In-cell structures of conserved supramolecular protein arrays at the mitochondria–cytoskeleton interface in mammalian sperm

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Mitochondria–cytoskeleton interactions modulate cellular physiology by regulating mitochondrial transport, positioning, and immobilization. However, there is very little structural information defining mitochondria–cytoskeleton interfaces in any cell type. Here, we use cryofocused ion beam milling-enabled cryoelectron tomography to image mammalian sperm, where mitochondria wrap around the flagellar cytoskeleton. We find that mitochondria are tethered to their neighbors through intermitochondrial linkers and are anchored to the cytosome through ordered arrays on the outer mitochondrial membrane. We use subtomogram averaging to resolve in-cell structures of these arrays from three mammalian species, revealing they are conserved across species despite variations in mitochondrial dimensions and cristae organization. We find that the arrays consist of boat-shaped particles anchored on a network of membrane pores whose arrangement and dimensions are consistent with voltage-dependent anion channels. Proteomics and in-cell cross-linking mass spectrometry suggest that the conserved arrays are composed of glycerol kinase-like proteins. Ordered supramolecular assemblies may serve to stabilize similar contact sites in other cell types in which mitochondria need to be immobilized in specific subcellular environments, such as in muscles and neurons.

In many cell types, mitochondria collectively form a dynamic network whose members divide, fuse, and communicate with one another (1–3). Through interactions with the cytosome, mitochondria are transported—sometimes across large distances—and positioned in response to dynamic stimuli (4, 5). Interactions with the cytosome can also restrict mitochondria to specific subcellular regions. In neurons, axonal mitochondria can be immobilized by interactions with the microtubule or actin cytosome (6–8). In cardiac and skeletal muscle, mitochondrial distribution is regulated by interactions with myofibrils and intermediate filaments (9, 10). However, despite the prevalence of intermitochondria and mitochondria–cytoskeleton interactions and their integral roles in cellular function, there is very little information on the molecular architectures of these interaction sites in any cell type. One of the most striking mitochondrial configurations occurs in amniote sperm, in which mitochondria are arranged in a spiral around the axoneme, defining a region called the midpiece (11, 12). Mitochondria are among the few organelles retained in sperm throughout their maturation, during which they otherwise lose most of their cytoplasm and organelles en route to becoming highly streamlined cells specialized for finding and fusing with the egg. The extensive mitochondrial sheath in amniote sperm may be an adaptation needed to power the large, long flagellum in these lineages. Variations in midpiece morphology affect sperm motility and competitiveness (13, 14), and different species rely on energy from mitochondrial respiration to different extents (15, 16), warranting comparative studies of mitochondrial structure across species.

The core of the midpiece is the flagellar cytosome, composed of the microtubule-based axoneme and accessory elements called outer dense fibers (ODFs). A poorly characterized network of cytosome filament called the submitochondrial network of membrane pores whose arrangement and dimensions are consistent with voltage-dependent anion channels. Proteomics and in-cell cross-linking mass spectrometry suggest that the conserved arrays are composed of glycerol kinase-like proteins. Ordered supramolecular assemblies may serve to stabilize similar contact sites in other cell types in which mitochondria need to be immobilized in specific subcellular environments, such as in muscles and neurons.

mitochondria–cytoskeleton contacts | cryoelectron tomography | cryo-FIB milling | cross-linking mass spectrometry | subtomogram averaging

Significance

Spatial organization of mitochondria is vital for cellular function. In many specialized cell types, mitochondria are immobilized at specific subcellular loci through interactions with the cytosome. One of the most striking mitochondrial configurations occurs in mammalian sperm, where mitochondria wrap around the flagellum. Malformation of the mitochondrial sheath causes infertility, but the molecular structures underlying this intricate arrangement are unknown. Here, we analyzed the mitochondrial sheath in sperm from three mammalian species. We find that although mitochondrial dimensions and cristae architecture vary across species, molecular assemblies mediating intermitochondria and mitochondria–cytoskeleton interactions are conserved. These findings yield important insight into sperm physiology and evolution and are relevant for other polarized cell types, such as muscles, neurons, photoreceptors, and hair cells.

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The mitochondrial sheath occurs late in spermiogenesis and involves an intricately choreographed series of events (23, 24). Initially, spherical mitochondria are broadly distributed in the cytoplasm. Mitochondria are then recruited to the flagellum, forming ordered rows along the flagellar axis. Finally, mitochondria elongate and twist around the axoneme. While our understanding of the molecular details of these processes is cursory at best, studies on gene-disrupted mice have implicated several proteins in mitochondrial sheath morphogenesis. For instance, mice expressing mutant forms of kinesin light chain 3 (KLC3) have malformed midpieces, hinting at a role for microtubule-based transport (25). Other examples are the voltage-dependent anion channels (VDACs), which are highly abundant mitochondrial proteins that mediate transport of metabolites, ions, and nucleotides like ATP across the OMM (26). Male mice lacking VDAC3 are infertile, and their sperm cells have disorganized mitochondrial sheaths (27), so VDACs may also have unappreciated roles in mitochondrial trafficking: indeed, KLC3 binds mitochondria through VDAC2 (25). Similarly, disrupting sperm-specific isoforms of glyceral kinase (GK) leads to gaps in the mitochondrial sheath despite proper initial alignment of spherical mitochondria (28, 29). Mice lacking spermatogenesis-associated protein 19 (SPATA19) (30) and glutathione peroxidase 4 (GPX4) (31, 32) also have structurally abnormal mitochondria.

