Case Report: Expanding the Digenic Variants Involved in Thyroid Hormone Synthesis—10 New Cases of Congenital Hypothyroidism and a Literature Review

Rulai Yang†, Yijun Lu‡, Chenxi Yang‡, Xiaoyu Wu‡, Junqi Feng†, Ling Zhu†, Qiang Shu†, and Pingping Jiang†,‡,*

†The Children’s Hospitals, Zhejiang University School of Medicine, National Clinical Research Center for Child Health, Hangzhou, China, ‡Institute of Genetics and Department of Human Genetics, Zhejiang University School of Medicine, Hangzhou, China, *Zhejiang Provincial Key Laboratory of Genetic and Developmental Disorders, Hangzhou, China

Congenital hypothyroidism (CH) is the most common neonatal metabolic disorder. Although it has been understood to be a monogenic disease, some CH patients are reported to carry two or more variants at different genes. Here, ten permanent congenital hypothyroidism (PCH) patients were retrospectively reviewed, with elevated levels of serum thyroid-stimulating hormone and levothyroxine dependence during follow-up between 2015 and 2019. Each affected individual carried digenic variants, which were heterozygous at two of pathogenic genes. In total, five pathogenic genes, TSHR, TG, TPO, DUOX2 and DUOXA2, were simultaneously identified in subjects that were involved in the same metabolic pathway: thyroid hormone biosynthesis. There were digenic variants at TSHR and DUOX2 combined in three patients, DUOX2 and TG combined in two patients, DUOX2 and DUOXA2 combined in two patients, TG and DUOXA2 combined in two patients, and TG and TPO combined in one patient. Additionally, seven novel variants, TSHR c.679G>A, DUOX2 c.127A>T, c.608-619del, c.959T>C, TG c.2307G>A, and c.6759_6765del, and DUOXA2 c.93T>G, were identified in these PCH patients. Along with a literature review on digenic variants in patients with CH, our findings illustrated the complexity of genetic etiology in CH.

Keywords: digenic variants, thyroid hormone synthesis, congenital hypothyroidism, genetic counseling, oligogenic cases

BACKGROUND

Congenital hypothyroidism (CH) is the most common neonatal metabolic disorder. It has an incidence ranging from 1:1,400 to 1:2,800 live births in many countries (Wassner and Brown, 2015), and it results in severe neurodevelopmental impairment if not treated early and effectively. Primary CH is usually classified into two categories by pathogenesis: thyroid dysgenesis, a defect in thyroid gland development in which a few cases were caused by FOXE1, NKX2-1, NKX2-5, and PAX8, and thyroid dyshormonogenesis (DH), an intrinsic defect of thyroid hormone biosynthesis caused by DUOX2, DUOXA2, IYD (DEHAL1), TG, TPO, SLC26A4 (PDS), SLC26A7, SLC5A5 (NIS), and TSHR (Cangul et al., 2018; Kwak, 2018). Based on the newborn screening (NBS) program...
and clinical diagnosis, thyroid dyshormonogenes dominate compared to thyroid dysgenesis in the Chinese population upon increased molecular diagnosis (Long et al., 2018; Sun et al., 2018). Whereas thyroid dysgenesis still accounts for more than 69% of primary CH worldwide (Wassner and Brown, 2015; Peters et al., 2018). The inheritance of CH is controversial. Although it has been understood to be autosomal recessive (biallelic) in most cases as a monogenic disorder, a few CH cases appear to be monoallelic in one gene (Nicholas et al., 2016, Fugazzola et al., 2003), or 2 or more variants in different genes (Sripripradang et al., 2011; Satoh et al., 2015; Makretskaya et al., 2018; Yamaguchi et al., 2020). Here, we report 10 permanent congenital hypothyroidism (PCH) cases carrying digenic variants in which each affected individual is heterozygous at two of pathogenic genes simultaneously as well as the identification of seven novel genetic variants.

