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Shamim Naghdi  
Thomas Jefferson University

Péter Várnai  
Semmelweis University

György Hajnóczky  
Department of Pathology and Cell Biology, Thomas Jefferson University, Philadelphia, PA 19107, USA

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Motifs of VDAC2 required for mitochondrial Bak import and tBid-induced apoptosis

Shamim Naghdi,a Péter Várnai,b and György Hajnóczkya,1

MitoCare Center for Mitochondrial Imaging Research and Diagnostics, Department of Pathology, Anatomy, and Cell Biology, Thomas Jefferson University, Philadelphia, PA 19107; and Department of Physiology, Faculty of Medicine, Semmelweis University, Budapest 1094, Hungary

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Voltage-dependent anion channel (VDAC) proteins are major components of the outer mitochondrial membrane. VDAC has three isoforms with >70% sequence similarity and redundant roles in metabolite and ion transport. However, only Vdac2−/− (V2−/−) mice are embryonic lethal, indicating a unique and fundamental function of VDAC2 (V2). Recently, a specific V2 requirement was demonstrated for mitochondrial Bak import and truncated Bid (tBid)-induced apoptosis. To determine the relevant domain(s) of V2 involved, VDAC1 (V1) and V2 chimeric constructs were created and used to rescue V2−/− fibroblasts. Surprisingly, the commonly cited V2-specific N-terminal extension and cysteines were found to be dispensable for Bak import and high tBid sensitivity. In gain-of-function studies, V2 (123–179) was the minimal sequence sufficient to rendered V1 competent to support Bak insertion. Furthermore, in loss-of-function experiments, T168 and D170 were identified as critical residues. These motifs are conserved in zebrafish V2 (zfV2) that also rescued V2-deficient fibroblasts. Because high-resolution structures of zfV2 and mammalian V1 have become available, we could superimpose these structures and recognized that the critical V2-specific residues help to create a distinctive open "pocket" on the cytoplasmic surface that could facilitate Bak recruitment.

mitochondria | VDAC2 | Bak | tBid | apoptosis

Voltage-dependent anion channel (VDAC) proteins provide the primary pathway for metabolite and ion flux through the outer mitochondrial membrane (OMM) (1, 2). The VDAC has three isoforms in vertebrates, which show >70% sequence similarity and have a broad and overlapping tissue distribution. The three isoforms can substitute for each other in the metabolite and ion flux (2). However, they also have nonredundant roles in cell function. Striking evidence of an isoform-specific role of VDAC2 (V2) is that embryos of homozygous deletion of Vdac2 (V2−/−) mice are embryonic lethal, indicating a unique and fundamental function of VDAC2 (V2). Recently, a specific V2 requirement was demonstrated for mitochondrial Bak import and truncated Bid (tBid)-induced apoptosis. To determine the relevant domain(s) of V2 involved, VDAC1 (V1) and V2 chimeric constructs were created and used to rescue V2−/− fibroblasts. Surprisingly, the commonly cited V2-specific N-terminal extension and cysteines were found to be dispensable for Bak import and high tBid sensitivity. In gain-of-function studies, V2 (123–179) was the minimal sequence sufficient to render V1 competent to support Bak insertion. Furthermore, in loss-of-function experiments, T168 and D170 were identified as critical residues. These motifs are conserved in zebrafish V2 (zfV2) that also rescued V2-deficient fibroblasts. Because high-resolution structures of zfV2 and mammalian V1 have become available, we could superimpose these structures and recognized that the critical V2-specific residues help to create a distinctive open "pocket" on the cytoplasmic surface that could facilitate Bak recruitment.

Recent evidence suggests that V2 is involved in Ca2+ transport by cardiac mitochondria (6–8), and specifically interacts with several endogenous cytoplasmic and mitochondria-associated proteins, including steroidogenic acute regulatory protein (StAR) (9), glycerol synthase kinase 3 beta (GSK3β) (10), O-GlcNAc transferase (11), ryanodine receptor 2 (8), tubulin (12), viral protein 5, (13, 14), and drugs [erastin (15) and esfevin (7)]. These interactions facilitate mitochondrial targeting, transport, andsignaling processes, and, interestingly, they seem to have striking effects on cell survival. However, the most investigated interaction of V2 is with Bak, a proapoptotic Bcl-2 family protein that is central to OMM permeabilization, allowing the release of intermembrane space proteins like cytochrome c (cyto c) and AIF to the cytosol to induce apoptosis (16, 17).

