Dissecting the role of the mitochondrial chaperone mortalin in Parkinson’s disease: functional impact of disease-related variants on mitochondrial homeostasis

Lena F. Burbulla¹,²,³, Carina Schelling¹,², Hiroki Kato⁴, Doron Rapaport⁴, Dirk Woitalla⁷, Carola Schiesling¹,², Claudia Schulte¹,², Manu Sharma¹,², Thomas Illig⁸, Peter Bauer⁵, Stephan Jung⁶, Alfred Nordheim⁶, Ludger Schöls¹,², Olaf Riess⁵ and Rejko Krüger¹,²,*

¹DZNE, German Center for Neurodegenerative Diseases, Tübingen, Germany, ²Department of Neurodegenerative Diseases, Hertie-Institute for Clinical Brain Research, ³Graduate School of Cellular & Molecular Neuroscience, ⁴Interfaculty Institute for Biochemistry, ⁵Medical Genetics and ⁶Proteome Center Tübingen, Interfaculty Institute for Cell Biology, University of Tübingen, Tübingen, Germany and ⁷Department of Neurology, St Josef-Hospital, Ruhr-University Bochum, Bochum, Germany and ⁸Institute of Epidemiology, Helmholtz Zentrum München, German Research Center for Environmental Health, Neuherberg, Germany

Received July 21, 2010; Revised August 20, 2010; Accepted August 26, 2010

The mitochondrial chaperone mortalin has been linked to neurodegeneration in Parkinson’s disease (PD) based on reduced protein levels in affected brain regions of PD patients and its interaction with the PD-associated protein DJ-1. Recently, two amino acid exchanges in the ATPase domain (R126W) and the substrate-binding domain (P509S) of mortalin were identified in Spanish PD patients. Here, we identified a separate and novel variant (A476T) in the substrate-binding domain of mortalin in German PD patients. To define a potential role as a susceptibility factor in PD, we characterized the functions of all three variants in different cellular models.

In vitro import assays revealed normal targeting of all mortalin variants. In neuronal and non-neuronal human cell lines, the disease-associated variants caused a mitochondrial phenotype of increased reactive oxygen species and reduced mitochondrial membrane potential, which were exacerbated upon proteolytic stress. These functional impairments correspond with characteristic alterations of the mitochondrial network in cells overexpressing mutant mortalin compared with wild-type (wt), which were confirmed in fibroblasts from a carrier of the A476T variant. In line with a loss of function hypothesis, knockdown of mortalin in human cells caused impaired mitochondrial function that was rescued by wt mortalin, but not by the variants. Our genetic and functional studies of novel disease-associated variants in the mortalin gene define a loss of mortalin function, which causes impaired mitochondrial function and dynamics. Our results support the role of this mitochondrial chaperone in neurodegeneration and underscore the concept of impaired mitochondrial protein quality control in PD.

INTRODUCTION

Parkinson’s disease (PD) is the second most common neurodegenerative disorder after Alzheimer’s disease. Although for the majority of patients the underlying cause of the disease is still unknown, existing data suggest that genetic susceptibility factors acting together with environmental risk factors are contributing to the sporadic form of the disease. Following
linkage studies in rare familial forms of PD and the screening of large samples of sporadic PD patients, to date, 16 genetic disease loci have been identified including several genes that allowed the first insight into molecular pathways leading to neurodegeneration (1,2). The identification of PD patients with variants in nuclear encoded mitochondrial proteins was the first genetic support for various biochemical findings that had previously implicated impaired mitochondrial function in PD pathogenesis (3–5). A specific and selective loss of mitochondrial complex I activity in the substantia nigra of PD patients reflects an important role of mitochondrial pathology in PD (6). Furthermore, mitochondrial homeostasis plays a crucial role in aging and programmed cell death. Nevertheless, the intramitochondrial signaling pathways involved in cellular stress response and initiation of cell death mechanisms are currently poorly understood.

Variants in the DJ-1 gene have established an important link between mitochondrial impairment and the pathogenesis of PD. Oxidation of DJ-1 and its subsequent translocation to the mitochondrial matrix were identified as crucial for the maintenance of mitochondrial homeostasis (7–9). DJ-1 encodes a mitochondrial protein that acts as a sensor of cellular oxidative stress and exerts a crucial role in protecting cells against stress-induced cell death (8). Known loss-of-function DJ-1 variants decrease the protective capacities against neuronal cell death and can play a critical role in the susceptibility to neurodegeneration (10,11).

Recently, the mitochondrial heat shock protein mortalin (also known as GRP75, mthsp70 or PBP74) was identified as a novel mitochondrial DJ-1-interacting protein, also involved in the oxidative stress response (12,13). Mortalin is a 679 amino acid protein that has been found in multiple subcellular localizations such as the endoplasmic reticulum, cytoplasmic vesicles and the cytosol (14,15). However, the majority of mortalin is located within the mitochondrial matrix. The protein reaches this location after its import via the translocases of the mitochondrial outer and inner membranes (16,17). Moreover, mortalin also takes an active role in the import of other proteins via the translocases of the mitochondrial inner membrane channels. It has been identified as the only ATPase component of the preprotein mitochondrial import complex and is therefore essential for effective export of nuclear encoded proteins into mitochondria (18,19). Interestingly, in the brain, mortalin primarily localizes to neurons, but is observed in glial cells upon pathological activation (20–22).

As a lifespan-regulating protein and a member of the Hsp70 family of chaperones, mortalin is also involved in the regulation of cellular senescence and immortalization. Lifespan-regulating proteins directly affect mitochondrial function, including energy metabolism and reactive oxygen species (ROS) production (23,24). Importantly, stress response and aging are recognized as major risk factors for neurodegenerative diseases such as PD (23,25–27). Impaired mitochondrial function is critically linked to imbalanced dynamic fusion and fission events of mitochondria and to energetic depression, which may subsequently result in the activation of programmed cell death mechanisms. Overexpression of mortalin leads to an extended lifespan in nematodes and in human cells (28,29). On the other hand, it serves as a major target for oxidation and was shown to be involved in aging of the human brain, including PD (30). Since mortalin interacts with many proteins (31), its modifications in response to oxidation-related stress and damage are likely to generate pleiotropic effects (32).

Recently, the following observations suggested a role for mortalin in PD pathogenesis: (i) mortalin binds the PD-associated gene DJ-1, (ii) it plays an important role in the maintenance of mitochondrial homeostasis as critically related to neurodegeneration and (iii) coding variants in the mortalin gene in PD patients have been identified. Considering these associations, we first performed a mutational screening in a large sample of German PD patients, followed by functional mutation analyses including all PD-associated mortalin variants. Our observation that the loss of mortalin performance can cause impaired mitochondrial function and morphology provides a mechanism for the pathogenic role of mortalin in PD.

