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## ATP7B variant spectrum in a French pediatric Wilson disease cohort<sup>☆</sup>

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### ABSTRACT

**Background/aim:** The spectrum of ATP7B variants varies significantly according to geographic distribution, and there is insufficient data on the variants observed in the French population.

**Methods:** Clinical data of 113 children included in the French WD national registry were gathered from March 01, 1995 to July 01, 2020. Data included epidemiological, clinical, laboratory, genetics.

**Results:** Diagnosis was made at a mean age of 11.0 ± 4.1 years (range 1–18 years). At diagnosis, 91 patients (79.8 %) had hepatic manifestations, 18 (15.8 %) presented neurological manifestations, and 4 patients (3.5 %) were asymptomatic. Only 29 patients (25 %) were homozygous for a variant. We have found a total of 102 different variants including 14 novel variants. Recurrent variant p.His1069Gln was the most prevalent, n = 31 alleles (14,2%), with only seven homozygous; in contrast 55% of variants are identified in only one family. 45% were truncating variants. In respect of mutated exon, the three most prevalent were exon 14 (16.5%), exon 8 (13.8%), and exon 3 (11.5%). When considering patients with two Nonsense / Frameshift variants as a group and those with two Missense variants, we found significantly lower ceruloplasmin for the former: 2.8 ± 0.7 mg/dl vs 8.4 ± 5mg/dl (p<0.05).

**Conclusion:** p.His1069Gln is the most frequent variant (14,2%) and exons 14, 8, and 2 of the ATP7B gene account for 41.7% of total variants. However, there is significant heterogeneity in the French population concerning the other ATP7B variants. Nonsense / Frameshift variants were associated with lower ceruloplasmin levels.

**Abbreviations:** WD, Wilson disease; ALF, acute liver failure; UCE, urinary copper excretion; H, hepatic manifestations; N, neurologic manifestations; AS, asymptomatic; KF ring, Kayser-Fleischer ring; MRI, magnetic resonance imaging.

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## Authors' contribution

EC wrote the manuscript and performed statistical calculations, SB, TS, CP, and MB helped in writing the manuscript, OG performed statistical calculations and revised the manuscript, DH, DD, TL, PB, AF, CV, RS, FG, LBH, AP: contributed to data and revision of the manuscript. AB: collected the data, ASB and AL conceived and developed the project. MB: manuscript revision and approval.

## 1. Introduction

Wilson's disease (WD) is a rare autosomal recessive genetic disorder associated with *ATP7B* gene variants and causing an impaired biliary copper elimination and consequently a copper accumulation, mainly in the liver and the brain. Clinical expressions are variable; the hepatic manifestations prevail during childhood, and the first neurological signs classically appear in the second and third decades of life as the copper accumulation progresses (Schilsky, 2017; Socha et al., 2018). The prevalence of WD is considered as 1:30,000, although the existing data on WD epidemiology and genetics are somewhat contradictory (O'Brien et al., 2014; Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020). According to the recent data from the European population, WD's prevalence may account for 1:10,000 or 1:7,000, being much more frequent than the earlier estimations (Coffey et al., 2013). Such a contradiction may occur due to poor diagnosis of WD, the presence of latent forms, various methodological issues and presence of consanguinity which increases disease frequency.

The diagnosis of WD is based on a pattern of clinical signs, laboratory results, and genetic analysis (Socha et al., 2018; Ferenci et al., 2003). To establish a WD diagnosis can be particularly challenging in paediatric cases since young children often do not display clinical and laboratory hallmarks (elevated urinary copper excretion, Kayser-Fleischer ring, abnormal brain MRI) (Socha et al., 2018), thus enhancing the importance of concomitant genetic analysis.

To date, more than 900 WD-causing variants in the *ATP7B* gene have been identified in WD patients, and most of them have been identified in compound heterozygote patients. Despite considerable research efforts, this broad spectrum of phenotypic and genotypic WD heterogeneities has hampered the study of phenotype-genotype correlation.

The spectrum of *ATP7B* variants varies significantly between different geographical populations. p.His1069Gln is considered the most frequent variant in the European population whereas p.Arg778Leu and c.del441-427 are the most common in the Asian population and Sardinia respectively (GGV, 2016). In France, data on the observed variants are insufficient, with only one previous publication in 2012 reporting the distribution of *ATP7B* variants in French WD patients (Bost et al., 2012).

Therefore, in our study, we aim to provide a detailed analysis of the *ATP7B* variants from a paediatric cohort of the French WD registry and their possible correlation with clinical and laboratory data.

## 2. Patients and methods

### 2.1. Subjects and sample collection

Clinical data of children included in the French WD national registry were gathered from March 01, 1995 to July 01, 2020 (retrospective collection between 1995 and 2005 and prospective collection since 2006).

The patients were followed by paediatricians, hepatologists and neurologists in the two reference centers (Paris and Lyon) and the eight centers of competence.

The diagnosis of WD was based on clinical manifestations, biochemical parameters, and/or genetic analysis as previously

published (Schilsky, 2017; Socha et al., 2018; Ferenci, 2017; European Association for, 2012). Collected data included: clinical data, presence of a Kayser-Fleischer ring (KF ring), liver function tests, immuno-nephelometric measurement of serum ceruloplasmin levels, serum copper level, determination of 24-h urinary copper excretion (UCE), genetics. Patients without genetic information were excluded. In family cases, only the index case was included. All patients included in the registry had a clinical score  $\geq 4$  using the score proposed by Ferenci et al. (2003).

Depending on the primary clinical manifestations, cases were classified as hepatic (H), neurological (N), or asymptomatic (AS); we considered as asymptomatic only those with normal hepatic function tests and no neurological symptom.