OMM permeabilization can occur when death receptors like TNF-α receptors are engaged and trigger caspase-8 activation to cleave Bid, a BH3-only proapoptotic Bcl-2 family member. Relocation of truncated Bid (tBid) (18, 19) to the OMM and interaction with Bcl-2 family proteins (either proapoptotic and/ or antiapoptotic) lead to a conformational change and subsequent homo-oligomerization or hetero-oligomerization of Bak (19, 20) and another proapoptotic Bcl-2 family protein, Bax (21), providing the means of exit for intermembrane space proteins across the OMM (22–24). At least in several cell types, the rapid kinetics of tBid-induced OMM permeabilization are primarily dependent on Bak (25), presumably because Bak usually resides in the OMM, whereas the mostly cytoplasmic Bax first needs to translocate to the mitochondria (26, 27). Distribution of Bak among individual mitochondria is uneven in cells lacking the OMM fusion proteins Mfn1/2, indicating that OMM fusion is required for Bak distribution (28). However, the availability of Bak in the OMM is primarily controlled by insertion of Bak from cytoplasm to the OMM.

Bak is a tail-anchored protein, but it is different from other tail-anchored proteins like Bcl-2 or Bcl-xL in that unmasking of its C-terminal tail domain and insertion to the mitochondria require V2 (25, 29). In the OMM, Bak was also shown to associate with V2 and to be part of a large complex (3). The interplay between Bak and V2 is central to mitochondrial apoptosis and might contribute to the vital significance of V2 (3, 25, 30–32).

Significance

Voltage-dependent anion channel 2 (VDAC2), like other VDAC proteins, transports solutes across the outer mitochondrial membrane, but it is the only isoform that supports mitochondrial import of cell death-executing proteins, Bak/Bax. To address the underlying mechanism, we created chimeras of VDAC2 and VDAC1 and tested their competency to rescue Bak insertion and apoptosis in VDAC2-deficient cells. Our gain-of-function studies revealed that the middle domain (123–179 aa) is required for rescue. Loss-of-function studies identified two critical residues (T168, D170). Finally, applying these clues to recently solved structures we identified a discrete site on the cytoplasmic side of VDAC2’s pore that likely supports Bak insertion and apoptosis.

Author contributions: S.N., P.V., and G.H. designed research; S.N. performed research; P.V. contributed new reagents/analytic tools; S.N. and G.H. analyzed data; and S.N. and G.H. wrote the paper.

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1To whom correspondence should be addressed. Email: gyorgy.hajnoczky@jefferson.edu.

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Differently, however, the biophysical studies show a β-barrel formed by 19 β-sheets with the N-terminal α-helix lying inside the pore. The α-helix is highly flexible and is likely to be associated with VDAC channeling activity for the transfer of metabolites like ATP (40, 41). The literature about the orientation of the VDACs is divergent; however, a recent work by De Pinto and coworkers (42) shows that the C terminus faces the intermembrane space. The only crystallography study of V2 used zebrafish V2 (zfV2) that had >90% sequence similarity to mammalian V2. This work revealed few differences between V1 and V2, mostly confined to the cytoplasmic loops (37).

In the present project, we have studied the domain(s) of V2 that are necessary and/or sufficient for Bak targeting to the OMM and Bak-mediated cytochrome c release. To this end, the few sequence differences between mammalian V1 and V2 were systematically analyzed by creating a series of chimeric constructs. The functional activity of each construct was tested by evaluating the rescue of Bak’s mitochondrial insertion and tBid-induced OMM permeabilization in Vdac<sup>2<sup>−/−</sup></sup> (V2<sup>−/−</sup>) mouse embryonic fibroblasts (MEFs). Our results demonstrate that previous suspicions like the N-terminal extension and high cysteine (C) content of V2 are irrelevant. Unexpectedly, the sequences of amino acids 168–171 and 123–179 represent the minimum domain of V2, which is necessary and sufficient to provide for Bak import and rapid tBid-induced apoptosis, respectively. In combination with the results of recent structural studies, our findings allow us to propose a mechanism of how the apparently minor sequence differences between V1 and V2 can result in stark functional differences.

**Results**

**Genetic Rescue of V2<sup>−/−</sup> MEFs Provides an Approach for Studying the Requirements of Mitochondrial Bak Import and Cytochrome c Release.** We and others have provided evidence that Bak insertion to the OMM and tBid-induced OMM permeabilization are supported only by V2 among VDAC isoforms (25, 29, 30). To establish a paradigm for determining the molecular basis for this nonredundant function of V2, we used V2<sup>−/−</sup> MEFs and compared their Bak level and their sensitivity to tBid-induced cell death under rescued and nonrescued conditions. First, V2<sup>−/−</sup> MEFs were infected with V2-badenovirus (avV2), and 36–48 h after the infection, the presence of V2 was confirmed in the cells by immunoblotted membrane lysates using a polyclonal anti-V2 antibody (Fig. 1A). This antibody detects two bands at around 32–34 kDa in MEFs expressing V2; however, only the upper band is relevant because it is the one that is absent in the V2<sup>−/−</sup> MEFs and is restored in the genetically rescued cells (arrow in Fig. 1A). Immunoblotting also showed Bak in WT cells, which was low in V2<sup>−/−</sup> cells and was restored upon rescue with avV2. Sensitivity of the cells to tBid was monitored by studying cytochrome c release from mitochondria to cytosol. Blotting of the membrane and cytosolic fractions against cytochrome c showed that 25 nM tBid for 5 min fails to release cytochrome c in V2<sup>−/−</sup> MEFs but causes release in the rescued cells (Fig. 1B). Digitonin (600 μg/mL) was