RESULTS

Unbiased proteomic approach identifies mortalin as a binding partner of DJ-1

Using a glutathion S-transferase (GST) pull-down assay with recombinant GST-tagged DJ-1 protein, we screened for DJ-1-interacting proteins in lysates from human dopaminergic neuroblastoma cells (SK-N-BE). Selected protein bands on a Coomassie gel were in-gel-digested and submitted to LC-ESI-MS/MS analysis. In comparison with publicly available databases, we identified mortalin as a DJ-1-interacting protein in lysates from human dopaminergic SK-N-BE cells (Fig. 1). Processing mass spectrometry data by using OpenMS and Mascot search engines, we identified mortalin as a DJ-1-interacting protein, with a probability-based mouse score of 139 indicating a high degree of homology (33). This interaction was subsequently confirmed by co-immunoprecipitation of wild-type (wt) DJ-1-myc and mortalin-V5 in HEK293 cells (data not shown). Therefore, our results confirm previous studies using either a candidate approach or an alternative
proteomic approach, i.e. stable isotope labeling by amino acids in cell culture, to define the interaction between DJ-1 and the mitochondrial chaperone protein mortalin (12, 13).

**Identification of a novel mortalin variant in German PD patients**

Initially, we performed a detailed mutational analysis of the coding region of the *mortalin* gene in a sample of 286 German PD patients (mean age: 65.1 ± 10.5 years; males: 51.9%, females: 48.1%). Using denaturating high-performance liquid chromatography (dHPLC) analysis of amplified PCR fragments for high-throughput mutational screening and subsequent direct sequencing, we were able to define sequence variations of German PD patients from that of a group of 290 healthy German controls. In total, we identified five sequence variations in the coding region of the *mortalin* gene. Most of them were silent base pair exchanges representing known polymorphisms without evidence of disease association (Table 1). However, a novel c.1426G>A variant in exon 12 was identified in one apparently sporadic PD patient, leading to an amino acid exchange from alanine to threonine in position 476 (Supplementary Material, Fig. S1A). The A476T variant is located in a highly conserved region of the protein sequence that forms part of the substrate-binding domain of the protein (Fig. 2A and B). This variant, which was identified in the heterozygous state in the carrier, was not present in 580 chromosomes of population and age-matched healthy controls.

Consequently, we screened an independent cohort of 1008 German PD patients (mean age of patients at onset 56.0 ± 12.1 years; males: 59.0%, females: 41.0%) and 1342 population-based matched controls (mean age of controls at examination 55.6 ± 11.7 years; males: 58.7%, females: 41.3%) for the identified novel variant in the German population and two recently identified *mortalin* variants from Spanish PD patients [c.376C>T and c.1525C>T; (34)]. The latter variants were each identified in individual Spanish PD patients and affect highly conserved regions of the protein, which was discovered by an interspecies comparison (Fig. 2C and D).

The variants identified in Spanish PD patients were absent in our cohorts, as all patients and controls were homozygous for the respective wt allele. The heterozygous A476T exchange was subsequently identified in four additional PD patients and six individuals from the population-based controls. Interestingly, four of the control individuals carrying the A476T variant

**Table 1. Frequency of the identified polymorphisms in the coding region of the mortalin gene in patients and controls**

<table>
<thead>
<tr>
<th>Polymorphism</th>
<th>Detection method</th>
<th>Patients</th>
<th>Controls</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ala81Ala (exon 4)</td>
<td>dHPLC, sequencing</td>
<td>2.1% (6/286)</td>
<td>2.6% a</td>
</tr>
<tr>
<td>Glu258Gln (exon 8)</td>
<td>dHPLC, sequencing</td>
<td>3.15% (9/286)</td>
<td>4.8% a</td>
</tr>
<tr>
<td>Lys316Lys (exon 9)</td>
<td>dHPLC, sequencing</td>
<td>54% (85/157)</td>
<td>42% (116/276)</td>
</tr>
<tr>
<td>Leu645Leu (exon 16)</td>
<td>dHPLC, sequencing</td>
<td>46% (129/279)</td>
<td>45% (85/191)</td>
</tr>
</tbody>
</table>

a[http://www.ensembl.org/Homo_sapiens/Transcript/ProtVariations?db=core; g=ENSG00000113013;r=5:137918923-137939014;t=ENST00000297185](http://www.ensembl.org/Homo_sapiens/Transcript/ProtVariations?db=core; g=ENSG00000113013;r=5:137918923-137939014;t=ENST00000297185).
were reported with previously defined extrapyramidal symptoms that did not fulfill the diagnosis of PD (mean age 72.6 ± 5.3 years), whereas the other two asymptomatic carriers of the A476T variant were significantly younger (mean age 44.5 ± 9.5 years). Collectively, this leads to a frequency of the A476T variant of 0.0069 in individuals with extrapyramidal symptoms and a frequency of 0.0012 in healthy controls. We speculated that the A476T substitution may act as a risk factor for PD upon aging and therefore included it together with the two Spanish PD-associated variants in our functional studies.

**Clinical phenotype of A476T variant**

The mean age of disease onset in carriers of the A476T variant was 53 years (SD ± 11.7). All patients presented with typical dopa-responsive Parkinsonism and displayed no family history leading to the diagnosis of idiopathic PD. The most frequent genetic cause of sporadic PD in Caucasians, the G2019S mutation in the LRRK2 gene, was excluded in all asymptomatic carriers of the A476T variant in the mortalin gene (35). One patient, whose full medical records were available, showed first symptoms at the age of 61 with general slowing of movements, shuffling of gait, intermittent tremor of both upper limbs and reduced olfactory sense. Magnetic resonance imaging revealed slight microcalcifications in the basal ganglia as occasionally observed in asymptomatic persons; however, there is no evidence for a symptomatic Parkinsonian syndrome, i.e. due to vascular lesions. The diagnosis of PD was made by a neurologist experienced in movement disorders. The symptoms responded well to dopaminergic therapy. At the age of 63, the patient reported worsening of motor symptoms with additional difficulties in concentration and dysarthria so that dopaminergic treatment was increased up to a levodopa-equivalent dose of 850 mg/day. No levodopa-related complications were reported, specifically no levodopa-induced psychosis occurred. Neuropsychological testing revealed a cognitive decline beginning at the age of 63. The patient was first tested at this age, when he reported a decline in cognitive capacity. A follow-up test at the age of 66 confirmed mild cognitive deficits without progression over time. Seven years after the diagnosis of PD at the age of 66, the patient unexpectedly died of cardiac failure following an acute myocardial ischemia.

