Spatiotemporal regulation of ATP and Ca$^{2+}$ dynamics in vertebrate rod and cone ribbon synapses

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Purpose: In conventional neurons, Ca$^{2+}$ enters presynaptic terminals during an action potential and its increased local concentration triggers transient exocytosis. In contrast, vertebrate photoreceptors are nonspiking neurons that maintain sustained depolarization and neurotransmitter release from ribbon synapses in darkness and produce light-dependent graded hyperpolarizing responses. Rods transmit single photon responses with high fidelity, whereas cones are less sensitive and exhibit faster response kinetics. These differences are likely due to variations in presynaptic Ca$^{2+}$ dynamics. Metabolic coupling and cross-talk between mitochondria, endoplasmic reticulum (ER), plasma membrane Ca$^{2+}$ ATPase (PMCA), and Na$^+$/Ca$^{2+}$ exchanger (NCX) coordinately control presynaptic ATP production and Ca$^{2+}$ dynamics. The goal of our structural and functional studies was to determine the spatiotemporal regulation of ATP and Ca$^{2+}$ dynamics in rod spherules and cone pedicles.

Methods: Central retina tissue from C57BL/6 mice was used. Laser scanning confocal microscopy (LSCM) experiments were conducted on fixed-frozen vertical sections. Primary antibodies were selected for their tissue/cellular specificity and ability to recognize single, multiple or all splice variants of selected isoforms. Electron microscopy (EM) and 3-D electron tomography (ET) studies used our standard procedures on thin- and thick-sectioned retinas, respectively. Calibrated fluo-3-Ca$^{2+}$ imaging experiments of dark- and light-adapted rod and cone terminals in retinal slices were conducted.

Results: Confocal microscopy showed that mitochondria, ER, PMCA, and NCX1 exhibited distinct retinal lamination patterns and differential distribution in photoreceptor synapses. Antibodies for three distinct mitochondrial compartments differentially labeled retinal areas with high metabolic demand: rod and cone inner segments, previously undescribed cone juxtanuclear mitochondria and the two plexiform layers. Rod spherule membranes uniformly and intensely stained for PMCA, whereas the larger cone pedicles preferentially stained for NCX1 at their active zones and PMCA near their mitochondria. EM and ET revealed that mitochondria in rod spherules and cone pedicles differed markedly in their number, location, size, volume, and total cristae surface area, and cristae junction diameter. Rod spherules had one large ovoid mitochondrion located near its active zone, whereas cone pedicles averaged five medium-sized mitochondria clustered far from their active zones. Most spherules had one ribbon synapse, whereas pedicles contained numerous ribbon synapses. Fluo-3 imaging studies revealed that during darkness rod spherules maintained a lower [Ca$^{2+}$] than cone pedicles, whereas during light adaptation pedicles rapidly lowered their [Ca$^{2+}$] below that observed in spherules.

Conclusions: These findings indicate that ATP demand and mitochondrial ATP production are greater in cone pedicles than rod spherules. Rod spherules employ high affinity/low turnover PMCA and their mitochondrion to maintain a relatively low [Ca$^{2+}$] in darkness, which increases their sensitivity and signal-to-noise ratio. In contrast, cone pedicles utilize low affinity/high turnover NCX to rapidly lower their high [Ca$^{2+}$] during light adaptation, which increases their response kinetics. Spatiotemporal fluo-3-Ca$^{2+}$ imaging results support our immunocytochemical results. The clustering of cone pedicle mitochondria likely provides increased protection from Ca$^{2+}$ overload and permeability transition. In summary, these novel studies reveal that several integrated cellular and subcellular components interact to regulate ATP and Ca$^{2+}$ dynamics in rod and cone synaptic terminals. These results should provide a greater understanding of in vivo photoreceptor synaptic terminal exocytosis/endocytosis, Ca$^{2+}$ overload and therapies for retinal degenerations.

During an action potential, Ca$^{2+}$ enters the presynaptic terminal of conventional neurons through voltage-gated channels that cluster near the synaptic vesicle release sites or active zones. The increased subplasmalemmal Ca$^{2+}$ concentration in these microdomains plays a key role in transient exocytosis [1-5]. Metabolic coupling and cross-talk between mitochondria, endoplasmic reticulum (ER), the plasma membrane Ca$^{2+}$ ATPase (PMCA), and Na$^+$/Ca$^{2+}$ exchanger (NCX), and Ca$^{2+}$ channels must coordinately control ATP production and Ca$^{2+}$ dynamics in the presynaptic terminals to ensure recovery for the next action potential [1,2,4-9]. Thus, the spatiotemporal control of the synaptic terminal ATP production and Ca$^{2+}$ concentration are essential for regulating the probability and rate of neurotransmitter release, quantal content, and kinetics.
of vesicle exocytosis and endocytosis [1,2,5]. For example, reciprocal cooperation between mitochondria and ER controls presynaptic Ca\textsuperscript{2+} handling and neurotransmitter release in lobster neuromuscular junction and bullfrog sympathetic ganglia [10,11], but not in the mouse calyx of Held [7], mouse neuromuscular junction [8] or goldfish mixed rod-cone bipolar (Mb1) cell [12]. Moreover, it is reported that mitochondria in the goldfish Mb1 cell terminals, which cluster far from the active zones [13], do not buffer presynaptic Ca\textsuperscript{2+} but instead primarily provide ATP to the PMCA to extrude intraterminal Ca\textsuperscript{2+} [12-14].

In contrast, vertebrate rod and cone photoreceptors are nonspiking neurons that maintain sustained depolarization and neurotransmitter release from ribbon synapses in darkness and produce light-dependent graded hyperpolarizing responses [15,16]. Following membrane depolarization, the kinetics of rapid exocytosis are faster in cones than rods, which is likely due to differences in presynaptic Ca\textsuperscript{2+} dynamics [17,18]. For example, salamander rods have one pool of vesicles that are released rapidly and linearly by submicromolar to micromolar concentrations of Ca\textsuperscript{2+} [17,19]: consistent with the single photon sensitivity and high fidelity of rods [20,21]. During darkness, photoreceptors maintain a sustained depolarization and continuously release glutamate [15,16,22,23]. To date, no comprehensive electrophysiological and pharmacological experiments examining the regulators of ATP production and Ca\textsuperscript{2+} homeostasis, similar to those conducted in several different neurons and goldfish retinal Mb1 cells [7,8,10-14,24], have been performed on rod or cone photoreceptor synaptic terminals. However, it has been reported that PMCA is the predominant Ca\textsuperscript{2+} extrusion mechanism in photoreceptor synaptic terminals [25] and that a ryanodine-dependent amplification mechanism is coupled to the release of glutamate in dark-adapted salamander rods [26,27]. In addition, studies with amphibian and mammalian retinas have demonstrated the presence of ER [28-31] and PMCA [25,32] in photoreceptor synaptic terminals.

Synaptic release of neurotransmitters at conventional synapses is rapid and transient [1,3,6-9]. The ATP-dependence of neurotransmitter uptake into synaptic vesicles, exocytotic vesicle priming and replenishment as well as the Ca\textsuperscript{2+}-dependence of vesicle fusion and neurotransmitter exocytosis at synaptic membranes of neurons is widely accepted [1,3,4,33,34]. In contrast to conventional chemical synapses, the presynaptic terminals of retinal photoreceptors and bipolar cells possess ribbon synapses [35,36]. Ribbon synapses are specialized organelles located in nerve terminals that maintain a high rate of sustained transmitter release during a graded depolarization [35,36]. In addition, the photoreceptor and bipolar cell synaptic terminals differ from one another in their ribbon synaptic unit morphology, Ca\textsuperscript{2+} sensitivity for exocytosis, and kinetics of exocytosis and endocytosis [9,17,36,37]. Despite the kinetic differences of neurotransmitter release between conventional and ribbon synapses as well as among photoreceptor and bipolar cell ribbon synapses, studies suggest that mitochondrially-generated ATP is required for neurotransmitter uptake by synaptic vesicles, vesicle priming and vesicle competence, but not for vesicle fusion or neurotransmitter release [8,33,34,37,38].

Neuronal mitochondria play a fundamental role in regulating ATP production and intracellular Ca\textsuperscript{2+} for processes related to cellular metabolism, ionic homeostasis, neurotransmitter uptake/release and cell survival [1,2,39,40]. Energy demand drives mitochondrial ATP production [41]. ATP production is modulated by the total number of mitochondria per cell, size of the mitochondria, volume of the mitochondria per cell, and total surface area of the inner mitochondrial cristae membranes [42,43]. Neuronal somata are separated from their presynaptic terminals by relatively long axonal processes. This necessitates that mitochondria be differentially distributed within neurons to serve compartment-specific demands such as oxidative metabolism, neurotransmitter release, and Ca\textsuperscript{2+} homeostasis [2,6-8,44]. To develop a comprehensive knowledge and understanding of mammalian photoreceptor synaptic terminal ATP production, Ca\textsuperscript{2+} homeostasis and neurotransmitter kinetics, integrative immunocytochemical, ultrastructural and Ca\textsuperscript{2+} imaging studies on photoreceptor synaptic terminals, mitochondria and their interrelation with the proteins that utilize ATP and regulate Ca\textsuperscript{2+} concentrations are needed.

The retina has one of the highest rates of mitochondrial oxygen consumption and energy demand of any tissue [45]. The rod and cone photoreceptors contain at least 75% of the total retinal mitochondria, have the highest retinal cytochrome oxidase (COX) activity, and consume 2 fold to 3 fold more oxygen than the inner (proximal) retina [46-51]. To date, mitochondria have been localized to the photoreceptor inner segments (IS) and synaptic terminal regions [45-55]. Extralfoveal primate cone photoreceptor IS (CIS) have more mitochondria, higher COX activity, and stain more intensely for the Na\textsuperscript{+}, K\textsuperscript{+}-ATPase (Naka) than rod photoreceptor IS (RIS) [46,47,51]. Murine CIS have two-fold more mitochondria than RIS and have higher COX activity than RIS [55]. In addition, CIS mitochondria have narrower cristae junctions (CJs), greater cristae interconnectivity, and approximately 3 fold greater cristae membrane surface area than RIS mitochondria [55]. Furthermore, cones have significantly more synaptic ribbons in their terminals, possess unique basal junctions on their pedicles compared to rods, and have a higher concentration of ATP than rods [47,56-64]. Collectively, these biochemical, structural and substructural studies indicate that mammalian cones have an overall higher bioenergetic demand and ATP production than rods [55,56].

The overall goal of these studies was to develop a comprehensive structural and functional understanding of rod spherule and cone pedicle ribbon synaptic terminals. The specific aims of these laser scanning confocal microscopy (LSCM), electron microscopy (EM), three-dimensional electron tomography (ET), and Ca\textsuperscript{2+}-fluor imaging studies were four-fold. The first was to determine the cellular distribution and spatial interrelation of mitochondria, ER, PMCA, and NCX in the retina and especially in the photoreceptor synaptic terminals. The second was to determine if the cellular and subcellular ATP and Ca\textsuperscript{2+} regulatory mechanisms differed between rod spherule and cone pedicle synaptic terminals. The third
was to determine if the relative [Ca\textsuperscript{2+}] in rod spherules and cone pedicles differed during darkness and light-adaptation. The fourth goal was to determine and discuss the functional significance of our results in relation to rod and cone synaptic terminal bioenergetics, Ca\textsuperscript{2+} homeostasis and neurotransmitter release. Here we show that mitochondria, ER, PMCA and NCX each exhibited a distinct retinal lamination pattern. At higher spatial resolution, unique functionally related distributions of mitochondria, ER, PMCA and NCX were observed in both rod spherules and cone pedicles. Rod spherule and cone pedicle mitochondria also had marked functionally related differences in the location, total number, size, volume and total surface area of the inner mitochondrial cristae membranes. Furthermore, rod spherules maintained a lower [Ca\textsuperscript{2+}] than cone pedicles during darkness, whereas cone pedicles lowered their intraterminal [Ca\textsuperscript{2+}] faster and to a greater degree than rod spherules during light adaptation. These results provide a more comprehensive understanding of the spatiotemporal control of Ca\textsuperscript{2+} concentrations and mitochondrial ATP production in rod and cone ribbon synaptic terminals as they relate to exocytosis and endocytosis. Moreover, they provide important groundwork for further understanding compartmental differences in photoreceptors and offer new insight into strategies that will be necessary to treat visual deficits that result from rod and/or cone photoreceptor degeneration.

**METHODS**

*Materials:* All chemicals were purchased as analytical or molecular biology grade from Sigma Chemical Co. (St. Louis, MO) or Fisher Scientific (Pittsburgh, PA) unless otherwise noted. The pH of all solutions was 7.40 at indicated temperatures.

*Experimental animals:* All experimental and animal care procedures complied with the principles of the American Physiological Society, the NIH Guide for the Care and Use of Laboratory Animals and Maintenance (NIH publication No. 85-123, 1985) and were approved by the Institutional Animal Care and Use Committee of the University of Houston. Wild-type C57BL/6J mice (Harlan Sprague Dawley, Indianapolis, IN), from litters bred at our facility, were maintained on a 12:12 light:dark cycle (10-20 lux cage luminance) with food and water available ad libitum.

