar (43). Thus D-II activity increases T3 production in brain and pituitary, and consequently also the local production of serotonin. In addition, D-III activity might be expected to decrease the local concentration of T3 and indirectly of serotonin in brain tissues.

Studies of the influence of several known antidepressive drugs on rat D-II and D-III enzyme activities have revealed a similar pattern: lithium (44), des-imipramine (45), carbamazepine (46), and fluoxetine (47) have been shown to enhance the activity of the D-II enzyme resulting in increased local T3 concentrations in brain tissues. Also sleep deprivation of rats, which is a known treatment modality for depression, has been shown to increase the D-II activity (48). In contrast, lithium, carbamazepine, and fluoxetine have been shown to decrease the activity of the D-III enzyme (44, 46, 47), also resulting in increased local T3 concentration (for overview, see Fig. 1). Although this is an attractive pattern, a contradictory study has demonstrated that lithium decreases the activity of D-II in mouse neural and pituitary tissues (49). One must also be aware that the above mentioned changes are not necessarily found in the same regions of the brain, and furthermore it is not known whether these regions have any relevance to depression.

The consequence of the effect of the antidepressive drugs on the D-II and D-III enzyme activities is an increased local T3 concentration in brain tissues and, as discussed above, this might increase local serotonin concentration, and thus might form an additional biological basis for the positive effect of these drugs in alleviating the depressive state.

Thus the hypothesis that a major pathogenic factor for depression is serotonin deficiency is sufficient to explain the changes seen in the HPT-axis. However, another hypothesis, put forward many years ago, has been that the depressed patient suffers a state of local hypothyroidism in the brain. This hypothesis, originally based on the similarities of symptoms seen in hypothyroidism and depression, seems supported by the finding that T3 treatment alleviated the depressive symptoms (50). Reduced T3 content in brain tissues leads to serotonin deficiency in brain tissues, and thus is sufficient to explain the findings in the HPT-axis seen in depression (Fig. 1).

Both hypotheses can explain that (i) patients with thyroid insufficiency are especially sensitive to develop depression (51, 52). (ii) T3 treatment is beneficial also in euthyroid, depressed patients, (iii) pharmacological stimulation of the D-II and inhibition of the D-III brain enzyme could be expected to have a beneficial effect on the depression, and (iv) T3 treatment might be at least as effective as T4 treatment. Only the serotonin deficiency hypothesis might explain the low normal serum TSH and low Δmax TSH seen during depression, and that patients with continued low Δmax TSH after apparent recovery from depression are subject to early relapse. This parameter can thus be regarded as a marker of continued active disease.

Another fruitful hypothesis developed for the biological explanation of depression suggested that depression was due to a state of relative brain deficiency of catecholamines, especially norepinephrine (53). This has formed the basis of the development of another group of useful antidepressive drugs, norepinephrine re-uptake inhibitors. In 1981 Whybrow & Prange Jr (54) hypothesized that thyroid hormones, by enhancing beta-adrenergic receptor function, promote transmission in central noradrenergic pathways and accelerate recovery. Later, the same group (55) demonstrated that T3 levels in rat brain synaptosomes are much higher than whole brain levels, and that T3 but not T4 can be released from depolarized synaptosomes. Recently Rozinov & Dratman (56) found increased T3 concentrations in nuclei and projection sites of rat central noradrenergic systems. These studies suggest that T3 may play a neuromodulatory or neurotransmitter role in the noradrenergic central nervous system. Norepinephrine stimulates the release of both TRH and TSH (23), and therefore norepinephrine deficiency cannot explain the changes in the HPT-axis seen in depression, especially the increased CSF TRH levels. However the hypothesis is fully compatible with the beneficial effect of T3 treatment in depression.

Treatment of depression with hormones of the HPT-axis

The similarities between symptoms seen in depression and untreated hypothyroidism have lead to several clinical trials in which thyroid hormones have been given to depressed patients, either alone or in combination with other antidepressive treatment modalities.

Treatment without other treatment modalities

TRH In the 1970s a number of studies demonstrated a mild but transient improvement in mood in depressed patients treated with TRH for a few days or at most 3 weeks. Later other studies including controlled double-blind designs, failed to demonstrate any beneficial effect of TRH (reviewed in 57).

T3 T3 given as the only drug to depressed patients has been evaluated in only two studies. Feldmesser-Reiss (50) treated 24 patients with depression or melancholia with 10–15 μg T3 daily, and observed considerable improvement in 10 patients. Wilson et al. (58) gave 25 μg, increasing to 62.5 μg, of T3 daily for 9 days to nine patients, which apparently was as effective on depression as imipramine, as evaluated by the Hamilton rating scale. Imipramine plus T3 had no additional effect. Unfortunately this otherwise randomized, double-blind study had no placebo group. Thus in
these two studies T₃ given alone seems to have an antidepressive effect, but placebo-controlled studies are lacking.