Here, we use cryofocused ion beam (cryo-FIB), milling-enabled cryo-electron tomography (cryo-ET) to image the mitochondrial sheath in mature sperm from three mammalian species. We take advantage of the uniquely multiscale capabilities of cryo-ET to unveil aspects both of the overall organization of the mitochondrial sheath and of the molecular structures important for its assembly. We find that mitochondria are tethered to their neighbors through intermitochondrial linkers and to the underlying cytoskeleton through conserved protein arrays on the OMM. These arrays were first described by deep-etch, freeze-fracture EM in guinea pig sperm (20). Here, we resolve the three-dimensional structures of the OMM arrays in a near-native state, we imaged sperm thinned by cryo-FIB milling (Fig. 1 G–L). This revealed unexpected diversity in the internal ultrastructure of mitochondria across mammalian species, especially in terms of cristae morphology. Horse sperm mitochondria have an expanded intermembrane space and a condensed matrix (Fig. 1 I and J). Mouse sperm mitochondria have an expanded matrix, with a narrow intermembrane space and thin cristae (Fig. 1 K and L). Pig sperm mitochondrial morphology is intermediate (Fig. 1 G and H), and although the mitochondrial matrix was dense, we could identify individual complexes that resembled ATP synthase on cristae of FIB-milled mitochondria (SI Appendix, Fig. S2A and B), which was confirmed by subtomogram averaging (SI Appendix, Fig. S2B).

Interspecies differences in cristae morphology correlate with measurements of matrix volume relative to mitochondrial volume (SI Appendix, Fig. S2D). In this regard, horse sperm mitochondria resemble “condensed” mitochondria, which correlate with higher rates of oxidative activity in a number of different cell types, including developing germ cells, neurons, and liver (36–38). Indeed, horse sperm are dependent on oxidative phosphorylation (39), whereas pig (16) and mouse sperm (40, 41) are thought to rely largely on glycolytic mechanisms.
Fig. 1. Mitochondrial dimensions and cristae organization vary across species. (A–F) Slices through VPP cryotomograms (Left) and corresponding three-dimensional segmentations (Right) of mitochondria from the start (A–G) or middle (D–J) of the midpiece from pig (A and D), horse (B and E), and mouse (C and F) sperm. (G–L) Slices through cryotomograms of FIB-milled pig (G and H), horse (I and J), and mouse (K and L) sperm midpieces. Right panels show digital zooms of the regions boxed out in the Left panels. The OMM is traced in green, the inner mitochondrial membrane in yellow, and the plasma membrane in blue. Arrowheads indicate intermitochondrial linker complexes. Labels: nuc, nucleus; sc, segmented columns; m, mitochondria; odf, outer dense fibers; dc, distal centriole; ax, axoneme; mtd, microtubule doublets; cpa, central pair apparatus; and pm, plasma membrane. (Scale bars: [A–L] Left panels, 250 nm, and [G–L] Right panels, 100 nm.)

Ordered Protein Arrays at the Mitochondria–Cytoskeleton Interface Are Conserved across Species. To determine how mitochondria interact with the flagellar cytoskeleton, we imaged the mitochondria–cytoskeleton interface in guinea pig sperm (20) and mouse sperm (21) and thin-section EM on golden hamster sperm (19) uncovered an ordered array of particles arranged in a ladder-like pattern on the axoneme-facing surface of sperm mitochondria. We found these arrays on the axoneme-facing surface of the OMM in our tomograms as well (Fig. 2A, yellow box) across all three species and along the entire midpiece (SI Appendix, Fig. S3 A–F). We observed direct interactions between the arrays and either the ODFs or the cytoskeletal filaments surrounding the ODFs (Fig. 2B and C and SI Appendix, Fig. S4 A–L), indicating that these arrays tether mitochondria to the midpiece cytoskeleton. The highly ordered arrays are absent from the plasma membrane surface (Fig. 2A, red box), which is consistent with freeze-fracture EM studies that detect only very short rows scattered randomly across this surface (20).

We then aligned and averaged subvolumes containing the protein arrays and the underlying OMM from the axoneme-facing surface (Fig. 3 and SI Appendix, Table S1). Our averages revealed ∼22-nm-long, twofold-symmetric, boat-shaped structures connected via four densities to a porous membrane (Fig. 3 and SI Appendix, Fig. S3 G–I). Each boat-shaped particle rises ∼5 nm above the membrane and consists of two tîle-shaped densities arranged end-to-end. The boat-shaped structures form rows in which each particle is related to its closest neighbors by a ∼10-nm translation perpendicular to the particle long axis and a ∼6-nm shift along this axis, yielding a center-to-center spacing of 10 nm.
of ∼12 nm (Fig. 3 D–F). Each row is oriented ∼120° to the long axis of the flagellum, and adjacent rows are spaced ∼12 nm apart, forming extensive arrays on the axoneme-facing surface of the OMM (Fig. 3G). Remarkably, the averages we obtained from the three species were highly similar, both in terms of individual particle dimensions and in terms of their supramolecular arrangement (Fig. 3 and SI Appendix, Fig. S3). This conservation suggests that these arrays are a crucial structural element of the mitochondrial sheath.