For most studies, 21 and 60 day old female mice were decapitated between one and two hours after light-onset and their eyes were rapidly removed and immersed in ice-cold phosphate buffered saline (PBS). For a few studies, 60 day old female mice were dark-adapted overnight and decapitated two hours after scheduled light-onset under dim red light (λ >650 nm). The corneas were gently punctured at the limbus. Then the eyes were either immersion-fixed in room temperature 4% paraformaldehyde in 0.1 M cacodylate buffer for 30 min for LSCM studies or in ice-cold 3% glutaraldehyde, 2% paraformaldehyde and 0.1% CaCl\textsubscript{2} in 0.1 M cacodylate buffer (Karnovsky’s fixative) for 12 h at 4 °C for conventional EM or ET studies as described [40,55,56,64]. Three to seven retinas from different mice were used for each independent analysis. There were no age-dependent differences on any analysis.

*Antibodies and lectins:* The primary antibodies and lectin used in these studies were selected carefully for their tissue and cellular specificity, ability to recognize single or multiple protein isoforms, ability to recognize all splice variants of selected isoforms, and commercial availability (Table 1). We conducted extensive preliminary experiments to ensure that every antibody utilized in these studies had the appropriate specificity and penetration. All antibodies utilized in this study were titrated through a broad range of dilutions (most over 3 orders of magnitude) to determine optimal working dilutions. In addition, the concentration of Triton X-100 in the blocking agent was titrated to ensure optimal penetration without a significant loss in epitope. Immunolabeling specificity was confirmed by processing retinal sections as described below in the absence of the primary antibodies, by substituting normal rabbit or goat serum for polyclonal antibodies, or by using immunizing peptides for neutralization experiments. These procedures eliminated all specific labeling and revealed no false-positive labeling.

An extensive panel of well-characterized primary antibodies directed against cell- and organelle-specific markers in the retina was used in double and triple labeling experiments. These were antibodies for rhodopsin, M-opsin, S-opsins, M-cone arrestin, vesicular glutamate transporter 1 (VGluT1), protein kinase C α (PKCα), synaptotagmin 1, and peanut agglutinin (PNA) [58,65-72]. The details about the other antibodies, previously not used for retinal immunocytochemistry studies, are immediately below. COX is the terminal electron transport complex of the mitochondrial respiratory chain and standard activity-dependent inner boundary membrane (IBM) and cristae marker [41,47]. The anti-COX subunit IV (COX IV) mouse monoclonal 20E8 antibody (Molecular Probes, Eugene, OR) is a molecular marker of the inner membrane system [73] and detects a single 16 kDa band on Western blots [40,105]. The anti-voltage-dependent anion channel (VDAC: mitochondrial porin) rabbit polyclonal antibody Ab-5 (Calbiochem, San Diego, CA) is an established marker of the outer mitochondrial membrane (OMM), was raised against amino acids 185-197 of the human VDAC, recognizes all three VDAC isoforms and detects a single 31 kDa band on Western blots [74]. We did not use the anti-VDAC mouse monoclonal antibody 31HL, used in retinal studies by Gincel et al. [104], since it only recognizes the VDAC1 isoform [75]. The anti-mitochondrial DNA polymerase-γ (POLG) rabbit polyclonal antibody Ab-1 (Lab Vision, Fremont, CA) is a nuclear-encoded protein responsible for mitochondrial DNA (mtDNA) repair and replication that is located in the mitochondrial matrix, was raised against amino acids 714-1061 of the human POLG, and detects a single 140 kDa band on Western blots [76]. POLG expression and message level are maintained regardless of the mtDNA status [77]. The anti-calreticulin rabbit polyclonal antibody AB3825 (Chemicon, Temecula, CA) is specific for the Ca\textsuperscript{2+}-binding chaperone located in the lumen of all ER that actively modulates Ca\textsuperscript{2+} transport across the ER membrane [78]. It was raised against amino acids 412-417 of the C-terminus of calreticulin, does not cross react with other ER proteins and detects a single
60 kDa band on Western blots [79]. The anti-pan-sarcoplasmic-endoplasmic reticulum Ca\(^{2+}\) ATPase isoform 3 (SERCA3) rabbit polyclonal antibody PA-1-910A (Affinity BioReagents, Golden, CO) was raised against amino acids 29-39 of the mouse and rat SERCA3 isoform, recognizes all splice variants of human and rodent SERCA3, does not cross react with other SERCA isoforms, and detects a single 97 kDa band on Western blots [80]. The anti-pan-PMCA mouse monoclonal antibody MA3-914 (Affinity BioReagents) was raised against amino acids 724-783 of the human erythrocyte Ca\(^{2+}\) pump, recognizes all four isoforms of PMCA and detects a 140 kDa band on Western blots [81]. The anti-NCX1 rabbit polyclonal antibody p11-13 (Swant, Switzerland) was raised against the full length canine cardiac NCX1, recognizes all splice variants of NCX1, does not cross react with other NCX isoforms, and detects the 120 and 160 kDa bands on Western blots [81,82].

**Laser scanning confocal microscopy studies:** Fixed eyes were rinsed in ice-cold PBS for 10 min. Fixed and washed eyes were cryoprotected in 30% sucrose/PBS solutions. The anterior segments were removed, eyecups were embedded in Tissue-Tek® OCT mounting media (Electron Microscopy Sciences, West Grove, PA) or Alexa Fluor 488 (Molecular Probes) immersion in liquid nitrogen. Retinas were sectioned along the vertical meridian on a cryostat at a thickness of 10-15 µm, collected onto Superfrost®/Plus microscope slides (Fisher Scientific) and stored at -20 °C until used.

For all LSCM experiments, sections were fixed and immunolabeled in parallel to insure identical processing. All analyzed sections were obtained 200-400 µm from the optic nerve head. Immunofluorescent labeling of frozen sections was essentially as described [79]. Briefly, sections were thawed for 60 min before use and postfixed by immersion in 4% paraformaldehyde for 15 min to improve tissue adherence to the slides. Sections were rinsed in nanopure water (npH\(_2\)O), treated with 1% sodium borohydride to reduce nonspecific tissue autofluorescence, and immediately rinsed in npH\(_2\)O. Sections were rinsed in PBS and treated for two hours at RT with 5% bovine serum albumin, 1% fish gelatin, 10% normal goat serum and 0.1-0.3% Triton X-100 in ice-cold PBS to block non-specific immunolabeling. Primary antibodies were applied for two days at 4 °C. For double and triple labeling experiments, primary antibodies from different host animals were applied simultaneously.

After incubation in primary antibody, the sections were rinsed three times in PBS and blocked for 30 min. Dilutions (1:500) of Cy3- or Cy5- (Jackson ImmunoResearch Laboratories, West Grove, PA) or Alexa Fluor 488 (Molecular Probes) antibodies were applied simultaneously. Immunolabeled sections were rinsed three times in PBS before mounting on Superfrost®/Plus slides with Vectorshield containing DAPI (Vector Laboratories, Burlingame, CA) and stored at 4 °C.

**TABLE 1. Cell-specific primary antibodies and lectin**

| Primary antigen or lectin | Structure labeled | Host       | Source             | Dilution | References |
|--------------------------|------------------|------------|--------------------|----------|------------|
| Calreticulin             | ER               | Rabbit     | Chemicon           | 1:100    | 78,79      |
| Middle wavelength-sensitive cone arrestin (M-CAr) | Cones | Rabbit | Chemicon | 1:1000 | 109 |
| Cytochrome oxidase subunit IV (COX IV) | Mitochondrial inner membrane system | Mouse | Molecular Probes | 1:500 | 40,73,94 |
| Kinesin KIF3A | Photoreceptor ribbon and synaptic vesicles | Mouse | BD Biosciences | 1:100 | 71 |
| Na\(^+\)/Ca\(^{2+}\) exchanger isoform 1 (NCX1) | Synaptic terminals | Rabbit | Swant | 1:100 | 81,82 |
| Middle wavelength-sensitive opsin (M-opsin) | Cones | Rabbit | Kind gift from Cheryl Craft | 1:1000 | 65 |
| Short wavelength-sensitive opsin (S-opsin) | Cones | Rabbit | Kind gift from Cheryl Craft | 1:1000 | 65 |
| pan-Plasma membrane Ca\(^{2+}\) ATPase (PMCA) | Synaptic terminals | Mouse | Affinity BioReagents | 1:100 | 81 |
| Mitochondrial DNA polymerase-γ (POLG) | Mitochondrial matrix | Rabbit | Lab Vision | 1:500 | 76 |
| Protein kinase C-α (PKCα) | Rod bipolar cells | Rabbit | Sigma | 1:1000 | 68,69 |
| Rhodopsin (ID4) | Rods | Mouse | Chemicon | 1:1000 | 72 |
| Sarcoplasmic-endoplasmic reticulum Ca\(^{2+}\) ATPase isoform 3 (SERCA3) | ER | Rabbit | Affinity BioReagents | 1:400 | 80 |
| Synaptotagmin 1 | Photoreceptor and bipolar synaptic vesicles | Mouse | Chemicon | 1:100 | 70,113 |
| pan-Voltage-dependent anion channel (VDAC) | Mitochondrial outer membrane | Rabbit | Calbiochem | 1:1000 | 74 |
| Vesicular glutamate transporter 1 (VGLUT1) | Photoreceptor and bipolar cell terminals | Guinea pig | Chemicon | 1:1000 | 69 |
| Peanut agglutinin (PNA)-Alexa Fluor 647 Conjugate | Cone outer segments and terminals | Molecular Probes | 1:50 | 58,69 |

The primary antigen or lectin, structure(s) they label, host, source, and reference are presented.
-conjugated secondary antibodies were applied and incubated for 60 min in the dark at RT. For double and triple labeling experiments, secondary antibodies directed against primary antibodies from different species were applied simultaneously. After incubation with secondary antibody, the sections were rinsed in PBS and npH2O. The immunolabeled slides were dried and cover-slipped with Vectashield® anti-fade mounting medium (Vector Laboratories, Burlingame, CA) and stored at 4 °C until visualized. For double and triple labeling experiments using PNA, PNA-Alexa Fluor 647 (1:50 dilution; Molecular Probes) was applied simultaneously with the secondary antibody/antibodies.

LSCM images were acquired using a Leica TCS SP2 LSCM (Leica Microsystems, Exton, PA). Stacks of images from different Z-planes were obtained using a step size of 0.3-0.5 µm. “Bleedthrough” of fluorescent signals from different channels was eliminated by adjusting laser power, detector sensitivity and by sequentially imaging each fluorescent channel. Confocal images were identically and minimally processed by importing them into Adobe Photoshop CS software (Adobe Systems, Inc., Mountain View, CA). The results shown are representative of three to six separate immunolabeling experiments from three to five different mouse retinas. In all double and triple labeling experiments, the voxel dimensions in the X-Y dimensions were smaller than in the Z-dimension. Epitopes were designated as “colabeled” when the fluorescent pixels overlapped in the images. For all figures, the designation colabeled implies that the epitopes were within 290-400 nm of each other.

Semi-quantitative assessment of immunolabeling intensity: The lamination-specific intensity of COX IV, VDAC, POLG, calreticulin, PMCA, and NCX immunolabeling was assessed by three independent viewers. Each viewer examined a minimum of five confocal immunofluorescent sections per retina from three to five mice and ranked the immunolabeling intensity on a relative five-point scale. The fluorescent labeling scale was intense (+++), strong (++), moderate (++), weak (+) or absent (0). The combined results had a 90-95% concordance between viewers and are presented in Table 2.

Conventional electron microscopy: The ultrastructure of mouse and rat photoreceptors has been described in several classic papers [28,53,83,84]. The fixation procedures used in these studies preserved the ultrastructure of the outer retina, although they were not optimal for maintaining the photoreceptor mitochondria ultrastructure and substructure as these were not the major goals of these studies. In contrast, one of our primary goals was to analyze and compare the ultrastructural and substructural features and characteristics of rod and cone photoreceptor mitochondria. Therefore, we used our well-validated fixation and embedding procedures for these endeavors, essentially as described [40,55,65,85]. Briefly, each eye was fixed overnight and a piece of the superior temporal retina 200-250 µm from the optic nerve was obtained. We chose this retinal area for two reasons. First, we used the same area of mouse retina for our previous ultrastructural and ET work on mouse cone inner segment mitochondria [55]. This allowed us to compare directly our results from rod and cone inner segment mitochondria to the current study on rod and cone synaptic terminal mitochondria. Second, this region contains mostly middle wavelength-sensitive (M) cones [169] and M cones in the mouse are similar to those in other mammals [86,87], which enables cross-species comparisons. Sections were dehydrated and embedded in Spurr’s or Araldite resin as described [40,55,65,88]. Ultra-thin vertical sections of the retina were stained with uranyl acetate and lead citrate before

Table 2. Retinal Lamination and Corresponding Staining Intensity of Mitochondria, Calcium Transporters and ER

| Retinal Area or Structure | Mitochondrial antibody | Calcium transporters |
|--------------------------|------------------------|----------------------|
|                          | COX IV | VDAC | POLG | PMCA | NCX | Calreticulin |
| ROS and COS              | 0      | 0    | 0    | 0    | 0   | 0          |
| RIS and CIS              | +++    | ++/+++ | ++++ | na   | na  | na         |
| Cone juxtanuclear mitochondria | ++++/++++ | +++/+++ | +++ | na   | na  | na         |
| Rod juxtanuclear mitochondria | +/+++ | ++ | ++ | na | na | na         |
| Overall ONL              | +/++   | ++/+ | +++ | ++ | ++ | ++         |
| OPL                      | +++    | +++  | +++ | +++ | ++ | ++/+++     |
| Distal INL somas         | ++ | +  | ++  | ++ | ++ | ++/+++     |
| Middle INL somas         | +/++ | +  | ++ | ++ | ++ | ++         |
| Proximal INL somas       | +/+  | +   | ++  | +  | ++ | ++/+++     |
| IPL sublamina-α          | ++++/+++ | ++/+ | +++ | ++ | +++ | 0/+       |
| IPL sublamina-β          | ++++/+++ | ++/+ | +++ | ++ | ++ | 0/+        |
| RGC                      | +++   | ++   | ++  | ++ | ++ | ++         |
| Müller glial end-feet     | +++   | 0/+  | ++  | 0/+ | 0/+ | 0/+        |

COX IV represents cytochrome oxidase subunit IV; VDAC represents voltage-dependent anion channel; POLG represents mitochondrial DNA polymerase-γ; PMCA represents plasma membrane Ca²⁺ ATPase; NCX represents Na⁺/Ca²⁺ exchanger isoform 1; ER represents endoplasmic reticulum ROS and COS represents rod and cone outer segments; RIS and CIS represents rod and cone inner segments; ONL and INL represents outer and inner nuclear layer; OPL and IPL represents outer and inner plexiform layer; GCL represents ganglion cell layer intensity staining key; ++++ represents intense; +++ represents strong; ++ represents moderate; + represents weak; 0 represents absent; na represents not applicable.
being examined in a JEOL 100-C or 1200EX transmission EM (Tokyo, Japan). The number of mitochondria per rod spherule and cone pedicle was calculated from three to five different grids from each of five different mice. The mean number from each mouse was determined and the overall mean±SEM was calculated.

