Kallmann syndrome in a patient with Weiss– Kruszka syndrome and a *de novo* deletion in 9q31.2

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Abstract

Patients with deletions on chromosome 9q31.2 may exhibit delayed puberty, craniofacial phenotype including cleft lip/palate, and olfactory bulb hypoplasia. We report a patient with congenital HH with anosmia (Kallmann syndrome, KS) and a *de novo* 2.38 Mb heterozygous deletion in 9q31.2. The deletion breakpoints (determined with wholegenome linked-read sequencing) were in the *FKTN* gene (9:108,331,353) and in a non-coding area (9:110,707,332) (hg19). The deletion encompassed six protein-coding genes (*FKTN, ZNF462, TAL2, TMEM38B, RAD23B*, and *KLF4*). *ZNF462* haploinsufficiency was consistent with the patient's Weiss–Kruszka syndrome (craniofacial phenotype, developmental delay, and sensorineural hearing loss), but did not explain his KS. In further analyses, he did not carry rare sequence variants in 32 known KS genes in whole-exome sequencing and displayed no aberrant splicing of 15 KS genes that were expressed in peripheral blood leukocyte transcriptome. The deletion was 1.8 Mb upstream of a KS candidate gene locus (*PALM2AKAP2*) but did not suppress its expression. In conclusion, this is the first report of a patient with Weiss–Kruszka syndrome and KS. We suggest that patients carrying a microdeletion in 9q31.2 should be evaluated for the presence of KS and KS-related features.

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Introduction

Congenital hypogonadotropic hypogonadism (CHH) is a rare genetic disease that prevents pubertal development and elicits infertility due to deficient secretion or action of gonadotropin-releasing hormone (GnRH) (1). CHH is called normosmic (nCHH) if the patient has a normal sense of smell, and Kallmann syndrome (KS) if the patient has an absent or deficient ability to smell (1). Both nCHH and KS may manifest with accompanying anomalies, such as bone abnormalities, hearing impairment and ear abnormalities, cleft lip or palate, and anophthalmia and/ or coloboma (1). These diseases present wide phenotypic and genetic heterogeneity, as over 60 genes underlying nCHH and KS have been identified to date (1, 2). However, several nCHH and KS genes remain to be discovered, since the currently known genes account for only half of all cases (1). nCHH and KS might be inherited in a Mendelian or oligogenic fashion depending on the causative gene; large cohort studies suggest that a significant proportion of the cases are oligogenic (1).

Recently, Weiss and Kruszka reported a series of patients with developmental delay, distinct craniofacial phenotypes, and hearing loss and showed that this

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syndrome is due to loss-of-function ZNF462 mutations or microdeletions in the 9q31.2 area (3). We report a patient with a complex phenotype including Weiss-Kruszka syndrome, KS, and a de novo deletion on chromosome 9q31.2. The deletion encompassed ZNF462, FKTN, TAL2, TMEM38B, RAD23B, and KLF4. To date, 25 patients with Weiss-Kruszka syndrome have been reported (3, 4, 5, 6), yet delayed puberty, anosmia, or CHH are not among the listed phenotypic features. On the other hand, deletions in the 9q31.2 chromosomal area have been continuously described since the 1970s (7, 8, 9, 10, 11, 12, 13, 14, 15), at least one of whom had CHH (11), olfactory bulb hypoplasia (14), or delayed puberty in multiple family members (15), and at least four had cleft lip or palate (8, 10, 13, 14), that is, phenotypic features reminiscent of KS. However, no clear link between the 9q31.2 deletions and complete KS currently exists. The closest connection comes from disruption or pathogenic variants in the closely located PALM2AKAP2 locus. Indeed, PALM2 and AKAP2, which can form fusion transcripts, constitute potential KS disease genes, since a female with KS and Graves' disease carried a missense mutation, which was predicted to be deleterious, in PALM2 (16), and a male with KS and bone anomalies carried a balanced chromosomal translocation that disrupted AKAP2 expression (17).

