Management of Graves’ Disease during Pregnancy: The Key Role of Fetal Thyroid Gland Monitoring

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Background: Fetuses from mothers with Graves’ disease may experience hypothyroidism or hyperthyroidism due to transplacental transfer of antithyroid drugs (ATD) or anti-TSH receptor antibodies, respectively. Little is known about the fetal consequences. Early diagnosis is essential to successful management. We investigated a new approach to the fetal diagnosis of thyroid dysfunction and validated the usefulness of fetal thyroid ultrasonograms.

Methods: Seventy-two mothers with past or present Graves’ disease and their fetuses were monitored monthly from 22 wk gestation. Fetal thyroid size and Doppler signals, and fetal bone maturation were determined on ultrasonograms, and thyroid function was evaluated at birth. Thyroid function and ATD dosage were monitored in the mothers.

Results: The 31 fetuses whose mothers were anti-TSH receptor antibody negative and took no ATDs during late pregnancy had normal test results. Of the 41 other fetuses, 30 had normal test results at 32 wk, 29 were euthyroid at birth, and one had moderate hypothyroidism on cord blood tests. In the remaining 11 fetuses, goiter was visualized by ultrasonography at 32 wk, and fetal thyroid dysfunction was diagnosed and treated; there was one death, in a late referral, and 10 good outcomes with normal or slightly altered thyroid function at birth. The sensitivity and specificity of fetal thyroid ultrasound at 32 wk for the diagnosis of clinically relevant fetal thyroid dysfunction were 92 and 100%, respectively.

Conclusion: In pregnant women with past or current Graves’ disease, ultrasonography of the fetal thyroid gland by an experienced ultrasonographer is an excellent diagnostic tool. This tool in conjunction with close teamwork among internists, endocrinologists, obstetricians, echographists, and pediatricians can ensure normal fetal thyroid function. (J Clin Endocrinol Metab 90: 6093–6098, 2005)

GRAVES’ DISEASE IS present in about 0.2% of pregnancies. Neonatal hyperthyroidism has been extensively studied, although clinical hyperthyroidism seems to occur in only 1% of neonates born to mothers with Graves’ disease (1–5). The relative abundance of information available on neonates contrasts sharply with the paucity of data on the fetal effects of maternal Graves’ disease. Evidence indicates a need for moving diagnostic efforts from the neonatal to the fetal period, to prevent fetal death or permanent neurological impairment (6–9). Fetal thyroid dysfunction precedes neonatal hyperthyroidism (2, 3, 9–11). The fetal thyroid gland starts secreting thyroid hormones at about wk 12 of development, and fetal TSH receptors become responsive to TSH and to TSH receptor antibodies around wk 20 (10). Whereas maternal T4 usually crosses the placenta in only minimal (but crucial for fetal development) amounts, and TSH does not cross the placenta at all, maternal TSH receptor antibodies freely cross the placenta and can cause overstimulation of the fetal thyroid gland during the second half of pregnancy (11). Because antithyroid drugs (ATDs) cross the placenta, administration of ATDs to the mother improves the fetal and neonatal prognosis (8, 9, 12–17). However, ATD administration to the mother can also jeopardize fetal development by inducing fetal hypothyroidism.

Few data on fetal abnormalities and outcomes are available to internists, endocrinologists, and obstetrician for devising guidelines on the management of pregnant women with Graves’ disease (18). Published guidelines deal with both managing Graves’ disease in the woman and predicting the risk of neonatal thyrotoxicosis. The objective of the present study was to assess fetal ultrasonography as a noninvasive tool for detecting fetal thyroid dysfunction, determining the appropriate treatment, and monitoring treatment effects during pregnancy in patients with active or treated Graves’ disease. The pregnant women were included and monitored prospectively by a multidisciplinary hospital team.

Subjects and Methods

We prospectively included 72 pregnant women (72 fetuses) managed between 1999 and 2002 at Robert Debré Teaching Hospital (Paris, France) who were not included in our previous study (17). The inclusion criterion was past or current history of Graves’ disease diagnosed by an
for 102 such women). Therefore, FT levels did not differ significantly from the upper limit of the normal range of pregnant women (our own data). Therefore, FT levels did not differ significantly between the two groups of pregnant women (see below) or between women with and without a history of thyroidectomy. Once a month, starting at 22 wk gestation (WG), fetal heart rate (FHR) was recorded, and a fetal ultrasound scan was performed for measurements of thyroid gland size (diameter and circumference; Fig. 1A), determination of fetal growth parameters, and evaluation of fetal bone maturation. An EVB 525 variable focus ultrasound machine (Hitachi, Hialeah, FL) with a 3.5-MHz sector transducer was used for all fetal sonograms.