**Three-dimensional electron microscope tomography:** Mouse retinas were prepared for ET essentially as described [40,55]. Briefly, the superior temporal retina 200-250 µm from the optic nerve was trimmed (vide supra) and the retinal sections were dehydrated, embedded in Durcupan resin, sectioned (500 nm thick) and imaged using the single- and double-tilt series techniques described by Perkins and co-workers [55,89,90]. Fiducial cues, consisting of 20 nm colloidal gold particles, were deposited on both sides of the section. For each reconstruction, a series of images at regular tilt increments was collected with a JEOL 4000EX intermediate-voltage EM operated at 400 kV. To limit anisotropic specimen thinning during image collection, the specimens were irradiated before each tilt series. Tilt series were recorded at 20,000X magnification with an angular increment of 2° from -60° to +60° about an axis perpendicular to the optical axis of the microscope. A computer-controlled goniometer accurately incremented the angular steps. A slow-scan CCD camera with pixel dimensions of 1960x2560 was used to collect images. The pixel resolution was 1.1 nm. Illumination was held to near parallel beam conditions and constant optical density was maintained constant by varying the exposure time. The IMOD package [91] was used for rough alignment with the fine alignment and reconstruction performed using the TxBR package [92].

Volume segmentation was performed by manual tracing in the planes of highest resolution with the program Xvoxtrace [90]. The mitochondrial reconstructions were visualized using Analyze (Mayo Foundation, Rochester, MN) or the surface-rendering graphics of Synu (National Center for Microscopy and Imaging Research, San Diego, CA) as previously described [55,93]. These programs allow one to step through a small volume glass bottom dishes coated with Matrigel (Collaborative Research, Palo Alto, CA) and incubated for 15 min at 27°C in HEPES buffer containing 1.5 mM CaCl₂ in order to restore the normal extracellular Ca²⁺ concentration and increase the esterase activity [96]. The HEPES buffer with CaCl₂ was aspirated and replaced with fresh buffer prior to the onset of Ca²⁺ imaging.

To confirm that the observed fluorescence signals reflected changes in internal Ca²⁺ levels, we conducted three different experiments. First, we added 1 mM Pb²⁺, a potent fluo-3 fluorescence quencher [Kd for fluo-3 is 6 pM: 94,97], to the Ca²⁺-containing HEPES buffer. Similar to our previous results [40,94], this significantly quenched the Ca²⁺-enhanced fluo-3 fluorescence in dark-adapted photoreceptor synaptic terminals (data not shown). Second, we incubated retinal slices in Ca²⁺-free HEPES buffer with 5 mM BAPTA-AM [1,2-bis(o-aminophenoxy)ethane-N,N',N'-tetraacetic acid] for 15 min at 27°C. The fluo-3 fluorescence measured in dark-adapted photoreceptor synaptic terminals, from these retinal slices, was not above background (data not shown). Third, we attempted an in vivo calibration of fluo-3 fluorescence in photoreceptor synaptic terminals using the standard ionomycin, Mn²⁺ and digitonin procedure for NIH 3T3 cells [98,99] and isolated cerebellar granule cells [97] as well as the Ca²⁺ ionophore A-23187. These calibration procedures saturated the fluo-3 fluorescent signal above the physiological range. However, they were unreliable because shortly after the fluo-3 fluorescent signal saturated the rod photoreceptors initiated apoptosis: as we reported [40,94]. Therefore, we established an in vitro relative fluorescent intensity (RFI) standard curve, as described [98,100], in order to estimate the free [Ca²⁺] in the photoreceptor synaptic terminals. An intracellular buffer (25 mM HEPES, 130 mM KCl, 5 mM NaCl, 3 mM MgCl₂, pH 7.2, 305±3 mOsm) that contained calibrated Ca²⁺ buffers (0-40 μM Ca²⁺; Molecular Probes) and 5 μM of the penta-ammonium salt of fluo-3 (Molecular Probes) was pipetted onto dual concave glass slides maintained in the dark at 27°C. Fluo-3 fluorescence was measured with the Zeiss LSM-410 confocal
Ca²⁺-fluor-3 in dark- and light-adapted rod spherules and cone pedicles were 200-250 nm and the Z resolution was 300 nm. The RFI of the image was a Z-stack maximum projection (single confocal section). The results show that the Ca²⁺ and fluo-3 fluorescence calibration curve.

Normalized intensity values were calculated, means±SEMs were determined, and the data was statistically analyzed. The mean RFI of dark-adapted rod spherules and cone pedicles was determined using NIH Image, version 1.62. To directly compare the RFI, and thus the relative free [Ca²⁺], in dark- and light-adapted rod and cone synaptic terminals, only experiments where relatively adjacent photoreceptor synaptic terminals were imaged are included in this data set. To normalize all the images a standard background subtraction was performed on each image. Then the overall RFI within each dark- and light-adapted rod spherule and cone pedicle was determined based on a grey scale with 256 levels. The light-adapted decrease in photoreceptor oxygen consumption stabilizes during the first minute of the light stimulation [49,50,101].

RESULTS

Topographically distinct mitochondrial antibodies reveal differential lamination and cellular distribution of retinal mitochondria: LSCM and EM studies: Photoreceptors contain about 75% of the retinal mitochondria, have the highest COX activity, and have significantly greater oxygen consumption than inner retina [46-51]. To test the hypothesis that the distribution of retinal mitochondria reflects cell-specific and synaptic-specific differences in bioenergetics, we examined the immunofluorescent staining pattern of three independent topographical markers of mitochondrial membranes and compartments: COX IV, VDAC, and POLG (Figure 1; Table 1). Figure 1 illustrates the three topographically distinct mitochondrial regions that were recognized by the selected antibodies: the OMM, inner membrane system that includes the IBM and cristal membranes, and mitochondrial matrix. Below, we describe the typical labeling patterns, from outer or distal retina to inner or proximal retina, for each of these three markers. No specific labeling of the rod or cone outer segments (ROS; COS) was observed with any mitochondrial antibody, consistent with the absence of mitochondria in this photoreceptor compartment [53,65]. Intense COX IV immunolabeling was present in both RIS and CIS mitochondria (Figure 2A and Figure 3) consistent with our previous results and the concentration of mitochondria in the IS [55]. At first glance, the outer nuclear layer (ONL) contained relatively few mitochondria. However, a more detailed analysis of the ONL revealed two distinct COX IV-positive mitochondrial populations in the ONL. The first population was located in the distal ONL where strong to intense COX IV-positive arc-shaped puncta were detected (Figure 2A: white arrowheads). Double labeling experiments, with a mixture of middle and short wavelength-sensitive cone opsins antibodies (Table 1), revealed that these previously undescribed pair of juxtanuclear mitochondria were located above and below the most distal cone nuclei (Figure 3: pairs of white arrowheads). Electron microscopy confirmed this result (Figure 4: white and black arrowheads). Double labeling experiments with COX IV and VDAC revealed that these two mitochondrial markers colocalized in the juxtanuclear mitochondria (Figure 1).
The second mitochondrial population in the ONL was a band of smaller, moderately labeled puncta located in the proximal ONL (Figure 2A,D and Figure 3: white arrows). These puncta are juxtanuclear mitochondria localized next to individual rod nuclei present in the proximal ONL, as confirmed by electron microscopy (Figure 4: white and black arrows). Moreover, COX IV and VDAC colocalized throughout the OPL (Figure 2C).

The IS and outer plexiform layer (OPL) intensely labeled for COX IV (Figure 2A and Figure 3), confirming that mammalian photoreceptor IS and synaptic terminals contain numerous highly active mitochondria [47,55,62]. A mixture of M- and S-cone opsin antibodies [65] labeled all the cones in the mouse retina [66]. In addition, the high magnification confocal image of COX IV and cone opsin double labeling revealed a laminar COX IV staining pattern in the OPL (Figure 3). That is, a single row of large opsin-positive cone pedicles was present in the proximal OPL that contained numerous COX IV-positive puncta (Figure 3: double white arrows). In contrast, the distal OPL had several rows of opsin-negative terminals that contained large round COX IV-positive spheres (Figure 3). These observations suggest that rod spherules contain a single large mitochondrion, whereas cone pedicles contain numerous, albeit smaller, mitochondria. To directly investigate this we conducted EM studies as described below. In summary, the overall staining pattern and intensity of COX IV immunofluorescence in photoreceptors indicates that most of
their aerobically-generated ATP is produced by IS and synaptic terminal mitochondria (Table 2).

The proximal (inner) retina also exhibited a differential distribution of COX IV immunolabeling. The inner nuclear layer (INL) somas had weak to moderate COX IV labeling with the strongest staining located in the distal INL (Figure 2A). The location of these distal somas and their Chx10-positive staining (data not shown; manuscript in preparation) indicates that they are bipolar cells [67,68,103]. A subpopulation of these Chx10-positive bipolar cell somas were COX IV- and PKCα-positive (Figure 2A: inset), confirming that they are rod bipolar cells [67,68]. Less intense staining was

Figure 2. Molecular markers for three separate mitochondrial compartments reveal distinct retinal distribution and lamination patterns. The abbreviations of the retinal layers are used for this and all subsequent figures. IS represents inner segments, ONL represents outer nuclear layer, OPL represents outer plexiform layer, INL represents inner nuclear layer (d: distal, m: middle, p: proximal), IPL-α represents inner plexiform layer sublamina-α (OFF lamina), IPL-β: inner plexiform layer sublamina-β (ON lamina), GCL: ganglion cell layer, and Mef: Müller glial end-feet. A: Confocal image of retina immunolabeled for COX IV. The pairs of white arrowheads identify numerous intensely labeled arc-shaped puncta located in the distal ONL, which are juxtanuclear mitochondria in the cone somas. Between the white arrows, in the proximal ONL, is a band of small circular puncta that are juxtanuclear mitochondria near rod nuclei. The inset shows colocalization of COX IV (green) and PKCα (red) in rod bipolar cells (white arrows): scale bar equal 20 µm. B: Confocal image of retina immunostained for VDAC. Note the intense labeling in the OPL. C: Confocal image of retina double labeled with antibodies against COX IV (red) and VDAC (green). The pseudocoloring in panels A and B was reversed in this panel to clearly show the colocalization (yellow-orange pixels) in the IS, cone juxtanuclear mitochondria (white arrowheads) and rod juxtanuclear mitochondria (white arrows). Punctate colocalization also is present throughout the OPL, IPL-α, and IPL-β. D: Confocal image of retina immunolabeled for POLG. POLG also labels the cone juxtanuclear mitochondria (white arrowheads) and rod juxtanuclear mitochondria (white arrows). The inset is a higher magnification (2X) view. Scale bar equal 40 µm for all panels.
detectable in the middle and proximal INL somas: the size and location of these somas suggested they were Müller glial somas and amacrine cells, respectively [68]. COX IV strongly to intensely labeled the inner plexiform layer (IPL), ganglion cell layer (GCL) and distal Müller glial end-feet. A clear demarcation between the OFF (sublamina-α: IPL-α) and ON (sublamina-β: IPL-β) IPL sublamina was observed (Figure 2A).

Figure 2B,C show that moderate to strong VDAC labeling was present throughout the IS and that VDAC and COX IV colocalized in this compartment. Thin VDAC-positive processes are evident throughout the ONL. These VDAC-positive processes surrounded the tightly packed rhodopsin-positive somas in the ONL (Figure 5), suggesting that they are Müller glial cell processes. EM studies confirmed this (Figure 4). Cone and rod juxtanuclear mitochondria strongly colabeled for VDAC and COX IV (Figure 2C). The most intense VDAC labeling occurred throughout the OPL and this colocalized with COX IV (Figure 2C). In contrast to the ONL, the INL weakly stained for VDAC. Moderate to strong VDAC labeling was evident throughout the IPL. However, stronger punctate labeling occurred in the most proximal band of IPL-β, where the rod bipolar cells terminate [67,68]. Double labeling experiments showed that VDAC and COX IV colocalized in these rod bipolar cell terminals (Figure 2C). Triple labeling experiments with VDAC, PKCα, and VGluT1 confirmed that these were rod bipolar cell terminals (data not shown; manuscript in preparation). The GCL and proximal Müller glial cell end-feet exhibited weak VDAC labeling.