Our averages revealed that the OMM underlying the protein arrays is studded with ∼3 to 4-nm pores arranged in a pseudo-lattice with a center-to-center spacing of ∼5 nm (Fig. 3 A–C and SI Appendix, Fig. S3 G–I). These pore sizes are consistent with the diameters of the VDACs, which are known to form ordered arrays in the OMM (45–48). VDACs are known to localize to the sperm midpiece (49, 50); indeed, our label-free quantitative proteomics experiments show that VDAC2 and VDAC3 are among the most abundant OMM proteins in pig sperm (SI Appendix, Table S2). Furthermore, the lattice dimensions in our averages closely match those of VDAC in purified Neurospora OMM (45, 51). The lattice can be modeled by fitting multiple copies of the VDAC2 crystal structure (52) (SI Appendix, Fig. S5A). We oriented VDAC2 in the membrane plane based on its known topology (53, 54); however, at the current resolution, we cannot determine the orientation around the pore axis. Thus, in our model, each boat-shaped particle stretches across eight VDAC molecules (Fig. 3).

**Fig. 2.** Ordered protein arrays on the OMM interact with the cytoskeleton. (A) Slice through a cryotomogram of an FIB-milled, horse sperm midpiece showing mitochondria (mito), the mitochondrial reticulum (smr) ODFs (pdf), microtubule doublets (mt), and the central pair apparatus (cpa). Note how individual complexes (like the radial spoke, rs) are visible in the raw tomogram. The ordered protein array is only found on the axoneme-facing surface (yellow) of midpiece mitochondria and not on the plasma membrane-facing surface (red). (B and C) Slices through a cryotomogram of an FIB-milled, horse sperm midpiece showing how the array directly interacts with the submitochondrial reticulum to anchor mitochondria to the flagellar cytoskeleton (arrowheads). In Right panels, the OMM is traced in green, the inner mitochondrial membrane in yellow, and the plasma membrane in blue. (Scale bars: [A] Left, 250 nm, inserts, 100 nm, and [B and C] 100 nm.)

**GK-Like Proteins Are Probable Constituents of the Conserved Arrays at the Mitochondria–Cytoskeleton Interface.** To search for possible constituents of the protein arrays on the OMM, we used in-cell XL-MS (55, 56) to find potential VDAC2/VDAC3 interaction partners on the OMM (Fig. 4). We treated pig sperm cells with the cross-linker disuccinimidyl sulfoxide (DSSO), which covalently links free lysines that are within ∼3 nm (Ca-Ca) of each other. To increase confidence, we screened for protein interactions supported with at least two cross-link spectral matches (CSMs) (Materials and Methods).

We first screened candidate proteins based on their known subcellular localizations (Fig. 4A). VDAC2/VDAC3 cross-linked to mitochondria-associated proteins as well as to sperm head–associated proteins. This is consistent with immunofluorescence studies localizing VDAC2/VDAC3 both to the midpiece and to the acrosome, a large vesicle capping the anterior sperm nucleus (49, 50, 57). Of the proteins in the mitochondria-associated interaction hub, four proteins are particularly noteworthy because they are known to localize to the OMM and because their disruption results in dysplasia of the mitochondrial sheath: armadillo repeat-containing protein 12 (ARMC12) (58), SPATA19 (30), GPX4 (31, 32), and GK (28, 29).

To distinguish among these candidates, we compared the location of the cross-links with the known topology of VDAC in the OMM (53, 54). GPX4 would interact on the side facing the intermembrane space, whereas SPATA19, ARMC12, and GK would interact on the cytoplasmic face. Although we detect ARMC12 in mature pig sperm, in the mouse, it is only present in developing spermatids and disappears from mature sperm (58), which is inconsistent with the fact that the OMM arrays are present in cauda epididymal mouse sperm. SPATA19 and GK are both highly abundant (SI Appendix, Table S2), as would be expected for proteins forming extensive arrays. Assuming an average protein density of ∼1.43 g/cm³ (59), which corresponds to ∼0.861 Da/Å³, we estimate that each boat-shaped particle in the array has a molecular weight of ∼250 kDa. SPATA19 is a small protein with an estimated molecular weight of ∼18 kDa. To fit into our EM densities, it must either be present in
multiple copies or form a complex with other proteins. In contrast, GK has an estimated molecular weight of ∼60 kDa and is known to form S-shaped dimers (∼120 kDa) that are conserved from bacteria (60, 61) to eukaryotes (62, 63).

To build a GK–VDAC model based on our subtomogram average, we used rigid-body fitting to place two GK dimers end-to-end into a boat-shaped density (SI Appendix, Fig. S5B and Fig. 4B). These fits defined a clear orientation for GK, with the N termini pointing upwards and the C-terminal helices facing the OMM (Fig. 4B). To validate our fits, we mapped the cross-linked lysines onto the resulting model (Fig. 4C). All cross-links were between the cytosolic face of VDAC2 and the OMM-facing surface of GK, which is consistent with the orientation expected from our fits. These fits show that the molecular dimensions and relative positions of candidate proteins are consistent with our subtomogram average maps, but we cannot define specific interaction sites at the current resolution. We also attempted to analyze interactions between the putative GK-like proteins and the underlying cytoskeleton, and although we detect cross-links on the putative cytoskeleton-facing side of GK (SI Appendix, Fig. S6), interpretation is complicated by the fact that the precise protein compositions of the submitochondrial reticulum and the ODFs are unclear.