To investigate the putative relationship between the proband's 9q31.2 deletion and KS, we first defined the exact deletion breakpoints with linked-read wholegenome sequencing. Next, we investigated if the deletion was associated with decreased expression of *PALM2AKAP2* and screened this gene in a set of Finnish KS patients. Since these investigations did not support the connection between *PALM2AKAP2* and KS, we performed wholeexome and RNA sequencing in the proband and his family members to exclude defects in genes implicated in KS and to reveal potential new candidate genes. Our results, together with the CHH and KS-related phenotypes in previous patients with 9q31.2 deletions suggest that microdeletions in this chromosomal region underlie KS.

Subjects and methods

Subjects

We investigated a Finnish family whose son (the proband) had been assessed for delayed development and subsequently for the absence of puberty. He was the second child of healthy nonconsanguineous parents (Fig. 1), and he had a healthy sister, who had normal timing of puberty since she reached menarche at the age of 12

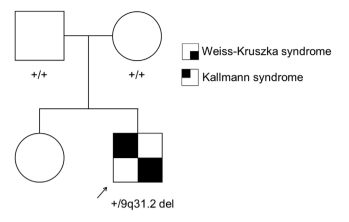


Figure 1

Pedigree of the family, in which the proband (*arrow*) carried a heterozygous *de novo* 2.38 Mb deletion in chromosome 9q31.2 that encompassed *FKTN*, *TAL2*, *TMEM38B*, *ZNF462*, *RAD23B*, and *KLF4*. The proband had Weiss–Kruszka syndrome due to *ZNF462* haploinsufficiency with ptosis, flat nasal tip, low set ears, mild developmental delay, and mild bilateral sensorineural hearing loss, and Kallmann syndrome (congenital hypogonadotropic hypogonadism, anosmia, and the absence of olfactory bulbs).

years. The mother's menarche age had been 11 years, and the father had no reported history of delayed puberty. Following an uneventful pregnancy, the proband was born at gestation week H42+1. His birth weight was 3240 g and his length was 50 cm. He received 4, 6, and 9 Apgar points at the ages of 1, 5, and 10 min, respectively. He was noted to have a muscular ventricular septal defect that closed spontaneously. His testes were normally descended. Before going to school, he was diagnosed with attention deficit disorder and mild developmental delay. He had distinctive facial features including ptosis, flat nasal tip, low set ears, and mild bilateral sensorineural hearing loss with normal semicircular canals in MRI. At the age of 6.5 years, he underwent comparative genetic hybridization (Agilent 44K), in which a 2.38 Mb deletion in the 9q31.2 area was detected, yet the precise breakpoints remained undetermined. The deletion was de novo, as his parents tested negative for the presence of the deletion.

At the age of 13.5 years, the proband was referred for the evaluation of delayed puberty and was noted to have a hypoplastic scrotum and testes. He had selfreported anosmia, which had been well appreciated in the family and absent olfactory bulbs in the MRI. He had no synkinesia, hand or foot deformities, pigmentation defects or missing teeth, nor he reported problems with balance. Abdominal ultrasound revealed two kidneys. His circulating reproductive hormone levels were low

(LH 0.1 IU/L, FSH 0.2 IU/L, testosterone 0.7 nM, and inhibin B level was very low, 14 ng/L). At the age of 14 years and 11 months, he was still prepubertal (Tanner stage G1) with pubic hair stage P3. His testes were very small, only 1 cm in length (corresponding to < 1 mL). His reproductive hormone levels were still low; serum LH 0.1 IU/L, inhibin B 23 ng/L, and testosterone 0.4 nM. Based on the very small testicular size, low reproductive hormone and inhibin B levels, anosmia, and the absence of olfactory bulbs in the MRI scan, he was prepubertal, and a diagnosis of Kallmann syndrome was set (1, 18). Induction of his puberty with low-dose testosterone was commenced. During the treatment, his testis size remained extremely small (assessed at the age of 16 years and 9 months) in the setting of low gonadotropin and inhibin B (23 ng/L at the age of 15 years and 9 months; 29 ng/L at 16 years 9 months, and 28 ng/L at 17 years 7 months) levels consistent with the KS diagnosis. As a part of the diagnostic procedure, he underwent BluePrint Genetics® Kallmann Syndrome Panel Plus (version 3), a diagnostic platform that covers the coding regions of the following genes: ANOS1, CHD7, FGF8, FGFR1, GNRHR, KISS1R, PROK2, PROKR2, and TACR3 up to 20 bp of the intronic sequence with single nucleotide changes, small indels up to 220 bp, and copy number variations defined as single exon or larger deletions and duplications.

