Structure of the human voltage-dependent anion channel

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The voltage-dependent anion channel (VDAC), also known as mitochondrial porin, is the most abundant protein in the mitochondrial outer membrane (MOM). VDAC is the channel known to guide the metabolic flux across the MOM and plays a key role in mitochondrially induced apoptosis. Here, we present the 3D structure of human VDAC1, which was solved conjointly by NMR spectroscopy and x-ray crystallography. Human VDAC1 (hVDAC1) adopts a β-barrel architecture composed of 19 β-strands with an α-helix located horizontally midway within the pore. Bioinformatic analysis indicates that this channel architecture is common to all VDAC proteins and is adopted by the general import pore TOM40 of mammals, which is also located in the MOM.

The outer membrane of mitochondria (MOM) contains three integral membrane protein families, two of which form channels as part of larger protein complexes (for review, see ref. 1). These two MOM complexes, the general import pore TOM and the SAM insertase, allow for the entire translocation and insertion of nearly all newly synthesized proteins destined to the mitochondrial organelle (2, 3). The third protein family of typically high abundance (~10,000 copies per mitochondrion) is termed voltage-dependent anion channels (VDACs), because of the voltage sensitivity of its open probability (4, 5). Together, this small number of protein families is sufficient for full communication between mitochondria with their cellular environment (1).

The VDAC channel was initially described as being reminiscent of bacterial porins and primarily responsible for the exchange of chemical energy equivalents between the cytosol and the mitochondrion (4, 6). Indeed, a variety of structural features (like barrel geometry and dimension) known from the bacterial precursors are maintained (7, 8). By contrast, a variety of functions have been ascribed to the VDAC isoforms among which the direct coupling of the mitochondrial matrix to the energy maintenance of the cytosol seems to be the most general function (9). The structure of VDAC is of interest because of a substantial body of evidence connecting VDAC to apoptosis. It is suggested that VDAC is a critical player in the release of apoptogenic factors from mitochondria of mammalian cells, and consequently several hypotheses describing the mechanism of mitochondria-mediated apoptosis involving VDAC have been proposed (for review, see ref. 10).

Results and Discussion

Structure Determination of hVDAC1: Combining NMR Spectroscopy and X-Ray Crystallography. In a parallel structural biology approach, we set out to characterize the structure of hVDAC1, the major isoform of this channel in mammalian tissues, by a combination of NMR spectroscopy and x-ray crystallography. The idea behind this project was to gain complementary structural information to have a solid basis for future studies, e.g., analysis of protein heterocomplex formation by NMR and crystal structures as a basis for drug target design. Only information derived from both methods and the application of an iterative structure calculation procedure allowed the structure determination of the protein (see Fig. 1).

To have large quantities of the channel available in manipulable form, we overexpressed the protein in Escherichia coli and refolded the channel to produce labeled protein for NMR and x-ray studies (11). For NMR studies, 15N-labeled hVDAC1 was refolded in detergents (LDAO or Cymal-5) and qualitatively analyzed by 13N-1H TROSY experiments. hVDAC1 refolded in LDAO showed ~250 well dispersed cross-peaks [supporting information (SI) Fig. S1A]. Using a combination of TROSY-type triple-resonance experiments, seven different amino acid specific 15N and 13N/13C-labeled samples, several 15N-resolved NOESY spectra and 20 different point mutations combined with paramagnetic spin labeling (SI Text, Fig. S2), the backbone resonances of 192 of the 282 residues of hVDAC1 (excluding the N-terminal methionine) were assigned. Ca secondary chemical shifts and slow hydrogen-deuterium exchange of amide protons revealed the topology of 19 β-strands with 65 interstrand NOEs unambiguously determining the register of all 19 β-strands (Fig. 1B and Fig. S3). At the N terminus, a combination of NMR parameters identified an α-helix that binds to the barrel wall (Fig. 1A and B and Table S1).

To obtain the crystal structure by x-ray diffraction, hVDAC1 was subjected to an extensive detergent screening. Native crystals were obtained with the hanging drop method by mixing 1.5-μl protein–Cymal-5 detergent solution containing 5–15 mg/ml protein with 0.75 μl of reservoir solution (30% PEG 400: 0.1 M Na Hepes, pH 7.5; 0.2 M magnesium chloride). The trigonal crystals (space group P321) diffraacted up to 4 Å (12). Multilength wavelength anomalous diffraction phasing with data from Se-Met crystals was used to approach the phase problem. Together with phases from a Pt-derivative and after careful density modification, an initial low-resolution electron density map was obtained that clearly showed the protein envelope and the overall barrel dimensions. This initial density was further improved through enforcing of a barrel-like solvent envelope leading to an electron density that allowed the tracing of eight β-strands and the N-terminal helix (see Fig. 1E).

