Protein Structure and Folding:
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High Resolution Structure and Double Electron-Electron Resonance of the Zebrafish Voltage-dependent Anion Channel 2 Reveal an Oligomeric Population*

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Background: Biochemical characterization of voltage-dependent anion channel 2 (VDAC2) is limited due to an inability to obtain functional protein.

Results: The crystal structure of VDAC2 suggests a dimer interface that is confirmed by double electron-electron resonance and cross-linking.

Conclusion: zfVDAC2 has a fractional dimeric population.

Significance: VDAC isoforms are structurally similar, but this study has identified a number of hot spots that require further exploration.

In recent years, there has been a vast increase in structural and functional understanding of VDAC1, but VDAC2 and -3 have been understudied despite having many unique phenotypes. One reason for the paucity of structural and biochemical characterization of the VDAC2 and -3 isoforms stems from the inability of obtaining purified, functional protein. Here we demonstrate the expression, isolation, and basic characterization of zebrafish VDAC2 (zfVDAC2). Further, we resolved the structure of zfVDAC2 at 2.8 Å resolution, revealing a crystallographic dimer. The dimer orientation was confirmed in solution by double electron-electron resonance spectroscopy and by cross-linking experiments disclosing a dimer population of ~20% in lauryldimethane amine oxide detergent micelles, whereas in lipidic bicelles a higher population of dimeric and higher order oligomers species were observed. The present study allows for a more accurate structural comparison between VDAC2 and its better-studied counterpart VDAC1.

Mitochondria are often depicted as the cell’s “powerhouse” because they generate ATP from ADP and inorganic phosphate through the oxidative phosphorylation process. It is now firmly established that mitochondria have a far more expansive role in cellular function, including signaling, cell death/survival, and differentiation (1–3). As such, mitochondria are central to normal human physiology as well as disease. The most abundant proteins in the outer mitochondrial membrane, the voltage-dependent anion channels (VDACs), are large pore-forming proteins that mediate the efficient exchange of metabolites and ions between the cytosol and the mitochondrial intermembrane space (IMS). In mammals, there are three VDAC isoforms (VDAC1–3), which have a high degree of sequence homology, yet their relative ratios and distribution between cell types are varied.

The sequence identity/similarity between human VDAC1 (hVDAC1), hVDAC2 and hVDAC3 is 75%/91% and 67%/85%, respectively (Fig. 1). Undoubtedly, VDAC1 is the best characterized isoform. After decades of struggle, three groups independently resolved the high resolution structure of VDAC1 (4–6). Using the lipidic bicelle crystallization technique (6–8), we solved a high resolution x-ray structure of mouse VDAC1 (mVDAC1) at a resolution of 2.3 Å. All reported high resolution structures of VDAC1 revealed a novel fold of a 19-stranded β-barrel with an N-terminal α-helix lying inside the pore. This helix is aligned nearly parallel to the membrane plane, causing a partial narrowing of the pore. Numerous studies have established the functional importance of the N-terminal α-helix in regulating the flux of metabolites through the channel (9, 10), and the observation that it is localized within the pore makes it

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*The abbreviations used are: VDAC, voltage-dependent anion channel; hVDAC, zVDAC, and mVDAC, human, zebrafish, and mouse VDAC, respectively; CHAPS, 3-[N-(3-cholamidopropyl)dimethylammonio]-1-propanesulfonic acid; IMS, intermembrane space; DEER, double electron-electron resonance; LDAO, lauryldimethane amine oxide; CW, continuous wave; DEF, dipolar evolution function.

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ideally suited to regulate the passage of ions and metabolites through the pore. Based on molecular dynamics simulations and a series of cross-linking studies, the structure is believed to reside in the "open" conformation, facilitating the passage of ATP and other nucleotides (11, 12).

In contrast to VDAC1, structural and biochemical characterization of VDAC2 and -3 is rather limited, yet these isoforms have distinct physiological functions. In particular, VDAC2 null mice are embryonic lethal (13), whereas null mice of VDAC1 and -3 isoforms display mild phenotypes (14, 15). Furthermore, VDAC2 has been shown to interact and inhibit the proapoptotic protein BAK, thereby hampering mitochondrial apoptosis (13). In cardiomyocytes, VDAC2 specifically interacts with the ryanodine receptor and mediates Ca\(^{2+}\) transfer from the sarcoplasmic reticulum into mitochondria (16), implicating a role for VDAC2 in cardiac Ca\(^{2+}\) homeostasis. Additionally, the small quinazolinone agent erastin, an anti-tumor compound, interacts with VDAC2 to induce cell death in tumor cells expressing mutant RAS (17, 18). In total, there is significant evidence that despite the high sequence identity in this family, VDAC2 has a specific phenotype, which is distinct from those of the other two VDAC isoforms.