For the genotype assessment, we hypothesized that the following variables would be associated with variant type: age at onset of first symptoms, neurological *versus* hepatic form of the illness, ceruloplasmin level, serum copper values, and 24h urine copper level, and the presence of KF rings. Many comparisons were made: first for the homozygous for the most common *ATP7B* variants ( $\geq 2$ ), then between groups composed of homozygous and compound heterozygous for the most common variants. We also compared patients with homozygous missense variants (MSM) *versus* patients with frameshift-nonsense variants.

### 2.2. Genetic analysis of *ATP7B* gene

Genetic analyses after 1994 were performed for patients and their families by bi-directional sequencing on the 21 exons and intron-exon boundary regions of *ATP7B* gene. For patients with only one *ATP7B* variant detected by sequencing, a Multiplex Ligation-dependent Probe Amplification (MLPA) assay was performed using the SALSA MLPA P098 for Wilson Disease kit (MRC-Holland, Amsterdam, Netherlands) to detect large deletions. Since 2017, next-generation sequencing (NGS) was used for *ATP7B* variant detection in WD patients with a bioinformatic pipeline allowing the detection of SNV and copy number variation.

### 2.3. Ethics and regulatory aspects

The study received approval from the INSERM (Institut National de la Santé et de la Recherche Médicale) Ethics evaluation committee (IRB 00003888, authorization No.19-550/January 24, 2019). Written informed consent was obtained from parents or adult patients.

This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

### 2.4. Statistical analysis

Normally distributed quantitative variables were expressed as mean  $\pm$  SD; non-normally distributed quantitative variables were expressed as median [IQR], and were compared using the Student t-test or non-parametric tests (Mann-Whitney, Kruskal-Wallis) when appropriate. Qualitative variables were expressed as count (percentage), and compared using the  $\chi^2$  test or Fischer's exact test when appropriate. Statistical analysis was performed with SPSS version 23.0 (IBM, New York, USA); p-values lower than 0.05 were considered statistically significant.

## 3. Results

### 3.1. Study population

A total of 113 children (aged 0–18 years) were included in the register analysis. Demographic and laboratory parameters are summarized in Table 1.

Gender was quite equally distributed M/F = 56/57. Diagnosis was made at a mean age of  $11.0 \pm 4.1$  years (range 1–18 years). At diagnosis, 91 patients (79.8 %) had hepatic manifestations (H), 18 (15.8 %) presented neurological manifestations (N), and 4 patients (3.5 %) were asymptomatic (AS).

Mean age at diagnosis was  $10.4 \pm 4.0$  years (range 2–18) for H patients,  $14.6 \pm 1.9$  years (range 11–18) for N patients ( $p < 0.001$ ), and  $8.3 \pm 6.0$  years (range 1–17) for AS patients.

### 3.2. Biochemical results at diagnosis

The mean serum ceruloplasmin level was  $7.8 \pm 5.5$  mg/dL. A total of 8 patients (5.5 %) had serum ceruloplasmin levels  $\geq 20$  mg/dL.

At diagnosis, the median urinary copper excretion (UCE) was 4.1  $\mu\text{mol}/24$  h (IRQ 5.2).

The median urinary copper excretion was significantly lower for AS patients 1.3  $\mu\text{mol}/24$  h (IRQ 2.1) compared to H patients 3.8  $\mu\text{mol}/24$  h (IRQ 5.6) and to N patients 5.0  $\mu\text{mol}/24$  h (IRQ 2.5),  $p < 0.001$ .

### 3.3. Variant analysis

*ATP7B* gene analysis was performed in 113 WD patients. Two variants within the *ATP7B* gene were identified in 98 patients, just one variant in 13, and 2 patients had three variants (patient 1: variants p.Asn41Ser and variant p.Ile1021Val in *cis* and variant p.Arg1319\* in *trans*; patient 2: variants p.Lys175Serfs\*28 and variant p.Gln260Profs\*10 in *cis* and variant p.Gly1266Arg in *trans*). MLPA allowed to detect large exon deletions in 4 patients. Only 29 patients (25 %) were homozygous for a variant. We have found a total of 102 different variants; details about variants are listed in Table 2. p.His1069Gln was the most prevalent,  $n = 31$  alleles (14.2 %), with only seven homozygous. The frequency of the three other most prevalent variants were: p.Gln111\* (3.7 %), c.1708-1G > A, p.Arg1319\*, p.Met769Hisfs\*26 and p.Val890Met (2.3 % each). The allele frequency of other variants was  $\leq 2$  %; 55 % of variants were present in only one family, confirming genetic heterogeneity of WD in France. 14 novel variants were identified and submitted to Clinvar. Data used to classified novel missense variants and variants previously classified as variants of unknown significance are detailed in Table 3.

In respect of mutated exon, the 3 most prevalent were exon 14 (16.5 %), exon 8 (13.8 %), and exon 3 (11.5 %). Very few variants were identified in exons 7, 3, 12, 9 and 21. 12.4 % of variants were located in intron-exon boundary regions of *ATP7B* gene; introns 4, 5, 8, and 12 were the most frequently involved (Fig. 1). This point is important according epidemiological level.

Concerning the type of variant, the repartition was as follow ( $n$

alleles - %): Missense (127–58.2 %), Splice site (30–13.8 %), Frameshift (30–13.8 %), Nonsense (24–11 %), Deletion (7–3.2 %). Overall, 45 % of variants identified were truncating variants.

Variant distribution in the *ATP7B* protein domains was as follow: ATP binding domain (29.4 %), transmembrane domain (23.9 %), copper-binding domain (22.9 %), phosphatase domain, and phosphorylation domain (4.1 % and 1.8 %, respectively), combined domains accounted for the rest of variants (17.9 %).

