

## ORIGINAL ARTICLE

# A frequent oligogenic involvement in congenital hypothyroidism

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## Abstract

Congenital hypothyroidism (CH), the most frequent form of preventable mental retardation, is predicted to have a relevant genetic origin. However, CH is frequently reported to be sporadic and candidate gene variations were found in <10% of the investigated patients. Here, we characterize the involvement of 11 candidate genes through a systematic Next Generation Sequencing (NGS) analysis. The NGS was performed in 177 unrelated CH patients (94 gland-in-situ; 83 dysgenesis) and in 3,538 control subjects. Non-synonymous or splicing rare variants (MAF < 0.01) were accepted, and their functional impact was predicted by a comprehensive bioinformatic approach and co-segregation studies. The frequency of variations in cases and controls was extended to 18 CH-unrelated genes. At least one rare variant was accepted in 103/177 patients. Monogenic recessive forms of the disease were found in five cases, but oligogenic involvement was detected in 39 patients. The 167 variations were found to affect all genes independently of the CH phenotype. These findings were replicated in an independent cohort of additional 145 CH cases. When compared to 3,538 controls, the CH population was significantly enriched with disrupting variants in the candidate genes ( $P = 5.5 \times 10^{-7}$ ), but not with rare variations in CH-unrelated genes. Co-segregation studies of the hypothyroid phenotype with multiple gene variants in several pedigrees confirmed the potential oligogenic origin of CH. The systematic NGS approach reveals the frequent combination of rare variations in morphogenetic or functional candidate

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genes in CH patients independently of phenotype. The oligogenic origin represents a suitable explanation for the frequent sporadic CH occurrence.

## Introduction

Congenital hypothyroidism (CH) is the most common endocrine developmental disorder and preventable cause of mental retardation (1). A tendency toward an increased CH incidence or detection has been described in various socio-economically developed countries (2–4). Both functional and developmental defects can account for CH but its etiology is still poorly understood (5,6).

It is classically described to be sporadic but several findings in humans and experimental models support a relevant genetic origin. CH is currently considered as a puzzle of monogenic diseases and the investigations on its pathogenesis were driven by phenotype and performed by Sanger sequencing (5–10). Indeed, pathogenic variations in candidate genes have been found in <10% of the cases (5), but genetic studies targeting specific phenotypes or particular ethnic groups yielded higher mutation detection rates (11–14). On these bases, CH can be classified as a disease with a strong genetic component, but with a largely missing heritability (15).

Next generation sequencing (NGS) allows the simultaneous and systematic analysis of candidate genes in unselected disease populations, likely opening novel perspectives in the diagnosis and classification of several diseases (15,16), but no data are available on the application of this strategy in a large number of CH patients.

Here, we systematically analyzed by NGS a panel of 11 candidate genes in 177 unrelated patients with CH, the identified variations were pathogenically classified by a comprehensive bioinformatic approach and genotype-phenotype co-segregation studies in families. The frequency of accepted gene variants (AVs) was then checked in a large control sample of 3,538 subjects from the same geographic area and in an independent CH cohort ( $n = 145$ ).