Determination of the deletion breakpoints and presence of structural variants in KS genes with whole-genome linked-read sequencing

DNA of all family members was extracted from peripheral blood leukocytes (PBL). The proband and parent RNA were isolated with the QIAamp RNA Blood Mini Kit (QIAGEN) from PBLs. The whole-genome linked-read sequencing of the proband was performed at FIMM (Institute for Molecular Medicine Finland) according to 10x Genomics Chromium library preparation (Chromium Genome Reagent Kits v2 RevB; 10× Genomics, Pleasanton, CA, USA). The sample was sequenced with NovaSeq 6000 system (Illumina Inc., San Diego, CA, USA) using S4 flow cell and XP workflow. Read length for the paired-end run was 2×151. The reads were aligned to the GRCh37 (hg19) reference genome. The data were analyzed with Longranger 2.2.2 WGS pipeline with default parameters and GATK for variant calling and visualized with Loupe 2.1.1 (both programs by 10× Genomics). The genome was covered with a mean depth of 34.7×, and a mean molecule length of 26.8 kb. Loupe interactive visualization tool 2.1.1 was employed to define the exact 9q31.2 deletion

breakpoints. In addition, as MLPA (multiplex ligationdependent probe amplification) is available for only a fraction of known KS genes, we verified the absence of genomic structural variants of over 40 bp in size in the coding regions of 32 known KS genes (ANOS1, FGFR1, FGF8, FGF17, PROK2, PROKR2, CHD7, NSMF, HS6ST1, WDR11, SEMA3A, SEMA7A, PLXNA1, SOX10, IL17RD, FEZF1, NDNF, TCF12, TUBB3, DCC, SMCHD1, KLB, NTN1, SPRY4, PTCH1, FLRT3, AMH, DUSP6, PLXNA3, NRP1, SPRY2, and NRP2) with Loupe 2.1.1.

Investigation of *PALM2AKAP2* expression with RT-qPCR

The RNA extracted from the proband and his parents were converted into cDNA using the iScript cDNA Synthesis Kit (Bio-Rad) according to the manufacturer's instructions in a regular thermocycler with 1 μ g of total RNA. The synthesized cDNAs were used as templates to assess the mRNA expression with a ready-to-use qPCR mix (Solis Biodyne) in a quantitative PCR machine (Roche LC480 II). The expression levels of *PALM2AKAP2* were normalized to *GAPDH*. The relative expression levels were standardized against a healthy unrelated control sample, which was given an arbitrary value of 1.0. The PCR conditions and primers are available on request.

Sanger sequencing of *PALM2AKAP2* in Finnish Kallmann syndrome patients

We examined a set of 16 Finnish KS patients (15 men, 1 woman) without mutations in ANOS1, FGFR1, FGF8, PROK2, PROKR2, CHD7, and WDR11 (19, 20). The exons and exon-intron boundaries of PALM2 (NC_000009.11, NM_053016.6, GRCh37) and AKAP2 (NC_000009.11, NM 001198656.1, GRCh37) were amplified with PCR from the genomic DNA. The PCR conditions and primers are available upon request. The PCR products were purified with ExoProStar treatment (GE Healthcare Life Sciences), and sequenced from both directions using the ABI BigDyeTerminator Cycle Sequencing Kit (v3.1) and ABI Prism 3730xl DNA Analyzer automated sequencer (Applied Biosystem). The DNA sequences were aligned and read with Sequencher® 4.9 software (Gene Codes Corporation, AnnArbor, MI, USA). As one of the patients harbored a variant NC_000009.11:g.112918586C>T, in intron 2 of AKAP2, his RNA was converted into cDNA by using the SuperScript® III First-Strand Synthesis System for RT-PCR kit (Invitrogen by Life Technologies) according to the manufacturer's instructions. Part of