![Image](https://example.com/image1.png)

**Fig. 1.** Genetic rescue studies in V2<sup>−/−</sup> MEFs. Unique N-terminal extension in V2 is neither necessary nor enough for Bak import and cytochrome c release. (A) Western blot verification of the presence of V2 and Bak in the membrane fraction of WT and V2<sup>−/−</sup> cells rescued with avV2. The arrow shows the relevant band of V2. (B) Cytochrome c release was monitored by detecting the cytochrome c level in the membrane and cytosolic fraction of nonrescued (−) and avV2-rescued (+) V2<sup>−/−</sup> permeabilized MEFs 5 min after treatment with solvent [tBid (25 nM) or digitonin (Dig, 600 μg/mL)]. TC, time control. Hsp70 and actin were used as loading controls. (C) Representative traces for the time course of ΔΨ<sub>m</sub> recorded in cell suspensions using a fluorometer. The experiment was carried out in permeabilized V2<sup>−/−</sup> (black line) or avV2-rescued V2<sup>−/−</sup> (red line) cells treated with tBid or Dig (added at 300 s, marked by an arrow). FCCP (5 μM) was added at the end of the recordings to dissipate the remaining ΔΨ<sub>m</sub>. Data are presented as the percentage of the initial ΔΨ<sub>m</sub>. TCs are the nontreated samples. (D) HA and Bak Western blot of the membrane fraction of V2<sup>−/−</sup> cells transfected (tf) with HA-tagged V2, V2(Δ1–12), V1, and V2(1–12)V1. Immunoblot (E) and quantitative analysis (F) of cytochrome c in the cytosolic fraction of V2<sup>−/−</sup> cells transfected with V2, V2(Δ1–12), V1, or V2(1–12)V1 and treated with tBid or Dig after plasma membrane permeabilization. In response to tBid, the cytochrome c band was normalized to the response to Dig in each condition (n = 3).
**Unique N-Terminal Extension in V2 Is Not Necessary for Bak Import and Cyto c Release.** To identify the motifs of V2 that are necessary and/or sufficient for Bak import and tBid-induced OMM permeabilization, the protein sequence of V2 was systematically compared with V1 and VDAC3 isoforms. The comparison showed that four of the Cαs (48, 77, 104, and 134) were mutated to asparagine (N) [called V2(48N)] (Fig. S2D). V2ΔC was added to dissipate the ΔΨm completely. These results suggest that V2ΔC MEFs provide a model allowing mutual transfection of these reagents into cells transfected with V2 and V2ΔC(-/−) and V1 (Fig. S1C). Altogether, the data confirm that only N-terminal extension of V2 supports Bak targeting and tBid-induced cyto c release. Furthermore, these findings extend the genetic evidence on the critical role of V2 in mitochondrial Bak import and tBid-induced rapid cyto c release.

**V2-Specific Cαs Are Dispensable for Bak Import and tBid-Dependent OMM Permeabilization.** Comparison of amino acid sequences in V1, V2, and VDAC3 revealed similar values for all residues except Cα, which is exclusively higher in mammalian V2 (11 vs. 2 in V1 and 6 in V3). Localization of the Cαs in the published biophysical model of VDAC (35–38) showed that four of the Cαs (48, 77, 104, and 134; shown by blue arrows in Fig. 2A) in V2 are in the loop between β-sheets. They are seemingly at the same side of the V2 molecule and might be involved in protein–protein interaction (Fig. 2A). To test their possible role in Bak recruitment, the four linker-localized Cαs (48, 77, 104, and 134) were mutated to asparagine (N) [called V2(48AC)] (Fig. S2B, Upper). Furthermore, based on similarities between structures of V1 and V2 isoforms, we also localized the Cαs in the so-called “biochemical model” of V1 (1, 34). In this model, two of the Cαs (211 and 228) are in an external loop of V2 (Fig. 2B). We hypothesized that a disulfide bond between neighboring Cαs in the loop might stabilize a suitable docking site for Bak or any relevant mediator protein.
Therefore, in another mutant, C228 was replaced with its V1-specific counterpart, which is N (N216) (Fig. S2B, Lower). Expression and the presence of the described mutants in the membrane fraction of V2\(^{\text{-/-}}\) cells expressing V2(4AC) and V2(C228N) were confirmed using an anti-V2 antibody (Fig. 2C). V2\(^{\text{-/-}}\) cells expressing V2(4AC) or V2(C228N) showed recruitment of Bak protein to the membrane (Fig. 2C) and were sensitized to tBid (Fig. 2D), indicating that the C's are not needed for the competence of V2 to support Bak recruitment/tBid-induced cyto c release.