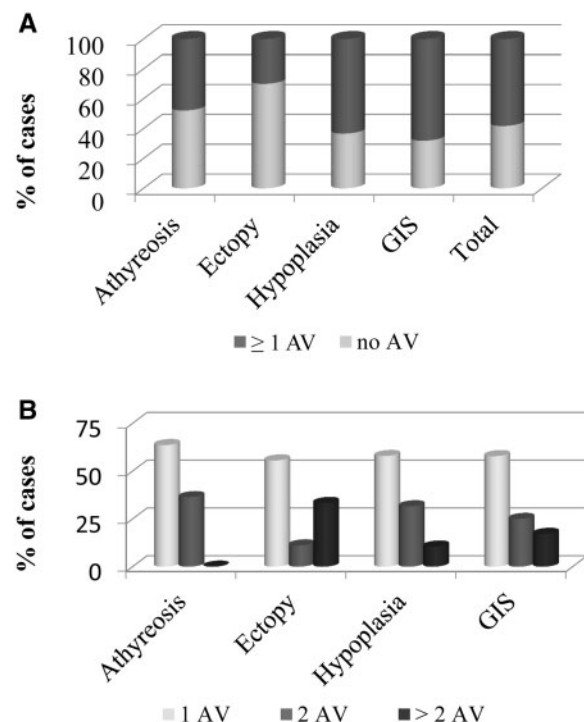
## Results

The NGS analysis of the 11 candidate genes in 177 unrelated CH patients identified 167 allelic variants (116 AVs with  $MAF < 0.01$  and 51 novel AVs; 23 AVs were found more than once in homozygosity or in combination with other variations)(Supplementary Material, Table S2). At least one variant was identified in 103/177 patients (58.2%). Variations were found in all CH categories, but patients with gland-in-situ (GIS) and hypoplasia tended to be more frequently affected (Fig. 1A). A significant number of patients have  $\geq 2$  AVs in the same or in different candidate genes (44; 24.8%) (Fig. 1B). Five of these cases had biallelic (homozygous or compound heterozygous) gene variants with a phenotype corresponding to a recessive mode of inheritance: one hypoplasia associated with a compound heterozygosity in *FOXE1* (p.S152W/p.Y192LfsX37), another hypoplasia with a compound heterozygosity in *TSHR* (p.I640L/p.R310H), and three GIS: one with a homozygous *TG* variant (p.T411HfsX80), one with a compound heterozygosity in *DUOX2* (p.R516H/p.D870A), and one with compound heterozygosity in *TPO* gene (p.A489T/p.E799K). These five patients with monogenic disease were excluded from further analyses. The list of identified variants is reported in Supplementary Material, Table S2, and their distribution in CH carriers is reported in Supplementary Material, Figure S1. A variable number of AVs were detected in the

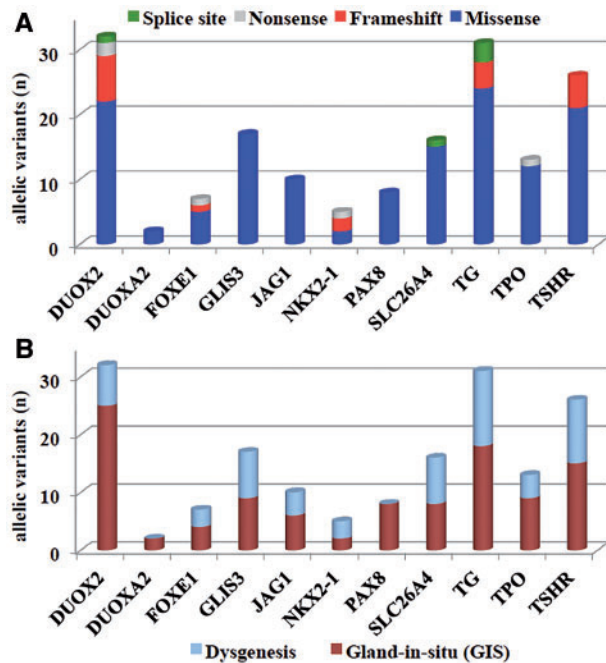
11 candidate genes and most of them were missense variants (Fig. 2A). The genes affected with the higher number of AVs were *DUOX2*, *TG* and *TSHR*. The eight *PAX8* and two *DUOX2* AVs were found exclusively in 10 patients with GIS, but all other genes, independently of their role in thyroid morphogenesis or function, were associated either with the dysgenesis or GIS phenotype (Fig. 2B). The combinations of gene variants appear to occur casually (Supplementary Material, Fig. S1), and biallelic variations in two functional genes, such as *TG* or *SLC26A4*, were associated with thyroid dysgenesis in CH cases #7 (athyreosis) and #14 (ectopy) that are also affected with extra-thyroid manifestations (Supplementary Materials, Fig. S1 and Table S3).

The distribution of extra-thyroid manifestations among the patients with or without AVs is reported in Supplementary Material, Table S3. As expected, choreoathetosis and/or hypotonia were found in three carriers of *NKX2-1* AVs and heart malformations in two carriers of *JAG1* defects; however, *SLC26A4* variations were surprisingly prevalent in CH patients with concomitant heart defects (7/20 cases).