### 3.4. Genotype-phenotype

Missense variants (p.His1069Gln, p.Val890Met, p.Lys1020Arg) – Splice site (c.1708-1G > A).

The 7 homozygous for p.His1069Gln were compared with 3 other homozygous variants with a frequency of at least 2 patients (p.Val890Met, p.Lys1020Ar, c.1708-1G > A). We didn't observe any statistically significant difference for clinical phenotype, age at diagnosis, presence of KF ring, ceruloplasmin levels, serum copper values and 24h urine copper excretion. When adding compound heterozygous for these variants, only p.His1069Gln was enlarged with 13 patients and only one for c.1708-1G > A. Comparison between p.His1069Gln homozygous with compound heterozygous did not find significant statistical differences for any of the variables previously mentioned. Details are shown in Table 4.

### 3.5. Nonsense/frameshift variants

For Nonsense/Frameshift variants, there were only compound heterozygotes and no homozygous. When considering patients with two Nonsense/Frameshift variants as a group and compared with those with two Missense variants, we found significant lower ceruloplasmin for the former:  $2.8 \pm 0.7$  mg/dl vs  $8.4 \pm 5$  mg/dl ( $p < 0.05$ ). No differences were found in age at diagnosis, serum copper values or 24h urine copper excretion. Details are shown in Table 4.

**Acute liver failure:** 8 patients had acute liver failure at diagnosis of which only 3 were homozygous, each one of a different type hence no comparisons were made.

## 4. Discussion

The p.His1069Gln variant was the most frequent variant in our cohort and the prevalence was very similar to the reported by Bost et al., (2012) (Bost et al., 2012), this prevalence is one of the lowest ones reported in Central and East Europe where ranges of prevalence are between 15 % and 72 % (GGV, 2016).

The distribution of mutated exons could give the spectrum of *ATP7B*

**Table 1**

Characteristics of the study population ( $n = 113$ ).

	Asymptomatic	Hepatic	Neurologic	All
Total patients n (%)	4 (3.5 %)	91 (79.8 %)	18 (15.8 %)	113
Male/Female	2/2	45/46	9/9	56/57
Age at diagnosis (mean $\pm$ sd)	$8.3 \pm 6.0$	$10.4 \pm 4.0$	$14.6 \pm 1.9$	$11.0 \pm 4.1$
Kayser-Fleischer ring (n,%)	0	13 (14.2 %)	15 (83.3 %)	28 (24.8 %)
Serum ceruloplasmin mg/dl (mean $\pm$ sd)	$8.0 \pm 5.5$	$8.5 \pm 5.6$	$4.6 \pm 2.7$	$7.8 \pm 5.5$
Urinary copper excretion, $\mu\text{mol}/24$ h - median (IQR)	1.3 (2.1)	3.8 (5.6)	5.0 (2.5)	4.1 (5.2)
Serum copper $\mu\text{mol}/L$ - median (IQR)	3.94 (3.31)	4.88 (6.14)	6.30 (2.52)	5.04 (5.2)

Reference ranges: ceruloplasmin (20–35 mg/dL) - serum copper (12.6–31.5  $\mu\text{mol}/L$ ) - urinary copper excretion <0.6  $\mu\text{mol}/24\text{h}$ .

**Table 2**

Variants detected in children with Wilson disease in this study.

Protein change	cDNA change	ClinVar accession	Variant classification (source)	Exon or Intron	Variant type	Protein domain	Variant allele frequency (%)	Age of onset mean (range)	Clinical form (n)
p.His1069Gln	c.3207C > A	VCV000003848	5 (CV, HGMD)	E 14	MS	Ph	14,22 %	12,3 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019; Stapelbroek et al., 2004)	H = 21 N = 9 AS = 1
p.Gln111*	c.331C > T	VCV000188899	5 (CV, HGMD)	E 2	NS	Cu1	3,67 %	12,2 (Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019)	H = 8
	c.1708-1G > A	VCV000370195	5 (CV, HGMD)	I 4	Spl	Cu6	2,29 %	13,8 (GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019)	H = 5
p.Met769HisFs*26	c.2304dupC	VCV000456552	5 (CV, HGMD)	E 8	FS	TM4	2,29 %	10,4 (Schilsky, 2017; Socha et al., 2018; O'Brien et al., 2014; Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010)	H = 3 AS = 2
p.Val890Met	c.2668G > A	VCV000189121	5 (CV, HGMD)	E 11	MS	Td/TM5	2,29 %	8,2 (Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004)	H = 5
p.Arg1319*	c.3955C > T	VCV000035728	5 (CV, HGMD)	E 19	NS	TM7	2,29 %	6,4 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017)	H = 5
p.Arg226Trp	c.676C > T	VCV000312397	4 (This study)	E 2	MS	Cu2/ Cu3	1,83 %	10,7 (Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012)	H = 2 N = 2
p.Met645Arg	c.1934T > G	VCV000003862	5 (CV, HGMD)	E 6	MS	Cu6/ TM1	1,83 %	9 (Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004)	H = 4

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Table 2 (continued)

