Clinical Study

Newborn TSH concentration and its association with cognitive development in healthy boys

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Abstract

Objective: An association between thyroid function during pregnancy or infancy and neurodevelopment in children has been demonstrated. We aimed to investigate whether newborn TSH concentrations are related to subsequent neurocognitive development.

Design: We conducted a longitudinal study on 178 children from a general population birth cohort in Granada (Spain) born in 2000–2002.

Methods: TSH concentrations were measured in umbilical cord blood, and cognitive functions were assessed at 4 years of age using the McCarthy’s scales of children’s abilities (MSCA). Organochlorine (OC) compound concentrations and the combined oestrogenicity (total effective xeno-oestrogenic burden (TEXB)) were also determined in the placentae.

Results: Mean newborn TSH was 3.55 mU/l (range 0.24–17 mU/l). In multivariate regression analyses, adjusting for maternal and child characteristics, higher newborn TSH concentrations showed a decrease of 3.51 and 3.15 points on the MSCA general cognitive and executive function scores respectively and were associated with a higher risk of scoring below the 20th percentile (P20) (odds ratio (OR) = 2.64). Children with TSH in the upper quartile (4.19–17.0 mU/l) were at higher risk of scoring > P20 on span memory (OR = 5.73), whereas children with TSH in the second quartile (2.05–2.95 mU/l) were at lower risk of scoring < P20 on the verbal scale (OR = 0.24). Neonatal TSH status was also associated with general cognitive and executive function outcomes when controlling for prenatal exposure to OCs or placental TEXB.

Conclusions: Newborn thyroid hormone status expressed by TSH in cord blood may adversely affect later cognitive function. A more thorough screening for neonatal thyroid deficiency is warranted.

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Introduction

Thyroid hormones (THs) are essential for the fetal and postnatal human development and for the regulation of neuropsychological function in children and adults (1). THs regulate the processes of neurogenesis, myelination, dendrite proliferation and synapse formation (2, 3). Although THs are required throughout gestation, the fetal thyroid gland does not produce its own TH in appreciable amounts until the third trimester (4, 5). Accordingly, an increasing number of epidemiological studies and case reports have strongly supported the notion that impaired maternal thyroid function during early gestation may result in poor fetal neurodevelopment (6–10).

Iodine deficiency (ID), which compromises adequate production and secretion of thyroxine (T4) by the thyroid, remains the most frequent cause of maternal and fetal TH deficit and therefore of preventable mental retardation (5). Over the past two decades, attention has been drawn to the sub-optimal cognitive or behavioural functioning (which may be sub-clinical) observed in children born to mothers with even mild or moderate ID (10–16) or the developmental implications for children with slight neonatal elevations of TSH (17, 18). Thyroid deficiency during the last two trimesters of pregnancy and the first few months post delivery can also result in mental and physical retardation and sometimes neurological deficits, a condition known as cretinism (19). Neurological features are less severe in neonatal hypothyroidism than in prenatal hypothyroidism, although deficits in memory and intelligence quotient may persist (2). Overall, few studies have addressed whether subsequent development can be influenced by moderate thyroid dysfunction in neonates or even by variations within the normal range of TH levels.
Recent studies indicate that exposure to certain environmental contaminants may also interfere with maternal thyroid status during pregnancy and with thyroid function in newborns (20–26). Hence, it has been speculated that some of the neurotoxic effects of early exposure to the environmental chemicals may result from thyroid disruption (27, 28). Nonetheless, the influence of early exposure to endocrine disruptors (ED) on thyroid function and therefore children’s neurodevelopment remains to be elucidated.

In a previous study of a mother–child cohort, we analysed the influence on neonatal TH status of placental exposure to certain organochlorine (OC) pesticides with known ED activity (C Freire, M J Lopez-Espinosa, M F Fernández, J M Molina-Molina, R Prada and N Olea, unpublished observations). In this study, we examined the association of newborn TSH concentrations with cognition at 4 years of age in the same cohort (in Granada, Southern Spain). We hypothesised that infants with poorer thyroid status, manifested by higher TSH levels, would have lower scores in subsequent cognitive testing. This investigation is part of the ‘Infancia y Medio Ambiente (Environment and Childhood) (INMA) Project’, a prospective multicentre study in Spain (29).