The mother of this patient suffered from a stroke at the age of 85 and died about 10 years later due to cardiac insufficiency. The father suffered from cardiac asthma and died at an early age.

The A476T variant was also identified in one child of the index patient aged 49 years that was clinically unaffected at the time of the neurological examination. Furthermore, we excluded the variant in another child of this patient that displayed no signs of motor impairment at an examination at the age of 51 (Supplementary Material, Fig. S1B and C).

**Wt mortalin and mortalin variants in HEK293 and SH-SY5Y cells display intact mitochondrial targeting**

Mortalin is a nuclear encoded mitochondrial protein that requires posttranslational transport across the mitochondrial membranes to reach the mitochondrial matrix. To confirm the ability of all mortalin variants to be imported to mitochondria, we studied their subcellular localization by immunocytochemistry in HEK293 cells as well as in SH-SY5Y cells transiently transfected with wt mortalin or one of the variants. Using immunocytochemical techniques, we found mitochondrial localization of the recombinant protein for wt mortalin and all three variants in HEK293 cells (Supplementary Material, Fig. S3) and in SH-SY5Y cells (Supplementary Material, Fig. S4). These results cannot exclude the possibility that mortalin variants are only associated on the surface of mitochondria rather than being inside the mitochondrial matrix. To test the import capacity of the variants in more detail, we performed an in vitro import assay employing radiolabeled mortalin variants and mitochondria isolated from HeLa cells. The results demonstrate that both, the final efficiency and the kinetic of the import of the variant proteins, are not altered compared with that of wt mortalin (Fig. 3, lanes 2–5). As expected, in the presence of the respiratory chain uncoupler, carbonyl cyanide m-chlorophenylhydrazone (CCCP), which acts as a mitochondrial stressor, neither wt nor variant mortalin proteins were imported into mitochondria (Fig. 3, lane 6, asterisk).

Taken together, our results suggest that, as observed previously for wt mortalin (16), the mutated versions are also efficiently imported into mitochondria.

**Cells overexpressing mortalin variants or wt mortalin have differential effects on intramitochondrial ROS levels**

Dysfunction of mitochondria is reflected by impaired oxidative phosphorylation, which leads to the generation of ROS. Mortalin as a mitochondrial chaperone can protect the organelle against such ROS production. Therefore, we monitored ROS levels in dopaminergic SH-SY5Y cells stably overexpressing wt, A476T, R126W or P509S mortalin and in cells transfected with an empty vector as a control. The intramitochondrial ROS production was quantified by flow cytometry using MitoSOX Red, a fluorogenic dye that is targeted to the mitochondria and is readily oxidized by superoxide species. ROS production was monitored under basal conditions and under the induction proteolytic stress (Fig. 4A). Under basal conditions, analysis of MitoSOX Red fluorescence demonstrated significantly lower amounts of ROS in...
cells overexpressing wt mortalin compared with control cells (P < 0.05; Fig. 4A, left panel). This reduction of ROS was less pronounced for all mortalin variants, with significantly different levels observed for P509S mortalin-overexpressing cells compared with wt mortalin-overexpressing cells.

Next, we examined whether the application of the proteasomal inhibitor MG-132 to SH-SY5Y cells stably overexpressing wt mortalin or mortalin variants might modulate the intramitochondrial ROS production. When cells were subjected to proteolytic stress induced by MG-132 (Fig. 4A, right panel),
both, cells overexpressing one of the variants of mortalin and control cells, showed a significantly increased MitoSOX Red fluorescence after treatment compared with cells overexpressing wt mortalin \( P < 0.01 \) for A476T mortalin; \( P < 0.001 \) for R126W mortalin, P509S mortalin and control). Of note, the different variants of mortalin do not protect SH-SY5Y cells from proteolytic stress as efficiently as the wt protein.

**Cells overexpressing mortalin variants or wt mortalin have differential effects on mitochondrial membrane potential**

Next, we examined changes in mitochondrial membrane potential (MMP) as a further marker of impaired mitochondrial function. MMP was examined by flow cytometry in SH-SY5Y cells stably overexpressing wt mortalin and its variants. Using tetramethylrhodamine ethyl ester perchlorate (TMRE) fluorescence to monitor the MMP, we were able to find significant differences in cells with altered MMP under basal conditions. In comparison with empty vector control, cells overexpressing wt mortalin showed a significantly improved MMP \( P < 0.01 \) (Fig. 4B, left panel). This change in the MMP was not observed for the mortalin variants showing similar MMP levels like the empty vector control.

Next, we examined whether proteolytic stress affects the MMP of the various cells (Fig. 4B, right panel). As seen above, treatment of cells with MG-132 resulted in a significant alteration of the MMP. These changes in the MMP were significantly less pronounced for cells overexpressing wt mortalin compared with empty vector control or cells overexpressing one of the mortalin variants \( P < 0.05 \).

**Wt mortalin but not mortalin variants rescues functional impairments in knockdown models**

To assess the effect of loss of mitochondrial mortalin on mitochondrial function, we measured MMP and ROS production by fluorescent-activated cell sorting (FACS) analysis in HEK293 cells depleted partially of their endogenous mortalin by an siRNA-mediated process. Mortalin protein was efficiently downregulated as demonstrated by immunoblotting (Fig. 6A). Densitometric analysis revealed a knockdown efficiency of endogenous mortalin protein of \( > 50\% \) compared with the status of cells treated with control siRNA (Fig. 6B). The levels of mortalin in these cells could be restored to \( > 80\% \) of the expression level in control cells.

We first tested whether siRNA-mediated knockdown of endogenous mortalin modifies the levels of ROS production and whether transfection of wt mortalin or disease-associated variants would restore the phenotype. For this, we utilized the FACS technique using MitoSOX staining. Knockdown of endogenous mortalin caused a significant increase of intramitochondrial ROS production compared with control siRNA-treated cells \( P < 0.01 \); Fig. 7A). We then investigated whether these functional abnormalities induced by mortalin knockdown could be restored by reintroduction of plasmid-encoding wt mortalin or one of the disease-associated variants. Reintroduction of wt mortalin, but not R126W mortalin or P509S mortalin, restored intramitochondrial ROS levels \( P < 0.05 \). Cells transfected with A476T mortalin showed a similar tendency; however, the statistic significance of the results was low \( P = 0.056 \).