The frequency of potential pathogenic variations within the same 11 candidate genes was assessed in a cohort of Italian individuals. The analysis of the principal components revealed that the case and control populations were comparable (data not shown). When considering all the AVs, a relevant number of control subjects (43.5%) resulted to be carrier of  $>1$  rare variant, however only a smaller fraction (11.7%) was carrier of multiple



**Figure 1.** Panel A: Percent distribution of patients with or without accepted variants (AVs) according to the CH category. The percentage of positive cases tends to be lower in the patients with thyroid ectopy or athyreosis. Panel B: Percent distribution of AVs according to the type of CH and their association in single cases. Multiple gene involvements are seen in all CH categories.



**Figure 2.** Panel A: number of AVs with deleterious effect or prediction (loss-of-function by nature: frameshift, nonsense, splicing variants; and missense variants predicted to be deleterious by  $\geq 5$  out of 7 algorithms) identified in each candidate gene. Variants were indeed found more frequently in larger genes, such as TG, DUOX2 or TSHR. Panel B: number of variants identified in each single gene according to the CH phenotype. Genes typically associated with functional defects (TG, TPO, SLC26A4, TSHR, DUOX2) were also found mutated in cases with thyroid dysgenesis, and viceversa genes typically associated with dysgenesis (FOXE1, JAG1, NKX2-1) were found mutated in cases with GIS.

AVs, if compared to the CH group (23.2%). The overall burden of variants resulted statistically different between cases and controls ( $P = 1.2 \times 10^{-6}$ ), consistent with an enrichment of the rare variations within the CH population (Fig. 3). Moreover, a significantly higher number of CH patients resulted to be a carrier of at least one disruptive AV (frameshift, nonsense and splicing variants, or missense variants previously reported to be loss-of-function in functional tests or familial linkage, or predicted to be deleterious by at least 5 out of the 7 used algorithms) in comparison with controls (32.9 vs 19.1%;  $P = 5.5 \times 10^{-7}$ , Fig. 3).

To confirm the specificity of these results, we performed a parallel analysis of 18 CH-unrelated genes in which again the distribution of rare variants was compared between cases and controls. As expected in this analysis, no significant difference in the mutation burden between the two groups was detected ( $P = 0.1982$ , Fig. 3).

The analysis of the AVs in nine familial settings reveals variable combinations of inherited and de novo variations in the CH probands that are consistent with an oligogenic model of the disease. Inherited multiple rare variants in distinct genes from the two parents were either affected with mild forms of non-autoimmune hypothyroidism or apparently unaffected (see details in Fig. 4).