Inclusion criteria were defined as a thyroid circumference equal or superior to the 95th percentile for gestational age, according to the normative data established by Ranzini et al. (21) and to our own unpublished normogram established by E.V. in 250 patients, which is consistent with Ranzini’s curves (data not shown). When fetal goiter was found, color flow Doppler of the thyroid was performed as previously described with a velocity of 13 cm/sec (17, 22); a Doppler signal throughout the gland was considered suggestive of fetal hyperthyroidism, and a Doppler signal confined to the periphery of the gland was considered suggestive of fetal hypothyroidism (Fig. 1, B and C) (17, 22).

Bone maturation was evaluated on the sonogram obtained at 32 WG. Normally, the distal femoral ossification center is undetectable before 28 WG, is dot-like around 32 WG, is smaller than 3 mm before 33 WG, and is consistently visible after 35 WG. Accelerated bone maturation was defined as the presence of the distal femoral ossification center before 31 WG, and delayed bone maturation was defined as the absence of the center after 33 WG (23).

Fetal tachycardia was defined as an FHR continuously greater than 160 beats/min (24).

TSH, free T3 (FT3), and FT4 were measured using a chemiluminescence immunoassay with the ACS-180SE system (Bayer Diagnostics, Westwood, CA). The values were interpreted according to gestational or postnatal age (25). TRAK were measured by RIA with second-generation antibodies (RIA-2 Dynotest TRAK human, BRAHMS Diagnostica GmbH, Berlin, Germany) (19). A positive result (TRAK*) was defined as an antibody titer greater than 2 U/liter. The results are reported as multiples of the upper limit of normal. TRAK assay results and ATD treatment were used to divide the mothers into two groups. The high-risk group, in which the fetuses were considered at risk for thyroid dysfunction, comprised mothers with at least one TRAK* (20) and/or ATD treatment in the third trimester of pregnancy. The low-risk group was composed of mothers with consistently negative TRAK assays (TRAK-) and no ATD treatment, indicating a low risk for fetal thyroid dysfunction (26). The thyroid function did not differ between the two groups of mothers, indicating adequate ATD and/or T4 treatment in those treated (see below).

In neonates with an age at birth of 36 WG or more, normal ranges for cord blood values were defined as follows: FT4, 10.4–16.4 pmol/liter, and TSH, 2.6–11.8 mU/liter (25). Hypothyroidism was defined as an FT4 level below the 2.5th percentile and a TSH value greater than the 97.5th percentile. Hyperthyroidism was defined as an FT4 value greater than the 97.5th percentile and a TSH value below the 2.5th percentile. FT3 levels in cord blood are normally very low and consequently unhelpful for diagnosing hypothyroidism, although they can assist in the diagnosis of hyperthyroidism. We therefore used only cord blood FT4 and TSH for diagnosing fetal thyroid dysfunction.

Decisions to perform fetal blood sampling (FBS) were based on the treatment of the mother, Doppler signal pattern, and TRAK positivity; maternal consent was required before the procedure. FBS was not performed when the results of other tests discriminated between hypothyroidism and hyperthyroidism or when the pregnancy was sufficiently advanced to allow induction of labor within a reasonable period (27).

Once the type of fetal thyroid dysfunction was determined, the treatment consisted of an ATD dosage increase when the diagnosis was fetal hyperthyroidism [with maternal levothyroxine (l-T4) therapy if needed] and of ATD withdrawal or dosage reduction when the diagnosis was fetal hypothyroidism, as permitted by the maternal endocrine status. When early-onset fetal hypothyroidism was diagnosed, intratracheal l-T4 was administered by amniocentesis, as previously described (17). Of note, this approach has been used sparingly and should only be discussed by teams used to this type of prenatal care.

Fig. 1. Fetal thyroid and ultrasonography. A, Fetal thyroid gland with normal circumference and increased diameter. Note the central echo-free disc corresponding to the trachea and carotid arteries visualized by color Doppler. B, Fetal goiter and hypothyroidism; peripheral flash on the color Doppler. C, Fetal goiter and hyperthyroidism; central flash on the color Doppler. The measurement of fetal thyroid size has been validated and can be used in routine prenatal care. The color Doppler is still an experimental method that should be validated in more cases.
At delivery, cord blood was retrieved for thyroid function tests, the newborn was examined by a pediatric endocrinologist, and ultrasonography of the thyroid gland was performed (28).