**zfV2 Rescues Bak Import and tBid-Induced OMM Permeabilization in V2\(^{\text{-/-}}\) MEFs.** To examine further the effect of both the N-terminal extension and the high numbers of C's in the V2 sequence on the OMM targeting of Bak, we used zfV2 (37). The zfV2 has 82%/92% sequence identity/similarity with mouse V2 (mV2), but it lacks the N-terminal extension and contains only one C at the position of 127. HA-tagged mV2 and zfV2 were transfected in V2\(^{\text{-/-}}\) cells, and 48 h later, the proteins were detected in the membrane fractions using HA antibody (Fig. 2E). Bak levels in the membrane fraction were similarly elevated by zfV2 and mV2 rescue, indicating that zfV2 is capable of recruiting Bak. Furthermore, treatment of the cells with 25 nM tBid caused similar release of cyto c in the V2\(^{\text{-/-}}\) cells expressing zfV2 and mV2 (Fig. 2F). Thus, studies of zfV2 further support that the N-terminal extension and C's are not critical for Bak targeting to the OMM.

**Gain-of-Function Studies Using Chimeras of V1 and V2.** As explained above, the biochemical model (Fig. 2B) predicts a long cytoplasmic loop in the C-terminal third of V1 (34), which might be well positioned to support unmasking of the tail of Bak. To find out whether this domain is sufficient or required for Bak targeting, four chimeras were generated. To test whether the loop is sufficient, chimera 1 (Chi1) with the first two-thirds of V1 (1–185) linked to the last third of V2 (198–295) was generated. To test whether the loop is necessary, Chi3 with the first two-thirds of V2 (1–188) linked to the last third of V1 (177–283) was created. Chi2 was created by adding the 12 N-terminal residues to Chi1. Chi4 was created by deletion of this segment from Chi3 (Fig. S2C). V2\(^{\text{-/-}}\) MEF cells were transfected with the described chimeras, and the presence of relevant proteins was confirmed in the membrane fraction (Fig. 3A). The Bak level did not change in V2\(^{\text{-/-}}\) cells expressing Chi1 and Chi2, but it increased in the cells expressing Chi3 and Chi4. Furthermore, rescue of tBid-induced cyto c release was apparent only in the cells expressing Chi3 and Chi4 (Fig. 3B). This evidence shows that the last one-third of V2 (198–295), including the long external loop, is neither enough nor required for the tBid/Bak OMM permeabilization pathway. On the other hand, this result indicates that the domain relevant for Bak targeting to the OMM is located between amino acid 12 and amino acid 188 of V2. Therefore, in the next step, we generated two chimeras, Chi5 and Chi6, by substituting one-half of the V2-specific sequence of Chi4 with the relevant sequence of V1 (Fig. S2D). In Chi5, V2 (13–118) of Chi4 was replaced with the relevant sequence of V1, and in Chi6, V2 (119–188) of Chi4 was replaced with the relevant sequence of V1. The presence of the HA-tagged chimeric proteins in the membrane fraction of V2\(^{\text{-/-}}\) MEF cells expressing Chi5 and Chi6 was confirmed (Fig. 3C). The Bak level increased in the membrane fraction of cells expressing Chi5, but it was unaltered in the cells expressing Chi6 (Fig. 3C). Consistently, tBid released cyto c from V2\(^{\text{-/-}}\) cells expressing only Chi5 (Fig. 3D and E). This set of data suggests that the domain that mediates targeting of Bak to OMM is located in the middle domain of V2 (amino acids 119–188).

**V2 (123–179) Is the Shortest Sequence Sufficient for Bak Recruitment.** To narrow the region sufficient for Bak import further, the sequence of V2 in Chi5 was compared with the relevant sequence of V1. Based on clusters of different and not so similar amino acids (using Clustal W nomenclature) in the V2 sequence, we generated three constructs. In each of them, we substituted one-third of the V2-specific middle domain in Chi5 with the corresponding sequence of V1. Specifically, in Chi 7–Chi 9, amino acids 119–137, 138–161, and 162–188 of the V2 sequence were replaced with their V1 counterparts, respectively (Fig. S2E). Expression and proper targeting of the chimeras were confirmed in V2\(^{\text{-/-}}\) MEF cells expressing Chi 7–Chi 9 (Fig. 4A). In the cells expressing Chi7 and Chi8, the Bak level increased in the membrane fraction, but it did not change in the cells expressing Chi9. The tBid-induced release of cyto c also occurred in the cells expressing Chi7 and Chi8, but no release appeared in the cells expressing Chi9 (Fig. 4B). These data indicate that amino acid residues in the domain of 161–188 are needed for the Bak/cyto c release pathway. To test whether this domain is sufficient for Bak recruitment, Chi10 was generated with V2 (162–188) substituted for the relevant fragment in V1 (Fig. S2F). This mutant was expressed in the V2\(^{\text{-/-}}\) cells, but it failed to support Bak recruitment (Fig. 4A) and cyto c release after treatment with tBid (Fig. 4B). Thus, V2 (162–188) is essential, but not sufficient, for Bak targeting to the OMM. To determine the N-terminal and C-terminal ends of the region sufficient to substitute for Chi5, new chimeras were generated with various lengths of V2 in Chi5 swapped with the equivalent of V1. In Chi11, the main cluster of differences in the N-terminal end of the V2-specific sequence (119–133) in Chi5 was taken out and was replaced with the corresponding sequence of V1. In Chi12, differences in the C-terminal end of the V2 sequence of Chi5 (180–188) were eliminated by replacement with the relevant part of V1. In Chi13, both V2 (119–133) and V2 (18–188) were cut out and replaced with the relevant fractions of V1 (Fig. S2F).

