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# DYT6 Dystonia: Review of the Literature and Creation of the UMD Locus-Specific Database (LSDB) for Mutations in the *THAP1* Gene

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**ABSTRACT:** By family-based screening, first Fuchs and then many other authors showed that mutations in *THAP1* (THAP [thanatos-associated protein] domain-containing, apoptosis-associated protein 1) account for a substantial proportion of familial, early-onset, nonfocal, primary dystonia cases (DYT6 dystonia). *THAP1* is the first transcriptional factor involved in primary dystonia and the hypothesis of a transcriptional deregulation, which was primarily proposed for the X-linked dystonia-parkinsonism (DYT3 dystonia), provided thus a new way to investigate the possible mechanism underlying the development of dystonic movements. Currently, 56 families present with a *THAP1* mutation; however, no genotype/phenotype relationship has been found. Therefore, we carried out a systematic review of the literature on the *THAP1* gene to colligate all reported patients with a specific *THAP1* mutation and the associated clinical signs in order to describe the broad phenotypic continuum of this disorder. To facilitate the comparison of the identified mutations, we created a Locus-Specific Database (UMD-*THAP1* LSDB) available at <http://www.umd.be/THAP1/>. Currently, the database lists 56 probands and 43 relatives with the associated clinical phenotype when available. The identification of a larger number of *THAP1* mutations and collection of high-quality clinical information for each described mutation through international collaborative effort will help investigating the structure-function and genotype-phenotype correlations in DYT6 dystonia.

**KEY WORDS:** dystonia; DYT6; *THAP1*, database; UMD

## Introduction

Dystonia defines a heterogeneous group of movement disorders due to central nervous system (CNS) dysfunction leading to sustained involuntary muscle contractions that cause abnormal postures and twisting movements. Since secondary dystonia is often linked to striatal injuries, it has been hypothesized that hereditary dystonia also could be due to a dysfunction of these structures and more broadly of the basal ganglia. Accordingly, many studies have reported electrophysiological, metabolic, and structural abnormalities in these brain areas [Vidailhet et al., 2009]. However, basal ganglia are not the only brain region that has been found to be altered in hereditary dystonia. Indeed, dysfunction of cerebral cortex, cerebellum, spinal cord, and other structures that are mainly involved in motor function has also been reported [Breakefield et al., 2008; Defazio et al., 2007]. Therefore, the concept that dystonia is a disorder linked to a specific CNS structure appears to be too restrictive and it should rather be considered as the result of a general disturbance of the motor circuit functions.

Hereditary dystonias are clinically and genetically heterogeneous. To date, 20 different genetically forms are known (DYT1 to DYT21; with DYT14 = DYT5) and mutations in nine genes have been involved [Breakefield et al., 2008; Norgren et al., 2011]. All the inheritance modes are found (autosomic recessive, autosomic dominant, and X-linked) but no epistatic relation between these genes was discovered except for DYT1 and DYT6 forms [Gavarini et al., 2010; Kaiser et al., 2010]. Since their products have different cell functions, it is very difficult up to now to propose a scheme of cell dysfunction common to all the dystonia forms and particularly to propose a unitary pathophysiological mechanism even if links with dopamine deregulation are often found [Blanchard et al., 2010; Park et al., 2005].

The last identified gene, *THAP1* for THAP [thanatos-associated protein] domain-containing, apoptosis-associated protein 1, was first found to be mutated in Amish-Mennonite families with primary dystonia and its implication was then extended to other

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populations [Almasy et al., 1997; Fuchs et al., 2009]. *THAP1* mutations cause DYT6 dystonia, an autosomal dominant primary form with about 60% penetrance. The main clinical characteristics of this form are early onset (mean = 17.8 years of age [calculated from 78 patients with available data]) and symptoms often localized to one upper limb at the beginning with tendency to extend to other body regions.

Fifty-three different mutations in the *THAP1* gene have been reported so far in 56 families (Table 1). These mutations are mainly private, essentially missense or out-of-frame deletions, usually non-recurrent, and widely distributed throughout the gene. To date, no genotype/phenotype relationship has been observed. From this perspective, we have compiled a database that gather all the available information on this gene and on the patients with *THAP1* mutations and created a software package to facilitate the mutational analysis of *THAP1*. This database is available at <http://www.umd.be/THAP1/>.

## THAP1 Expression

Two spliced mRNA variants that produce functional proteins have been reported (*THAP1a*: CCDS6136 and *THAP1b*: CCDS6137) [Girard et al., 2010]. The first 2.2-kb isoform contains three exons (Fig. 1), whereas the second corresponds to an alternatively spliced isoform that lacks exon 2 (2-kb mRNA). This second isoform encodes a truncated *THAP1* protein without the C-terminus of the THAP domain. The two isoforms are expressed in many tissues, suggesting that *THAP1* has a widespread (although not ubiquitous) distribution in humans.

In mouse brain tissue, immunoblot analysis revealed a highest concentration in embryonic whole brain tissue (at E16), which declines after birth in the different tested brain regions (at P60) [Gavarini et al., 2010].

*THAP1a*, which contains the THAP domain with DNA-binding properties, is the most studied.

## THAP1 Is a Transcription Factor

*THAP1* was first identified by two-hybrid screen of a high endothelial venule cell cDNA library using a cytokine bait [Roussigne et al., 2003]. *THAP1* is a 213 amino-acid-long protein characterized by an N-terminal THAP domain (amino acids 1 to 81) [Bessiere et al., 2008] with DNA-binding properties, followed by a proline-rich region (amino acids 90 to 110), and a nuclear localization signal (amino acids 146 to 162) (Fig. 1A).

The protein is only found in the nucleus where it can display both a diffuse distribution and a discrete localization in nuclear dots. *THAP1* colocalization with DAXX, a well-characterized protein that is expressed in promyelocytic leukemia (PML) nuclear bodies (NBs), indicates that these subnuclear regions are PML NBs. PML NBs are spheres of 0.1–1.0  $\mu\text{m}$ , heterogeneous in composition, mobility, and function, that are present in most mammalian cell nuclei [Lallemand-Breitenbach and de The, 2010]. The PML protein is the key organizer that recruits an ever-growing number of proteins, residing constitutively or more often transiently. PML NBs facilitate partner protein posttranslational modification (acetylation, sumoylation, or phosphorylation) resulting in partner sequestration, activation, or degradation. They are proposed to fine-tune a wide variety of processes: induction of apoptosis and cellular senescence, inhibition of proliferation, maintenance of genomic stability, and antiviral responses [Bernardi and Pandolfi, 2007].

*THAP1* interacts with the prostate apoptosis response-4 (PAR-4, HGNC-approved gene symbols Pawr), a pro-apoptotic

protein that acts as a transcriptional regulator [Roussigne et al., 2003]. PAWR is present in low amounts in dendrites and synapses throughout the brain under normal conditions and its amount increases rapidly in neurons subjected to several triggers of apoptosis including trophic factor deprivation, oxidative stress, and excitotoxins [Duan et al., 1999a; Guo et al., 1998]. Pawr induction has been linked to neuronal death in various neurodegenerative diseases as Parkinson's [Duan et al., 1999b], Alzheimer [Guo et al., 1998], or amyotrophic lateral sclerosis [Pedersen et al., 2000]. More interestingly, Pawr has been shown to serve other important function in nervous system. It interacts with D2 dopamine receptor (D2DR) suggesting its potential role in modulating calcium-mediated dopaminergic signaling [Park et al., 2005].

