What is the benefit of screening for thyroid function in pregnant women in the detection of newly diagnosed thyropathies?

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**Background.** Thyroid gland disturbances are the most common endocrine disorders in pregnancy. There are some particular recommendations for the investigation of women in risk groups, but no consensus guidelines for general screening exists at present in the Czech Republic.

**Aim.** The aim of our study was to determine whether universally conducted screening of pregnant women would reveal a significant number thyropathies.

**Material and Methods.** We examined 592 pregnant women for thyroid-stimulating hormone (TSH) and free thyroxine (fT4) levels and for autoantibodies against thyroperoxidase (antiTPO) in the 6th - 10th week of their pregnancy.

**Results.** Levels of TSH, fT4 or antiTPO beyond laboratory reference limits were found by gynaecologists in 214 women (36.1%) and 141 of whom (23.8%) underwent endocrinological examination. In the women without known risk factors (n=91) we found undiagnosed autoimmune thyroiditis in 20 cases (22 %) and in 7 cases (7.7%) some degree of subclinical hypothyroidism was confirmed. Finally, 18 (19.8%) women had hypothyroxinemia in the 1st trimester (fT4 average 8.76 pmol/L) with normal TSH levels. Altogether, a total of 45 women were successfully identified (49.5% of the endocrinologically examined group without risk factors, i.e. 7.6% of the whole screened group) who warranted monitoring.

**Conclusions.** Of 592 women in the 6th - 10th week of pregnancy who underwent thyropathy screening, we newly diagnosed 3.4% of women with autoimmune thyroiditis, 1.2% with subclinical hypothyroidism and 3% with hypothyroxinemia, for whom no thyropathy risk factor had been evident. Thyropathies were identified in 7.6% of probands. We believe that our results support the importance of universal screening in pregnancy.

**Key words:** thyropathy, pregnancy, screening

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**INTRODUCTION**

The relationship between thyroid gland function, conception, course of pregnancy, and foetal development has been known for a long time. The most common thyroid gland dysfunction in pregnancy is hypothyroidism, the prevalence of which is estimated at 1.5 - 4.4% of pregnant women in the population. As for the etiology an autoimmune inflammation of the thyroid gland, chronic thyroiditis is the most common cause. The other reason might be the inadequate supply of iodine and conditions after surgery on the thyroid gland or radioactive iodine treatment. Hypothyroidism is linked to reduced fertility in women. If a woman with hypothyroidism becomes pregnant then the lack of thyroid gland hormones leads during intrauterine development to irreversible damage to the foetus (failure of differentiation of nerve cells, inadequate central nervous system development and psychomotor function, increased risk of perinatal defects). The other threats are related to risk of premature birth, spontaneous abortion, placental abruption and post-natal bleeding, pre-eclampsia. A recent study confirmed no improvement in the cognitive function of children born to women who were given thyroxine treatment on the basis of hypothyroidism detected through screening in the first trimester.

Serious complication may also be a danger for euthyroid women with positive autoantibodies especially against thyroperoxidase. A vast number of these women suffer from autoimmune thyroiditis (AIT). The thyroid gland is then unable to react to the increased demands for the production of thyroxine in pregnancy. Patients with autoimmune thyroiditis have twice or three times the risk of miscarriage in the 1st trimester, which can be positively influenced by timely prescription of thyroxine.

A rather rare condition in pregnant women is hyperthyroidism: Both Graves - Basedow disease and autoimmune thyroiditis in its hyperfunctional form represent a complication in the course of pregnancy. The negative impact on the development of the foetus (foetal growth retardation, pre-eclampsia, miscarriage, premature birth) is evident. It is important to differentiate between this illness and simple TSH suppression or temporary thyrotoxicosis...
in pregnancy, which recedes spontaneously without treatment after the 14th week of pregnancy\textsuperscript{2,9}.

Despite the known complications of undiagnosed and untreated disorders of thyroid gland function for mothers and babies, there is still ongoing discussion about the extent of thyropathy screening. Universal examination of all women is not recommended\textsuperscript{9-11}, the position supported by the American Thyroid Association (ATA) (Ref.\textsuperscript{12}).