Fig. 3. Ordered protein arrays at the mitochondria–cytoskeleton interface are conserved across species. (A–C) Subtomogram averages of the protein arrays and underlying OMM after applying twofold symmetry (note that density is black). (D–F) Isosurface renderings of the subtomogram averages in A–C with boat-shaped particles in gray and the OMM in green. (G, Left) Segmentation of the tomogram shown in Fig. 2A, with the OMM in green, the IMM in yellow, microtubule doublets in blue, and the cpa in pink. Subtomogram averages of boat-shaped particles are colored gray and plotted back into their positions and orientations in the tomogram. (Right) Rotated and zoomed-in view of the axoneme-facing surface of a mitochondrion. The axoneme is oriented horizontally, so the ladder-like arrays are oriented ∼120° to the flagellar long axis, and individual particles within the array are oriented ∼60° to this axis. (Scale bars, 10 nm.)
 Assigning GK-like proteins as constituents of the ordered OMM arrays at the mitochondria–cytoskeleton interface is also supported by recent genetic studies. Sperm from mice lacking sperm-specific GK isoforms have disorganized mitochondrial sheaths (28, 29). In these mice, spherical mitochondria properly align along the flagellum but fail to properly elongate and coil around it, leaving gaps along the midpiece where the ODFs are exposed (29). This phenotype is consistent with our data showing contacts between GK protein arrays and the underlying cytoskeleton (Fig. 2B and C and SI Appendix, Fig. S4A–L) and by experiments showing that the submitochondrial reticulum remains attached to the OMM in detergent-extracted sperm (18). Although sperm-specific GK isoforms do not show GK activity in vitro (64), we cannot completely exclude that they function in mitochondrial metabolism in situ, especially since knockout mice have higher ATP levels than wild-type (28). If the arrays that we observe indeed consist of GK-like proteins and if these proteins are in fact metabolically active, then their preferential orientation toward the flagellar cytoskeleton could enhance metabolite shuttling between mitochondria and the axoneme.

**Discussion**

In this study, we used cryo-FIB milling–enabled cryo-ET to image the sperm mitochondrial sheath in three mammalian species. Our data reveal that overall, mitochondrial dimensions are remarkably consistent in sperm from the same species (Fig. 1 and SI Appendix, Fig. S1). This contrasts with other mitochondria-rich tissues such as muscle, in which there are massive variations in mitochondrial size and morphology within individual cells (65). In addition, we did not observe mitochondrial nanotunnels in any of the species we examined, in contrast to their relative abundance in muscle tissue (3, 65). However,
we do observe discrete linker proteins that tether neighboring mitochondria in all three species. It will be important to investigate the precise molecular identity of these linkers and to define whether they serve purely structural roles or also act as conducting channels that coordinate metabolite shuttling, energy production, and calcium signaling across the entire midpiece. Hints at possible interorganellar coordination, we observed transmembrane cristae alignment at specialized junctions between neighboring mitochondria in mouse sperm (Fig. 1). Thus, the concept of an interconnected mitochondrial reticulum that occurs in muscle (2) may also be applicable to the mammalian sperm midpiece and warrants further consideration.

In this study, we imaged either ejaculated or mature cauda epididymal sperm. However, mammalian sperm must undergo a plethora of biochemical and morphological changes in the female reproductive tract before they become fertilization competent. These processes are collectively known as capacitation and cause extensive metabolic changes in sperm (66). To cope with increased energy demands, rates of both glycolysis and oxidative phosphorylation increase during capacitation in mouse sperm (67). Future work could use cryo-ET to establish whether these metabolic changes relate to any structural changes in midpiece mitochondria, for instance at the level of cristae architecture or intermitochondrial junctions.

Our data also show that mitochondrial dimensions and cristae architecture vary across species (Fig. 1), providing possible structural bases for interspecific differences in mitochondrial energetics. Horse sperm mitochondria appear to take on a more condensed appearance than their counterparts in pig and mouse, which may correlate with increased reliance on oxidative phosphorylation in horse sperm. Although studies on horse sperm physiology are limited (68), condensed mitochondrial profiles are suggested to be a sign of calcium overload in other species (69, 70), so cross-species studies exploring the link between mitochondrial morphology and possible differences in mitochondrial calcium homeostasis are clearly necessary. In general, the natural diversity of sperm enables comparative studies of how mitochondrial structure varies with sperm metabolism, which will undoubtedly contribute to our broader understanding of how mitochondrial form relates to function.

Our data also demonstrate that the diversity of OMM underpinnings of mitochondrial sheath architecture are conserved, at least in mammals. Specifically, we identified intermitochondrial linkers that tether adjacent mitochondria (Fig. 1 and SI Appendix, Fig. S2). We also describe ordered protein arrays on the axoneme-facing surface of sperm mitochondria, which were first identified by freeze-fracture EM. Here, we characterize the three-dimensional architecture of these arrays in a near-native state and at molecular resolution, revealing how they are anchored on the OMM and how they interact with the flagellar cytoskeleton. In-cell subtomogram averaging reveals that these arrays consist of boat-shaped particles anchored on a lattice of OMM pores (Figs. 2 and 3). Proteomics and in-cell XL-MS suggest that these arrays consist of GK-like proteins anchored on VDAC lattices in the OMM (Fig. 4). Given that VDACs are ubiquitous OMM proteins, our findings motivate efforts to explore whether they also regulate mitochondria–cytoskeleton interactions in other cell types.

The OMM arrays may function to regulate the precise elongation and coiling of mitochondria, contributing to the striking consistency within the mitochondrial sheath. In mature sperm, these arrays may help maintain the integrity of mitochondria–cytoskeleton contacts, stabilizing them against shear stresses during sperm motility and hyperactivation. However, it is unclear what determines the organization of these arrays in the first place. Our averages do not hint at direct interactions between boat-shaped particles. Instead, their spacing may be defined by the organization of the underlying VDAC lattice. Another intriguing possibility is that the arrays are organized by their cytoskeletal binding partners; the periodicity of relevant motifs on the sub mitochondrial reticulum could dictate the spacing of the OMM arrays.