Figure 2D shows that POLG intensely labeled the IS, OPL, distal and proximal INL somas, proximal IPL-β, and GCL. Similar to the COX IV staining pattern in the ONL, there were strong POLG-positive arc-shaped puncta in the distal ONL (white arrowheads) and circular puncta in the proximal ONL (white arrows). Except for the presumed rod bipolar cell terminals in IPL-β, the remainder of the IPL exhibited relatively moderate staining. The COX IV, VDAC, and POLG labeling patterns were similar in sections obtained from the superior and inferior central retina.

*Endoplasmic reticulum are differentially distributed throughout the retina:* Several LSCM, EM, and ET studies have shown that ER and mitochondria are in close apposition and that microdomains of high Ca2+ are shared between these organelles [2,37,89,93,104,105]. To directly test the hypothesis that ER are closely apposed to retinal mitochondria, and especially those in the photoreceptor synaptic terminals, we examined the staining pattern of two independent ER markers in combination with COX IV or VGluT1 (Figure 6; Table 1 and Table 2) and conducted EM studies (vide infra). Figure 6A shows that calreticulin, a molecular marker of all ER [75], intensely labeled the entire IS region and that the OPL, INL, and GCL were moderately to strongly labeled. In contrast, the ONL and IPL weakly labeled for calreticulin. Calreticulin and COX IV colocalized in multiple retinal areas (Figure 6B: punc-
tate yellowish pixels), indicating that these two proteins were in close apposition. This interrelation was especially prominent throughout the photoreceptor ellipsoid region. Moreover, calreticulin was in close apposition to the cone juxtanuclear mitochondria (white arrowheads) and mitochondria in the OPL. Labeling in the latter revealed that calreticulin was located close to COX IV-positive synaptic terminal mitochondria (see OPL in Figure 3). Calreticulin and COX IV also colocalized in the INL as evidenced by the diffuse orange-colored pixels. Figure 6C,D further illustrate that calreticulin is expressed intensely in CIS and cone pedicles (Figure 6D: white arrows and purple pixels).

To determine if another ER antibody colocalized in photoreceptor synaptic terminals, single and double label experiments were conducted with the SERCA3 and VGluT1 antibodies. We did not utilize the SERCA N89 polyclonal rabbit antibody as it labels COS and photoreceptor IS, but only sparsely labels photoreceptor synaptic terminals in adult mouse, rat and monkey [31]. Figure 6E shows that the SERCA3 moderately to strongly labeled ER in the RIS and CIS (white arrows), closely apposed to the cone juxtanuclear mitochondria (white arrowheads), and in the OPL. Thus, the SERCA3 labeling pattern in the outer (distal) retina was similar to that of calreticulin. Double labeling experiments with the pan-SERCA3 and VGluT1 revealed that they completely and strongly colocalized in both rod spherules and cone pedicles (Figure 6F: purple pixels).

**PMCA and NCX1 differentially and selectively label the rod spherules and cone pedicles, respectively:** PMCA and NCX share a complementary role in regulating the presynaptic Ca2+ concentration. PMCA has a high affinity and low turnover rate, whereas NCX has a low affinity and high turnover rate for Ca2+ extrusion [75,76]. Few studies have examined the location or functional roles of the two major presynaptic Ca2+ transporters in neurons [6,25,106-108]. The kinetics of exocytosis is 10 fold faster in cones than in rods [17,18], suggesting that PMCA and NCX may be differentially distrib-
uted or located in rod and cone synaptic terminals. To directly test this hypothesis, the staining pattern of PMCA and NCX1 in combination with markers of photoreceptor cell terminals [69] or cones [57,58] was examined (Figure 7; Table 1 and Table 2). Previous work showed that mouse retinal neurons exhibit cell-specific expression of the four different PMCA isoforms [32], however, this study did not examine differential PMCA expression in rod and cone synaptic terminals or present the IS results. A preliminary study with a 1:1000 dilution of the same anti-NCX1 rabbit polyclonal antibody we used at 1:100 dilution suggested that NCX was weakly expressed in rat cone, but not rod, photoreceptors and was more abundant in the inner retina [25]. However, no co-labeling experiments were performed to confirm this observation and

Figure 5. High magnification confocal images of outer retina double labeled for VDAC (red) and rhodopsin (green). A: To reveal the voltage-dependent anion channel (VDAC) labeling of the IS (Figure 2B) the prominent rhodopsin labeling in the OSs and most IS was removed using Photoshop. Minimal colocalization of VDAC and rhodopsin occur in the ONL, highlighting the lack of mitochondria in most rod somas. However, the Müller glial cell processes that surround the rod somas are VDAC positive (data not shown). B: Stratification of rod spherule and cone pedicle mitochondria in the OPL. Double labeling for VDAC and rhodopsin shows the distal-proximal stratification of rod spherules (white arrows) and cone pedicles and their associated mitochondria. Tiers of rod spherules, each with one large mitochondrion, overly the rhodopsin-positive region. The more proximal cone pedicles contain clusters of mitochondria (white arrowheads) and are located in the rhodopsin-negative region. The inset is a higher magnification (2X) view of the OPL. Scale bar equal 20 µm for both panels.
there are no studies on NCX expression in mouse retina.

Overall, Figure 7A shows that pan-PMCA intensely labeled the synaptic terminals (plexiform layers) of the retina: as described with different PMCA antibodies [82,83]. In the outer retina, PMCA weakly labeled the IS and ONL and intensely labeled the OPL (Table 2). Double label experiments with VGluT1 and the selective cone antibody M-CAr [109] established the location of the large dome-shaped cone pedicles in the OPL (Figure 7B: purple pixels indicate colocalization). Figure 7C revealed that the intensely PMCA-positive rod spherules have a horseshoe-like or “V”-like [25] appearance. In contrast, the cone pedicles stained weakly and diffusely for PMCA (Figure 7C,H), except for a band around the distal part of the pedicle plasma membrane (Figure 7C: white arrowheads) and where the cone axon descends into the pedicle (Figure 7B: white arrowheads). Double labeling with PMCA and VGluT1 confirmed that PMCA extensively and uniformly labeled almost the entire rod spherule plasma membrane, whereas it only selectively labeled a region of the cone pedicle (Figure 7D: white arrowheads). Since PMCA2 labeled rod bipolar terminals in the IPL [32], we examined whether the dendrites in the OPL labeled with pan-PMCA. Pan-PMCA did not label the somas or dendrites of PKCα-positive rod bipolar cells, although PMCA and PKCα were in close apposition.

**Figure 6. Molecular markers of endoplasmic reticulum (ER) reveal distinct retinal distribution and lamination patterns.** A-D: Confocal image of retina triple labeled for calreticulin, COX IV and PNA. Scale bar equal 20 μm. A: Retina stained for calreticulin, a Ca2+-binding protein present in all ER lumen. Note the intense labeling in the IS, OPL and INL. B: Location of calreticulin (red) and COX IV (green). These proteins are in close apposition in the ellipsoid region of photoreceptor IS (white arrows and yellow pixels), juxtanuclear mitochondria associated with cone somas (white arrowheads) and in the OPL (punctate yellow-orange pixels). C: Retina stained with peanut agglutinin (PNA), a relatively selective marker for cones, shows distinct labeling in the IS and OPL. D: Calreticulin (red) and PNA (blue) colocalize in CIS and near the pedicles (white arrows; bright purple pixels). E: Pan-Smooth ER Ca2+ ATPase isoform 3 (SERCA3) immunolabeling. This antibody labels in close apposition to the RIS and CIS (white arrows), juxtanuclear mitochondria associated with cone somas (white arrowheads) and the mitochondria in the OPL. F: Colocalization of pan-SERCA3 (red) and VGluT1 (blue) throughout the OPL, indicating the presence of ER in the photoreceptor synaptic terminals. Scale bar for panels E and F equal 40 μm.
Figure 7. Lamination patterns and differential compartmentation of pan-plasma membrane Ca\(^{2+}\) ATPase and Na\(^{+}-\text{Ca}\(^{2+}\) exchanger isoform 1 in rod spherules and cone pedicles.  

A: Confocal image of retina immunolabeled for pan-plasma membrane Ca\(^{2+}\) ATPase (pan-PMCA). The double white arrowheads identify strongly labeled PMCA bands in each IPL sublamina. The scale bar applies to A, D and represents 40 \(\mu m\).  

B-D: PMCA preferentially labels rod spherules. OPL double labeled with markers for PMCA (green), vesicular glutamate transporter 1 (VGlut1: blue) and/or M-cone arrestin (M-CAr: red). Scale bar equal 20 \(\mu m\). B: VGlut1, which labels photoreceptor terminals and M-CAr, which labels cones, colocalize in the OPL. This reveals the large dome-shaped cone pedicles (purple pixels) and some cone axons (white arrowheads). C: This high magnification image shows the horseshoe-like appearance of the PMCA-positive rod spherules. In contrast, the retina double labeled with PMCA and M-CAr shows that cone pedicles stain weakly and diffusely for PMCA, except for a discrete band at the top of the pedicle (white arrowheads: yellow pixels). D: A retina double labeled with PMCA and VGlut1 confirms that PMCA extensively labels rod spherules, but only sparsely labels cone pedicles (white arrowheads).  

E: PKC\(\alpha\)-positive rod bipolar cells (red) are pan-PMCA-negative (green), although they are in close apposition around the rod spherules (yellow pixels). The scale bar represents 20 \(\mu m\).  

F: Retinal localization of Na\(^{+}-\text{Ca}\(^{2+}\) exchanger isoform 1 (NCX1). NCX1 intensely labels IS, cone axons and cone pedicles (white arrowheads). G-I: NCX1 preferentially labels cone pedicles. OPL from triple labeled experiments with markers for NCX1 (red), PMCA (green), and/or PNA (blue). Scale bar equal 20 \(\mu m\). G: PNA stains selected flat contact regions on the large cone terminal (white arrows). NCX1 intensely labeled the entire cone pedicle and axon terminals, whereas rod spherules were diffusely labeled. H: PMCA labels the rod spherule membranes with a horseshoe-like appearance, but does not prominently colocalize with PNA-positive cone pedicle membranes (white arrows). I: Retina double labeled with PMCA and NCX1 shows that these proteins colocalized in cone pedicles near the axon junction (white arrows: yellow pixels).  

J, K: High magnification confocal images showing preferential labeling of cone pedicles by NCX1 and rod spherules by PMCA. OPL from triple labeled experiments with markers for NCX1 (red), PMCA (green), and/or PNA (blue). Scale bar equals to 10 \(\mu m\). J: Double labeled retina shows that NCX1 intensely labels the entire cone pedicle and axon terminals, but only diffusely labels the rod spherules. This high magnification image also shows that PNA stains selected flat contact regions on the large cone terminal, whereas invaginating synaptic regions (white arrows) are not labeled by PNA. K: Double labeled retina shows that PMCA and NCX1 colocalize in cone pedicles near the axon junction (white arrows: yellow pixels).
around the synaptic terminal region of the rod spherules (Figure 7E: yellow pixels).

In the inner retina, PMCA moderately labeled the INL, GCL, and Müller end feet (Figure 7A). Recently, it was shown that constant illumination increases Ca\(^{2+}\) in the proximal Müller glial cell processes [110]. Our results suggest that PMCA likely extrudes this Ca\(^{2+}\) from the Müller end feet. In addition, PMCA strongly labeled the entire IPL and more intensely labeled a band in each IPL sublamina (Figure 7A). These bands were identified previously as PMCA1-positive [32,67] and PMCA2-positive [32]. The location of the INL somas and dendrites of these cells, are similar to the PMCA1- and Ca\(^{2+}\)-binding protein 5 (CaB5)-labeled cells identified as ON- and OFF-cone bipolar terminals [67]. In addition, the PMCA2-labeled cells [32] have a lamination pattern in the IPL similar to cholinergic amacrine cells [69,111].

Low and high magnification confocal images of immunostained for NCX1, PMCA and/or peanut agglutinin (PNA) are presented in Figure 7F-K. NCX1 did not stain OS, showing that this NCX1 antibody did not cross-react with the ROS Na\(^+\)/Ca\(^{2+}\)-K\(^{+}\) (NCKX1) or COS (NCKX2) exchangers [112], NCX1 intensely labeled IS, cone axons descending through the ONL (Figure 7F-K), the OPL (Figure 7G,J) and complete cone pedicles (Figure 7F-K,G,J: white arrows; Table 2). Weak to moderate labeling outlined somas in the INL and GCL. Diffuse labeling occurred throughout the IPL, although the IPL-\(\alpha\) labeled more strongly than IPL-\(\beta\). High magnification confocal images revealed that NCX1 diffusely labeled rod spherules (Figure 7G and Figure 7J). In cones, PMCA and NCX1 only colocalized in small bands at the top of the cone pedicle (Figure 7I and Figure 7K: yellow pixels); similar to that seen with M-CAr in Figure 7C.

