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A Phenotype of Atypical Apraxia of Speech in a Family Carrying SQSTM1 Mutation

Claire Boutoleau-Bretonnière^{a,b,c,*}, Agnès Camuzat^{d,e,f}, Isabelle Le Ber^{d,e,f,g,h}, Kawtar Bouya-Ahmed^{d,e,f}, Rita Guerreiroⁱ, Anne-Laure Deruet^{a,b}, Christelle Evrard^{a,b}, José Brasⁱ, Estelle Lamy^{a,b}, Elisabeth Auffray-Calvier^j, Amandine Pallardy^k, John Hardyⁱ, Alexis Brice^{d,e,f}, Pascal Derkinderen^{a,b,l}, and Martine Vercelletto^{a,b}

^aCHU Nantes, Centre de Mémoire et de Ressource et Recherche (CM2R), Nantes, France

bInserm, CIC 04, Nantes, France

^cLaboratoire d'études des mécanismes cognitifs, EA 3082, Université Lyon 2, Bron F-69500, France

dInserm, UMR_S1127, CRICM, F-75013, Paris, France

eUPMC Univ Paris 06, UMR S975, F-75013, Paris, France

^fCNRS UMR 7225, F-75013, Paris, France

⁹AP-HP, Hôpital de la Pitié-Salpêtrière, Centre de Référence des Démences Rares, F-75013, Paris, France

^hAP-HP, Hôpital de la Pitié-Salpêtrière, Département de Neurologie, F-75013, Paris, France

Department of Molecular Neuroscience, UCL Institute of Neurology, London, UK

^jService de Neuroradiologie, CHU de Nantes, Nantes, France

^kService de Médecine Nucléaire, CHU de Nantes, Nantes, France

¹Inserm, UMR 913, Nantes, France

Abstract

SQSTM1 mutations, coding for the p62 protein, were identified as a monogenic cause of Paget disease of bone and of amyotrophic lateral sclerosis. More recently, SQSTM1 mutations were identified in few families with frontotemporal dementia. We report a new family carrying SQSTM1 mutation and presenting with a clinical phenotype of speech apraxia or atypical behavioral disorders, associated with early visuo-contructional deficits. This study further supports the implication of SQSTM1 in frontotemporal dementia, and enlarges the phenotypic spectrum associated with SQSTM1 mutations.

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^{*}Correspondence to: Claire Boutoleau-Bretonnière, Centre Mémoire Recherche et Ressources (CMRR), Clinique Neurologique CHU Nord, 44093 Nantes, France. Tel.: +33 240 165 422; Fax: +33 240 165 397; claire.boutoleau-bretonniere@chu-nantes.fr.

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Keywords

Amyotrophic lateral sclerosis; apraxia of speech; behavioral variant of FTD; frontotemporal lobar degeneration; non fluent variant of primary progressive aphasia; p62; Paget disease of bone; progressive non-fluent aphasia; progressive supranuclear palsy; SQSTM1

INTRODUCTION

Frontotemporal degeneration (FTD) defines a heterogeneous group of disorders that are interrelated. Clinical consensus criteria defined three variants according to the presenting symptoms: progressive behavioral changes in the behavioral variant of FTD (bvFTD), and prominent linguistic impairment in agrammatic (nonfluent/agrammatic primary progressive aphasia, PNFA) and semantic variants [1, 2]. The definite diagnosis is based on three major pathological subtypes of frontotemporal lobar degeneration (FTLD) characterized by the presence of TDP43 (FTLD-TDP), tau (FTLD-tau), or FUS-positive neuronal inclusions [3].

Thirty percent of FTD patients have a positive family history of the disease with an autosomal dominant inheritance. Mutations in the *SQSTM1* gene, coding for the p62 (sequestosome 1) protein, were initially identified as a cause of Paget's disease of bone (PDB) and, recently, as a cause of familial amyotrophic lateral sclerosis (ALS) [4]. More recently, *SQSTM1* mutations were also identified in few families with autosomal dominant bvFTD [5, 6]. We described here a new clinical phenotype associated to *SQSTM1* mutation.

PATIENTS AND METHODS

Patients

A French family has been studied (Fig. 1). Three siblings developed FTD at age ranging from 70 to 75 (patients 007, 008, 009). Their parents died at the age of 58 and 44 without known neurological disease. Family history was censored in both parental branches.