This study was approved by the Paris-Saint Louis ethics committee for biomedical studies in humans and was conducted in compliance with French law. Written informed consent was obtained from the pregnant women.

Results

Description of the cohort of women with Graves’ disease (n = 72)

The mean age in the overall population was 33 yr (range, 26–43 yr). The gestational age at inclusion was 17 WG (range, 10–28 WG). Of the 72 women, 41 were in the high-risk group, and 31 were in the low-risk group. Details are given in Table 1. Overall, 45 women received no treatment or only L-T4 and 27 received ATD therapy (with or without L-T4). Of the 41 high-risk women, 33 had at least one TRAK+ result during pregnancy, and eight received ATD therapy during the last trimester, but had consistently TRAK− tests; these eight patients were treated because of recent onset or exacerbation of Graves’ disease, in keeping with the widely recognized need for giving ATD therapy for 18 months and not more (20). Of the 26 women with a history of total (n = 3) or subtotal (n = 23) thyroidectomy, 9 (34%) had at least one TRAK+ test during pregnancy.

Fetal thyroid function and its relation to maternal thyroid function

In the low-risk group (n = 31), fetal sonograms showed no evidence of goiter. Cord blood thyroid function tests were normal at delivery in all 31 neonates. Thyroid dysfunction did not develop in any of these neonates during the first postnatal month (Fig. 2). In the high-risk group (n = 41), 11 of the 41 fetuses had ultrasonogram evidence of goiter at 32 WG and, consequently, were evaluated for thyroid dysfunction (Fig. 2). The results showed hypothyroidism in seven fetuses and hyperthyroidism in four fetuses at 32 WG (Table 2).

Fetal hyperthyroidism was usually associated with low maternal TRAK titers, high ATD dosages (=150 mg/d propylthiouracil, =15 mg/d methimazole, or =100 mg/d benzylthiouracil), and/or delayed fetal bone maturation. The Doppler signal was positive throughout the fetal thyroid gland in two cases and at the periphery of the gland in four cases; it was not detected in one case and was not documented in one case. Delayed bone maturation was noted in two of these seven fetuses. Fetal blood was obtained in four cases for thyroid function tests, which confirmed the diagnosis of fetal hypothyroidism. The ATD dosage was reduced in all seven cases, and L-T4 was injected into the amniotic fluid in three fetuses after confirmation of hypothyroidism by fetal blood sampling. In one case (no. 4, Table 2), FBS was not performed because the low TRAK level led us to postulate that the fetus had hypothyroidism. This prompted us to decrease the ATD dose. This appeared subsequently to be correct, because a decrease in thyroid size occurred 2 wk later. Indeed, after treatment, sonograms performed at 2-wk intervals consistently showed a decrease in the size of the thyroid gland; in two cases, the gland returned to normal size.

Thyroid function did not differ between the low-risk and high-risk groups of mothers. In the low-risk group, mean ± SD FT4 and FT3 levels during the third trimester were, respectively, 12.4 ± 1.3 and 4.1 ± 0.3 pmol/liter, not different from those in the high-risk group (13.9 ± 2.5 and 4.8 ± 1.1 pmol/liter). Even the maternal FT4 and FT3 levels in the three women with fetuses with the more severe hypothyroidism (Table 2) did not differ from those in the other women of the high-risk group (mean of the third trimester: case 1: FT4, 12.2 pmol/liter; FT3, 4.1 pmol/liter; case 5: FT4, 11.2 pmol/liter; FT3, 3.9 pmol/liter; case 7: FT4, 15.3 pmol/liter; FT3, 5.2 pmol/liter; Table 2). However, at the time of FBS and ultrasonography, FT4 and FT3 in those three women were: case 1: FT4, 8.6 pmol/liter; FT3, 3.2 pmol/liter; case 5: FT4, 9.7 pmol/liter; FT3, 3.9 pmol/liter; and case 7: FT4, 16.1 pmol/liter; FT3, 5 pmol/liter (Table 2).

At birth, cord blood tests showed normal FT4 levels in six babies and a borderline low FT4 in one (Table 2). Within the first postnatal week, hyperthyroidism requiring ATD therapy developed in three of these seven neonates (Table 2, upper part). The mothers of these seven fetuses had normal thyroid function.