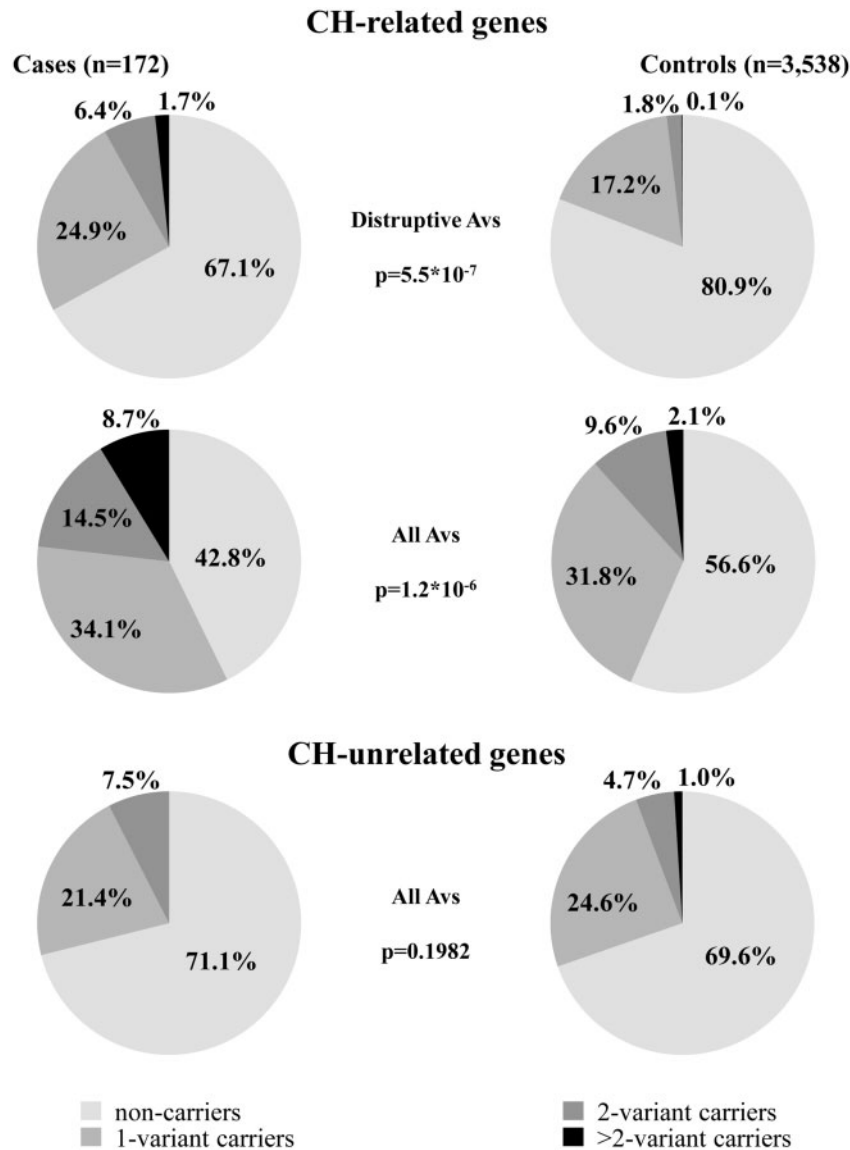
Finally, the NGS analysis was replicated in an independent cohort of 145 CH patients: we found 91 carriers (62.7%) of  $\geq 1$  AVs in the 11 genes. At least two AVs were detected in 41/145 CH patients (28.3%); since 3/41 had monogenic defects, oligogenic defects were found in 38/145 patients (26.2% of total).

## Discussion

The systematic NGS analysis of 11 known candidate genes provides evidence that inheritable variations with a proven or predicted loss-of-function effect can be found in a large portion of CH patients. According to what was anticipated in a mouse model (17), we show an oligogenic involvement in 22% of total cases, with a likelihood of oligogenicity among mutated patients of about 42.7%. Importantly, these results were replicated in an independent CH cohort and CH candidate gene variations were found also in the control population, but with a significantly lower prevalence. Thus, our results indicate that the pathogenesis of CH can frequently be due to the sum of rare (MAF  $< 0.01$ ) alleles (15) that generate minor functional impairments and are not expected to be associated individually with an overt phenotype. This oligogenic origin of CH is confirmed by the co-segregation of variants with CH or non-autoimmune hypothyroid state in several families. Therefore, our findings give a significant contribution to the missing heritability of CH, and may constitute a suitable explanation for the variable modes of inheritance described in several pedigrees (18), as well as for the minor glandular defects detected in several relatives of CH patients (19).