*THAP1* and PAWR exhibit pro-apoptotic activities (they increase the apoptosis sensitivity of mouse 3T3 fibroblasts to TNF- $\alpha$  and serum withdrawal when overexpressed) and they are both indirectly recruited to PML NBs by PML [Roussigne et al., 2003]. Roussigne and coworkers proposed that PML NBs regulate the functions of *THAP1* and PAWR. They may be the site of assembly and/or post-translational modifications of the *THAP1*/PAWR complex or they may serve as a site of storage or degradation, thus regulating the free nucleoplasmic pool of PAWR and *THAP1*.

Extensive database search using the THAP domain identified about 100 THAP proteins among seven model animal organisms [Clouaire et al., 2005]. Interestingly, some of these proteins are tightly linked to the pRB (retinoblastoma protein)/E2F pathway, which regulates commitment of mammalian cells to DNA replication. The THAP domain is present in the zebrafish ortholog of E2F6 and in five *Caenorhabditis elegans* proteins (LIN-36, LIN-15B, LIN-15A, HIM-17, and GON-14), which are known to be genetically linked to LIN-35, the only Rb ortholog found in *C. elegans*. Based on these data, Cayrol et al. [2007] investigated whether *THAP1* could be linked to the pRB/E2F pathway. They found that silencing as well as overexpression of *THAP1* led to cell cycle arrest at the G1/S transition. The similar effects following *THAP1* silencing/overexpression suggest that, like many other proteins, *THAP1* needs an optimal range of concentration to fulfill its physiological roles. DNA microarray analysis showed in both cases (upregulation or downregulation of *THAP1*) a reduction of the mRNA levels of cell cycle regulators and about 40 pRB/E2F-target genes [Cayrol et al., 2007], including *RRM1* (ribonucleotide reductase M1), *BIRC5* (survivin), *CDC2* (cell division cycle 2), *CCNB1* (cyclin B1) [Lacroix, 2007]. *RRM1* is activated at the G1/S transition and contains two predicted *THAP1*-binding sites (THABS) in its promoter. DNase I footprinting, electrophoretic mobility shift assay (EMSA) protein/DNA binding, and chromatin immunoprecipitation (ChIP) in proliferating endothelial cells demonstrated that endogenous *THAP1* directly binds to the promoter of *RRM1* [Cayrol et al., 2007]. Moreover, Mazars et al. [2010] showed that *THAP1* recruits HCF-1 (Host cell factor 1) to the *RRM1* promoter to allow its expression, thus adding further evidence for a role of *THAP1* in cell cycle regulation. Although the list of cofactors that participate in *THAP1* regulation of target genes is certainly not exhaustive, the available data clearly link *THAP1* to apoptosis (through its interaction with PAWR) and to cell cycle (through HCF-1).

## THAP1 Structure

The most extensive structural studies have focused on the THAP domain of *THAP1* and implicitly serve as a model for the other hundred THAP proteins discovered in the animal kingdom. In many aspects, the THAP domain, which is about 80-amino-acid long,

**Table 1. Mutations Reported in the THAP1 Gene in Each Family/Proband**

Mutation name	Nomenclature (protein)	Wild Type codon	Mutant codon	Mutational event	Amino acid position	Exon no.	Structure	HCD	Relatives in the database	Reference no.
<b>Inindel mutation</b>										
c.135_139delinsGGGTTTA, <sup>a</sup>	p.Phe45Leu5X28	CCC	Indels	Indels stop at 73	47	2	L3-THAP dom.		25	[Fuchs et al., 2009; Bressman et al., 2009]
<b>Missense mutations</b>										
c.1A>G	p.Met1?	ATG	GTG	A->G	1	1	THAP dom.		1	[De Carvalho Aguiar et al., 2010]
c.161T>C	p.Ser6Pro	TCC	CCC	T->C	6	1	L1-THAP dom.		1	[Clot et al., 2010]
c.17C>T	p.Ser6Phe	TCC	TTC	C->T	6	1	L1-THAP dom.		0	[Houlden et al., 2010]
c.23A>G	p.Iyr8Cys	ATG	TGC	A->G	8	1	L1-THAP dom.		0	[Houlden et al., 2010]
c.25G>T	p.Gly9Cys	GGC	TGC	G->T	9	1	L1-THAP dom.		1	[Xiao et al., 2010]
c.36C>A	p.Asn12Lys	AAC	AAA	C->A	12	1	L1-THAP dom.		0	[Bressman et al., 2009]
c.38G>A	p.Arg13His	CGC	CAC	G->A	13	1	L1-THAP dom.		1	[Zittel et al., 2010]
c.50A>G	p.Asp17Gly	GAC	GGC	A->G	17	1	L1-THAP dom.		0	[Xiao et al., 2010]
c.61T>A	p.Ser21Thr	TCT	ACT	T->A	21	1	L1-THAP dom.		0	[Bressman et al., 2009]
c.68A>C	p.His23Pro	CAC	CCC	A->C	23	1	BS1-THAP dom.		0	[Kaiser et al., 2010]
c.77C>G	p.Pro26Arg	CCT	CGT	C->G	26	2	L2-THAP dom.	DNA binding	0	[Houlden et al., 2010]
c.86G>A	p.Arg29Gln	CGA	CAA	G->A	29	2	L2-THAP dom.	DNA binding	0	[Paisan-Ruiz et al., 2009]
c.86G>A	p.Arg29Gln	CGA	CAA	G->A	29	2	L2-THAP dom.	DNA binding	0	[Paisan-Ruiz et al., 2009]
c.86G>C	p.Arg29Pro	CGA	CCA	G->C	29	2	L2-THAP dom.	DNA binding	0	[Bressman et al., 2009]
c.89C>G	p.Pro30Arg	CCC	CGC	C->G	30	2	L2-THAP dom.		1	[Jech et al., 2011]
c.95T>A	p.Leu32His	GTT	CAT	T->A	32	2	L2-THAP dom.	AA interactions	0	[Schneider et al., 2011]
c.115G>A	p.Ala39Thr	GCT	ACT	G->A	39	2	AH1-THAP dom.	AA interactions	0	[Bressman et al., 2009]
c.161G>A	p.Cys54Tyr	TGT	TAT	G->A	54	2	BS2-THAP dom.	C2CH motif	0	[Gavarini et al., 2010]
c.169C>A	p.His57Asn	CAC	AAC	C->A	57	2	AH2-THAP dom.	C2CH motif	0	[Sohn et al., 2010]
c.176C>T	p.Thr59Ile	ACT	ATT	C->T	59	2	THAP dom.		0	[Groen et al., 2010]
c.215T>G	p.Leu72Arg	CTG	CGG	T->G	72	2	L4-THAP dom.		0	[Clot et al., 2010]
c.224A>T	p.Asn75Ile	AAT	ATT	A->T	75	2	L4-THAP dom.		1	[Cheng et al., 2010]
c.241T>C	p.Phe81Leu	TTT	CTT	T->C	81	2	AH4-THAP dom.	AA interactions	3	[Fuchs et al., 2009]
c.247T>C	p.Cys83Arg	TGT	CGT	T->C	83	2			0	[Bressman et al., 2009]
c.266A>G	p.Lys89Arg	AAG	AGG	A->G	89	2			0	[Xiao et al., 2010]
c.395T>C	p.Phe132Ser	TTC	TCC	T->C	132	3			0	[Houlden et al., 2010]
c.407A>G	p.Asn136Ser	AAC	AGC	A->G	136	3		HCF-1 binding	1	[Groen et al., 2010]
c.408C>G	p.Asn136Lys	AAC	AAG	C->G	136	3		HCF-1 binding	0	[Sohn et al., 2010]
c.410A>G	p.Tyr137Cys	TAT	TGT	A->G	137	3	Coiled-coil dom.		0	[Sohn et al., 2010]
c.427A>G	p.Met143Val	ATG	GTG	A->G	143	3	NLS		1	[Xiao et al., 2010]
c.446T>C <sup>b</sup>	p.Ile149Thr	AIT	ACT	T->C	149	3	NLS		0	[Cheng et al., 2010]
c.449A>C	p.His150Pro	CAT	CCT	A->C	150	3	Coiled-coil dom.		0	[Xiao et al., 2010]
c.496G>A	p.Ala166Thr	GCA	ACA	G->A	166	3	Coiled-coil dom.		0	[Houlden et al., 2010]
c.506G>A	p.Arg169Gln	CGA	CAA	G->A	169	3	Coiled-coil dom.		0	[Bonetti et al., 2009]
c.508T>C	p.Cys170Arg	ATG	CGC	T->C	170	3	Coiled-coil dom.		0	[Xiao et al., 2010]
c.559C>A	p.Gln187Lys	CAG	AAG	C->A	187	3	Coiled-coil dom.		0	[Xiao et al., 2010]
c.574G>A	p.Asp192Asn	GAC	AAC	G->A	192	3			0	[Sohn et al., 2010]