On the other hand, there are studies which recommend universal screening\textsuperscript{13}. One study conducted in 2010 confirmed twice the incidence of thyropathies in universal screening of pregnant women than in the case of examining women at risk\textsuperscript{2}.

In our study we focused on the benefit of universal examination of pregnant women in the Olomouc region (Czech Republic) and women’s interest in the option of examination.

MATERIALS AND METHODS

In cooperation with gynaecologists, blood samples were taken over a period of 2 years (2010, 2011) from pregnant women within prenatal screening in the 6th - 10th week of pregnancy to determine the levels of TSH, fT4 and antiTPO. This was a total of 592 pregnant women who signed the written form to undergo examination and the processing of results.

All the pregnant women were advised to increase their dietary intake of iodine at the time of their first visit to the gynaecologist and they were offered supplements in the form of 100 µg iodine per day or in the form of multivitamin with iodine for pregnant women. Commencement of iodine supplements varied, mostly after blood sampling.

Pregnant women, whose tests determined levels of TSH, fT4 or antiTPO outside the laboratory reference limits in any of the categories, were advised to undergo endocrinological examination. This applied to 214 (36.1%) women, of which 141 (23.8%) underwent examination.

Sonographic examination of the thyroid gland was carried out, personal and family history of thyropathy was recorded along with personal history of other autoimmune illnesses. We retrospectively evaluated the available results of laboratory tests for women who did not undergo examination (73 women, 12.3% of the screened group).

In determining thyroid-stimulating hormone (TSH), free thyroxine (fT4) and autoantibody against thyroperoxidase (antiTPO) the Architect analyser was used (Abbott reagent) in the CMIA method (Chemiluminiscent Microparticle Immunoassay).

Laboratory reference limits for levels of TSH were 0.3 - 4.0 mU/L, laboratory reference limits for fT4 were 9.55 - 23 pmol/L, the laboratory normal limit for antiTPO was up to 5.61 kU/L.

With regard to TSH and HCG interference in the first trimester we reassessed TSH results obtained by gynaecologists according to recommendations for pregnant women (TSH 0.06 or 0.03 - 3.67 mU/L) (Ref.\textsuperscript{9,14}).

RESULTS

Of 592 women screened a total of 141 (23.8%) pregnant women underwent endocrinological examination on the recommendation of a gynaecologist (Fig. 1). Of these, 8 (5.8%) women had a personal history of thyropathy and a further 42 (29.8%) had positive family history of thyroid gland illness. None of them had Type 1 diabetes mellitus or any other known autoimmune illness and 91 (64.5%) of the pregnant women examined by an endocrinologist had no known thyropathy risk factors.

In the women with a positive family history (42 women) we found autoimmune thyroiditis with positive antiTPO and corresponding to sonographic findings in 13 (31%) women. A further 5 (11.9%) women had positive autoantibodies with normal sonographic findings. 14 (33.3%) women were found to have pathological ultrasonography results, but we found no higher antiTPO levels. Of the women with a positive family history as a risk factor there were 32 (76.2%) women requiring endocrinological observation, while 18 women (42.9%) had positive antiTPO (Fig. 2).

In women with no known thyropathy risk factor (91 women), there were 20 (22%) patients with newly diagnosed autoimmune thyroiditis - 14 euthyroid cases and another 6 subclinical hypothyroidism cases with path-
Fig. 2. Women with family history risk (n=42).

Fig. 3. Newly identified thyropathy in women with negative family history and with no known risk factors (n=91). Reference range of TSH 0.06 - 3.67 mU/L.

Fig. 4. Levels of TSH beyond recommended reference limits in women with positive (FH+) and negative (FH-) family history.


**Table 1. Risk groups of women advised to undergo examination of thyroid function in pregnancy.**

1. Women with a personal history of thyropathy or with previous irradiation of head and neck
2. Women with a family history of thyroid gland illness
3. Women with clinically known illness, goitre or symptoms
4. Women with positive antiTPO or antiTBG autoantibodies
5. Women with Type 1 DM or other autoimmune illness
6. Women examined for sterility within examination protocol

Adapted according to Abalovich et al. 2007 (ref.9).