So far, monogenic defects had been described in rare familial forms of CH, mainly associated with biallelic LOF mutations in functional genes (5–7,9,13,20–22). Instead, heterozygous mutations in thyroid transcription factor genes were associated with thyroid defects with variable penetrance and expressivity, but several phenotype driven studies with previous screening methods (such as denaturing gel gradient electrophoresis or Sanger sequencing) in large CH cohorts reported a poor rate of positive results (5–7,23–25). Our approach allowed the identification of classic recessive monogenic CH forms in five cases, but unexpected variations in genes that would have been excluded a priori from screening on the basis of the clinical phenotype. The AVs involve all the examined genes with different frequencies and independently if they were previously linked to dysgenetic or functional defects, thus justifying the previous low rate of variant detection following the classic phenotypical approach. It is worth noting that we identified biallelic mutations of FOXE1 in a CH case affected with thyroid hypoplasia but not with the classic manifestations of Bamforth-Lazarus syndrome (athyreosis, cleft palate and spiky hair), and monoallelic FOXE1 AVs were indeed found in GIS cases; in both of this conditions the analysis of FOXE1 would have been excluded a priori. Furthermore, the association of biallelic AVs in functional genes such as SLC26A4 (CH14 in Supplementary Material, Table S2) and TG (CH7 in Supplementary Material, Table S2) with thyroid dysgenesis is particularly surprising. A previous study (26) proposed a ‘toxic’ effect produced by SLC26A4 inactivation on thyroid follicles as a possible explanation for this association. However, the co-existence of extra-thyroid defects, such as Arnold Chiari and severe short stature, in cases #14 and #7 suggests the possible combination with still uncovered defects in developmental genes. Moreover, we found SLC26A4 variants equally distributed among cases with thyroid dysgenesis or GIS, and frequently associated with mild congenital heart defects. This latter finding is particularly intriguing because SLC26A4 transcripts are expressed also in the heart tissue under the control of steroid hormones (27).

As recently reported (28,29), variations in candidate genes were indeed detected more frequently by NGS than previously observed. But these studies were conducted in small series of familial CH with gland-in-situ (<49 subjects from 34 ethnically



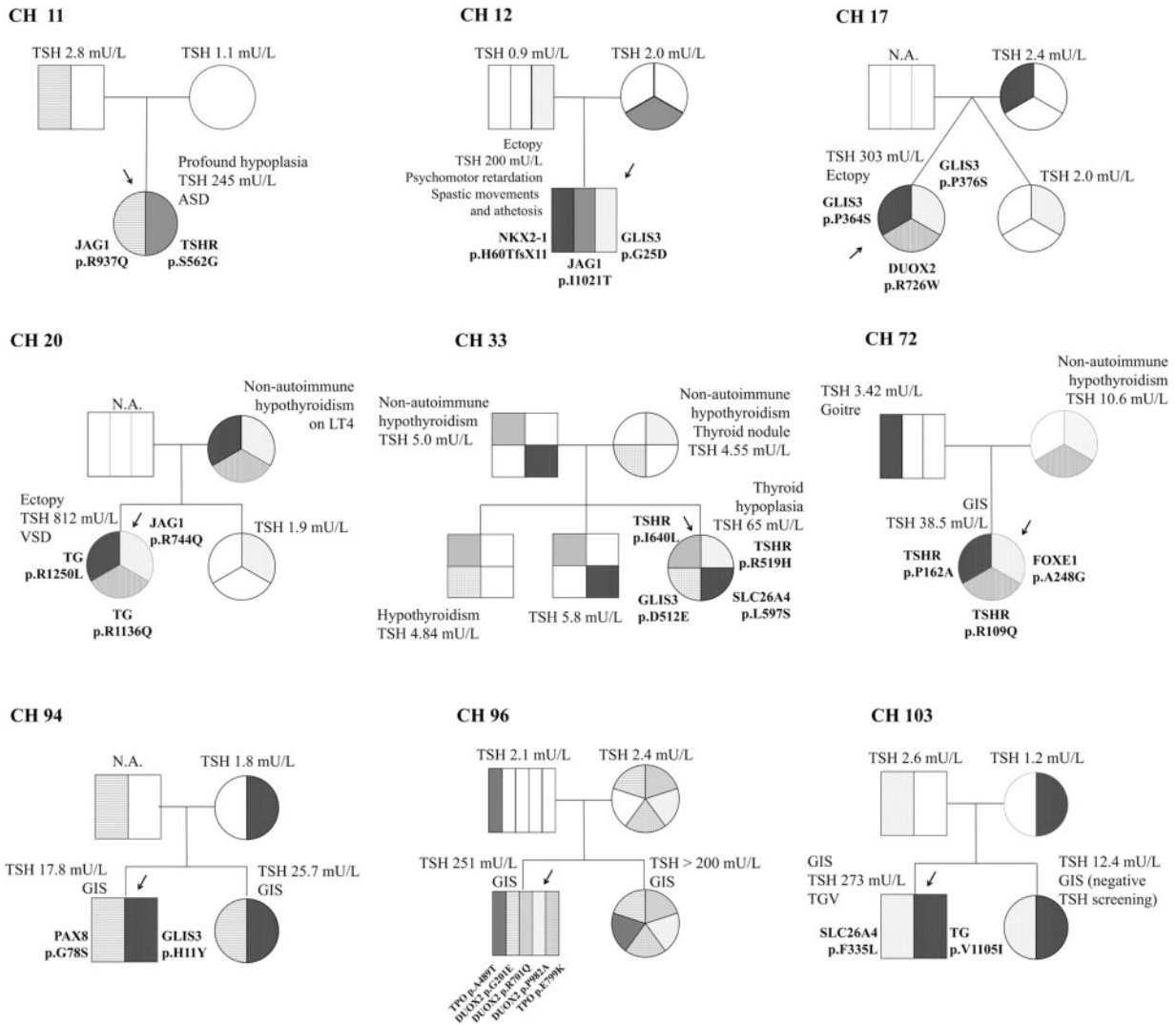
**Figure 3.** Distribution of accepted variants (AVs) in 11 CH-related genes (*NKX2-1*, *PAX8*, *FOXE1*, *GLIS3*, *JAG1*, *TSHR*, *SLC26A4*, *TG*, *TPO*, *DUOX2*, *DUOXA2*) and 18 CH-unrelated genes (*AIP*, *CDKN1B*, *FGFR1*, *FOXL2*, *FSHR*, *GDF9*, *GH1*, *GHRHR*, *GNRH1*, *GNRH2*, *GNRHR*, *MEN1*, *NRSA1*, *PROK2*, *PROKR2*, *PROP1*, *POU1F1*, *RET*) among the CH patients and the control population. The CH patients are highly significantly enriched only with variations in CH-related genes. The significance is even stronger when variants with a demonstrated functional impact or predicted to be deleterious in  $\geq 5/7$  in silico programmes (disruptive AVs) are considered.

