Subclinical hypothyroidism in atopic South Italian children

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Abstract

AIM: To verify if subclinical hypothyroidism (SCH) could be associated to atopy in children.

METHODS: Seven hundred and thirty-two Caucasian children from South Italy presenting symptoms of allergic disease were enrolled and submitted to atopy, obesity, chronic low grade inflammation, and SCH work up.

RESULTS: Four hundred and forty-five out of 705 (63.12%) children affected by allergic disease were diagnosed as atopic and 260 (36.88%) as not atopic. The SCH prevalence was 6.3%. Significant higher prevalence of SCH among atopic children with average (group 2) and high (group 3) low grade chronic inflammation compared to atopic children with mild (group 1) low grade chronic inflammation was present. Moreover, group 1 and group 2 presented an OR to show SCH of 2.57 (95%CI: 1.55-6.26) and 2.96 (95%CI: 1.01-8.65), respectively. Both in atopic and not atopic children we found C3 serum levels significantly higher in group 3 respect to group 2 and group 1 children. Noteworthy, among atopic patients, also total immunoglobulin E (IgE) serum levels, were significantly higher in group 3 compared to group 2 and group 1 children. In atopic children, C3 and total IgE serum values increased in parallel with the increase of C-reactive protein values, while in not atopic children this phenomenon was not evident.

CONCLUSION: The possibility exists that an increasing atopic inflammation contributes to SCH occurrence. So far this is the first report in literature showing an
association between SCH and atopy but further studies are needed to confirm our data.

**Key words:** Thyroid derangement; Atope; Children; Low grade chronic inflammation; Subclinical hypothyroidism

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Core tip: A high subclinical hypothyroidism (SCH) prevalence has been associated in childhood to obesity and the chronic low grade inflammation found obese children has been involved in this relationship. In our population, obesity does not influence SCH prevalence. Interestingly, we found a SCH prevalence twice higher compared to all other patients and a significant higher risk to show SCH in atopic children affected by the highest C-reactive protein values characterizing low-grade inflammation.

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INTRODUCTION

Subclinical hypothyroidism (SCH) is defined as a thyroid stimulating hormone (TSH) serum level above the statistically defined upper limit of the reference range despite normal serum free T4 (fT4) and free T3 (fT3) concentration. In addition to genetic cause, SCH has been associated to childhood obesity[1], with some slight myocardial dysfunction[2].

It has been recently demonstrated that atopy can be related to childhood obesity and that chronic low grade inflammation related to it was involved in this relationship[3-5].

The aim of our study is to verify if, as well as to obesity, SCH could be also associated to atopy in children.

MATERIALS AND METHODS

Seven hundred thirty-two Caucasian children from South Italy (age in years 6.03 ± 3.63) presenting symptoms of allergic disease (erythema, pruritus, eczematous rash, wheals, ocular and nasal pruritus and wheezing) attending consecutively from January 2013 to July 2015 the Department of Pediatrics of the Second University of Naples were enrolled. None showed any clinical symptoms of thyroid disease. All the enrolled children were submitted to atopy, obesity, chronic low grade inflammation, and SCH work up. Body mass index (BMI) was calculated by dividing weight in kilograms by height squared in meters (kg/m²). Obesity was defined by a BMI ≥ 95th percentile[6].

In all patients, after an overnight fasting, a blood sample was obtained to evaluate C-reactive protein (CRP) and Complement C3 serum levels, measured using an Olympus AU 560 apparatus by an enzymatic colorimetric method, total and specific IgE by a fluorenszymeimmunoassay (ImmunoCap and ImmunoCap 0-100), thyroid hormones (TSH, fT3, and fT4) and anti-thyroid peroxidase antibodies (TPO-Ab) and anti-thyroglobulin antibodies (Tg-Ab), determined by high-specific solid-phase technique-chemiluminescence immunoassays (Perkinelmer, Turku, Finland).