Materials and methods

Subject recruitment

From 2000 to 2002, 700 eligible mother–son pairs registered at the San Cecilio University Hospital were enrolled at delivery, establishing the INMA-Granada cohort, with the initial aim of investigating chronic exposure to ED and urogenital malformations in newborn boys. Exclusion criteria were the maternal presence of serious chronic disease, such as diabetes, hypertension or thyroid disease; a pregnancy complication that could affect growth or development and a non-residence or thyroid disease; a pregnancy complication that could affect growth or development and a non-residence in the hospital referral area (30). In 2005–2006, 220 of the 700 boys aged 4 years and their mothers were randomly invited to participate in the physical examination and cognitive testing (31, 32).

Approval by committee for human subjects

A written informed consent was obtained from parents before the study, which was approved by the Ethics Committee of the San Cecilio University Hospital, in accordance with the Code of Ethics of the World Medical Association (Declaration of Helsinki) for experiments involving humans.

The study was approved by the Institutional Ethical Committee of the Hospital, and signed informed consent was also obtained from the women who agreed to participate.

TSH determination

TSH was measured in a cord blood sample spotted on a filter paper (Schleicher & Schuell no. 2992), which is routinely obtained at delivery for the screening programme of neonatal congenital hypothyroidism (33). TSH concentrations were determined by using time-resolved sandwich fluoroimmunoassay (Auto-DELFIA, Perkin Elmer/Wallac, Turku, Finland) at the Centre for the Early Detection of Metabolopathies in Neonates in San Juan de Dios Hospital (Granada, Spain). The limit of detection (LD) was 0.01 mU/l. A cord blood level ≥ 14 mU/l is established in the Centre’s laboratory to trigger the protocol for the study and for the confirmation of neonatal hypothyroidism.

Quantification of OC pesticides and oestrogenicity in the placenta

Placentas were collected at delivery from the cohort, and 17 OC pesticides (dichlorodiphenyltrichloroethane (DDT) isomers and metabolites, endosulphan isomers and metabolites, aldrin, endrin, dieldrin, lindane, hexachlorobenzene, methoxychlor and mirex) were extracted from tissue samples by a previously described method, which was developed to separate natural oestrogens (α fraction) from more lipophilic xeno-oestrogens (β fraction) without destroying either (34, 35).

OC pesticide concentration was determined in 308 randomly selected placenta samples by gas chromatography (GC) with electron-capture detection. The compounds were confirmed by GC and mass spectrometry (30). The LD for the studied chemicals ranged from 0.1 to 3.0 ng/ml. For levels below the LD, we considered a value of half the LD.

The total effective xeno-oestrogenic burden (TEXB) of the α fraction (α-TEXB) and the β fraction (β-TEXB) was estimated in the placenta samples by using the E-Screen bioassay (34, 35). The α-TEXB can be considered a marker of the TEXB of environmental organohalogenated oestrogens (35). The LD of TEXB was defined as the concentration needed to produce a significantly different proliferative effect from that observed in the control cells.

Cognitive testing

The neurocognitive evaluation of the children was performed by two specifically trained psychologists in the Paediatrics Department of our Hospital. Cognitive and motor abilities were assessed using a Spanish adaptation of the McCarthy scales of children’s abilities (MSCA) (36), which gives standardised test scores for five domains (quantitative, verbal, memory, perceptual performance and motor). A general cognitive score, which estimates global intellectual function, was calculated by combining the verbal, perceptual...
performance and quantitative scores. A strict protocol was applied to avoid inter-observer variability (37), which was < 5%. Psychologists involved in the cognitive testing of the children were unaware of the design sequence of the study.

At the same time as the children were evaluated, the parents completed a self-reported questionnaire on parent-to-infant attachment and another questionnaire on mental health, considered as effect modifiers on infant mental development (38). The parent-to-infant attachment questionnaire consisted of 19 items that assessed the emotional bond of affection experienced by the parent towards the infant (39). The 12-item version of the general mental health questionnaire was used to identify psychological distress and short-term changes in parental mental health.