In summary, rod spherule membranes intensely stained for PMCA, but only weakly and diffusely stained for NCX1. In contrast, cone pedicle membranes intensely stained for NCX1, but only the apical portion of the pedicle stained for PMCA. Additionally, NCX1, but not PMCA, labeled the RIS and CIS. These results confirm our hypothesis that the presynaptic Ca\(^{2+}\) transporters differentially distribute and localize in rod and cone photoreceptor terminals and strongly suggest that these differences have important functional implications relating to Ca\(^{2+}\) dynamics and neurotransmitter release.

Based on their kinetic properties (i.e., Kd for Ca\(^{2+}\) and kcat) and the dependence of PMCA on ATP as a substrate, a second hypothesis was that NCX1 would be closely apposed to the active zones, whereas mitochondria would be closely apposed to PMCA in photoreceptor synaptic terminals. Figure 8A, a high magnification confocal image of a NCX1 and VGluT1 double label experiment, clearly shows that NCX1 intensely labeled cone pedicles (white arrows; purple pixels), but only diffusely labeled rod spherules. To determine whether NCX1 localized to the synaptic ribbon region (active zone), two different markers for rod and cone synaptic vesicles—synaptotagmin 1 [70,113] and kinesin KIF3A [71] were used in colocalization experiments. Figure 8B shows that cone pedicles had large clusters of double labeled NCX1- and synaptotagmin 1-positive puncta (white arrowheads: yellow-orange pixels), whereas only small NCX1- and synaptotagmin 1-positive puncta were present in rod spherules (white arrows: yellow pixels). The kinesin KIF3A-labeled rod spherules had an arc-shaped appearance (Figure 8C), consistent with the strong kinesin KIF3A labeling of the photoreceptor ribbon matrix [71]. When retinas were double labeled with NCX1 and kinesin the cone pedicles had large visible clusters of double labeled NCX1- and kinesin-positive puncta (white arrows: yellow-orange pixels), whereas rod spherules contained only small yellow puncta. Double labeling with PMCA and synaptotagmin1 (data not shown) or syntaxin 3 [32] revealed that PMCA did not colocalize with either marker, indicating that PMCA was not present at the active zone of rod spherules.

A set of triple labeling confocal experiments determined the spatial interrelation of synaptic terminal mitochondria to NCX1-labeled (Figure 8D-F) and PMCA-labeled (Figure 8G,H) synaptic membranes. Triple labeling with NCX1, PNA and COX IV revealed that cone pedicles contained multiple mitochondria that were not located at the ribbon synaptic unit (Figure 8E: white ellipsoids). In high spatial resolution images, it is evident that the mitochondria in the same six cone pedicles are located relatively far from the active zone (Figure 8F: white arrowheads). Triple labeling experiments with PMCA, VDAC and VGluT1 (Figure 8G) or PMCA, VDAC and PNA (Figure 8H) suggest that rods contain a single large mitochondrion (white arrowheads) located close to the horse-shoe-shaped PMCA-labeled synaptic membranes. In contrast, the cone pedicles contain multiple mitochondria (white arrows). However, the mitochondria in cone pedicles cluster close to the PMCA-labeled membranes located near the apical portion of the pedicles. As described below in more detail, electron microscopic studies confirmed these observations (Figure 9).

In summary, these results confirmed our second hypothesis that NCX1 localizes to active zones, whereas mitochondria are closely apposed to PMCA in photoreceptor synaptic terminals. These results expand our understanding of how the two major presynaptic Ca\(^{2+}\) transporters can differentially regulate rod spherule and cone pedicle Ca\(^{2+}\) levels during sustained depolarization in darkness and thereby participate in the kinetic regulation of neurotransmitter (glutamate) release. Our high-resolution immunocytochemical results on the presynaptic localization of NCX1 and PMCA in photoreceptors are not entirely consistent with the epifluorescent microscopy study of the rat retina, which states that NCX1 staining was weak in rat cones and absent in rods [25]. However, this difference is understandable since 10 fold less NCX antibody was used in the rat study, which resulted in no NCX labeling in the IS and very weak NCX labeling in the OPL [25].

**Cones pedicles have more ATP production capacity than rod spherules:** conventional electron microscopy and tomography of mouse outer plexiform layer and photoreceptor synaptic terminal mitochondria: A number of electrophysiological, biochemical and morphological observations indicate that cone synaptic terminals, compared to rods, have a higher ATP demand [18,47,114-120]. To test the functionally-based hypothesis that the ATP production capacity of cone pedicles is
Figure 8. NCX1 localizes to ribbon synaptic units of cone pedicles and mitochondria closely associate with PMCA in photoreceptor terminals. 

A: NCX1-positive cone pedicles (red) and VGluT1 (blue) colocalize in the proximal ONL (white arrowheads: purple pixels). B: Synaptotagmin 1 (green) labels photoreceptor synaptic vesicles. Small NCX1- and synaptotagmin 1-positive puncta colocalize in rod spherules (white arrows; yellow-orange pixels). Scale bar equal 20 µm. 

C: Kinesin KIF3A (green) labels photoreceptor ribbons and docked synaptic vesicles. The kinesin-labeled rod spherules have an arc-shaped appearance and colocalize with diffusely located NCX1 (small yellow puncta). In contrast, cone pedicles have large clusters of double labeled NCX1- and kinesin-positive puncta (white arrows; yellow-orange pixels). Scale bar equal 20 µm. 

D-F: Mitochondria cluster away from the active zone in cone pedicles. Scale for D and E equal 20 µm and for F equal 10 µm. D: NCX1 (red) and PNA (blue) colocalize in cone pedicles (purple pixels). E: Triple labeling with NCX1, PNA and COX IV (green) reveals that the cone pedicles (white ellipsoids) contain multiple mitochondria that are located away from the ribbon synaptic unit. F: Higher magnification image of the same six pedicles in E reveals the COX IV and NCX1 colabeling (white arrowheads) and the distance of the COX IV-positive mitochondria from the active zone. G, H: Mitochondria closely associate with PMCA in the rod spherules and cone pedicles. Triple labeling with PMCA (green), VDAC (red), and with either VGluT1 (blue) or PNA (blue). Rods contain a single large mitochondrion (arrowheads) located close to the PMCA-labeled membranes. Cone pedicles contain multiple mitochondria (white arrows) clustered close to PMCA-labeled membranes, which are located away from the active zones. Scale bar equal 20 µm.
greater than that of rod spherules, we used EM and ET to examine the number, size, volume and total inner membrane system surface area of mitochondria [42,43] in the synaptic terminals of rods and cones. The high magnification electron micrographs of rod and cone synaptic terminals, presented in Figure 9A-D, reveal numerous important morphological differences between rod spherules and cone pedicles and their associated mitochondria. First, Figure 9A illustrates a fundamental organizational feature of the OPL. That is, three to five tiers of rod spherules are located above the larger more electron lucent cone pedicles (also see Figure 4). Second, rod spherules contained a single very large ovoid mitochondrion, as described [28]. It is especially worth noting that the rod mitochondrion is located close to the synaptic ribbon complex (Figure 9A,C and Figure 10), consistent with our LCSM studies. Second, cone pedicles have four to six closely grouped mitochondria (per pedicle=SEM=5.2±0.2) that are located in the distal portion of the pedicle relatively far from the synaptic ribbon complexes (Figure 9A and Figure 10B): as shown in the confocal studies (Figure 8). This is reminiscent of the goldfish retinal Mb1 cell where multiple mitochondria cluster away from the active zone [13]. Third, the rod spherule mitochondrion is larger (1.8-2.0 μm diameter) than any of the cone pedicle mitochondria (0.8-1.5 μm diameter). The rod mitochondrion occupies 25-30% of the spherule volume, whereas the cone pedicle mitochondria collectively occupy 10-15% of the pedicle volume (Figure 9A-C and Figure 10). Fourth, the average volume of the rod spherule is 3.0-3.5 μm³, whereas the average volume of the cone pedicle is about 10 times larger. Fifth, rod spherules usually contain one ribbon synapse (Figure 9A-C). It was estimated that about 5% contain two ribbons [120,121], as illustrated in the rod spherule labeled r* in Figure 9A. Occasionally, we noticed that the most proximal rod nuclei had a ribbon synapse at the base of its soma, as described [94], and the spherule lacked a mitochondrion (data not shown). Sixth, cone pedicles contain 6 to 14 synaptic ribbons (Figure 9B-D; [52,53]; data not shown). The invaginating processes of two lateral horizontal cells (h) and a central bipolar cell (b), the classic triad [122], were seen at the short synaptic ribbon of the cone and the longer synaptic ribbon of the rod (Figure 9A). The arciform density was noticeable at the base of each of several ribbons.

In addition to their differences, rod and cone pedicles and their mitochondria share some similarities. First, the mitochondria in rod and cone synaptic terminals were in the orthodox conformation (Figure 10). This configuration is characterized by a relatively large matrix volume and small intracrystal space, as the mitochondrial IBM is closely apposed to the OMM [123]; Figure 1. Second, mitochondria in both the spherules and pedicles are relatively large compared to the IS mitochondria [55] and those in other neural tissues [89,93,124], although the rod spherule mitochondria are larger and rounder. Third, cisternae of endoplasmic reticulum (ER) were observed near the mitochondria and synaptic ribbon of both the rod spherule and cone pedicle (Figure 9A,C). In some instances, the ER was likely contiguous with or closely apposed to individual mitochondrion, as observed in rat spherules [28]. ER also were observed in frog rod and cone synaptic terminals [29,30], where ATP markedly enhanced the Ca²⁺ uptake by rod spherule ER [29].

Analysis of the three-dimensional structural features of rod spherule and cone pedicle mitochondria: To determine the total inner membrane system surface area of mitochondria, a quantitative analysis and comparison of the ultrastructure and substructure of rod spherule and cone pedicle mitochondria using the high-resolution 3-dimensional tools of ET was required (Figure 11 and Figure 12; Table 3). Since there is a paucity of detailed structural information about rod spherule and cone pedicle mitochondria, another goal was to compare and contrast the structural features of photoreceptor synaptic terminal mitochondria with our published ultrastructural and substructural results of mouse RIS and CIS mitochondria [40,55,56]. Volume segmentation provided an analysis of individual cristae. Their 3-dimensional shapes and membrane architecture were visualized in varying orientations to classify the structural motifs of lamellae, tubes, and constrictions (narrow connections between crista segments) per each crista. Surface rendering and measurements of volume and surface elements permitted a comparison of substructures inside mitochondria (Figure 12; Table 3).

Three-dimensional ETs revealed that rod spherule and cone pedicle mitochondria contain a very large number of cristae (Figure 11 and Figure 12). The means±SEM number of cristae per mitochondrial volume for rod spherules and cone pedicles was 1,337±192 and 1,057±209 μm⁻³, respectively. The cristae, presented after segmentation and surface rendering, are represented by different colors (Figure 11D,H). Most of the cristae in rod and cone synaptic terminal mitochondria were tubular, however, a few had lamellar compartments. Although the rod spherule mitochondria had 26% more cristae per unit volume than the cone pedicle mitochondria, this was not statistically different (Figure 12A). However, there were significantly more cristae membranes in a rod spherule, compared to a cone pedicle, mitochondrion. That is, the cristae surface/mitochondrial surface (i.e., amount of cristae membranes) and cristae volume/mitochondrial volume were 2.2-2.5 fold larger in the rod spherule mitochondrion, compared to a cone pedicle mitochondrion (Figure 12B,C). Overall, and importantly, the four to six cone pedicle mitochondria collectively contain twice as many cristae membranes as the single rod spherule mitochondrion. In contrast, the number of cristae segments per volume (Figure 12A), number of segments per crista (Figure 12B) and fraction of cristae with multiple segments (Figure 12C) were similar in rod spherule and cone pedicle mitochondria. A detailed examination of Figure 12 reveals that there are distinct and marked differences in the rod and cone cristae of mitochondria located in the synaptic terminal compared to those in the inner segment [55].