Both normal and B-factor sharpened (13) density maps were used for model building. In general, the B-factor sharpened density maps


The authors declare no conflict of interest.

Data deposition: The coordinates and structure factors have been deposited in the Protein Data Bank (PDB ID code 2jx4).

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after density modification) gave a clearer indication of the position and direction for the β-strands. Although the electron density of the strands in the N- and C-terminal parts was readily interpretable, the quality of the intermediate part remained problematic. This could be due to various reasons: (i) the lack of very good low-resolution reflections because of overloads or beamstop shadow; (ii) the general high nonisomorphism between different datasets from different crystals, making it difficult to combine phase information from different datasets in SHARP (19); (iii) the relatively high anisotropy of the reflection data (see footnote, Table S2); and (iv) the low resolution of the x-ray data (Fig. 1E).

The quality of the electron density was not sufficient to allow the determination of the hVDAC1 structure by standard crystallographic methods alone. Therefore, we pursued a hybrid approach in which we integrated complementary information from NMR spectroscopy and x-ray crystallography. These calculations were done iteratively by cycling between real space ensemble generation using the Inferential Structure Determination (ISD) program (16) and reciprocal space refinement using the BUSTER-TNT (17) program (compare with Fig. 1).

In the ISD real space calculations, the crystallographic data are introduced as positional restraints. To generate the initial structure ensemble, the crystallographic data were used to dock the model into the density map and to structurally align the model and the Cα-trace. The Se-Met positions were used to dock the model into the density map and to structurally align the model and the Cα-trace. The distance, dihedral angle and positional restraints as input an initial structure ensemble was generated with the ISD software (16). The highest-probability conformation served as a starting structure in a subsequent reciprocal space refinement with BUSTER-TNT (17). In addition to the structure factor amplitudes and experimental phases from SHARP (18) (H), secondary structure distance restraints (D) were imposed in all BUSTER-TNT refinements. The structure calculation proceeded by cycling between ensemble generation and crystallographic refinement. In the conformational sampling, the positional restraints (E) were used only in the initial iteration. In subsequent calculations, the last structure obtained by crystallographic refinement served as a template (positional restraints to all Cα-coordinates). The distance and dihedral angle restraints were imposed in all ISD structure calculations.

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ensemble, positional restraints were derived from the four SeMet atoms that could be assigned unambiguously and from eight β-strands that were built manually into the electron density (see Fig. 1 F). In later iterations, the positional restraints were derived from the structure obtained by reciprocal space refinement. The four SeMet sites in the mutant L10M, V184M are at positions 10, 129, 155, and 184.

The positional restraints were combined with standard NMR restraints: 95 distance restraints were obtained from the NOEs, 368 backbone and 170 side chain dihedral angle restraints were predicted from the chemical shifts. Moreover, H/D exchange rates in combination with the topology model were used to derive 150 hydrogen bonds that were also introduced as distance restraints into the ensemble calculation. Finally, we applied additional Ca-Cα distance restraints within the β-strand regions. These restraints improved the convergence of the calculation but were not essential for the overall structure. Test calculations showed that virtually the same ensemble is obtained without imposing the Ca intrastrand restraints.

After each ensemble generation, a joint NMR/x-ray refinement was carried out by using BUSTER-TNT. The highest-probability model of the ensemble was docked into the electron-density envelope generated by the x-ray crystallographic phases and refined against the spatial constraints given by the diffraction data (Fig. 1G). Distance restraints based on the NMR secondary structure assignment were used for all β-sheet residues and for the N-terminal helix (21). Cyclic rebuilding and addition of further restraints into the model calculation improved the initial phases to yield an electron density map that allowed tracing of the entire model against the spatial constraints given by the diffraction data (Fig. 1F, crystallographic data: Tables S2 and S3). The improvement of the electron density from the initial experimental map after refinement with BUSTER-TNT is shown in Fig. S4.

**3D Structure of hVDAC1.** The channel pore of slightly concave shape (Fig. 2A) has dimensions of ~3.5 × 3.1 nm in the horizontal and ~4 nm in the vertical directions (Fig. 2B). Similar dimensions for VDAC proteins in the native state have also been obtained by high resolution AFM investigations (3.8 × 2.7-nm diameter) (22) and electron microscopy studies (diameter of ~3 nm) (7), both of which derived from *Saccharomyces cerevisiae* in natural membrane composition. hVDAC1 reconstituted in artificial membranes showed identical dimensions (3.7-nm diameter × 4.3-nm height) (23). The inner diameter of the pore is ~1.5 × 1 nm and therefore leaves space for diffusion of small metabolites (Fig. 2B and F).