To further improve our understanding of the specific functions associated with neonatal TSH, the MSCA items were reorganised into the following new outcomes for tasks highly associated with specific neurocognitive functions: verbal memory (items 3 and 7II), working memory (items 5 and 14II), memory span or short-term memory (items 6, 7I and 14I), gross motor (items 9, 10 and 11), fine motor (items 12 and 13) and executive function (items 2, 5, 6, 14II, 15, 17 and 18) (32, 37).

**Covariates**

The attending paediatrician and trained interviewers gathered information at delivery and at the 4-year visit respectively on maternal age, alcohol consumption and cigarette smoking during pregnancy, reproductive history, parity, pre-pregnancy body mass index (BMI), duration of breastfeeding, maternal and paternal education, marital status and area of residence (urban: city of Granada; metropolitan: towns of > 20 000 inhabitants in city residential belt; sub-urban: towns of 10 000–20 000 inhabitants; rural: < 10 000 inhabitants). Information on gestational age and anthropometric measurements at birth were obtained from medical records. Covariates considered for inclusion in the statistical analysis were expressed as shown in Table 1.

Complete cognitive outcome data and information on cord blood TSH concentration and covariates were available for 178 subjects from the cohort (n = 220). Information on the former and prenatal exposure to OCs and TEXB in the placenta was available for a subset of 101 of the 178 children (57%) in this study. No and TEXB in the placenta was available for a subset of 178 subjects from the cohort (n = 15) to homogenise the scales. A cut-off point corresponding to the 20th percentile (P20) was used to categorise the outcomes with a non-normal distribution. We used simple linear regression or ANOVA to examine the relationship of covariates with general cognitive scores.

TSH values were transformed into natural logarithms (log transformed) to improve the normality. We used adjusted general additive models (GAM) to evaluate the linearity of the relationship between general cognitive scores and TSH levels, comparing models with TSH in a linear and a non-linear manner (a cubic smoothing spline with 2–4 degrees of freedom) by means of likelihood ratio tests (Fig. 1). Because no significant improvement in the model was obtained with non-linear models, we first treated TSH as a continuous variable. In a second analysis, TSH was categorised into quartiles.

The strength of the unadjusted and adjusted associations between the outcome scores and TSH levels was measured by calculating coefficients (β) and odds ratios (ORs) for linear and logistic regression models. Variables associated with the general cognitive score at a significance level of P < 0.20 in the bivariate analysis or whose inclusion in the models changed TSH effect estimates by > 10% were considered confounders. All multivariate models controlled for maternal age (≥ 32 years) and gestational age (continuous), regardless of their statistical significance. The potential confounding of exposure to OC and TEXB levels in the placenta was examined in a further regression analysis of the association of TSH with general cognitive and executive function outcomes. Concentrations of OCs and TEXB values were categorised using the LD cut-off point, except for p,p′-dichlorodiphenylchloroethylene (p,p′-DDE) (detected in ≥ 90% of the placenta extracts), which was introduced as continuous (ng/g of placenta). In all the cases, the level of significance was P ≤ 0.05. STATA version 9.2 (STATA Corporation, College Station, TX, USA) was used for the analyses.

**Results**

Mean (s.d., range) newborn TSH was 3.55 (2.54, 0.24–17) mU/l, and only three newborns had a TSH level above 14 mU/l (laboratory reference value). Repeated measurement analyses confirmed the TSH levels of these three babies, none of whom were diagnosed with thyroid disorder. Mean (s.d.) infant birth weight was 3289 (487) g, and gestational length was 39 (1.8) weeks; maternal age at delivery was 31 (5) years, and pre-pregnancy BMI was 23.5 (4.0) kg/m². Around 22% of women smoked during pregnancy, 47% were primiparous, 85% breastfed the child, 15% of mothers and 16% of fathers had university education and 64% of the families lived in urban or metropolitan settings. TSH concentrations were slightly higher in infants with lower birth weight (by −0.66 mU/l per kg; 95% confidence interval (CI) = −1.4, 0.11;
$P=0.09$), and infants of fathers with university education were substantially more likely to have lower TSH (by $\text{K}1.27 \text{mU/l}; 95\% \text{CI} \text{K}2.20, \text{K}0.34; P=0.007$) (data not shown). Newborn TSH levels did not differ by maternal age, gestational age, parity or other covariates. Mean (S.D., range) age of the boys at psychological testing was 51 (2, 47–58) months, and their general cognitive score was 100.1 (14.5, 56.9–131) points. This score was significantly higher among older children, urban children, those from primiparous mothers and those from parents with university education, and it was positively associated with gestational age, birth length and mother-to-infant attachment score at 4 years of age (Table 1). Perceptual performance scores ranged between 53.1 and 135; the range of the executive function was 58.3–133; verbal, 57.2–133 (22.5, <P20); quantitative, 61.5–155 (23.6, <P20); memory, 60.4–146 (19.7, <P20); motor, 45.7–131 (20.2, <P20); span memory, 59.2–133 (15.7, <P20); working memory, 76.8–159 (42.7, <P20); verbal memory, 54.4–161 (29.2, <P20); gross motor, 69.2–137 (15.2, <P20) and fine motor, 52.8–134 (15.7, <P20) (data not shown).