There are two classes of contact sites: classical and bridge. The classical contact site is defined by the OMM and IBM pinching together (Figure 1), whereas the bridge contact site spans across the intermembrane space without a change in distance between the outer and inner membranes [55,93,125]. Tomographic reconstructions showed that both types of con-
Figure 9. Ultrastructure of rod and cone photoreceptor synaptic terminals. A: The rod spherules (r) form two to four tiers that overly the larger more electron lucent cone pedicles (cone). Two rod nuclei (RN) are present. The rod spherules contain a single large mitochondrion (m), whereas the cone pedicles contain several mitochondria. Cisterna of smooth endoplasmic reticulum (white arrowheads) are often located adjacent to individual mitochondrion. Both terminals are filled with round synaptic vesicles. The invaginating processes of two lateral horizontal cells (h) and a central bipolar cell (b), the classic triad [122], are seen at the short synaptic ribbon (black arrow) of the cone and the longer synaptic ribbon (black arrow) of the rod in the upper right corner of the figure. The rod spherule on the left labeled r* appears to be a rod with two synaptic ribbon units as described [121]. B: Rod spherule and axon; cone pedicle. On the left portion of the panel, a descending rod axon (black and white arrowheads) and its expansion into a rod spherule (rod) is visible for two rods. An elongated mitochondrion (m; black arrowheads) is visible in the thin axon (0.20-0.25 µm) as it enlarges into the spherule. The rod spherules are filled with 25-35 nm electron dense round synaptic vesicles and a single large mitochondrion. A tangential view of an adjacent rod spherule with a synaptic ribbon (black arrow) is shown. A larger more electron lucent cone pedicle has two visible synaptic ribbons (black arrows) at invaginating synapses in this section. A rod nucleus (RN) in the most proximal row of the ONL is present. C: Rod spherule and cone pedicle. The rod spherules contain a single large mitochondrion (m) and an invaginating process with two lateral horizontal cells (h) and a central bipolar cell that is just out of the section. The rod has a long synaptic ribbon (black arrow) and a visible arciform density at its base. The most proximally located cone pedicle has two classic triads visible with two lateral horizontal cells (h) and a central bipolar cell (b). Other horizontal cell processes also are visible and the arciform density is evident. Cisterna of smooth endoplasmic reticulum (white arrowheads) are more numerous in cone pedicles than rod spherules. D: Two cone pedicles and a rod spherule. Each large cone pedicle has multiple synaptic ribbons (black arrows) at invaginating synapses. Several lateral horizontal cell processes, of varying size and intensity, are visible. The enlarged axonal region of rod spherule, with its darker matrix, contains a mitochondrion (m) that is just out of the section. Scale bars for all panels equal 1 µm.
tact sites were present in rod spherule and cone pedicle mitochondria. A crista junction (CJ) is the site where a crista connects to the IBM [89,126]. CJs are tubular openings that are invariably narrow in orthodox mitochondria, remarkably uniform in diameter, and connect the intracristal space with the intermembrane space (Figure 11B-D and Figure 10F-H). The de novo formation of tubular CJs is energetically favorable as these are dynamic structures [127]. The average CJ diameter in brain mitochondria is 14-16 nm, whereas it is 12 nm in CIS mitochondria and 17 nm in RIS mitochondria [55,93] (Table 3). The usual narrow opening and uniform size of the CJ likely restricts the diffusion of cytochrome c and ADP/ATP out of the intracristal compartments, adjacent to the membranes where the respiratory complexes are concentrated [124,126,128]. Unexpectedly, the mean CJ in cone pedicle mitochondria was 9 nm (Figure 11F). This is significantly smaller than the 12 nm diameter measured in rod spherule mitochondria (Figure 11B) and smaller than any other known CJ (Table 3).

In summary, these novel ET results revealed that rod and cone synaptic terminal mitochondria share some unique substructural features that are different from mitochondria in the rod and cone inner segments (Figure 12) and other neurons (Table 3). In addition, the rod and cone synaptic terminal mitochondria possess significantly different and uniquely distinguishing characteristics (Figure 12; Table 3).

Ca²⁺ imaging and correlative electron microscopy of rod and cone synaptic terminals in dark-adapted and light-adapted whole retinas: The above results, in concert with the demonstrated sensitivity and kinetics of rod and cone synaptic transmission [9,17-21], predict that the free [Ca²⁺] will be lower in dark-adapted mouse rod spherules compared to cone pedicles. In addition, they suggest that mouse cone pedicles would exhibit faster and larger spatiotemporal decreases in free [Ca²⁺] during light adaptation than rod spherules. To test these two functionally-based hypotheses, fluo-3 Ca²⁺ imaging and L SCM experiments were conducted in retinal slices during darkness (Figure 13A,C) and one min after the onset of a rod saturating stimulus (Figure 13B,D). This time period was used because the light-adapted decrease in photoreceptor oxygen consumption stabilizes during the first minute of light stimulation [49,50,101]. Figure 13 presents representative pseudocolored images of free [Ca²⁺] from two adjacent rod spherules (Figure 13A,B) and a nearby cone pedicle (Figure 13C,D). For higher resolution...
Figure 11. Three-dimensional electron tomograms of a rod spherule and a cone pedicle mitochondrion. 

A-D: Three-dimensional imaging of a rod spherule mitochondrion. A: A representative 2.2 nm slice through a tomographic volume of a rod spherule (from Figure 9A) that shows a large mitochondrion with many cristae. The inset (enlarged 3X) illustrates an example of a classical contact site, defined as the location where the OMM and IBM join (arrow). Scale bar for A and B equal 200 nm. B: Another slice through the volume showing two crista junctions (CJs: boxed) that have tubular openings, which connect the cristae with the intermembrane space. The inset at the bottom shows the openings (white arrowheads) of the two CJs enlarged 2X. C: Side view of the inner membrane of the segmented volume displayed with left lighting. CJ openings are invariably narrow, tubular and remarkably uniform in diameter. There are fifteen numbered CJ openings in this view. D: Top view of the segmented volume showing the outer membrane (blue) and a subset of cristae (various colors). Most of the cristae are tubular. However, some cristae possess lamellar compartments, which connect to the intermembrane space via CJs.

E-H: Three-dimensional imaging of a cone pedicle mitochondrion. E: A representative 2.2 nm slice through a tomographic volume of a cone pedicle (from Figure 9B) that shows a typical medium- to large-sized mitochondrion typical at this terminal. As illustrated, the abundance of cristae membranes is significantly smaller in each cone pedicle, compared to rod spherule mitochondria. The inset (enlarged 3X) illustrates an example of a classical contact site (arrow). Scale bar for E and F equal 200 nm. F: Another slice through the volume showing two CJs (boxed). The inset at the bottom shows the openings (white arrowhead) of the two CJs expanded 2X. The openings of these crista junctions are significantly smaller than their counterparts in the rod spherule mitochondria (Table 3). G: Side view of the inner membrane of the segmented volume displayed with left lighting. The eight CJ openings seen in this view are numbered. H: Top view of the segmented volume showing the outer membrane (blue) and a subset of cristae (various colors). Most of the cristae are tubular, as in the rod spherule mitochondrion, although a small proportion have lamellar compartments. The cristae surface area and volume are significantly smaller in cone pedicle, compared to rod spherule, mitochondria (B, C).
resolution and better visualization of the Ca^{2+} microdomains, the pseudocolored images were transformed using an advanced image synthesis algorithm (Figure 13G-K). The corresponding electron micrographs of these rod spherules and cone pedicle are shown in relatively the same orientation (Figure 13E,F). Considering the various preparation procedures and length of time that the retinas were incubated in different buffers, the mitochondria and synaptic terminals are intact and relatively well preserved compared to our immediately fixed tissue (Figure 9). In addition, the Ca^{2+} and fluo-3 calibration curve, used to estimate free [Ca^{2+}] in the photoreceptor synaptic terminals, is presented in Figure 13G.

Table 4 presents the Ca^{2+}-fluo-3 RFI values and the estimated mean free [Ca^{2+}] for the dark- and light-adapted rod spherules and cone pedicles. The estimated free [Ca^{2+}] in dark-adapted rod spherules was 3.2 fold lower (about 2.2 µM) than in cone pedicles (about 6.8 µM), which is consistent with our hypothesis that dark-adapted rod spherules maintain a lower mean resting [Ca^{2+}] than cone pedicles. In contrast, the estimated free [Ca^{2+}] in light-adapted cone pedicles was 3.2 fold lower (about 0.2 µM) than in light-adapted rod spherules (about 0.7 µM). Thus, during light adaptation the estimated free [Ca^{2+}] decreased 2 fold in rod spherules and almost 30 fold in cone pedicles compared to darkness. These latter results reveal that during light adaptation cone pedicles rapidly and efficiently lowered their intraterminal [Ca^{2+}] compared to rod spherules. This is consistent with our immunocytochemical, EM and ET findings that cone pedicles possess structural and functional mechanisms for the more rapid removal of intraterminal free Ca^{2+} than rod spherules (vide supra).

The microdomains of high [Ca^{2+}] were larger and closer to the plasma membrane in dark-adapted (depolarized) cone pedicles (Figure 13C,H) compared to rod spherules (Figure 13E,F). Considering the various preparation procedures and length of time that the retinas were incubated in different buffers, the mitochondria and synaptic terminals are intact and relatively well preserved compared to our immediately fixed tissue (Figure 9). In addition, the Ca^{2+} and fluo-3 calibration curve, used to estimate free [Ca^{2+}] in the photoreceptor synaptic terminals, is presented in Figure 13G.

Table 4 presents the Ca^{2+}-fluo-3 RFI values and the estimated mean free [Ca^{2+}] for the dark- and light-adapted rod spherules and cone pedicles. The estimated free [Ca^{2+}] in dark-adapted rod spherules was 3.2 fold lower (about 2.2 µM) than in cone pedicles (about 6.8 µM), which is consistent with our hypothesis that dark-adapted rod spherules maintain a lower mean resting [Ca^{2+}] than cone pedicles. In contrast, the estimated free [Ca^{2+}] in light-adapted cone pedicles was 3.2 fold lower (about 0.2 µM) than in light-adapted rod spherules (about 0.7 µM). Thus, during light adaptation the estimated free [Ca^{2+}] decreased 2 fold in rod spherules and almost 30 fold in cone pedicles compared to darkness. These latter results reveal that during light adaptation cone pedicles rapidly and efficiently lowered their intraterminal [Ca^{2+}] compared to rod spherules. This is consistent with our immunocytochemical, EM and ET findings that cone pedicles possess structural and functional mechanisms for the more rapid removal of intraterminal free Ca^{2+} than rod spherules (vide supra).

The tissue or cell type, number of measurements, mean crista junction diameter±SD, and reference are presented.

![Figure 12](http://www.molvis.org/molvis/v13/a97/) A-C: Comparison of cristae measurements in rod and cone synaptic terminal and inner segment mitochondria. The reconstructed portion of the mitochondrion was segmented along membranes: outer membrane, inner boundary membrane and cristae membranes. The programs Synuarea and Synuvolume calculated the volume and surface area values for the outer membrane (used for “mito” in the denominator) and each cristae. The values for individual cristae were summed to provide the number per volume (A) or numerator of the ratios (B and C). The mean values (±SEMs) for mitochondria in rod spherules (blue bars) and cone pedicles (red bars) are presented in each plot. In addition, our previously published mean (±SEMs) values for rod inner segment (black bars) and cone inner segment (green bars) mitochondria are plotted for ease of comparison [55]. Measurements were conducted on the tomographic reconstructions from three rods and three cones from three different mice. A two-tailed Student’s t-test determined significant differences: *p<0.05; **p<0.01.
In dark-adapted retinas, we estimated that the Ca$^{2+}$ microdomains were 300-400 nm in rod spherules (Figure 13A) and 600-700 nm in cone pedicles, where they overlapped in the vicinity of the synaptic ribbons (Figure 13C). Furthermore, the RFI appeared higher in the presumed location of the rod spherule mitochondria than in the cone pedicle area where mitochondria cluster. This suggests that rod mitochondria maintain a higher matrix [Ca$^{2+}$] than cone mitochondria, which is consistent with the location of rod spherule mitochondria and our suggestion that rod spherule mitochondria actively participate in intraterminal Ca$^{2+}$ buffering.

| Table 4. Ca$^{2+}$-Fluo-3 Relative Fluorescence Intensity Values in Dark-Adapted and Light-Adapted Rod and Cone Photoreceptor Synaptic Terminals |
|-----------------|-----------------|-----------------|
| Rod Spherules   | 180.5±4.9       | 138.2±6.7       |
| Cone Pedicles   | 214.8±9.7       | 82.7±4.6        |

All means±SEM relative fluorescence intensity values are scaled from 0 to 255 as described in the Materials and Methods section and as shown in Figure 13.

Figure 13. Ca$^{2+}$ imaging and correlative electron microscopy of rod and cone synaptic terminals in dark-adapted and light-adapted retinas. A-D: Pseudocolored Ca$^{2+}$-fluo-3 confocal images obtained from an adjacent pair of rod spherules (A, B) and a single cone pedicle (B and D) during dark adaptation (A and C) and light-adaptation (B and D). Scale bar equal 1 μm. The pseudocolor scale represents pixel intensity from 0 to 255 gray levels. The RFI values and estimated free [Ca$^{2+}$] in dark-adapted rod spherules were lower than in cone pedicles, whereas the RFI values and estimated free [Ca$^{2+}$] in light-adapted cone pedicles were markedly lower than in light-adapted rod spherules (Table 4). During light adaptation, the cone pedicles lowered their intraterminal [Ca$^{2+}$] about 3 fold more than rod spherules (Table 4). G-K: For higher resolution and better visualization of the Ca$^{2+}$ microdomains, the pseudocolored images were transformed using an advanced image synthesis algorithm. The microdomains of high [Ca$^{2+}$] were larger and closer to the plasma membrane in dark-adapted cone pedicles (C and H) compared to rod spherules (A and G). The scale bar and pseudocolor scale are as presented in D. E and F: The electron micrographs of the rod spherules E and cone pedicle F correspond to the rod spherules in panels A and B and cone pedicle in C and D, respectively. RN: rod nuclei, m: mitochondrion. Scale bar equal 1 μm for both panels. I: A fluo-3 Ca$^{2+}$ calibration curve used to estimate the free [Ca$^{2+}$] in the photoreceptor synaptic terminals (see Table 4).
The presynaptic terminals of the bullfrog saccular and axolotl lateral-line hair cells contain dense bodies that reversibly bind fluo-3 with a Kd of 550 nM [129]. We conducted high magnification electron microscopy studies on dark-adapted mouse retinas to determine whether the Ca\(^{2+}\) microdomains in our dark-adapted photoreceptor synaptic terminals might be due to the binding of fluo-3 to presynaptic dense bodies. Consistent with the findings of Vollrath et al. [130], we did not see any dense spheres in the photoreceptor terminals of our dark-adapted mouse retinas. Thus, we conclude that the Ca\(^{2+}\) microdomains do not result from fluo-3 binding to this synaptic organelle.

