The MitCHAP-60 Disease Is Due to Entropic Destabilization of the Human Mitochondrial Hsp60 Oligomer

Avital Parnas‡, Michal Nadler§, Shahar Nisemblat‡, Amnon Horovitz§, Hanna Mandel¶, and Abdussalam Azem‡

Department of Biochemistry, George S. Wise Faculty of Life Sciences, Tel Aviv University, 69778 Tel Aviv, the Department of Structural Biology, Weizmann Institute of Science, Rehovot 76100, and the Rappaport Faculty of Medicine and Research Institute, Technion-Israel Institute of Technology and Metabolic Disease Unit, Rambam Health Care Campus, Haifa 31096, Israel

Received for publication, June 10, 2009, and in revised form, August 9, 2009. Published, JBC Papers in Press, August 25, 2009, DOI 10.1074/jbc.M109.031997

The 60-kDa heat shock protein (mHsp60) is a vital cellular complex that mediates the folding of many of the mitochondrial proteins. Its function is executed in cooperation with the co-chaperonin, mHsp10, and requires ATP. Recently, the discovery of a new mHsp60-associated neurodegenerative disorder, MitCHAP-60 disease, has been reported. The disease is caused by a point mutation at position 3 (D3G) of the mature mitochondrial chaperonin, mHsp10, and requires ATP. Recently, the discovery of the homologous bacterial protein in Escherichia coli (Magen, D., Georgopoulos, C., Bross, P., Ang, D., Segev, Y., Goldsher, D., Nemirovski, A., Shahar, E., Ravid, S., Luder, A., Heno, B., Gershoni-Baruch, R., Skorecki, K., and Mandel, H. (2008) Am. J. Hum. Genet. 83, 30–42) of the mitochondrial Hsp60 protein, which renders it unable to complement the deletion of the homologous bacterial protein in Escherichia coli (Magen, D., Georgopoulos, C., Bross, P., Ang, D., Segev, Y., Goldsher, D., Nemirovski, A., Shahar, E., Ravid, S., Luder, A., Heno, B., Gershoni-Baruch, R., Skorecki, K., and Mandel, H. (2008) Am. J. Hum. Genet. 83, 30–42). The molecular basis of the MitCHAP-60 disease is still unknown. In this study, we present an in vitro structural and functional analysis of the purified wild-type human mHsp60 and the MitCHAP-60 mutant. We show that the D3G mutation leads to destabilization of the mHsp60 oligomer and causes its disassembly at low protein concentrations. We also show that the mutant protein has impaired protein folding and ATPase activities. An additional mutant that lacks the first three amino acids (N-del), including Asp-3, is similarly impaired in refolding activity. Surprisingly, however, this mutant exhibits profound stabilization of its oligomeric structure. These results suggest that the D3G mutation leads to entropic destabilization of the mHsp60 oligomer, which severely impairs its chaperone function, thereby causing the disease.
spastic paraplegia SPG13, a neurodegenerative disorder associated with two independent mutations in the gene encoding mHsp60 (22, 23). Recently, a large kindred including 23 patients suffering from MitCHAP-60 disease, an autosomal recessive neurodegenerative disorder, has been identified (24). Magnetic resonance imaging of the brains of the patients showed diffuse hypomyelination and leukodystrophy, in which myelin is not formed properly. The disease-causing mutation was identified as a homozygous missense mutation in the human HSPD1 gene encoding the mHsp60 protein (24), namely D3G in the mature protein. Initial studies showed that, in contrast to wild-type mHsp60, the mutant, together with mHsp10, was not able to fully complement a deletion of the bacterial homologues, GroEL and GroES, in E. coli (24). The mechanism by which the D3G mutation may compromise the function of the mHsp60 has not been reported. In this study, we suggest that the D3G mutation impedes the function of mHsp60 by entropic destabilization of the oligomeric structure of the molecule.