Table 2 shows the unadjusted associations between the cord blood TSH levels and children’s cognitive test scores. In accordance with our hypothesis, a higher TSH was related to lower MSCA scores. The risk of scoring
<P20 on the quantitative scale was greater in children with higher TSH levels (OR = 2.25; 95% CI = 1.20, 4.25; \( P = 0.01 \)), but no significant association was found with any MSCA outcome score in crude analyses. By contrast, multivariate models adjusted for maternal and child characteristics showed an association between elevated newborn TSH levels as a continuous measure and poorer characteristics showed an association between elevated newborn TSH levels as a continuous measure and poorer general cognitive scores (by \( P = 0.03 \)).

Figure 1 depicts the GAM for the relationship between log-transformed TSH concentrations and general cognitive scores, showing a positive linear trend. With the exception of \( p,p' \)-DDE, detected in 92.1% of the placenta samples and with a mean (s.d.) concentration of 3.09 (6.50) ng/g placenta, the percentage detection of OC pesticides was <90% (n = 101), ranging from 24.8% (dieldrin) to 84.2% (lindane) (data not shown). The TEXB of the \( \alpha \) and \( \beta \) fractions were above the LD in 67.3 and 83.2% of the placenta extracts respectively. Table 4 shows that the negative associations of cord blood TSH status with the general cognitive score (by \( -6.34; 95\% \text{ CI} = -12.32, -0.36; \( P = 0.04 \)) and executive function score (by \( -7.85; 95\% \text{ CI} = -14.04, -1.67; \( P = 0.009 \)) were also present after simultaneous adjustment for prenatal exposure to the 17 OCs measured. After controlling for TEXB, the association remained significant for the general cognitive score (by \( -5.35, 95\% \text{ CI} = -10.24, -0.45; \( P = 0.03 \)) and was marginally significant for executive function score (by \( -5.05; 95\% \text{ CI} = -10.27, 0.18; \( P = 0.06 \)). However, contrary to expectations, the magnitude of the effect of TSH on cognitive functions was strengthened after adjustment for prenatal OC exposure (change in regression coefficient >10%).

Table 2 Crude coefficients and odds ratios (95% confidence intervals) for the association between cord blood TSH levels and MSCA outcomes, INMA-Granada cohort, 2000–2006 (n = 178).

<table>
<thead>
<tr>
<th>MSCA outcomes at 4 years of age</th>
<th>TSH (mU/l)</th>
<th>( P ) value</th>
</tr>
</thead>
<tbody>
<tr>
<td>General cognitive (( \beta ))</td>
<td>-1.71 (-4.90, 1.49)</td>
<td>0.29</td>
</tr>
<tr>
<td>Verbal (OR)</td>
<td>1.11 (0.62, 1.99)</td>
<td>0.72</td>
</tr>
<tr>
<td>Perceptual performance (( \beta ))</td>
<td>-0.04 (-3.30, 3.22)</td>
<td>0.98</td>
</tr>
<tr>
<td>Quantitative (OR)</td>
<td>2.26 (1.20, 4.25)</td>
<td>0.01</td>
</tr>
<tr>
<td>Memory (OR)</td>
<td>1.17 (0.63, 2.16)</td>
<td>0.61</td>
</tr>
<tr>
<td>Span (OR)</td>
<td>1.19 (0.61, 2.31)</td>
<td>0.62</td>
</tr>
<tr>
<td>Verbal memory (OR)</td>
<td>1.40 (0.82, 2.39)</td>
<td>0.22</td>
</tr>
<tr>
<td>Working memory (OR)</td>
<td>1.55 (0.95, 2.55)</td>
<td>0.08</td>
</tr>
<tr>
<td>Motor skills (OR)</td>
<td>0.71 (0.39, 1.29)</td>
<td>0.26</td>
</tr>
<tr>
<td>Fine motor skills (OR)</td>
<td>1.12 (0.57, 2.18)</td>
<td>0.75</td>
</tr>
<tr>
<td>Gross motor skills (OR)</td>
<td>1.19 (0.58, 2.44)</td>
<td>0.63</td>
</tr>
<tr>
<td>Executive function (( \beta ))</td>
<td>-1.62 (-4.79, 1.56)</td>
<td>0.32</td>
</tr>
</tbody>
</table>