Next, we measured the effect of mortalin knockdown on MMP by TMRE staining. FACS analysis revealed that transfection with mortalin siRNA resulted in a reduction of intact MMP in 25% of the HEK293 cells compared with cells transfected with non-targeting control siRNA \( P < 0.05 \); Fig. 7B). The impaired MMP after mortalin knockdown could be rescued by wt mortalin, but not by the R126W or P509S mortalin variants \( P < 0.01 \). Again, cells transfected with A476T mortalin showed a similar tendency; however, the changes in MMP were not significant \( P = 0.069 \). These observations are consistent with a loss of the physiological function in mortalin variants.

**Human fibroblasts from the carrier of the A476T variant display changes in mitochondrial morphology**

Finally, we extended our morphological studies of mitochondria to human fibroblasts. The mitochondrial morphology of A476T, R126W or P509S mortalin in SH-SY5Y cells is significantly less potent in changing mitochondrial morphology.
Figure 5. Effect of mortalin on mitochondrial morphology in SH-SY5Y cells. (A) Mitochondrial morphology in living SH-SY5Y cells transiently cotransfected with Mito-DsRed (red) and wt mortalin, its disease-associated variants or the empty vector in the ratio 1:4 was analyzed by live-cell-imaging microscopy (Cell Observer Z1, Zeiss) at 37 °C using ApoTome® optical slides with 0.240–0.300 μm z-stacks. Forty-eight hours after cotransfection, nuclei were stained with Hoechst 33342 (blue). The respective stainings were merged and mitochondria were analyzed using Image J 1.41o software for area, perimeter, major and minor axes based on binary images, which only have two possible values (black/white) for each pixel. On the basis of these parameters, the AR of a single mitochondrion and its FF were calculated. (B) Mitochondrial branching as indicated by the FF was significantly increased in SH-SY5Y cells cotransfected with Mito-DsRed and wt mortalin compared with cotransfection with empty vector or one of the mortalin variants (P < 0.001; left panel). Mitochondrial length as indicated by the AR was significantly increased in cells overexpressing wt mortalin compared with R126W mortalin or P509S mortalin (P < 0.001) as well as in cells transfected with the empty vector (P < 0.05; right panel). Images from 222 individual cells were analyzed from three independent experiments. Data are presented as mean ± SE of three independent experiments.
fibroblasts from a heterozygous carrier of the A476T variant and from a sibling control carrying the homozygous wt allele was monitored (Fig. 8A). We found that control fibroblasts exhibited large interconnected mitochondria, whereas fibroblasts from the carrier of the A476T variant were significantly shorter and fragmented (FF; $P < 0.001$ and AR; $P < 0.001$; Fig. 8B). Therefore, our analysis of human fibroblasts from the carrier of the A476T variant was monitored (Fig. 8A). We found that control fibroblasts exhibited large interconnected mitochondria, whereas the R126W variant affects the ATPase domain of mortalin, and P509S variants are located in the substrate-binding domain, which harbors the ATPase domain (46). The A476T and the P509S variants are located in the substrate-binding domain, whereas the R126W variant affects the ATPase domain of the protein. Irrespective of their localization, our results underscore a role for mortalin in the regulation of mitochondrial morphology, similar to our observation in SH-SY5Y cells overexpressing mortalin variants (Fig. 5).

**DISCUSSION**

Aging as a major risk factor for PD shares typical biochemical alterations with neurodegenerative processes due to impaired mitochondrial function, i.e. increased levels of oxidative damage and impaired cellular energy supply. Indeed, the role of unbalanced mitochondrial homeostasis in PD has been recently supported by the identification of variants in nuclear encoded proteins that are responsible for mitochondrial dysfunction and impaired dynamics, i.e. Parkin (PARK2), PINK1 (PARK6), DJ-1 (PARK7) and Omi/HtrA2 (PARK13) (43–40). The mitochondrial chaperone protein mortalin has been linked to PD pathogenesis based on reduced levels in affected brain regions of sporadic PD patients and its interaction with the PD-associated protein DJ-1 (12,30). Here, we confirm this interaction and provide first functional evidence for a direct contribution of aberrant mortalin to impaired mitochondrial function and dynamics in PD.

Using an unbiased proteomic approach to screen for DJ-1-interacting proteins, we confirmed previous reports of the interaction of DJ-1 with mortalin using independent proteomic techniques (12,13). Therefore, we hypothesized that mortalin may also act in signaling pathways responsible for mitochondrial stress response and the maintenance of mitochondrial integrity similar to that of the redox-sensing protein DJ-1 (8).

All currently identified disease-associated variants of the *mortalin* gene were observed in the heterozygous state. Although heterozygosity of these variants in the *mortalin* gene suggests an autosomal dominant effect, due to the lack of a reported family history in the respective patients, it is more likely a genetic trait with reduced penetrance and argues in favor of a genetic susceptibility factor for PD. Since the complete loss of mortalin function is not compatible with cell survival as observed in different models from yeast to vertebrates (41), heterozygous variants in the *mortalin* gene would be in line with the concept of a risk factor contributing to a late onset of neurodegenerative disease. Indeed, genetic studies for susceptibility factors in the common late onset form of PD showed that the sporadic appearance of the disease does not preclude an involvement of genetic factors in the pathogenesis of PD (42). Further support for a potential role of mortalin in PD pathogenesis is derived from a substantial number of genomic screens that have consistently shown a susceptibility locus for PD on the long arm of chromosome 5, which harbors the *mortalin* gene located at 5q31.1 (42–45).

First evidence for the pathogenic relevance of the novel variants in the mortalin protein came from interspecies comparisons of the mortalin amino acid sequence that showed complete conservation of the respective amino acid residues from vertebrates to yeast (Fig. 2B–D). The functional domains of the mortalin protein include the ATPase domain and the substrate-binding domain (46). The A476T and the P509S variants are located in the substrate-binding domain, whereas the R126W variant affects the ATPase domain of the protein. Irrespective of their localization, our results argue in favor of a loss of protective mortalin function in the mitochondria. In fact, we found that cells overexpressing one of the mortalin variants were more susceptible to increased mitochondrial oxidative stress compared with wt...
mortalin-overexpressing cells. Furthermore, using a knockdown model of mortalin in human cells, we found that the reported variants were unable to restore the observed mitochondrial phenotype, thus confirming the loss-of-function hypothesis. Indeed, the observed \(\sim 50\%\) knockdown of wt mortalin in these cells is compatible with a model of haploinsufficiency for mortalin function. This model is also in line with the mitochondrial phenotype, which we observed in fibroblasts from the heterozygous carrier of the A476T variant and is characterized by a reduced mitochondrial connectivity.

