## REVIEW



# Molecular genetic diagnostics of hypogonadotropic hypogonadism: from panel design towards result interpretation in clinical practice

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# Abstract

Congenital hypogonadotropic hypogonadism (CHH) is a clinically and genetically heterogeneous congenital disease. Symptoms cover a wide spectrum from mild forms to complex phenotypes due to gonadotropin-releasing hormone (GnRH) deficiency. To date, more than 40 genes have been identified as pathogenic cause of CHH. These genes could be grouped into two major categories: genes controlling development and GnRH neuron migration and genes being responsible for neuroendocrine regulation and GnRH neuron function. High-throughput, next-generation sequencing (NGS) allows to analyze numerous gene sequences at the same time. Nowadays, whole exome or whole genome datasets could be investigated in clinical genetic diagnostics due to their favorable cost–benefit. The increasing genetic data generated by NGS reveal novel candidate genes and gene variants with unknown significance (VUSs). To provide clinically valuable genetic results, complex clinical and bioinformatics work are needed. The multifaceted genetics of CHH, the variable mode of inheritance, the incomplete penetrance, variable expressivity and oligogenic characteristics further complicate the interpretation of the genetic variants detected. The objective of this work, apart from reviewing the currently known genes associated with CHH, was to summarize the advantages and disadvantages of the NGS-based platforms and through the authors' own practice to guide through the whole workflow starting from gene panel design, performance analysis and result interpretation. Based on our results, a genetic diagnosis was clearly identified in 21% of cases tested (8/38).

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# Congenital hypogonadotropic hypogonadism (CHH)

# Congenital hypogonadotropic hypogonadism (CHH) as a clinically heterogeneous entity

Congenital hypogonadotropic hypogonadism (CHH) is a genetic condition characterized by incomplete or absent puberty and infertility due to central (tertiary or hypothalamic) hypogonadism caused by gonadotropic hormonereleasing hormone (GnRH) deficiency. Three clinical forms are distinguished by the European consensus: (1) GnRH deficiency with defective sense of smell (Kallmann syndrome, KS), (2) isolated GnRH deficiency (normosmic CHH) and the third form when KS/CHH is part of a complex genetic syndrome (Boehm et al. 2015). CHH has an incidence of 1:125,000 in female and 1:30,000 in males indicating the male predominance (Stamou and Georgopoulos 2018). CHH has a heterogeneous clinical appearance, and lately the constitutional delay of growth and puberty (CDGP), the adultonset hypogonadotropic hypogonadism and the hypothalamic amenorrhea are also considered as a milder end of the spectrum (Stamou and Georgopoulos 2018). Most patients are diagnosed in adolescence due to delayed puberty. In male, neonate cryptorchidism and micropenis can be considered as signs of CHH, but there are no specific signs of CHH in female neonates (Young et al. 2019). Prepubertal testes and undervirilized secondary sexual features are the most common symptoms in males, while absence of breast development and primary amenorrhea occur in females as a consequence of CHH (Young et al. 2019). The disease can be diagnosed in adulthood as well by low libido, infertility, bone loss and fractures when it is untreated (Young et al. 2019). Interestingly, in 10-20% of the cases, CHH is reported reversible, however, the pathophysiology behind this is not clearly revealed (Stamou and Georgopoulos 2018; Young et al. 2019). To establish the biochemical diagnosis in infants is challenging as GnRH neurons are active only during mini-puberty (4-8 weeks after birth) and after that, their activity becomes quiescent until puberty. In adolescence, results of biochemical tests (basal and stimulated blood levels of sex hormones and gonadotropins), brain imaging for examination of olfactory bulbs, assessment of smell and evaluation of family history are parts of the routine medical investigations. (Naturally, additional work-ups, i.e. evaluation of bones, kidneys and sexual organs, are also required for diagnosis and for differential diagnostic purposes (Young et al. 2019). Constitutional delayed of growth and puberty (CDGP) is defined as the lack of the start of sexual maturation at an age > 2 SDs above the mean for a given population (Stamou and Georgopoulos 2018). There is no identifiable cause behind and finally puberty occurs. 50-80% of CDGP individuals have positive family history of the phenomenon, and approximately 10% of CHH patients have relatives with CDGP (Stamou and Georgopoulos 2018). Differentiating CDGP and CHH in adolescence is challenging as to date no gold-standard diagnostic test is known for this purpose (Young et al. 2019).

There are non-reproductive features as well that are commonly recognized in patients with CHH. Midline facial defects (cleft lip or palate), dental agenesis, unilateral renal agenesis, short metacarpals, hearing loss, synkinesia, cerebellar ataxia can appear additionally to CHH (Young et al. 2019). Furthermore, the disease can occur as part of complex genetic syndromes summarized in Table 1.

Diagnostics and genetic counseling is important in CHH as effective therapies are available for the development of secondary sexual features and fertility (Maione et al. 2018; Young et al. 2019).

Adult onset of hypogonadotropic hypogonadism is a rare form of CHH. It is a non-reversible, long-lasting condition but the etiology and pathogenesis have to be investigated and demonstrated. The diagnosis can be made when all other acquired causes of hypogonadotropic hypogonadism (e.g. structural anomalies, infiltrative/inflammatory origin, pituitary/CNS tumors etc.) have been excluded (Stamou and Georgopoulos 2018).

# **Genetic background of CHH**

CHH is heterogeneous not only clinically but also genetically. To date, more than 40 genes have been identified as pathogenic cause in the background of the disease (Boehm et al. 2015; Maione et al. 2018; Stamou and Georgopoulos 2018). Analysis the individual CHH genes (Table 1) one by one exceeds the goal of our study, but these are excellently reviewed in recent papers (Topaloglu and Kotan 2016; Topaloğlu 2018; Maione et al. 2018). Genes implicated in the pathogenesis of CHH are usually divided into two major categories (Boehm et al. 2015; Topaloğlu 2018; Maione et al. 2018; Stamou and Georgopoulos 2018). The first group consists of genes that control development and GnRH neuron migration. Therefore, the pathogenic variants of these genes are frequently associated with anosmia and midline developmental anomalies (Table 1). The second group of genes is responsible for neuroendocrine physiology and GnRH neuron function (either by afferent modulators or by regulating GnRH secretion), these can be detected in normosmic CHH forms. Although there are genes with multiple roles that participate in both mechanisms, their mutations can be often identified in both anosmic and normosmic forms (Boehm et al. 2015; Maione et al. 2018; Stamou and Georgopoulos 2018) (Table 1).

Autosomal dominant, autosomal recessive and X-linked inheritance have been identified, however, with the availability of high-throughput next-generation sequencing at least 20% of CHH cases have thought to be di- or oligogenic. In these cases, two or more gene variants can be identified in the same patient (Boehm et al. 2015) (Table 1). Still, in more than half of the CHH cases, there is no pathogenic mutation identified. Among the main genetic forms of CHH, the most common autosomal recessively inherited types are caused by GNRHR, KISS1R and TACR3 variants (Maione et al. 2018). Kallmann syndrome caused by ANOS1 gene mutations is inherited by X-linked recessive trait as it is located on chromosome X. FGFR1 and PROK2/PROKR2 lead to autosomal dominantly inherited type of CHH (Boehm et al. 2015; Maione et al. 2018). Regarding FGFR1, nearly half, regarding PROK2/PROKR2, nearly twothird of the cases exhibit incomplete penetrance and variable expressivity that complicate the determination of inheritance (Maione et al. 2018). Recently, a normosmic CHH patient was reported who inherited a pathogenic variant in GNRHR gene in a homozygous form due to the occurrence of uniparental isodisomy (Cioppi et al. 2019). (Uniparental disomy-UPD is a non-Mendelian inheritance pattern when an individual has inherited two copies of a specific chromosome (or part of it) from a single parent. When a chromosomal pair inherited from the same parent, it is called uniparental heterodisomy, when

Gene name	Inheritance	Gene function contributing in symp- toms	Clinical phenotype based on expert consensus (Boehm et al. 2015) and OMIM database	OMIM number ID or Ref
ANOSI (KALI) AXL	XR, contributes to oligogenicity AD	Migration of GnRH neurons Migration of GnRH neurons	Kallmann syndrome, CHH reversal Kallmann syndrome; CHH	308700 Salian-Mehta et al. (2014)
BBS1, BBS2, ARL6, BBS4, BBS5, MKKS, BBS7, TTC8, BBS9, BBS10, TRIM32, BBS12	AD or AR	Maintenance and function of cilia (cell movement, perception of sensory input such as sight, hearing, and smell)	Bardet-Biedl syndrome 1–12	209900
CHD7	AD	Migration of GnRH neurons	Kallmann syndrome; CHH with or without reversal; CHARGE sy	612370
CPE	AR	As part of a syndrome	CHH, obesity, diabetes mellitus type 2	Alsters et al. (2015)
DCAF17	AR	As part of a syndrome	Woodhouse-Sakati syndrome	241080
DMXL2	AD or AR	As part of a syndrome	CHH; PEPNS (polyendocrine defi- ciencies and polyneuropathies)	616113
DUSP6	AD	Migration of GnRH neurons	CHH with or without anosmia	615269
FEZF1	AR	Migration of GnRH neurons	Kallmann syndrome	616030
FGF17	AD, contributes to oligogenicity	Hypothalamus/pituitary development	Kallmann syndrome; CHH; Dandy- Walker sy	615270
FGF8	AD, contributes to oligogenicity	Hypothalamus/pituitary development,migration of GnRH neurons, neuroendocrine regulation	Kallmann syndrome; CHH; Com- bined pituitary hormone deficiency	612702
FGFR1	AD, contributes to oligogenicity	Hy pothalamus/pituitary development, neuroendocrine regulation	Kallmann syndrome; CHH with or without reversal; Combined pituitary hormone deficiency; Septo- optic dysplasia; Hartsfield sy; Split hand/foot malformation	147950
FLRT3	AD	Migration of GnRH neurons	CHH with anosmia	615271
FSHB	AR	Hypothalamus/pituitary development	CHH without anosmia	229070
GLCE	AD, AR	Hypothalamus/pituitary development	KS, CHH	Stamou and Georgopoulos (2018)
GNRH1	AR	neuroendocrine regulation	СНН	614841
GNRHR	AR, contributes to oligogenicity	Neuroendocrine regulation	CHH with or without reversal	146110
HDAC8	XR	Transcriptional regulation, cell cycle progression and development	Cornelia de Lange syndrome 5	300882
HESX1	AD or AR	Hy pothalamus/pituitary development	Kallmann syndrome; Combined pituitary hormone deficiency, Septo- optic dysplasia	182230
HFE	AD or AR	Controlling iron absorption by regulat- ing the interaction of the transferrin receptor with transferrin	Hereditary hemochromatosis	235200