**DISCUSSION**

In 2004 examination was recommended for pregnant women and women planning pregnancy with the aim of preventing the unfavourable result of unrecognised subclinical hypothyroidism17. In 2007 a recommendation was drawn up on the basis of consensus of thyroid associations8. The recommendation deals with illness of the thyroid gland in pregnancy, its detection and treatment, ideally also setting preconception diagnosis. Examination of TSH is recommended, possibly fT4 and antiTPO for patients in risk groups (Table 1). Consensus guidelines for thyroid disorder screening have been published in the Czech Republic16.

In our opinion, of the stated risk factors it is easy to identify women with a personal history of thyropathy and with irradiation of head and neck. Information on family history is more problematic - we repeatedly came across data on negative family history only for the patient to change the original information during a check-up after the examination. Clinical symptoms of thyroid gland illness - tiredness, inefficiency, sleepiness etc. are not typical, and coincide with general symptoms of pregnancy. This is also given by the fact that, in the Czech Republic where iodine intake is sufficient, we are dealing with subclinical forms of hypothyroidism17. In our experience, palpable goitres are uncommon in young women, evaluation of the size of the thyroid gland without sonographic examination is often problematic. Autoantibodies (antiTPO and antiTBG) are specifically determined in women with suspected thyropathy.

In our study group we found 50 women (8.4%) in risk groups - 8 women with personal history of thyropathy, a further 42 women had positive family history of thyropathy. The high percentage of women with diagnosed thyropathy with positive family history indicates the importance of monitoring this case-history information, even if it is not always completely valid information.

Through universally conducted examination in the group of women with no known risk of thyropathy we found 45 women (7.6%) who require endocrinological treatment.

Even in this risky period, where we would expect greater interest in good health and a successful course of pregnancy, there were a total of 55 women (9.3% of the whole screened group) whose screening tests indicated pathologies, but who did not come for endocrinological examination despite this being recommended.

It is impossible to speak of the risk of these women who should be monitored and treated. They run the risk of complications due to increased antiTPO level, practically half of them have subclinical hypothyroidism and run the risk of postnatal thyroiditis and hypothyroidism18.

The incidence of positive autoantibodies against thyroperoxidase was high among women with positive family history, overall we detected them in 10.1% of the women, which corresponds with published data8,14.

Our observation found 7.6% newly diagnosed thyropathy in women with no risk factor, which corresponds with a pilot study in CZ (ref.19). Another study showed that if only patients in risk groups are examined about a third.
of women with hypothyroidism will not be detected\textsuperscript{20}. A similar conclusion was reached by a further study - here as much as 55% of women with thyropathy, who had no risk factors, would not be examined\textsuperscript{2,13}. Universally conducted screening revealed 132 women, i.e. 22.3% of all women examined, who should be monitored and treated for thyropathy. It can be stated that, from every 100 women without risk factors who are screened, we would find almost 17 women with newly detected thyropathy. In terms of the effectiveness of the expenses incurred we consider this approach to be appropriate\textsuperscript{6,21}.

CONCLUSIONS

On the basis of the results of our study we believe that 7.6% newly diagnosed thyropathy in women with no known risk factor is a sound argument for introduction of simple universal screening of pregnant women. We believe that evaluation of TSH and FT\textsubscript{4} levels in the 5th-10th week of pregnancy is necessary to determine thyroid function. Without evaluation of antiTPO levels 1.9% of autoimmune thyroiditis with pathological sonography could not be diagnosed in our study. Attention should be given to another 12.3% of women in our study who did not undergo endocrinological examination despite recommendation and 9.3% who should be observed and treated further, the risk factors for this group are not known.

It is essential to improve cooperation between gynaecologists, endocrinologists and GPs. It is also essential that the public is better informed on the importance of examination.

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CONFLICT OF INTEREST STATEMENT

Author’s conflict of interest disclosure: None declared.

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