To gain insight into the structure and function of VDAC2, we expressed and purified zfVDAC2. The purified protein forms active channels in lipid planar bilayers, and we were able to determine the structure at 2.8 Å resolution. The overall structure is very similar to VDAC1, revealing a dimer interface between \(\beta_{17}, \beta_{18}, \beta_{19}, \beta_{1}, \) and \(\beta_{3}\). Using double electron-electron resonance (DEER) measurements and cross-linking experiments, we confirmed the dimer topology in solution and found a fractional dimer content of \(\sim 20\%\) in detergent micelles and \(\sim 40\%\) in lipidic bicelles.

**EXPERIMENTAL PROCEDURES**

**Cloning of pQE60-zfVDAC2\(^{C127A}\) and Single-cysteine Mutants—**
The full-length zfVDAC2 ORF was purchased from Open Biosystems and PCR-amplified. PCR primers were used to add an Ncol site (5') and a His\(_{6}\) tag followed by a HindIII site (3') for subsequent Ncol/HindIII cloning into the pQE60 expression plasmid. For DEER experiments, we created construct pQE60-zfVDAC2\(^{C127A/S232C}\). Three individual PCRs were used to create three overlapping fragments of the zfVDAC2 ORF and to introduce the C127A and S232C mutations, respectively, by mutagenesis primers. Individual PCR fragments were fused by
splicing by overlap extension-PCR, and the resulting product was ligated into the pQE60 backbone using Ncol/HindIII. All PCR-amplified products were fidelity-checked by sequencing (Laragen, Inc., Culver City, CA).

To create single-Cys mutants for cross-linking experiments, the Cys-less backbone pQE60-zfVDAC2C127A was created by fusion of base pairs 427–870 of the ORF from pQE60-zfVDAC2 into pQE60-zfVDAC2C127A using PstI and HindIII, thereby eliminating the S232C mutation. Single-cysteine mutants were subsequently generated in the pQE60-zfVDAC2C127A backbone by mutagenesis PCR. For pQE60-zfVDAC2C127A/D50C, pQE60-zfVDAC2C127A/S232C, and pQE60-zfVDAC2C127A/T83C, the S232C mutation primers were used to introduce the respective mutations in two independent PCRs, which were then fused in a splice overlap extension PCR. Final PCR products were ligated into pQE60-zfVDAC2C127A using XhoI and PstI. For pQE60-zfVDAC2C127A/Q196C, pQE60-zfVDAC2C127A/I257C, pQE60-zfVDAC2C127A/S232C, and pQE60-zfVDAC2C127A/I255C, mutagenesis PCR was performed as described above, but PCR products were ligated into pQE60-zfVDAC2C127A using PstI and Nhel. All PCR-amplified products were fidelity-checked by sequencing (Eurofins MWG Operon, Ebersberg, Germany).

Expression and Purification of zfVDAC2—Cloning, expression, and purification of zfVDAC2 followed the previously reported protocol for mVDAC1 (6). In short, M15 Escherichia coli cells carrying pQE60-zfVDAC2 were grown in Luria broth with agitation at 37 °C, induced at an OD of 1 with 0.4 mM isopropyl-1-thio-β-D-galactopyranoside and grown for an additional 4–5 h. Cells were harvested by centrifugation and resuspended in 50 mM Tris-HCl (pH 8.0), 300 mM NaCl, 50% PEG 2000, 0.1% LDAO. 2% LDAO was added to the protein solution prior to the second dialysis step. Refolded protein was concentrated, ultra centrifuged (355,000 × g, 30 min) to remove aggregated protein, and passed over a Superdex 200 column in 150 mM NaCl, 0.1% LDAO, 20 mM Tris-HCl (pH 8.0) to obtain a homogenous protein population (Fig. 2, A and B). For crystallization, fractions containing the non-aggregated zfVDAC2 peak were collected, concentrated, and buffer changed to 50 mM NaCl, 0.1% LDAO, 20 mM Tris-HCl (pH 8.0) using an Amicon Ultra-30k (Millipore) concentrator.