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Table 1 (continued)				
Gene name	Inheritance	Gene function contributing in symp- toms	Clinical phenotype based on expert consensus (Boehm et al. 2015) and OMIM database	OMIM number ID or Ref
HS6ST1	AD, contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome; CHH with or without reversal	614880
IL17RD (SEF)	AD, AR or digenic dominant, contributes to oligogenicity	Migration of GnRH neurons	CHH with or without anosmia	615267
IRF2BPL (EAP1)	Not known	Neuroendocrine regulation	СНН	Mancini et al. (2019)
KISS1	AR	neuroendocrine regulation	CHH	614842
KISS1R	AR, contributes to oligogenicity	Neuroendocrine regulation	СНН	614837
LEP	AR	Neuroendocrine regulation	CHH, obesity	Strobel et al. (1998)
LEPR	AR	Neuroendocrine regulation	CHH, obesity	Hannema et al. (2016)
LHB	AR	Hypothalamus/pituitary development	CHH with or without anosmia	228300
LHX3	AR	Hypothalamus/pituitary development	Combined pituitary hormone defi- ciency	221750
NDN	AD	Neuroendocrine regulation	Prader-Willi syndrome	176270
NR0B1 (DAX1)	XR	Hypothalamus/pituitary development	CHH, congenital adrenal hypoplasia	300200
NSMF (NELF)	AD, contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome; CHH with or without reversal	614838
OTUD4	AR	As a deubiquitinase it negatively regulates inflammatory and patho- gen recognition signaling in innate immune response	CHH; Gordon-Holmes sy	212840
PCSK1	AR	Hypothalamus/pituitary development	CHH, obesity	600955
PNPLA6	AR	Regulation of neurite outgrowth and process elongation during neuronal differentiation	CHH; Boucher-Neuhauser syndrome	215470
POLR3A	AR	As a DNA-dependent RNA polymer- ase it catalyzes the transcription of DNA into RNA	Leukodystrophy, hypomyelinating, seven, with or without oligodontia and/or hypogonadotropic hypog- onadism	607694
POLR3B	AR	As a DNA-dependent RNA polymer- ase it catalyzes the transcription of DNA into RNA	Leukodystrophy, hypomyelinating, eight, with or without oligodontia and/or hypogonadotropic hypog- onadism	614381
PROK2	AD, contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome; CHH	610628
PROKR2	AD, contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome; CHH with or without reversal; Combined pitui- tary hormone deficiency; Morning glory syndrome	244200

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Table 1 (continued)				
Gene name	Inheritance	Gene function contributing in symp- toms	Clinical phenotype based on expert consensus (Boehm et al. 2015) and OMIM database	OMIM number ID or Ref
PROP1	AR	Hypothalamus/pituitary development	Combined pituitary hormone defi- ciency	262600
RAB18	AR	Eye and brain development, regulating membrane trafficking in organelles and transport vesicles	Warburg micro syndrome 3	614222
RAB3GAP1	AR	Eye and brain development, regulated exocytosis of neurotransmitters and hormones	Warburg micro syndrome 1	600118
RAB3GAP2	AR	Eye and brain development, regulated exocytosis of neurotransmitters and hormones	Martsolf syndrome (cataract, mental- retardation, hypogonadism)	212720
RBM28	AR	Splicing regulator	Alopecia, neurologic defects, and endocrinopathy syndrome	612079
RNF216	AR	Ubiquitination, protein degradation by the proteasome, regulation of TNF-, IL1- and NFKB signaling	CHH; Gordon-Holmes sy	212840
SEMA3A	AD, contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome	614897
SEMA3E	AD	Migration of GnRH neurons	KS, CHH	Cariboni et al. (2015)
SEMA7A	contributes to oligogenicity	Migration of GnRH neurons	Kallmann syndrome; CHH	Känsäkoski et al. (2014)
SOX10	AD	Hypothalamus/pituitary development	Kallmann syndrome; Waardenburg syndrome; PCWH syndrome (peripheral demyelinating neuropa- thy, central demyelination, Waarden- burg syndrome, and Hirschsprung disease)	611584
SOX2	AD	Hypothalamus/pituitary development	CHH, optic nerve hypoplasia and abnormalities of the central nervous system	206900
SOX3	XR	Hypothalamus/pituitary development	CHH	Izumi et al. (2014)
SPRY4	AD	Migration of GnRH neurons	Kallmann syndrome	615266
SRA1	AD	Neuroendocrine regulation	CHH	Kotan et al. (2016)
STUB1	AR	Ubiquitination, protein degradation by the proteasome	Spinocerebellar Ataxia 16, Autosomal Recessive	615768
TAC3	AR, contributes to oligogenicity	Neuroendocrine regulation	CHH with or without reversal	614839
TACR3	AR, contributes to oligogenicity	Neuroendocrine regulation	CHH with or without reversal	614840
TBCID20	AR	Regulation of vesicle-mediated transport, GTPase-activating protein specific for Rab1 and Rab2	Warburg micro syndrome 4	615663

Gene nameInheritanceGene function contributing in symp- consensus (Boehm et al. 2015) and OMIM databaseOMIM number ID or R consensus (Boehm et al. 2015) and OMIM databaseOMIM databaseTUBB3ADMigration of GnRH neurons"TUBB3 E410K syndrome"Chew et al. (2013); Pat without reversal; Combined pitui- tary hormone deficiency	Table 1 (continued)				
TUBB3     AD     Migration of GnRH neurons     "TUBB3 E410K syndrome"     Chew et al. (2013); Pat       WDR11     AD, contributes to oligogenicity     Migration of GnRH neurons     Kallmann syndrome; CHH with or     614858       WDR11     AD, contributes to oligogenicity     Migration of GnRH neurons     Kallmann syndrome; CHH with or     614858       WDR11     AD, contributes to oligogenicity     Migration of GnRH neurons     Kallmann syndrome; CHH with or     614858	Gene name	Inheritance	Gene function contributing in symp- toms	Clinical phenotype based on expert consensus (Boehm et al. 2015) and OMIM database	OMIM number ID or Ref
	TUBB3 WDR11	AD AD, contributes to oligogenicity	Migration of GnRH neurons Migration of GnRH neurons	"TUBB3 E410K syndrome" Kallmann syndrome; CHH with or without reversal; Combined pitui- tary hormone deficiency	Chew et al. (2013); Patel et al. (2017) 614858

two identical chromosomes are inherited it is called uniparental isodisomy (iUPD). This discovery further complicates the inheritance pattern of CHH and raises the possibility of the same phenomenon in case of other genes as well.

Individual relevance of genes in oligogenic CHH cases are needed to be interpreted with cautions. For instance, another candidate CHH gene *NSMF* (earlier *NELF*), listed in the expert consensus statement, has now a controversial role (Spilker et al. 2016). Several publications reported *NSMF* variants in CHH patients alone or in combination with a mutation in another gene (Miura et al. 2004; Pitteloud et al. 2007; Xu et al. 2011) underlining again its questionable role (Spilker et al. 2016).

Interestingly, rare variants of *TAC3*, *TACR3* and other genes are suggested to be linked with CHH reversal that further raises the possibility of therapy discontinuation from time to time to test the reversibility of CHH in these carriers (Gianetti et al. 2010; Boehm et al. 2015).

Variants of known CHH genes have been investigated and identified in CDGP and in cases with hypothalamic amenorrhea too. This suggests that the time of menarche and menopause are genetically determined which is strongly supported by family histories (Stamou and Georgopoulos 2018).

In CDGP, Zhu et al. identified that variants in CHH genes were enriched in CDGP family members compared to unaffected family members suggesting the genetic link between CHH and CDGP (Zhu et al. 2015). This is further supported by variants identified in CDGP patients in *TAC3*, *TACR3*, *IL17RD*, *GNRHR*, *PROKR2*, *HS6ST1*, *FGFR1*, *FEZF1*, *AXL* genes (Gianetti et al. 2012; Tusset et al. 2012; Zhu et al. 2015; Hietamäki et al. 2017; Cassatella et al. 2018). However, results of Cassatella et al. demonstrated that CDGP and CHH have distinct genetic profiles that may facilitate the differential diagnosis in patients presenting with delayed puberty (Cassatella et al. 2018).