OR, logistic regression odds ratio; \( \beta \), linear regression coefficient. For ORs, the cut-off points correspond to the 20th percentile (reference group >P20). Normal distributed data are standardised (mean: 100; s.d.: 15).

Each row represents a model adjusted for child’s age and school term and psychologist administering the test.

Log-transformed TSH levels.
Table 3 Adjusted regression coefficients and odds ratios (95% confidence intervals) for the effect of cord blood TSH levels (mU/l) on MSCA outcomes, INMA-Granada cohort, 2000–2006 (n=178)a.

<table>
<thead>
<tr>
<th>MSCA outcomes at 4 years of age</th>
<th>Continuous TSHc</th>
<th>2.05–2.95</th>
<th>2.96–4.18</th>
<th>4.19–17.0</th>
</tr>
</thead>
<tbody>
<tr>
<td>General cognitive (β)</td>
<td>−3.52 (−6.81, −0.23)*</td>
<td>−0.36 (−6.29, 5.67)</td>
<td>−3.03 (−8.78, 2.74)</td>
<td>−5.42 (−11.30, −0.61)*</td>
</tr>
<tr>
<td>Verbal (OR)</td>
<td>1.48 (0.66, 3.30)</td>
<td>0.24 (0.05, 1.05)*</td>
<td>0.55 (0.15, 2.05)</td>
<td>1.48 (0.38, 5.39)</td>
</tr>
<tr>
<td>Perceptual performance (β)</td>
<td>−1.41 (−4.87, 1.91)</td>
<td>−1.12 (−7.28, 4.99)</td>
<td>−4.04 (−9.99, 1.84)</td>
<td>−2.17 (−8.31, 3.92)</td>
</tr>
<tr>
<td>Quantitative (OR)</td>
<td>2.64 (1.16, 5.54)c</td>
<td>1.21 (0.31, 4.25)</td>
<td>1.28 (0.32, 4.73)</td>
<td>4.92 (1.30, 16.47)c</td>
</tr>
<tr>
<td>Memory (OR)</td>
<td>1.61 (0.60, 3.80)</td>
<td>0.83 (0.13, 3.60)</td>
<td>0.92 (0.18, 3.74)</td>
<td>3.51 (0.65, 15.26)</td>
</tr>
<tr>
<td>Span (OR)</td>
<td>1.51 (0.52, 3.94)</td>
<td>0.53 (0.07, 3.23)</td>
<td>0.32 (0.05, 2.50)</td>
<td>5.73 (0.72, 2.46)*</td>
</tr>
<tr>
<td>Verbal memory (OR)</td>
<td>1.72 (0.86, 3.41)</td>
<td>0.69 (0.21, 2.29)</td>
<td>1.22 (0.39, 3.71)</td>
<td>1.98 (0.62, 6.30)</td>
</tr>
<tr>
<td>Working memory (OR)</td>
<td>1.35 (0.76, 2.49)</td>
<td>1.03 (0.35, 2.93)</td>
<td>1.34 (0.50, 3.77)</td>
<td>1.47 (0.54, 4.27)</td>
</tr>
<tr>
<td>Motor skills (OR)</td>
<td>0.65 (0.32, 1.47)</td>
<td>0.39 (0.11, 1.87)</td>
<td>0.42 (0.13, 1.73)</td>
<td>0.80 (0.24, 3.10)</td>
</tr>
<tr>
<td>Fine motor skills (OR)</td>
<td>1.48 (0.63, 3.66)</td>
<td>1.68 (0.35, 9.12)</td>
<td>2.14 (0.54, 9.28)</td>
<td>2.31 (0.53, 11.39)</td>
</tr>
<tr>
<td>Gross motor skills (OR)</td>
<td>0.69 (0.26, 1.99)</td>
<td>4.27 (0.69, 28.84)</td>
<td>0.24 (0.02, 2.55)</td>
<td>2.49 (0.44, 16.67)</td>
</tr>
<tr>
<td>Executive function (β)</td>
<td>−3.15 (−6.66, −0.19)*</td>
<td>−0.30 (−6.51, 5.96)</td>
<td>−3.37 (−9.46, 2.56)</td>
<td>−4.29 (−10.56, 1.86)</td>
</tr>
</tbody>
</table>