Crystallization—Crystals of ~0.6 × 0.2 × 0.1 mm were grown by the hanging drop vapor diffusion method using the Mosquito crystallization robot. A 12 mg/ml protein solution was mixed (1:1) with a well solution of 100 mM Tris-HCl (pH 8.0), 100 mM KCl, and PEG 2000 ranging between 19 and 24%. Subsequent optimization was performed by the addition of n-undecyl-β-D-thiomaltopyranoside at its critical micelle concentration (0.01%). Crystals were cryoprotected by transferring them for 10 s into a solution containing 100 mM Tris-HCl (pH 8.0), 100 mM KCl, 50% PEG 2000, 0.011% n-undecyl-β-D-thiomaltopyranoside prior to flash freezing in liquid nitrogen.

Lipid Planar Bilayer—Lipid bilayer experiments were performed as described previously (6). In brief, recordings were performed in diphytanoyl phosphatidylcholine bilayers across a 0.1-mm hole in 10 mM Tris-HCl (pH 7.4), 1 mM KCl, 5 mM CaCl2. After channel insertion, currents were recorded in response to test pulses between −50 and +50 mV. Total point histograms at each applied voltage were plotted to calculate the conductance (G) of open and closed states (Fig. 2C).

X-ray Data Collection and Processing—X-ray diffraction data were collected to 2.7 Å on flash-frozen crystals (~180 °C) at the Advanced Light Source (Berkeley, CA) (beam line 5.0.2). Indexing and integration were performed with XDS (19), using data up to 2.8 Å, and the space group was determined to be P3_121 with the following unit cell dimensions: a = 72.47 Å, b = 72.47 Å, and c = 177.95 Å (α = β = 90° and γ = 120°). One molecule is present per asymmetric unit with an estimated solvent content of 70.4% (V_m = 4.15 Å³/Da) based on the Matthews probability calculator (20, 21). The structure was solved by molecular replacement using the murine VDAC1 coordinates (Protein Data Bank code 3EMN) as the search model in PHENIX (21). After generation of the initial model, it was manually rebuilt using Coot (22), followed by iterative rounds of refinement in PHENIX (21). The final model (Protein Data Bank code 4BUM) contained 283 amino acids, six water molecules, and one LDAO molecule with a final R-factor and R-free of 0.246 and 0.282, respectively. Data collection and refinement statistics are listed in Table 1. All figures containing structures were generated using PyMOL (23) or Chimera (24).

Spin Labeling, Continuous Wave (CW) EPR Spectroscopy, and DEER Experiments—For CW and DEER experiments, DTT was removed during the size exclusion chromatography step, and the protein was incubated with a 5-fold molar excess of 1-oxyl-2,2,5,5-tetramethylpyrroline-3-methyl methanethiosulfonate for 16 h at 4 °C to generate the R1 side chain (25). Excess spin label was removed by repeated washes using an Amicon Ultra-30k (Millipore) concentrator in 20 mM Tris-HCl (pH 8.0), 50 mM NaCl, 0.1% LDAO, 20% glycerol. For zfVDAC2C127A/S232C, the fraction of spin-labeled protein was >98% as estimated from a 4,4'-dithiodipyridine titration for free thiol after the spin labeling reaction (26). A fraction of the spin-labeled zfVDAC2C127A/S232C was reconstituted in bilayers by mixing the protein with bilayers in a 2.5:1 ratio with a 35% (2.8:1) 1-α-dimyristoylphosphatidylcholine/CHAPSO bilayer solution.

The CW EPR spectra of zfVDAC2 232R1 in LDAO and bilayers were recorded at room temperature on a Varian E-109 spectrometer fitted with a two-loop one-gap resonator; samples of 5 μl were contained in a borosilicate capillary tube (0.64-mm inner diameter × 0.84-mm outer diameter; VitroCom Inc.). For LDAO, a second spectrum was recorded with the sample containing 30% (w/w) sucrose solution. The spectra were recorded using 2-milliwatt microwave power and 1-G field modulation at 100 kHz.
Four-pulse DEER experiments (27) at Q band were conducted at 80 K on a Bruker ELEXSYS 580 spectrometer fitted with a Bruker EN5107D2 resonator. The protein concentration during the DEER experiments was maintained at or below 200 μM. 20 μl of spin-labeled protein reconstituted in LDAO or bicelles containing 20% (v/v) glycerol were loaded into a borosilicate capillary tube (1.4-mm inner diameter × 1.7-mm outer diameter; VitroCom) and subsequently flash-frozen in liquid nitrogen. A 36-ns π-pump pulse was set at the maximum absorption spectra, and the observer /H11032 using a 4,4'-dithiodipyridine assay (26). From the depth of modulation and using /H9004/H18528/H20849/H20850/H20850 where