![Fig. 3. Middle domain of V2 is required for mitochondrial Bak import and tBid-induced cyto c release.](image-url)

(A) HA Western blot (Upper) for the V2\(^{\text{-/-}}\) cells transfected with HA-tagged V2 or Chi1 and Bak Western blot (Lower) for the cells expressing V2 or Chi1–Chi4. (B) Cyto c Western blot of the V2\(^{\text{-/-}}\) cells transfected with V2 or Chi1–Chi4. (C) HA and Bak Western blot of the V2\(^{\text{-/-}}\) cells expressing V2, Chi5, or Chi6. V2\(^{\text{-/-}}\) expressing nontagged Chi1 was used as a negative control in an HA Western blot. Western blot (D) and quantification (E) of cyto c release in V2\(^{\text{-/-}}\) cells expressing V2, Chi5, or Chi6 are shown.
Loss-of-Function Mutations in V2: T168 and D170 Are Critical Amino Acids for Bak Targeting to the OMM. In the next step, we tried to find the critical residues supporting Bak import. Cutting the C-terminal part of V2 in Chi5 (162–188) in Chi9 totally abrogated tBid-induced cyto c release; however, cutting of V2 (180–188) in Chi12 did not change the response. That result led us to hypothesize that the most critical domain must be located in V2 (162–179). The location of this sequence at the edge of a β-sheet (β-sheet 10) in the biophysical model (Fig. 2A) strengthens this idea. In addition, cells expressing Chi7 and Chi13 show a partial response, and that result led us to assume that there might be some effective motifs in the N-terminal part of the V2 sequence in Chi5. Therefore, based on the differences between V2 and V1, which were conserved in all species, we made short mutations in the amino acid 122–179 segment of V2. To test the importance of the C terminus, M1, M2, and M3 were generated in which amino acids 168–176, 168–171, and 175–176 in V2 were replaced with the equivalent sequence of V1 in order. On the other hand, to study the effect of the N-terminal segment, M4 was made by the replacing of V2 (123–124) with the relevant sequence in V1 (Fig. S2G). The presence of mutant proteins in the membrane fraction of V2 cells expressing V2 (M1–M4) was validated (Fig. 4E and Fig. S1E). Bak showed an increase in V2Δψm cells expressing M3 (Fig. 4E) and M4 (Fig. S1E). By contrast, it did not show any alteration in the cells expressing M1 and M2. The tBid-induced cyto c release was significant in V2Δψm cells expressing M3 (similar to the cells rescued by V2). It was moderate in cells expressing M4, and it was abolished in the cells expressing M1 and M2. Also, tBid induced significant depolarization in the V2Δψm cells rescued with M3; however, it failed to change Δψm in the cells expressing M2 (Fig. 4G and H). All of the evidence showed that a small sequence of threonine (T), phenylalanine (F), aspartic acid (D), serine (S), or “TFDS” in the position of 168–171 in V2 is critical for Bak insertion and that amino acids serine/glycine (S/G) in the location of 123/124 are also relevant.

In the sequence of TFDS, amino acid F was present both in V1 and V2, so only three amino acids remained to examine. In M5–M7, T, D, and S were replaced with their equivalent in V1, which were asparagine (N), glutamic acid (E), and T (Fig. S2H). The residence of the mutants in the OMM was confirmed in V2Δψm cells expressing M5–M7 (Fig. S4). To check whether each mutant was inserted into the mitochondria, the subcellular distribution of the mutants was analyzed using immunofluorescence staining for the HA tag (red in Fig. 5B). V2Δψm cells were cotransfected with GFP-OMP25, an OMM marker (green in Fig. 5B). In all of the conditions, >90% of the red signal overlapped with the area of green signal, which means that the HA-tagged V2 mutant proteins were colocalized with the OMM marker (Fig. 5C). Bak was increased in the V2Δψm MEFs expressing M7 (Fig. 5B); however, it was similar to the negative control in cells expressing M5 and M6. Furthermore, tBid rapidly released cyto c and caused collapse of the ΔΨm in cells expressing only M7 (similar to rescued cells with V2) (Fig. 5D and E). These data demonstrated that two amino acids, including T168 and D170, are the most critical amino acids in mV2-mediated support of the tBid/Bak cell death pathway.

