Identifying hypothyroidism’s psychiatric presentations

Is the thyroid causing your patient’s symptoms?

Thomas D. Geraciotti, Jr. MD
Research service
Cincinnati Veterans Affairs Medical Center
Department of psychiatry
University of Cincinnati College of Medicine

Hypothyroidism—even when occult or subclinical—can cause subtle or frank changes in energy, mood, anxiety level, or cognition. Some patients’ affective symptoms remit with thyroid hormone replacement or with antidepressants only after a euthyroid state is established.

To help you recognize hypothyroidism in patients presenting with psychiatric illnesses and provide effective treatment, this article describes:

• hypothyroidism’s signs and symptoms
• primary and subclinical hypothyroidism, thyroiditis, central hypothyroidism, and thyroid hormone resistance
• laboratory screening for thyroid dysfunction in patients with psychiatric symptoms.

OVERLAPPING CLINICAL SIGNS
Thyroid hormone is required for the metabolic activity of every cell in the body. When patients experience symptoms related to abnormal functioning of the hypothalamic-pituitary-thyroid axis (Figure), psychiatrists often are the first professionals they consult.

Diagnosis of thyroid disorders is based on biochemical and clinical data (Box 1, page 100), which might not be congruent. Clinical symptoms of hypothyroidism, for example, are notoriously variable. Severe biochemical hypothyroidism may be associated with mild clinical symptoms, whereas mild biochemical hypothyroidism may be associated with severe symptoms.

Patients with thyroid disturbance and psychiatric symptoms most often are diagnosed with a depressive-spectrum syndrome. Most common are:

• atypical depression (which may present as dysthymia)
• bipolar spectrum syndrome (including manic-depression, mixed mania, bipolar depression, rapid-cycling bipolar disorder, cyclothymia, and premenstrual syndromes)
• borderline personality disorder
• or psychotic disorder (typically paranoid psychosis).

Psychiatric symptoms of hypothyroidism (Table 1, page 105) are often prominent or even primary. Patients commonly show:
The degree of impairment may depend on the patient’s normal functional level. For example, a well-educated patient of mine with thyroiditis-related hypothyroidism reported word-finding...
difficulties. Instead of asking her husband to take a bottle of wine from the rack, she asked him to take a bottle of wine from the “thing.”

Loss of vitality, fatigue, lethargy, hypersomnia (especially if sleep apnea is present), and depressed mood also are commonly seen.

**Depressive symptoms.**

Hypothyroid patients usually meet several criteria for a major depressive episode—such as concentration difficulties, lassitude, low libido, and sometimes pessimism or sadness—and symptoms improve after sustained thyroid hormone replacement therapy.¹ Women with mild hypothyroidism who screen negative for a psychiatric syndrome show statistically significant mood improvement and improved verbal fluency after 6 months of levothyroxine replacement therapy.¹

In some patients with no clear evidence of a biochemical or clinical thyroid disorder, mood symptoms nevertheless respond to thyroid hormone augmentation of antidepressants.¹

**Anxiety symptoms.** Occasionally thyroid dysfunction is seen in patients with anxiety disorders, including panic disorder, agoraphobia, social phobia, performance anxiety, post-traumatic stress disorder, and generalized anxiety disorder.⁴

It may seem counterintuitive, but hypothyroidism is probably as common as hyperthyroidism in extremely anxious patients. Both hypothyroidism are seen much more often in patients with panic-level anxiety than in the general population.

In a sample of 144 consecutive female psychiatric patients with a lifetime history of panic disorder and/or agoraphobia:

- 27% had a history of thyroid disorder
- 17% had hypothyroidism
- 8% had hyperthyroidism.¹

**Hypothyroid symptoms.** Associated symptoms of hypothyroidism (Table 2, page 105) may include cold intolerance, lack of or reduced perspiration, dry skin, constipation, lethargy, psychomotor slowness, and subjective paresthesias and muscle pains.

Edema is often present. The face typically is swollen or “puffy” in the morning, but by evening the lower legs (and not the face) are edematous.