(Continued)

**Table 1. Continued**

Mutation name	Nomenclature (protein)	Wild Type codon	Mutant codon	Mutational event	Amino acid position	Exon no.	Structure	HCD	Relatives in the database	Reference no.
<b>Nonsense mutations</b>										
c.7C>T	p.Gln3X	CAG	TAG	C->T	3	1	L1-THAP dom.		0	[Houlden et al., 2010]
c.85C>T	p.Arg29X	CGA	TGA	C->T	29	2	L2-THAP dom.	DNA binding	0	[Bressman et al., 2009]
c.150T>G	p.Iyr50X	TAT	TAG	T->G	50	2	L3-THAP dom.	DNA binding	0	[Houlden et al., 2010]
c.370C>T	p.Gln124X	CAG	TAG	C->T	124	3			0	[Söhn et al., 2010]
<b>Small out-of-frame deletions</b>										
c.2delT	p.Met1?	ATG	del1b	Stop at 72	1	1	THAP dom.		0	[Bressman et al., 2009]
c.20_33del	p.Ala7GlufsX23	GCC	del14b	Stop at 29	7	1	L1-THAP dom.		0	[Clot et al., 2010]
c.174delT	p.Phe58LeufsX15	TTT	del1c	Stop at 72	58	2	THAP dom.	DNA binding	0	[Houlden et al., 2010]
c.197_198delAG	p.Glu66ValfsX19	GAG	del2b	Stop at 84	66	2	L4-THAP dom.		0	[Söhn et al., 2010]
c.236delC	p.Thr79LysfsX41	ACA	del1b	Stop at 119	79	2	AH4-THAP dom.	AVPTIF motif	0	[Houlden et al., 2010]
c.377_378delCT	p.Pro126ArgfsX2	CCT	del2b	Stop at 127	126	3			1	[Blanchard et al., 2011]
c.388_389delTC	p.Val113PhefsX3	TCA	del2a	Stop at 133	130	3			0	[Söhn et al., 2010]
c.388_389delTC <sup>b</sup>	p.Val131PhefsX3	TCA	del2a	Stop at 133	130	3			0	[Djarmati et al., 2010]
c.436_443del	p.Arg146AspfsX9	CGG	del8a	Stop at 154	146	3	NLS		0	[Clot et al., 2010]
c.460delC	p.Gln154SerfsX27	CAG	del1a	Stop at 180	154	3	NLS		0	[Bressman et al., 2009]
c.474delA <sup>b</sup>	p.Lys158AsnfsX23	AAA	del1c	Stop at 180	158	3	NLS		2	[Djarmati et al., 2010]
<b>Small in-frame deletion</b>										
c.207_209delCAA	p.Asn69_Asn69del	AAC	del3c	In-frame del	69	2	L4-THAP dom.		0	[Groen et al., 2010]
c.207_209delCAA	p.Asn69_Asn69del	AAC	del3c	In-frame del	69	2	L4-THAP dom.		2	[Clot et al., 2010]
<b>Small out-of-frame insertion</b>										
c.514dup	p.Arg172LysfsX7	AGG	ins1b	Stop at 178	172	3	Coiled-coil dom.		0	[Blanchard et al., 2011]

For each mutation, the nucleotide and protein names are given according to the nomenclature guidelines (<http://www.hgvs.org/>).

<sup>a</sup>Note that the c.134\_135insGGGT;137\_139delAAC Amish mutation has been renamed in c.135\_139delinsGGTTTA to follow the guideline for "insdel" mutation (<http://www.hgvs.org/>).

<sup>b</sup>Same patient described in the two given references; "THAP dom.": THAP domain;

"L1-THAP dom.": THAP-domain loop 1;

"L2-THAP dom.": THAP-domain loop 2;

"L3-THAP dom.": THAP-domain loop 3;

"L4-THAP dom.": THAP-domain loop 4;

"AH1-THAP dom.": THAP-domain  $\alpha$ -helix 1;

"H2-THAP dom.": THAP-domain 3<sup>o</sup>helix 2;

"H3-THAP dom.": THAP-domain 3<sup>o</sup>helix 3;

"H4-THAP dom.": THAP-domain 3<sup>o</sup>helix 4;

"Coiled-coil dom.": coiled-coil domain;

"NLS": nuclear localization sequences;

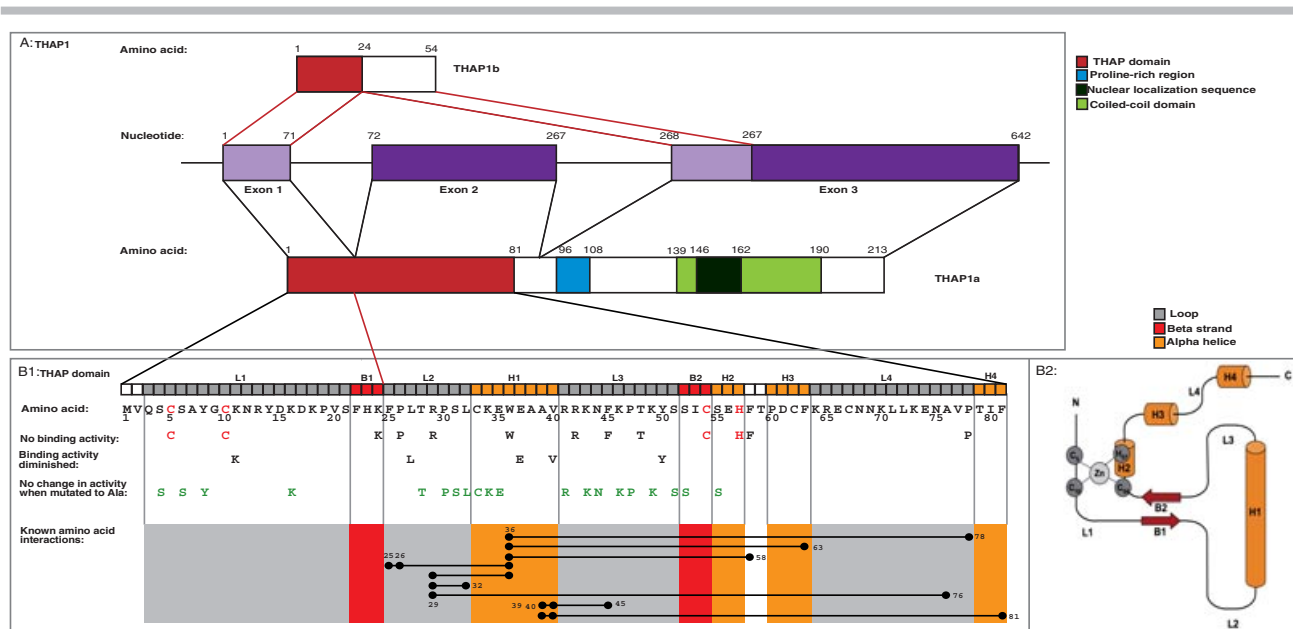
"DNA binding": amino acids proven to be involved in DNA-binding activity;

"C2CH motif": amino acids involved in the zinc-dependent DNA-binding activity of THAP1 through the C2CH motif;

"HCF-1 binding": amino acids involved in the interaction with HCF-1;

"AA interactions": interactions between THAP1 amino acids as described in Figure 1B. Example: THAP1 amino acid 39 is involved in interactions with THAP1 amino acids 40, 45, and 81.