Structural Implications of the Domains Identified by Mutagenesis. The structure of a mammalian V2 has not been solved. Fortunately, the zfV2 structure has recently become available (37), and our data showed that zfV2, like mV2, supports the Bak/tBid pathway (Fig. 2). The regions corresponding to the sufficient motifs and necessary residues of mV2 are shown in magenta and yellow, respectively, in the zfV2 structure [Protein Data Bank (PDB) ID code 4bum] (Fig. 6A and B). The side view in Fig. 6A shows that T168 (S156 in zfV2) is almost at the end of β-sheet 10 and D170 (D158 in zfV2) is located in the beginning of the loop between β-sheets 10 and 11. The top view shows that both of the

expression and membrane localization of Chi1–Chi3 were confirmed (Fig. 4C). The Bak level in the membrane fraction and cyto c release were fully restored in the cells expressing Chi2, whereas only partial rescue took place in the cells expressing Chi1 and Chi3 (Fig. 4C and D). Time lapse monitoring of ΔΨm in adherent cells showed the same pattern of depolarization for V2Δψm cells expressing Chi2 or V2 after treatment with tBid (Fig. 4H and Fig. S1D). Altogether, these data show that V2 (123–179) is the shortest sequence that is fully supporting the recruitment of Bak and tBid-induced OMM permeabilization.
Thr167 and Glu169 are the necessary residues in mV2 for mitochondrial Bak import and tBid-induced apoptosis. (A) HA and BAK Western blot of cells expressing M5–M7. (B) Localization analysis of V2 mutants in V2−/− cells coexpressing M5–M7 and GFP-OMP25 (as an OMM marker, green signal) using anti-HA immunofluorescence (red signal). An endoplasmic reticulum (ER)-targeted protein with an HA tag (ER-HA) was used as a negative control. Scale bar on top left image is equal to 5 um. (C) Colocalization of HA immunofluorescence with mitochondria presented as the percentage of the overlap between the green and red areas and the total area of the red signal. (D) Cyto c Western blot of the V2−/− cells expressing V2 or M5–M7. (E) ΔΨm recorded as described for Fig. 4J for the V2−/− cells expressing M5–M7. Arrows show the time of the addition of tBid.

necessary residues are on the cytosolic side of the channel and the side chains of both of these amino acids are facing the inner side of the pore. The positioning of these amino acids supports the possibility that the interaction with any other proteins (either
directly with Bak or a third party needed for Bak) might be affected from the inner side of the channel. Further along this line, the S171 (T159 in zfV2) that was not critical for Bak recruitment projects to the outer surface.

Next, the crystal structure of zfV2 (111–168) was aligned with mV1, which cannot support Bak recruitment and has a crystal structure determined at a similar resolution to zfV2 (36). Comparison of the crystal structure of zfV2 with mV1 (PDB ID code 3emn) showed that N156 and E158 in V1 form extensions oriented toward each other to close the gap between them, whereas S156 and D158 diverge to create an open space in zfV2 (Fig. 6C, Upper Right). The presence of S156 and D158 in zfV2 also causes differences in the proximity of adjacent residues (Fig. 6C, Upper Left). In Fig. 6C, sticks represent the amino acids that are within a distance of 4 Å from either S156 or D158 in zfV2 (blue carbons) or either N156 or E158 in V1 (orange carbons). These adjacent amino acids have been indicated in the sequences in Fig. 6C (Lower). Based on the distance between residues in 3D structures, it seems the loops between β-sheets 10 and 11 and between β-sheets 9 and 8 in zfV2 are in closer vicinity than the equivalent loops in mV1. Based on the greater accessibility of the space defined by the V2-specific critical residues and the differential arrangement of their neighbors, it is tempting to assume that V2 might provide a more suitable docking site for Bak recruitment. Notably, the small sequence difference in the highlighted region seems to result in differential plane positioning of the loop between β-sheets 10 and 11 relative to the lipid bilayer (Fig. S3) [predicted by the Orientations of Protein in Membranes (OPM) database (44)], providing another factor for differential isoform-specific interaction with other proteins.

Discussion

In this study, we identified by mutational analysis the domain of V2, which plays a role in mitochondrial Bak import and tBid-Bak–mediated apoptosis. Although the N-terminal extension and the higher C content were the suspects previously, we demonstrated that these unique features of the V2 structure are dispensable. Our gain-of-function studies revealed that V2 (123–179) replacement in V1 is sufficient to support the Bak pathway. Furthermore, loss-of-function studies demonstrated that even a small change at T168 or D170 of V2 is sufficient to arrest the Bak pathway and an alteration at S123 or G124 also suppresses this pathway. Based on the available crystal structure, we predict that these residues might help to create a potential docking site for Bak or an adaptor protein. Thus, our findings come together...
to support an unexpected model for the distinctive, and potentially vital, apoptotic function of V2.