**EXPERIMENTAL PROCEDURES**

**Proteins**—mHsp10 from mouse (25) carrying a hexahistidine tag was purified using a nickel agarose column in 25 mM potassium phosphate buffer (pH 8) with 100 mM NaCl and 5 mM imidazole and then further purified using a gel filtration column (Superdex 200, GE Healthcare) equilibrated with 50 mM Tris-HCl (pH 7.7) buffer containing 300 mM NaCl, 10 mM MgCl₂, and 5% glycerol (supplemental Fig. 1S). GroEL was purified as described previously (26).

**Purification of mHsp60**—The nucleotide sequence encoding the mature human mHsp60 was amplified using PCR and inserted between restriction sites for BamHI and NotI in a modified pET21d (27). The engineered plasmid overexpresses mHsp60 containing an octa-histidine tag at its amino terminus that can be removed after treatment with tobacco etch virus protease. The Rosetta™ E. coli strain (Novagen) was used as an overexpression host for all constructs in this study. The complete purification protocol will be described elsewhere. Briefly, mHsp60 monomers were first purified using a nickel-agarose column. Following removal of the histidine tag and further purification on an ion exchange column, oligomers were reconstituted as described previously (28) (supplemental Fig. 1S). Protein concentrations mentioned in this study refer to monomer concentrations and were determined using the bicinchoninic acid kit (Sigma catalog number BCA1). Due to cloning constraints, the various constructs used in this study contain minor differences at their amino termini. Fig. 1 details the various constructs examined in this study and the sequence of their amino termini.

**RESULTS AND DISCUSSION**

**Cross-linking of mHsp60 Oligomers**—Cross-linking with glutaraldehyde was carried out to verify the oligomeric state of mHsp60. For this purpose, a given concentration of mHsp60 was incubated with 0.1% glutaraldehyde for 10 min at 25 °C in buffer A (50 mM Na-HEPES (pH 7.5) 100 mM KCl, 30 mM MgCl₂) or buffer B (100 mM Na-HEPES (pH 7.5), 10 mM KCl, 10 mM MgCl₂). The cross-linking reaction was stopped by the addition of 3× SDS sample buffer containing 1 μl urea and boiled for 5 min. The cross-linking products, containing 9 μg of protein, were separated on 2.4–12% acrylamide gradient SDS-PAGE and stained with Coomassie Blue.

**Protein Folding Assay**—The refolding assay of HCl-denatured MDH was carried out as described previously (29).

**ATPase Assays**—Initial (steady-state) rates of ATP hydrolysis were measured by monitoring time-resolved changes in fluorescence emission at 465 nm of coumarin-labeled phosphate-binding protein upon excitation at 430 nm using an ISS PCI spectrofluorometer (ISS, Inc.) (30, 31). The reactions were carried out in 50 mM Tris- HCl buffer (pH 7.5) containing 10 mM MgCl₂, 10 mM KCl, and 1 mM diithiothreitol at 25 °C.

**FIGURE 1. Amino-terminal sequences of chaperonins used in this study.** The mutated amino acid that causes the MitCHAP-60 disease is marked in boldface type. The underlined letters signify amino acids that have been added due to cloning constraints. WT, wild type.
D3G mutant was completely dissociated into monomers, whereas a significant fraction of the wild-type mHsp60 was found as heptamers. As shown in Fig. 4, the dissociation of the D3G oligomer is also time-dependent. At 10 μM protein, complete dissociation of the D3G mutant was reached after 60 min, at which time only ~30% of the wild-type protein was monomeric (Fig. 4A). At a protein concentration of 2.5 μM, dissociation of the oligomer to monomers was much more rapid. Ten minutes after dilution of proteins, all the D3G protein was found as monomers, whereas ~30% of the wild-type protein was still oligomeric even after a 3-h incubation (Fig. 4B). Purified wild-type mHsp60 and the D3G mutant exhibited similar CD spectra (supplemental Fig. 4S), which suggests that the observed rapid dissociation of the D3G oligomers is not due to a drastic change in the secondary structure of the protein. We conclude that one main effect of the D3G mutation on mHsp60 is destabilization of the oligomeric structure of the protein.