Protein change	cDNA change	ClinVar accession	Variant classification (source)	Exon or Intron	Variant type	Protein domain	Variant allele frequency (%)	Age of onset mean (range)	Clinical form (n)
p.Arg778Trp	c.2332C > G	VCV000456553	5 (CV, HGMD)	E 8	MS	TM4	1,83 %	7,75 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017)	H = 4
	c.2865+1G > A	VCV000157942	5 (CV, HGMD)	I 12	Spl	TM5/ TM6	1,83 %	12,75 (Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010)	H = 3 N = 1
p.Lys1020Arg	c.3059A > G	pending	5 (HGMD)	E 13	MS	Ph/ATP Loop	1,83 %	11 (Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005)	H = 2 N = 2
p.Asn1270Ser	c.3809A > G	VCV000003859	5 (CV, HGMD)	E 18	MS	ATP Hinge	1,83 %	9,5 (O'Brien et al., 2014; Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019; Stapelbroek et al., 2004)	H = 3 N = 1
p.Ser1365CysFs*12	c.4092_4093del	VCV000371438	5 (CV, HGMD)	E 20	FS	TM8	1,83 %	12 (Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005)	H = 2 N = 2
	c.51+4A > T	VCV000312401	5 (CV, HGMD)	I 1	Spl	before Cu1	1,38 %	10 (Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911)	H = 3
	c.1869+2T > C	pending	5 (HGMD)	I 5	Spl	Cu6	1,38 %	12,33 (Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010)	H = 2 AS = 1
p.Asp765Asn	c.2293G > A	VCV000003855	5 (CV, HGMD)	E 8	MS	TM4	1,38 %	8 (Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016)	H = 3
p.Val845SerFs*28	c.2532delA	VCV000188883	5 (CV, HGMD)	E 10	FS	Td	1,38 %	10 (Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911)	H = 2 N = 1
p.Thr977Met	c.2930C > T	VCV000035710	5 (CV)	E 13	MS	Ch/TM6	1,38 %	8,66 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017)	H = 3

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Table 2 (continued)

Protein change	cDNA change	ClinVar accession	Variant classification (source)	Exon or Intron	Variant type	Protein domain	Variant allele frequency (%)	Age of onset mean (range)	Clinical form (n)
p.Ile1148Thr	c.3443T > C	VCV000037122	5 (CV, HGMD)	E 16	MS	ATP Loop	1,38 %	10,3 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005)	H = 2 N = 1
p.Asp1267Ala	c.3800A > C	VCV000552417	5 (CV, HGMD)	E 18	MS	ATP Hinge	1,38 %	12,6 (Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019)	H = 3
	del_E4	NA	5 (This study)	E 4	Del	Cu5/ Cu6	1,38 %	11,6 (Ferenci, 2017; European Association for, 2012)	H = 3
p.Leu168Pro p.Gln260ProFs*10	c.503T > C	VCV000967323	5 (CV, HGMD)	E 2	MS	Cu2	0,92 %	3	H = 2
	c.778dupC	VCV000188938	5 (CV, HGMD)	E 2	FS	Cu2/ Cu3	0,92 %	8 (Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016)	H = 2
p.Gln289*	c.865C > T	VCV000003864	5 (CV, HGMD)	E 2	NS	Cu3	0,92 %	14 (European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005)	N = 2
p.Ala553Glu p.Thr569del	c.1658C > A	pending	5 (HGMD)	E 4	MS	Cu5/6	0,92 %	9	H = 2
	c.1705_1707 + 10del	pending	5 (HGMD)	E 4	Del	Cu5/6	0,92 %	17	N = 2
p.Gly710Ser	c.2128G > A	VCV000156281	5 (CV, HGMD)	E 8	MS	TM2	0,92 %	11 (GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004)	H = 2
p.Gly711Arg	c.2131G > A	VCV000495405	5 (CV, HGMD)	E 8	MS	TM2	0,92 %	10 (Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004)	H = 2
p.Thr766Arg	c.2297C > G	VCV000003861	5 (CV, HGMD)	E 8	MS	TM4	0,92 %	5,5 (O'Brien et al., 2014; Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003)	H = 2
p.Met769Val	c.2305A > G	VCV000035706	5 (CV)	E 8	MS	TM4	0,92 %	10 (Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012)	H = 2
p.Arg778Gly	c.2332C > G	VCV000156283	5 (CV, HGMD)	E 8	MS	TM4	0,92 %	14,5 (European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019)	H = 1 N = 1
p.Trp779*	c.2336G > A	VCV000156284	5 (CV, HGMD)	E 8	NS	TM4	0,92 %	8 (Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012)	H = 2
p.Gly836Glu p.Ile857Thr p.Asp918Asn	c.2356-2A > G	pending	5 (HGMD)	I 9	Spl	TM4/Td	0,92 %	11	N = 2
	c.2507G > A	pending	5 (HGMD)	E 10	MS	Td	0,92 %	8	H = 2
	c.2570T > C	VCV000377538	5 (CV, HGMD)	E 10	MS	Td	0,92 %	6	H = 2
	c.2752G > A	VCV000552229	5 (HGMD)	E 12	MS	Td/TM5	0,92 %	15	H = 2

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Table 2 (continued)

