## Running title: A spectrum of MED12-related disorders

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#### Abstract

MED12 is a member of the large Mediator complex that controls cell growth, development, and differentiation. Mutations in MED12 disrupt neuronal gene expression and lead to at least three distinct X-linked intellectual disability (XLID) syndromes (FG, Lujan-Fryns, and Ohdo). Here we describe six families with missense variants in MED12 (p.(Arg815Gln), p.(Va1954Gly), p.(Glu1091Lys), p.(Arg1295Cys), p.(Pro1371Ser) and p.(Arg1148His), the latter being firstly reported in affected females) associated with a continuum of symptoms rather than distinct syndromes. The variants expanded the genetic architecture and phenotypic spectrum of MED12-related disorders. New clinical symptoms included brachycephaly, anteverted nares, bulbous nasal tip, prognathism, deep set eyes, and single palmar crease. We showed that MED12 variants, initially implicated in X-linked recessive disorders in males, may predict a potential risk for phenotypic expression in females, with no correlation of the X chromosome inactivation pattern in blood cells. Molecular modeling (Yasara Structure) performed to model the functional effects of the variants strongly supported the pathogenic character of the variants examined. We demonstrated that molecular modeling is a useful method for in silico testing of potential functional effects of MED12 variants and thus can be a valuable addition to the interpretation of the clinical and genetic findings.


## KEY WORDS

MED12, X-linked intellectual disability, FG syndrome, Lujan-Fryns syndrome, Ohdo syndrome, molecular modeling

## INTRODUCTION

The Mediator is a large multiprotein complex that regulates gene expression in all eukaryotes and interacts with RNA polymerase II. The role of the complex involves transcriptional elongation and termination, mRNA processing, noncoding RNA activation, super enhancer formation, and epigenetic regulation. Thus, the Mediator was called an '"integrative hub" for transcriptional processes and emerged as a master coordinator of cell growth and homeostasis, development, and differentiation ${ }^{1}$. The Mediator complex (MED) consists of 31 subunits (MED1-MED31) and can be divided into four distinct modules termed as the head, middle, tail, and CDK8 kinase module. The latter module contains CDK8, cyclin C, MED12 and MED13. MED12 is a critical transducer of regulatory information essential for organogenesis. At least three different intellectual disability (ID) conditions that are associated with MED12 mutations have been described. These conditions include OpitzKaveggia syndrome (FG syndrome type 1, FGS1), Lujan-Fryns syndrome (LFS), and Xlinked Ohdo syndrome (XLOS, OHDOX) ${ }^{2}$. These syndromes are allelic disorders that share clinical findings including ID, hypotonia and some physical features, such as tall prominent forehead, open mouth or high narrow palate. Probands with non-syndromic X-linked ID (XLID), including females with variable cognitive impairment, have also been described as harboring MED12 mutations. Here, we present clinical and genetic details on six families with missense variants in the MED12 gene, as well as molecular modeling results for four variants, and expand the phenotypic spectrum of MED12-related disorders.

## MATERIALS AND METHODS

Six ID families with MED12 variants (NM_005120) were included in the study (Fig.1). Patients' phenotypes are described in Supplementary material 1 and summarized in Supplementary material 2. Families 1 to 4 were collected as part of the X -exome resequencing project of unresolved families with assumed XLID, while Families 5 and 6 were sequenced in routine diagnostics. For none of the families a clinical suggestion had been made prior to next generation sequencing (NGS). The procedures employed were reviewed and approved by the appropriate institutional review committees. DNAs were isolated from peripheral blood according to standard procedures. Written informed consent was obtained for all individuals participating in this study. For the index probands NGS was performed in three laboratories (Berlin, Tübingen, and Warsaw) as described in Supplementary material 3. Segregation analysis was performed by Sanger sequencing for the other available family members. The pathogenicity of variants obtained from NGS was assessed by molecular modeling (Yasara Structure) as described in Supplementary material 3.

## RESULTS

We report on 13 affected males and 2 affected females from 6 families with variants in the MED12 gene (Fig. 1). Clinical symptoms of the affected males and females along with a comparison with symptoms described in published MED12-related syndromes are presented
in Table 1. All probands had ID and developmental delay, most of them had macrocephaly, long narrow face, prominent forehead, small ears, dental abnormalities, prognathism and hypertelorism. The remaining clinical features were variable. The new clinical findings associated with MED12 variants (present in at least 2 families) were as follows: brachycephaly, anteverted nares, bulbous nasal tip, prognathism, deep set eyes, and single palmar crease. In the families we identified 3 novel missense variants in MED12 (p.(Val954Gly), p.(Glu1091Lys), p.(Pro1371Ser)), 1 known variant but firstly reported in affected females (p.(Arg1148His)), as well as 2 variants (p.(Arg815Gln) and p.(Arg1295Cys)) which have already been described as a cohort data by the authors of the current paper, but without clinical details of the presented families ${ }^{3,4}$.