The diagnosis of atopy suspected for clinical history and symptoms, was confirmed by levels of serum total IgE (normal value from 25 to 60 mo of age < 81 kU/L, from 61 to 156 mo of age < 101 kU/L) and specific immunoglobulin E (IgE) assay (> 0.36 kU/A/L) as well as by Skin Prick Tests (SPTs). None had taken steroids or received immuno-suppressive therapy for at least 3 mo before investigation. Antihistamine therapy had been stopped at least 2 wk before SPTs were performed and serum samples were collected.

According to atopy diagnosis the children enrolled in the study were divided in two groups: Atopic and not atopic children.

Chronic low grade inflammation is diagnosed by slightly raised concentrations of inflammatory markers in the systemic circulation[7]. Therefore, to better assess the chronic low grade inflammation status we applied both in atopic and not atopic children the cut-off point described by Pearson et al[8] for assessing cardio vascular disease (CVD) risk. Thus we divided children in three different CRP serum values gradation groups: group 1 low grade (CRP < 0.1 mg/dL), group 2 average grade (CRP 0.1-0.3 mg/dL) and group 3 high grade (CRP > 0.3- < 1 mg/dL) of chronic low grade inflammation.

Twenty-seven children with CRP serum values greater than 1 mg/dL were excluded from the study given the possibility of an ongoing infection.

SCH was diagnosed when TSH value was higher than 5 mUI/mL, as indicated by the “European Thyroid Association Guidelines for the Management of Subclinical Hypothyroidism in Pregnancy and in Children”[9,10], with fT3, fT4, and antithyroid antibodies values in the normal range for age and no clinical signs or symptoms of hypothyroidism[10].

Informed consent was obtained from all enrolled children and their parents and in accordance with the approved protocol from the Institutional Review Board at the University of Naples.

Skewness and Kurtosis tests were used to evaluate if the distribution of continuous variables was normal. According to distribution, the values were expressed as mean ± SD or median and minimum-maximum values. To analyze categorical variables was used t test for unpaired data. A χ² test was also used to analyze the differences between the frequencies. Mann-Whitney U test was used for comparison of continuous variables which did not exhibit normal distribution. A P value < 0.05 was considered significant.
Odds ratio (OR) was calculated to evaluate the association of Atopy and SCH prevalence. OR was considered significant when a 95%CI excluded unity. Statistical analysis were performed using Stat-Graphics Centurion 3.0 for Windows.

RESULTS

On the basis of atopy work-up 459 out of 705 (62.7%) children affected by allergic disease were diagnosed as atopic and 273 (37.3%) as not atopic. The overall prevalence of SCH was 6.3%.

Table 1  Clinical and laboratory differences between atopic and not atopic children affected by allergic disease

|                      | Atopic children        | Not atopic children   | P      |
|----------------------|------------------------|-----------------------|--------|
| Patients, n          | 459/732 (62.7%)        | 271/732 (37.3%)       |        |
| Age expressed in years (mean age ± SD) | 6.24 ± 3.55           | 5.82 ± 3.71           | 0.128  |
| Gender (male)        | 245/459 (53.37%)       | 137/274 (50%)         | 0.37   |
| Family history of atopy | 365/420 (86.9%)   | 195/235 (80%)         | 0.17   |
| Family history of thyroid disease | 191/421 (45.36%) | 94/235 (40%)         | 0.18   |
| BMI (kg/m²)          | 16.6 (10.9-36.8)       | 16.4 (10.7-30.17)     | 0.069  |
| SCH affected (%)     | 33/445 (7.41%)         | 13/260 (5%)           | 0.21   |
| CRP (mg/dL)          | 0.06 (0-10.3)          | 0.08 (0-7.9)          | 0.123  |
| TSH (UI/mL)          | 2.33 (0.35-12.2)       | 2.24 (0.52-6.98)      | 0.269  |

P value < 0.05 was considered significant. BMI: Body mass index; SCH: Subclinical hypothyroidism; CRP: C-reactive protein; TSH: Thyroid stimulating hormone.