Molecular analyses

Blood samples of the three patients (007, 008, 009) were obtained; all participants signed an informed consent for genetic studies. This study was approved by the local research Ethics Committee (Ethics Committee from *Assistance-Publique Hôpitaux de Paris*). DNA was extracted from blood samples. Mutations in known FTD (*C90RF72*, *MAPT*, *PGRN*, *VCP*, *CHMP2B*) and ALS genes (*S0D1*, *TARDBP*, *FUS/TLS*, *PFN1*, *UBQLN2*, *hnRNPA2/B1*, *hnRNPA1*, *MATR3*) were initially excluded by Sanger sequencing or by repeat-primed PCR (for *C90RF72*). The whole exome was sequenced on DNA samples of the three patients. Briefly, genomic DNA of patients was prepared according to Illumina's TruSeq Sample Preparation v3 (Illumina, CA, USA) and sequence capture, enrichment, and elution was performed according to manufacturer's protocols (Illumina's TruSeq Exome Enrichment). Sequencing was performed on Illumina's HiSeq2000 using 100 bp paired-end reads. Sequence alignment and variant calling was performed against the reference human genome (UCSC hg19) using bwa and the Genome Analysis Toolkit. Polymorphisms found in the public databases and in 50 in-house exomes of controls were excluded. Secondarily,

mutations identified in *SQSTM1* gene were validated by Sanger sequencing according to previously described methods [6]. The entire coding sequence was also sequenced in 350 French age-matched healthy controls.

RESULTS

Molecular analyses

A heterozygous mutation, c.714 716delGAA, p.Lys238del, was identified in the exon 5 of *SQSTM1* gene in the three affected siblings (007, 008, 009). The mutation was validated by the Sanger method in the three patients (Fig. 1). The Lys238 residue is conserved across species. The mutation was previously identified as a disease-causing mutation in ALS patients [4] and was predicted to be deleterious by SIFT/PROVEAN *in silico* software (score: –11.7). Furthermore, this mutation was absent from 350 French controls, from 6,503 individuals of the 'exome variant server' (http://evs.gs.washington.edu/EVS/), dbSNP (http://www.ncbi.nlm.nih.gov/projects/SNP/) and 1000Genomes (http://www.1000genomes.org/) databases, thus strongly supporting its deleterious effect.

Clinical features

Patients' neuropsychological scores are summarized in Table 1.

Proband 007

This 77-year-old right-handed patient was evaluated in our Department after two years of language difficulty. He had endoprothetic treatment of the left hip at 67 years old and a right femoral fracture at the age of 75, without any obvious radiological or biological signs of PDB. Examination showed apraxia of speech characterized by an articulation planning deficit, dysarthria with hypokinetic features and buccofacial apraxia, associated with semantic and phonetic paraphasia. Phonemic and semantic fluency sores were low (respectively 12 and 2 words in two minutes). Syntactic and single-word comprehensions were spared. Mild grammatical deficit appeared in his spoken picture description and in writing. Neuropsychological testing showed executive dysfunction associated with severe visuo-constructional impairment when copying the Rey figure but preservation of memory (Fig. 2, Table 1). Careful interview of the patient's caregiver revealed the absence of behavioral disorders. Deep tendon reflexes were increase in all limbs and plantar reflex was flexor bilaterally. He had no motor symptoms, no gaze limitation, and no parkinsonian signs. Brain magnetic resonance imaging (MRI) showed major diffuse cortical atrophy more marked on the frontal and temporal lobes. ECD single photon emission computed tomography (SPECT) of the brain revealed severe hypoperfusion of frontal and temporal lobes, predominantly on the right side, and hypoperfusion of basal ganglia (Fig. 3). Five years later, the patient displayed severe phonetic disintegration with worsening of agrammatism but comprehension was spared. There were no clinical or electromyographic signs of ALS.