Hypothalamic amenorrhea is also a reversible dysfunctional feature that can be triggered by nutritional deficit, extensive exercise or psychological stress. Genetic variants have been identified in *FGFR1*, *PROKR2*, *GNRHR* and *ANOS1* genes suggesting that these mutations may contribute to the variable functional changes in GnRH secretion (Caronia et al. 2011).

CHH can be also part of complex genetic syndromes which are summarized by Boehm et al. and genetic background are summarized in Table 1 (Boehm et al. 2015).

# Genetic testing and genetic counseling in CHH

# **Testing strategies**

Although high-throughput screening can be recommended, targeted panel testing, prioritization and gene selection based on clinical data are also possible (Boehm et al. 2015; Topaloğlu 2018; Stamou and Georgopoulos 2018). The first step is to exclude the presence of genetic syndromes based on clinical findings. When a clinical geneticist based on the whole clinical presentation indicates a specific syndrome (e.g. CHARGE sy., Bardet-Biedl sy., Gordon-Holmes sy., see details in Table 1) targeted gene testing is recommended. When complex syndromes can be excluded additional associated signs and symptoms can increase the probability of finding casual mutations (Boehm et al. 2015). For instance, besides anosmia/hyposmia, bimanual synkinesia or renal agenesis can associate with ANOS1 mutation (Fig. 1). Cleft palate/lip, dental agenesis and digital bone anomalies were frequently associated with CHH caused by mutations in genes of FGF8 signaling (FGFR1, FGF8, HS6ST1) (Costa-Barbosa et al. 2013; Boehm et al. 2015). Hearing impairment commonly appeared with CHH in CHD7, SOX10 or IL17RD mutation carriers (Costa-Barbosa et al. 2013; Boehm et al. 2015). Additionally, early onset of morbid obesity with CHH could suggest variants in LEP, LEPR or PCSK1 genes (Jackson et al. 1997; Farooqi and O'Rahilly 2008). If CHH is associated with severe adrenal insufficiency congenital adrenal hypoplasia caused by NR0B1 (DAX1) is likely.

Combined pituitary hormone deficiency (CHPD) should also be clinically investigated/excluded as CHH and CHPD have overlapping genetic etiologies. If isolated CHPD is diagnosed, genetic testing of genes encoding the pituitary transcription factors (*PROP1*, *POU1F1*, *LHX4*, *LHX3* and *HESX1*) should be recommended (Fang et al. 2016). However, lately, variants of certain CHH genes including *CHD7*, *PROKR2*, *WDR11*, *FGFR1* and *FGF8* have also been implicated in CPHD (Raivio et al. 2012; Fang et al. 2016). Similar to CHH, CPHD is suggested to be a multifactorial disease as symptoms frequently present incomplete penetrance even harboring the very same mutations (Raivio et al. 2012).

Genetic testing starts with evaluation of the inheritance pattern using pedigree analysis. However, Mendelian inheritance have been described for the majority of genes associated with CHH, some genes show different inheritance patterns (e.g. *FGFR1*: AD/AR/oligogenic/de novo; *PROK2/PROKR2*: AD/AR/oligogenic), see Table 1 (Boehm et al. 2015; Maione et al. 2018).

Parallel with revolution of molecular genetic technologies for patients with CHH multi-gene panel testing can be recommended, because there is a wide overlap between both symptoms and genetic background (Boehm et al. 2015; Maione et al. 2018). Since expert consensus have been published in 2015 (Boehm et al. 2015) several high-throughput multi-gene panel studies were carried out. However, there is no consented gene list that should be offered for patients providing an accurate diagnosis for the majority of cases. After publication of the expert consensus 7 CHH gene panel testing studies were reported (Table 2) (Quaynor et al. 2016; Aoyama et al. 2017; Wang et al. 2017; Cassatella et al. 2018; Zhou et al. 2018; Amato et al. 2019; Kim et al. 2019). In these studies, 25-261 genes were included as susceptibility genes of CHH. The positive detection rate varied between 33 and 56% (Table 2). Mutations in the FGFR1 gene were found the most commonly (in all eight studies), ANOS1 (in seven studies) and CHD7, PROKR2, TACR3 and IL17RD variants were also frequently detected (in six studies) among different groups (Table 2). Analyzing the detection rate by patient number FGFR1 variants were detected most commonly, in an average of 11.4% of all investigated patients, CHD7, PROKR2 and ANOS1 in 8.4, 6.4 and 5.7% of patients, respectively, across all studies. All other gene



Fig. 1 Genetic testing strategies in CHH (based on Boehm et al. 2015; Stamou and Georgopoulos 2018; Topaloğlu 2018)

Table 2	High-throughput N	GS studies	investigating	CHH patients
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	Quaynor et al. (2016)	Wang et al. (2017)	Aoyama et al. (2017)	Cassatella e	et al. (2018)	Zhou et al. (2018)	Kim et al., (2019)	Amato et al. (2019	Current study
Nr of CHH patients	48	51	22	116 CHH	72 CDGP	153	28	130	38
Nr of genes investi- gated	261	164	27	25		83	69	36	41
Nr of patients with identified variant	NA	26 (51%)	12 (54.5%)	59 (51%)	5 (7%)	87 (56%)	11 (39%)	43 (33%)	22 (57%)
Ratio of cases with di-/ oligogenic back- ground	19%	9.8%	0%	15%	1.4%	19%	not reported	6.9%	21%
Genes with variants identified (frequency of detec- tion in the partciular cohort)	AXL (6.3%)	PROKR2 (17.6%)	CHD7 (18.2%)	FGFR1 (15.5%)	AXL (1.4%)	RELN (20.3%)	FGFR1 (14.3%)	CHD7 (10.8%)	FGFR1 (12.5%)
	FGFR1 (6.3%)	FGFR1 (13.7%)	ANOS1 (18.2%)	CHD7 (13.8%)	FGFR1 (1.4%)	PROKR2 (17.6%)	CHD7 (7.1%)	FGFR1 (8.5%)	GLI3 (7.5%)
	GLI3 (4.2%)	CHD7 (7.8%)	FGFR1 (13.6%)	PROKR2 (5.2%)	HS6ST1 (1.4%)	CHD7 (9.8%)	TACR3 (7.1%)	IGSF10 (5.4%)	NOTCH1 (7.5%)
	AMN1 (2.1%)	IL17RD (5.9%)	TACR3 (4.5%)	SOX10 (4.3%)	PROKR2 (1.4%)	ANOS1 (7.2%)	PROKR2 (3.6%)	GNRHR (5.4%)	MASTL (7.5%)
	CCKBR (2.1%)	ANOS1 (5.9%)		AXL (3.4%)	TAC3 (1.4%)	ERBB4 (6.5%)	ANOS1 (3.6%)	WDR11 (4.6%)	PROKR2 (5%)
	CRY1 (2.1%)	FGF17 (2%)		GNRHR (3.4%)	FEZF1 (1.4%)	FGFR1 (6.5%)	SOX3 (3.6%)	ANOS1 (4.6%)	AMH (5%)
	CXCR4 (2.1%)	KISS1R (2%)		SEMA3A (2.6%)		EGFR (5.9%)		TACR3 (3.8%)	JAG1 (5%)
	FGF13 (2.1%)	PROK2 (2%)		IL17RD (2.6%)		LHB (5.9%)		PROK2 (3.8%)	IL17RD (5%)
	GAP43 (2.1%)	SEMA3A (2%)		TACR3 (2.6%)		PLXNB1 (0.59%)		DMXL2 (3.1%)	PDE3A (5%)
	GNRH1 (2.1%)	SPRY4 (2%)		ANOS1 (1.7%)		SEMA4D (5.9%)		PROKR2 (2.3%)	ANOS1 (5%)
	GNRHR (2.1%)			FGF8 (1.7%)		EGF (4.6%)		POLR3B (2.3%)	GNRHR (5%)
	IL17RD (2.1%)			HS6ST1 (1.7%)		NRP2 (4.6%)		IL17RD (2.3%)	TAC3 (2.5%)
	JAG1 (2.1%)			WDR11 (1.7%)		B3GNT1 (3.9%)		SPRY4 (1.5%)	TACR3 (2.5%)
	(2.1%) MASTL (2.1%)			GNRH1 (1.7%)		(3.9%) IL17RD		SOX10 (1.5%)	AMHR2
	(2.1%) NOS1			(1.7%) KISS1		NOS1 (3.9%)		SEMA7A	(2.5%) KISS1R
	(2.1%) NOTCH			FGF17		ROBO3 (3.9%)		(1.5%) SEMA3A	(2.5%) SPRY4 (2.5%)
	NRP2 (2.1%)			PROK2 (0.9%)		DCC (3.3%)		POLR3A (1.5%)	(2.570)

Table 2 (continued)