Table 2  Atopic and not atopic children divided into three groups on the basis of the C-reactive protein serum levels

|                      | CRP (mg/dL)          | TSH (UI/mL)          | P      | CRP (mg/dL)          | TSH (UI/mL)          | P      |
|----------------------|----------------------|----------------------|--------|----------------------|----------------------|--------|
|                      | < 0.1                | > 0.3                |        | < 0.1                | > 0.3                |        |
| Atopic children      | 18/272 (6.2%)        | 8/52 (15.38%)        |        | 8/151 (5.29%)        | 2/28 (7.14%)         |        |
|                      | (6.62%)              | (5.78%)              |        | (5.29%)              | (3.70%)              |        |
|                      | 115 (20-174)         | 138 (103-192)        |        | 116.5 (10-170)       | 126 (73-177)         |        |
|                      | 122 (50-172)         |                      |        | 139 (109-170)        |                      |        |
|                      | 123.5 (100-4328)     |                      |        |                      |                      |        |
|                      | 162.1 (124-5000)     |                      |        |                      |                      |        |
| Not atopic children  | 110 (0.1-2270)       | 116.5 (10-170)       | 1.86e-8 | 110 (0.1-2270)       | 116.5 (10-170)       | 1.86e-8 |
|                      | 126 (73-177)         | 126 (73-177)         | 0.027  | 126 (73-177)         | 126 (73-177)         | 0.000053 |
|                      | 139 (109-170)        | 139 (109-170)        | 0.0021 | 139 (109-170)        | 139 (109-170)        | 0.0024  |
|                      | 174.5 (1.9-98)       | 224.5 (1.9-93)       | 0.00017| 174.5 (1.9-98)       | 224.5 (1.9-93)       | 0.00045 |
|                      | 224.3 (1.9-88.9)     | 224.3 (1.9-88.9)     | 0.024  | 224.3 (1.9-88.9)     | 224.3 (1.9-88.9)     | 0.024   |

1Group 1 vs group 3; 2Group 1 vs group 2; 3Group 2 vs group 3. SCH: Subclinical hypothyroidism; CRP: C-reactive protein.

Figure 1  Complement C3 and total IgE serum levels values according to low grade inflammation in both atopic (A) and not atopic (B) children affected by allergic disease. CRP values < 0.1 mg/dL identify children with low grade of chronic low grade inflammation (group 1). CRP values between 0.1 and 0.3 mg/dL identify children with average grade of chronic low grade inflammation (group 2). CRP values > 0.3 and < 1 mg/dL identify high grade of chronic low grade inflammation (group 3). In atopic children, C3 and total IgE serum values increased in parallel with the increase of CRP values, while in not atopic children this phenomenon is not evident. A: For C3, group 1 vs group 3, \( P = 1.86 \times 10^{-8} \); group 1 vs group 2, \( P = 0.027 \); group 2 vs group 3, \( P = 0.000021 \). For total IgE, group 1 vs group 3, \( P = 0.000017 \); group 1 vs group 2, \( P = 0.05 \); group 2 vs group 3, \( P = 0.000045 \); B: For C3, group 1 vs group 3, \( P = 0.000053 \); group 1 vs group 2, \( P = 0.0085 \); group 2 vs group 3, \( P = 0.024 \). For total IgE, group 1 vs group 3, \( P = 0.05 \); group 1 vs group 2, \( P = 0.05 \); group 2 vs group 3, \( P = 0.05 \).

Odds ratio (OR) was calculated to evaluate the association of Atopy and SCH prevalence. OR was considered significant when a 95%CI excluded unity. Statistical analysis were performed using Stat-Graphics Centurion 3.0 for Windows.
group 1 (CRP < 0.1 mg/dL), group 2 (CRP 0.1-0.3 mg/dL) and group 3 (CRP > 0.3<- 1 mg/dL). First we compared the prevalence of SCH among all the groups and no significant differences were found.

On the contrary, among atopic children, significant higher prevalence of SCH in group 3 and group 2 compared to group 1 was found ($P = 0.033$ and $P = 0.04$, Table 2). Moreover, group 3 and group 2 atopic children presented an OR to show SCH of 2.57 (95%CI: 1.55-6.26) and 2.96 (95%CI: 1.01-8.65) respectively.

Both in atopic and not atopic children we found C3 serum levels significantly higher in group 3 respect to group 2 and group 1 (Table 2). Noteworthy, among atopic patients, also total IgE serum levels, were significantly higher in group 3 compared to group 2 and group 1 children.