diverse families) of variable geographic origin. At variance, our study has been performed in a larger cohort of unrelated CH patients, including either dysgenetic and functional thyroid defects thus representing the general CH population.

An added value of our study is the comparison of the frequency of gene variations in a large Italian control population. This analysis reveals that the CH population is indeed significantly enriched with rare/low frequency alleles in the 11 CH-related genes, and the frequency of multiple gene involvement is 2- to 4-fold higher than in the control population. Importantly, the distribution difference between cases and controls: a) is higher when only disruptive variants in the CH related genes are taken into account, and b) disappears when the variations in 18 CH-unrelated genes are considered. It is then conceivable that variants with a modest functional impairment can produce a negligible effect on thyroid function when expressed alone, but the sum of minor alleles, even acting at

different levels (thyroid morphogenesis or hormonogenesis), can justify the birth of a child with CH in families with a history of minor thyroid defects or without any previously recognized thyroid disease (Fig. 4). Our findings extend the relevance of the genetic components of CH, supporting the hypothesis that sporadic CH may be the result of the combination of multiple defects in different CH genes. This oligogenic model may indeed constitute another suitable explanation for the variable expressivity and penetrance of genetic defects previously reported in several CH familial settings (5,6,9,18,19,23), and well fits the previously proposed animal model of a multifactorial pathogenesis of CH (17). On this line, we recently reported the involvement of *JAG1* loss-of-function variants in CH pathogenesis (10). In this context, *JAG1* may act as a gene modifier contributing to uncover a mild functional defect due to the concomitant presence of rare alleles in 'specific thyroid genes'. This is outlined in the pedigrees of CH cases #11 and #20, and may explain the first