The D3G Mutant Exhibits Reduced ATPase and Protein Refolding Activities—The function of chaperonin 60 requires hydrolysis of ATP that is dependent upon the oligomeric structure of the protein for optimal function. We, therefore, examined the ATPase activity of the D3G mutant when compared with that of the wild-type mHsp60. Taking into consideration the above mentioned observed effect of protein concentration on the oligomeric state of the protein, we examined the ATPase activity at different protein concentrations (Fig. 5A). Wild-type mHsp60 was found to have ATPase activity with a rate constant of 0.44 min⁻¹. This rate was the same at all protein concentrations examined (from 1 to 30 μM) and independent of protein concentration. Notably, the ATPase activity of the D3G mutant was found to differ from that of the wild-type protein in (i) its significantly lower rate and (ii) the dependence of the rate on the protein concentration. Even at high protein concentrations, when the D3G mutant was expected to assemble into oligomers, its ATPase activity was only ~40% of that observed for...
the wild-type protein. This result suggests that the D3G mutation severely impairs the ATPase activity of mHsp60 in addition to destabilizing the oligomeric structure of the protein.

We next asked how the altered structural and ATP hydrolysis properties of the D3G mutant affect its ability to refold denatured proteins. For this purpose, we examined the chaperone properties of D3G and wild-type mHsp60 using MDH as a model substrate (Fig. 6). When HCl-denatured MDH was refolded with the help of either GroEL or wild-type mHsp60, in the presence of mHsp10, the yields of refolding after 2 h were 82 and 88%, respectively. The refolding yield in the presence of the D3G mutant and mHsp10 was found to be smaller (∼60%). Notably, although the refolding yield that is obtained with GroEL in the absence of co-chaperonin and ATP was close to background levels and did not increase over time (not shown), the yields obtained with wild-type mHsp60 and the D3G mutant, in the absence of ATP and mHsp10, were higher and increased with increasing incubation time. Thus, after a 2-h incubation, the yields in the case of the wild type and the D3G mutant were 25 and 30%, respectively. The latter result can be explained by a “minichaperone effect” (35, 36) due to the instability of the oligomeric structures of the D3G mutant and, although to a lesser extent, of the wild-type mHsp60. In other words, the chaperonins are able to bind unfolded proteins, but upon their dis-oligomerization, the unfolded substrate is released into solution in a form that is able, in vitro, to refold spontaneously. This explanation is supported by the observation that in the presence of ATP, which stabilizes the oligomeric state of mHsp60, the spontaneous refolding did not increase with increasing incubation time. Finally, another difference between the D3G mutant and wild-type mHsp60 is related to the rate of their protein folding activity. In the presence of both ATP and mHsp10, the maximum yield in the case of the wild-type protein was reached after 15 min, whereas in the case of the D3G mutant, it was reached after ∼45 min. In summary, the defective protein folding activity of the D3G mutant is reflected in both the yield and the rate when compared with the wild-type mHsp60.

Deletion of the First Three Amino Acids of mHsp60 Stabilizes the Oligomeric State of mHsp60 but Impairs Its Refolding Activity—The most obvious explanation for the effect of the D3G mutation on the structural stability of mHsp60 is that Asp-3 participates in interactions that are important for the stabilization of the oligomeric structure of the D3G mutant and, although to a lesser extent, of the wild-type mHsp60. In other words, the chaperonins are able to bind unfolded proteins, but upon their dis-oligomerization, the unfolded substrate is released into solution in a form that is able, in vitro, to refold spontaneously. This explanation is supported by the observation that in the presence of ATP, which stabilizes the oligomeric state of mHsp60, the spontaneous refolding did not increase with increasing incubation time. Finally, another difference between the D3G mutant and wild-type mHsp60 is related to the rate of their protein folding activity. In the presence of both ATP and mHsp10, the maximum yield in the case of the wild-type protein was reached after 15 min, whereas in the case of the D3G mutant, it was reached after ∼45 min. In summary, the defective protein folding activity of the D3G mutant is reflected in both the yield and the rate when compared with the wild-type mHsp60.