CASE PRESENTATION

During January 2015 and December 2019, the mean incidence of CH was 1:1,093 based on the NBS program in the Children's Hospital, Zhejiang University. CH screening strategies are designed to detect elevated levels of TSH and/or decreased concentrations of thyroxine (T4) (Group for Newborn Screening Society of Child Health Chinese Preventive Medicine Association, 2011). Total 2647 CH cases were diagnosed, of which 148 cases were offered genetic tests, and 66 cases (44.6%) had clear genetic confirmation, either carrying one P/LP variant in a dominate gene or two P/LP variants in a recessive gene. However, another 10 CH patients carrying digenic variants were retrospectively reviewed. They were clinically diagnosed to be PCH with a defect of thyroid hormone biosynthesis based on careful evaluation of clinical features and levothyroxine treatment during follow-up. As shown in Table 1, all patients had initially elevated TSH levels (>9 µIU/mL), ranging from 9.15 to 25.5 µIU/mL, and were proven to be permanent by receiving a trail off levothyroxine (LT4) at 2–3 years of age. Additionally, the influences of preterm, low-birthweight, and autoimmune thyroid disease on these cases were excluded. The detailed clinical information of the patients was listed in Table 1. With LT4 treatment with a dose of 12.5–33.3 µg per day, all patients had normal ASQ (Ages & Stages Questionnaires) and maintained serum TSH levels ranging from 1 to 10 (mIU/L) with a normal level of free thyroxine (FT4) between 9.01 and 19.05 (pmol/L) cutoff during follow-up. Cases #5 has a goiter by ultrasound during NBS with compensatory goiter recorded in Case #5 after 1 year with the LT4 supplement.

Identification of Digenic and Novel Variants

Identification of causative gene via whole-exome sequencing (WES) using peripheral blood was performed for 10 patients. The DNA library was prepared by an Agilent SureSelect Inherited Disease Capture Kit and sequenced using an Illumina HiSeq 2500 platform. All sequencing reads were mapped to the human reference genome (GRCh37) by BWA (Li and Durbin, 2010) and annotated by ANNOVAR (http://annovar.openbioinformatics.org). A series of automatic tools (SIFT, Polyphen, MutationTaster, etc.) were used to predict the functional significance of variants (Table 2). DNA samples from family #1, #3, #4, #5, #7, #8, and #10 were further verified by Sanger sequencing (Supplementary Figure 1).

The majority of dyshormonogenesis has an identifiable genetic basis since there are more than 10 genes reported to be involved in thyroid hormone biosynthesis (Kwak, 2018). All identified genes and variants were summarized in Figure 1. Five causative genes, TSHR, TG, TPO, DUOX2, and DUOXA2, were identified among the 10 patients. DUOX2 was detected in seven patients, followed by TG in five patients, DUOXA2 in four patients, TSHR in three patients, and TPO in one patient. There were digenic variants involving TSHR and DUOX2 in Case #1 (TSHR c.679G>A and DUOX2 c.127A>T), #2 (TSHR c.1574T>A and DUOX2 c.608-619del), and #3 (TSHR c.733G>A and DUOX2 c.3516_3531del), DUOX2 and TG in Case #4 (DUOX2 c.2654G>T and TG c.6759_6765del) and #5 (DUOX2 c.3516_3531del and TG c.2307G>A), DUOX2 and DUOXA2 in Case #6 (DUOX2 c.4027C>T and DUOXA2 c.738G>A) and #7 (DUOX2 c.959T>C and DUOXA2 c.738G>C), TG and DUOXA2 in Case #8 (TG c.3040G>A and DUOXA2 c.937G>T) and #9 (TG c.3808C>T and DUOXA2 c.413dupA), and TG and TPO in Case #10 (TG c.5791A>G and TPO c.2647C>T). In total, 18 variants were identified: 12 missense, 2 nonsense, and 4 frameshifts. These resulted from three deletions and one duplication. Five truncating proteins were observed in 7 cases, including DUOX2 p.K1174S fs*12 (c.3516_3531del) in Cases #3 and #5, TG p.S2254M fs*88 (c.6759_6765del) in Case #4, p.W769* (c.2307G>A) in Case #5, DUOXA2 p.Y246* (c.738C>G) in Cases #6 and #7, and p.Y138* (c.413dupA) in Case #9. Usually, truncating variants were pathogenic based on American College of Medical Genetics (ACMG) guidelines. Variants were mostly transmitted from both parents, although two heterozygous variants of Case #2 and #9 were solely from the mother (Figure 1). Moreover, a de novo variant resulting in a truncated protein, TG c.2307G>A (p. W769*), was detected in Case #5 (Supplementary Figure 1). Among these 18 variants, 7 were novel, identified as TSHR c.679G>A (p.G227R); DUOX2 c.127A>T (p.N43Y), c.608-619del (p.I203P207delinsP) and c.959T>C (p.L320P); TG c.2307G>A (p.W769*) and c.6759_6765del (p.S2254Mfs*88); and DUOXA2 c.937G>T (p.F31L). All the novel variants were localized in highly conserved regions of each protein (Figure 1) and predicted to be potential pathogenic variants by
functional consequences annotation through multiple software (Table 2).