Deep tendon reflexes usually relax slowly after initial stimulation. Vascular resistance is increased, but hypertension is not usual. Noradrenergic systems become more active in a compensatory, counter-regulatory manner; however, bradycardia—when present—is sometimes profound. Weight gain can occur but often is conspicuously absent.

continued on page 105

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**Box 1**

**Thyroid dysfunction: Psychiatric patients show increased risk**

**Psychiatric patients**—particularly those with mood disorders—are more likely to exhibit biochemical evidence of frank or subclinical hypothyroidism, hyperthyroidism, and autoimmune thyroiditis than the general population. In a study of 17,533 Americans,¹ approximately 12% had thyroid abnormalities and 80% of these were hypothyroid. An additional 17% of women and 9% of men tested positive for antithyroid antibodies.

Biochemical hypothyroidism was defined as TSH >4.5 mIU/L, with low total T4, and subclinical hypothyroidism as TSH >4.5 mIU/L with normal T4 levels. Population studies and other data have led some endocrinologists to regard serum TSH levels >2.5 mIU/L as abnormally high.
Severe hypothyroidism presents with paradoxical tremendous agitation, paranoia, and aggressiveness. The skin is leathery, and facies are characteristically rough. Myxedema is fairly common, even in high-functioning patients. I have seen only one case of so-called “myxedema madness;” the female patient’s hyperarousal, yelling, cursing, grossly poor cognitive ability, and loosely conceived paranoid delusions are unforgettable.

Galactorrhea (related to hyperprolactinemia) can be a symptom of severe hypothyroidism, presumably from increased hypothalamic thyrotropin-releasing hormone (TRH) drive. TRH is the main known secretagogue for pituitary prolactin secretion. Infertility, oligomenorrhea, or amenorrhea could be part of the hypothyroid clinical picture.

Other symptoms. Macroglossia and hypotrophy of the uvula are possible; in a recent report, dysarthria resulting from these oral changes was the only presenting symptom of a hypothyroid man.6 Dysarthria promptly corrected after levothyroxine replacement.

Hypothyroidism is a primary cause of central sleep apnea caused by dysfunction of ventilatory control and/or reduction in airway aperture.

**Hypothyroidism’s psychiatric signs and symptoms**

<table>
<thead>
<tr>
<th>Cognitive changes</th>
<th>Impaired memory, psychomotor slowing, reduced attention span</th>
</tr>
</thead>
<tbody>
<tr>
<td>Vegetative symptoms</td>
<td>Hypersomnia, sleep apnea, fatigue, lethargy, apathy, anergia, low libido</td>
</tr>
<tr>
<td>Mood changes</td>
<td>Depression, mood instability, mania, anxiety</td>
</tr>
<tr>
<td>Other</td>
<td>Psychosis</td>
</tr>
</tbody>
</table>

**Hypothyroidism’s other signs and symptoms**

<table>
<thead>
<tr>
<th>Metabolic</th>
<th>Low basal body temperature/cool skin, diminished perspiration, weight gain or difficulty losing weight</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cardiovascular</td>
<td>Bradycardia, dizziness</td>
</tr>
<tr>
<td>Dermatologic</td>
<td>Dry, rough, or scaly skin; brittle nails; coarse or thinning hair (especially in women); pallor; dependent edema; myxedema/skin mucinosis (classically pretibial)</td>
</tr>
<tr>
<td>Digestive</td>
<td>Nausea, constipation, enlarged tongue</td>
</tr>
<tr>
<td>Reproductive</td>
<td>Oligomenorrhea, amenorrhea, infertility, miscarriage, delayed ejaculation</td>
</tr>
<tr>
<td>Musculoskeletal and peripheral nervous system</td>
<td>Muscle cramps, joint pain, paresthesias, numbness, weakness, reduced exercise tolerance, delayed ankle reflex</td>
</tr>
<tr>
<td>Sensory</td>
<td>Upper eyelid drooping, dysarthria, hoarseness, diminished hearing, diminished taste (hypogustia)</td>
</tr>
</tbody>
</table>

**CAUSES OF THYROID DISORDERS**

**Primary hypothyroidism** results from thyroid gland failure. The many causes include iodine deficiency, but most cases of adult-onset or acquired hypothyroidism are attributed to autoimmune thyroiditis (see page 110). In primary hypothyroidism, replacement begins with a single oral dose of levothyroxine; subsequent adjustments are made with T4/TSH levels.
Hypothyroidism

continued from page 105

**Box 2**

**When to try thyroid hormone therapy for psychiatric symptoms**

**Target psychiatric symptoms** for prescribing replacement thyroid hormone are depression; mood cycling or instability; low energy, fatigue, or lethargy; cognitive impairment (Table 1, page 108); and psychosis, if present. Because these symptoms are not specific to thyroid dysfunction, institute thyroid hormone only when biochemical evidence of compromised or suboptimal thyroid function is also present.