Fetal hyperthyroidism (n = 4) was usually associated with high maternal TRAK titers and/or accelerated fetal bone

### TABLE 1. Treatment during pregnancy and TRAK status in the women

<table>
<thead>
<tr>
<th>TRAK+</th>
<th>TRAK−</th>
</tr>
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<tbody>
<tr>
<td>ATD</td>
<td>16</td>
</tr>
<tr>
<td>ATD + L-T4</td>
<td>3</td>
</tr>
<tr>
<td>l-T4</td>
<td>6</td>
</tr>
<tr>
<td>None</td>
<td>8</td>
</tr>
<tr>
<td>Total</td>
<td>33</td>
</tr>
</tbody>
</table>

Data are shown for 41 high risk women, 33 from the TRAK+ column and eight from the TRAK− column treated with ATDs (in bold). All but three were TRAK+ after 20 WG. The three remaining patients were TRAK+ in the first trimester and TRAK− subsequently.

FIG. 2. Ultrasonography for fetal thyroid monitoring according to maternal status and cord blood thyroid function test results. Hypot, Hypothyroidism; Eut, euthyroidism; Hypert, hyperthyroidism. 1) In one mother, FT4 (20.2 pmol/liter) was at the upper limit of normal with suppressed TSH (<0.05 mU/liter) despite the high maternal dose of propylthiouracil (PTU). 2) FT4, Mean of FT4 in all of these newborns with 1 SD; TSH, mean TSH in all of these newborns with 1 SD.
maturation. The Doppler signal was positive throughout the fetal thyroid gland in all four cases. Advanced bone maturation was noted in two fetuses. In two cases, after a multidisciplinary evaluation, fetal blood tests were performed and confirmed the diagnosis, showing a very high T4 value (51.7 pmol/liter) and a minimally elevated value (14.8 pmol/liter), but a high T3 level (7.5 pmol/liter) in this case and both cases with low TSH levels (<0.05 mU/liter; Table 2). The ATD dosage was increased in all four mothers. The size of the fetal thyroid gland decreased within the next 2 wk; it returned to normal in one fetus. However, one fetus died in utero at 35.5 WG from congestive heart failure with cardiomyopathy. This was a late-referred case with very high TRAK values and a preexisting fetal goiter, as shown by a peripheral color Doppler signal at 22 WG. ATD therapy was increased up to 60 mg methimazole/d to no effect. FBS confirmed the diagnosis of severe hyperthyroidism, and the fetal blood TRAK titer was massively elevated (325 ULN). Of the four mothers, two had hyperthyroidism and two had normal thyroid function.

The remaining 30 fetuses in the high-risk group had no goiter and consequently were not considered at risk for thyroid dysfunction (Fig. 2). All had normal thyroid values in cord blood at delivery, except one neonate who had moderate hypothyroidism (FT4, 9.6 pmol/liter; TSH, 20.6 mU/liter) that resolved spontaneously without treatment. Retrospectively, this case could have been detected, because a peripheral color Doppler signal was noted at 38 WG, when thyroid gland size parameters were at the upper limit of normal.

Based on the results of thyroid cord blood testing at delivery and the therapeutic interventions in the treated fetuses, the sensitivity and specificity of fetal thyroid ultrasound at 32 WG in screening for clinically relevant (leading to therapeutic action) fetal thyroid dysfunction were 92% and 100%, respectively (Table 3). This clearly indicates a major role for fetal thyroid ultrasonography in diagnosing fetal thyroid dysfunction.

<table>
<thead>
<tr>
<th>TABLE 2. Fetuses at 32 WG: seven with hypothyroidism (cases 1–7) and four with hyperthyroidism (cases 8–11)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Case no.</strong></td>
</tr>
<tr>
<td>---</td>
</tr>
<tr>
<td>1</td>
</tr>
<tr>
<td>2</td>
</tr>
<tr>
<td>3</td>
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<tr>
<td>4</td>
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<tr>
<td>5</td>
</tr>
<tr>
<td>6</td>
</tr>
<tr>
<td>7</td>
</tr>
</tbody>
</table>

* Number of times the upper limit of normal.