Quaynor et al. (2016)	Wang et al. (2017)	Aoyama et al. (2017)	Cassatella et al. (2018)	Zhou et al. (2018)	Kim et al., (2019)	Amato et al. Current stu (2019	ıdy
PALM2 (2.1%)			KISS1R (0.9%)	MTOR (3.3%)		NSMF (1.5%)	
PDE3A			TAC3	SEMA7A		IGFALS	
(2.1%)			(0.9%)	(3.3%)		(1.5%)	
PLEHKA5				DLX5 (2.6%)		GNRH1	
(2.1%)				CNDUD		(1.5%)	
RD3 (2.1%)				(2.6%)		(1.5%)	
TRAPPC9 (2.1%)				IGF1 (2.6%)		TAC3 (0.8%)	
TSPAN11				KISS1R		RNF216	
(2.1%)				(2.6%)		(0.8%)	
				PAX6 (2.6%)		PNPLA6 (0.8%)	
				AXL (2%)		OTX2 (0.8%)	
				CNTN2 (2%)		IGSF1 (0.8%)	
				EBF2 (2%)		FLRT3 (0.8%)	
				EFNA5 (2%)		EBF2 (0.8%)	
				MET (2%)		FGF17 (0.8%)	
				PLXNA1 (2%)			
				SEMA3A (2%)			
				SLIT2 (2%)			
				TACR3 (2%)			
				FEZ1 (1.3%)			
				CCKAR (1.3%)			
				DCAF17 (1.3%)			
				EDNRB (1.3%)			
				EPHA5 (1.3%)			
				GHR (1.3%)			
				HGF (1.3%)			
				NRP1 (1.3%)			
				WDR11 (1.3%)			
				CASR (0.7%)			
				GH1 (0.7%)			
				GNRH1			
				(0.7%)			
				LEI $(0.7\%)$			
				NELE(NSME)			
				(0.7%)			
				PROK2 (0.7%)			
				STS (0.7%)			
				TLE4 (0.7%)			

Table 2 (continued)

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Quaynor et al. (2016)	Wang et al. (2017)	Aoyama et al. (2017)	Cassatella et al. (2018)	Zhou et al. (2018)	Kim et al., (2019)	Amato et al. (2019	Current study
				TYRO3 (0.7%)			
					-		

variants were found less than an average of 3% in these patients (see details in Table 2). Di- and oligogenic cases occurred approximately between 10 and 20% of all cases.

# Genetic counseling

Genetic screening is essential in CHH as it can be treated and patients could have a good reproductive prognosis upon treatment (see details in (Boehm et al. 2015; Maione et al. 2018). Genetic counseling should give information on heritability for other family members too, and also required before family planning (Maione et al. 2018).

In certain cases, heritability can be determined relatively easily. For instance, in case of *GNRH1/GNRHR*, *TAC3/TACR3*, *KISS1/KISS1R*, autosomal recessive inheritance pattern is characteristic, while *ANOS1* is inherited as an X-linked trait (Maione et al. 2018). However, for genes of which variants inherited by an autosomal dominant way, the penetrance and expressivity can be variable. In case of *FGFR1* nearly half, regarding *PROK2/PROKR2* nearly twothird of the cases exhibit incomplete penetrance and variable expressivity complicating the determination of the inheritance pattern (Maione et al. 2018). Regarding certain genes (e.g. *FGFR1*), de novo mutations are also relatively common that has to be taken into consideration when analyzing pedigrees.

Additionally, together with the availability of NGS, the main challenge is to distinguish true oligogenicity from rare variants which appear as incidental findings and are not related to the phenotype. In determination of oligogenicity, genotype-phenotype co-segregation should be assessed by investigating both the affected and healthy family members. In addition, in diagnosis of oligogenicity, Maione et al. (2018) suggested that oligogenic load has to be correlated with phenotype severity. There are several complicating factors (small families, not available or not compliant family members, incomplete penetrance and variable expressivity) in segregation analysis, still, it is one of the most important way to identify the closest evidence of pathogenicity clinically besides in vitro and in vivo studies (Oliver et al. 2015; Maione et al. 2018). Additionally, in clinical interpretation of variants of unknown significance (VUSs), clinical data (genotype-phenotype segregation) are of utmost important.

Once heritability is assessed, risk of disease transmission can be discussed according to the Mendelian rules.

Prognosis has also to be discussed as approximately 20% of the cases appear to be spontaneously reversible. From

genetic point of view, to date *TAC3* and *TACR3* loss of function variants were described to be associated with CHH reversal (Gianetti et al. 2010), but with the increasing data provided by high-throughput NGS platforms, the number of genes connected to this phenomenon will probably increase as well.

# Next-generation sequencing allows evaluation of sequence variants of several genes at the same time in a cost-effective way

Formerly, genetic testing was confined to rare genetic disorders due to their complexity, labour intensity and cost. Now, NGS-based methods are widely available allowing to test even hundreds of genes at the same time. Therefore, NGS has been rapidly integrated into laboratory diagnostics workflows for identification of germline mutations in inherited diseases. Due to its time and cost effectiveness, it is especially useful in cases when several genes have been identified in the background of a certain genetic condition such as CHH.

# NGS-based platform options for clinical genetic diagnostics

Although the technology allows to investigate the sequence of the whole genome (WGS, whole genome sequencing) or exome (WES, whole exome sequencing) currently, the most prevalent applications of NGS in clinical practice are the evaluation of certain genes using targeted gene panels (Di Resta et al. 2018).

As WGS covers the whole genome (coding and noncoding regions) it may seem the most preferable choice in identification of pathogenic gene mutations in inherited diseases. The advantage of WGS is that library preparation is straightforward as it does not require target enrichment. Additionally, data obtained from WGS can easily be used for detection of CNVs. However, among NGS approaches it gives the least average depth of coverage and it is still a costly technology (Di Resta et al. 2018). Also, from clinical point of view, the interpretation of noncoding variants and variants of unknown significance (VUSs) make its utility limited.

WES aims to cover all coding regions in the genome. Exome contains all of the protein-coding regions of genes and it comprises  $\sim 1-2\%$  of the genome, yet contains  $\sim 85\%$  of known disease causing mutations. Its cost is also more preferable and it is a more feasible option comparing to WGS (Di Resta et al. 2018). Usually, the average exome coverage of a WES test is 90–95% due to sequence complexity. WES is sometimes used by clinical laboratories by interpreting only genes which have been already associated with any disease. When mutation has not been identified data analysis can be extended to the remaining exome regions. It has been shown that WES provides diagnosis in approximately of 11–40% of cases where the clinical diagnosis were uncertain (Sawyer et al. 2016). Furthermore, because the depth of coverage for WES is not uniform the sensitivity is usually lower compared to those observed in case of targeted disease panels.

Customized targeted gene panels offer the ability to perform fast and low-cost screening option, therefore, currently, it is the most widely used NGS approach in clinical practice (Di Resta et al. 2018; Wang et al. 2018; Graziola et al. 2019). By focusing on a limited set of genes selected for certain clinical condition, it is able to provide high coverage that increases analytical sensitivity even in detection of mosaicism. Furthermore, because the role of genes included in these panels are known to be associated with the particular condition the detection rate (positive finding) is also higher compared to WES (Di Resta et al. 2018; Wang et al. 2018; Graziola et al. 2019). Targeted panels give the advantage to avoid incidental, secondary findings and to decrease the number of VUSs detected.

Therefore, when the genetic background is well-defined, targeted testing of a gene panel could offer at a relatively low-cost sensitive detection of genetic variants responsible for a disease. However, when no suspect genes stand behind the clinical phenotype, exome sequencing may provide a wider screening option, but in these cases, trio sequencing would allow a more comprehensive result compared to the "only" individual sequencing.

# Workflow of an NGS-based genetic analysis

NGS-based sequencing analysis comprises of three steps: (1) library preparation, (2) parallel sequencing and (3) data analysis and variant interpretation (Oliver et al. 2015).

Molecular genetic analysis is routinely performed using DNA extracted from peripheral blood or buccal mucosa. In our example, we used DNA samples of 38 consecutive patients and 2 family members with hypogonadotropic hypogonadism referred to our diagnostics laboratory. Fourteen patients developed the disease  $\leq 18$  years [2 girls and 12 boys with an average age of 16.2 year ( $\pm 2.1$  years)]. Twenty-four patients developed disease in adult age (3 females, 21 males with an average age of 31.8 years ( $\pm 12.7$  years). Patient characteristics, clinical findings, laboratory results and imaging studies are included in Supplementary Table 1. Our

study was approved by the Scientific and Research Committee of the Medical Research Council of Ministry of Health, Hungary (67/PI/2012). All samples were obtained after acquiring written informed consent from all adult patients and permissions were given by parents of all minors. For NGS-based technologies, the amount and quality of input DNA is an essential factor. Degradation or low concentration of DNA may jeopardize the analysis.

For any NGS-based strategy, library preparation is a key step in the laboratory workflow. The instrumentation determines the library preparations, because high-throughput instruments allow larger analysis. Barcodes (unique, short sequences) are used to label different samples enabling pooling patients' samples into one reaction and decreasing the per-sample cost. Library preparation methods can be grouped into two main categories by principle used for gene amplifications: (1) PCR-based and (2) hybridization-based methods (Butz and Patócs 2019). Although processes using hybridization-based capture are more time consuming and labour intensive, those have the advantage of having greater tolerance against sequence variations (sequence variants and copy-number alteration).

The sequencing characteristics (read length, output read number, cost and run time) of each platform can be different that are needed to be taken into consideration.

For an in-house panel design (gene selection), the recommendation of the European Society of Human Genetics should be followed. Only genes with known relationship between genotype and phenotype should be included in the analysis for diagnostic purposes (Matthijs et al. 2016). Also, the guideline states that "to avoid irresponsible testing, for the benefit of the patients, 'core disease gene list' should be established by the clinical and laboratory experts" (Matthijs et al. 2016). Therefore, consensus statements and guidelines, OMIM (Online Mendelian Inheritance in Man) database and literature search should be assessed to assemble genes in a diagnostic panel. For CHH, there is an available European Consensus Statement (Boehm et al. 2015) which was used as a primary guide during our panel design too.