**Figure 4.** Distribution of accepted gene variants in nine families. The filled sections in the squares and circles outline the variants. All the CH probands (indicated by an arrow) were carriers of >1 variant (up to five variants in case #96 and his sister, both affected with CH) that occurred de novo or were inherited either from the mother or father. The probands #11, #12 and #20 had been previously reported (10,35). The combinations of AVs in morphogenetic and functional genes represent a suitable explanation for the thyroid dysgenesis seen in cases #11, 12 and #17. The variable combinations of multiple variants are associated with CH, whereas milder forms of non-autoimmune hypothyroidism were found in the relatives carrying only part of the variants present in the family (as in cases #20, #33, #72 or #103). Familiarity for CH was known in cases #94 and #96 and both affected siblings carry the same combinations of AVs. The young sister of proband #103 (2 years old), the two brothers (12 and 2 years old) and both parents of proband #33, as well as the mother of case #72 were found to be hypothyroid after positive genetic screening. The co-segregation of CH with multiple gene defects supports the oligogenic model of CH. ASD: atrial septum defect; VSD: ventricular septum defect; TGV: transposition of great vessels.

association of *NKX2-1* mutations with thyroid ectopy in case #12 (30). Therefore, we propose that beside rare monogenic forms, CH may more frequently arise in sporadic cases as a multifactorial disease with a relevant genetic predisposition. Indeed, environmental factors such as endocrine disruptors (or iodine deficiency) could contribute to endocrine dysfunction by influencing gene expression (31,32) and generating a more profound phenotype in carriers of rare genetic variants.

The NGS approach was predicted to become a useful tool for a more precise diagnosis and classification of affected patients (6). Its application to congenital hypothyroidism may overcome the current phenotypical classification based on cumbersome clinical and biochemical investigations (scintigraphy, perchlorate discharge test, ultrasound, biochemical thyroid function tests) and offer the possibility to increase the number of positive cases and predict the risk of CH or other

forms of hypothyroidism in affected families. In the present experience, we could identify predisposing variants in the majority of CH babies and the study of their segregation in families revealed several relatives with a previously unknown hypothyroidism (Fig. 4). The extension of the NGS panel to other candidates will indeed increase the performance of the NGS approach.

In conclusion, the first systematic NGS analysis performed in a large CH population reveals a frequent oligogenic origin of CH that may represent a suitable explanation for the missing heritability of CH and the prevalent sporadic presentation of the disease. Further studies are needed to prove if these rare alleles may contribute a significant portion of the functional and structural thyroid defects in the general population that are not justified by autoimmunity or iodide deficiency (33,34).

**Table 1.** Clinical and biochemical characteristics of the 177 CH patients. Most of them had severe forms of CH at diagnosis. Thyroid diseases in the relatives were frequent and included nodular and autoimmune diseases in most of them. Instead familiarity for CH was reported in six CH cases, consistent with the prevalent sporadic presentation of the disease

Characteristics	N (%)
Thyroid dysgenesis	83 (46.9)
-athyreosis	23 (13.0)
-ectopy	30 (16.95)
-hypoplasia	30 (16.95)
Gland-in-situ (GIS)	94 (53.1)
Severe CH (high TSH, low FT4)	151 (85.3)
Mild CH (high TSH, normal FT4)	26 (14.7)
Thyroid disease(s) in the family	88 (49.7)
Associated abnormalities:	55 (31.1)
-Nervous system	17 (9.6)
-Cardiovascular system	20 (11.3)
-Growth/skeletal	22 (12.4)
-Others	17 (9.6)

## Materials and Methods

### Subjects

The patients were enrolled in several Italian referral centers within a specific research protocol (RF-2010-2309484) that was approved by the Ethics Committees of the involved institutions. Informed consent was obtained from the parents of each CH patient. The analysis was applied to the first 177 eligible subjects enrolled after the approval of the protocol.

The inclusion criteria were: positive neonatal TSH screening (dry blood spot TSH  $\geq 10$  mU/l) with a diagnosis of primary CH confirmed by serum thyroid function tests at 1–3 weeks of age (serum TSH range: 9.2–951 mU/l; normal range: 0.4–7.5; serum FT4: 0.1–1.69 ng/dl; normal range: 1.5–2.4).

The exclusion criteria were: i) high urinary iodine; ii) anti-TSH receptor antibody in serum at neonatal age; iii) premature birth (<35 weeks of gestation).