We find that mitochondria–cytoskeleton contact sites in the sperm midpiece can be quite variable (SI Appendix, Fig. S4). Occasionally, OMM arrays make direct contact with the ODFs (SI Appendix, Fig. S4 A–C, E, F, and I); sometimes, there are considerable gaps between them that are bridged either by the sub mitochondrial reticulum (SI Appendix, Fig. S4 D, G, H, and J) or by unidentified linkers (SI Appendix, Fig. S4 K and L). This may be explained by variability in both the sizes and shapes of the ODFs themselves (SI Appendix, Fig. S4 M–R). Defining the precise mechanisms by which the OMM arrays tether to the underlying cytoskeleton is an important avenue for future research, in part because it has important implications for understanding sperm motility. In mammalian sperm, forces from axoneme bending are transmitted to the connecting piece through the ODFs, which are physically coupled to the microtubule doublets along most of the principal piece but not in the midpiece (71, 72). The observation that mitochondria are anchored to the ODFs through extensive ordered arrays adds another level of complexity to the layered sliding mechanics of the mammalian sperm flagellum, affecting, for example, the extent to which the ODFs can slide relative to each other or to the axoneme. Indeed, the organization of these specific GK isoforms appears to be caused by motility defects arising from the fragmented mitochondrial sheath (28, 29). Intermitochondrial linkers that physically tether neighboring mitochondria may also affect the extent to which the midpiece can bend during movement. On a larger scale, interspecies variations in mitochondrial dimensions and packing coupled with known differences in overall midpiece length could result in subtle changes in sperm motility. Furthermore, the stiffness of the midpiece changes during sperm maturation, becoming less rigid as sperm transit through the epididymis (73, 74). Tracing how the molecular structures that we describe here form and change during spermiogenesis and epididymal maturation are likely to provide further insight in this respect. Ultimately, combining structural information from cryo-ET with motility analysis and mathematical modeling could help illuminate how midpiece morphometry affects swimming behavior.

The methods we use here can also be used to study other specialized organelle configurations in the highly-streamlined sperm cell. For example, it is unclear how exactly the sperm head anchors to the tail or how the acrosomal vesicle is laminated to the nucleus. Beyond sperm, such methods could be used to gain molecular insight into the organization of organelles in other terminally differentiated and polarized cell types like neurons, photoreceptors, or hair cells. For example, exploring whether similar arrays are present at the mitochondria–cytoskeleton interface in other differentiated cell types—and whether they use a similar pool of protein components—is an area ripe for study. In striated muscle, proper mitochondrial positioning is critical for muscle function and depends on direct associations between mitochondria and intermediate filaments (9, 75). Similarly, in skin cells, mitochondrial organization depends on keratin (76). The structural bases for these associations are unknown, but cryo-ET and in-cell XL-MS may prove useful in these contexts as well.

Materials and Methods

Sperm Collection and Preparation. Pig sperm samples were obtained from Varkens KI Nederland and prepared for imaging within a few hours of delivery. Sperm were layered onto a discontinuous gradient consisting of 4 ml of 35% Percoll (GE Healthcare) underlaid with 2 ml of 70% Percoll both in Heps-buffered saline (HBS: 20 mM Heps, 137 mM NaCl, 10 mM glucose, 2.5 mM KCl, 0.1% kanamycin, and pH 7.6) and centrifuged at 750 × g for 15 min.
at room temperature (RT). Pelleted cells were washed once in phosphate-buffered saline (PBS: 137 mM NaCl, 3 mM KCl, 8 mM Na2HPO4, 1.5 mM KH2PO4, and pH 7.4) and resuspended in PBS for further processing. All solutions were iso-osmotic at RT (290 to 300 mOsm/kg).

Horse semen was collected from mature Warmblood stallions using a Hanover artificial vagina in the presence of a teaser mare. Semen was filtered through gauze and kept at RT until further processing. Semen was diluted in INRA96 (IMV Technologies) to a sperm concentration of 30 × 10^6 cells/mL. Sperm were then centrifuged through a Percoll gradient (as described above for the pig) for 10 min at 700 × g followed by 10 min at 750 × g (77). The pellet was resuspended in 1 mL of PBS, centrifuged for 5 min at 750 × g, and finally resuspended in PBS.

Mouse sperm were obtained from the cauda epididymis of adult male C57BL/6 mice as described in ref. 78. Male mice were caged as described in ref. 79, and the cauda epididymides were dissected with the vas deferens attached. Tissues were placed in a 500-µL droplet of modified Biggers, Whitten, and Whittingham media (BBW, 20 mM Hepes, 91.5 mM NaCl, 6.6 mM KCl, 1.7 mM CaCl2, 0.27 mM sodium pyruvate, 44 mM sodium lactate, 5 U/mL penicillin, and 5 µg/mL streptomycin, adjusted to pH 7.4 and an osmolality of 300 mOsm/kg). Sperm were gently pushed out from the vas deferens, after which two incisions were made with a razor blade in the cauda. Spermatozoa were allowed to swim out of the cauda into the BBW over a period of 15 min at 37°C, after which the tissue was removed and sperm were loaded onto a 27.5% Percoll density gradient and washed by centrifugation at 400 × g for 15 min. The sperm pellet was resuspended in BBW and centrifuged at 400 × g for 2 min to remove excess Percoll, then finally resuspended in BBW.