\[ \Delta = \frac{\ln(1 - \Delta)}{C} \]

where \( \Delta \) is the “depth of modulation” (shown in Fig. 7, B and C), and C is a constant that depends on the pump pulse profile, the resonator, and the spectral line shape of the nitroxide spin label (28). In the present experiments, C was determined using double cysteine mutants of T4 lysozyme (N68C/T109C and T109C/V131C) known to be quantitatively labeled with R1 as judged by determination of free thiols after the labeling reaction using a 4,4'-dithiodipyridine assay (26). From the depth of modulation and using \( n_m = 2 \), C is obtained from Equation 1; the values obtained from the two T4 lysozyme mutants agreed to within 1%. From the experimentally determined depth of modulation and C, the value of \( n_m \) for VDAC2 in LDAO or bicelles can be determined.

Cross-linking Experiments—Disulfide bonds between single-cysteine mutants were induced with the oxidizing reagent dichloro(1,10-phenanthroline)Cu(II). zfVDAC2 mutants were diluted to 15 μM in buffer containing 150 mM NaCl, 20 mM Tris-HCl (pH 8.0), 5 mM DTT, and 0.1% LDAO. The oxidizing reaction was performed with 5 mM dichloro(1,10-phenanthroline)Cu(II) at 4 °C for 2 min and subsequently quenched with 10 mM EDTA and 10 mM N-ethylmaleimide. The results were analyzed on 12% SDS-polyacrylamide gels stained with a Coomassie-based InstantBlue reagent (Expedeon).

RESULTS

Protein Purification and Lipid Bilayer Conductance—Native recombinant zfVDAC2 protein with a His tag fused to the C terminus was prepared using a two-step column purification that consisted of 1) solubilized inclusion bodies that were applied to a Talon metal affinity column and eluted with 150 mM imidazole and 2) concentrated protein that was refolded and applied to a Superdex 200 column to remove aggregation and obtain a homogenous protein population (Fig. 2, A and B). Purified zfVDAC2 protein was inserted in planar lipid bilayers, where it exhibited a single channel conductance of 4.2 ± 0.5 nanosiemens at 10 mV in 1 M KCl (open state; Fig. 2C). As the applied voltage was increased to ≥30 mV, the channel presented distinct gating behavior between the open and closed states with a conductance of 1.7 ± 0.2 nanosiemens (Fig. 2C). This behavior is in good agreement with recordings from VDAC1 isolated from rats (29) and indicates that zfVDAC2, from our preparation, forms active channels. To further understand the functional aspects of ion and metabolite trafficking as well as the relation of zfVDAC2 to previously reported VDAC1, we pursued a high resolution structure of zfVDAC2.

Structural Overview—Optimized crystals of zfVDAC2 belong to the triggular space group P3 21 with one molecule per asymmetric unit. The structure was determined by molecular replacement using the original mVDAC1 coordinates (Protein Data Bank code 3EMN) as a search model. The only part of the model that required manual rebuilding following the molecular replacement procedure was the loop between β-strands 1 and 2. The model was ultimately refined from merged data to a resolution of 2.8 Å with an R-free of 24.6% and an R-work of 28.2% (Table 1). 282 of the 283 amino acids of zfVDAC2 were assigned along with six water molecules and one copy of LDAO. Further details regarding the crystallization and structure determination can be found under “Experimental Procedures.” A refined electron density map is shown in Fig. 3C.

The structure of zfVDAC2 is similar to that of mVDAC1 (Protein Data Bank code 3EMN), forming a β-barrel composed of 19 β-strands (Fig. 3). All of the strands exhibit an antiparallel pattern with the exception of strands 1 and 19, which associate in a parallel manner to seal the barrel. The maximal height and
width of the β-barrel are 35 and 40 Å, respectively. The correct orientation of the protein within the outer membrane of the mitochondria is still debated, with multiple studies producing conflicting results on the endogenous position of the N and C termini (30–32); however, a recent study has shown the C terminus to reside in the IMS (33). The N terminus of the protein forms a distorted α-helix that is aligned nearly parallel to the plane of the membrane, causing a partial narrowing of the pore (Fig. 3). The pore forms a large pathway that traverses the entire length of the protein. There have now been a number of studies confirming that this conformation probably resides in the “open” state of the channel (11, 12).