We have confirmed the mitochondrial localization of wt mortalin by an in vitro import assay and immunocytochemistry. All disease-associated mortalin variants showed the same in vitro import kinetics and these results are supported by our immunocytochemical studies in human cells. Thus, the various mutations in the variants do not affect the

Figure 7. Effect of downregulation of mortalin on mitochondrial function. (A) ROS production as indicated by the level of MitoSOX Red fluorescence signal was significantly increased in mortalin siRNA-transfected HEK293 cells relative to control siRNA-transfected cells \((P < 0.01)\). Transfection of siRNA-treated cells with wt mortalin showed a complete rescue of this mitochondrial phenotype. Rescue with A476T mortalin was partially achieved, whereas transfection with R126W or P509S mortalin variants failed to restore this mitochondrial phenotype \((P < 0.05\) compared with control siRNA-treated cells). (B) Measurement of TMRE red fluorescence signal revealed a significant increase of altered MMP in HEK293 cells transfected with mortalin siRNA compared with control \((P < 0.05\). Restoration of this signal could be observed by transfection of wt mortalin, but not with R126W or P509S mortalin variants. Partial rescue was observed by transfection of A476T mortalin. Data are indicated as mean \(\pm\) SD of three independent experiments.
The subcellular localization of the protein, indicating that the observed differences in the levels of ROS and MMP compared with wt mortalin were not due to defective import of the mortalin variants into mitochondria, but rather due to a loss-of-functional mortalin within the mitochondrial matrix. Indeed, due to their high metabolic activity, mitochondria are notably exposed to ROS and are at risk of oxidative damage to their DNA, lipids or proteins. Specifically for dopaminergic neurons that are reported to have a constitutively lower mitochondrial mass (47), the integrity of a functional protein clearance machinery is critical for mitochondrial function. In mitochondria, effective refolding of damaged proteins by chaperones or targeting for proteolytic cleavage is mediated by specific proteins, of which Omi/HtrA2 has been recently identified as an important part of the mitochondrial protein quality control (48). Mortalin was reported to be involved in protein refolding in the mitochondrial matrix compartment due to coupling with the mitochondrial Hsp60 protein (49). Moreover, the yeast homologue of mortalin is known to target misfolded proteins to mitochondrial proteases for degradation (50). Indeed, we found that overexpression of wt mortalin is protective in paradigms of proteolytic stress as assessed by intramitochondrial ROS and maintenance of the MMP. In contrast, the disease-related mortalin variants were devoid of this protective function and did not differ from controls.

Figure 8. Effect of mortalin on mitochondrial morphology in human fibroblasts. (A) Mitochondrial morphology in living human fibroblasts from a carrier of the A476T variant and a healthy sibling control were analyzed by live-cell-imaging microscopy at 37°C using ApoTome® optical slides with 0.240–0.300 z-stacks. Mitochondria were stained with 200 nM of the specific mitochondrial dye MitoTracker® green FM (green) for 15 min at 37°C; nuclei were stained with Hoechst 33342 (blue). The respective stainings were merged and mitochondria were analyzed using Image J 1.41o software for area, perimeter, major and minor axes based on binary images. The calculation of the AR and FF was done as described in Figure 5. (B) Mitochondrial branching as indicated by the FF was significantly reduced in fibroblasts from the carrier of the A476T variant in comparison with control fibroblasts ($P < 0.001$) (left panel). Mitochondrial length as indicated by the AR was significantly reduced in fibroblasts from the carrier of the A476T variant in comparison with control cells (right panel). Eighty-two individual cells were analyzed from three independent experiments. Data are indicated as mean ± SE of three independent experiments.
Therefore, we speculate that misfolding of mortalin and/or impaired interaction with intramitochondrial substrates and interaction partners might interfere with the protective role of mortalin in maintaining mitochondrial function. Indeed, increased proteolytic stress has been shown to predispose to misfolding of mortalin, resulting in defective protein import into mitochondria and mitochondrial morphology in yeast (51). More interestingly, the loss of function of Ssc1p, the yeast homologue of mortalin, resulted in the disintegration of the mitochondrial network and the formation of aggregates. This was not due to impairments of mitochondrial import, but rather due to the effects of the yeast mortalin homologue within the matrix (52). These results support our findings and reveal a similar morphological phenotype of loss of mortalin function in yeast and human cells. Therefore, our results underscore the crucial involvement of mortalin in the maintenance of mitochondrial morphology due to its function in the mitochondrial matrix.

Several proteome-based studies from postmortem brains of PD patients and controls revealed reduced levels of mortalin in affected brain regions (22,30,53). In addition, the amount of mortalin not only turned out to be a marker for sites of neurodegeneration in PD, but also a correlation of protein levels with disease progression was reported with a further decrease of mortalin in advanced stages of PD (53). Together with the loss of function variants in the mortalin gene described here in PD patients, the critical role of functional mortalin in neurodegeneration in PD becomes evident. Therefore, large genetic studies in different populations of PD patients and subsequent functional studies are required to further dissect the contribution of sequence variations in the mortalin gene to the pathogenesis of PD.

MATERIALS AND METHODS

Patients and controls
A detailed mutation analysis of the mortalin gene in a large sample of 286 German sporadic and familial PD patients was performed after obtaining informed consent. All patients were evaluated by a neurologist and were diagnosed as idiopathic PD, based on the UK Parkinson’s Disease Brain Bank criteria (54). The mean age of disease onset was 57.2 years (±11.3). Ethical approval was obtained by the Ethics Committee of the University of Tübingen. A total of 290 healthy German individuals closely matched for age and gender served as controls after a standardized neurologic examination to exclude participants with clinical signs of PD or any other extrapyramidal disorder (mean age: 72 ± 4.3 years; males: 52%, females: 48%).

The three mortalin variants A476T, R126W and P509S were subsequently genotyped in an independent cohort consisting of 1008 German sporadic PD patients and 1342 German population-based controls. The PD cases were collected by movement disorder specialists of the Universities of Munich and Tübingen, who established the diagnosis according to the UK Brain Bank criteria. The mean age at onset was 56.0 ± 12.1 years. All subjects gave written informed consent and the study including DNA collection was approved by the local ethics committees. The control individuals were selected from the KORA survey (Cooperative Health Research in the Region of Augsburg, www.helzt-muenchen.de/kora), a population-based study. The samples were genotyped using the matrix-assisted laser desorption/ionization time-of-flight mass spectrometry method (MassArray system, Sequenom).