OR, logistic regression odds ratio; β, linear regression coefficient. For ORs, the cut-off points correspond to the 20th percentile (reference group >P20).
*P<0.05; †P<0.01. Normal distributed data are standardised (mean: 100; s.d.: 15).
*aEach row represents two models: one using continuous TSH and the other using TSH quartiles. All models are adjusted for birth length, gestational age, maternal age, parity, breastfeeding, maternal and paternal education, mother-to-infant attachment, child’s age and school term and psychologist administering the test.
*bLog-transformed TSH levels.

discussion
This study of 178 children in Southern Spain born with normal thyroid function yielded evidence of an association between neonatal TSH and the cognitive development of children at 4 years of age, supporting our study hypothesis. Thus, MSCA general cognitive, quantitative and executive function scores appeared to be impaired by higher TSH cord blood levels. Limited data are available on the effect of newborn thyroid status on neurodevelopment, and most reports of associations have described the influence of thyroid status during pregnancy, specifically in relation to reduced T4 levels (5). In addition, infants of fathers with university education had lower TSH, suggesting that neonatal thyroid status may be affected by social conditions. Paternal education and cord blood TSH levels contributed to predict cognitive performance at 4 years of age, consistent with the findings in research on ID that maternal education has a protective role in infant development (40).

The relationship between TH or TSH and cognitive function has mainly been studied in children with congenital hypothyroidism, children of mothers with low TH concentrations during pregnancy or children living in ID areas (41). A number of case–control studies have reported associations between decreased maternal or neonatal T4 and/or triiodothyronine (T3) levels and poorer neurodevelopment in children born to mothers with hypothyroxinaemia during pregnancy or in ex-preterm infants, among others (11, 12, 42, 43). Detected neurocognitive deficits include attention deficits (11, 43), reduced mental and motor development scores (12, 42, 43), impaired intelligence and language skills and difficulties in school performance at later ages (11). Interestingly, a cross-sectional analysis of 334 healthy children from two general population cohorts at 4 years of age (from Menorca and Ribera d’Ebre; INMA study) found an association between higher serum TSH levels (2.43–5.01 mU/l) and delayed general cognitive, quantitative, memory, verbal and
perceptual performance MSCA outcomes (44), consistent with the present findings.

By contrast, other authors reported that higher newborn T\textsubscript{4} was unexpectedly associated with lower scores on the visual recognition memory test at the age of 6 months but not with scores for verbal abilities, intelligence or visual motor abilities at the age of 3 years (45). Another study observed no neurological impairment in infants aged < 1 year born to mothers with elevated TSH during pregnancy (46). Furthermore, neonatal T\textsubscript{4} levels were not associated with the risk of a heterogeneous group of developmental diagnoses in 5–12-year-old children, including attention deficit disorder, autism spectrum disorder, behavioural disorder, cognitive disorder, developmental delay, emotional disorder, learning disability and speech/language disorder (47). A recent case-control study in Southern Spain observed a superior psychometric and behavioural development among children whose mothers had received iodine supplementation compared with the children of non-supplemented mothers: cord blood TSH was significantly higher in infants of supplemented mothers (16). A study of Sicilian children reported that a mild to moderate ID was associated with a reduced IQ and attention deficit hyperactivity disorder (13). Finally, increases in the risk of delay in gross and fine motor coordination and socialisation in 18-month infants were found to result from a period of isolated hypothyroxinaemia in pregnant women from a coastal region in Spain with mild ID (15).