**DISCUSSION**

Most cases of CH are common endocrine disorders caused by biallelic or monoallelic variants in one gene. With the widespread use of newborn screening programs and the application of genetic testing, some cases were found to carry two or more variants at different genes (Satoh et al., 2015; Nicholas et al., 2016; Sun et al., 2018; Yamaguchi et al., 2020), indicating the complexity of genetic etiology in CH. Cases with two or more variants in different genes were usually understood to be oligogenic cases, compared to those in biallelic and monoallelic cases (Yamaguchi et al., 2020). Here, we present 10 PCH cases carrying digenic variants in genes involved in thyroid hormone biosynthesis. Similar to our findings, another 58 cases harboring digenic variants were reported elsewhere (Supplementary Table 1). A total of 24 cases had digenic variants in TSHR and DUOX2, including 5 cases out of 220 Chinese CH (Fang et al., 2019) and 6 cases in Japanese patients (Abe et al., 2018; Yamaguchi et al., 2020). The coexistence of heterozygous variants in TSHR and DUOX2 was also revealed in Caucasian cases (Makretskaya et al., 2018; Sasivari et al., 2019). More digenic variants were heterozygous in two causative genes, including combined DUOX2 and TG in 13 patients (Löf et al., 2016; Fan et al., 2017; Long et al., 2018; Sun et al., 2018; Yamaguchi et al., 2020), DUOX2 and DUOX2A in 4 patients (Zheng et al., 2016; Yamaguchi et al., 2020), DUOX2 and TPO in 3 patients (Matsuo et al., 2016; Long et al., 2018; Makretskaya et al., 2018), TG and TPO in 6 patients (Nicholas et al., 2016; Makretskaya et al., 2018; Yamaguchi et al., 2020), and TG and SLC26A4 in 2 patients (Löf et al., 2016; Sun et al., 2018). Moreover, it was also demonstrated that 23% of Italian CH patients harbored pathogenic variants in more than one gene (Filippis et al., 2017), indicating that there was no ethnicity limiting the digenic form but rather a frequency of dyshormogenogenesis-associated variants. As shown in Supplementary Table 1, eight genes (TSHR, TG, DUOX2, DUOX1, DUOX2A, TPO, IYD, and SLC26A4) were present in those oligogenic cases. The higher frequency genes were DUOX2 (35.3%), TSHR (22.8%), and TG (22.8%). This was consistent with prior studies showing that DUOX2 and TSHR variants were more prevalent in Chinese, Japanese, and Korean patients (Jin et al., 2014; Fu et al., 2016; Fang et al., 2019; Yamaguchi et al., 2020). Higher frequent TG variants were detected in the Sudanese population (Bruehlemann et al., 2020). However, only two cases harbored variants of IYD that one individual combined with TG (Makretskaya et al., 2018) and the other one with DUOX1 (Sun et al., 2018). The digenic variants thereby seemed to be common in CH, but it is somewhat challenged in the variant interpretation by the dominant effect of some of these variants. For example, there were monoallelic variants reported in DUOX2 (Moreno et al., 2002), and later this turned out to be associated with transient hypothyroidism (Wang et al., 2014; Matsu et al., 2016).