the *AKAP2* transcript was PCR-amplified with cDNAspecific primers and subsequently sequenced. Effect of the *AKAP2* variant NC_000009.11:g.112918586C>T was also predicted with Human Splicing Finder (https://www. genomnis.com/access-hsf) (21), MutationTaster (http:// www.mutationtaster.org/) (22), NetGene2 (http://www.cbs. dtu.dk/services/NetGene2/) (23, 24), and BDGP NNSPLICE (https://www.fruitfly.org/seq_tools/splice.html) (25), and its clinical interpretation was performed with InterVar (http://wintervar.wglab.org/) (26).

Whole-exome sequencing

The whole-exome sequencing (WES) of the proband's family was performed with Illumina Novaseq S2 PE100 technology. First, the adapter was trimmed from the reads, as well as any low-quality nucleotides from the 5' or 3' ends of the read, removing pairs with less than 36 bp. The reads were aligned to the GRCh37 (hg19) reference genome with the BWA (Burrows–Wheeler Aligner). Non-unique read pairs and non-unique single reads were removed and GATK Base Recalibrator was used to clean the alignment. Any potential PCR duplicates were removed using Picard MarkDuplicates, and GATK IndelRealigner was used for indel sites. The mpileup from the SAMTOOLS package was used for variant calling (27). The sequencing yielded a mean target coverage of 222× and 98% of 20× coverage.

Data analysis

First, we verified that the coding exons of 32 KS-associated genes (ANOS1, FGFR1, FGF8, FGF17, PROK2, PROKR2, CHD7, NSMF, HS6ST1, WDR11, SEMA3A, SEMA7A, PLXNA1, SOX10, IL17RD, FEZF1, NDNF, TCF12, TUBB3, DCC, SMCHD1, KLB, NTN1, SPRY4, PTCH1, FLRT3, AMH, DUSP6, PLXNA3, NRP1, SPRY2, and NRP2) were covered in WES by using the BasePlayer analysis and visualization tool (28). Subsequently, the proband's VCP file, provided by FIMM (27), was annotated using ANNOVAR (https:// doc-openbio.readthedocs.io/projects/annovar/en/ latest/). We defined that a potentially causative variant should be in the consensus splice-site or it should be nonsynonymous and not classified as benign by any of the ten applied in silico tools (SIFT, LRT, MutationTaster, MutationAssessor, FATHMM, PROVEAN, MetaSVM, MetaLR_pred, M-CAP, and fathmm-MKL). Additionally, a potentially causative variant(s) should (i) occur only in the proband (de novo) (and have a minor allele frequency MAF <0.1%); (ii) be biallelic (autosomal

recessive inheritance) (MAF < 2%); (iii) be monoallelic and inherited from one parent and be absent in the sister (autosomal dominant) (MAF <0.1%); or (iv) be monoallelic and inherited in the X chromosome (X-linked recessive) (MAF <2%). The frequency criteria were applied for all ethnic subpopulation frequencies in gnomAD, ExAC, 1000 Genomes, and Exome Variant Server (provided in the ANNOVAR annotation) and in dbSNP (https://www.ncbi.nlm.nih.gov/snp/) (29) (manually verified). Finally, to be accepted for further analyses, a potentially causative variant in WES had to be present in the RNA sequencing data and/or the linked-read WGS data. We confirmed the genotype and segregation of the two such variants in all family members with Sanger sequencing. The primers were designed for sequences around candidate variants in RIMBP3C (NC_000022.10, NM_001128633.2) and SARS1 (NC_000001.10, NM_006513.4) (GRCh37). The PCR conditions and primers are available upon request.

RNA sequencing

The quality and integrity of the RNA were assessed with high sensitivity D5000 Screen Tape® System (Agilent). Following the quality control, RNASeq libraries were prepared using NEBNext Ultra II Directional Poly (A) Capture Kit. RNA sequencing was carried out as pairedend sequencing with Illumina NextSeq 500 Mid Output System. RNA quality-testing, library preparation, and sequencing were performed in the Functional Genomics Unit (FUGU) at the University of Helsinki, Finland.