A structural alignment between the crystal structure of zfVDAC2 and mVDAC1 (Protein Data Bank code 3EMN) showed a root mean square deviation of 0.98 Å (Fig. 4A). The well aligned structures, including the N-terminal segment, indicate that these proteins share many similar properties and

| Data collection | Crystallography statistics for zfVDAC2 |
|-----------------|---------------------------------------|
|                 | Crystallography statistics for zfVDAC2 |
|                 | P3_21                                 |
|                 | Cell dimensions                        |
|                 | a, b, c (Å)                            |
|                 | 72.47, 72.47, 177.95                   |
|                 | α, β, γ (degrees)                      |
|                 | 90, 90, 120                            |
|                 | Resolution (Å)                        |
|                 | 51.29–2.701                           |
|                 | R_{merge}, or R_{merge}                |
|                 | 0.08 (0.76)                           |
|                 | Completeness (%)                      |
|                 | 97.98 (97.3)                          |
|                 | Redundancy                             |
|                 | 3.5 (3.4)                             |

| Refinement | Refinement |
|------------|------------|
| Resolution (Å) | 50–2.8 |
| No. of reflections | 53,149 |
| R_{merge}/R_{merge} | 24.55/28.22 (31.23/35.34) |
| No. of atoms | 2151 |
| Protein | 2133 |
| Ligand/ion | 12 |
| Water | 6 |
| B-Factors | |
| Protein | 51.0 |
| Ligand/ion | 58.3 |
| Water | 24.4 |
| Ramachandran favored (%) | 94 |
| Ramachandran outliers (%) | 0.71 |
| Root mean square deviations | 0.004 |
| Bond lengths (Å) | 0.87 |

* Values in parentheses are for the highest resolution shell.

**FIGURE 4. Superposition of zfVDAC2 and VDAC1.** A. zfVDAC2 in blue and mVDAC1 (Protein Data Bank code 3EMN) in red were aligned with a C_{α} root mean square deviation of 0.98 Å. There is a large displacement (~12 Å) in the loop between β-strands 1 and 2. B, an IMS view of the loop between β-strands 1 and 2, including zfVDAC2 in blue, mVDAC1 in red, hVDAC1 (Protein Data Bank code 2JK4) in green, and a representative NMR conformer of hVDAC1 (Protein Data Bank code 2K4T) in salmon.

**FIGURE 3. Overall structure of zfVDAC2.** Shown are cartoon representations of zfVDAC2 viewed perpendicular to the membrane plane (A) and in the membrane plane (B). The protein backbone is rainbow-colored from the N terminus (blue) to the C terminus (red). A 2F_{o} − F_{c} map contoured at 1.5σ is displayed in a blue wire frame with the zfVDAC2 model shown in sticks (C).
functions. The sole exception is a large displacement (~12 Å) of the loop between strands 1 and 2 (Fig. 4B). Further comparison using Columbic electrostatic calculations (Fig. 5) revealed a continuous electronegative patch on the cytosolic lip of mVDAC1 extending from the C terminus of the protein (β19) and within the loops connecting β1 to β2, β3 to β4, and β5 to β6. In zfVDAC2, the loop connecting β1 to β2 is electronegative, disrupting the electronegative continuity. In addition, there are a number of other subtle differences. Residues 61–65 in β-strand 3 contain four positive residues in zfVDAC2 (KYKRS) as opposed to two positive residues in mVDAC1 (KYRWT); residues 280–282, at the C terminus of the protein, contain two negative residues in zfVDAC2 (Glu-Leu-Glu) as opposed to one in mVDAC1 (Glu-Phe-Gln); and Glu-36, which resides in mVDAC1 in the mobile loop connecting β-strands 1 and 2, is substituted to an Ala in zfVDAC2. Additional hydrophobicity analysis did not show any significant difference between the two isoforms.

**Dimer Interface**—The oligomeric organization of VDAC has been strongly correlated with its physiological function. Numerous studies have demonstrated that VDAC can achieve a number of oligomeric populations, depending on the cell’s environment, including monomers, dimers, trimers, tetramers, hexamers, and even higher order oligomers (34). More recently, this arrangement has been shown to be dynamic and further influenced by the lipidic environment (35). As was done with hVDAC1 (5), a zfVDAC2 dimer was constructed using crystal symmetry operators. zfVDAC2 forms crystals of a space symmetry group P3221, which leads to three distinct zfVDAC2 symmetry-related interfaces. The most extensive surface, with a buried surface area of 554 Å² and a solvation free energy gain of ΔG = −14.9 kcal/mol, is formed by buried hydrophobic residues in strands β17 (Arg-252 and Leu-251), β18 (Leu-259, Leu-257, and Ile-255), β19 (Leu-281, Leu-279, Leu-277, and Leu-275), β1 (Val-31, Leu-29, and Val-27), and β3 (Val-54, Ser-52, Asp-50, Thr-49, and Ser-47) (Fig. 6).