For each proband, the number of relatives included in the database is given.



**Figure 1.** *THAP1* and its THAP domain. **A:** Schematic representation of the *THAP1* gene and its two isoforms. Isoform 1 (*THAP1a*) and isoform 2 (*THAP1b*) result from alternative splicing of exon 2; **B1:** Schematic representation of the THAP domain with amino acid positions for each structure. The upper figure represents the  $\alpha$ -helix H1, the three  $3_{10}$  helices (H2, H3, and H4) (all in orange), the two  $\beta$ -sheets (B1 and B2) (in red), and the four loops (L1 to L4, in gray). “No binding activity”: when these amino acids of the THAP domain are mutated to alanine, *THAP1* shows loss of zinc-dependent DNA-binding activity; amino acids that are part of the C2CH motif are in red; “Binding activity diminished”: when these amino acids are mutated to alanine, there is partial loss of DNA-binding activity; “No binding activity modification when mutated to Ala”: mutation of these amino acids to alanine does not change the DNA-binding activity of *THAP1* [Bessiere et al., 2008; Clouaire et al., 2005]; “Known amino acid interactions”: structural representation of the known amino acid interactions [Bessiere et al., 2008]; **B2:** Topology diagram of the THAP domain of human *THAP1* showing the secondary structure elements: the  $\alpha$ -helix H1, the three  $3_{10}$  helices (H2, H3, and H4) (all in orange), the two  $\beta$ -sheets (B1 and B2) (in red), and the location of the four loops (L1 to L4). The zinc finger and the four ligands are shown in gray. Modified from Bessiere et al. [2008].

exhibits some specific features (Fig. 1B1). It is characterized by a C2CH signature (Cys- $X_{2-4}$ -Cys- $X_{35-53}$ -Cys- $X_2$ -His) associated with four invariant residues (Pro<sub>26</sub>, Trp<sub>36</sub>, Phe<sub>58</sub>, Pro<sub>78</sub>) in the *THAP1* sequence. Direct mutagenesis of each of these eight amino acids showed their critical identity for the zinc-dependent specific binding of the THAP domain to a precise DNA sequence (TXXXGGCA: THABS consensus sequence) [Clouaire et al., 2005] (Fig. 1B2). An AVPTIF box (Ala<sub>76</sub>-Phe<sub>81</sub>, in the *THAP1* sequence), also essential for DNA binding, is located at the C-terminus of the THAP domain.

The NMR solution structure of the THAP zinc finger of *THAP1* [Bessiere et al., 2008] indicates that this domain consists of a first loop (L1: Gln<sub>3</sub>-Ser<sub>21</sub>), followed by two short antiparallel  $\beta$ -sheets (B1: Phe<sub>22</sub>-Lys<sub>24</sub> and B2: Ser<sub>52</sub>-Cys<sub>54</sub>) separated by a loop (L2: Phe<sub>25</sub>-Lys<sub>32</sub>)-helix (H1: Cys<sub>33</sub>-Val<sub>40</sub>)-loop (L3: Arg<sub>41</sub>-Ser<sub>51</sub>) structure (Fig. 1B2). The four residues of the C2CH motif (Cys<sub>5</sub>-Cys<sub>10</sub>-Cys<sub>54</sub>-His<sub>57</sub>) are three-dimensionally brought together to form the zinc-dependent DNA-binding site. After the second  $\beta$ -strand, there are three  $3_{10}$  helices: H2 (Ser<sub>55</sub>-His<sub>57</sub>), H3 (Arg<sub>60</sub>-Phe<sub>63</sub>), and H4 (Thr<sub>79</sub>-Phe<sub>81</sub>). H4 contains part of the AVPTIF motif and is preceded by a last mobile loop (L4: Lys<sub>64</sub>-Pro<sub>78</sub>).

The alpha-helix H1 can be considered as an important interaction hub and it is essential for the maintenance of the three-dimensional (3-D) shape of the THAP domain (Fig. 1B1). Indeed, it makes contacts with many residues distributed throughout the domain allowing a spatial connection between H1 (Trp<sub>36</sub>, Ala<sub>39</sub>, Val<sub>40</sub>), L2 (Phe<sub>25</sub>, Pro<sub>26</sub>, Arg<sub>29</sub>), L3 (Phe<sub>45</sub>),  $\sim$ H2 (Phe<sub>58</sub>), H3 (Phe<sub>63</sub>), L4 (Pro<sub>78</sub>), and H4 (Phe<sub>81</sub>). Residue Trp<sub>36</sub>, which is located in the core of H1, seems to be a key amino acid for this function. It makes hydrophobic contacts with four amino acids (Phe<sub>25</sub>, Arg<sub>29</sub>, Phe<sub>58</sub>,

and Phe<sub>63</sub>), and NMR data strongly suggest the presence of interaction also with the two highly conserved prolines (Pro<sub>26</sub> and Pro<sub>78</sub>). Four of these residues have been functionally tested (Pro<sub>26</sub>, Arg<sub>29</sub>, Phe<sub>58</sub>, Pro<sub>78</sub>) and their replacement by alanine strongly decreases the THAP domain ability to bind to DNA, suggesting that they have a critical role in the maintenance of *THAP1* 3-D shape [Clouaire et al., 2005].

The 3-D structure of the DNA-THAP domain complex has recently been solved [Campagne et al., 2010], showing that the THAP domain contacts DNA by insertion of its two antiparallel  $\beta$ -sheets into the major DNA groove. This ensures the connection of the second half of loop L3 (residues Lys<sub>46</sub> to Ser<sub>51</sub>) with the core of the THABS sequence. Upon binding to DNA, the folding of *THAP1* is changed, enabling loop L4 to contact the DNA minor groove. Based on their nonconservation in the THAP family, specific base-contact with the core of the THABS motif and chemical behavior with non-specific DNA, some THAP domain residues have been proposed to play a crucial role in the specificity of DNA recognition: Tyr<sub>50</sub> and Ser<sub>51</sub> of loop L3, Ser<sub>52</sub> of B2, Gln<sub>3</sub> and Ser<sub>4</sub> of the N-terminal tail and, finally Arg<sub>65</sub> of L4. Other residues are involved in the nonspecific interaction with DNA, but their role remains absolutely necessary for the targeting of *THAP1* to the THABS sequence. For example, Thr<sub>48</sub>, which is located in the middle of the DNA-interaction area of loop L3, shows a chemical shift perturbation when the THAP domain is incubated with an unrelated DNA sequence, suggesting that Thr<sub>48</sub> is involved in nonspecific interactions. However, mutation of this residue led to total loss of the DNA-binding activity of the THAP domain and it has been proposed that, with Lys<sub>46</sub>, it may be important for positioning the protein onto DNA.