**DISCUSSION**

This study tested several functionally-based hypotheses related to the metabolic coupling and cross-talk between rod and cone synaptic terminal mitochondria, ER, PMCA and NCX and their potential roles in generating/updating ATP and regulating Ca\(^{2+}\) dynamics for neurotransmitter release. Three novel and significant sets of results were obtained. First, we characterized the spatial interrelation between mitochondria, ER, Ca\(^{2+}\) transporters and active zones in ribbon synapses as well as determined these details in the retina and photoreceptor synaptic terminals. Second, we determined the distribution, number and structure of mitochondria in rod spherules and cone pedicles as well as their substructure by utilizing ET. Third, we determined the mean levels of free [Ca\(^{2+}\)] in rod and cone synaptic terminals of whole isolated retinas during darkness and light adaptation.

Overall, our LSCM results revealed that retinal mitochondria exhibit laminar, cellular and compartmental segregation in different retinal neurons: see Table 2 and schematic summary Figure 14. For example, a strong-intense COX IV staining pattern was evident in mitochondria located in IS, OPL, IPL-\(\alpha\), IPL-\(\beta\), GCL and Müller glial cell end-feet, whereas the INL only had weak-moderate COX IV labeling. The photoreceptors displayed three separate compartments of strong to intense COX IV labeling: IS, portions of the ONL, and the OPL. The CIS labeled more intensely for COX IV than the RIS, consistent with their two-fold higher mitochondrial content [55]. In contrast, bipolar cells exhibited differential staining such that there was moderate COX IV staining in their somas while the terminals labeled strongly. The intense labeling of IPL-\(\alpha\) (ON-sublamina) and IPL-\(\beta\) (OFF-sublamina) with...
all three mitochondrial markers likely reflects the high amount of ATP required for glutamate uptake into synaptic vesicle and synaptic vesicle priming at ribbon synapses [1,33,34]. The above observations are consistent with functional results showing that mitochondrial oxygen consumption is higher in photoreceptors than in the inner retina during darkness and illumination [49,97] and that the mean inner retinal oxygen consumption is similar during dark and light adaptation [49,50,101]. In addition, the COX IV and POLG staining patterns were coincident in all retinal layers, except the INL. This suggests that mitochondria located in regions of high metabolic demand, where they might readily produce reactive oxygen species and initiate apoptosis, also might have an increased capacity for mitochondrial DNA replication and repair.

The overall VDAC immunolabeling pattern was similar to that of COX IV and POLG: illustrated by the areas of COX IV and VDAC colocalization. However, VDAC only weakly-moderately labeled IS. This lower IS immunofluorescence, using the pan-VDAC antibody, is not due to antibody specificity or tissue penetration problems, since this antibody recognizes a conserved sequence in all three VDAC isoforms (BLAST) and intensely stained mitochondria in the OPL. The weaker VDAC labeling might result from a low concentration of ER in the IS, since VDAC is found in ER closely apposed to mitochondria [131]. However, this is not likely since the calreticulin, SERCA3 and EM findings reveal an extensive amount of ER in RIS and CIS. Thus, VDAC protein expression is relatively low in mouse IS: a conclusion supported

Figure 14. (Legend previous page).
by in situ hybridization results [132]. This raises the intriguing possibility that lower levels of VDAC2 in IS and particularly RIS, increase the susceptibility of rods to mitochondrially mediated apoptosis [40,94,133] as VDAC2 inhibits BAK activation and apoptosis [134]. Finally, the strong to intense VDAC labeling and colocalization with VGluT1 in the plexiform layers suggests another possible level of metabolic coupling between mitochondria and synaptic transmission, since physiological concentrations of glutamate modulate VDAC channel activity [135].

During the LSCM and EM studies, we unexpectedly found distinct and highly localized COX IV-, VDAC- and POLG-positive mitochondria associated with cone and rod nuclei. First, in the most distal ONL large juxtanuclear mitochondria were located above and below the cone nuclei. The mitochondrion below the cone nucleus is in the axon, as described [53]. Second, in the most proximal ONL numerous single, smaller juxtanuclear mitochondria were situated above each rod nucleus. It is likely that these juxtanuclear mitochondria are involved with somal ATP synthesis, phosphotransfer networks and nucleocytoplasmic communication [136]. For example, cone nuclei contain larger amounts of electron lucent, transcriptionally active euchromatin compared to rods ([53]; Figure 3 and Figure 8A). This suggests that transcription and other ATP-requiring nuclear processes [136,137] are more active in cones than rods. In addition, cone nuclei are located in the most distal portion of the ONL, such that their somas are further from their synaptic terminals than those of rods. Moreover, glycolytic and glucose-6-phosphate dehydrogenase enzyme activities are low in the ONL compared to other photoreceptor compartments [138] and ATP has a short diffusion distance [139]. Thus, local ATP synthesis is particularly important for these cone nuclei. The proximal rod somas likely are dependent upon these juxtanuclear mitochondria because they lack a classic spherule that contains a single mitochondrion, although they possess a ribbon synapse and active zone at the base of their somas [122]. Interestingly, these rod juxtanuclear mitochondria and somas appear selectively vulnerable to Ca2+ overload or oxidative stress [39,140,141].

The mechanisms and molecular signals that direct retinal mitochondria to their different cellular and compartmental sites are unknown. Neuronal mitochondria, compared to other organelles, have several unique properties with respect to axoplasmic transport. First, they undergo both anterograde and retrograde transport. Second, their motility is characterized by saltatory movements and prolonged stationary phases. Third, the net direction of movement varies with the physiological status of the axon. Fourth, mitochondria utilize microtubules and actin-myosin based microfilament systems for motility, and neurofilament associations for static phases within axons [142-144]. The molecular signals that direct mitochondria to their cellular positions include: high local ADP:ATP in areas of high ATP demand [145], GTP hydrolysis by small GTP-binding proteins [146], actin-myosin-dependent nerve growth factor/TrkA/PI 3-kinase signaling [147], and various kinases [148,149]. Intrinsically mitochondrial proteins, such as OPA1, also may contribute to the selective distribution, clustering and stability of mitochondria in retinal neurons in addition to their role in mitochondrial fission and fusion, genome maintenance, and regulation of cristae morphology and substructure [150,151].

Eight new anatomical, ultrastructural and/or substructural differences between mouse rod spherules and cone pedicles were found. First, EM studies showed that there were three to five rows of rod spherules and a single row of larger, more electron lucent cone pedicles in the OPL. High magnification double labeling LSCM experiments, using either COX IV and M- and S-opsins or VDAC and rhodopsin, yielded complimentary results. Second, the cone pedicle volume is about 10 times larger than the rod spherule volume. Third, rod spherules contain a single very large ovoid mitochondrion that occupies 20-25% of the spherule volume, whereas cone pedicles contain an average of five medium-sized mitochondria that occupy 10-15% of the pedicle volume. Fourth, the rod spherule mitochondrion is located close to synaptic vesicle release sites, whereas cone pedicle mitochondria cluster in the distal part of the terminal. Fifth, the abundance of cristae membranes and correspondingly the cristae volume are two-fold greater in the rod spherule mitochondrion compared to an individual cone pedicle mitochondrion. Overall, however, the total amount of cristae membranes is 2 fold higher in cone pedicles than in rod spherules. Sixth, mitochondria CJ openings are 25% smaller in cone pedicle mitochondria than in rod spherule mitochondria. Seventh, networks of ER are in rod, as seen in rats [28], and cone synaptic terminals, although they are larger and denser in cone pedicles. Eighth, rod spherules uniformly and intensely stain for PMCA, whereas cone pedicles preferentially stain for NCX1 at their active zones. As reported by others, we found that mouse rod spherules mostly (about 95%) contain one ribbon synapse, but can contain two ribbon synaptic units [152,153], whereas cone pedicles contain 6-14 ribbon synapses [52,53; data not shown].

A comparison of our new substructural data on synaptic terminal mitochondria with that previously obtained on IS mitochondria [40,55], reveals three significant differences (see Figure 11). First, rod spherule and cone pedicle mitochondrial cristae segments (i.e., tubes and lamellae) do not exhibit high connectivity, whereas IS cristae segments are highly connected to each other so that a single crista can have many segments. CIS connectivity is maze-like [55]. Moreover, the number of segments per crista in CIS is significantly greater than in RIS. As opposed to IS mitochondria that connect many cristae segments together, rod spherule and cone pedicle mitochondria accommodate their greater amount of inner membrane by increasing the number of cristae. The mean number of cristae per mitochondrial volume for rod spherules and cone pedicles was about 1,300 and about 1,100, respectively, whereas it was about 570 and about 520 for RIS and CIS, respectively. Second, the cristae surface to mitochondrial surface ratio and the cristae volume to mitochondria volume ratio are 1.4- to 2.0 fold larger in CIS compared to RIS [55]. In marked contrast, the abundance of cristae membranes and correspondingly the cristae volume are 2.5 fold and 2.2 fold, respectively, greater in rod spherule mitochondria, than in each cone pedicle mito-
The CJ opening in cone pedicle mitochondria is smaller than any previously measured in normal tissue or cells. Based on results that model the effects of matrix outward pressure on CJ diameter [155], we suggest that these differences in rod and cone mitochondria CJ diameter reflect small differences in matrix pressure. For example, when the matrix volume of mitochondria is experimentally increased, CJs become smaller and can pinch off leaving some cristae detached from the IBM [126,127]. The putative bioenergetic role of CJ remains to be determined. One hypothesis, consistent with the computer simulations, suggests that CJs partially regulate ATP generation by restricting the diffusion of ADP and ATP into and out of cristae [126]. That is, larger CJ openings facilitate and increase the capacity for ATP production. Alternatively, smaller CJs may restrict the diffusion of cytochrome c from the cristae lumen and thereby minimize diffusion distances that limit rates of electron transport needed for ATP synthesis. This would confer an energetic advantage to cone pedicle mitochondria and provide them with increased resistance to mitochondrially mediated apoptosis [40,156].

The biological and functional implications of the different number and position of mitochondria in rod spherules and cone pedicles are three-fold. The first relates to the optimal generation and utilization of ATP for various cellular processes and synaptic transmission. The second involves the coordinated regulation of presynaptic Ca\(^{2+}\) levels necessary for synaptic transmission. The third is associated with potential Ca\(^{2+}\) overload and cell survival. The first consideration addresses two interrelated questions: (i) Is the ATP demand higher in a rod spherule or cone pedicle? (ii) Does a rod spherule or cone pedicle produce more ATP under normal physiological conditions? A number of observations strongly indicate that mouse (mammalian) cone pedicles have a higher ATP demand than rod spherules. Mouse cone pedicles have about 10 times more ribbons than rods (vide supra). Primate cone pedicles can have up to 50 synaptic ribbons [114,115]. Mammalian cone ribbons dock and tether approximately five-fold more vesicles than rod ribbons [116]. These vesicles require ATP for glutamate uptake and priming [33,34]. Cones have a faster initial component of exocytosis [18] and by extension a more rapid need to refill and prime the vesicle pool [34,117]. In addition, cones operate over a much larger dynamic range [118,119]. Biophysical and modeling experiments reveal that ATP demand logarithmically increases as the rate of luminance information transferred to second-order neurons increases [120], indicating that the metabolic cost of information transfer is higher in cones than rods. Photoreceptor NAKA, which has a specific activity three times higher than the whole retina [97,157] and consumes greater than 50% of the cell’s ATP [158], is higher in cone pedicles than rod spherules [47]. One major reason for the higher Na\(^+\) pump activity and associated ATP demand in cone pedicles is that pedicles preferentially utilize NCX to extrude Ca\(^{2+}\) from their active zones (this paper; see schematic in Figure 14), and NCX activity is mostly controlled by NAKA activity [106]. In addition, selected areas of the cone axon and synaptic terminal utilize PMCA to extrude Ca\(^{2+}\), whereas this is the major Ca\(^{2+}\) extrusion mechanism in rod spherules (this paper). Moreover, our detailed results are different from those obtained in the calyx-type presynaptic nerve terminals of the chick ciliary ganglia [6,106], cultured mouse brain astrocytes [6,106] and rabbit skeletal muscle [107]. Interestingly, the presynaptic distribution of NCX and PMCA in mouse cone pedicles is opposite to that in the chick ciliary ganglia [106]. This difference may relate to the presence of N-type Ca\(^{2+}\) channels in the presynaptic terminal of the chick ganglia [159] as opposed to L-type Ca\(^{2+}\) channels in photoreceptor synapses [161,162].

ATP production in cone pedicle mitochondria is also likely greater than that in rod spherules. This is similar to our findings on RIS and CIS mitochondria [55,56]. Although the rod spherule mitochondrion has 2 fold more cristae surface area than each cone mitochondrion, cone pedicles have five mitochondria per synaptic terminal and thus twice the total amount of cristae membranes. In addition, the CJ openings are smaller in cones, which could lead to more effective coupling of the electron transport chain to ATP synthesis [126]. Together, these should facilitate increased ATP production as evidenced by the higher COX activity in cone pedicles, compared to rod spherules [47]. Moreover, mammalian cone pedicles, but not rod spherules, contain glycogen and the phosphorylase isozyme necessary to convert it to glucose [162,163]. In contrast, the cone pedicle volume is large and the mitochondria cluster some distance from active zones. These factors will decrease the overall effective cellular ATP concentration at the active zones since the ATP diffusion distance is small [139]. Nevertheless, and likely more important for regulating the pedicle Ca\(^{2+}\) concentration, these clustered mitochondria will provide ATP for the nearby PMCA (see high magnification LSCM Figure 8D,E).