RNA-sequencing data analysis

Raw adaptor-trimmed FASTQ files were subjected to quality assessment with the FASTQC tool (Simon Andrews, Babraham Bioinformatics). Based on the overall data quality and the individual reads, additional trimming was unnecessary. The sequences were aligned to the GRCh37 human genome version with the STAR package (30). Splicing events of known KS genes were visually verified with the Sashimi plot feature of the MISO framework (31) in the IGV browser from Broad Institute (32).

Statistics and the probability estimate for the occurrence of a 9q31.2 microdeletion and KS

The relative expression values of *PALM2AKAP2* in the PBLs of the proband (n = 4) and his parents (n = 4) were compared with one-way ANOVA, and followed by Tukey's

HSD *post-hoc* analysis. P-value < 0.05 was accepted to indicate statistical significance. To evaluate the role of the 9q31.2 microdeletion in our proband, we utilized the CNV data obtained from the UK Biobank (33, 34, 35), in which 4 deletions, 1 Mb or larger in size, in a set of 472 734 people, overlapped the 24 Mb region, which is depicted with a rectangle on chromosome 9 in Fig. 2 (chr9:102,253,143-126,253,089, GRCh37). Thus, the estimated probability for our proband not to harbor a causal mutation in known KS genes and to carry such a microdeletion by chance would be very small (1:2 \times 4:472 734=4.23e-06). Moreover, at the population level, the probability to encounter a Finnish male with KS (incidence 1:30 000) (19) without a causative mutation in KS genes, who would carry such a microdeletion would be even several magnitudes smaller (i.e. $1:30\ 000 \times 1:2 \times 4:472\ 734 = 1.41e-10$).

Results

We investigated the genetic and phenotypic features of a patient with KS and Weiss-Kruszka syndrome who carried a 2.38 Mb deletion on chromosome 9q31.2. A diagnostictargeted sequencing panel for KS that covered the coding exons of genes ANOS1, CHD7, FGF8, FGFR1, GNRHR, KISS1R, PROK2, PROKR2, and TACR3 up to 20 bp intronic sequence and structural variants was negative for likely pathogenic variants.

Determination of the 9q31.2 deletion breakpoints

To define the exact deletion breakpoints, we performed a whole-genome linked-read sequencing of the proband's DNA. He carried a 2.38 Mb deletion on chromosome 9:108,331,353-110,707,332, excluding PALM2AKAP2 locus that is located at 9:112,542,589-112,934,792 (hg19) (Fig. 2). Instead, the deletion disrupted FKTN, which contains ten exons, at 9:108,331,353 in the first intron, and no other known gene at 9:110,707,332. The deletion encompassed six protein-coding genes (FKTN, TAL2, TMEM38B, ZNF462, RAD23B, and KLF4), none of which has been implicated in KS.