To date, all reported genes with digenic variants are involved in the same metabolic pathway: thyroid hormone biosynthesis. As shown in Figure 2, the thyroid hormone is synthesized at the apical surface of polarized thyroid follicular cells, where the initial step is the binding of TSH to its receptors (TSHR) in the basolateral membrane, activating TG expression. To date, only 5 oligogenic cases carried heterozygous TSHR and TG variants in the Chinese and Japanese population (Fu et al., 2016; Yamaguchi et al., 2020). However, most cases of heterozygous TSHR (77.4%, 24/31) were combined with heterozygous DUOX2 as shown in Supplementary Table 1 (involved in steps 1 and 2). Subsequently, TG, TPO, and the DUOXs (DUOX2 and DUOX1) and their accessory protein DUOX2A are involved in iodide oxidation to form T4 and T3 (Kwak, 2018). Here, 7 out of 10 cases were caused by two of the five genes in this step. Additionally, overall 32 oligogenic cases (47%, 32/68) carried the combination of two heterozygous variants in two genes in this step, indicating that iodide organization defects may be more common in CH patients. Genes, IYD/DEHAL1, NIS/SLC5A5 and PDS/SLC26A4.

### Supplementary Table 1

| Cases# | Initial TSH (>9 μIU/ml) | TSH (mIU/L) | T3 (nmol/L) | T4 (nmol/L) | FT3 (nmol/L) | FT4 (pmol/L) | Levothyroxine (μg/day) |
|--------|------------------------|-------------|-------------|-------------|--------------|--------------|------------------------|
| 1. 6 years | 9.15 | 8.61 | 1.99 | 87.47 | 5.62 | 15.36 | 25 |
| 2. 4 years | 13.2 | 3.662 | 1.88 | 126.81 | 5.95 | 15.62 | 12.5 |
| 3. 5 years | 11.8 | 3.891 | 1.71 | 76.08 | 4.78 | 135.1 | 12.5 |
| 4. 6 years | 14.3 | 3.787 | 1.85 | 144.23 | 5.98 | 16.73 | 33.3 |
| 5. 4 years | 25.5 | 2.662 | 2.51 | 136.62 | 6.8 | 14.5 | 12.5 |
| 6. 2 years | 15.1 | 2.322 | 2.28 | 150.34 | 6.55 | 16.16 | 12.5 |
| 7. 3 years | 10.8 | 2.962 | 2.08 | 129.48 | 6.27 | 15.14 | 16.7 |
| 8. 3 years | 13.7 | 7.827 | 2.76 | 96.44 | 7.32 | 12.17 | 12.5 |
| 9. 6 years | 14.1 | 3.313 | 2.29 | 120.27 | 8.19 | 16.38 | 16.7 |

TSH, thyroid-stimulating hormone; T3, Triiodothyronine; T4, Thyroxine; FT3, free Triiodothyronine; FT4, free Thyroxine; †, ‡. years.
TABLE 2 | Genetic variants and their prediction on protein function.