Protein change	cDNA change	ClinVar accession	Variant classification (source)	Exon or Intron	Variant type	Protein domain	Variant allele frequency (%)	Age of onset mean (range)	Clinical form (n)
p.Ala1018Val	c.3053C > T	VCV000188722	5 (CV, HGMD)	E 13	MS	Ph/ATP Loop	0,92 %	5 (Schilsky, 2017; Socha et al., 2018; O'Brien et al., 2014; Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016)	H = 1 AS = 1
p.Lys1028SerFs*40	c.3083_3085delinsG	VCV000633073	5 (CV, HGMD)	E 14	FS	Ph	0,92 %	12,5 (Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010)	H = 2
p.Ser1067Arg <sup>a</sup>	c.3201T > G <sup>a</sup>	pending	4 (This study)	E 14	MS	ATP Loop	0,46 %	18	H = 1
p.Ala1135GlnFs*13	c.3402delC	VCV000088958	5 (CV, HGMD)	E 15	FS	ATP Loop	0,92 %	8 (Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012)	H = 2
p.Ala1183Thr	c.3547G > A	pending	5 (HGMD)	E 16	MS	ATP Loop	0,92 %	13 (Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005)	H = 2
p.Gly1186Ser	c.3556G > A	VCV000188900	5 (CV, HGMD)	E 16	MS	TM6	0,92 %	13,5 (GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004; N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005; Kluska et al., 2019; Stapelbroek et al., 2004)	H = 2
p.Gly1266Arg	c.3796G > A	VCV000003849	5 (CV, HGMD)	E 18	MS	ATP Hinge	0,92 %	9	H = 2
	c.3904-2A > G	VCV000371387	5 (CV, HGMD)	I 19	Spl	ATP Loop	0,92 %	3	H = 2
	c.4125-1G > T <sup>a</sup>	pending	5 (This study)	I 20	Spl	TM8	0,92 %	8,5 (Collet et al., 2018; Poujois et al., 2018; Sandahl et al., 2020; Coffey et al., 2013; Ferenci et al., 2003; GGv, 2016; Bost et al., 2012; Ferenci, 2017; European Association for, 2012; Panagiotakaki et al., 2004)	H = 2
p.Glu1382* <sup>a</sup>	c.4144G > T <sup>a</sup>	pending	5 (This study)	E 21	NS	after TM8	0,92 %	15 (Merle et al., 2010)	N = 2
p.Asn41MetFs*26 <sup>a</sup>	c.122delA <sup>a</sup>	pending	5 (This study)	E 2	FS	Cu1	0,46 %	10	H = 1
p.Asn41Ser	c.122A > G	VCV000157928	5 (CV, HGMD)	E 2	MS	Cu1	0,46 %	3	AS = 1
p.Ser105*	c.314C > A	VCV000189193	5 (CV, HGMD)	E 2	NS	Cu1	0,46 %	14	H = 1
p.Glu127LysFs*26	c.379del	pending	5 (HGMD)	E 2	FS	Cu1	0,46 %	18	H = 1
p.Lys175SerFs*28	c.524_525delAA	VCV000188995	5 (CV, HGMD)	E 2	FS	Cu2	0,46 %	9	H = 1
p.Tyr187SerFs*16 <sup>a</sup>	c.560_561del <sup>a</sup>	pending	5 (This study)	E 2	FS	Cu2/3	0,46 %	11	H = 1
p.Gly379AlaFs*2	c.1136delG	pending	5 (HGMD)	E 2	FS	Cu4	0,46 %	8	H = 1
p.Arg414SerFs*9 <sup>a</sup>	c.1242_1243del <sup>a</sup>	pending	5 (This study)	E 2	FS	Cu4	0,46 %	12	H = 1
	c.1286-2A > G	pending	5 (HGMD)	I 2	Spl	Cu4	0,46 %	10	H = 1
	c.1369C > T	pending	5 (HGMD)	E 3	NS	Cu4/5	0,46 %	10	H = 1
p.Gln457*	c.1512dupT	VCV000370348	5 (CV, HGMD)	E 3	NS	Cu5	0,46 %	7	H = 1
p.Asn505*	c.1543+1G > T	VCV000590806	5 (CV, HGMD)	I 3	Spl	Cu5	0,46 %	14	H = 1
	c.1708-34G > A <sup>a</sup>	pending	4 (This study)	I 5	Spl	Cu6	0,46 %	14	H = 1
	c.1740del <sup>a</sup>	pending	5 (This study)	E 5	FS	Cu6	0,46 %	7	H = 1
p.His580GlnFs*3 <sup>a</sup>	c.1745_1746del	VCV000189112	5 (CV, HGMD)	E 5	MS	Cu6	0,46 %	8	H = 1
p.Ile582ArgFs*25	c.1772G > A	VCV000210482	5 (CV, HGMD)	E 5	MS	Cu6	0,46 %	4	H = 1
p.Gly591Asp	c.1811C > A	pending	4 (This study)	E 5	MS	Cu6	0,46 %	12	H = 1
	c.1869 +	pending	5 (HGMD)	I 6	Spl	Cu6	0,46 %	7	H = 1
	5_1869+8del								
p.Gly626Ala	c.1877G > C	VCV000157930	5 (HGMD)	E 6	MS	Cu6/TM1	0,46 %	18	H = 1
p.Leu641Ser	c.1922T > C	VCV000420002	5 (HGMD)	E 6	MS	Cu6/TM1	0,46 %	16	H = 1

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Table 2 (continued)