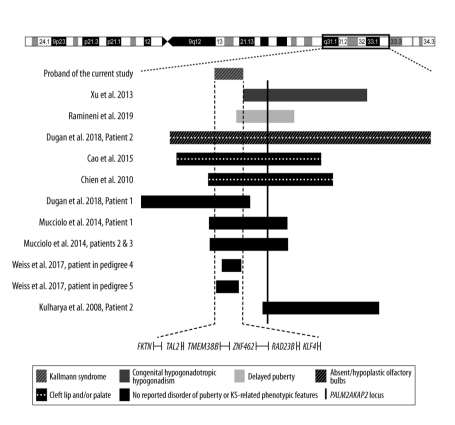


Figure 2

Schematic representation of the proband's 9q31.2 deletion in relation to other informative patients with deletions encompassing this area. The dashed vertical lines indicate our proband's 2.38 Mb deletion (chr9:108,331,353-110,707,332) (hg19), which encompassed six protein-coding genes (FKTN, TAL2, TMEM38B, ZNF462, RAD23B, and KLF4). Rectangles indicate relative locations and sizes of deletions in various patients, their colors and patterns highlight important Kallmann syndrome-related phenotypes and the vertical line indicates a Kallmann syndrome candidate gene locus (PALM2AKAP2). A summary of the patients is presented in Supplementary Table 2. The figure is modified from (14). The deletion coordinates are presented as GRCh37 (hg19). For Kulharya et al. (patient 2), Chien et al., and Ramineni et al. patients the UCSC genome coordinate conversion tool https://genome.ucsc.edu/ cgi-bin/hgLiftOver was employed to convert hg18 coordinates to hg19.

PALM2AKAP2 expression in peripheral blood leukocytes

Given that *PALM2AKAP2* has been implicated in KS and our proband's deletion laid approximately 1.8 Mb upstream of the *PALM2AKAP2* locus (Fig. 2), we investigated whether the expression of *PALM2AKAP2* was altered in peripheral blood leukocyte RNA. However, the proband's *PALM2AKAP2* transcript was expressed at a higher level than in the father (Fig. 3), indicating that the deletion does not suppress *PALM2AKAP2* expression.

Sanger sequencing of *PALM2* and *AKAP2* in 16 Finnish KS patients

To further probe the possible role of *PALM2AKAP2* in KS, we screened the coding exons and exon-intron boundaries of both *PALM2* and *AKAP2* in 16 Finnish KS patients

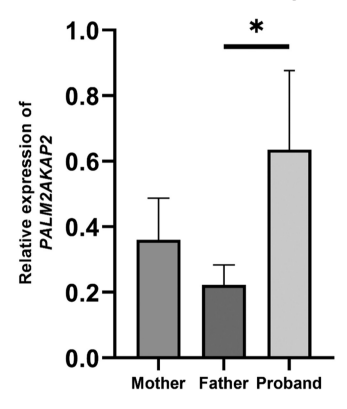


Figure 3

Relative expression of *PALM2AKAP2* in the peripheral blood leukocytes of the proband with Kallmann syndrome, Weiss– Kruszka syndrome, and a heterozygous *de novo* deletion in 9q31.2, and his healthy parents. The bars represent four replicates of *PALM2AKAP2* expression measured by qPCR, normalized by a housekeeping gene (*GAPDH*), and adjusted against an unrelated control sample. *P < .05 (ANOVA followed by Tukey HSD *post-hoc* analysis). without mutations in established KS-causing genes (19, 20). In the intron 2 of AKAP2, we found one heterozygous variant, NC_000009.11:g.112918586C>T (rs777796314), with a frequency of 0.00004267 in Finns (gnomAD, https://gnomad.broadinstitute.org/) (36). This variant was classified as likely benign according to the 2015 ACMG/ AMP guidelines (InterVar, (http://wintervar.wglab.org/)) (26). The carrier was diagnosed with KS at the age of 14 years. He had sensorineural hearing loss in the right ear, micropenis, normal MRI, and alopecia at the age of 5. He had no family history of delayed puberty. Given that two out of four *in silico* tools predicted the variant to affect the acceptor site of exon 3 (MutationTaster: disease-causing; NetGene2: alteration of acceptor sites), we confirmed in vitro that the splicing was normal (data not shown). All other encountered PALM2 and AKAP2 variants had MAFs above 1% among gnomAD Finns.

Whole-exome sequencing, RNA sequencing, splicing analysis, and identification of potentially causative variants