The final protein structure contained all residues with the N and C termini of the protein oriented to the same side of the barrel (Fig. 2 A, C, and D). In contrast to bacterial outer membrane proteins, which archetypically show structures with an even number of strands and a distribution of long loops pointing to the extracellular and short turns to the periplasmic side (24), hVDAC1 shows an uneven number of 19 strands and 18 loop-like connections with a less distinct distribution of shortened loop structures to one side of the membrane (see Fig. 2C). Adjacent to the loops, which are dynamic on the pico- to nanosecond time scale according to NMR spectroscopic relaxation measurements (Fig. S1C), the formation of two incomplete aromatic girdles separated by a distance of only ~1.5 nm on an axis parallel to the membrane normal is visible (data not shown). The unequal number of strands in the barrel requires one parallel interaction of two adjacent and slightly twisted terminal β-strands ending on the same side (β1 and β19) of the membrane (Fig. 2 D and E). The average inclination of the β-strands relative to the membrane normal is ~1.5 nm on an axis parallel to the membrane normal is visible (data not shown). The unequal number of strands in the barrel requires one parallel interaction of two adjacent and slightly twisted terminal β-strands ending on the same side (β1 and β19) of the membrane (Fig. 2 D and E). The average inclination of the β-strands relative to the membrane normal is ~1.5 nm on an axis parallel to the membrane normal is visible (data not shown).
to the barrel axis is 37°, varies between 27 and 46°, and forms a barrel of almost circular shape (see Fig. 2F), similar, e.g., to the monomeric porin OmpG (25, 26).

VDACs within a certain species, e.g., mammals or fungi, share a high sequence identity (typically >80%) (data not shown), suggesting that all VDAC proteins share the same fold. In addition to the VDAC isoforms, the translocase of the outer mitochondrial membrane (human TOM40) (2) shows a striking homology to hVDAC1 in terms of both sequence and predicted secondary structure (data not shown), suggesting that hTOM40 also folds into a 19-stranded β-barrel.

**N-Terminal Helix Is Located Inside the Pore.** The N-terminal helix comprising residues Tyr-7 to Val-17 is folded horizontally inside the barrel wall approximately at the midpoint of the hydrophobic portion of the membrane (Fig. 2F and G). Although this arrangement has no “precursors” in bacterial porins, the position and the influence of the helix onto the overall barrel architecture is reminiscent of the long loop L3 present in bacterial porins (24), which folds in a similar position halfway perpendicular to the membrane normal into the barrel lumen (Fig. 2B). Whereas the preceding N-terminal residues (Val-3–Tyr-7) are rather loosely attached to the barrel wall and point toward the pore outlet, the N terminus with an overall charge of zero is amphiphatic and more strongly connected to the barrel by a short stretch of residues running almost in parallel to the angle of the β-sheets. Positively charged residues face the barrel interior, whereas negatively charged aspartates interact with the barrel wall. The counter residues on the barrel wall (His-181, Ser 193, Tyr-195, Lys-224) are predominantly hydrophilic and are conserved among most VDAC proteins (data not shown).

**Dimerization of VDAC via a Small Hydrophobic Interface.** Previous studies have indicated that VDAC from several species can exist in different oligomerization states from monomers to dimers, trimers, tetramers, hexamers, and higher oligomers (27–29). A transient dimerization is in agreement with chemical cross-linking experiments where dimers and higher-order oligomers were found in diluted solutions of hVDAC1. NMR relaxation measurements of VDAC in LDAO, which correspond to a much lower local concentration than that of VDAC in the mitochondrial membrane indicate a global correlation time of the VDAC-micelle assembly of 60 ns, corresponding to a monomer/dimer equilibrium (see SI Text).

Using crystallographic symmetry operators, a dimer was constructed that coincides with electron micrographs and implies the formation of a parallel dimer (Fig. 3A) (23). In analogy to OMPs, this interface is formed by the four β-strands (β18, β19, β1, β2) with the lowest inclination values. The interface provided by residues in β-strands β1 (Ile-27, Leu-29), β2 (Glu-50, Thr-51), β18 (Leu-257, Leu-259), and β19 (Leu-277) covers 550 Å², suggesting that it might be much more dynamic than in bacterial porins (Fig. 3B). It should be noted that residues showing both enhanced (β1, β2) and slow exchange rates (β18, β19) with water (see below) are involved in the dimer interface excluding transient dimerization as the reason of broadening of NH resonances of strands β1 and β2.