Accordingly, our panel was designed during the first half of 2017. Some CHH-related genes were left out, mostly those which have been already introduced earlier into clinical practice in our laboratory (e.g. genes responsible for combined pituitary deficiency or adrenal diseases) (Halász et al. 2006; Bertalan et al. 2019) or due to the capacity of the applied method. Genes associated with complex syndromes were not present either in our selection owing to our patient profile. Finally, 41 genes were analyzed (see in Supplementary Table 2 and in Table 3).

We selected NimbleGene approach to create the appropriate hybridization capture probe set for our gene list using NimbleDesign Software (https://sequencing.roche.com/en/ products-solutions/by-category/target-enrichment/software/ nimble-design-software.html) targeting the region of interests (exons + /- 30 bp/exon). Capture probe synthesis was done by the supplier. Library was prepared following double capture; the library quantification was performed following the manufacturer's instructions (NimbleGen SeqCap EZ

Table 3 CHH gene list and panel performance indicated by coverage (mean read/base  $\pm$  SD)

Gene name	Ensembl gene ID	Covered region (bp)	Avg coverage/ base + SD
KISS1	ENSG00000170498	594	$66 \pm 14$
RD3	ENSG00000198570	765	$77 \pm 15$
CXCR4	ENSG00000121966	1128	$90 \pm 23$
NRP2	ENSG00000118257	3813	$86 \pm 18$
IL17RD	ENSG00000144730	2997	$82 \pm 18$
GAP43	ENSG00000172020	1062	$80 \pm 18$
GNRHR	ENSG00000109163	1164	$79 \pm 18$
TACR3	ENSG00000169836	1695	$76 \pm 17$
SPRY4	ENSG00000187678	1146	$74 \pm 12$
GLI3	ENSG00000106571	5640	$80 \pm 15$
SEMA3A	ENSG0000075213	3333	$64 \pm 15$
FGF17	ENSG00000158815	948	$80 \pm 15$
GNRH1	ENSG00000147437	456	$70 \pm 17$
FGFR1	ENSG0000077782	3639	$83 \pm 17$
TRAPPC9	ENSG00000167632	5058	$75 \pm 17$
PALM2	ENSG00000243444	1557	$79 \pm 17$
NOTCH1	ENSG00000148400	9705	$76 \pm 14$
NSMF	ENSG00000165802	2421	$72 \pm 13$
MASTL	ENSG00000120539	3354	$65 \pm 15$
FGF8	ENSG00000107831	1092	$75 \pm 13$
CCKBR1	ENSG00000110148	1641	$80 \pm 15$
FSH	ENSG00000131808	567	$70 \pm 15$
PLEKHA5	ENSG0000052126	5082	$61 \pm 14$
PDE3A	ENSG00000172572	4383	$69 \pm 13$
TSPAN11	ENSG00000110900	1239	$71 \pm 12$
AMN1	ENSG00000151743	1194	$51 \pm 13$
AMHR2	ENSG00000135409	2379	$83 \pm 17$
TAC3	ENSG00000166863	723	$78 \pm 15$
DUSP6	ENSG00000139318	1323	$74 \pm 12$
CRY1	ENSG0000008405	2538	$64 \pm 14$
NOS1	ENSG0000089250	6144	$87 \pm 17$
CDH7	ENSG0000081138	3075	$72 \pm 18$
KISS1R	ENSG00000116014	1494	$49 \pm 19$
AMH	ENSG00000104899	1980	$42 \pm 10$
AXL	ENSG00000167601	3882	$85 \pm 17$
LHB	ENSG00000104826	603	$32 \pm 7$
PROKR2	ENSG00000101292	1332	$94 \pm 20$
JAG1	ENSG00000101384	3834	$78 \pm 16$
FLRT3	ENSG00000125848	2067	$79 \pm 20$
ANOS1	ENSG0000011201	2880	$41 \pm 18$
FGF13	ENSG00000129682	1035	$39 \pm 16$

Library protocol). Sequencing runs were done on Illumina MiSeq instrument using MiSeq Reagent Micro Kit v2.

## Sequencing data processing, performance analysis

During NGS, huge amount of data is produced which require special bioinformatics handling and analysis (Biesecker and Green 2014), therefore, appropriate hardware, software and expert personnel are required for data analysis (Oliver et al. 2015). Currently, there is no gold standard, freely available tool or filtering settings for bioinformatics analysis related to clinical applications of NGS. Each laboratory has to develop and validate its own pipeline (Oliver et al. 2015).

First step of sequencing data analysis is base calling that is integrated into the instrument's software. During the next step, raw sequence reads are aligned to the reference human genome (Sayitoğlu 2016). Quality filtering of read alignment defines sensitivity and specificity of the test. Using very strong filtering could lead to loss of variants, while inclusive filters can minimize false negative results but it will increase the burden of confirmatory analysis. Both coverage depth and uniformity are important regarding detection accuracy. In germline testing, a minimum of 20 reads/ alleles are required for diagnostic purposes. On the other hand, as read/error ratio increases with the increase of coverage practically 300-500 reads/target has been suggested to be enough for diagnostics (Strom 2016; Deans et al. 2017; Butz and Patócs 2019). Even if the coverage is adequate, it is important to evaluate coverage uniformity in order not to miss certain regions falling below the detection cut-off, because variants not detected will not be further analyzed (Rizzo and Buck 2012). In certain cases, due to sequence complexity, 1-2% of the targeted region may not be covered (Rizzo and Buck 2012).

Variant calling is performed to identify alterations compared to the reference sequence (Oliver et al. 2015). In this step, false sequence variants are omitted by investigating variant allele frequency (VAF) (Lee et al. 2014; Deans et al. 2017). (VAF is the percentage of sequence reads divided by the overall coverage of the particular locus. In germline testing, VAF represents diploid zygosity (near 0 and 100% for homozygosity and near 50% for heterozygosity). Unfortunately, results of different variant calling algorithms do not correlate well, therefore, to maintain technical validity, confirmatory tests are recommended (Trubetskoy et al. 2015; Matthijs et al. 2016; Muller et al. 2016). In germline NGS applications, Sanger sequencing is generally accepted for validation.

As In Vitro Diagnosis (IVD) proved NGS-based assays are not widely available, each laboratory has to develop and validate their own protocols including from sample and library preparation, bioinformatics analysis and quality assurance (Rehm et al. 2013). In our analysis, we followed the Genome Analysis Tool Kit (GATK) Best Practices guideline using the germline short variant discovery (SNPs + Indels) algorithm (DePristo et al. 2011). A minimum coverage of 20 reads was applied as detection filter. In our gene panel, all regions were covered by  $71 \pm 14$  reads/base (see details regarding each gene in Table 3).

The accuracy depends on the depth of sequence coverage therefore NGS gene panels show the highest diagnostic accuracy (Oliver et al. 2015). Indeed, in a recent study, comparing different exome sequencing platforms found that 93.2% of the investigated regions were covered > 10 reads (Kong et al. 2018) (of the covered regions the sensitivity was reported 97.5–99.99%). Comparably, in our panel, 97.2% of the investigated regions (86,329 bp) was covered > 20 read/base. Of the investigated region, 14,532 bp was assessed by Sanger sequencing as well, and all detected variants were identified by both approach, therefore the specificity of our panel was 100%.

### Variant interpretation

Variant interpretation are guided by expert recommendations for clinical diagnostics [American College of Medical Genetics and Genomics (ACMG), European Society of Human Genetics (ESHG)] which should be followed for all laboratories offering NGS-based diagnostics (Rehm et al. 2013; Richards et al. 2015; Matthijs et al. 2016).

WGS usually identifies 3–4 million, while WES detects usually 15,000–20,000 variants. Therefore, variant prioritization and interpretation are needed to determine the one or the few pathogenic variants responsible for disease (Fig. 2). First step is to assess the prevalence of certain variants in general population-based databases to filter out frequent variants assuming that pathogenic variants are not common in the broad population. However, in oligogenic diseases, relatively frequent variants can have additional or genetic modifier effect on the phenotype (Maione et al. 2018).

Analyzing the functional consequence of a certain variant may help the interpretation. Using various algorithms

Primary bioinformatic analysis (base calling)	<pre>multiplex quality filtering</pre>
Secondary bioinformatic analysis (variant calling)	$\longleftarrow$ multiplex quality filtering
+	
Identified variants	
1	
Tertiary bioinformatic analysis (variant filtering and prioritization)	Bioinformatic tools/databases
<ul> <li>Frequency filtering (minor allele frequency, databases)</li> </ul>	e.g. 1000 Genome Project, Exome Aggregation Consortiu gnomAD, Exome Sequencing Project
<ul> <li>Structural based filtering (variant impact on codon and gene structure)</li> </ul>	• e.g. SnpEff, VEP
<ul> <li>Prediction based filtering (predicted variant impact based on sequence conservation, structural characteristics and splicing)</li> </ul>	SIFT, PolyPhen, DANN, CADD, PROVEAN, MutationTaster, NNSplice, Splice Predictor, ASSA, et
<ul> <li>Evidence based filtering         (variant, gene, disease specific databases,         and peer-reviewed literature data)</li> </ul>	Variants - e.g. HGMD, Ensemble.org, NCBI ClinVar ← Locus specific - e.g. LOVD, UMD-LSDB Disease specific - e.g. OMIM
<ul> <li>In vitro functional testing (laboratory assays)</li> </ul>	published functional consequences (literature data)
Clinical interpretation	
<ul> <li>Clinical data</li> <li>Pedigree analyis</li> </ul>	Genetic counseling
<ul> <li>Inheritance (phenotype coherence with zygosity)</li> </ul>	

Fig. 2 Process of molecular genetic testing by NGS from NGS data analysis to variant interpretation. See details in the text

(Fig. 2), variants can be classified to have low, moderate or high impact on protein function. It has to be kept in mind that even synonymous variants could sometimes influence splicing and, therefore, amino acid composition of the mature protein resulting in a pathogenic variant (Gianetti et al. 2010; Courage et al. 2019). After classification, further gene, variant and disease specific databases together with peer-reviewed literature data (Fig. 2) can help the accurate interpretation (Richards et al. 2015).