**Table 2. Clinical Characteristics of Carriers of *THAP1* Mutations**

	Amish patients <sup>a</sup> N = 25	Non-Amish patients N = 81	All patients <sup>b</sup> N = 106
<b>Sex:</b>	F 15 (60%)	F 46 <sup>c</sup> (57%)	F 61 <sup>c</sup> (58%)
<b>Age at onset (y):</b>	Median 14.5 (5–38)	Median 13 (2–62)	Median 13 (2–62)
<b>Age at last examination (y):</b>	Median 40 (10–66)	Median 44 (7–86)	Median 44 (7–86)
<b>Family history:</b>	100%	58.5%	61.4%
<b>Site at onset:</b>	<b>25 patients</b>	<b>77 patients<sup>d</sup></b>	<b>102 patients<sup>d</sup></b>
Upper limb	11 (44%) <sup>e</sup>	36 (47%) <sup>e</sup>	48 (47%)
Lower limb	1 (4%) <sup>e</sup>	14 (18%) <sup>e</sup>	17 (17%)
Cervical	5 (20%) <sup>e</sup>	17 (22%) <sup>e</sup>	23 (23%)
Cranial	8 (32%)	17 (22%)	25 (25%)
<b>Site at examination:</b>	<b>25 patients</b>	<b>81 patients</b>	<b>106 patients</b>
Upper limb	22 (88%)	60 (74%)	82 (77%)
Lower limb	12 (48%)	39 (48%)	51 (48%)
Cervical	14 (56%)	59 (73%)	73 (69%)
Cranial	18 (72%) <sup>e</sup>	40 (49%) <sup>e</sup>	58 (55%)
Speech	17 (68%) <sup>e</sup>	48 (59%) <sup>e,f</sup>	65 (61%) <sup>e,f</sup>
<b>Distribution:</b>	<b>25 patients</b>	<b>81 patients</b>	<b>106 patients</b>
FD: 3 (12%)		FD: 16 (20%)	FD: 19 (18%)
SD: 10 (40%)		SD: 20 (25%)	SD: 30 (28%)
MD: 4 (16%)		MD: 5 (6%)	MD: 9 (8%)
GD: 8 (32%)		GD: 40 (49%)	GD: 48 (45%)

Gender: F, female; M, male. Dystonia distribution: “GD”: generalized dystonia;

“FD”: focal dystonia;

“MD”: multifocal dystonia;

“SD”: segmental dystonia.

<sup>a</sup>Four families with common ancestry, 25 patients. Data from Fuchs et al. [Söhn et al., 2010].

<sup>b</sup>Thirty-five families, 22 sporadic, total 106 patients.

<sup>c</sup>Sex is unknown for one patient from Söhn et al. [Söhn et al., 2010].

<sup>d</sup>For some patients, data were unavailable [Bressman et al., 2009; Gavarini et al., 2010; Kaiser et al., 2010].

<sup>e</sup>Data were estimated from Bressman et al. [2009] and Fuchs et al. [2009] when possible.

<sup>f</sup>Of note, two patients from Groen et al. [2010] have oromandibular dystonia without knowing if there was speech disturbance or not.

## THAP1 Is Involved in DYT6 Dystonia

DYT6 dystonia has been first described in the genetically isolated Amish–Mennonite population [Almasy et al., 1997] as an autosomal dominant trait with incomplete penetrance. In two Amish–Mennonite families, the site of onset of the disease was localized with similar frequency to upper limb, cervical, or cranial muscles and symptoms began between 5 and 38 years of age (average 18.1) with a tendency to progressive involvement of other body regions. Hence, DYT6 dystonia was named “idiopathic torsion dystonia of ‘mixed’ type.” These families were large enough to identify by linkage analysis the morbid locus, which maps in both families to chromosome 8p21–22.

The locus was further narrowed by the discovery of other affected Amish–Mennonite families [Saunders–Pullman et al., 2007] and, in 2009, Fuchs et al. [2009] demonstrated the involvement of a *THAP1* mutation in DYT6 dystonia: an insertion/deletion founder mutation found in four related Amish–Mennonite families. A point mutation was then identified in an unrelated non-Amish–Mennonite German family (family S, c.214T>C), indicating that DYT6 dystonia was present also in populations of different ancestry. Indeed, the complex insertion/deletion mutation was later identified also in a German family with no known Amish–Mennonites origin but sharing the same haplotype [Bressman et al., 2009].

Several teams have reported to date 53 different mutations in the coding sequence of *THAP1* in 56 families (56 probands and 43 relatives) of American [Bressman et al., 2009; Xiao et al., 2010], Chinese [Cheng et al., 2010; Xiao et al., 2010], Czech [Jech et al., 2011], Dutch [Groen et al., 2010], English [Houlden et al., 2010], French [Blanchard et al., 2011; Clot et al., 2010; Houlden et al., 2010], German [Bressman et al., 2009; Djarmati et al., 2009; Fuchs et al., 2009; Houlden et al., 2010; Söhn et al., 2010; Zittel et al., 2010],

Greek [Bonetti et al., 2009; Houlden et al., 2010], Iranian [Schneider et al., 2011], Jewish [Houlden et al., 2010], Mauritian/Indian [Houlden et al., 2010], Spanish [Paisan-Ruiz et al., 2009], and Brazilian [De Carvalho Aguiar et al., 2010] ancestry. Unfortunately, clinical data are not available for 11 probands implemented in the database [Bressman et al., 2009; Gavarini et al., 2010; Kaiser et al., 2010] and insufficient clinical and molecular data for 10 relatives do not allow to implement them in the database [Bressman et al., 2009]. One of the mutation has been described in an asymptomatic individual control (c.197\_198delAG) [Söhn et al., 2010].

The enlargement of the group of patients with *THAP1* mutations (108 up to now without the asymptomatic carrier) has provided new data for delineating the phenotype of DYT6 dystonia, which nevertheless has hardly changed since the original description (Table 2). DYT6 dystonia has been reported slightly more in females than males. The age distribution at onset is broad, ranging from 2 to 62 years (mean 17.8 years [calculated from 78 patients with available data]). Onset before 10 years of age occurs in 35.8% of the patients, and onset after 28 years of age in one-sixth (16%) of the patients. In almost half (47%) of the patients, the first symptoms of dystonia concern an upper limb, while cranial or cervical onset is almost equally observed in the other half of the patients. Onset in a lower limb is less common (17/102 patients reported in the literature). DYT6 dystonia tends to spread slowly to other body regions. Although the symptom distribution varies widely, the involvement of the cranial region is frequent, and disability mainly results from cranial dystonia with speech difficulties. At last follow-up, almost half of the reported patients with DYT6 dystonia suffered from generalized dystonia, but most of them remained ambulatory. To date, 9 patients underwent deep brain stimulation; one good response is reported in one patient [Jech et al., 2011], although deep brain stimulation (DBS) resulted in only mild or moderate improvement

in the other patients, with poor effect on speech or dysarthria [Clot et al., 2010; Djarmati et al., 2009; Groen et al., 2010; Jech et al., 2011; Zittel et al., 2010].

Early onset in a limb and progressive generalization in few years are shared features between DYT6 and DYT1 dystonias [Valente and Albanese, 2010]. Nevertheless, several differences emerge. In DYT6 dystonia, cervical or cranial muscles are involved most of the time, whereas lower limbs are frequently spared. The DYT6 clinical spectrum is more variable, and some adult-onset cases with stable focal dystonia are reported as well. Onset (mean age) is somewhat later and with a wider age range in DYT6 than in DYT1 patients. The penetrance in DYT6 is difficult to evaluate; it was performed only in the Amish–Mennonite population (around 60% [Saunders-Pullman et al., 2007]).

Recently, *TOR1A* has been demonstrated as a direct target of *THAP1* by EMSA, ChIP, and luciferase reporter gene assays [Gavarini et al., 2010; Kaiser et al., 2010]. Specific modulation of torsinA expression was not reported in nonneuronal cells after *THAP1* knockdown or overexpression, nor in fibroblasts or lymphoblast cells from DYT6 patients [Gavarini et al., 2010; Kaiser et al., 2010]. The lack of effect could be explained by tissue and/or developmental stage specificity. However, *THAP1* is predicted to potentially regulate numerous gene targets [data not published] and *TOR1A* deregulation in DYT6 patients should then be responsible of only one part of the phenotype. Nevertheless, human wild-type torsinA can lead to a cellular dysfunction of mice neurons when overexpressed at high levels (abundant inclusion-like structures, abnormalities of the NE ultrastructure, behavioral abnormalities, significant reduction of dopamine levels, serotonin, and homovanillic acid) as shown in transgenic mice line hWT24 [Grundmann et al., 2007].