Protein change	cDNA change	ClinVar accession	Variant classification (source)	Exon or Intron	Variant type	Protein domain	Variant allele frequency (%)	Age of onset mean (range)	Clinical form (n)
p.Asp642His	c.1924G > C	VCV000189109	5 (CV, HGMD)	E 6	MS	Cu6/TM1	0,46 %	7	H = 1
	c.1946+6T > C	VCV000495403	5 (HGMD)	I 6	Spl	Cu6/TM1	0,46 %	14	N = 1
p.Gly691Arg	c.2071G > A	VCV000003866	5 (CV, HGMD)	E 7	MS	TM2	0,46 %	18	H = 1
	c.2122-8T > G	VCV000035705	5 (CV, HGMD)	I 8	Spl	TM2	0,46 %	15	N = 1
p.Leu708Pro	c.2123T > C	VCV000003865	5 (CV, HGMD)	E 8	MS	TM2	0,46 %	6	AS = 1
p.Pro768Leu	c.2303C > T	VCV000370887	4 (CV)	E 8	MS	TM4	0,46 %	14	N = 1
p.Met769ThrFs*26	c.2304dup	VCV000456552	5 (CV, HGMD)	E 8	FS	TM4	0,46 %	15	N = 1
p.Arg778Gln	c.2333G > A	VCV000550914	5 (CV, HGMD)	E 8	MS	TM4	0,46 %	16	H = 1
p.Trp779Cys	c.2337G > C	pending	4 (This study)	E 8	MS	TM4	0,46 %	5	H = 1
p.Leu795Phe	c.2383C > T	VCV000188814	5 (CV, HGMD)	E 9	MS	TM4/Td	0,46 %	7	H = 1
p.Gly836Arg	c.2506G > A	pending	5 (HGMD)	E 10	MS	Td	0,46 %	4	H = 1
p.Lys838Glu	c.2512A > G	VCV000860738	4 (This study)	E 10	MS	Td	0,46 %	14	H = 1
p.Gly881*	c.2637_2650del	pending	5 (This study)	E 11	NS	Td/TM5	0,46 %	15	N = 1
p.Thr991Ala	c.2971A > G	pending	4 (This study)	E 13	MS	TM6/Ph	0,46 %	12	H = 1
p.Ala1003Pro	c.3007G > C	pending	4 (This study)	E 13	MS	TM6/Ph	0,46 %	16	H = 1
p.Ala1003Val	c.3008C > T	VCV000188781	5 (CV, HGMD)	E 13	MS	TM6/Ph	0,46 %	10	H = 1
p.Ile1021Val	c.3061-12T > A	VCV000557116	5 (CV, HGMD)	I 13	Spl	Ph	0,46 %	16	H = 1
	c.3061A > G	VCV000430276	5 (HGMD)	E 14	MS	Ph	0,46 %	3	AS = 1
p.Met1025Lys	c.3074T > A	pending	5 (HGMD)	E 14	MS	Ph	0,46 %	10	H = 1
p.Arg1041Trp	c.3121C > T	VCV000312384	5 (CV, HGMD)	E 14	MS	ATP	0,46 %	8	H = 1
p.Gly1099Ser	c.3295G > A	VCV000370820	5 (CV, HGMD)	E 15	MS	Loop	0,46 %	6	AS = 1
						ATP			
p.Ile1102Thr	c.3305T > C	VCV000430725	5 (CV, HGMD)	E 15	MS	Loop	0,46 %	18	H = 1
						ATP			
p.Val1106Leu	c.3316G > C	pending	5 (HGMD)	E 15	MS	Loop	0,46 %	15	H = 1
						ATP			
p.Thr1178Ala	c.3532A > G	VCV000959197	5 (CV, HGMD)	E 16	MS	Loop	0,46 %	17	H = 1
						ATP			
p.Gln1210_Ser1211del	c.3627_3632del	pending	5 (HGMD)	E 7	Del	loop	0,46 %	12	H = 1
						ATP			
p.Val1217_Leu1218del	c.3649_3654del	VCV000189015	5 (CV, HGMD)	E 17	Del	Loop	0,46 %	11	H = 1
						ATP			
p.Thr1220Met	c.3659C > T	VCV000035725	5 (CV, HGMD)	E 17	MS	ATP	0,46 %	10	H = 1
						Bind			
p.Thr1232Pro	c.3694A > C	VCV000555144	5 (CV, HGMD)	E 17	MS	ATP	0,46 %	15	N = 1
						Bind			
p.Ala1274Thr	c.3820G > A	pending	5 (HGMD)	E 18	MS	ATP	0,46 %	12	H = 1
						Hinge			
p.Gly1281Asp	c.3842G > A	VCV000550301	5 (HGMD)	E 18	MS	ATP	0,46 %	18	H = 1
						Hinge			
p.Val1282LysFs*22	c.3843dup	VCV000958649	5 (CV, HGMD)	E 18	FS	ATP	0,46 %	12	H = 1
						Hinge			
p.Ala1283GlyFs*21	c.3845dupT	pending	5 (This study)	E 18	FS	ATP	0,46 %	16	N = 1
						Hinge			
p.Gln1351*	c.4051C > T	VCV000188947	5 (CV, HGMD)	E 20	NS	TM7	0,46 %	12	H = 1
p.Leu1373Pro	c.4118T > C	VCV000552788	5 (HGMD)	E 20	MS	TM8	0,46 %	11	H = 1
	del_E1	NA	5 (This study)	E 1	Del	before	0,46 %	3	H = 1
						Cu1			

Variant nomenclature based on Refseq NM\_000053.3.

Variant classification according to ACMG guidelines. 5: pathogenic. 4: likely pathogenic. 3: unknown significance. 2: likely benign. 1: benign.

Variant classification source. CV: ClinVar database (searched on 21/04/21). HGMD: Human Gene Mutation Database Professional (searched on 21/04/21). This study: for variants not referenced in ClinVar nor HGMD Pro, or referenced with uncertain significance or conflicting interpretation, authors interpretation based on ACMG recommendations.

Variant location and type. E: exon. I: Intron. MS: missense. NS: nonsense. FS: frameshift. Spl: splice site. Del: deletion.

Protein domain. Cu: copper-binding domain. TM: transmembrane domain. Td: Transduction domain ATP Bind: ATP binding domain. Ph: Phosphorylation domain. Clinical form. H: hepatic. N: neurologic. AS: asymptomatic.

variants in the French population since exons 14, 8 and 2 account for 42.6 % of total variants. However, 80 % of the variants in this population span at least 14 exons contrary to just eight exons in the UK population according to Coffey et al. (2013).

The considerable diversity of variants and the low percentages of homozygous patients, make it difficult to have more conclusive results on a genotype-phenotype correlation in our cohort. This correlation has

been extensively studied over the years in WD with no definitive results.

We followed similar approaches as those used by Panagiotakaki et al. (2004), Nicastro et al. (N et al., 1911) and Merle et al. (2010), but we could only verify lower ceruloplasmin levels associated with Frame-shift/Nonsense variants. Those studies founded an association of p. His1069Gln homozygous variant with neurologic and later presentation (Panagiotakaki et al., 2004); early onset of the disease, lower



**Table 3**

Data used to classified novel missense variants and variants previously classified as variants of unknown significance.