To estimate the pathogenicity of variants of uncertain significance (VUS) is more challenging. Multiple sources of information (variant frequency, in silico predictions of variant effect on protein function and subsidiary functional studies) are needed to be taken into account in order to follow recommendations of guidelines in categorization of a particular variant (Richards et al. 2015; Matthijs et al. 2016). In this framework, in vitro and in vivo functional assays are not always available. These experiments are labour intensive, need longer time, and typically performed as the part of research.

The molecular genetic laboratory report should focus on containing the clinically relevant information for clinicians together with a brief description of all NGS quality metrics (technical characteristics, bioinformatics pipelines, validation), variant annotations and classification (Richards et al. 2015; Matthijs et al. 2016). Disease-specific statements and/ or recommendation can greatly guide the interpreter in variant evaluation. The raw data and the full report should also be available upon request.

In our case, for variant filtering, the following parameters were used: minor allele frequency (MAF) cut-off 1%, coding properties (synonymous variants were omitted), and variants' effects were evaluated by prediction softwares (SNPeffect—Cingolani et al. 2012 and DANN). Variant interpretation was done following the ACMG recommendation (Richards et al. 2015). Additionally, the European Consensus Statement on congenital hypogonadotropic hypogonadism, Human Gene Mutation Database (HGMD) and peer-viewed articles were searched to categorize the detected variants.

All identified Class V, IV and III variants (pathogenic, likely pathogenic and variant of unknown significance, VUS) were validated using conventional bidirectional Sanger sequencing on an Applied Biosystems 3130 Genetic Analyzer System. Following Sanger validation of all pathogenic, likely pathogenic and VUS variants we found 100% of concordance between NGS and Sanger results.

# Pathogenicity of the identified variants, genotypephenotype correlation

After publication of the CHH expert consensus recommendation 7 NGS panel studies have been published about the molecular genetic analysis of CHH. Including our current study, a total of 588 patients with CHH and 72 patients with CDGP were evaluated. Using various NGS approaches of these patients 262 (44%) with CHH and 5 (5.5%) with CDGP diagnosis carried a pathogenic variant (Table 2).

Regarding phenotype-genotype correlation some authors reported inconclusive results and little co-segregation by analyzing pedigrees in their cohorts (Aoyama et al. 2017; Zhou et al. 2018), probably due to the complex genetic background of CHH. However, differences in genetic profile among populations are indicated in Chinese and Japanese cohorts (Aoyama et al. 2017; Zhou et al. 2018). Zhou et al. reported that in Chinese population, cryptorchidism was the most common accompanying feature in addition to CHH, but no single gene in their panel showed association with this abnormality (Zhou et al. 2018). Wang et al. reported that the frequency of PROKR2 mutations was higher in dual CHH patients (showing hypothalamic and/or pituitary defects with testicular hypoplasia) when compared to other CHH cases. The authors suggested that testicular development are affected in early life reflecting the results of animal experiments where the loss of *Prokr2* compromised the integrity of the testicular vasculature (Wang et al. 2017).

In Kallmann syndrome, anosmia/hyposmia is part of the clinical picture, and ANOS1, CHD7, FGFR1, PROK2, PROKR2, and SEMA3A variants were reported to be involved in isolated congenital anosmia (Alkelai et al. 2017). The genetic background of CHH reversal is still unclear, however, the recently identified IGSF10 and GNRHR variants in addition to previously reported TAC3 and TACR3 variants need further studies for clarification of their pathogenic role (Amato et al. 2019).

The genetic background of CDGP and CHH share common aspects, they also have distinct profiles. In CHH patients, both mutations and oligogenicity of CHH genes have been more commonly identified compared to CDGP (Cassatella et al. 2018). In turn, the genetic profile of CDGP resembled more closely to those founded in control cohort. No pathogenic alterations, but frequent (MAF 1.0–2.5%) genetic variants have been more commonly detected in CDGP compared to controls suggesting their genetic modifier's role (Cassatella et al. 2018).

In CHH, oligogenicity was reported between 0 and 19% (Table 2). Interestingly, in Japanese populations Aoyama et al. did not find any patients with CHH caused by di-/ oligogenic mutations (Aoyama et al. 2017) while Quaynor et al. described that the majority of the suggested di- and oligogenic background could be supported by pedigree analysis (9/11 pedigrees) in their cohort (Quaynor et al. 2016). Nevertheless, others suggested that with the increase of the numbers of genes investigated the detection rate of oligogenicity will increase (Amato et al. 2019) making difficult to prove the true role of di/oligogenic findings (Maione et al. 2018).

Table 4	Varia	nts identifie	d by targete	ed NGS panel	in 38 CHH probands								
Patient	Sex	Clinical	Gene_name	Inheritance	Variant identified			ACMG clas-	Minor allele fre	quency (MAF)	Prediction	Gene	Variant Reference
∃		diagnosis			HGVS cdna	HGVS prot	Zygosity	sification	Total	Eu (non- Finnish)	(DANN)	Keterence	(HGMD ID, Ket)
Probands	(n = 38)	(											
с,	<u>г.</u>	KS, anosmia (primer amenor- rhea)	PROKR2	AD, con- tributes to oligogenic- ity	NM_144773.3:c.518 T > G	p.Leu173Arg	Heterozy- gous	Likely patho- genic	0.00229	0.00347	0.9976	Boehm et al. (2015)	Dodé et al. (2006), Cole et al. (2008), Monnier et al. (2009), Abreu et al. (2010), Reynaud et al. (2012), Cassatella et al. (2018), Amato et al. (2019)
Ś	М	KS	AMH	AR (Malone et al. 2019)	NM_000479.4:c.1556C>T	p.Ala519Val	Heterozy- gous	SUV	0.00159	0.00252	0.9582	Malone et al. (2019)	Novel in IHH (It was deteceted in a DSD case by Hughes et al. (2019))
			JAG1	AD (Quaynor et al. 2016)	NM_000214.3:c.3109G > A	p.Asp1037Asn	Heterozy- gous	NUS	0.00000795	0.0000176	0.9918	Quaynor et al. (2016)	Novel in IHH
10	М	KS, anosmia, pubertal delay	FGFR1	AD, con- tributes to oligogenic- ity	NM_001174067.1:c.417delC <sup>4</sup>	p.Ser140Argf- sTer43	heterozy- gous	Pathogenic	Absent	Absent	na	Boehm et al. (2015)	Novel in IHH
			IL.17RD	AD, AR or digenic dominant, contributes to oligo- genicity	NM_017563.5:c.1696C>T	p.Pro566Ser	Heterozy- gous	vus	0.0144	0.0214	1866.0	Boehm et al. (2015)	Novel in IHH [Amato et al. (2019) found a pathogenic variant in the same codon: c.1697C>T; p.Pro566Leu]
12	X		PDE3A	Digenic (Quaynor et al. 2016)	NM_000921.4:c.293C>A	p.Ala98Glu	Heterozy- gous	SUV	0.000431	0.000884	0.7627	Quaynor et al. (2016)	Novel in IHH
13	M	KS	ISONA	XR, con- tributes to oli gogenic- ity	NM_000216.4:c.1700G>A	p.Gly567Asp	Hemizy- gous	Likely patho- genic	Absent	Absent	0.9959	Boehm et al. (2015)	Novel in IHH
15	Σ	KS, anosmia, parosmia	GNRHR	AR, con- tributes to oligogenic- ity	NM_000406.2:c.416G>A	p.Arg139His	Heterozy- gous	Likely patho- genic	0.000144	0.000168	0.9994	Boehm et al. (2015)	Costa et al. (2001), Topaloglu et al. (2006), Beneduzzi et al. (2014)