## The UMD-THAP1 Database

We have used the Universal Mutation Database (UMD) software [Beroud et al., 2000, 2005] to create a computerized locus-specific database that contains information about the published mutations of the *THAP1* gene. Codons were numbered (e.g., +1 = A of ATG) based on the cDNA sequence of the *THAP1* variant 1 (isoform with three exons) (accession number NM\_018105.2, “Homo sapiens THAP domain containing, apoptosis associated protein 1 (*THAP1*), transcript variant 1, mRNA”) obtained from the GenBank database. Intron–exon boundaries as well as intronic sequences were defined by matching the cDNA sequence to the corresponding genome sequence (NT\_167187.1) and to the module organization from SwissProt (accession number Q9NVV9). The database follows the guidelines on mutation databases of the Hugo Mutation Database Initiative including the latest nomenclature (<http://www.hgvs.org/>).

The *THAP1* mutation database can list point mutations, large and small deletions, insertions, indels, and mutations affecting splicing (intronic mutations) in the *THAP1* gene. It cannot accommodate mutations from the UTR and promoter regions. In addition, two mutations that affect the same allele are entered as two different records linked by the same sample ID. It allows to link specific polymorphisms identified in the course of sequencing to one mutation for a specific patient. Separate entries are made for each probands and for each relative (with different sample ID) in order to describe specific phenotypes. If the same mutation in a single patient has been reported in different papers (example: the same patients 7021, L-2257, L-3736, and L-3637 were first reported in [Djarmati et al., 2009] and subsequently in [Zittel et al., 2010]), only one entry was made. If the same mutation has been reported in apparently unrelated patients (example c.207\_209delCAA identified in French

and Dutch patients [Clot et al., 2010; Groen et al., 2010]), separate entries were made for each patient as recurrent mutations, in the absence of haplotypes demonstrating common ancestor. For each mutation, different types of information are provided: genetic (exon and codon number, wild type and mutant codon, mutational event, mutation name), protein (wild type and mutant amino acids, affected domain, mutation name), clinical data (age of onset, site of onset, dystonia distribution, sites involved, genealogic tree when available), and experimental data.

We have annotated the *THAP1* sequence with indirect arguments in order to determine whether potential missense mutations are really causative using the UMD-Predictor<sup>®</sup> tool [Frederic et al., 2009]. The UMD-Predictor<sup>®</sup> tool is dedicated to the analysis of nucleotide substitutions in cDNA sequences. It provides a combinatorial approach that includes localization of the mutation in the protein, conservation, biochemical properties of the mutant and wild type residue, and potential impact of the variation on mRNA (<http://www.umd.be>). The indirect arguments, listed in the “Highly conserved domain” (HCD) of UMD-Predictor<sup>®</sup> are: (1) amino acids involved in the zinc-dependent DNA-binding activity of *THAP1* through the C2CH motif [Clouaire et al., 2005]; (2) amino acids involved in DNA binding of the THAP domain of human *THAP1* [Bessiere et al., 2008; Clouaire et al., 2005]; (3) amino acids involved in the interaction with HCF-1 (HBM consensus sequence) [Mazars et al., 2010]; and (4) *THAP1* amino acids involved in interactions with other *THAP1* amino acids [Bessiere et al., 2008] (Fig. 1B1).

Mutations that potentially affect splicing can be highlighted with dedicated algorithms that have been recently implemented in the Human Splicing Finder (HSF) tool [Desmet et al., 2009]. HSF is also available for any gene as a web resource (<http://www.umd.be/HSF/>) and is dedicated to the prediction of the impact of a mutation on splicing signals and/or to the identification of these motifs in any human sequence. It contains all available matrices for auxiliary sequences prediction, as well as new position weight matrices to evaluate the strength of 5' and 3' splice sites and of branch point sequences.

The UMD-THAP1 Database can be used with different routines described at <http://www.umd.be/THAP1/>. It is possible to analyze the mutation distribution by exon or by mutation type and the mechanism for deletion/insertion. With the “HCD” annotation, it is possible to know whether an amino acid has a known reported function in the protein. The UMD software has already been used successfully since 1994 for the analysis of multiple genes implicated in genetic diseases, such as *LDLR* [Villegier et al., 2002], *TGFBR2* [Frederic et al., 2008], *DYS* [Krahn et al., 2009], and in cancer, such as *TP53* [Cariello et al., 1994a, b; Hamroun et al., 2006] or *MEN1* [Wautot et al., 2002]. Twenty-nine UMDs are available through the web. More information concerning the UMD software is available at <http://www.umd.be>.

The current database and the ensuing updated versions are available online at <http://www.umd.be>. Notifications of omissions and errors in the current version as well as specific phenotypic data will be gratefully received by the corresponding author. The software will be expanded as the database grows and according to the users' requirements, and new functions may be implemented.

## Mutation Analysis

To date, a total of 53 different *THAP1* mutations are known (Table 1) since two of the three patients and their specific mutation reported by Zittel et al. [2010] were already described by Djarmati



et al. [2009], and the patient reported by Van Gerpen et al. [2010] is part of the cohort described by Xiao et al. [2010]. These mutations are mainly private, missense, and widely distributed throughout the gene. Only three apparently recurrent mutations have been described. For the c.86G>A, described in two patients by Paisan-Ruiz et al. [2009], no other polymorphism has been described ruling out common ancestor [Coro Paisan-Ruiz, personal communication]. The c.388\_389delTC mutation [Djarmati et al., 2009; Söhn et al., 2010] and the c.207\_209delCAA mutation [Clot et al., 2010; Groen et al., 2010] have been characterized by different teams, but until haplotype analyses become available, it is unclear whether these are truly recurrent mutations or whether they result from a founder effect. These mutations have then been considered as recurrent. It is noteworthy that two mutations, c.407A>G [Houlden et al., 2010] and c.95T>A [Schneider et al., 2011] have been found to be homozygous.

### Insel Mutations

The heterozygous Amish mutation, described as a 5-base (GGGTT) insertion followed by a 3-base deletion (AAC) (c.134\_135insGGGTT;137\_139delAAC) [Fuchs et al., 2009], has been detected in four related Amish–Mennonite families (families M, C, R, and W). This mutation was later identified also in a German family without known Amish–Mennonite ancestry (family 2) but sharing the Amish haplotype [Bressman et al., 2009]. This patient has then been implemented as a relative. To follow the international guideline for “insdel” mutation nomenclature (<http://www.hgvs.org/>), the name of the mutation has been corrected to c.135\_139delinsGGGTTTA, that is deletion of five bases (TAAAC) and insertion of seven bases (GGGTTTA), which results in the same predicted mutant protein.

### Nonsense and Missense Mutation

Four reported mutations are nonsense mutation: c.7C>T [Houlden et al., 2010], c.85C>T [Bressman et al., 2009], c.150T>G [Houlden et al., 2010], and c.370C>T [Söhn et al., 2010]. Thirty-seven are missense mutations. Five mutations are predicted to impair THAP1 DNA-binding activity: c.77C>G [Houlden et al., 2010]; the two recurrent mutations c.86G>A [Paisan-Ruiz et al., 2009]; c.86G>C [Bressman et al., 2009]; and c.169C>A in the C2CH motif [Söhn et al., 2010]. Three mutations are located in the sequence involved in THAP1 binding to HCF-1: c.407A>G [Houlden et al., 2010], c.408C>G [Groen et al., 2010], and c.410A>G [Söhn et al., 2010]. One mutation (c.241T>C) is located in the AVPTIF motif [Fuchs et al., 2009]. The homozygous mutation c.407A>G [Houlden et al., 2010] creates a potential acceptor splice site in exon 3, but its strength and the presence of many upstream good candidate acceptor splice sites make it unlikely that it will be used.