Bak and Bax use their C-terminal α-helix (α9) to target the OMM. Bak is constitutively present in the OMM and mediates the acute phase of Bid-induced OMM permeabilization, whereas Bax, which is predominantly cytosolic in healthy cells, enters and oligomerizes at the OMM more slowly (25, 45). Differently from other C-terminal anchored proteins, Bax requires V2 for insertion of its hydrophobic C terminus to the OMM (29), and Bax targeting also seems to depend on the presence of either Bak (the hetero-oligomerization partner of Bax) or V2 (30). Furthermore, fusion of the isolated C terminus of Bak to the C terminus of GFP results in mitochondrial targeting of GFP independent of V2. Thus, V2 might interact with a remote site on Bak to unmask a9 that would otherwise be unavailable for insertion to the OMM in the context of the full-length Bak (29, 46). However, how the structure of V2 supports the interaction with Bak and insertion of this protein to the OMM had not been investigated.

A mystery, and a unique opportunity for mutational analysis, was presented by the fact that the three VDAC isoforms display high sequence similarity, but only V2 can facilitate Bak localization to the OMM. Thus, to ensure that the β-barrel channel structure is maintained in the constructs and to avoid protein misfolding, we did not use deletion and created chimeric constructs by V2-specific substitutions in V1 and by V1-specific substitutions in V2. In addition, because of the lack of an antibody that can recognize the chimeras, we decided to use C-terminally HA-tagged constructs. Although a previous work described mitochondrial toxicity, fragmentation, and a drop in ΔΨm when C-terminally modified V1 was expressed and raised concern about C-terminal tagging of β-barrel channels (47), we did not observe any of the above-described problems and, most importantly, we had proper mitochondrial localization and rescue of function with expression of either V2-HA or nontagged V2 in V2α−/− MEFs.

We first focused on the most striking sequence differences between V2 vs. V1 and VDAC3, which are the extended N terminus and the relatively high C content of V2. The possible role of these V2-specific features was also suggested by the facts that (i) based on biophysical model, the N terminus is in the pore (35–39, 41) and might interact with cytoplasmic proteins, and (ii) the C's have been implicated in strengthening the channel interaction with their environment and in stabilization of the channel (48). However, we found that the N-terminal extension and the surplus C's are dispensable for Bak import and tBid-mediated apoptosis (Fig. 2).

The remaining scattered and subtle sequence differences were evaluated initially with gain-of-function studies, which concluded that 123–179 is the minimum sequence of V2 sufficient to lend competency to V1 to target Bak to the OMM (Figs. 3 and 4). If either the N-terminal or the middle third of this sequence is switched back to V1, the gain of function is partial, whereas reversing back 161−179 to V1 precludes any Bak recruitment and tBid-induced cyto c release (Fig. 4). When the few isoform-specific residues in this sequence were further tested in loss-of-function studies, replacement of V2 (123–124) with the corresponding sequence of V1 caused partial loss of function, whereas replacement of V2 (168–171) led to complete loss of function (Fig. 4). Finally, point mutations at position T168 or D170 of V2 caused essentially complete loss of function (Fig. 5). Thus, multiple residues in the 123–179 sequence contribute to formation of the relevant conformation, and T168 or D170 is an essential residue.

Because this study is, to our knowledge, the first systematic mutational analysis of V2, there is little literature to provide a context for our results. To predict how the essential residues contribute to a potential protein-protein interaction site, we considered the available structures. Although mammalian V2 remains to be studied, the structure of zfV2 has just been solved, and we found that zfV2 rescues Bak import and tBid sensitivity in V2−/− MEF mitochondria. In zfV2, the sufficient sequence spans from 111 to 167 and the essential residues are S156 and D158 (Table 1). As Fig. 6 shows, S156 is almost at the end of β-sheet 10 and D158 is located at the beginning of the loop between β-sheets 10 and 11. This loop is at the cytosolic side of the channel (42), and the side chains of both of these amino acids are facing the inner side of the pore, indicating possible interaction with cytoplasmic proteins (either directly with Bak or with an adaptor) from the inner side of the channel. Consistent with the proposed relevance of the inner surface, mutation of S171 in mV2, the equivalent of T159, whose side chain in mV1 is directed to the outer surface of the channel, did not affect Bak recruitment. Interestingly, S123 and G124 (S111 and G112 in zfV2), the replacement of which by V1-specific residues caused partial loss of Bak-related activity (M4 mutant in Fig. 4), are also at the cytosolic end of a β-sheet (β-sheet 7) (Fig. 6d, marked as 111) and so are well positioned to contribute to the formation of a surface domain that supports Bak recruitment.

Because the structures of both zfV2 and mV1 are available at very similar resolutions, we could examine the structural difference caused by the critical residues. Despite the similarity between zfV2 and mV1 in the critical amino acids, comparison of the superimposed structures showed that N156 and E158 in V1 form a closed area, whereas S156 and D158 in zfV2 form the edge of an open “cup.” The wider space created by the side chains of S156 and D158 might favor protein–protein interaction. In addition, this small sequence difference might lead to a change in positioning of the loop between β-sheets 10 and 11 relative to the lipid membrane [as has been predicted by the OPM database (44)], and thereby might affect the affinity for interacting with other proteins. Notably, a recent computational study concluded that A294 in V2 thereby might affect the affinity for interacting with other proteins.