<table>
<thead>
<tr>
<th><strong>Bone</strong></th>
<th><strong>Cord blood</strong></th>
<th><strong>Neonatal hyperthyroidism</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Bone</td>
<td>Cord blood</td>
<td>Neonatal hyperthyroidism</td>
</tr>
<tr>
<td>3</td>
<td>10.8</td>
<td>No</td>
</tr>
<tr>
<td>4</td>
<td>13.5</td>
<td>No</td>
</tr>
<tr>
<td>5</td>
<td>12</td>
<td>No</td>
</tr>
<tr>
<td>6</td>
<td>2.5</td>
<td>No</td>
</tr>
<tr>
<td>7</td>
<td>6</td>
<td>Yes</td>
</tr>
<tr>
<td>8</td>
<td>0.05</td>
<td>Death</td>
</tr>
<tr>
<td>9</td>
<td>160</td>
<td>Normal</td>
</tr>
<tr>
<td>10</td>
<td>51.7</td>
<td>Yes</td>
</tr>
<tr>
<td>11</td>
<td>10</td>
<td>Yes</td>
</tr>
</tbody>
</table>

* Normal ranges for cord blood values were: FT4, 10.4–16.4 pmol/liter; and TSH, 2.6–11.8 mU/liter (25).

**TABLE 3. Fetal thyroid ultrasonogram in screening for fetal thyroid dysfunction**

<table>
<thead>
<tr>
<th>Goiter</th>
<th>TP = 11</th>
<th>FP = 0</th>
</tr>
</thead>
<tbody>
<tr>
<td>No goiter</td>
<td>FN = 1</td>
<td>TN = 60</td>
</tr>
</tbody>
</table>

Sensitivity, 92%; specificity, 100%; positive predictive value, 100%; negative predictive value, 98%. These values were calculated with our criteria at 32 WG when screening for clinically relevant (leading to therapeutic action) fetal thyroid dysfunction and are based on the values of the thyroid cord blood testing at delivery and the therapeutic interventions in the treated fetuses. FP, False positive; FN, false negative; TN, true negative; TP, true positive.

* Measured in cord blood at delivery.
thyroidectomy were TRAK−, indicating that TRAK assays should be performed in this patient subgroup, at least at the beginning of pregnancy. This later point is in line with European Thyroid Association recommendations (18).

Propylthiouracil was the most widely used ATD among the women who took ATD therapy during pregnancy. This drug is currently preferred over methimazole based on its lower teratogenic potential, even if its limited transplacental passage has been shown to be similar to that of methimazole in an in vitro assay using the isolated perfused placenta lobule (29). Indeed, methimazole given during organogenesis has been reported to cause aplasia cutis and tracheoesophageal fistula as well as a characteristic embryopathy (30, 31).

The second important finding of our study is that 11 (27%) of the 41 fetuses from high-risk mothers had goiter by fetal ultrasonography. Normograms for fetal thyroid size were published by Ranzini et al. in 2001 (21) and were validated in our setting in 250 euthyroid pregnant women (our own unpublished observations), thus allowing a precise diagnosis of fetal goiter. Monthly fetal thyroid ultrasonograms ensured the detection of all cases of fetal goiter and therefore of all cases of clinically significant (requiring therapeutic modifications) thyroid dysfunction with 100% specificity. Indeed, it has been reported that at birth, several offspring of mothers with Graves’ disease treated with ATD had transient elevated TSH levels and no goiter (32, 33). However, no neonatal thyroid ultrasound was performed; therefore, the presence of subtle thyroid enlargement cannot be ruled out (32, 33). Moreover, this technique can be acquired by ultrasonographists, thus allowing its widespread use. Nachum et al. (34) recently reported data from 18 women managed over a period of 10 yr. In these patients, fetal blood sampling was extensively practiced even in the absence of fetal goiter and occasionally revealed fetal thyroid dysfunction. This discrepancy with our results can be explained by the fact that fetal sonograms were less standardized in the study by Nachum et al. (34). Our data establish that fetal blood sampling is unnecessary in the absence of fetal goiter. Similarly, Kilpatrick (35) stated that isolated high TRAK titers are not sufficient to recommend routine FBS. In a series of 20 women, including five with goiter in the fetus, Cohen et al. (36) also obtained convincing evidence that ultrasonography is an effective noninvasive screening tool for detecting fetal goiter. However, they did not assess maternal TRAK titers, so the interpretation of fetal goiter in their study is difficult.