**Subtomogram Averaging of ATP Synthase and Ladder-Like Arrays.** Subtomographic averaging with mixed redundancy compensation was performed using Particle Estimation for Electron Tomography (PET) 1.130 (90, 91) on defocus-contrast tomograms of cryo-FIB-milled lamellae. Resolution was estimated using the Fourier shell correlation at a cutoff of 0.5 (90). Alignments were generally performed first on binned data, after which aligned positions and orientations were transformed to less-binned data using scripts provided by Davin Vasishtan. Details of acquisition parameters and particle numbers are summarized in SI Appendix, Tables S1 and S2. Alignment strategies for mitochondrial complexes were designed to take advantage of their defined orientations relative to the membrane plane. Particles were picked manually, and their initial orientations were defined using stakkinit. Initial references were either a randomly chosen particle (for ladder-like arrays) or an average of all particles after roughly aligning them based on their initial orientations (for ATP synthase). Independent alignments using independent initial references were performed for datasets from different species. Alignments allowed for large rotational search ranges around the particle long axis (defined as the y-axis, perpendicular to the membrane plane), with limited search ranges around the x- and z-axes (the membrane plane).

All initial alignments were performed without symmetry. After visual inspection of the maps, twofold symmetry was applied for ladder-like arrays. Symmetrization involved using the aligned positions from the unsymmetrized raw particle scores and rotating particles around the axis of symmetry to cluster virtual particles. A symmetrized volume was generated by averaging all particles and virtual particles and used as a reference for a final, restricted alignment.

PlotBacks were generated in IMOD by first running createAlignedModel to generate model files reflecting updated particle positions and orientations after alignment. The relevant subtomogram average was then thresholded for visualization and saved as an isosurface model, which was then placed back into the tomograms using clonemodel.

**Measurements and Quantification.** All measurements of mitochondrial width were performed in IMOD on VPP tomograms filtered with a SIRT-like filter. Mitochondrial width was measured in the nonmimicking wedge direction at five points along the length of each mitochondrion. Only mitochondria that were entirely in the field of view were included in the measurements. Tomograms and corresponding measurements were then grouped based on their locations relative to the connecting piece, which were determined based on low-magnification images used for targeting during data acquisition.

Internal mitochondrial ultrastructure was quantified from tomograms from cryo-FIB-milled lamellae. The volume occupied by the matrix ($V_{matrix}$), the volume enclosed by the inner mitochondrial membrane ($V_{IMM}$), was measured relative to the volume occupied by the entire mitochondrion ($V_{total}$, the volume enclosed by the OMM). Mesh volumes were extracted from segmentations using imodMesh. Because neural network-based segmentation often resulted in gaps, mitochondrial membranes were segmented manually in IMOD for quantification. Only slices in which both the IMM and OMM were clearly defined were used for segmentation.

**Cross-Linking, Lysis, Digestion, and Peptide Fractionation.** All proteomics and XL-MS experiments were performed on Percoll-washed pig sperm prepared as described above. For cross-linking, ∼300 × 10^6 cells were used, each from three different animals. Briefly, pelleted sperm cells were resuspended in 540 µL of PBS supplemented with DSSO (Thermo Fisher Scientific) to a final concentration of 1 mM. The reaction mix was incubated for 30 min at 25°C with 700 rpm shaking in a ThermoMixer C (Eppendorf) and subsequently quenched for 20 min by adding Tris HCl (final concentration 50 mM). Cross-linked cells were spun down at 13 × 800 × g for 10 min at 4°C, after which the supernatant was removed. Cells were then lysed according to a protocol modified from ref. 92. Cells were resuspended in 1 mL of lysis buffer (100 mM Tris HCl pH 8.5, 8 M Urea, 1% Triton X-100, 5 mM tris(2-carboxyethyl)phosphine (TCEP), 30 mM chloroacetamide (CAA), 10 U/ml Dnase I, 1 mM MgCl2, and 1% benzoxane (Merck Millipore, Darmstadt, Germany), 1 mM sodium orthovanadate, phosphoSTOP phosphatases inhibitors, and Complete Mini ethylenediamine tetraacetic acid-free protease inhibitors). Cells were sonicated on ice for 2 min using an ultrasonic processor (UP100H, Hielscher) at 80% amplitude. The proteins were then precipitated according to ref. 93 and the dried protein pellet resuspended in digestion buffer (100 mM Tris HCl pH 8.5, 1% sodium deoxycholate (Sigma-Aldrich), 5 mM TCEP, and 30 mM CAA). Trypsin and Lys-C proteases were added to a 1:25 and 1:100 ratio (weight/weight), respectively, and protein digestion performed overnight at 37°C. The final peptide mixtures were desalted with solid-phase extraction C18 columns (Sep-Pak, Waters) and fractionated with an Agilent 1200 HPLC pump system (Agilent) coupled to a
strong cation exchange (SCX) separation column (Luna SCX 5 μm to 100 Å particles, 50 × 2 mm, Phenomenex), resulting in 25 fractions.