| Case# | Gene | cDNA and amino acid change | ExonicFunc.refGene | Resource | ACMG interpretation | ACMG classification | Allele Frequency (ExAC ALL) | Prediction | SIFT | Polyphen2 | LRT | Mutation Taster | FATHMM |
|-------|------|-----------------------------|-------------------|----------|---------------------|--------------------|--------------------------|------------|------|----------|-----|----------------|--------|
| 1     | TSHR | c.679G>A (p.G227R)          | Nonsynonymous SNV | Maternal | PM1+PM2+PP3         | VUS                | –                        | D          | D    | D        | D   | D             | D      |
|       | DUOX2| c.127A>T (p.N43Y)           | Nonsynonymous SNV | Paternal | PM2+PP3             | VUS                | 0.09225%                 | D          | D    | D        | D   | D             | D      |
| 2     | TSHR | c.1574T>C (p.F525S)         | Nonsynonymous SNV | Maternal | PM1+PM2+PP3+PR3P    | 0.1%               | T                        | D          | D    | D        | D   | T             | T      |
|       | DUOX2| c.608_619del (p.L203_P207delinsP) | Nonframeshift deletion | Maternal | PM2+PP2+PM4        | P                  | –                        | .          | .    | .        | .   | .             | .      |
| 3     | TSHR | c.733G>A (p.G245S)          | Nonsynonymous SNV | Maternal | PM2+PP3+PP5         | VUS                | 0.1%                     | D          | D    | D        | D   | D             | D      |
|       | DUOX2| c.3516_3531del (p.K1174Sfs*12) | Nonframeshift deletion | Paternal | PM2+PP2+PP5        | P                  | 0.008266%                | .          | .    | .        | .   | .             | .      |
| 4     | DUOX2| c.2654G>T (p.R886L)         | Nonsynonymous SNV | Maternal | PM2+PP3+PP5         | VUS                | 0.3%                     | D          | D    | D        | D   | T             | T      |
|       | TG   | c.675_676del (p.S2254Mfs*68) | Frameshift deletion | Paternal | PM2+PP2+PP5        | P                  | –                        | .          | .    | .        | .   | .             | .      |
| 5     | DUOX2| c.3516_3531del (p.K1174Sfs*12) | Frameshift deletion | Paternal | PM2+PP2+PP5        | P                  | 0.008266%                | .          | .    | .        | .   | .             | .      |
|       | TG   | c.2307G>A (p.W769*)         | Stopgain de novo  | Paternal | PM2+PP2+PP5        | P                  | –                        | .          | .    | .        | D   | A             | .      |
| 6     | DUOX2| c.4027C>T (p.L1343F)        | Nonsynonymous SNV | Maternal | PM1+PM2+PP3+PR3P    | 0.5%               | T                        | P          | D    | D        | D   | T             | T      |
|       | DUOX2| c.738C>G (p.Y246*)          | Stopgain          | Paternal | PM1+PM2+PP5        | P                  | 0.2%                     | .          | .    | N        | D   | .             | .      |
| 7     | DUOX2| c.4959T>C (p.L320P)         | Nonsynonymous SNV | Maternal | PM2+PP3             | VUS                | 0.03304%                 | D          | P    | N        | D   | T             | T      |
|       | DUOX2| c.738C>G (p.Y246*)          | Stopgain          | Paternal | PM2+PP3             | VUS                | 0.2%                     | .          | .    | N        | D   | .             | .      |
| 8     | TG   | c.3040G>A (p.D1014N)        | Nonsynonymous SNV | Paternal | PM1+PM2+PP5        | VUS                | 0.02472%                 | T          | B    | N        | N   | T             | T      |
|       | DUOX2| c.93T>G (p.F31L)            | Nonsynonymous SNV | Maternal | PM2+PP3             | VUS                | 0.2%                     | D          | D    | D        | D   | T             | T      |
| 9     | TG   | c.3808C>T (p.R1270C)        | Nonsynonymous SNV | Maternal | PM2+PP3             | VUS                | 0.2%                     | D          | D    | D        | N   | N             | T      |
|       | DUOX2| c.413dupA (p.Y138*)         | Stopgain          | Paternal | PM2+PP3             | VUS                | 0.2%                     | .          | .    | D        | A   | .             | .      |
| 10    | TG   | c.5791A>G (p.I1931V)        | Nonsynonymous SNV | Maternal | PM2+PP5             | VUS                | 0.2%                     | T          | B    | N        | N   | T             | T      |
|       | TPO  | c.2847C>T (p.R883S)         | Nonsynonymous SNV | Paternal | PM2+PP5             | VUS                | 0.5%                     | T          | B    | N        | N   | T             | T      |

The reference gene version is GRCh37/hg19. DUOX2 (NM_014080.4), dual oxidase 2; DUOX2A (NM_207581.4), dual oxidase maturation factor 2; TSHR (NM_000369.2), TSH receptor; TG (NM_003235.4), thyroglobulin; TPO (NM_000547.5), thyroid peroxidase.