**Exceptions** to this rule may include patients with target psychiatric symptoms and:

- other hypothyroidism symptoms (Table 2, page 105)
- circulating T3, free T3, or free T4 levels near the low margin of the normal range and/or TSH >3.5 to 4.0 mIU/mL.
- a history of inadequate response to psychopharmacologic interventions
- symptoms of and a strong family history of thyroid disease.

The decision to treat subclinical hypothyroidism with thyroid hormone replacement is less controversial in psychiatry than in endocrinology. Psychiatric patients with subclinical hypothyroidism—especially those with incomplete responses to psychotropic therapy—should usually be treated with thyroid hormone (Box 2).

Free T3 levels in the lower 20% of the laboratory’s normal range are cause for pause in a patient with a mood or psychotic disorder and any of hypothyroidism’s clinical stigmata, even if thyroxine and TSH concentrations are normal.

**Thyroiditis** is characterized by thyroid gland inflammation. The thyroid may be painful or nonpainful, enlarged, fleshy, goitrous, normal in size, or atrophic and fibrotic (especially late in the course).

Postulated precipitants include viruses and other infectious agents, vaccines, iodine excess, lithium therapy (Box 3), tobacco smoke, environmental chemicals or toxins, irradiation, and—arguably—cortically-mediated (psychological) stress in vulnerable individuals.

Clinically, thyroiditis syndromes often have a long prodromal phase, wax and wane in severity, have an insidious and sometimes silent course, and can be serially associated with hyperthyroidism (especially early in the course), euthyroidism, or hypothyroidism (especially late in the course). Most thyroiditis syndromes appear to resolve spontaneously, but many become chronic or show evidence of subtle thyroid dysfunction years after the first occurrence or diagnosis.

Most presentations are nonspecific; symptoms may be limited to lethargy, fatigue, and depression. Increased antithyroid antibody titers have been linked with psychotic and depressive syndromes in borderline personality disorder.
In early thyroiditis, thyroxine and triiodothyronine secretion is often elevated, with low or suppressed TSH. However, antithyroid antibody production is associated with a significantly increased risk of eventual subclinical or frank hypothyroidism. Permanent hypothyroidism develops eventually in at least one-half of women with histories of postpartum thyroiditis.\textsuperscript{11} \textbf{Central hypothyroidism} stems from TSH deficiency. Both pituitary thyrotrophic failure and hypothalamic failure—secondary and tertiary hypothyroidism, respectively—are considered central (or “secondary”). Hypothyroidism of pituitary and hypothalamic origins are lumped together as “central” because it is often very difficult to differentiate these pathologies. \textbf{Thyroid hormone resistance}, in which end-organ or cellular resistance to thyroid hormone signals is seen, is an increasingly recognized syndrome in clinical medicine. The typical case is an euthyroid or hypothyroid individual with elevated T4 and T3 and nonsuppressed or even frankly elevated TSH. Inappropriately elevated TSH combined with high thyroid hormone levels also can be seen in TSH-secreting pituitary tumors, although the clinical picture in this case is one of hyperthyroidism.

Thyroid hormone resistance ranges from euthyroid and clinically transparent to profoundly hypothyroid, and different organs in the same patient may show different sensitivities to thyroid hormone. Clinical features vary, depending on the strength of thyroid hormone resistance.

Early emergence of resistance (as would be expected in someone with an inherited thyroid hormone receptor abnormality) leads to developmental problems, including:

- short stature
- mental or learning disabilities (including attention deficits).

In a study of 18 families with strong history of generalized resistance to thyroid hormone, 70\% of affected children met diagnostic criteria for attention-deficit/hyperactivity disorder.\textsuperscript{12}

\textbf{Box 3} \textbf{Long-term lithium therapy linked with hypothyroidism risk}

\textbf{Patients taking lithium} for mood stabilization will likely need supplemental thyroid hormone eventually because lithium is thyrotoxic. Also, patients with bipolar mood symptoms often have coexisting thyroid abnormalities, and giving supraphysiologic thyroid hormone dosages sometimes converts those who do not respond to mood stabilizers into responders.