Liquid Chromatography with MS. Approximately 1,000 ng of peptides from each biological replicate before SCX fractionation were first injected onto an Agilent 1290 Infinity ultra-high-performance liquid chromatography system (Agilent) on a 50-cm analytical column packed with C18 beads (Dr. Maisch ReproSil C18-3 μm) coupled online to an Orbitrap HX-FT (Thermo Fisher Scientific). For this classical bottom-up approach, we used the following liquid chromatography with MS/MS (LC-MS/MS) parameters: after 5 min of loading with 100% buffer A (H2O with 0.1% formic acid), peptides were eluted at 300 nL/min with a 95-min gradient from 13 to 40% of buffer B (80% acetonitrile and 20% H2O with 0.1% formic acid). For MS acquisition, we used an MS1 Orbitrap scan at 60,000 resolution, automatic gain control (AGC) target of 10 5 ions and maximum inject time of 20 ms for MS1 at 7,100× 1,600 m/z; the 10 most-intense ions were submitted to MS2 Orbitrap scan at AGC target of 2 × 10 4 ions and maximum inject time of 50 ms (isolation window of 1.4 m/z, normalized collision energy at 27%, and dynamic exclusion of 16 s). The SCX fractions were analyzed with same Agilent HPLC and the same nano-column coupled on-line to an Orbitrap Lumos mass spectrometer (Thermo Fisher Scientific). For these runs, we used a gradient from 6 to 39% buffer B over 100 min with MS settings for DSS cross-links: survey MS1 Orbitrap scan at 60,000 resolution from 375 to 1,500, AGC target of 4 × 10 4 ions and maximum inject time of 50 ms and MS2 Orbitrap scan at 30,000 resolution, AGC target of 5 × 10 4 ions, and maximum inject time of 100 ms for detection of DSS signature peaks (difference in mass of 37.972 Da). The four ions with this specific difference in mass were selected for further analysis at AGC target of 2 × 10 4 ions, maximum inject time of 150 ms for sequencing selected signature peaks (representing the individual peptides).

Data Processing. The three raw files obtained with classical bottom-up approach were analyzed with MaxQuant version 1.6.17 with all the automatic settings adding Deamidation (N) as dynamic modification against the S. scrofa reference proteome (Uniprot version of 08/2020 with 49,795 entries). With this search, we were able to calculate intensity-based absolute quantification values and created a smaller FASTA file to use for analysis of cross-linking experiments. Raw files for cross-linked cells were analyzed with Proteome Discover software suite version 2.4.1.15 (Thermo Fisher Scientific) with the incorporated XlinkX node for analysis of cross-linked peptides as described in ref. 94. Data were searched against the smaller FASTA created in house with “MS1, MS3 acquisition strategy.” For the XlinkX search, we selected full tryptic digestion with three maximum missed cleavages, 10 ppm error for MS1, 20 ppm for MS2, and 0.5 Da for MS3 in Ion Trap. For modifications, we used static Carbamidomethyl (C) and dynamic Oxidation (M), Deamidation (N), and Met- loss (protein N-term). The crosslinked peptides were accepted with a minimum score of 40, minimum score difference of 4, and maximum false discovery rate set to 5%; further standard settings were used.

Interactive Analysis, Homology Modeling, and Cross-Link Mapping. The cross-link map for VDAC1 protein was generated in R (95) using the graph package (version 1.2.4.2). Only protein interactions supported with at least two CSMs were included in the final network. Homology models of pig GK and pig VDAC2 were generated in Robetta (96) and fitted into subtomogram average maps by rigid body fitting in ChimeraX. Cross-links were mapped onto the resulting models using ChimeraX.

Data Availability. Subtomogram average maps have been deposited to the EM Data Bank (EMDB) with the following accession numbers: EMDB-12354-EMDB-12357. The model of putative GK-like proteins anchored on a VDAC-4 array has been deposited to the Protein Data Bank (PDB) with the accession number PDB ID 7NIE. MS data have been deposited to the ProteomeXchange Consortium via the Proteomics Identification Database (PRIDE) partner repository with the dataset identifier PXD025562.