PCR conditions and restriction length polymorphism analysis
Based on the published genomic sequence of the human mortalin gene on chromosome 5q31.1 (NCBI accession number NM_004134), we performed a detailed mutational screening of the coding sequence that includes 17 exons and adjacent intronic regions. Primers to amplify the respective sequences of the mortalin gene were generated using the online software Primer3 (http://frodo.wi.mit.edu/primer3/input.htm; Table 2). PCR reactions were carried out in a Thermocycler PTC Dyd 220 (MJ Research, Biozym, Germany) using the following conditions: 100–500 ng DNA were amplified in a final volume of 26 µl in the presence of PCR Buffer (Promega, USA), 200 µM of each dNTP, 10 pmol of each PCR primer and 0.5 units of Taq Polymerase (Promega). Standard cycling conditions were established using a touchdown protocol: 5 min at 94°C; 10 times: 30 s at 94°C, 30 s at 65°C (−1°C per cycle), 30 s at 72°C; 30–35 times: 30 s at 94°C, 30 s at 55°C, 30 s at 72°C; followed by a final elongation for 6 min at 72°C.

For the subsequent genotyping of the c.1426G>A variant in exon 12 of the mortalin gene, the entire PCR product was digested by adding 15 units of the restriction enzyme SatI (MBI Fermentas, USA) to the reaction mixture in a final volume of 45 µl. Restriction fragments were visualized on a 2% agarose gel stained with ethidium bromide.

dHPLC analysis and sequencing
Prepared DNA samples from PCR were screened for variants using dHPLC analysis on the WAVE™ DNA Fragment Analysis System (Transgenomic, USA). For the mutational screening, the resulting PCR products from two patients were pooled at a 1:1 ratio followed by a denaturation step at 94°C and subsequent renaturation over 15 min by cooling down the sample to 10°C gradually. This allows the formation of heteroduplexes even in the presence of homozygous sequence variations. Column temperatures and running conditions were established using Wavemaker™ Software (Version 4.1.31, Transgenomic) and are given in Table 2. DNA samples exhibiting curve patterns suspicious for heteroduplexes were sequenced on an ABI Prism™ Sequence Detecting System (3100 Genetic Analyzer) using the ABI BigDye® Terminator v3.1 Cycle Sequencing Kit (Applied Biosystems, USA) with primers used for exon amplification.

GST pull-down assay and mass spectrometry
Screening for novel interacting proteins of DJ-1 was performed by a GST pull-down assay. Recombinant GST-DJ-1 protein was produced by cultivation of Escherichia coli cells transformed with pGEX-6P-3-DJ-1. Cells were collected by
<table>
<thead>
<tr>
<th>Exon</th>
<th>Primer, forward</th>
<th>Primer, reverse</th>
<th>Product length (bp)</th>
<th>Annealing temperature (°C)</th>
<th>Column temperature (dHPLC) (°C)</th>
<th>Nucleotide substitution</th>
<th>Amino acid substitution</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>5′-TGG TTG GAG GTT TCC AGA AG-3′</td>
<td>5′-AAT TCA AAC CCT AAA GGG CG-3′</td>
<td>275</td>
<td>a</td>
<td>63.5; 66.6</td>
<td>c. –31T&gt;C; c.81+6G&gt;A</td>
<td>n</td>
</tr>
<tr>
<td>2</td>
<td>5′-TTC TCT TTT TCC TCC CAG GAT-3′</td>
<td>5′-TTC CCT CTC AAA GGA AAT GA-3′</td>
<td>152</td>
<td>a</td>
<td>56.0; 58.0</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>5′-TCC AGT GAC TTG GGT CTA TCA AGA-3′</td>
<td>5′-AAA GGG CAA ATC AGG TTC TC-3′</td>
<td>246</td>
<td>60</td>
<td>55.2; 57.4</td>
<td>c.243C&gt;T</td>
<td>Ala81Ala</td>
</tr>
<tr>
<td>4</td>
<td>5′-CTC TCT ATC ACA TTT TGG GAG TTT-3′</td>
<td>5′-CAT GCT GAG CCT CTT CTC TG-3′</td>
<td>278</td>
<td>a</td>
<td>58.2; 61.4</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>5′-TGG TAT GTG ATT TTG TCA TGG-3′</td>
<td>5′-CTA GTG ATC TCA CAG GAA TTA TCG-3′</td>
<td>259</td>
<td>a</td>
<td>54.6; 58.0</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>6–7</td>
<td>5′-GCT GCA AAA GGA TGA CAC AG-3′</td>
<td>5′-CAG TAA AAG CAC TGT AAA AGG CTC-3′</td>
<td>447</td>
<td>a</td>
<td>55.0; 57.3</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>5′-ATC CGT GTG ATA GGT TTT GTT C-3′</td>
<td>5′-AAG AGT ATC TGT GTC TAG AAT AAG GGG-3′</td>
<td>348</td>
<td>a</td>
<td>54.6; 58.0</td>
<td>c.744G&gt;A; c.879+80C&gt;T</td>
<td>Gln258Gln</td>
</tr>
<tr>
<td>9</td>
<td>5′-TGG GTT GTT CCA CCT TAT TAC TGC-3′</td>
<td>5′-TTT TAA ATG CCC TCT AGG GGC TG-3′</td>
<td>235</td>
<td>a</td>
<td>57.3; 59.2</td>
<td>c.948G&gt;A; c.880–53delITTA</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>5′-AAA TGT AAC CGT CAT TTG GC-3′</td>
<td>5′-GGG CAT ATA TTT GTG CCA CC-3′</td>
<td>350</td>
<td>55</td>
<td>57.8</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>11</td>
<td>5′-ACA GTG GCC TCT TGT CT-3′</td>
<td>5′-GGG TGC AAT TAC ATG CAG C-3′</td>
<td>368</td>
<td>a</td>
<td>56.5; 60.0</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>12</td>
<td>5′-TGG TAT GAT GTA AAC CTT TGG C-3′</td>
<td>5′-GAG TAT GTA TGT GTA TGC CAG GAC-3′</td>
<td>305</td>
<td>a</td>
<td>55.7; 57.7</td>
<td>c.1426G&gt;A; c.1411–16G&gt;T</td>
<td>Ala476Thr</td>
</tr>
<tr>
<td>13</td>
<td>5′-AAC ACT ATG AGA CCC CTC TG-3′</td>
<td>5′-GGG TCT ATT CCC AAG ACC TCC-3′</td>
<td>234</td>
<td>a</td>
<td>55.8; 57.8</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>14</td>
<td>5′-TGG TAT GTT GAT GGA AGA AGA GG-3′</td>
<td>5′-CCC AAA CTC CCA CTG TCA AG-3′</td>
<td>229</td>
<td>a</td>
<td>55.9</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>15</td>
<td>5′-TGG TAT GAT GTA AAC CTT TGG C-3′</td>
<td>5′-CCTA TCA TCA AAA CCC ACA GA-3′</td>
<td>225</td>
<td>a</td>
<td>56.2</td>
<td>n</td>
<td></td>
</tr>
<tr>
<td>16</td>
<td>5′-TGG TAT AGG AGA CCT GGA AAA-3′</td>
<td>5′-AAA ATC CAC TGG AGG CTT TG-3′</td>
<td>213</td>
<td>a</td>
<td>56.0; 58.4</td>
<td>c.1933C&gt;T</td>
<td>Leu645Leu</td>
</tr>
<tr>
<td>17</td>
<td>5′-AAC GTT AAA TCT GAG TGG CTC T-3′</td>
<td>5′-TGT TGT CCT TGC TGG TTC AA-3′</td>
<td>166</td>
<td>a</td>
<td>56.5; 58.5</td>
<td>n</td>
<td></td>
</tr>
</tbody>
</table>