![FIGURE 5. Steady-state ATPase activities of mHsp60 variants. A, initial rate measurements of ATP hydrolysis by wild-type mHsp60 ( ), D3G ( ), and N-del ( ) mutants were performed at different concentrations of mHsp60 and a constant concentration of 2 mM ATP. The deviation from linearity observed for the D3G mutant reflects its dissociation into monomers at low protein concentrations. B, the effect of mHsp10 on the initial rates of ATP hydrolysis by wild-type (WT) mHsp60 and the D3G and N-del mutants was also determined and is presented as a percentage of change relative to the rate in the absence of mHsp10. These reactions were carried out in the presence of 30 μM mHsp60, 2 mM ATP, with or without 30 μM mHsp10.](image)

![FIGURE 6. Time-dependent refolding activity of mHsp60 variants. Refolding of HCl-denatured MDH by GroEL (black diamonds), wild-type mHsp60 (blue squares), D3G (red circles), and N-del (green triangles) was carried out in the presence (filled symbols) or absence (empty symbols) of ATP and mHsp10. Red plus signs indicate refolding of MDH by the D3G mutant in the presence of ATP but without co-chaperonin.](image)
latter explanation is correct, then the deletion should not lead to destabilization of the mHsp60 oligomer, and its chaperone activity will be maintained. If, however, the former explanation is correct, then the deletion should result in a destabilizing effect, as in the case of the D3G mutation. The results presented in Fig. 2 show clearly that deletion of the first three amino acids of mHsp60 (N-del) creates a very stable oligomer, even when compared with the wild-type protein. Cross-linking products corresponding to subheptameric forms of the protein are barely observed. The N-del oligomer was also found to be more resistant to heat-induced aggregation in comparison with wild-type mHsp60 and the D3G mutant (supplemental Fig. 3S) and to have a CD spectrum exhibiting slightly more secondary structure than the wild type and the D3G mutant (supplemental Fig. 4S). Interestingly, the yield of refolded MDH that is obtained with the N-del protein was close to that obtained with the D3G mutant (Fig. 6). In summary, we found that deletion of the first three amino acids in mHsp60 leads to unexpected stabilization of the protein but reduced chaperone activity (~50% of the wild-type activity), ruling out the possible participation of Asp-3 in stabilizing interactions.

The Deletion Mutant of mHsp60 Shows Unusual ATPase Activity—To gain more insight into the effect of the deletion mutant (N-del), we examined the ability of this mutant to hydrolyze ATP. When present alone, N-del was found to have a very low rate constant of ATP hydrolysis (Fig. 5A) of 0.036 min⁻¹, which is ~8 and 21% of those of the wild-type protein and the D3G mutant, respectively. Examining the ATPase activity of the wild type and the two mutants in the presence of mHsp10 yielded a very surprising result (Fig. 5B). In the case of the wild-type protein, the addition of mHsp10 resulted in 30% inhibition of its hydrolysis rate, whereas in the case of the D3G mutant, the ATPase activity was increased by 70% (although the co-chaperonin is expected to inhibit the ATPase activity by 50% (37)). Previous studies showed that mHsp10 stabilizes the oligomeric structure of the unstable mitochondrial and chloroplast chaperonins and, therefore, can cause an indirect activation of the ATPase activity as observed in the case of the D3G mutant (14, 29). Interestingly, the rate of ATP hydrolysis by the N-del mutant was enhanced by 65% when mHsp10 was included. As this mutant has a very stable oligomeric structure, this increase in ATPase activity cannot be explained by the stabilization effect of mHsp10 on the N-del mutant. How can deletion of the first three amino acids cause such a profound effect on the ATP hydrolysis properties of mHsp60? The allosteric regulation of ATP hydrolysis by mHsp60 requires a fine tuning of the strength of contacts between the subunits of the mHsp60 oligomer. Thus, it could be that the deletion leads to a very strong association between the subunits of the oligomer in a less active T state (38) that switches to the more active R state only upon the addition of ATP and mHsp10. Consequently, deletion of the three amino-terminus amino acids results in impaired ATP hydrolysis only in the absence of mHsp10.