D, Damaging/Deleterious/disease causing; P, possibly damaging; N, Neutral; A, disease causing automatic; T, Tolerated.

Truncating protein as a result of stopgain or deletion is considered pathogenic according to ACMG guidelines. Novel variants are in bold. *, stopgain or truncated protein.
FIGURE 1 | Variants in CH patients and their conservative analysis. (A) Genotypes of CH pedigrees; (B) Conservative analysis of seven novel variants in species. The reference sequences for TSHR, DUOX2, TG, and DUOXA2 in species are followed as: TSHR: Homo sapiens, NP_000360.2; Pan troglodytes, XP_009426511.1; DUOX2: Homo sapiens, NP_003482.1; Pan troglodytes, XP_009426511.1; TG: Homo sapiens, NP_000360.2; Pan troglodytes, XP_009426511.1; DUOXA2: Homo sapiens, NP_003482.1; Pan troglodytes, XP_009426511.1; TPO: Homo sapiens, NP_000360.2; Pan troglodytes, XP_009426511.1; Rattus norvegicus, XP_009426511.1; Mus musculus, XP_009426511.1; Bos taurus, XP_009426511.1; Danio rerio, XP_009426511.1. (Continued)
involved in recycling of T4, T3, iodide and tyrosine (Spitzweg et al., 2000). Limited by the number of cases, there was no variant detected in NIS as elsewhere (Long et al., 2018). Only a few variants have been reported to date in IYD and PDS. As shown in Supplementary Table 1, there were only three cases carrying heterozygous PDS and two cases carrying heterozygous IYD. Theoretically, any defects of these eight proteins in substrates, enzymes, and transport molecules in the same metabolic pathway led to thyroid dyshormonogenesis. In fact, our data and recent evidence revealed that the combinations of two pathogenic genes predominantly happened in iodide organification and then in TG expression.

The limitation here is the lack of a parental phenotype, and all our cases are simplex cases. Especially in Case#2 and #9, the parental phenotype is critical in understanding the functional effects of digenic variants. Unfortunately, the mothers carrying the same two variants refused to test their TSH levels. Moreover, the digenic variants in Case#1, #8, and #10 were classified to be VUS according to the ACMG guidelines, which need more cases or further functional experiments to evaluate their damage prediction.

Summarily, we reported here 10 PCH cases with digenic variants involved in the same metabolic pathway: thyroid hormone biosynthesis. To date, 68 CH patients have been reported harboring digenic variants in this metabolic pathway, including genes TSHR, TG, DUOX2, DUOX1, DUOXA2, TPO, IYD, and SLC26A4 with a high frequency of DUOX2, TSHR, and TG. The data present here will extend our awareness of the complexity of genetic etiology in CH.
DATA AVAILABILITY STATEMENT

The original contributions presented in the study are publicly available in NCBI using accession number PRJNA734721.

ETHICS STATEMENT

The studies involving human participants were reviewed and approved by the Research Ethics Committees, Children's Hospital of Zhejiang University School of Medicine. Written informed consent to participate in this study was provided by the participants' legal guardian/next of kin. Written informed consent was obtained from the individual(s), and minor(s)' legal guardian/next of kin, for the publication of any potentially identifiable images or data included in this article.

AUTHOR CONTRIBUTIONS

PJ and QS performed the conception, analysis, and interpretation of data. RY and LZ recruited patients and performed clinical evaluation. YL, CY, XW, and JF conducted the mutational sequencing and data analysis. PJ, YL, RY, and QS drafted and revised the article. All authors contributed to the article and approved the submitted version.

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Data Availiability Statement

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fgene.2021.694683/full#supplementary-material
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