**T₃ and T₄** We are not aware of any studies using TSH or T₄ as the only treatment of depressed patients.

**Treatment in combination with other treatment modalities**

Several studies have evaluated the effect of the hormones of the HPT-axis. These studies can be separated into two typical designs: (i) either combination therapy with thyroid hormone and some kind of antidepressive treatment from the beginning, or (ii) adjunction of thyroid hormones to patients refractory to conventional antidepressive medication (nonresponders).

**Initial combination therapy with T₃ or TSH in depression** Concerning the initial combination therapy, only the effect of T₃ has been evaluated. Repeatedly in randomized, double-blind placebo-controlled studies, 20 to 50 µg T₃ per day in combination with TCA has been shown to shorten the period of depression as compared with TCA alone (59–62), but the final number of recovered patients was not influenced (59, 60, 62). However, in two other well-conducted trials, but with fewer patients, no additional effect of T₃ with TCA could be demonstrated (63, 64). The studies demonstrating a beneficial effect of addition of T₃ to TCA included in total 112 patients, whereas the two negative studies included only 29 patients. Furthermore the majority of patients studied were females.

The effect of T₃ does not seem to be due to changed metabolism of TCA, and might thus be regarded as an independent factor (65).

Unfortunately no studies are available concerning the effect of the addition of T₃ to the modern serotonin re-uptake inhibitory drugs.

In one study T₃ or placebo was given to patients receiving ECT (66). Patients treated with T₃ needed significantly fewer number of ECTs, and demonstrated less damage to memory functions, in comparison to the placebo group.

One study demonstrated a beneficial effect of TSH over placebo (in a randomized, double-blind fashion) when given to nine patients as intramuscular injections (10 units of TSH on day 1 and day 8) in adjunction to TCA (67). This study has unfortunately not been repeated, but it seems likely that the effect of TSH is due to an increased thyroid secretion of T₄ and T₃.

**T₃ and T₄ in refractory depression (nonresponders)** T₃ treatment of the depressed patient resistant to TCA was first reported beneficial in an open study without a placebo group (68). Since then the positive effect of 25–50 µg T₃ daily as an adjunctive therapy has been confirmed in many studies, including a recent meta-analysis (69), which included four randomized double-blind trials (in total 69 patients) and three unblinded studies using historical controls (in total 185 patients). This analysis also discussed in detail those studies which did not find any beneficial effect of T₃. Overall the addition of T₃ to TCA increased the response rate significantly from 24 to 57%.

In another randomized double-blind study Joffe et al. (70) compared the ability of T₃ and lithium to convert nonresponders to TCA into responders, i.e. reduce the depressive symptoms on a Hamilton Rating Scale. T₃ was equally effective as lithium, and both drugs were superior to placebo.

No formal studies (only case reports) concerning the effect of the addition of T₃ in depressed patients resistant to therapy with monoamine oxidase inhibitors or serotonin re-uptake inhibitors exist.

One study has compared the effect of physiological doses of T₄ (150 µg daily) and T₃ (37.5 µg daily) as adjunctive therapy to nonresponders after 4 weeks on TCA. They found that T₃ was more effective than T₄ (71). Although randomized and double-blind, no placebo group was included, thus it is uncertain whether T₄ is superior to placebo.

Interestingly, an open, uncontrolled trial on nine L-T₄-substituted hypothyroid patients (median 150 µg daily) with depression and no response to TCA treatment, reported that seven of these patients recovered from depression after the addition of T₃ (72), suggesting a decreased deiodination of T₄ to T₃ in brain.

In another open trial on six nonresponding patients (to TCA) with nonrapid cycling bipolar affective disorders, Baumgartner (73) gave supraphysiological doses of T₄ (250–500 µg daily) resulting in a reduced number of relapses. The results of this preliminary study are also compatible with a reduced deiodination of T₄ to T₃.

In the rapid cycling form of bipolar affective disorders, T₄ adjunctive therapy has, in an open design without placebo controls (in total eight patients), been found to reduce both manic and depressive symptoms as well as number of relapses when given in supraphysiological doses (up to 500 µg daily) (74, 75).

These studies are all compatible with an inhibition of the D-II deiodinase or a stimulation of the D-III deiodinase in brain tissues (resulting in a reduced local T₃ concentration) in affective disorders.

**Antidepressive drugs – effect on thyroid hormones**

Changes in thyroid hormone and TSH levels during treatment of depressed patients with TCA and serotonin re-uptake inhibitors seem a result of the underlying psychiatric disorder, or its epiphenomena, i.e. increased...
serum cortisol and starvation, and not due to a direct effect of the drug on hormone secretion, plasma protein-binding and turnover (1, 17, 76).