"Touch-down protocol: 94°C → 5°; [94°C → 30°, 65°C → 30° (−1°C per cycle), 72°C → 30°] 10×; (94°C → 30°, 55°C → 30°; 72°C → 30°) 30×; 72°C → 6°; 8°C → ∞."
centrifugation and the pellet was resuspended using lysozyme and protease inhibitor cocktail Complete® (Roche, Germany). After complete ultrasonic lysis of the bacteria (cycle continued, output 70%, Bandelin Sonoplus GM70, Bandelin, Germany) and centrifugation, the supernatant contained the fusion proteins that were then bound to glutathione agarose (Molecular Probes, USA). After washing, the recombinant DJ-1 proteins were eluted from the agarose in several steps. Fractions of the elution that contained the recombinant protein were unified and the concentration was determined by a Bradford assay. Then 1 mg of enriched GST-DJ-1 fusion protein was incubated with lysates from SK-N-BE cells, and, as a control, the same cell lysate was incubated with glutathione agarose alone. Samples were run on a sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) and stained with Coomassie. Bands of interest underwent tryptic digestion. Resulting peptides were separated via reverse-phase liquid chromatography (Dionex Ultimate, Dionex, Idstein, Germany) coupled online to a mass spectrometer (QStar Pulsar i, Applied Biosystems, Germany) with the following settings: peptide and fragment mass tolerance 0.25, carbamidomethyl as fixed modifications and methionine oxidation as variable modifications. An in-house-curated database (IPI_human 3.56; 2009-03-19) containing reverse peptides as well as a list of typical contaminants was used for peptide identification with an expected cut-off score >0.05.

Cloning of wt mortalin and mortalin variants
cDNA of mortalin was obtained from the German Resource Center for Genome Research (RZPD; IRAUp969A034D) and cloned into the BamHI and XhoI sites of a pcDNA3.1/V5-His A vector (Clontech, USA) using 5′-CCAGCCAATGAGACGCTTG-3′ primer pairs were used for the insertion of the P509S variant: 5′-GTATTCTCTACTGCCACTGATGGTCAAACGCAA-3′. SH-SY5Y cells were cultured in a 5% CO2 humidified atmosphere in Dulbecco’s modified Eagle’s medium (DMEM) containing 1% penicillin and streptomycin and 15% fetal calf serum (FCS). Transfections were performed using Lipofectamine™ 2000 Reagent (Invitrogen, USA) according to the manufacturer’s instructions. To generate polyclonal stable cell lines, 700 000 SH-SY5Y cells were transfected with the pcDNA3.1/V5-His A vector containing wt mortalin or one of its variants (A476T, R126W, P509S) or an empty vector as a control and maintained in medium containing 1 mg/ml G418 (Biochrom, Germany) for selection. Stable expression of wt mortalin and mortalin variants was confirmed by immunoblotting.

Moreover, fibroblasts from offspring of the index patient carrying the A476T variant in the mortalin gene were included in our analyses. Skin biopsies were taken from one offspring carrying the heterozygous A476T variant as well as a sibling control carrying the homozygous wt allele. The carrier of the A476T variant did not show signs of PD at the time of the biopsy. The study was approved by the ethics committee of the University of Tübingen. All patients and controls gave written and informed consent. Primary fibroblast cell lines were maintained in RPMI medium with 15% FCS supplemented with 100 IU/ml penicillin, 100 mg/ml streptomycin and 1 mM pyruvate. For live-cell-imaging experiments, passage number of fibroblasts <10 were used. Only fibroblasts with the same passage number were taken for analyses.

Western blot analyses
WB analyses were conducted as described previously (4). In the present study, proteins were detected using antibodies against mortalin (anti-GRP75, Santa Cruz Biotechnology, USA), V5 (anti-V5, Chemicon, USA) and β-actin (anti-β-actin, Sigma, Germany). Secondary antibodies were purchased from GE Healthcare (UK) and either diluted 1:10 000 for anti-rabbit or 1:5000 for anti-mouse in Tris-buffered saline (TBS) with 0.1% Tween®20 (Roth, Germany). The densitometry analysis from WBs was performed using the Image J 1.41o software (Wayne Rasband; National Institutes of Health, USA).

Isolation of mitochondria from HeLa cells and in vitro import assay
HeLa cells were cultured in DMEM supplemented with 10% FBS, 50 μg/ml penicillin and 50 μg/ml streptomycin at 37°C under 5% CO2. The cells in a 10 cm culture dish were washed with phosphate-buffered saline (PBS) and collected by centrifugation at 800g for 5 min. From here on, all steps were performed at 4°C. The collected cells were resuspended in buffer A (20 mM HEPES–KO H, 220 mM Mannitol, 70 mM sucrose, 1 mM EDTA, 2 mg/ml BSA and 0.5 mM PMSF, pH 7.6) and were homogenized by being passed 20 times through a 27-gauge needle, and then this mixture was centrifuged at 800g for 5 min. The supernatant fraction was centrifuged at 10 000g for 10 min to obtain the mitochondrial fraction. The mitochondria were resuspended in 1 ml buffer B (20 mM HEPES–KO H, 220 mM Mannitol, 70 mM sucrose, 0.5 mM PMSF, pH 7.6) and were centrifuged again at 10 000g for 10 min. This mitochondria pellet was then used for in vitro import assays.