The timing of TH action is crucial for neurodevelopment, and the effects of TH status may therefore differ among pregnant women, neonates and children. In the neonatal and postnatal periods, neurological development still depends on THs, whose supplies to the brain are entirely derived from the child and are critical for continuing maturation (6). It has been demonstrated that THs during early pregnancy influence later child development, although the neurological effects of THs may be less severe in neonates. This study demonstrated that higher newborn T\textsubscript{4} levels within a normal reference range were related to lower intelligence, as measured by the general cognitive score, and to impairment of higher psychological processes (executive function) at the age of 4 years. These findings support the view that moderate alterations in neonatal thyroid status may play a role in subsequent neurodevelopment. Even subtle cognitive delays at this age may lead to sub-clinical but permanent decreases in IQ and to long-lasting effects on educational and social development (48). Hence, they should be considered clinically relevant, since early identification of sub-optimal cognitive functioning is necessary to adopt preventive measures.

The study limitations include the fact that only boys were studied; the non-assessment of behavioural or psychopathological outcomes such as attention deficit, social or emotional disorders, which have also been associated with early thyroid status (15, 43, 44) and the absence of measurements of newborn T\textsubscript{4}, T\textsubscript{3} or thyroid axis hormones other than TSH, which would have yielded complete information on the newborn’s thyroid regulatory system. In fact, recent studies have reported the unexpected finding of a positive correlation between TSH and free T\textsubscript{4} levels in cord blood correlate (14, 49), suggesting that TSH elevations should not necessarily be interpreted as indicating a potentially harmful effect on the child. There is a progressive modulation of the set point for T\textsubscript{4} negative feedback regulation of TSH secretion in infants, which implies that TSH production is overstimulated during gestation, decreasing from postnatal age of around 2 weeks (49). Owing to this decline in TSH levels over the first days of life, newborns with elevated TSH should be evaluated for congenital hypothyroidism with repeat TSH and free T\textsubscript{4} measurements. However, it is recognised that TSH measurement offers higher sensitivity to detect thyroid dysfunction in comparison with T\textsubscript{4} or T\textsubscript{3} testing, since subtle alterations in T\textsubscript{4} or T\textsubscript{3} within the normal reference range may result in an amplified alteration of TSH (50).

The study strengths include the longitudinal design, the considerable number of covariates considered (e.g. breastfeeding and parental attachment) and, most importantly, the fact that we examined the association between TSH and neurodevelopment in typically developing children from the general population. In addition, this is the first report to evaluate the potential confounding effect of prenatal exposure to a wide range of ED, environmental chemicals and their combined oestrogenic effect on the association under study.

To date, epidemiological studies have described associations of early exposure to OC compounds with TH levels (20–26) and with neurodevelopment impairment (51–53). However, a consistent influence on thyroid status and neurodevelopment of many OC compounds at environmental background levels has not been established yet. Humans may be exposed to mixtures of these and numerous other compounds, hampering the prediction of effects on TH levels. We have previously demonstrated the ubiquity of exposure to OC xeno-oestrogens in the INMA-Granada cohort (54). In this study, prenatal exposure to 17 OC pesticides or the xeno-oestrogenic burden in the placentas were observed to modify the impact of neonatal TSH on neurodevelopment, in agreement with the suspected capacity of OCs to interfere with the thyroid system (20–26).

In conclusion, this study of a birth cohort in Southern Spain revealed an impaired mental development at 4 years of age in children with higher neonatal TSH levels compared with children with lower neonatal TSH levels within the normal reference range. These findings indicate that a more thorough screening for neonatal thyroid deficiency is required to prevent long-term developmental effects. Further research is warranted into the influence on neurodevelopment of marginally altered TSH concentrations in newborns and into...
causal relationships between ED, environmental chemicals and TH status.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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References
between thyroid hormone levels and 4,4'-DDE concentrations in pregnant women (Valencia, Spain). Environmental Research 2009 109 479–485. (doi:10.1016/j.envres.2009.02.003)


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