To confirm the dimer interface of zfVDAC2 in solution and the relative abundance of zfVDAC2 oligomers, we generated a cysteine-less construct by replacing the endogenous cysteine residue with alanine (C127A) and subsequently engineered a cysteine at position 232, which is the N-terminal residue of β16. This mutant was then spin-labeled with 1-oxyl-2,2,5,5-tetramethylpyrroline-3-methyl methanethiosulfonate reagent to generate an R1 side chain, and the interspin distance distribution was measured using DEER spectroscopy in LDAO and in lipidic bicelles. The DEER experiment measures the dipolar interaction between unpaired electrons in the range of 17–70 Å (27) and thus can be used to detect the presence of homodimers or higher oligomers in a solution of singly labeled proteins. Residue 232 was selected for this study because modeling of the R1 side chain at site 232 in the crystallographic dimer shown in Fig. 6 suggests a distance between R1 labels of ~31 Å (Fig. 7A), which is well within the range of interspin distance that can be reliably measured with DEER spectroscopy.

In the DEER experiment, magnetic interactions between randomly distributed spins in different molecular entities give rise to an exponentially decaying “background” signal (red traces in the left-hand panels of Fig. 7, B and C). The experimental DEER signals (black and cyan traces in the left-hand panels of Fig. 7, B and C) deviate from the background at short evolution times, unequivocally demonstrating the presence of specific oligomers in solutions of both LDAO and bicelles. Removal of the background signal from the raw data gives the corresponding DEFs (right-hand panels of Fig. 7, B and C).
which can be fit to obtain a probability distribution of interspin distances (Fig. 7D). In LDAO, the distance distribution is broad with a full width at half-height of nearly 20 Å, which exceeds the width anticipated from a distribution of R1 rotamers (34) and suggests flexibility in the region of H925. The broad distribution may also arise from heterogeneity regarding the relative orientation of each molecule in the dimer. However, some of the local maxima in the broad distribution could arise from distinct rotamers of R1. The median of the distribution is 30.5 Å, close to that determined from modeling of R1 in the crystallographic dimer.

In bicelles, the median of the distribution is 32.4 Å (the most probable distance is 30.3), again close to that expected from the crystallographic dimer. However, there is a small population near 55 Å (arrow in Fig. 7D). The shape of the distribution is not well determined due to the limited data collection time of 2.5 μs, but a detectable population at long distance is required to fit the data. A distance in this range cannot correspond to any possible homodimer but must originate from higher order oligomers. The depth of modulation of the DEF (Δ; Fig. 7, B and C) is <4% for VDAC2 232R1 in either LDAO or bicelles, compared with ~10% for the doubly labeled T4 lysozyme pairs used for determination of the constant C under identical conditions (see “Experimental Procedures”). The low depth of modulation could be due to incomplete labeling of the protein and/or due to a small fraction of protein in oligomeric form (28). Because labeling was found to be quantitative (see “Experimental Procedures”), it is concluded that only a small fraction of the protein exists in oligomeric form, consistent with recent spectroscopy studies on hVDAC1 (35).

The distance distribution for 232R1 in LDAO clearly supports the formation of a homodimer. Within the context of this model, the fraction of VDAC2 in dimer form, fD, can be estimated from the modulation depth of the DEER signal according to Refs. 24 and 25 (\( f_D = n - 1 = \ln(1 - \Delta)/C \)), which gives ~18%.

The depth of modulation for VDAC2 232R1 in bicelles is larger than for the protein in LDAO (compare delta values in panels B and C of Fig. 7), which indicates a higher fraction of total oligomeric species (~40%). The larger depth of modulation arises due to the existence of higher order oligomers (indicated by the 55 Å population in the distance distribution) and possibly a larger value for fD. An increase in the oligomeric population in bicelles compared with LDAO micelles is in agreement with previous studies, which reported that the percentage of dimer is influenced by detergent or lipidic environments (35).