Figure 1 shows in both atopic (A) and not atopic (B) children complement C3 and total IgE serum levels values according to the different grading of CRP values.

In atopic children, C3 and total IgE serum values increased in parallel with the increase of CRP values, but no significant correlation was found. C3 and total IgE serum values in not atopic children showed a completely independent trend.

**DISCUSSION**

In our study atopic children showed a 7.41% prevalence of SCH while not atopic children allergic found in alike disease affected a 5% SCH prevalence. These frequencies are in line with the estimated prevalence of this thyroid disorder in the pediatric population[14,15].

A high SCH prevalence has been associated in childhood to obesity and the chronic low grade inflammation found obese children has been involved in this relationship[13]. In our population, obesity does not influence SCH prevalence.

Interestingly, we found a SCH prevalence twice higher (15.38%) compared to all other patients and a significant higher risk to show SCH [2.57 (95%CI: 1.55-6.26)] in atopic children affected by the highest CRP values characterizing low grade inflammation (group 3 atopics).

Atopy and the associated allergic disease are now regarded as systemic inflammatory disease that could affect the risk of atherosclerosis[14], impaired glucose tolerance[15], and coronary artery disease[16]. The chronic low grade inflammation involved in the pathogenesis of localized allergic disease causes a systemic inflammatory response that potentially could promote also SCH.

Moreover, in our atopic children the highest low grade chronic inflammation (CRP > 0.3<- 1 mg/dL) seems to correspond to the highest atopic inflammation as measured by total serum IgE values (Figure 1). The possibility exists that an increasing atopic inflammation contributes to SCH occurrence. A limitation of our study could be a recruitment bias because the patients enrolled were affected severe allergic disease needing of a specialist evaluation.

So far this is the first report in literature showing an association between SCH and atopy but further studies are needed to confirm our data.

**REFERENCES**

1. Grandone A, Santoro N, Coppola F, Calabrò P, Perrone L, Del Giudice EM. Thyroid function derangement and childhood obesity: an Italian experience. BMC Endocr Disord 2010; 10: 8 [PMID: 20441588]
2. Brienza C, Grandone A, Di Salvo G, Corona AM, Di Sessa A, Pascotto C, Calabrò R, Toraldo R, Perrone L, del Giudice EM. Subclinical hypothyroidism and myocardial function in obese children. Nutr Metab Cardiovasc Dis 2013; 23: 896-902 [PMID: 22748710 DOI: 10.1016/j.numecd.2012.04.006]
3. von Mutius E, Schwartz J, Neas LM, Dockery D, Weiss ST. Relation of body mass index to asthma and atopy in children: the National Health and Nutrition Examination Study III. Thorax 2001; 56: 835-838 [PMID: 11641506 DOI: 10.1136/thorax.56.11.835]
4. Ford ES. C-reactive protein concentration and cardiovascular disease risk factors in children: findings from the National Health and Nutrition Examination Survey 1999-2000. Circulation 2003; 108: 1053-1058 [PMID: 12925465 DOI: 10.1161/01.CIR.0000080913.81393.B8]
5. Cibella F, Cuttica G, La Grutta S, Melis MR, Bucchi S, Viggio G. A cross-sectional study assessing the relationship between BMI, asthma, atopy, and eNO among schoolchildren. Ann Allergy Asthma Immunol 2011; 107: 330-336 [PMID: 21962093 DOI: 10.1016/j.anai.2011.08.001]
6. Cacciari E, Milani S, Balsamo A, Spada E, Boni G, Cavallo L, Cerutti F, Gargantini L, Greggio N, Tonini G, Ciocognani A. Italian cross-sectional growth charts for height, weight and BMI (2 to 20 yr). J Endocrinol Invest 2006; 29: 581-593 [PMID: 16957465 DOI: 10.1007/BF03344156]

**COMMENTS**

**Background**

Recent studies demonstrate that atopy can be associated to childhood obesity and that chronic low grade inflammation could be involved in this relationship. The aim of the study was to verify if subclinical hypothyroidism (SCH) could be also associated to atopy in children.