The c.266A>G mutation [Bressman et al., 2009] is located in the penultimate nucleotide of exon 2. This mutation does not impair the donor acceptor splice site, neither auxiliary splicing sequences (exonic sequence enhancer [ESE] or silencer [ESS]).

For the 24 remaining mutations, the mutated codon does not seem to be clearly involved in a particular function. Among all the missense mutations, only nine were not predicted by the UMD-Predictor<sup>®</sup> tool to be pathogenic based on a combinatorial approach [Frederic et al., 2009] (Table 3): c.38G>A [Zittel et al., 2010], c.61T>A [Bressman et al., 2009], c.169C>A [Söhn et al., 2010], c.241T>C [Fuchs et al., 2009], c.266A>G [Bressman et al., 2009], c.427A>G [Söhn et al., 2010], c.506G>A [Houlden et al., 2010], c.559C>A [Xiao et al., 2010], and c.574G>A [Söhn

et al., 2010]. Nevertheless, mutation c.241T>C, reported in dbSNP without frequency data, shows a decreased DNA-binding activity in vitro [Fuchs et al., 2009]. EMSA experiments for c.61T>A show that the mutation abolishes binding of THAP1 to *TOR1A* [Bressman et al., 2009; Gavarini et al., 2010], while mutation c.38G>A is shown to significantly reduce THAP1-mediated repression of *TOR1A* (57% of residual activity) [Kaiser et al., 2010; Zittel et al., 2010].

UMD-Predictor<sup>®</sup> tool has a very high positive predictive value and a high negative predictive value. Mutations experimentally demonstrated by gel-shift assays (Table 1) [Bessiere et al., 2008; Clouaire et al., 2005] to have (1) no binding activity (p.Cys5Ala, p.Cys10Ala, p.Lys24Ala, p.Pro26Ala, p.Arg29Ala, p.Trp36Ala, p.Arg42Ala, p.Phe45Ala, p.Thr48Ala, p.Cys54Ala, p.His57Ala, p.Phe58Ala, and p.Pro78Ala) are all predicted as pathogenic; (2) a diminished binding activity were also all predicted as pathogenic (p.Glu37Ala, p.Lys11Ala, p.Leu27Ala, and p.Tyr50Ala) or probably pathogenic (p.Val40Ala); (3) no change in activity were either predicted: as pathogenic (p.Ser4Ala, p.Tyr8Ala, p.Lys16Ala, p.Leu32Ala, p.Arg41Ala, p.Asn44Ala, p.Lys46Ala, p.Lys49Ala); as probably pathogenic (p.Thr28Ala, p.Pro30Ala, p.Glu35Ala, p.Lys34Ala, p.Lys43Ala); as polymorphism (p.Ser55Ala); or as probable polymorphism (p.Ser6Ala, p.Pro47Ala, p.Ser31Ala, p.Cys33Ala, p.Ser51Ala, p.Ser52Ala). Nevertheless, these gel-shift assays demonstrate no modification of the DNA-binding activity “within the limits of detection of the present assay” and do not exclude “a role in the binding affinity and selectivity” [Bessiere et al., 2008]. Amino acids Ser<sub>4</sub>, Ser<sub>51</sub>, and Ser<sub>52</sub> are described to be involved in the specificity of DNA recognition and Lys<sub>46</sub> proposed, with Trp<sub>48</sub>, to be important for positioning the protein onto DNA [Campagne et al., 2010]. Therefore, the predictions are in full agreement with in vitro studies for pathogenic mutations (18/18). For the alanine substitutions classified as “non modifying binding activities,” some discrepancies are observed but their interpretation is difficult as they could result from wrong predictions by UMD-predictor or wrong interpretation of experimental data.

These experimental results show that in spite of the high efficiency of the UMD-predictor tool [Frederic et al., 2009] and other prediction tools [SIFT, PolyPhen, data not shown] for several genes (*FBN1*, *FBN2*, *TGFBR1*, *TGFBR2* [Frederic et al., 2009], *DYS* [Krahn et al., 2009], *CDKN2A* [Kannengiesser et al. 2009], and *ALAS2* [Ducamp et al. 2011]), more functional data will have to be collected to improve the prediction of missense variations in *THAP1* gene.

### Insertion/Duplication

Only one reported mutation is an insertion that corresponds to the duplication of a single base (A<sub>514</sub>). This insertion is predicted to create a premature termination codon (PTC), codon seven amino acids downstream of the mutation (position 178) [Blanchard et al., 2011].

### Deletions

Among the 13 small deletions, 11 create a PTC and two are in-frame deletions resulting in loss of the ASN<sub>69</sub> residues (c.207\_209delCAA). Two single-base deletions, c.174delT [Houlden et al., 2010] and c.474delA [Djarmati et al., 2009], might be the result of slipped mispairing. Five mutations are deletions of a repeated sequence: c.197\_198delAG [Söhn et al., 2010], the recurrent mutations c.388\_389delTC [Djarmati et al., 2009; Söhn et al., 2010] and c.207\_209delCAA [Clot et al., 2010; Groen et al., 2010]. One mutation, c.436\_443del, is flanked by direct repeats [Clot et al., 2010]. For the five other mutations, the mechanisms are unknown