To obtain information on the relative mobility of the 232R1 side chain in LDAO and bicelles, CW EPR spectra were recorded for the protein in the two environments. In both cases, the spectra show a dominant component reflecting restricted mobility of the R1 side chain (Fig. 7E, top spectra, I), and a second component reflecting higher mobility (m). The two spectral components can arise from slow exchange (microsec-
ond to millisecond time scale) between protein conformational
substates or from rotameric equilibria of R1 (36). It has been
shown that stabilizing osmolytes, such as sucrose, shift protein
conformational equilibria but have little effect on rotameric
equilibria of R1. For 232R1, the addition of sucrose shifts the
conformational equilibria but have little effect on rotameric
substates or from rotameric equilibria of R1 (36). It has been
shown that stabilizing osmolytes, such as sucrose, shift protein
conformational equilibria but have little effect on rotameric
equilibria of R1. For 232R1, the addition of sucrose shifts the
conformational equilibria but have little effect on rotameric
substates or from rotameric equilibria of R1 (36).

To further validate the data obtained by DEER, we employed
cross-linking experiments of single-cysteine mutants in an
effort to better define the dimer interface in solution. We
generated three cysteine mutants that were part of the putative
dimer interface close to the cytoplasmic side of VDAC at posi-
tions Val-27, Asp-50, and Ser-52 and another set of three
mutants close to the IMS at positions Leu-251, Ile-255, and
Leu-257 (Figs. 6 and 8). These mutants were designed in loop
regions and on β-strands at S–S distance ranges of 3.7–11.1 Å.
In addition, two cysteine mutants were generated at sites that
should reside outside of the putative dimer interface at posi-
tions Thr-83 and Gln-196 (Fig. 8A). Upon the addition of
dichloro(1,10-phenanthroline)Cu(II), significant cross-linking
was observed at all six sites that are part of the putative dimer
interface (Fig. 8, B and C), indicative of disulfide formation
between monomeric proteins as opposed to the control sites
that did not display appreciable cross-linking (Fig. 8D).

DISCUSSION

In recent years, there has been a vast increase in structural
and functional understanding of VDAC1, but VDAC2 and -3
have been understudied. zfVDAC2 is a good structural model
for mammalian VDAC2 due to a high degree of sequence
homology (e.g. 83.0% identity and 93.0% similarity) between
zfVDAC2 and hVDAC2 with no sequence gaps (Fig. 1 shows a
zfVDAC2 alignment against selected mammalian VDAC
sequences). Most of the minor sequence differences reside in
the exposed loop regions where the transmembrane segments
remain conserved (Fig. 9, top row). Interestingly, mammalian
VDAC2s have a small 11-amino acid N-terminal extension that
is not present in zfVDAC2. However, this N-terminal extension
was shown not to be essential in mediating VDAC2-specific
properties of the channel, such as tBid-induced cytochrome c
release (37). Most of these minor sequence differences reside in
the exposed loop regions where the transmembrane segments
remain conserved (Fig. 9, top row). In total, zfVDAC is a good
model for mammalian VDAC2.

VDAC2 has a number of unique and specific protein-interac-
ting partners, including the proapoptotic protein BAK (13),
the ryanodine receptor (16), and erastin and tubulin (38). A
logical assumption is that these specific interactions would be
governed by structural differences. However, a structural com-
parison between VDAC1 and -2 reveals few differences on a
positional (Fig. 9, bottom row) or electrostatic basis (Fig. 5). The
sole exceptions are large displacements of the loop between strands 1 and 2 (amino acid sequence 32–40) and the loop connecting -strands 5 and -strands 6 (amino acid sequence 88–95) as well as a change in the electrostatic properties on the cytosolic lip of VDAC. Indeed, differences between VDAC1 and VDAC2 accounting for the VDAC2-specific tBid-induced cytochrome c release were previously mapped to the N-terminal two-thirds, namely amino acids 13–188 (39). The loop connecting -strand 1 and -strand 2 is extended 12 Å away from the pore in the zfVDAC2 structure as compared with mVDAC1 (Protein Data Bank code 3EMN). In the hVDAC1 x-ray structure (Protein Data Bank code 2JK4), this loop occupies a position quite similar to that in the zfVDAC2, and the NMR structure of hVDAC1 has this loop even farther away from the mVDAC1 position (Fig. 4B). Interestingly, there is an amino acid divergence at position 36 located at the apex of this mobile loop. For VDAC1, there is a highly conserved Glu, but in VDAC2 the residue is primarily an Ala or a Cys. To date, this residue, as far as we know, has not been subjected to functional analysis.

The loop connecting -strands 5 and 6 in the human x-ray structure is located ~13 Å internally from the position in mVDAC1 and zfVDAC2 x-ray structures. In the NMR model, this loop resides in the middle of these two positions. In the case of zfVDAC2, this loop is pushed away due to the presence of a symmetry-related molecule.