All patients had a phenotypical classification by neck ultrasound and/or scintiscan. A thyroid dysgenesis was described in 83 CH patients, whereas a GIS of normal or enlarged size was described in 94 cases (Table 1). Additional information on the possible existence of thyroid disease in members of the family or associated malformations/diseases was collected in all cases. In case of positive genetic investigations, we extended whenever possible the biochemical and genetic studies to the available first-degree relatives. The data describing the clinical and biochemical characteristics of the 177 unrelated patients are reported in Table 1. The genetic investigations were replicated in an independent cohort of 145 unrelated CH patients (40.7% dysgenesis; 59.3% GIS).

The control population ( $n = 3,538$ ) was recruited within the ‘Atherosclerosis, Thrombosis, and Vascular Biology Italian Study Group’, as previously described (35).

### Genetic analyses

The genomic DNA of each patient was extracted from peripheral blood lymphocytes using Gene Catcher gDNA 96x10ml Automated Blood kit (Invitrogen, Life Technologies™). Genetic analyses were performed by NGS of a panel of 29 genes, including 11 CH candidate genes (NKX2-1, PAX8, FOXE1, GLIS3, JAG1,

TSHR, SLC26A4, TG, TPO, DUOX2, DUOXA2) (Supplementary Material, Table S1) and 18 CH-unrelated genes. The TruSeq Custom Amplicon assay (Illumina, San Diego, CA) was designed by the GenomeStudio software, having 681 amplicons covering exons with 25 bases of the flanking introns; the total coverage of the target genes by the designed amplicons was 80%. All regions not sequenced were recovered by Nextera® DNA Library Preparation kit (Illumina, San Diego, CA). The TruSeq NGS libraries were prepared according to the manufacturer’s instructions (Truseq® Custom Amplicon Library Preparation, Illumina, San Diego, CA). Pooled libraries were sequenced on MiSeq Reagent kit v.3 on an Illumina MiSeq sequencer (Illumina, San Diego, CA). Basic data analysis was performed according to the default parameters of the Illumina’s MiSeq Reporter software. For each subject, a .vcf file containing variant calls was generated, further reviewed, and filtered. For every equivocal call in the .vcf files, visual inspection of the mapped data was performed using the Integrated Genomics Viewer 2.3 software (IGV; Broad Institute, Cambridge, MA, USA). All variants of interest were verified by conventional dideoxy sequencing using BigDye® Terminator v.3.1 Cycle Sequencing Kit (Life Technologies, Carlsbad, CA, USA) on a 3100 DNA Analyzer from Applied Biosystems (Foster City, CA). Whole-exome sequencing (WES) of the control cohort was performed at the Broad Institute (Boston, MA). Sequencing, exome capture methods, variant annotation, and data processing of the samples were previously described (36). Since data in the case and control cohorts were generated by means of different sequencing platforms, the same quality filters for strand bias, coverage, and mapping quality were applied.

### Bioinformatics and statistical analyses

Only nonsense, frame-shift, splice-site (including the first six and the last three nucleotides of each intron), and missense variants with a minor allele frequency (MAF) <0.01 were considered in the analysis (accepted variant, AV), i.e. rare variants (15). Computational analyses were carried out to predict the possible pathogenic role of missense variants, through the use of the dbNSFP database (37). In detail, we considered as disruptive the nonsense, frame-shift, splice-site mutations and the non-synonymous missense variants annotated as deleterious by at least 5 out of 7 algorithms of the dbNSFP database, i.e.: SIFT, PolyPhen2, MutationTaster, MutationAssessor, LRT (Likelihood Ratio Test), and FATHMM (Functional Analysis Through Hidden Markov Model). And, for the intronic variants close to the splicing sites (-3/+6 nucleotides from the boundary) we used: NetGene2v.2.4 ESEfinder2.0, BDGP (see Supplementary Material, Table S2).

The frequency and the functional annotation of the identified variants were checked in public and licensed databases (Ensembl, UCSC Genome browser, 1000 Genome project, ExAC Browser, NCBI, HGMD professional).

Principal component analysis was performed on case and control populations using the SmartPCA program contained in the EIGENSOFT package (38,39).

Frequency and distribution of the variants in cases and controls were compared by means of Fisher’s exact test and R software (40).

## Supplementary Material

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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