**Table 3. Analysis of Missense Mutations with the UMD-Predictor<sup>®</sup> Tool**

Nomenclature c.	Nomenclature p.	Structure	HCD	Conservation	SIFT probability	BLOSUM 62	Biochemical values	ESE modification	Splice site	Pathogenicity	Conclusion
c.17C>T	p.Ser6Phe	L1-THAP dom.		0.71	0.06	-2.00	0.33		No impact	82	Pathogenic
c.23A>G	p.Tyr8Cys	L1-THAP dom.		0.71	0.08	-2.00	0.38		PASSC [73.14]	76	Pathogenic
c.25G>T	p.Gly9Cys	L1-THAP dom.		0.71	0.02	-3.00	0.46		No impact	93	Pathogenic
c.50A>T	p.Asp17Gly	L1-THAP dom.		0.71	0.05	-1.00	0.29		No impact	79	Pathogenic
c.68A>C	p.His23Pro	BS1-THAP dom.		0.5	0.00	-2.00	0.25		No impact	88	Pathogenic
c.77C>G	p.Pro26Arg	L2-THAP dom.	DNA binding -	0.79	0.02	-2.00	0.17	-SC35 [2.73]	No impact	100	Pathogenic
c.86G>C	p.Arg29Pro	L2-THAP dom.	DNA binding -	0.79	0.04	-2.00	0.17		PASSC [75.5]	100	Pathogenic
c.89C>G	p.Pro30Arg	L2-THAP dom.		0.79	0.02	-2.00	0.17		No impact	87	Pathogenic
c.95T>A	p.Leu32His	L2-THAP dom.	L2-THAP dom	0.79	0.01	-3.00	0.33		No impact	100	Pathogenic
c.115G>A	p.Ala39Thr	AH1-THAP dom.	L2-THAP dom	0.79	0.19	0.00	0.29	-SF2/ASF [2.48]	No impact	88	Pathogenic
<b>c.161G&gt;A*</b>	p.Cys54Tyr	BS2-THAP dom	C2CH motif.	0.79	0.01	-2.00	0.38		No impact	99	Pathogenic
c.215T>G	p.Leu72Arg	L4-THAP dom.		0.79	0.13	-2.00	0.25		No impact	76	Pathogenic
c.224A>T	p.Asn75Ile	L4-THAP dom.		0.71	0.02	-3.00	0.33		No impact	93	Pathogenic
c.247T>C	p.Cys83Arg			0.86	0.35	-3.00	0.25		No impact	82	Pathogenic
c.395T>C	p.Phe132Ser			1	0.77	-2.00	0.33		No impact	76	Pathogenic
c.408C>G	p.Asn136Iys		HCF-1 binding	1	0.06	0.00	0.46		No impact	76	Pathogenic
c.410A>G	p.Tyr137Cys		HCF-1 binding	1	0.10	-2.00	0.38		No impact	88	Pathogenic
c.446T>C	p.Ile149Thr	NLS		0.93	0.02	-3.00	0.42		No impact	87	Pathogenic
c.449A>C	p.His150Pro	NLS		0.93	0.24	-2.00	0.25		No impact	76	Pathogenic
c.508T>C	p.Cys170Arg	Coiled-coil dom.		0.86	0.00	-3.00	0.25		No impact	94	Pathogenic
c.1A>G	p.Met1?	THAP dom.		0.71	0.00	1.00	0.71		No impact	65	Probably pathogenic
c.16T>C	p.Ser6Pro	L1-THAP dom.		0.71	0.20	-1.00	0.38		No impact	71	Probably pathogenic
c.36C>A	p.Asn12Lys	L1-THAP dom.		0.71	0.22	0.00	0.46		No impact	65	Probably pathogenic
c.86G>A	p.Arg29Gln	L2-THAP dom.	DNA binding -	0.79	0.06	1.00	0.50		PASSC [71.63]	65	Probably pathogenic
c.176C>T	p.Thr59Ile	THAP dom.		0.79	0.07	-1.00	0.42		No impact	65	Probably pathogenic
c.407A>G	p.Asn136Ser		HCF-1 binding	1	0.13	1.00	0.63		PASSC [85.88]	71	Probably pathogenic
c.496G>A	p.Ala166Thr	Coiled-coil dom.		0.93	0.42	0.00	0.29		No impact	65	Probably pathogenic
<b>c.38G&gt;A*</b>	p.Arg13His	L1-THAP dom.		0.71	0.01	0.00	0.58		No impact	64	Probably pathogenic
c.169C>A	p.His57Asn	AH2-THAP dom.	C2CH motif	0.79	0.05	1.00	0.58		No impact	59	Probable polymorphism
<b>c.241T&gt;C*</b>	p.Phe81Leu	AH4-THAP dom.	AA interactions	0.79	0.45	0.00	0.71		No impact	59	Probable polymorphism
<b>c.61T&gt;A*</b>	p.Ser21Thr	L1-THAP dom.		0.71	0.00	1.00	0.79		No impact	47	Polymorphism
c.266A>G	p.Lys89Arg			0.79	0.34	2.00	0.96		PASSC [72.98]	35	Polymorphism
c.427A>G	p.Met143Val	Coiled-coil dom.		0.93	1.00	1.00	0.71		No impact	35	Polymorphism
c.506G>A	p.Arg169Gln	Coiled-coil dom.		0.86	0.18	1.00	0.50		No impact	47	Polymorphism
c.559C>A	p.Gln187Lys	Coiled-coil dom.		0.86	1.00	1.00	0.54		PASSC [78.26]	41	Polymorphism
c.574G>A	p.Asp192Asn			0.93	0.43	1.00	0.75		No impact	35	Polymorphism

The UMD-predictor<sup>®</sup> tool computes all elements (structure, biochemistry, splicing, and conservation) and provides for each of them a specific strength based on its relative impact.

“Conservation”: The score ranges from 0 for nonconserved to 1 for fully conserved.

“SIFT”: (sorting intolerant from tolerant): uses sequence homology to predict whether an amino acid substitution will affect protein function and, hence, potentially confer a phenotype [Ng and Henikoff, 2003]. Scores that are ≤0.05 are considered to be deleterious (in bold).

“BIOSUM62” is an amino acid substitution matrix based on local multiple alignments of an unselected protein set of related sequences [Henikoff and Henikoff, 1992]. Positive scores are associated with conservative changes and negative scores (in bold) with less conservative changes.

“Biochemical value” is an amino acid substitution matrix depending on 48 qualitative physicochemical properties describing side chain structure and functional groups, optical properties, hydrophobicity/charge/acid–base properties, and size (volume and side chain length) [Yu, 2001]. A value below 0.05 defines an invalid substitution concerning the physico-chemical properties (in bold).

“ESE” (exonic splicing enhancer) and “Splice site” modifications are evaluated by the UMD tool.

“PASSC”: potential acceptor splice site created; potential deleterious modifications are shown in bold.

“Pathogenicity”: the UMD-predictor<sup>®</sup> tool computes all the elements (structure, biochemistry, splicing, and conservation) providing for each a specific strength based on their relative impact. Predictions are normalized on a scale from 0 to 100.

In “Conclusion”: a value of less than 50 is associated with the prediction of a nonpathogenic mutation annotated as “polymorphism”; a value of 50–64 is associated with the prediction of a “probable polymorphism”; a value of 65–74 is associated with the prediction of a “probably pathogenic” mutation; while a value above 74 is associated with the prediction of a “pathogenic” mutation. For “Structure” and “HCD” descriptions see the legends to Table 1. For more details see Frederic et al. [2009].

\*These mutations have been studied in vitro [Fuchs et al., 2009; Gavarni et al., 2010; Kaiser et al., 2010].

and have to be determined by searching, among others, for the presence of quasi-palindromic sequences, inverted repeats, or symmetric elements that facilitate the formation of secondary-structure intermediates [Krawczak and Cooper, 1991].

## Splice-Site Mutations

Only one splice site variation is described in the literature: c.72-4T>C (or c.IVS1-4T>C) in association with the missense mutation c.77C>G [Houlden et al., 2010]. The splice site variation has been analyzed with the UMD algorithm implemented in HSF tool [Desmet et al., 2009]. The software computed the variation of the consensus value (CV) of the mutant “splice site” (CV = 87.25) relative to the wild-type site (CV = 86.59). As the variation between these values was not higher than 10%, it is highly probable that such CV variation does not disrupt the wild-type splice site and thus this variation corresponds to a polymorphism as suggested also by the authors [Houlden et al., 2010]. The pathogenic mutation is thus likely to be c.77C>G and consequently the c.72-4T>C (or c.IVS1-4T>C) polymorphism was associated to the c.77C>G mutation in the database.

## Conclusion

The UMD-*THAP1* database contains 53 different mutations (Table 1) in 56 probands and 43 relatives. These mutations are mainly private, essentially missense, usually nonrecurrent, and widely distributed in the *THAP1* domain and the rest of the *THAP1* gene. The currently reported *THAP1* mutations are mainly missense mutations (64.9%, 37/57) and small out-of-frame deletions (19.3%, 11/57), with a minority of other mutation types (7% of nonsense [4/57], 3.5% of small in-frame deletions [2/57], 1.8% of small out-of-frame insertions [1/57], and 3.5% of complex mutations). To date, no clear genotype/phenotype relationship has been identified. This is not surprising when considering past experience with other disease genes, such as the *Fibrillin-1* gene (*FBN1*), for which many mutations were accumulated before the genotype/phenotype relationships emerged. Indeed, the UMD-*FBN1* database was created in 1995 [Collod et al., 1996], and the first report about the correlations between genotype and phenotype was published in 2007 [Favre et al., 2007] and more followed [Detaint et al., 2010; Favre et al., 2007, 2008, 2009a, b, c; Stheneur et al., 2009]. From this perspective, the UMD-*THAP1* locus-specific database will facilitate the mutational analysis of *THAP1* gene as previously demonstrated for other genes.

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