It is well known that strong metabolic coupling and cross-talk between mitochondria, ER, NCX, PMCA, and Ca\(^{2+}\) channels regulate presynaptic Ca\(^{2+}\) dynamics and neurotransmitter release [1,2,5,37,164]. Our high magnification immunocytochemical studies revealed several significant, and presumably functionally relevant, differences in the location and distribution of mitochondria, PMCA and NCX in rod spherules and cone pedicles. As noted, the rod spherule mitochondrion is located close to synaptic vesicle release sites, whereas cone pedicle mitochondria cluster in the distal part of the terminal. Neuronal mitochondria have a low affinity, but high capacity for Ca\(^{2+}\) (Kd approximately 1-5 µM; 1-5 µmol mg protein\(^{-1}\)) [165,166]. NCX and PMCA are the two major Ca\(^{2+}\) transport-
ers in the presynaptic terminal [2,106]. We found that in the cone axon the active zone of pedicles preferentially and intensely stained for NCX, whereas rod spherules diffusely stained for NCX. In contrast, rod spherules uniformly and intensely stained for PMCA, whereas only the distal portion of the cone pedicle stained for PMCA. Triple labeling experiments with PMCA, VDAC and either VGluT1 or PNA revealed that the cone pedicle mitochondria were in close proximity to the PMCA labeled membranes. NCX utilizes the activity and gradient established by NAKA to maintain low intracellular Ca²⁺ levels, and is spatiotemporally associated with the two high ouabain affinity isozymes in neurons (α2: NAKA2 or α3: NAKA3). Immunocytochemical and biochemical experiments show that NAKA3 is present in rodent IS and synaptic terminals [97,167]. NCX has a low affinity and high turnover rate for Ca²⁺ extrusion (Kd approximately 1 μM; kcat approximately 30-250 s⁻¹), whereas PMCA has a high affinity and a low turnover rate for Ca²⁺ extrusion (Kd approximately 0.1 μM; kcat approximately 1-2 μM) [106,168]. This suggests that the synaptic terminal Ca²⁺ concentration is rapidly lowered by NCX to its Kd value, while the high affinity PMCA slowly lowers it to a dark-adapted value of 0.3-2 μM [19,26]. ER cisternae were widely distributed throughout the rod and cone terminals and often were in close apposition to mitochondria, as observed in other tissues [89,104,105]. SERCA3 extensively labeled the IS and OPL. It has a low affinity and relatively low turnover rate (Kd approximately 1-2 μM; kcat approximately 130-150 s⁻¹), but ER have a high capacity for Ca²⁺ uptake (0.5-1 μM) [169,170]. This is consistent with the role of ER Ca²⁺ uptake, which helps lower the intraterminal Ca²⁺ concentration with PMCA and NCX.

Our results, in concert with those from many other studies, enabled us to construct a spatiotemporal model of rod spherule and cone pedicle regulation of Ca²⁺ levels during darkness (see Figure 14 for illustration) and light adaptation. In the dark-adapted mouse retina, a low level of Ca²⁺ enters rod and cone synaptic terminals through Ca₁.4 and Ca₁.3 channels, respectively [160,161], and glutamate is continuously released. Although dark-adapted mammalian rods are slightly more depolarized than cones [61], it appears that the activation values for these voltage-gated calcium channels match their resting membrane potentials [161,171]. This should produce one or more, depending on the number of active zones, overlapping microdomains of elevated plasmalemmal Ca²⁺. By extension, this should result in higher Ca²⁺ concentrations around the cone active zones, compared to the rod active zone, which likely accounts for the 10 fold faster initial rate of exocytosis in salamander cones compared to rods [18]. In dark-adapted salamander rod terminals, the intraterminal Ca²⁺ concentration is estimated to range from 0.3-2 μM [19,26], whereas the microdomains likely contain 20-40 μM Ca²⁺ [17,172]. Some of this Ca²⁺ in the microdomain will bind to CaBP4, a Ca²⁺ binding protein in photoreceptor terminals that directly associates with Ca₁.4 channels [173], and synaptotagmin 1 in the active zone [70,113]. Following light adaptation, NCX should rapidly remove much of the remaining intraterminal Ca²⁺ in cooperation with the high capacity ER (SERCA3-positive): especially in cone pedicles where NCX localizes to the active zones. However, NCX will only lower the intraterminal Ca²⁺ until the ATP-dependent NAKA3 re-establishes the Na⁺ gradient. In cones, further Ca²⁺ extrusion must be facilitated by the slower ATP-dependent PMCA, which can maintain submicromolar levels of Ca²⁺ [17,175]. Presumably, in rods the PMCA are working continuously to maintain a spatially averaged Ca²⁺ concentration of 0.3-2 μM: the concentration estimated to maintain vesicle cycling in the salamander rod [19,26].

To test this spatiotemporal model of [Ca²⁺] regulation in dark- and light-adapted rod spherules and cone pedicles, we conducted Ca²⁺ imaging studies on retinal slices utilizing fluo-3: a high affinity fluorescent Ca²⁺ dye. These experiments produced three novel and functionally relevant results that validated several of our predictions. First, in dark-adapted retinas the estimated free [Ca²⁺] in rod spherules was about 2 μM, which was significantly (3.2 fold) lower than that in cones pedicles. These findings are consistent with the interpretation of our detailed immunocytochemical findings. That is, rod spherules predominantly possess high affinity/low turnover PMCA that maintains a low intraterminal [Ca²⁺] in order to increase the sensitivity and signal-to-noise ratio of rods [17-21]. Furthermore, the RFI appeared higher in the presumed location of the rod spherule mitochondria than in the cone pedicle area where mitochondria cluster. This observation is in agreement with our suggestion that rod spherule mitochondria actively participate in intraterminal Ca²⁺ buffering, which also increases their ATP production capacity. The recent findings of Krizaj and colleagues [176], who compared the morphology and light responses of retinas from control mice and deafwaddler dfw2J mice that lack the functional PMCA2 protein, confirm our suggestion and results that PMCA-mediated Ca²⁺ extrusion selectively modulates the rod spherules and not the cone pedicles. That is, the rod-mediated electroretinogram b-waves recorded from dfw2J mice were markedly smaller and slower than those in control mice, however, there were no alterations in the cone-mediated b-wave. At the light microscopy level, the retinas from both mice looked similar.

Second, during light adaptation the estimated free [Ca²⁺] in cone pedicles was 0.23 μM, which is significantly (3.2 fold) lower than that in rod spherules. This is consistent with our immunocytochemical, EM and ET findings that cone pedicles possess the structural components and functional mechanisms (e.g., low affinity/high turnover NCX) to more rapidly remove intraterminal free Ca²⁺ compared to rod spherules as well as PMCA to lower the free [Ca²⁺] below the Kd of NCX. Furthermore, it is compatible with the observation that the recovery and kinetics of exocytosis is faster in cones than rods [9,18,118,119].

Third, in dark-adapted retinas Ca²⁺ microdomains were observed adjacent to the plasmalemma and synaptic ribbons of rod spherules and cone pedicles. These Ca²⁺ microdomains were larger in cone pedicles than rod spherules and they overlapped in cone pedicles. Interestingly, the estimated size of the Ca²⁺ microdomains in dark-adapted cone pedicles were similar to those measured in dark-adapted goldfish Mb1 re-
nal cells [24]. Although the relation between Ca\(^{2+}\) microdomains in rod and cone synaptic terminals and the regulation of exocytosis remains to be determined, our Ca\(^{2+}\) imaging findings suggest that cone pedicles will release more synaptic vesicles and exhibit faster exocytosis than rods spherules. This is consistent with the physiology of these two photoreceptor terminals [9,17-21,36] and the goldfish MbI retinal cells [9,24,36]. Ca\(^{2+}\) microdomains, which cluster near Ca\(^{2+}\) channels, were visualized at the ribbon synapses of frog saccular hair cells, turtle cochlear hair cells and goldfish retinal MbI cells [24,177-179].

The mitochondria certainly provide the Mg-ATP necessary for NAKA3, SERCA3 and PMCA in both rod spherules and cone pedicles. The proximity of the rod spherule mitochondria to the active zone and its relatively high Ca\(^{2+}\) concentration (at least 20 µM) leads us to speculate that these mitochondria play two additional important and interrelated roles. First, they may participate in local Ca\(^{2+}\) buffering in order to limit the expansion of the Ca\(^{2+}\) microdomain [24] and thereby maintain the rod’s single photon sensitivity [20,21]. This likely operates because as the cytosolic ATP concentration decreases and ADP concentration increases, the electrogenic mitochondrial Ca\(^{2+}\) carrier (uniporter) increases its rate of Ca\(^{2+}\) uptake and retention of Ca\(^{2+}\) loads. When the cytosolic Ca\(^{2+}\) concentration returns to baseline, mitochondria release their stored Ca\(^{2+}\) loads [165,166]. Second, the activity of the mitochondrial dehydrogenases increases in response to the increase in the free mitochondrial matrix Ca\(^{2+}\) concentration. This results in an increase in the rate of electron transport and subsequently the amount of ATP production [180,181]. In summary, we suggest that reciprocal cooperation between mitochondria, ER and the Ca\(^{2+}\) transporters in rod spherules and cone pedicles during darkness control presynaptic Ca\(^{2+}\) handling, ATP production and glutamate release (Figure 14).

The third possibility, though not mutually exclusive from the first two, is that cone pedicle mitochondria cluster at a distance from active sites to protect themselves from Ca\(^{2+}\) overload during synaptic transmitter release. As noted, in dark-adapted retinas there are likely numerous overlapping microdomains of elevated Ca\(^{2+}\) close to the synaptic ribbons of the cone pedicle. Moreover, this steady Ca\(^{2+}\) influx could produce a Ca\(^{2+}\) induced Ca\(^{2+}\) release (CICR) from presynaptic rod and cone SER [28; this paper], which would preferentially produce Ca\(^{2+}\) overload in cone pedicle mitochondria. In the salamander rod synapse, there is a ryanodine-dependent amplification mechanism that couples CICR with continuous vesicle release [26,27]. Consistent with this concept, a recent report demonstrated that synaptic mitochondria are more susceptible to Ca\(^{2+}\) overload and mitochondrial permeability transition than non-synaptic mitochondria [182]. In addition, a similar phenomenon may occur in rod and cone synaptic terminals, like in RIS [40,94], as rod spherule mitochondria undergo permeability transition following postnatal only developmental lead exposure [data not shown, manuscript in preparation]. Moreover, both rod and cone synaptic mitochondria undergo permeability transition in an HRG4 mutant mice that models human cone-rod dystrophy [183].

In summary, rod spherules and cone pedicles have significantly different morphology, ultrastructure, mitochondrial density and location, patterns of protein expression, and regulatory mechanisms that control their [Ca\(^{2+}\)] during dark and light-adaptation. Compared to rod spherule mitochondria, individual cone pedicle mitochondria are smaller and more ovoid, reside more remotely from the active zone, possess less cris-tae surface and volume, and have a very small CJ diameter. These distinct characteristics play a fundamental role in the ability of rod and cone presynaptic terminals to regulate and produce ATP as well as to buffer Ca\(^{2+}\) microdomains. In addition, the ER, PMCA and NCX all play essential and interdependent roles in rod and cone presynaptic terminals. Their differences in cellular distribution and location, amount of protein expression, affinity and capacity for Ca\(^{2+}\) and their turnover rate of Ca\(^{2+}\) significantly contribute to the kinetics of exocytosis in rods and cones. The results from these LSCM, EM, ET, and Ca\(^{2+}\) imaging experiments can help design new electrophysiological, pharmacological, molecular/transgenic and imaging experiments that will enable us to gain a deeper and fuller understanding of the spatiotemporal control of vertebrate rod and cone synaptic terminal Ca\(^{2+}\) concentrations and mitochondrial ATP production as they relate to exocytosis and endocytosis. In addition, our results illustrate the need to identify and determine the spatial distribution, compartment-specific differences and interrelation of Ca\(^{2+}\)-handling proteins and mitochondria in other neural tissues so that the structure-function relations in different neurons and chemical synapses can be ascertained. Moreover, additional new retinal experiments will provide important data on the role and sensitivity of rod and cone synaptic terminal mitochondria to Ca\(^{2+}\) overload and cell injury as compared to RIS and CIS mitochondria [39,40,182]. Finally, we hope that the results from our work, the suggested experiments as well as future experiments will provide further groundwork for understanding, and treating, the pathophysiological effects and visual deficits that result from inherited, disease-related as well as chemically- and pharmacologically-induced retinal degenerations.

ACKNOWLEDGEMENTS

We thank Dr. David M. Sherry for expert technical advice and comments on an earlier version of the manuscript, and Dr. Bo Huang, Dr. Ann T. Poblenz and Ms. Margaret Gondo for technical assistance. Partially funded by NIH Grant ES012482 to DAF, and an NIH Core Grant EY07024 and Training Grant EY07088 to UHCO. Part of this work was performed at the National Center for Microscopy and Imaging Research supported by NIH Resource Grant RR04050 and NS14718 to MHE. Part of this data was presented by DAF during an invited talk at the Ninth Annual Vision Research Conference, entitled “Neuroimaging in the Retina”, held on April 29 and 30, 2005 in Ft. Lauderdale, FL. Other parts were presented on posters at the ARVO 2005, 2006, and 2007 annual meetings.
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