All MED12 variants identified in the families perfectly co-segregated with the phenotype, except for Family 5 in whom the MED12 variant present in the affected daughters was inherited from the healthy mother. Most of the variants were absent in the gnomAD database (except for $\mathrm{c} .2444 \mathrm{G}>\mathrm{A}$ present in 1 heterozygous female with a minor allele frequency (MAF) of $5.599 \mathrm{e}-6$ and $\mathrm{c} .4111 \mathrm{C}>\mathrm{T}$ present in 6 hemizygous individuals with a MAF of 0.00012 ). All the variants were predicted as damaging by Polyphen2, MutationTaster, as well as SIFT, except for $\mathrm{c} .4111 \mathrm{C}>\mathrm{T}$ identified in patient III: 1 from Family 6 which had conflicting interpretations of pathogenicity (Polyphen 2 - benign, Mutation Taster - disease causing, SIFT - tolerated). Additionally, this variant had been reported to ClinVar as likely benign. NGS performed subsequently in the patient revealed additionally a de novo heterozygous missense variant $\mathrm{c} .367 \mathrm{C}>\mathrm{T}$ in the PUF60 gene
(NM_078480.2) resulting in the p.(Arg123Trp) substitution. The variant was predicted as damaging (Polyphen2, MutationTaster, SIFT) and it was absent in gnomAD database.

Molecular modeling was applied to assess the functional effects of the MED12 (four of the six identified MED12 substitutions are located within the protein regions, the structure of which may be reasonably modeled by homology) and PUF60 variants identified by NGS. All the variants are expected to have substantial structural effects on proteins (Fig. 2).

X chromosome inactivation (XCI) analysis performed in Family 4 revealed that the mother (II:1) of the index patient (III:2) who also carried the mutation had mildly skewed XCI (83:17) in her blood cells. In Family 5 XCI analysis revealed a mildly skewed pattern in the mother (82:18) and a skewed pattern in her two affected daughters (100:0 and 85:15). In the three females, the same allele of the AR locus was inactive.

## DISCUSSION

The phenotypic spectrum of MED12-related disorders is still being expanded as new families with MED12 variants are being identified due to the improvement of new sequencing technologies. Moreover, it was postulated that succeeding syndromes should be defined to extend the list of the three allelic syndromes associated with MED12 mutations. However, as shown by the clinical data of our probands (Table 1) it seems that they do not manifest strict syndromes (FGS1, Lujan-Fryns or Ohdo), but they rather present, similar to recently published data ${ }^{5}$, a continuum of symptoms. Thus, it is more appropriate to indicate only a 'MED12-related disorder' than to attribute a definite syndrome for a majority of the subjects.

To date, over 20 MED 12 genetic variants have been reported which are evenly distributed with no apparent hot spots in specific exons (Fig. 1). All but one were missense variants causing FGS1, Lujan-Fryns, Ohdo syndrome or non-syndromic ID with different degrees of cognitive impairment. Many benign variants (456 in ClinVar) were also described as well as somatic variants implicated in cancer but not associated with germline mutations. The MED12 variants examined in the current study were defined as pathogenic by molecular modeling, the approach that is not only based on bioinformatic prediction, but also uses experimentally gained protein crystal structures. The results suggested a pathogenic effect of all four MED12 variants tested: p.(Va1954Gly), p.(Glu1091Lys), p.(Arg1295Cys), and p.(Pro1371Ser). The other two variants (p.(Arg815Gln) and p.(Arg1148His)) were not covered by the models obtained. Despite these results, the maternally inherited MED12 p.(Pro1371Ser) substitution identified in the sporadic male (Family 6) and also present in hemizygous individuals in gnomAD was found to be problematic in the case of the MED12 variant's pathogenicity. Because the MED12 variant was found in the proband by Xchromosome exome re-sequencing, we performed subsequent WES in order to exclude the possibility of other variants being engaged in ID pathogenesis. The WES analysis revealed a
de novo p.(Arg123Trp) variant in the PUF60 gene. PUF60 variants cause autosomal dominant Verheij syndrome which is clinically highly variable. Main clinical symptoms include ID, short stature, cerebral atrophy, microcephaly, as well as renal, skeletal, and cardiac defects ${ }^{6}$. With the current knowledge, the de novo PUF60 variant identified in this study only partially explains the phenotype of the proband from Family 6. The additional clinical findings such as macrocephaly, ptosis, Hirschsprung disease, genital hypoplasia, and anal anomaly fit in the clinical spectrum of MED12-related disorders and to the best of our knowledge have not been reported in probands with PUF60 variants so far. Both, the MED12 and PUF60 variants were predicted to be pathogenic by molecular modeling and combined they better explain the phenotype than each individually. Thus, it is possible that the proband from Family 6 presents a complex phenotype caused by these two variants. Such digenic inheritance has recently been postulated as an important cause of ID.