s         Clinical digensity         Gene. rank         Minor allele fragmenty (M)         Min	- 1												
Bits         Herodysic         LiGVS data         HGVS data         HGVS data         Total         Total         Eu (non- transis)           GL15         dgenic         M.400168.6c.2179G>A         D(JY27Ag         Herody- gous         VUS         0.00327         0.00327           JAG1         AD         AD         M.400168.6c.2179G>A         D(JY27Ag         Herody- gous         VUS         0.00327         0.00357           JAG1         AD         AD         M.400168.6c.2179G>A         D(JY27Ag         Herody- gous         VUS         0.00064         0.00163           Jacua         AD         M.400166.6c.2179G>A         D(JY27Ag         Patewor- gous         VUS         0.00064         0.00163           Jacua         AD         M.400168.6c.2179G>A         D(JY27Ag         Herody- gous         VUS         0.00064         0.00163           Jacua         AD         M.00168.6c.2179G>A         D(JY27Ag         Patewor- gous         VUS         0.00064         0.00163           Jacua         AD         M.00168.6c.2179G>A         D(JY27Ag         Herody- gous         VUS         0.00064         0.00163           Jacua         AD         MOUTAG         AD         MOUTAG         Patewor- gous         VUS         0.000	Clinical		Gene_name	Inheritance	Variant identified			ACMG clas-	Minor allele fr	equency (MAF)	Prediction	Gene P	Variant Reference
GL3         dgene (0uons) e.a.3005         M.000166.6.21790-A         Cu/727Ag         Herrory- gens         U/S         00027         00027         00027         00027           1AG1         AD (Quopes         M.000166.6.21790-A         b.8e1107b         peus- gens         VUS         Abent         Abent           1AG1         AD (Quopes         M.000166.6.21790-A         b.8e1107b         peus- gens         VUS         0.00004         0.00105           1         AD (Quopes         M.00166.6.21790-A         p.017317.5: 40405-A         p.017317.5: 40405-A         p.027         0.00004         0.00105           1         ADP (Quopes         M.00166.6.21790-A         p.031361.00         P.019         Puse         VUS         0.00004         0.00105           1         ADP (Quopes         M.00166.6.21790-A         p.031361.00         P.01573736         P.01573736         Puse         VUS         0.00004         0.00105           1         ADP (Quopes         DO (Quopes         DO (Quopes         P.0173736         P.01537374         P.0157374         P.015944         P.0105         P.010544	diagnosis				HGVS cdna	HGVS prot	Zygosity	suncauon	Total	Eu (non- Finnish)	(NAININ)	Kelerence	(HOMD ID, KEI)
	KS		GLI3	digenic (Quaynor et al. 2016)	NM_000168.6:c.2179G > A	p.Gly727Arg	Heterozy- gous	NUS	0.00527	0.00755	0.9994	Quaynor et al. (2016)	Novel in IHH (Radhakrishna et al. 1999 detected in post- axial polydactyly type-A/B)
NOTCH1         AD?         NM_017617.5c.4049G>T         pArg1350Leu         Heterory- gous         UU         000064         000103           al         Quaynor (Quaynor et al. 2016)         NM_00168.6c.2179G>A         p.Gly/27Arg         Heterory- gous         VUS         0.000537         0.000537           al         AD?         Quaynor (Quaynor et al. 2016)         NM_017617.5c.3860G>A         p.Gly/27Arg         Heterory- gous         VUS         0.000688         0.000101           al         AD?         Quaynor et al. 2016)         NM_01174067.1.c.1012deT*         p.Arg1287Hs         Heterory- gous         VUS         0.0000688         0.000048         0.000191           al         AD.con- oligogenic- in         NM_001174067.1.c.1012deT*         p.Arg1287Hs         Heterory- gous         VUS         0.0000688         0.000049         0.0000191           AB.con- oligogenic- in         AD.con- oligogenic- in         NM_00117303.2.c.2120A>T         p.His707Leu         Heterory- gous         VUS         0.000049         0.000049         0.000049           AB.con- oligogenic- in         AD.cit.         NM_001173303.2.c.2120A>T         p.His707Leu         Heterory- gous         VUS         0.000049         0.000049         0.0000049         0.000004           AB.con- oligogenic- in         NM_001172303.2.c.2			JAG1	AD (Quaynor et al. 2016)	NM_000214.3:c.3320C>T	p.Serl 107Phe	Heterozy- gous	SUV	Absent	Absent	0.9929	Quaynor et al. (2016)	Novel in IHH
GLI3         digenic (Quaynor et al. 2016)         NM_000168.6s.:2179G>A         p.Gly727Arg pous         Heteroy- gous         VUS         0.00527         0.00537           in         Quaynor (Quaynor al         AD; al. 2016)         NM_017617.5s.:3860G>A         p.Arg1287His         Heteroy- gous         VUS         0.000688         0.00191           in         MASTL         AD, con- nity         NM_001174067.1s.:1012deT*         p.Tyr338Mef*         Heteroy- gous         VUS         0.0000688         0.000191           al         Di, con- nity         NM_001172032.sc.2120A>T         p.His707Leu         Heteroy- gous         VUS         0.00004         0.00006           AS this         AD, di, nity         NM_001172303.2sc.2120A>T         p.His707Leu         Heteroy- gous         VUS         0.00004         0.00006           AS con- nity         NM_01172303.2sc.2120A>T         p.His707Leu         Heteroy- gous         VUS         0.000064         0.00006           AR, con- nity         NM_013251.4sc.248A>C         p.His83Pro         Peteroy- gous         VUS         0.000008         0.000068         0.000008           AR, con- nity         NM_013251.4sc.248A>C         p.His83Pro         VUS         0.000008         0.000008         0.000008	KS		NOTCH1	AD? (Quaynor et al. 2016)	NM_017617.5:c.4049G>T	p.Arg1350Leu	Heterozy- gous	SUV	0.000604	0.00103	0.6583	Quaynor et al. (2016)	Novel in IHH
Initial InitialMD7MD3NM_017617.5:c.3860G>Ap.Arg1287HisHeteroxy- gousVUS0.0006880.0001011a1Quaynor (Quaynor iy)AD, con- tributes to oligogenic- iy)NM_001174067.1:c.1012deTT*p.Tyr338Metf- sTer5Heteroxy- gousPathogenic1a1AD, cir, tributes to oligogenic- triputes to oligogenic- triputes to oligogenic- triputes to oligogenic- triputes to oligogenic-NM_001174067.1:c.1012deTT*p.Tyr338Metf- sTer5Heteroxy- gousPathogenic-1a1AD, di-, triputes to oligogenic- triputes to oligogenic- iy)NM_001172303.2:c.2120A>Tp.His707LeuHeteroxy- gousVUS0.000040.0000001a1TAC3AR, con- triputes to oligogenic- iy)NM_013251.4:c.248A>Cp.His83Proheteroxy- gousVUS0.0000080.000009	nCHH, puber delay	tal	GLI3	digenic (Quaynor et al. 2016)	NM_000168.6:c.2179G>A	p.Gly727Arg	Heterozy- gous	SUV	0.00527	0.00755	0.9994	Quaynor et al. 2016	Novel in IHH [Radhakrishna et al. (1999) detected in post- axial polydactyly type-A/B]
<ul> <li>FGFR1 AD, con- tributes to oligogenic- ity</li> <li>MASTL AD, di-, ity</li> <li>MASTL AD, di-, ity</li> <li>MASTL AD, di-, ity</li> <li>MASTL AD, di-, ity</li> <li>MASTL AD, di-, trigenic</li> <li>MASTL AD, di-, (000004 0.00004 gous</li> <li>TAC3 AR, con- tributes to oligogenic- ity</li> <li>TAC3 AR, con- tributes to</li> <li>MASTL AD, di-, stal. 2016</li> <li>MASTL AD, di-, gous</li> <li>MASTL AD, di-, gous<td>KS, anosn puber delay</td><td>nia, tal</td><td>NOTCH1</td><td>AD? (Quaynor et al. 2016)</td><td>NM_017617.5:c.3860G&gt;A</td><td>p.Arg1287His</td><td>Heterozy- gous</td><td>SUV</td><td>0.0000688</td><td>0.000191</td><td>0.9977</td><td>Quaynor et al. (2016)</td><td>Novel in IHH</td></li></ul>	KS, anosn puber delay	nia, tal	NOTCH1	AD? (Quaynor et al. 2016)	NM_017617.5:c.3860G>A	p.Arg1287His	Heterozy- gous	SUV	0.0000688	0.000191	0.9977	Quaynor et al. (2016)	Novel in IHH
MASTL         AD, di-, trigenic         NM_001172303.2:c.2120A>T         p.His707Leu         Heterozy- gous         VUS         0.00004         0.000000           al         Quaynor         Quaynor         gous         gous         0.000004         0.000000           TAC3         AR, con- tributes to oligogenic- ity         NM_013251.4:c.248A>C         p.His83Pro         heterozy- gous         VUS         0.000008         0.000009	KS, ano mia	\$	FGFR1	AD, con- tributes to oli gogenic- ity	NM_001174067.1:c.1012deT <sup>a</sup>	p.Tyr338Metf- sTer5	Heterozy- gous	Pathogenic	1	I	na	Boehm et al. (2015)	Novel in IHH
TAC3 AR, con- NM_013251.4:c.248A>C p.His83Pro heterozy- VUS 0.00008 0.00009 tributes to oligogenic- ity ity 0.00008 0.00009	nCHH, puberi delay	tal	MASTL	AD, di-, trigenic (Quaynor et al. 2016)	NM_001172303.2:c.2120A > T	p.His707Leu	Heterozy- gous	SUV	0.000004	0.000000	0.8767	Quaynor et al. (2016)	Novel in IHH
			TAC3	AR, con- tributes to oligogenic- ity	NM_0132514:c.248A>C	p.His83Pro	heterozy- gous	VUS	0.000008	600000.0	0.977	Boehm et al. (2015)	Novel in IHH [Cassatella et al. (2018) found a pathogenic variant in the same codon: c.248A>G; p.His83Arg]