In fetuses with goiter, the main clinical problem was to determine whether the cause was maternal treatment that was adequate to achieve normal maternal thyroid function but inadequate and excessive for the fetus and therefore responsible for fetal hypothyroidism or fetal thyroid stimulation by maternal Graves’ disease IgGs responsible for fetal hyperthyroidism. It is generally agreed that fetal FT4 levels correlate with maternal FT4 levels and that fetal euthyroidism can be achieved by maintaining maternal FT4 in the upper normal to mildly thyrotoxic range during antithyroid drug treatment. This fact is underlined by the present results, because maternal FT4 levels in the three cases with extremely severe hypothyroidism were in the normal range in case 7 and in the lower normal range in the two others (cases 1 and 5). Therefore, in these two last cases, the maternal FT4 level may have also helped us in the diagnosis of fetal hypothyroidism.

Nevertheless, we mainly used a combination of maternal criteria (TRAK titer and ATD use and dosage) and fetal criteria (thyroid Doppler signal, FHR, and bone maturation) to distinguish between fetal hypothyroidism and hyperthyroidism. FBS was only discussed after the patient’s agreement and if there was a fetal goiter with no possibility of distinguishing hypothyroidism and hyperthyroidism using maternal and fetal criteria. Fetal hyperthyroidism was diagnosed based on high maternal TRAK titers, accelerated fetal bone maturation, and an FHR greater than 160 beats/min; this last sign was uncommon, because it occurs late in the natural history of fetal hyperthyroidism. The limited usefulness of FHR for diagnosing fetal hyperthyroidism has also been underlined by Nachum et al. (34). Doppler examination of the fetal thyroid gland proved useful only when the flash was confined to the periphery of the gland, a pattern suggestive of fetal hypothyroidism. The prenatal response to treatment and the results of thyroid function tests of fetal blood when obtained and of cord blood at birth indicated that our criteria were effective in differentiating hypothyroidism and hyperthyroidism in all 11 fetuses with goiter.

Fetal blood was sampled from six of the 11 fetuses with goiter, and the results consistently confirmed the suspected diagnosis. Abortion, fetal bradycardia, and infection have been reported after FBS in 1% of cases (27). Consequently, FBS should be reserved for those cases in which intraamniotic T4 injection is considered or thyroid status is in doubt in a fetus whose mother has TRAK+ assays and receives ATD therapy.

Routine cord blood assays of FT4 and TSH at delivery allowed us to validate our prenatal strategy, but did not help us to predict subsequent neonatal thyroid dysfunction, as reported previously (37). Indeed, the remaining 30 fetuses in the high-risk group had no fetal goiter, and it should be noticed that the major evidence of the absence of fetal thyroid dysfunction is based on the results from cord blood at birth. Normal thyroid function in utero was not definitively demonstrated, because FBS appeared unnecessary and, therefore, unethical in this group of patients.

From our data we suggest the following strategy for the care of these pregnancies, which differs from that of the European Thyroid Association. It should now be validated in other data sets, taking into account our sample size. TRAK should be assayed routinely when the fetal thyroid is functioning. For practical reasons, we believe that it can be performed at the beginning of pregnancy in women with current or past history of Graves’ disease; indeed, it can be performed with other screening procedures offered to pregnant women at 12–13 WG, and a negative result will avoid unnecessary additional work-up. During pregnancy in women who are receiving ATD therapy and/or who have positive tests for TRAK, a fetal ultrasonogram should be performed monthly after 20 WG to screen for goiter and/or other evidence of fetal thyroid dysfunction. During pregnancy in a woman with negative tests for TRAK and not receiving ATD treatment, the usual ultrasonograms during pregnancy (recommended in France at 22 and 32 WG) should be performed. Of note, most fetal thyroid dysfunction (10 of 11 cases) arose in women with no history of thyroidectomy in our high-risk group, whereas only one case was observed in the offspring of the 26 women who underwent thyroidectomy. Therefore, the former group should be the target of careful prenatal follow-up.
In conclusion, our study shows that prenatal screening ultrasonography is a powerful tool for detecting fetal thyroid dysfunction in maternal Graves’ disease. Ultrasound monitoring of fetuses from mothers with Graves’ disease was extremely sensitive and specific for detecting intrauterine thyroid dysfunction and therefore allowed appropriate fetal management. In that it was a prospective and systematic study of fetal thyroid size, it is a new approach, which is not comparable to previous studies. This tool in conjunction with close teamwork among internists, endocrinologists, obstetricians, echographists, and pediatricians can ensure normal fetal thyroid function. Whether this systematic monitoring makes a difference in terms of ultimate psychomotor development of the children compared with more targeted investigations remains to be demonstrated.

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