Table 1. Alignment of the “sufficient” sequence of V2 from different species and mV1

<table>
<thead>
<tr>
<th>V2/V1 in different species</th>
<th>Sequences</th>
</tr>
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<tbody>
<tr>
<td>mV2</td>
<td>123 SGKIKSAFAYRSCINLGCUCVDFDFAGSPAIHSGAVGYGWLAGYQM</td>
</tr>
<tr>
<td>hvV2</td>
<td>122 SGKIKSKYRSCINLGCUCVDFDFAGPIAHDGAVGYGWLAGYQM</td>
</tr>
<tr>
<td>zfV2</td>
<td>111 SGKITYKSPFVNLCUCVDFDFAGPIIAHDGAVGYGWLAGYQM</td>
</tr>
<tr>
<td>mV1</td>
<td>111 NAKIGYSKRYREIMLCUCVDFDIAGPSIRGALYDGWLAGYQM</td>
</tr>
</tbody>
</table>

Blue highlight shows the necessary amino acids in V2s and their equivalent in mV1.

Because our experiments provided clearcut evidence that the V2-specific C terminus is not required for rescuing Bak targeting in V2−/− MEF mitochondria (Fig. 3). In addition to spatial positioning of the residues, posttranslational modification of T168 or D170 might be the reason for the competence of V2 to support mitochondrial Bak import. Indeed, Palaniappan et al. (11) showed that V2 might be a substrate for O-linked β-N-acetylglucosaminidation; however, our amino acids of interest have not been reported as the location for glycosylation. Furthermore, to study the posttranslational modification of V2, we used tandem MS of V2-HA enriched by immunoprecipitation. Unfortunately, the spectra recorded upon digestion with both trypsin and endoproteinase Asp-N did not reveal our segment of interest. Finally, the possibility of V2 phosphorylation
was tested by searching the phosphosite database (50). However, there are no data available for amino acids 132–180 of V2. Lack of data for this part of V2 might be because of a technical difficulty in protein enrichment or because conventional trypsin digestion does not provide fragments covering the sequence of interest. Thus, testing of the possible role of posttranslational modifications needs further refinement of the technology.

Collectively, our results have addressed the question of what V2-specific structural differences allow only this VDAC isoform to control mitochondrial Bak import and, in turn, the acute phase of Bid-mediated apoptosis. Beyond the fatal outcome of murine V2 deletion, the human disease relevance of V2 dysregulation is supported by multiple protein and gene expression datasets showing specific V2 up-regulation in human hepatocarcinomas. Therefore, the structural basis of the V2-Bak interplay might become important in identifying new drug targets. Furthermore, it is of interest to find out whether the V2-specific sequence supporting the Bak pathway is also relevant for the recently noticed interactions of V2 with other proteins, including StAR and GSK3β.

Materials and Methods

Plasmids. The mV2-HA and human VI (hV1)-HA were cloned into pEYFPN1. The rest of the constructs were generated using relevant restriction enzymes for V2 and V1 sequences (Fig. S4). Cloning was performed using standard molecular biology methods. All of the clones were confirmed with sequencing. All of the constructs have been linked to a chemiluminescent tag (human influenza HA), except V2(A2C) and V2(C228N). Due to the similarity between hV1 and mV1 (with only four not so different amino acids) and the accessibility of hV1, we used hV1 in our study. Mitochondrial matrix targeting sequence linked to YFP, Mitomatrix-YFP (Clontech) and GFP-OOMP25 targeting sequence of OOMP25 linked to GFP, GFP-OOMP25 (from Manuel Rojo, Université Pierre et Marie Curie, Bordeaux, France) used as mitochondrial markers and cyto c-GFP have been described previously (28).

Cell Culture, Transfection, and Electroporation. MEFs were cultured in DMEM supplemented with 10% (vol/vol) FBS, penicillin and streptomycin, and nonessential amino acids (Invitrogen). Constructs were introduced to the cells (4.5–6 × 10^5 cells per 300 μl) by electroporation using BTX-830 square-pulse generator (BTX) (25). Fluorescence-based sorting was performed 48 h after electroporation using 15–20 × 10^4 cells cotransfected with a VDAC construct or pcDNA and Mitomatrix-YFP or GFP-OOMP25. Infection with V2 adenovirus (Vector Biolabs) was done using 200 particles per cell for 48 h. 2 × 10^6 cells per well were plated onto glass-bottomed 96-well plates.

ΔΨm and Cyto c Release Assay in Suspensions of Permeabilized Cells. ΔΨm was measured in suspensions of permeabilized cells using a fluorometer (Delta \(25^\circ\)C) and V2(C228N). Due to the similarity between hV1 and mV1, we used hV1 in our study. Mitochondrial matrix targeting sequence linked to YFP, Mitomatrix-YFP (Clontech) and GFP-OOMP25 targeting sequence of OOMP25 linked to GFP, GFP-OOMP25 (from Manuel Rojo, Université Pierre et Marie Curie, Bordeaux, France) used as mitochondrial markers and cyto c-GFP have been described previously (28).

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