Variant	previous classification	authors classification	AF in GnomAD	In silico predictions	Variant associated in <i>trans</i>	Biochemical phenotype	Clinical phenotype	
p.Arg226Trp	c.676C > T	Unknown significance (CV, HGMD)	Likely pathogenic	max 0.039 % (all 0.0040 %)	prob. damaging (Polyphen), deleterious (SIFT)	c.2332C > T (p.Arg778Trp) c.2297C > G (p.Thr766Arg)  c.676C > T (p.Arg226Trp) homoz.	mild cytolysis cytolysis  NA	compensated cirrhosis fulminant hepatitis, liver transplantation neurologic form, KF ring. Liver transplantation compensated cirrhosis, hepatomegaly
/	c.1708-34G > A	ND	Likely pathogenic	Absent	creation of alternative consensus acceptor splice site	c.3800A > C (p.Asp1267Ala)	mild cytolysis	hepatitis, haemolytic anemia
p.Ala604Asp	c.1811C > A	ND	Likely pathogenic	absent	prob. damaging (Polyphen), deleterious (SIFT)	c.3843dup (p.Val1282LysFs*22)	Cp < 0,03	hepatitis, haemolytic anemia
p.Trp779Cys	c.2337G > C	ND	Likely pathogenic	absent	prob. damaging (Polyphen), deleterious (SIFT)	not identified	Persistent cytolysis, very low Cp, increased hepatic Cu	chronic hepatitis
p.Lys838Glu	c.2512A > G	Unknown significance (CV)	Likely pathogenic	max 0.017 % (all 0.0024 %)	prob. damaging (Polyphen), deleterious (SIFT)	c.51+4A > T	mild cytolysis, Cp = 0,12 g/L CuS = 5,75 µmol/L LCuEx = 0,69 µmol/L CuU = 14,4 µmol/24H	compensated cirrhosis, hepatomegaly
p.Thr991Ala	c.2971A > G	ND	Likely pathogenic	absent	prob. damaging (Polyphen), tolerated (SIFT)	c.1242_1243del (p.Arg414SerFs*9)	Cytolysis - Cp = 0,14 g/L - Cuu = 1,86 µmol/L - Cus = 4,40 µmol/L - CuEXC = 0,65 µmol/L - REC = 14,6 %	hepato-splenomegaly
p.Ala1003Pro	c.3007G > C	ND <sup>a</sup>	Likely pathogenic	absent	prob. damaging (Polyphen), deleterious (SIFT)	not identified <sup>b</sup>	Cp > 0,03 g/L CuS = 1,10 µmol/L CuEx = 0,38 µmol/L REC = 34,1 %	neurologic form, KF ring
p.Ser1067Arg	c.3201T > G	ND	Likely pathogenic	absent	prob. damaging (Polyphen), deleterious (SIFT)	c.3305T > C (p.Ile1102Thr)	Cp = 0,05 g/L - Cus = 6,9 µmol/L - CuEXC = 1,86 µmol/L - REC = 27 % - Cuu = 27,8 µmol/L	decompensated cirrhosis

Reference ranges for copper status explorations: Ceruloplasmin (Cp) > 0.2 g/L. Serum copper (CuS): 13–19 µmol/L. Exchangeable copper (CuEx): 0,38-1 µmol/L. Relative Exchangeable Copper (REC): 3–9%. Urinary copper (CuU) < 0.8 µmol/24h. <sup>a</sup> c.3007G > A (p.Ala1003Thr) reported pathogenic in CV and HGDM <sup>b</sup> Same genotype in affected sibling.

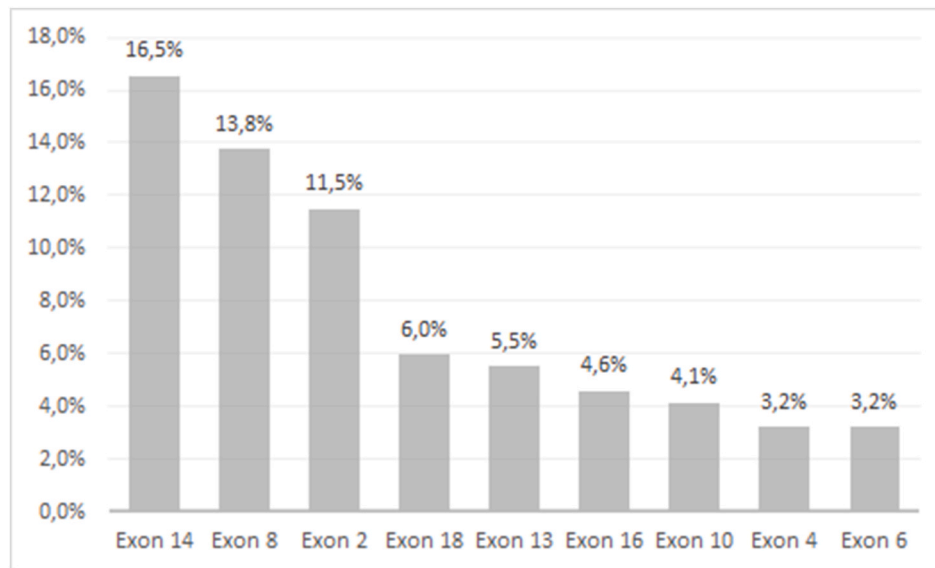


Fig. 1. Variant frequency of exons.

Table 4

Clinical and biochemical features of Patients Homozygous or Compound Heterozygous for Common Variants.