It would seem that loop displacements between -strands 1 and 2 and -strands 5 and 6, as observed here, would not suffice to explain the vast phenotypic differences between isoforms. However, the dynamic and electrostatic properties of these loops could play a key role in establishing protein-protein interactions and generate protein complexes or modulate gating properties and thus alter the function of VDACs. Indeed, binding of partner proteins was shown to vary substantially between VDAC isoforms, with compelling evidence that VDAC2 has a greater repertoire of binding partners (40–42).
Another aspect of VDAC physiology that has major implications, in particular for apoptotic events, is the protein’s oligomeric state. Current research has strongly implicated a dimeric organization of hVDAC1 by analysis of crystal packing (5, 7), chemical cross-linking (43), and fluorescence cross-correlation spectroscopy (35). However, these studies have been lacking for VDAC2.

Using crystallographic symmetry operators, we identified a parallel dimer interface for zfVDAC2 that is virtually identical to the one suggested for hVDAC1 with a buried surface area of 554 Å². The interface is formed by strands β9252, β9252, β9252, β1, and β3 (Figs. 6 and 8). Here we show that ~18% of the protein exists in a dimeric form in solution, as demonstrated by DEER spectroscopy for a singly R1-labeled mutant of zfVDAC2 reconstituted in LDAO; similar dimers exist in bicelles, apparently along with higher order oligomers (Fig. 7B). Remarkably, previous fluorescence cross-correlation spectroscopy studies done by Betaneli et al. (35) on hVDAC1 reconstituted in giant unilamellar vesicles showed a ~15% dimer population, comparable with that observed here for zfVDAC2 in LDAO.

The interspin distance distributions centered on ~30 Å observed in DEER are consistent with the interspin distance expected from modeling of the R1 side chain in the x-ray structure presented here, but the broad distributions indicate significant protein flexibility. The main probability distributions for VDAC2 232R1 in LDAO and bicelles are similar but not identical, showing that the structure of the dimer in both environments is similar.

The 55 Å population for the protein in bicelles, but not in LDAO detergent micelles, cannot arise from a homodimer. A possible oligomeric configuration could be drawn from crystal packing analysis of mVDAC1 crystals originating in bicelles (7). This structure reveals a more elaborate hexameric arrangement mimicking the native oligomeric packing observed in EM (44) and atomic force microscopy (45) images of the outer mitochondria membrane. Intramolecular distances between side chain 232R1 in this hexamer identify a likely origin for both the ~30 Å and 55 Å populations (Fig. 10).

Comparison of the CW EPR spectra of residue 232R1 in LDAO and bicelles shows an increase in the relative population of a more mobile state in LDAO (Fig. 7C), and osmolyte perturbation indicates that the two-component spectra probably arise from multiple conformational states in equilibrium. Thus, the small differences that do exist in the distance distributions for LDAO and bicelles may be due to differences in conformational flexibility of the protein at or near residue 232R1.

The cross-linking experiments strongly support the validity of the suggested dimer interface in zfVDAC2. The two cysteine mutants designed outside of the dimer interface displayed no cross-linking, whereas mutants in the dimer interface displayed distinct levels of cross-linking. The highest levels of cross-link-
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were observed for mutants in long flexible loop regions (D50C with S–S located 11.1 Å apart and S52C with an S–S distance of 6.8 Å and for an internal mutant located in the cytosolic side of β1 (V27C), which is the closest cysteine mutant in this set of cross-linking experiments (S–S distance of only 3.7 Å). The other three mutants on the mitochondrial IMS side that displayed lower cross-linking efficiencies were designed in rigid β-strands or just at the interface between short loops and β17 and β18 situated at relatively long S-S distance ranges of 6.4–8.9 Å.

Although the present study advances our structural understanding in the VDAC field, in particular VDAC2, there are still many unanswered questions that will need to be addressed. Perhaps the varied phenotype is not purely structural but depends on different affinities to partner proteins, expression patterns, subcellular localization of the distinct isoforms, or differences in the intrinsic flexibility of functionally important structural elements between the different isoforms. There is plenty of evidence for different physiological roles of the three isoforms, but the search for the underlying mechanisms has to be further explored. The structural characterization of VDAC2, however, provides blueprints to test many hypotheses and identifies a loop between strands 1 and 2 as a plausible dynamic structural element, as judged by changes in its relative position in structures of different VDAC isoforms, and identifies an area with a change in the electrostatic surface potential between zfVDAC2 and mVDAC1.

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