**Research frontiers**

Important areas of research related to the study are represented by the field of pediatric endocrinology and allergology. More in particular, the study aimed to hypothesize, throughout the verification of an association, the mechanisms by which atopy could lead to SCH.

**Innovations and breakthroughs**

This is the first report in literature showing an association between SCH and atopy with a possible causal link represented by chronic low grade inflammation.

**Applications**

The study can result in future researches confirming the authors’ findings and, moreover, understanding the pathophysiological basis underlining the association between atopy and SCH.

**Terminology**

SCH is defined as a thyroid stimulating hormone serum level above the statistically defined upper limit of the reference range despite normal serum free T4 and free T3 concentration.

**Peer-review**

The association among obesity, chronic inflammation and SCH is a very interesting topic.
Pedullà M et al. Subclinical hypothyroidism and atopy

7 Trayhurn P, Wood IS. Adipokines: inflammation and the pleiotropic role of white adipose tissue. Br J Nutr 2004; 92: 347-355 [PMID: 15469638 DOI: 10.1079/BJN20041213]

8 Pearson TA, Mensah GA, Alexander RW, Anderson JL, Cannon RO, Craigh M, Fadl Y, Fortmann SP, Hong Y, Myers GL, Rifai N, Smith SC, Taubert K, Tracy RP, Vinicor F. Markers of inflammation and cardiovascular disease: application to clinical and public health practice: A statement for healthcare professionals from the Centers for Disease Control and Prevention and the American Heart Association. Circulation 2003; 107: 499-511 [PMID: 12551878]

9 Lazarus J, Brown RS, Daumerie C, Hubalewska-Dydejczyk A, Negro R, Vaidya B. 2014 European thyroid association guidelines for the management of subclinical hypothyroidism in pregnancy and in children. Eur Thyroid J 2014; 3: 76-94 [PMID: 25114871 DOI: 10.1159/000362597]

10 Taubner K, Schubert G, Pulzer F, Pfäeffle R, Körner A, Dietz A, Thiery J, Kieß W, Kratzsch J. Serum concentrations of anti-thyroid peroxidase and anti-thyroglobulin antibodies in children and adolescents without apparent thyroid disorders. Clin Biochem 2014; 47: 3-7 [PMID: 24103918 DOI: 10.1016/j.clinbiochem.2013.09.017]

11 Gawlik A, Such K, Dejner A, Zachurzok A, Antonz A, Malecka-Tendera E. Subclinical hypothyroidism in children and adolescents: is it clinically relevant? Int J Endocrinol 2015; 2015: 691071 [PMID: 25882992 DOI: 10.1155/2015/691071]

12 Wu T, Flowers JW, Tudiver F, Wilson JL, Panayasvatsat N. Subclinical thyroid disorders and cognitive performance among adolescents in the United States. BMC Pediatr 2006; 6: 12 [PMID: 16623938]

13 Wärnberg J, Nova E, Romeo J, Moreno LA, Sjöström M, Marcos A. Lifestyle-related determinants of inflammation in adolescence. Br J Nutr 2007; 98 Suppl 1: S116-S120 [PMID: 17922948]

14 Knollach M, Kiechl S, Mayr A, Willeit J, Poewe W, Wick G. Allergic rhinitis, asthma, and atherosclerosis in the Bruneck and ARMY studies. Arch Intern Med 2005; 165: 2521-2526 [PMID: 16314550]

15 Wang Z, Zhang H, Shen XH, Jin KL, Ye GF; Qiu W, Qian L, Li B, Zhang YH, Shi GP. Immunoglobulin E and mast cell proteases are potential risk factors of impaired fasting glucose and impaired glucose tolerance in humans. Ann Med 2013; 45: 220-229 [PMID: 23110545 DOI: 10.3109/07853890.2012.732234]

16 Deliargyris EN, Upadhyya B, Sane DC, Dehmer GJ, Pye J, Smith SC, Boucher WS, Theoharides TC. Mast cell tryptase: a new biomarker in patients with stable coronary artery disease. Atherosclerosis 2005; 178: 381-386 [PMID: 15694948]

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