Sibling 008

This 71-year-old right-handed patient developed apragmatism over several months, without others behavioral troubles. On initial examination, he had a moderate dysarthria due to

buccofacial apraxia. There were no signs of upper or lower motor neurons dysfunctions. Neuropsychological testing showed executive dysfunction associated with visuoconstructional deficit, without aphasia. Brain MRI showed major diffuse atrophy more prominent on the frontal and parietal lobes. Brain SPECT revealed severe hypoperfusion of frontal and parietal lobes, particularly in bilateral sulcus centralis (Fig. 3). Eight years after the disease onset, the clinical features were highly consistent with apraxia of speech associated with visuo-constructional deficit. Executive functions also deteriorated (Table 1), but autonomy was still preserved with an overall slow progression.

Sibling 009

This 74-year-old right-handed patient presented with visual and auditory hallucinations by the age of 72. He had bilateral hip fractures at the age of 67 caused by falls, but had no obvious clinical, biological, or radiological signs of PDB. Neurological examination was normal. Neuropsychological testing showed preserved memory functions with severe executive syndrome and visuo-constructional disabilities (Fig. 2). CT scan showed bilateral frontal atrophy. Brain ECD-SPECT revealed cortical frontal and temporal hypoperfusion (Fig. 3). Severe behavioral disorders subsequently appeared, especially self-mutilation that required hand amputation. Cognitive deficit worsened over time leading to the diagnosis of bvFTD. The patient was finally institutionalized and died at 80 years.

DISCUSSION

We describe a new family carrying a heterozygous p.Lys238del mutation in *SQSTM1* gene segregating with FTD. This study further demonstrates the implication of *SQSTM1* in FTLD spectrum of diseases. Clinically, the three patients of the family were affected by atypical forms of FTD. Two patients had hip and femoral fractures, but none presented severe bone symptoms and deformations. The patients were not investigated specifically with biological testing or bone scintigraphy but no radiological signs of PDB were present on radiography. Furthermore, although the p.Lys238del mutation has been previously identified in ALS patients [4], motor neuron symptoms were absent in the three patients. ALS patients are often not tested for FTD symptoms, which could be the case for the ALS patient with the same mutation. This family thus illustrates the variability of bone and motor phenotype that can be moderate or absent in *SQSTM1* families. We propose that *SQSTM1* gene should be integrated in the 'multisystem proteinopathies' (MSP4), an emerging group of genetic diseases clinically characterized by variable association of FTD, ALS, PDB, and myopathy, and so far including *VCP* (MSP1), *hnRNPA2/B1* (MSP2), and *hnNPA2* genes (MSP3) [7].

The three patients exhibited a similar neuropsychological profile associating dysexecutive syndrome and early visuo-constructional disabilities. The episodic memory was for a long time preserved. The patients were mainly apathetic and one initially presented psychotic symptoms with hallucinations at onset. Brain imaging showed major opercular and perisylvian involvement. None of the three patients completely fit with the clinical criteria of bvFTD of Rascovsky [2] at onset. Patient 009 was the only one to fulfil these criteria after several years of evolution.

More interestingly, patient 007 presented apraxia of speech, with buccofacial apraxia dysarthria and anomia at onset. The patient 008 also had moderate symptoms of apraxia of speech. Apraxia of speech can be the initial sign of PNFA [2]. However, even if two patients presented with language problems, both of them presented early visuo-constructional disorders and, therefore, do not completely fit with the diagnosis of primary progressive aphasia according to the criteria of Gorno-Tempini et al. [2, 8]. They also showed different neuroimaging patterns, and neither of them presents the typical neuroimaging of nfv-PPA [2, 8].

Many genetic cases of FTD do not present with the typical features of sporadic FTD, or do not strictly meet the criteria for any of the clinical variants. Notably, in previous studies, all FTD patients carrying *SQSTM1* mutations presented a behavioral variant of FTD [5, 6]. This family demonstrates that the phenotypes associated with *SQTM1* mutations are more heterogeneous, and that this gene should also be studied in patients presenting with speech apraxia or behavioral disorders associated with visuo-constructive deficit at onset, even in absence of PDB or ALS.

Predicting the underlying pathology in patients with FTD is difficult but will be particularly important for future treatments targeting tau or TDP-43 proteins. A pathological study previously showed that *SQSTM1* mutation carriers had TDP-43 and p62-immunoreactive neuronal inclusions in the prefrontal cortex [9]. So far, PNFA and more precisely, apraxia of speech, was rather considered predictive of FTLD-tau [10–12]. Our study supports a possible link between speech apraxia and FTLD-TDP. This finding is supported by a recent pathological study in 11 patients showing that nonfluent/agrammatic primary progressive aphasia could be associated with FTLD-TDP pathology [13].