Genotype	n	Age		Phenotype								KF ring		Ceruloplasmin (mg/dL)		Serum copper ( $\mu\text{mol/L}$ )		Urinary copper ( $\mu\text{mol}/24\text{ h}$ )	
		(mean $\pm$ sd)		H	=	N	=	AS	=	AS	=	AS	=	AS	=	AS	=	AS	=
p.His1069Gln/p.His1069Gln	7	12,7 $\pm$ 3,5		H = 4		N = 1		AS = 2		29 %		12,0 $\pm$ 5,0		0,58 $\pm$ 0,2		4,6 $\pm$ 2,4			
c.1708-1G > A/c.1708-1G > A	2	15,0 $\pm$ 2,8		H = 2		N = 0		AS = 0		0 %		15,5 $\pm$ 3,5		0,73 $\pm$ 0,5		48,7 $\pm$ 65,3			
p.Val890Met/p.Val890Met	2	7,0 $\pm$ 0,0		H = 2		N = 0		AS = 0		0 %		11,5 $\pm$ 3,5		0,63 $\pm$ 0,1		4,2 $\pm$ 0,6			
p.Lys1020Arg/p.Lys1020Arg	2	11,0 $\pm$ 7,1		H = 2		N = 0		AS = 0		0 %		3,0 $\pm$ 0,0		0,89 $\pm$ 1,0		8,9 $\pm$ 6,1			
p.His1069Gln HMZ + compound HTZ	20	11,7 $\pm$ 3,5		H = 16		N = 2		AS = 2		15 %		12,1 $\pm$ 5,0		0,50 $\pm$ 0,3		4,8 $\pm$ 3,2			
c.1708-1G > A HMZ + compound HTZ	3	13,0 $\pm$ 2,8		H = 3		N = 0		AS = 0		0 %		11,0 $\pm$ 8,2		0,73 $\pm$ 0,4		34,2 $\pm$ 52,6			
p.Val890Met HMZ + compound HTZ	2	7,0 $\pm$ 0,0		H = 2		N = 0		AS = 0		0 %		11,5 $\pm$ 3,5		0,63 $\pm$ 0,1		4,2 $\pm$ 0,6			
p.Lys1020Arg HMZ + compound HTZ	2	11,0 $\pm$ 7,1		H = 2		N = 0		AS = 0		0 %		3,0 $\pm$ 0,0		0,89 $\pm$ 1,0		8,9 $\pm$ 6,1			
Missense	39	10,7 $\pm$ 4,7		H = 33		N = 4		AS = 2		18 %		8,2 $\pm$ 5,1		0,48 $\pm$ 0,4		4,9 $\pm$ 4,7			
Frameshift/Nonsense	8	10,8 $\pm$ 4,0		H = 7		N = 1		AS = 0		38 %		2,8 $\pm$ 0,7		0,33 $\pm$ 0,2		4,4 $\pm$ 3,6			

H: hepatic phenotype, N: neurological, AS: asymptomatic. KF: Kayser–Fleischer. HTZ: heterozygotes. HMZ: homozygotes.

ceruloplasmin levels, higher urinary copper excretion, and lower serum copper associated with Nonsense/Frameshift variants (N et al., 1911; Merle et al., 2010; Gromadzka et al., 2005). Type of variants have been associated with clinical phenotype and disease severity, but often other studies have opposite or not confirmatory results. A recent publication from Kluska et al. (2019) found an association with frameshift variant and a risk for neurological presentation. Neurological forms have been associated with p.His1069Gln homozygous variant according to some studies (Panagiotakaki et al., 2004; Stapelbroek et al., 2004; Maier-Dobersberger et al., 1997) but one study showed a more frequent hepatic presentation in patients with this variant (Mihaylova et al., 2012).

Many reasons could explain some of our differences with those studies. First of all, our population's extreme heterogeneity doesn't allow to perform very meaningful statistical calculations despite the important population since each of the multiple groups comprises only a few subjects. Being our cohort exclusively pediatric, the differences between the two main clinical forms might not yet be evident since hepatic forms are known to be predominant in this age group. Nevertheless, Nicastro et al. (N et al., 1911) reported a pediatric population

with differences in serum copper and ceruloplasmin levels related with genotype.

An impressive report of 1357 patients from Ferenci et al. (2019) found that clinical phenotype was more influenced by age and sex rather than by genotype, being that neurologic forms start later and are more common in men than the hepatic presentations. Other reports describe the association of age and sex (Kluska et al., 2019; Stapelbroek et al., 2004; Beinhardt et al., 2014; Litwin et al., 2012; Cheng et al., 2017) with WD's clinical forms. In our study, we've also found a later onset of neurologic forms comparing to asymptomatic and hepatic patients but no clear gender predominance. It has been speculated that the gender difference could be hormone-related. Following this line of thinking, this could also explain the lack of difference in our young cohort since estrogen exposure is much limited than of the women from adults cohorts.

Other factors that affect disease severity might include the copper levels in the diet, the amount of the metal in drinking water or other genetic modifying factors at nutritional and environmental levels.

Even considering the French population's extreme genetic heterogeneity, it would be useful to include adults in the analysis of the WD's

population to have a complete picture of phenotype evolution through age and its possible association with a genotype.

## 5. Conclusion

p.His1069Gln is the most frequent variant (14,2 %) and exons 14, 8, and 2 of the *ATP7B* gene account for 41.7 % of total variants. However, there is significant heterogeneity in the French population concerning the other *ATP7B* variants. Nonsense/Frameshift variants were associated with lower ceruloplasmin levels.

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**Update 1 of 2**

**European Journal of Medical Genetics**

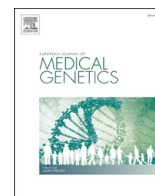
Volume 64, Issue 11, November 2021, Page

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## Corrigendum

## Corrigendum to “ATP7B variant spectrum in a French pediatric Wilson disease cohort” [Eur. J. Med. Genet. 64(10) (2021) 104305]



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**Update 2 of 2**

**European Journal of Medical Genetics**

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