**Conformational Instability of the N-Terminal Part of hVDAC1.** To probe the conformational stability and the influence of particular residues, we performed hydrogen/deuterium exchange coupled to NMR spectroscopy. Surprisingly, residues in β1-β4 of WT hVDAC1 solubilized in LDAO or in Cymal-5 showed rapid amide proton exchange (Fig. 4A and C and Fig. S5), whereas this was not observed for the other β-strands. This indicates that the N-terminal four β-strands are less stable than other regions of the VDAC barrel and switch between different conformations. As part of this conformational exchange, hydrogen bonds within the four N-terminal strands of the barrel are transiently broken. This is consistent with computational analyses that predict an increased flexibility for the N-terminal region (Fig. 4B). Importantly, mutants that affected the voltage sensitivity of Saccharomyces cerevisiae VDAC (scVDAC) are mainly located in the N-terminal domain: the α-helix (conserved Asp-16), the linker (conserved Lys-20), and β-strands β1-β5 (Lys-46, Lys-61, Lys-65, and Lys-84 in scVDAC) (Fig. 4C) (30). This suggests that the voltage gating of VDAC requires conformational flexibility.

The structural instability of the N-terminal part is particularly influenced by Glu-73. This residue is conserved in different isoforms of human VDAC and in VDAC from mammals and fungi (data not shown). In the 3D structure, E73 points to the membrane and interrupts the amphiphatic pattern in strand β4 (Fig. 1B and Fig. S1). We replaced Glu-73 by valine, a substitution that resulted in slower amide proton exchange in β-strands β1-β4 (Fig. 4A and C and Fig. S5), indicating that the conformation of the N-terminal part of the barrel is stabilized. Importantly, substitution of Glu-73 by glutamine has important consequences for the function of VDAC. This mutation abolished ruthenium red and hexokinase I-mediated inhibition of VDAC channel activity and ruthenium red and hexokinase I-mediated protection against apoptosis (33). In both E73V and E73Q, the charged residue that breaks the alternating pattern of hydrophobic and hydrophilic residues in strand β4 is removed. Thus, the β-barrel might also become more stable by
the E73Q mutation. This is consistent with the finding that ruthenium red is no longer able to close the pore of E73Q VDAC and the channel gating occurs at higher voltages for E73Q VDAC compared with WT murine VDAC (34). We conclude that the conformational flexibility of β-strands 1–4 is, in addition to the N-terminal helix, important for the voltage gating and for protein–protein interactions of VDAC in vitro and in vivo.

**VDAC Retains the Sidedness of Bacterial Porins.** In mitochondria, hVDAC1 faces the cytoplasm and the IMS by hydrophilic residue exposure (Fig. 3C). In the structure of hVDAC1, an asymmetry in the length of the loop regions is visible (Fig. 3), in agreement with AFM measurements that showed a rough and a flat side of VDAC (27). In addition, tyrosine residues are found almost exclusively at the water–membrane interface at the C-terminal end of every second strand (Fig. 1B) (35). Moreover, antibodies raised against the N terminus of scVDAC were able to bind to these membranes only when mitochondrial envelopes were disrupted (36). The combined data suggest that the C terminus of VDAC extends into the IMS, resembling the orientation found in all bacterial OMPs (24).

**Summary and Perspectives.** Our combined approach using data of both NMR spectroscopy and x-ray crystallography enabled the determination of the 3D structure of the first MOM protein, hVDAC1. The finding of a 19-stranded β-barrel is in strong contrast to evolutionary theories predicting bacterial β-barrels and related proteins to be formed by an ancient β-hairpin motif (37). The new channel architecture is likely to be a consequence of the differences in membrane environment, sorting signal and partner proteins experienced by integral membrane proteins of the outer membrane of bacteria and those of the MOM and most likely adopted by other MOM proteins. On the basis of the NMR resonance assignment and the 3D structure of the hVDAC1 channel, a variety of experiments and structural studies of other VDAC isoforms will be possible. The functional difference of the E73 mutant compared with WT could be rationalized based on the structure described in this work.

**Materials and Methods**
Expression, refolding, and purification of hVDAC1 were done as described. NMR spectra were recorded from a perdeuterated 15N and 15N/13C labeled...
samples containing 0.6 mM HVDAC, 25 mM BisTris, −250 mM Lauryledimethylamino-oxide (LDAO), and 10% D2O, pH 6.8. Assignment of backbone resonance of hVDAC1 was achieved through six independent types of NMR data: Cα chemical-shift connectivity, intrastrand and sequential HN-HN NOEs, amino acid-specific labeling, 20 point mutations, paramagnetic broadening induced by spin labels, and hydrogen/deuterium exchange experiments. Native crystals were obtained with the hanging-drop method by mixing 1.5 μl of protein-Cymal-5 detergent solution containing 5–15 mg/ml protein with 0.75 μl of reservoir solution (30% PEG 400; 0.1 M Na Heps, pH 7.5; 0.2 M magnesium chloride). The trigonal crystals (space group P321) diffracted up to 4 Å. Structure calculations were done iteratively by cycling between real space ensemble generation using the ISD method and refinement using the autoSHARP method.

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