In conclusion, our report widens the phenotypic spectrum of *SQSTM1* mutations, and further illustrates the variability of phenotypes associated with *SQSTM1* mutations thus supporting a pleiotropic effect of the mutations.

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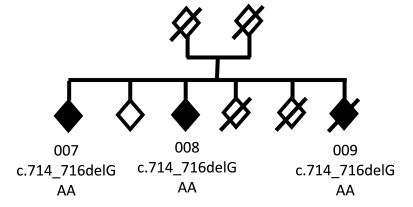


Fig. 1. Pedigree. The individuals are represented by diamonds for confidentiality. Black diamonds: FTD; white diamonds: non symptomatic individuals. Individuals 0007, 008, and 009 carried the c.714_716delGAA mutation.

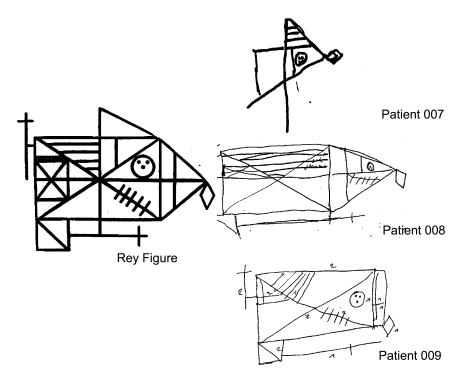


Fig. 2. Copy of Rey figure of the patients carrying *SQSTM1* mutations

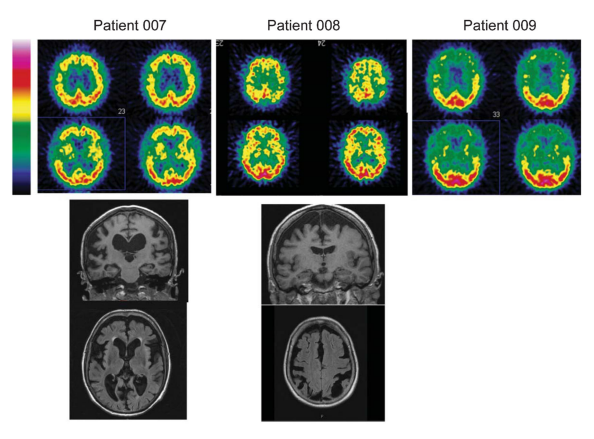


Fig. 3. Brain imaging. ECD-SPECT (top) and brain MRI (coronal and axial sections, bottom) of patients 007, 008, and 009.

Table 1

Neuropsychological scores of the patients

	Patient 007	Patient 008		Patient 009		
Age at evaluation, y	77	82	71	79	74	76
Disease duration, y	2	7	1	9	2	4
MMS/30	23	14	24	24	18	11
FCSRT						
Encoding/16	12*	13*	16	13*	14*	NA
Free recall 1,2,3/16	5/6/7	4/3/4*	6/9/9	6/7/8	5/5/9	NA
Cued recall 1,2,3/16	11/10/9	6/12/8	10/7/7	6/7/8	10/10/7	NA
Total recall/48	48	37*	48	42	46	NA
Recognition/16	16	16	16	16	16	NA
Delayed total recall	15	NA	NA	16	15	NA
Rey figure Copy/36	NA	NA	19	NA	17	18
Oral confrontation naming/80	69 [*]	61*	80	73	75	73
Verbal fluency 2'						
P letter	12*	NA	23	20	10*	9*
Animals	2*	NA	20	30	6*	4*
Frontal Assessment Battery/18	5*	5*	15*	16	9*	3*
Wisconsin Card Sorting Test						
Criteria/6	1*	NA	2*	2*	NA	NA
Errors	21*	NA	20*	13*	NA	NA
Perseverations	3*	NA	9*	5*	NA	NA
TMT A	NA	NA	C0.75	C50 - 0.75	C0.75	C0.75
TMT B	NA	NA	Impossible*	Impossible*	Impossible*	Impossible*

MMS, Mini Mental Score; FCSRT, Free and Cued Selective Recall Reminding Test; TMT, Trail Making Test; NA: not available,

^{*} abnormal value.