Table 4	(cont	tinued)											
Patient	Sex	Clinical	Gene_name	Inheritance	Variant identified			ACMG clas-	Minor allele fr	equency (MAF)	Prediction	Gene	Variant Reference
∃		diagnosis			HGVS cdna	HGVS prot	Zygosity	sification	Total	Eu (non- Finnish)	(DANN)	Keterence	(HGMD ID, Ket)
29	X	пСНН	GNRHR	AR, con- tributes to oligogenic- ity	NM_000406.2:c.350 T>G	p.Leu117Arg	Heterozy- gous	Likely patho- genic	absent	absent	0.9971	Boehm et al. (2015)	HGMD ID: CM128135 Gürbüz et al. (2012), Cassatella et al. (2018)
			GNRHR	AR, con- tributes to oligogenic- ity	NM_000406.2:c.158A > T	p.Asn53Ile	Heterozy- gous	SUV	Absent	Absent	0.9881	Boehm et al. (2015)	Novel in IHH
30	M	nCHH	NOTCHI	AD? (Quaynor et al. 2016)	NM_017617.5:c.2734C>T	p.Arg912Trp	Heterozy- gous	SUV	0.001710	0.002570	0.9989	Quaynor et al. (2016)	Novel in IHH [Digilio et al. (2019) reported in association with cardiovas- cular anomalies]
			PROKR2	AD, con- tributes to oligogenic- ity	NM_144773.3:c.701G>A	p.Gly234Asp	Heterozy- gous	SUV	0.00008	0.000018	0.9981	Boehm et al. (2015)	HGMD ID: CM1211225 Tommiska et al. (2013)
31	M	KS, hypos- mia	ANOS1	XR, con- tributes to oligogenic- ity	NM_000216.4:c.1246dupA <sup>a</sup>	p.Thr416Asnf- sTer29	Hemizy- gous	Pathogenic	I	1	па	Boehm et al. (2015)	Novel in IHH
			MASTL	AD, di-, trigenic (Quaynor et al. 2016)	NM_001172303.2:c.871C>T	p.Leu291Phe	Heterozy- gous	SUV	0.000540	0.000940	0.9989	Quaynor et al. (2016)	Novel in IHH
32	M	nCHH, pubertal delay	TACR3	AR, con- tributes to oligogenic- ity	NM_001059.2:c.1345G>T	p.Ala449Ser	Heterozy- gous	SUV	0.000162	0.000305	0.9737	Boehm et al. (2015)	Tusset et al (2012) reported in a case of CDGP
33	М	nCHH, pubertal delay	AMHR2	AR (Malone et al. 2019)	NM_020547.3:c.1484G>A	p.Arg495Gln	Heterozy- gous	Likely pathogenic <sup>a</sup>	0.000203	0.00009	0.9995	Malone et al. (2019)	Novel in IHH
35	М	pubertal delay	KISSIR	AR	NM_032551.5:c.581C>A	p.Ala194Asp	Heterozy- gous	SUV	0.000352	0.000882	0.977	Boehm et al. (2015)	Miraoui et al. (2013)
36	M	pubertal delay	MASTL	AD, di-, trigenic (Quaynor et al. 2016)	NM_001172303.2:c.1505A > G	p.Asp502Gly	Heterozy- gous	SUV	Absent	Absent	0.9219	Quaynor et al. (2016)	Novel in IHH

Table 4	(con	tinued)											
Patient	Sex	Clinical	Gene_name	Inheritance	Variant identified			ACMG clas-	Minor allele fre	quency (MAF)	Prediction	Gene	Variant Reference
9		diagnosis			HGVS cdna	HGVS prot	Zygosity	sification	Total	Eu (non- Finnish)	(DANN)	Keterence	(HGMD ID, Ket)
37	X	pubertal delay	SPRY4	AD	NM_030964.3:c.626G > A	p.Cys209Tyr	Heterozy- gous	۹SUV	0.002170	0.003450	0.9971	Miraoui et al. (2013)	HGMD ID: CM133832 Miraoui et al. (2013)
			НМА	AR (Malone et al. 2019)	NM_000479.4:c.761G>C	p.Arg254Pro	Heterozy- gous	۹SUV	0.000015	0.000000	0.9626	Malone et al. (2019)	Novel in IHH
39	M	KS	FGFR1	AD, con- tributes to oligogenic- ity	NM_001174067.1:c.367G>T	p.Asp123Tyr	Heterozy- gous	vus	0.000059	0.000000	0.9944	Boehm et al. (2015)	Novel in IHH (It was reported in deafness varia- tion database https://deafnessva riationdatabase .org/)
			GL13	digenic (Quaynor et al. 2016)	NM_000168.6:c.1163C>T	p.Pro388Leu	Heterozy- gous	SUV	Absent	Absent	0.7657	Quaynor et al. (2016)	Novel in IHH
40	M	KS	FGFR 1	AD, con- tributes to oligogenic- ity	NM_001174067.1:c.550C>A	p.Pro184Thr	Heterozy- gous	SUV	Absent	Absent	0.9974	Boehm et al. (2015)	Novel in IHH
Gene r	eferen	ices:											

Boehm et al. (2015), Malone et al. (2019), Quaynor et al. (2016)

# Variant references:

Dodé et al. (2006), Cole et al. (2008), Monnier et al. (2009), Abreu et al. (2010), Reynaud et al. (2012), Cassatella et al. (2018), Amato et al. (2019), Hughes et al. (2019), Costa et al. (2001), Topaloglu et al. (2006), Beneduzzi et al. (2014), Radhakrishna et al. (1999), Gürbüz et al. (2012), Digilio et al. (2019), Tommiska et al. (2013), Tusset et al. (2012), Miraoui et al. (2013)

<sup>a</sup>After family screening, we considered this variant not disease causing as the unaffected father carried the same variant in heterozygous form

<sup>b</sup>After family screening, we considered these variants not disease causing as the unaffected father carried both variants in heterozygous form

Our panel identified a total of 31 variants in 22 probands (1)13 patients with only 1 variant per individual; (2) 1 patient with compound heterozygous variants in the GNRHR gene; (3) 8 patients with 2 heterozygous variants in two different genes (digenic case). The digenic rate was 21% (8/38). The eight pathogenic/likely pathogenic variants, detected in eight patients, hence the genetic cause was clearly identified in 21% of cases tested. (8/38 = 21%). In three patients the pathogenic mutations were detected together with variants of unknown significance (VUSs) (Patient IDs: 10, 29 and 31). In addition, in 14 patients, only VUSs were identified (in 6 patients, 2 and in 8 patients, 1 VUS) which need further studies (Table 4). Grouping our patients into adult and pediatric groups, our data show that clearly pathogenic variants in adult patients were identified in 7 (29%; 7/24), while in 1 pediatric cases (7%; 1/14).

In pediatric cases where healthy parents were available for genetic test, we performed family screening. Healthy parents of patients with IDs "26", "33" and "37" were available for assessment of variant pathogenicity. By sequencing the particular variants in unaffected parents, we concluded that combination of SPRY4 p.Cys209Tyr and AMH p.Arg254Pro variants were probably not disease causing as the healthy father also carried the same genotype. Our family screening suggests that the originally predicted as likely pathogenic variant (AMHR2 p.Arg495Gln) is benign for CHH and might be VUS for delayed puberty because the unaffected father carried the same variant in heterozygous form. The co-existence of TAC3 p.His83Pro with MASTL p.His707Leu (Patient "26") could be potentially pathogenic as they were inherited from different parents. Naturally, in CDGP cases, follow-up time (onset of puberty could spontaneously occur) and further studies will possibly clarify the pathogenicity of these variants.

During the family screening of our case, Patient ID "9" the same pathogenic mutation was identified in his clinically affected brother (Patient ID "10"). There are phenotypes are similar, however, in Patient ID "10" anosmia was present.

It is noteworthy that some alterations (such as repetitive sequences, copy-number variations, long insertion-deletions, structural variants, aneuploidy or epigenetic alterations) are not well detectable by NGS methods Therefore, when these types of alterations are expected the appropriate method (such as multiplex ligation probe amplification (MLPA) or microarray-based comparative genomic hybridization (aCGH)) should be used.

# Summary and conclusion

Congenital hypogonadotropic hypogonadism has a heterogeneous clinical phenotype and genetic background. Especially in pediatric cases, even clinical diagnosis can be challenging (ie. pubertal delay vs. hypogonadism). Genetics data regarding hypogonadotropic hypogonadism with the wider availability of next-generation sequencing are increasing but appropriate tool and expertise are needed for correct interpretation of these results in clinical practice. Based on recent data, in more 50% of cases, the disease causing genetic alterations could be found. In house developed gene panels together with appropriate validation steps have at least the same diagnostic accuracy as the WES. The main challenge in NGS-based methods is the interpretation of variants with unknown significance. For clinical point of view, a great majority of data generated by exome and panel sequencing have still been waiting for clinical validation. The potentially new candidate genes and variants have to be further analyzed functionally (in vitro and in vivo animal experiments) together with thorough clinical genotype-phenotype investigations to prove their disease causing effects. The latter is especially challenging in CHH as the clinical phenotype cover a broad spectrum even in cases harboring the same mutation. In CHH, another great challenge is to distinguish true oligogenic inheritance from incidental, rare findings that are not in relation with CHH. The difficulty in determining inheritance due to non-complete penetrance and variable expressivity together with oligogenicity could mean a difficult situation for genetic counselors. However, over time with the increasing genetic data linked to clinical information will reveal the complex genetic landscape of CHH and eventually it will help variant interpretation.

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# **Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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