Given that the targeted KS gene panel, the deletion in 9q31.2 per se, or the adjacent candidate gene PALM2AKAP2 provided no equivocal answer to our question of whether this deletion underlies KS in the proband, we sequenced the whole exome in the proband, his healthy parents and sister. First, no rare sequence variants were found in the 32 genes implicated in KS (IL17RD, SPRY2, DUSP6, CHD7, FGFR1, SOX10, ANOS1, FGF8, FGF17, PROK2, PROKR2, NSMF, HS6ST1, WDR11, SEMA3A, SEMA7A, PLXNA1, FEZF1, NDNF, TCF12, TUBB3, DCC, SMCHD1, KLB, NTN1, SPRY4, PTCH1, FLRT3, AMH, PLXNA3, NRP1, and NRP2). Secondly, we investigated in the linked-read WGS data whether the proband carried genomic structural variants of over 40 bp in the coding regions of these 32 KS genes. However, none of the genes were disrupted by structural variants. As WES may not detect non-coding variants that affect splicing, we investigated the splicing events of KS genes in the proband and his parents with RNA sequencing and subsequent visualization of transcripts. Overall, 15 KS genes, FGFR1, PROK2, CHD7, NSMF, HS6ST1, WDR11, SEMA3A, SEMA7A, PLXNA1, TCF12, SMCHD1, PTCH1, and DUSP6, were expressed in PBLs, and none of them displayed aberrant splicing events (data not shown). Finally, we investigated whether the WES data would contain new potential KS candidate gene mutations. Two rare variants of uncertain significance emerged, one in RIMBP3C and the other in SARS1. An overview of the variants is shown in Supplementary Table 1 and Supplementary Material

(see section on supplementary materials given at the end of this article).

Overlapping genetic region with our proband and the previously reported patient with CHH

Our proband's deletion (9:108,331,353-110,707,332) and the deletion of the previously described patient with CHH (9:110,672,051–120,997,503) (11) overlapped in 9:110,672,051–110,707,332 (hg19). This region contained ribosomal pseudogene *RNA5SP293*. According to the UCSC Genome Browser (https://genome.ucsc.edu/), the following transcription factors were determined to have binding sites in this overlapping region: GATA3, MYC, JUND, CEBPB, TEAD4, GATA2, MAFF, USF2, USF1, BHLHE40, ZNF143, ATF3, and SIX5.

Discussion

We report a patient with KS, developmental delay, ptosis, ventral septal defect, craniofacial phenotype, and mild bilateral sensorineural hearing loss without coding region mutations in the established KS genes, which was assessed by using a targeted sequencing panel, linked-read whole-genome sequencing, RNA sequencing, and WES. Instead, he carried a heterozygous *de novo* microdeletion in 9q31.2, which we estimated to be extremely rare (AF below 1/200 000). The deletion encompassed loci of six protein-coding genes (*FKTN*, *TAL2*, *TMEM38B*, *ZNF462*, *RAD23B*, and *KLF4*).

The deletion was 2.38 Mb in size and the 5' breakpoint broke FKTN in the first intron. However, to the best of our knowledge, *FKTN* is not implicated in KS or CHH (37, 38). Haploinsufficiency of ZNF462 is associated with Weiss-Kruszka syndrome (3), the most typical symptoms of which include developmental delay, ptosis, metopic ridge, facial asymmetry, corpus callosum dysgenesis, downslanting palpebral fissures, arched eyebrows, epicanthal folds, short upturned nose with a bulbous tip, marked cupid bow/wide philtrum, low set ears, hypertelorism, and hearing loss (3, 5). Our proband, thus, clearly exhibited Weiss-Kruszka syndrome, and several putative links could connect ZNF462 and the reproductive system. For instance, fibroblast growth factor 8, required for GnRH neuron ontogeny, also induces Znf462 expression in the chick pre-placodal region, and hypothalamic expression of ZNF462 decreases at puberty in non-human primates (39, 40). On the other hand, mutations in ZNF462 have not been reported to cause delayed puberty or affect

the sense of smell (3, 4, 5, 6), suggesting that ZNF462 haploinsufficiency alone is not sufficient to explain the KS phenotype. In humans, the strongest GWAS signals that associated with the variation in the age at menarche were at 9q31.2 (rs2090409), the nearest genes to this SNP being TMEM38B, FKTN, TAL2, and ZNF462 (41), as well as SNPs rs12684013, rs4452860, and rs7028916 (42) in the intergenic regions in 9q31.2 (menarche, age at, quantitative trait locus 3, i.e. MENAQ3 locus; MIM 612883), which were all deleted in our proband. In a recent GWAS study, TMEM38B was associated with puberty timing in both males and females (43). Moreover, despite the putative association of the 9q31.2 region to age at menarche and puberty timing in GWAS studies, mutations in the genes deleted in our proband have not been implicated in CHH or KS.