In contrast lithium inhibits the secretion of thyroid hormones from the thyroid gland, resulting in an increase in serum TSH (77, 78) and thyroid auto-antibodies (79), which in susceptible patients might lead to overt hypothyroidism (79). This effect of lithium might result in a decreased supply of thyroid hormones to the brain, which according to the presented hypothesis is inconvenient. However the antidepressive effect of lithium has been demonstrated to be positively correlated to serum levels of T3 (80, 81) and negatively to serum TSH levels (82). The effects of lithium on the deiodinases D-II and D-III with the result of local increase of T3 (see above) might counteract the inappropriate effect of lithium on the thyroid hormone secretion.

Carbamazepine, an anti-epileptic drug, which also has some antidepressive effect, influences the HPT-axis in a complex manner. In addition to the earlier mentioned effect on the brain deiodinases (Fig. 1), carbamazepine seems to enhance the hepatic (D-I) deiodination of both T4 and T3 (83). Thus these patients often present with low to normal serum levels of T4 and T3, but normal TSH levels (83), suggestive of some inhibitory effect on the thyrotrhods as well.

Conclusions

It is hypothesized that the changes seen in the HPT-axis in untreated depression might be explained partly by a cerebral serotonin deficiency. These changes tend to increase serum levels of T3, which subsequently might increase serotonin content in the brain. Although this hypothesis is attractive, it is based on few and small studies, which clearly need confirmation.

Subjects with thyroid insufficiency seem especially susceptible to the development of depression.

Well established antidepressive treatment modalities influence the brain deiodinases D-II and D-III, which are responsible for the local T3 production in brain tissues, in a manner leading to increased T3 concentrations. Defects of the brain deiodinases might be a pathogenic factor in depression.

While the effect of the addition of physiological doses of T3 to TCA is well documented in depression, confirmatory studies are needed concerning the effect of T3 given alone, and T3 given in addition to ECT.

In refractory depressions, the addition of T3 to TCA has been shown to be equal to the addition of lithium, and superior to the addition of placebo or T4, to TCA in alleviating depressive symptoms. The superiority of T3 compared with T4 as adjunctive therapy must be further evaluated.

No studies exist on the effect of T3 as adjunctive therapy to the serotonin re-uptake inhibitors, nor on the effect of T3 in rapid cycling depression.

The effects of thyroid hormones, TSH and TRH, on the depressive syndrome have been extensively studied during the past three decades. Despite the fact that many of these studies are more than 20 years old, many fulfill modern criteria of good clinical science, in terms of using the randomized, double-blind placebo-controlled design. Throughout this review, we have made efforts to describe the design of the studies cited. However, the potential weaknesses of these studies is the relatively small number of patients enrolled in the different treatment modalities. This is especially relevant, since the depressive syndrome is a heterogenous group of disorders affecting a number of subjects; and furthermore the diagnostic criteria have changed over the years. Therefore previous findings should be re-evaluated, using modern scientific principles, and including a large number of patients. Furthermore, the effect of especially T3 in combination with various antidepressive treatment modalities (TCA, serotonin re-uptake inhibitors, ECT, sleep deprivation) should be studied.

The depressive syndrome is most likely a pathogenically heterogenous group. Patients are characterized phenomenologically, not by biological markers. Even phenomenologically identical groups might harbor different pathogenic subgroups of depression. Therefore future studies should also focus on identification of biological abnormalities in those patients who benefit from T3 treatment, in order to identify in which subgroups T3 treatment should be used.

In the light of the substantial amounts of data on the beneficial effect of T3 in depression, the restricted usage of T3 in this setting is surprising. T3 given in physiological doses is cheap and the monitoring is easy due to the new sensitive TSH assays. However, precautions should be taken in patients with concurrent heart disease, since T3 may sensitize the heart to the arrhythmogenic effects of TCA. Suicidal patients especially seem to be exposed if they take overdoses of T3 and TCA in combination.

References

3 Kirkegaard C, Korner A & Faber J. Increased production of thyroid hormone inappropriately elevated serum thyrotropin levels in endogenous depression. Biological Psychiatry 1990 27 472–476.
6 Doctor R, Krenning EP, de Jong M & Hennekamp G. The sick euthyroid syndrome: Changes in thyroid hormone serum


9 Kirkegaard C & Faber J. Influence of free thyroid hormone levels on the TSH response to TRH in endogenous depression. Psychoneuroendocrinology 1986 11 491–497.


Thyroid hormones in depression


51 Cowdry RW, Wehr TA, Zis AP & Goodwin FK. Thyroid abnormalities associated with rapid-cycling bipolar illness. *Archives of General Psychiatry* 1983 40 414–420.


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