Thyrotropin concentrations increase within 1 day after patients start taking lithium carbonate, but without commensurate increases in T3 or T4. More often, T3 and T4 concentrations decrease in the presence of lithium. Among 150 patients maintained on lithium for 10 years, hypothyroidism, autoimmune thyroiditis, or goiter developed at rates of 1.7\%, 1.4\%, and 2.1\% per year, respectively. The study authors suggested that long-term lithium may increase the risk for hypothyroidism in women and in patients with thyroid autoimmunity.\textsuperscript{13,14} This rhythm implies that the most accurate TSH measurement may be obtained by drawing blood in the

\textbf{LABORATORY SCREENING} \textbf{TSH and thyroid hormones}. The basic thyroid screen is a combination of serum levels of TSH (“sensitive TSH”) and free thyroid hormones.

TSH has a circadian rhythm, with a nocturnal surge amounting to a 50\% to 200\% increase over daytime levels, beginning at around 6 to 8 PM. Peak TSH pulsatile activity and levels are seen after sleep begins—usually after midnight—with trough levels and fewest TSH pulses in late morning and early afternoon.\textsuperscript{13,14} This rhythm implies that the most accurate TSH measurement may be obtained by drawing blood in the
morning before 9 AM. I have seen subclinical hypothyroidism missed when clinicians relied on afternoon TSH levels.

In general medicine, TSH alone frequently is used as a routine screening tool. In psychiatric practice, however, I recommend supplementing TSH with free T3 and free T4 because thyroid system dysfunction is frequent in psychiatric syndromes. If laboratory costs are a concern, free T4 with TSH usually suffices for initial screening.

Although the unbound, free fractions of T3 and T4 are of primary interest, total T4 and total T3 are necessary to assess the thyroid gland’s synthetic capacity. When clinical evidence suggests abnormal thyroid hormone function, order repeated or serial biochemical testing. Marginal biochemical results also mandate repeat thyroid function studies, expanded to include:

- total thyroid hormone concentrations (ideally T4 and T3)
- antithyroid antibodies
- serum cholesterol
- prolactin.

**Antithyroid antibodies.** Obtain antithyroglobulin and antithyroid microsomal antibody titers if:

- thyroid hormone indices are abnormal or marginal—in either direction
- or the patient now has, has had, or has a family history of autoimmune-mediated symptoms, such as lupus erythematosus or rheumatoid arthritis.

Negative or low antithyroid autoantibody titers do not rule out thyroiditis as a cause of hypothyroidism, as these titers are most likely to be generated during periods of active inflammation.

Many autoantibodies react with thyroid-related elements. Most clinical laboratories can quantify antibodies directed against thyroid peroxidase—also called antimicrosomal antibodies—and thyroglobulin. The presence of these antibodies is associated with thyroid inflammation and a risk of progression to thyroid failure and hypothyroidism.

**Other laboratory findings** of hypothyroidism include hypercholesterolemia, mild hyperprolactinemia, and types of anemia (including iron-deficiency anemia with low ferritin levels).

**References**


continued from page 112


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The Neuroleptic Malignant Syndrome Information Service (NMSIS) announces a competition to recognize promising new investigators, based on a scholarly paper addressing “New insights on psychotropic drug safety and side effects.”

Consistent with its mission to advance pharmacotherapy and patient safety, NMSIS offers these awards to promote education and research by early career psychiatrists. Two prizes of $2,500 and $1,500 will be awarded toward travel costs to attend the American Psychiatric Association annual meeting in May 2007.

• Papers should address specific issues related to the award theme and be no longer than 15 double-spaced, typed pages.
• Literature reviews, case reports, or original studies that are not in press or published are acceptable.
• Primary author must be a student, resident, fellow, or junior faculty member at or below the rank of assistant professor.
• Papers will be judged on originality, scholarship, relevance, and methodology.

Submit paper and the primary author’s curriculum vitae to Diane Van Slyke, 11 East State St., Sherburne, NY 13460, fax 607-674-7810, or via e-mail to diane@nmsis.org. Deadline is Feb. 1, 2007.

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