Conclusions—The aim of this study was to determine the molecular basis of the MitCHAP-60 disease caused by the D3G mutation. For this purpose, we studied the structural and functional properties of the purified D3G mutant and compared them with those of wild-type mHsp60 and a deletion mutant, in which the first three amino acids were removed. We demonstrated that at high protein concentrations, the D3G mutant can assemble into oligomers (heptamers and tetradecamers). However, the D3G oligomers are unstable when compared with wild-type mHsp60 and dissociate into monomers rapidly upon dilution. Interestingly, the deletion mutant was found to be much more stable than both the wild type and the D3G mutant. Taken together, our results suggest that the D3G mutation leads to entropic destabilization of the oligomer that may cause the MitCHAP-60 disease. Given, however, that the very stable deletion mutant has altered ATPase and protein folding activities, it is likely that the D3G mutation also has allosteric effects that remain to be identified. In summary, we suggest that the D3G mutation impairs the protein folding activity of mHsp60 by its profound destabilization effect on the mHsp60 oligomer and by disrupting its ATPase function by a yet to be determined mechanism.

Another neurodegenerative disorder associated with a point mutation (V72I) in the gene encoding mHsp60 is hereditary spastic paraplegia SPG13 (22). However, in contrast to the recessive inheritance of the disease studied here, the SPG13 disease was also observed in a heterozygote background. How can this difference in the inheritance pattern be explained? The dominant negative effect of the V72I mutation suggests that subunits with the V72I mutation form stable oligomers and produce inactive chaperonin 60 complexes, even when incorporated into mixed oligomers with wild-type subunits (39). In contrast, in the case of the MitCHAP-60 disease, we hypothesize that the mutated subunits (carrying the D3G mutation) do not form stable homo- or hetero-oligomers; consequently, they are rapidly degraded in the cell. As a result, a sufficient amount of wild-type oligomers is formed in the mitochondria. An alternative explanation is that mixed oligomers do retain activity. Hence, no dominant negative phenotype is observed in a heterozygote background.

Acknowledgment—We thank Dr. Celeste Weiss-Katz for useful discussions.

REFERENCES
1. Levy-Rimler, G., Bell, R. E., Ben-Tal, N., and Azem, A. (2002) FEBS Lett. 529, 1–5
2. Landry, S. J., Taher, A., Georgopoulos, C., and van der Vies, S. M. (1996) Proc. Natl. Acad. Sci. U.S.A. 93, 11622–11627
3. Landry, S. J., Zielstra-Ryalls, J., Fayet, O., Georgopoulos, C., and Gierasch, L. M. (1993) Nature 364, 255–258
4. Hartl, F. U. (1996) Nature 381, 571–579
5. Horwich, A. L., Farr, G. W., and Fenton, W. A. (2006) Chem. Rev. 106, 1917–1930
6. Horovitz, A. (1998) Curr. Opin. Struct. Biol. 8, 93–100
7. Lin, Z., and Rye, H. S. (2006) Crit. Rev. Biochem. Mol. Biol. 41, 211–239
8. Horwich, A. L., Fenton, W. A., Chapman, E., and Farr, G. W. (2007) Annu. Rev. Cell Dev. Biol. 23, 115–145
9. Horovitz, A., and Willison, K. R. (2005) Curr. Opin. Struct. Biol. 15, 646–651
10. Cheng, M. Y., Hartl, F. U., Martin, J., Pollock, R. A., Kalousek, F., Neupert, W., Hallberg, E. M., Hallberg, R. L., and Horwich, A. L. (1989) Nature 337, 620–625
11. Ostermann, J., Horwich, A. L., Neupert, W., and Hartl, F. U. (1989) Nature 341, 125–130
Entropic Destabilization of the Hsp60 Oligomer