Currently, reports on MED12-related clinical phenotypes in females are rather scarce. With the current knowledge, pathogenicity of the firstly reported MED12 p.(Arg1148His) substitution in affected females (Family 5) and assumed as causing Ohdo syndrome in males only, by clinical, genetic and functional findings remains to be established ${ }^{7}$. Segregation analysis revealed that the variant was inherited from their healthy mother. Although MED12 is subject to X -inactivation, there was no clear correlation between clinical phenotypes of the MED12 carrier females from Family 5 and their XCI pattern in blood cells, similar to the results published previously ${ }^{8}$. It seems that blood DNA-based X inactivation does not predict clinical outcome in MED12-related carrier females and different affection statuses in females might be related to varying X-inactivation in the brain. Thus, it is possible that similar to other genes initially implicated in X-linked recessive disorders in males, missense variants in MED12 may predict a potential risk for phenotypic expression in carrier females. This
assumption needs to be more firmly established in the future, and if confirmed would be important for genetic counseling and prognostic evaluation.

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## FIGURE LEGENDS

Fig. 1
A Pedigrees of the families with segregation of the MED12 variants (black) and the PUF60 variant (blue). Mutant alleles are represented by a plus (+) and wild-type alleles by a minus ( ). $\mathbf{B}$ The distribution of the MED12 variants in relation to the gene structure consisting of 45 exons and three functional domains: MED12, LCEWAV, and catenin-binding domain - PQL (domain positioning according to Pfam 31.0 database). Known pathogenic MED12 variants are indicated above the gene and novel variants below the gene. Asterisks denote MED12 variants present in affected females. C Images of MED12 probands from the presented families. Most of the probands presented with a long narrow face, prominent forehead, prominent nasal bridge, small ears, and hypertelorism. Novel facial dysmorphisms that expand the phenotypic spectrum of MED12-related disorders include deep set eyes (Families 1, 5), prognathism (Families 1, 3, 5, 6), anteverted nares (Families 4, 6), and bulbous nasal tip (Families 3, 5, 6).


## Fig. 2

A Representative models (Yasara Structure) of the MED12 segments 826Ala-1096Leu and 1278Tyr-1395Glu which harbor variants identified in the families, obtained based on the structures of homologous proteins. Glu1091 (E1091) forms a salt bridge with the proximal Lys1086, so the missense variants p.(Glu1091Lys) (E1091K), which replaces this interaction by an unfavorable Lys-Lys one (cation-cation), decreases protein stability. The amino acid substitution Val954Gly (V954G) decreases the protein stability distorting the hydrophobic core of the four helix bundle (blue). Arg 1295 (R1295) forms a salt bridge with Glu1302 (E1302) and/or Asp1303 (D1303) and displays pi-pi interactions with Trp1291 (W1291), thus stabilizing the relative orientation of two proximal helices. In this view, the p.(Arg 1295Cys) substitution will destroy all these interactions. It should be noted that three cysteine residues are proximal to $\operatorname{Arg} 1295$ (Cys1287, Cys1296 and Cys1301), so the Arg1295Cys replacement, which introduces another cysteine residue, enables formation of alternative patterns of CysCys bridges, all of which will interfere with the native one, thus affecting the protein folding. Pro1371Ser (P1371S) will increase the local flexibility of the short loop, thereby enabling reorientation of the flanking helices, and thus decreasing protein stability.


B The analysis (Yasara Structure) of the PUF60 p.(Arg 123Trp/Gly) amino acid substitutions. Arg123 (R123) is located just before the RRM1/RRM2 motifs of PUF60, in the proximity of the linker ${ }^{205}$ ProSerAsnIleGlyGln ${ }^{210}$ (magenta) that separates RRM domains (left). Arg123Gly (R123G) replacement only minutely affects the protein structure (middle). The change of the polar arginine to the highly hydrophobic tryptophan restricts the conformational flexibility of this linker (right), and the relative orientation of both RRM domains changes substantially upon RNA binding, thus most likely affecting the affinity of PUF60 to RNA.



Table 1. Phenotypic comparison between known MED12 related syndromes and the families presented in the study. Clinical symptoms specific for a given syndrome are coloured appropriately (FG, red; L-F, green; Ohdo, blue). Symptoms not described previously that expand the phenotypic spectrum of the MED12 variants are marked in bold. " + " in family columns denotes that at least one patient per family was diagnosed with the given feature, "blank" denotes that none of the family members presented the given feature.


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