Radioabeled precursor proteins were synthesized by the TNT-coupled reticulocyte lysate system (Promega) in the presence of 35S-methionine. The assay mixtures containing...
precursor proteins and 25 μg mitochondria were incubated in 50 μl of import buffer (10 mM HEPES–KOH, 1 mM ATP, 20 mM sodium succinate, 5 mM NADPH, 0.5 mM magnesium acetate, 220 mM mannitol, 70 mM sucrose, 0.5 mM PMSF, pH 7.4) at 30°C for the indicated time in the absence (Fig. 3, lanes 2–5) or presence of CCCP (Fig. 3, lane 6, asterisk). To degrade non-imported proteins, samples were treated at the end of the reaction with proteinase K (100 μg/ml) for 30 min on ice. The mitochondria were then isolated by centrifugation and analyzed by SDS–PAGE followed by autoradiography.

Immunocytochemistry and immunofluorescence microscopy
For immunocytochemistry, HEK293 cells were seeded on collagen-coated slides and transiently transfected with either wt mortalin or mortalin variants or the corresponding empty vector. Forty-eight hours following transfection, cells were incubated for 15 min with 100 nM MitoTracker® Red CMX-Ros (Molecular Probes) at 37°C in a humidified 5% CO2 atmosphere to visualize mitochondria via fluorescence microscopy. Then cells were washed with PBS, fixed with 10% (wt/vol.) paraformaldehyde (PFA) in PBS at room temperature for 15 min and permeabilized at −20°C with ice-cold methanol for 5 min. After washing, cells were incubated for 30 min in 10% (vol./vol.) FCS in PBS for blocking and subsequent incubation with rabbit V5 antibody (1:300) and visualization with a fluorescein isothiocyanate-conjugated anti-rabbit secondary antibody (1:5000; Dianova, Germany). Cell nuclei were stained with Hoechst 33342 (1:10 000; Molecular Probes). After a 3-fold wash step in PBS, the coverglass was mounted in Mowiol (Sigma). Mitochondrial localization of wt mortalin and mortalin variants was investigated using an epifluorescence microscope (Axioplan 2, Zeiss, Germany) with an ApoTome technique. Three slices with intervals between 0.230 and 0.280 μm were taken and the series of pictures were saved uncompressed. To determine the degree of colocalization, images were loaded into Axiovision 4.6 software and evaluated accordingly (Zeiss).

Live cell imaging
Mitochondrial morphology was analyzed by live cell imaging. SH-SY5Y cells or human fibroblasts were cultured in Lab-TekHII chambered coverglasses (number 155382; Nalge Nunc International). Mitochondria of fibroblasts were stained with 200 nM MitoTracker® Red CMX-Ros (Molecular Probes) for 20 min and washed twice with PBS. MMP was measured after incubation in a medium containing 100 nM TMRE for 30 min and washed twice with PBS. For proteolytic stress conditions, cells were treated with 10 μM protease inhibitor MG-132 (Sigma-Aldrich, Germany) for 16 h before harvesting the cells. For each measurement, at least 30 000 cells were analyzed for the corresponding fluorescence on a CyAnTM ADP apparatus (DakoCytomation) using the 488 nm argon laser and emission through the PE filter (575 nm). All experiments were performed in duplicate and repeated at least three times.

RNA interference-mediated downregulation of mortalin and retransfection with wt mortalin or mortalin variants
Mortalin gene silencing was established by transfection of mortalin HP GenomeWide siRNA or chemically unmodified non-targeting control siRNA from Qiagen (Germany) into human cells. As an efficient knockdown of mortalin in SH-SY5Y cells was linked with substantial cell death, either by the use of siRNA or by shRNA lentiviral transduction particles, we chose HEK293 cells as an alternate human cell line.

For the analysis of MMP and ROS production in cells with knockdown of endogenous mortalin and subsequent rescue experiments, 200 000 HEK293 cells were seeded in a 12-well plate and cultured in a 5% CO2 humidified atmosphere in DMEM containing 10% FCS, 1% penicillin and streptomycin. After 4–5 h, cells were transfected with 80 pmol of mortalin siRNA or non-targeting control siRNA using HiPerFect transfection reagent (Qiagen), leading to a reduction in mortalin protein of up to 58% in cells transfected with mortalin siRNA compared with control cells. For rescue experiments, 24 h after transfection with siRNA, cells were again transfected with 4 μg of DNA by using jetPEI transfection reagent according to the manufacturer’s instructions (Peqlab Biotechnologie GmbH, Erlangen, Germany). Following 24 h, cells were analyzed via a FACS method to determine the MMP or levels of ROS production as described above. Downregulation

Analysis of mitochondrial morphology
Fluorescence microscopy images were optimized by adjusting the contrast and subsequently binarized by conversion to 8 bit images. After reduction of unspecific noise of the fluorescence signal, a spatial filter (convolution filter) as well as a threshold was applied to the images to define mitochondrial structures. Using Image J 1.41o software, every single mitochondrion of the investigated cells was marked to analyze morphological characteristics such as its area, perimeter, major and minor axes. On the basis of these parameters, the AR of a mitochondrion (AR; ratio between the major and the minor axes of the ellipse equivalent to the object) and its FF (perimeter2/(4π×area)), consistent with the degree of branching, were calculated (58).
of mortalin and complementation efficiency were monitored by immunoblotting with the mortalin antibody.

Statistical analysis

Genetic data were evaluated for allele frequencies and genotype frequencies using the GenePop program as described previously (59).

Functional data were analyzed by a Student’s t-test; all statistical tests were two-sided, and those with a P-value < 0.05 were considered to be statistically significant. Data are expressed as means ± standard deviation (SD) or means ± standard error (SE) values.

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG online.

ACKNOWLEDGEMENTS

We are grateful to Dr Radhika Puttagunta for valuable comments during the preparation of the manuscript.

Conflict of Interest statement. None declared.

FUNDING

This work was supported by grants from the German Research Council (DFG, KR2119/3-2 to R.K. and RA1028/4-1 to D.R.); the Federal Ministry for Education and Research (BMBF, 01GS08134 to R.K. and O.R.); the Faculty of Medicine, University of Tübingen (Fortuene 1709-0-0 to R.K.) and from the charitable Hertie Foundation (Ph stipend award to L.F.B.). The KORA research platform (KORA: Cooperative Research in the Region of Augsburg; http://www.gsf.de/KORA) was initiated and financed by the German Federal Ministry of Education, Science, Research and Technology and by the State of Bavaria.

REFERENCES