KS-related phenotypes in our proband and representative previously reported patients with 9q31.2 deletions are shown in Fig. 2 and Supplementary Table 2. There are two previous deletions that overlap with the one encountered in our proband and are associated with delayed puberty or CHH (11, 15). Ramineni et al. reported a family, in which a deletion in 9q31.2 segregated with delayed puberty. The overlapping region between their deletion and that of ours encompassed only one proteincoding gene, KLF4. KLF4 is coexpressed with SOX10, a known KS gene, in the chick neural crest (44), and KLF4 interacts with β-catenin and inhibits Wnt signaling (45) by preventing β -catenin binding to TCF7L2 (46), a transcription factor implicated in the development of the hypothalamus and pituitary (47, 48). The deletion in our patient and in the family reported by Ramineni et al. also contained the SNP rs139300691, which is associated with the sense of smell (49). The second informative patient, originally classified to have delayed puberty by Xu et al., had the onset of puberty at the age of 18 years, and his Tanner stage was G2 at the age of 20 in the setting of normal gonadotropin levels (11). Thus, he fulfilled the criteria of CHH (1). The ~35 kb region shared by our proband and Xu's patient (11) contained only a ribosomal pseudogene RNA5SP293 (Fig. 2). This region also contains binding sites for ZNF143 (a transcription factor which interacts with a puberty-related factor, LIN28B, in neuroblastoma cells (50)), JUND (JunD binds to Gnrhr AP-1 site in mouse gonadotrope-derived α T3-1 cells (51) and is expressed in the mouse olfactory bulb (52)), GATA2 and GATA3 (crucial for neurogenesis and expressed in the olfactory bulbs (53)) and CEBPB (Cebpb in rats is expressed in the olfactory bulbs and olfactory ensheathing cells (54) and is known to be an upstream transcriptional regulator of

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Gnrh1 in mice (55)). However, the mechanism by which a loss of transcription binding site(s) alone would cause disease is difficult to decipher, especially as the closest KS candidate gene locus (*PALM2AKAP2*) is relatively far away, 1.8 Mb, downstream from the 3' breakpoint of the deletion. This locus in our patient with KS was, however, spared and expressed *PALM2AKAP2* efficiently (Fig. 3). Moreover, screening a set of Finnish KS patients revealed no pathogenic variants in *PALM2AKAP2*. Of note, the cases displayed in Fig. 2 showed wide phenotypic variability and we were able to verify KS-related features in only a proportion of them (Supplementary Table 2 and Supplementary Material).

Taken together, variation in the chromosomal region 9q31.2 at the population level associates with the variation in puberty timing and the sense of smell. Previous rare patients with deletions including 9q31.2 have exhibited CHH and KS-related phenotypes such as delayed puberty, an olfactory bulb defect, and cleft lip and/or palate. We describe the first patient with an extremely rare de novo deletion in 9q31.2 and KS. ZNF462 haploinsufficiency was consistent with the patient's Weiss-Kruszka syndrome but did not explain his KS. In further analyses, he did not carry rare sequence variants in known KS genes. Our results suggest that patients carrying a microdeletion in 9q31.2 should be evaluated for the presence of KS and KS-related features, and conversely, that KS patients with features of Weiss-Kruszka syndrome should be evaluated for the presence of a heterozygous 9q31.2 microdeletion.

Supplementary materials

Declaration of interest

The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of this study.

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Ethics committee approval

Author contribution statement

T R, V S, K V collected the study subjects and the clinical data and phenotyped the patients. J K, A-I P, V Y, H A, K V and T R analyzed the next-generation sequencing data, H A carried out ANNOVAR annotation, A-I P and K V performed Sanger sequencing data analyses. V Y performed *PALM2AKAP2* RT-qPCR analyses; A-I P, J K, V Y, K V and T R wrote the manuscript, and all authors critically revised the manuscript. T R coordinated and managed the study.

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