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33. Y. Fukuda, U. Laugks, V. Luìci, W. Baumeister, R. Danes, Electron cryomgnotography of vitrified cells with a Volta phase plate. J. Struct. Biol. 190, 143–150 (2015).
34. R. Danes, B. Buijse, M. Khoshouei, J. M. Plitzko, W. Baumeister, Volta potential phase plate for in-focus phase contrast transmission electron microscopy. Proc. Natl. Acad. Sci. U.S.A. 111, 15635–15640 (2014).
35. M. Chen et al., Convolutional neural networks for automated annotation of cellular organelles. Nat. Methods 14, 983–985 (2017).
36. C. De Martin et al., Morphological, histochemical and biochemical studies on germ cell mitochondria of normal rats. Cell Tissue Res. 196, 1–22 (1979).
37. G. A. Perkins, M. H. Ellisman, Mitochondrial configurations in peripheral nerve suggest differential ATP production. J. Struct. Biol. 173, 117–127 (2011).
38. P. H. Hanckenbrock, Ultrastructural bases for metabolically linked mechanical activity in mitochondria II. Electron transport linked ultrastructural transformations in mitochondria. J. Cell Biol. 37, 345–369 (1968).
39. M. P. Davila et al., Mitochondrial ATP is required for the maintenance of membrane integrity in stallion spermatozoa, whereas motility requires both glycosylation and oxidative phosphorylation. Reproduction 152, 683–694 (2016).
40. C. Muki, M. Okuno, Glycylase plays a major role for adenosine triphosphate supplementation in mouse sperm flagellar movement. Biol. Reprod. 71, 540–547 (2004).
41. F. Odet, S. Gabel, R. E. London, E. Goldberg, E. M. Eddy, Glycolysis and mitochondrial respiration in mouse HDNC-null sperm. Biol. Reprod. 88, 95 (2013).
42. M. Picard et al., Trans-mitochondrial coordination of crista at regulated membrane junctions. Nat. Commun. 6, 6209 (2015).
43. X. Huang et al., Kissing and nanotunneling mediate intermitochondrial communication in the heart. Proc. Natl. Acad. Sci. U.S.A. 110, 2846–2851 (2013).
44. M. Duvert, J. P. Mazat, A. L. Baret, Intermitochondrial junctions in the heart of the frog, Rana esculenta. A thin-section and freeze-fracture study. Cell Tissue Res. 241, 129–137 (1985).
45. X. W. Guo, C. A. Mannella, Conformational change in the mitochondrial channel, VDAC, detected by electron cryo-microscopy. Biophys. J. 64, 545–549 (1993).
46. B. W. Hoegenboom, K. Suda, A. Engel, D. Fotiadis, The supramolecular assemble of voltage-dependent anion channels in the native membrane. J. Mol. Biol. 370, 246–250 (2007).
47. R. P. Gonzalves, N. Bushinsky, V. Prima, J. N. Sturgis, S. Scheuring, Supramolecular assembly of VDAC in native mitochondrial outer membranes. J. Mol. Biol. 369, 413–418 (2007).
48. C. A. Mannella, Structure of the outer mitochondrial membrane: Ordered arrays of porelike subunits in outer-membrane fractions from Neurospora crassa mitochondria. J. Cell Biol. 89, 680–682 (1981).
49. W. S. Kwon, Y. J. Park, S. A. Mohamed, M. G. Pang, Voltage-dependent anion channels are a key factor of male fertility. Fertil. Steril. 99, 354–361 (2013).
50. A. Ercelay, A. M. Salicioni, E. Wertheimer, P. E. Visconti, Identification of proteins undergoing tyrosine phosphorylation during mouse sperm capacitation. Int. J. Dev. Biol. 52, 463–472 (2008).
51. C. R. Gonzalves, N. Bushinsky, V. Prima, J. N. Sturgis, S. Scheuring, Supramolecular assembly of VDAC in native mitochondrial outer membranes. J. Mol. Biol. 369, 413–418 (2007).
52. M. F. Tomasello, F. Guarino, S. Reina, A. Messina, V. De Pinto, The voltage-dependent anion channel 1 (VDAC1) topography in the mitochondrial outer membrane and their functional implications. EMBO J. 40, e70410 (2021).
53. K. A. Lesich, T. G. de Pinho, L. Dang, C. B. Lindemann, Ultrastructural evidence that motility changes caused by variations in ATP, Mg2+, and ADP correlate to conformational changes in reactivated bull sperm axonemes. Cryoletten (Hoboken) 71, 649–661 (2014).
54. C. Jeulin, L. M. Lewin, C. Chevrier, D. Schoevaert-Brossaut, Changes in flagellar movement of rat spermatozoa along the length of the epididymis: Manual and computer-aided image analysis. Cell Motil. Cytoskeleton 37, 147–161 (1996).
55. K. Miyata et al., Sperm calcium inhibition prevents mouse fertility with implications for male contraceptive. Science 350, 442–445 (2015).
56. P. Konecny et al., Myosin II integrity depends on desmin network targeting to Z-disks and costameres via distinct plectin isoforms. J. Cell Biol. 181, 667–681 (2008).
57. K. Steen et al., A role for keratins in supporting mitochondrial organization and function in skin keratinocytes. Mol. Biol. Cell 31, 1103–1111 (2020).
58. R. P. A. Harrison, B. Mairat, N. G. A. Miller, Flow cytometric studies of biconcave-mediated Ca2+ influx in boar sperm populations. Mol. Reprod. Dev. 39, 197–208 (1993).
59. H. Hutcheon et al., Analysis of the small non-protein-coding RNA profile of mouse spermatozoa reveals specific enrichment of piRNAs within mature spermatozoa. RNA 14, 1776–1790 (2008).
60. S. Meyers, E. Bulkeley, A. Foutouhi, Sperm mitochondrial regulation in motility and fertility in horses. Reprod. Domest. Anim. 54 (suppl. 3), 22–28 (2019).
61. G. W. Okunade et al., Targeted ablation of plasma membrane Ca2+-ATPase (PMCA 1 and 4) indicates a major housekeeping function for PMCA1 and a critical role in hyperactivated sperm motility and male fertility for PMCA4. J. Biol. Chem. 279, 33742–33750 (2004).
62. K. Jamil, I. G. White, Induction of acrosomal reaction in sperm with ionophore A23187 and calcium. Arch. Androl. 7, 283–292 (1981).
63. M. R. Leung et al., The multi-scale architecture of mammalian sperm flagella and implications for ciliary motility. EMBO J. 40, e70410 (2021).
64. Y. Pan, W. K. Deeker, A. H. M. Huu, W. J. Craigen, Retromotorsion of glyceral kinase-related genes from the X chromosome to autosomes: Functional and evolutionary aspects. Genomics 59, 282–290 (1999).
65. A. E. Vincent et al., Quantitative 3D mapping of the human skeletal muscle mitochondrial network. Cell Rep. 26, 996–1009.9 (2019).
66. M. S. Mounib, M. C. Chang, Effect of in utero incubation on the metabolism of rabbit spermatozoa. Nature 201, 143–144 (1964).
67. M. Balbach et al., Metabolic changes in mouse sperm during capacitation. Biol. Reprod. 70, 1031–801 (2004).
68. S. Meyers, E. Bulkeley, A. Foutouhi, Fertilization of mouse oocytes by spermatozoa after treatment with ionophore A23187 and calcium. Arch. Androl. 7, 283–292 (1981).
69. J. M. Heumann, A. Hoenger, D. N. Mastronarde, Clustering and variance maps for in-cell structures of conserved supramolecular protein arrays at the mitochondrion-cytoskeleton interface in mammalian sperm.