Visualization and quantification of cardiac mitochondrial protein clusters with STED microscopy

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Abstract

The visualization and quantification of mitochondria-associated proteins with high power microscopy methods is of particular interest to investigate protein architecture in this organelle. We report the usage of a custom-made STimulated Emission Depletion (STED) fluorescence nanoscope with ~30 nm lateral resolution for protein mapping of Percoll-purified viable mitochondria from murine heart. Using this approach, we were able to quantify and resolve distinct protein clusters within mitochondria; specifically, cytochrome c oxidase subunit 2 is distributed in clusters of ~28 nm; whereas the voltage dependent anion channel 1 displays three size distributions of ~33, ~55 and ~83 nm.

1. Introduction

Mitochondria participate in key metabolic reactions, regulate Ca2+ signaling, and play a central role in apoptosis (Dimmer and Scorrano, 2006; Liu et al., 1996; Yang et al., 1997). In the heart, mitochondria can determine myocardium survival and protection, whilst mitochondrial dysfunction contributes to heart disease (Baines, 2010). To understand the details of these mechanisms it is important to map mitochondria-associated proteins at nanometer resolution as localization is intimately related to functional performance.

STimulated Emission Depletion (STED) fluorescence nanoscopy with ~40–80 nm lateral resolution in biological samples offers the opportunity to scrutinize the distribution of proteins in subcellular compartments or organelles using common immunocytochemistry techniques and tagged proteins (Kellner et al., 2007; Neumann et al., 2010; Watanabe et al., 2011; Willig et al., 2006). STED has the advantage over electron microscopy that it reaches nanometer resolution while at the same time maintains the microscopic scale.

Only recently has STED been utilized to image a few mitochondrial proteins. Native hexokinase I was observed in mitochondria of an osteosarcoma cell line (U2OS) (Neumann et al., 2010); while ectopically expressed TOM20 was imaged delineating mitochondria periphery in a ring-like arrangement in Caenorhabditis elegans (Watanabe et al., 2011), and expressed voltage dependent anion channels (VDAC) 1–3 were visualized in U2OS cells with VDAC3 forming clusters of ~40 to 90 nm (Neumann et al., 2010).

The objective of this work was to map with STED nanoscopy two classical mitochondria proteins in cardiac mitochondria: VDAC1, an outer mitochondrial membrane protein that serves as an interface between cellular and mitochondrial metabolism (Shoshan-Barmatz and Ben-Hail, 2011); and cytochrome c oxidase (complex IV of the respiratory chain) located in the mitochondrial inner membrane and assembled by 13 subunits in humans. Specifically, we imaged cytochrome c oxidase's subunit 2 (Cox2) that forms part of the catalytic core of the enzyme (Brzezinski and Johansson, 2010; Mick et al., 2011).

2. Methods

2.1. Antibodies

Primary antibodies were used against VDAC1 (Ab14734, Abcam), Cox2 (A6407, Invitrogen), Cadherin (C1821, anti-Pan Cadherin...
antibody, Sigma), GM130 (ab1299, Abcam), Lamin B1 (ab16048, Abcam), GRP78 BiP (ab21685, Abcam), L-type Ca2+ channels (α1C) (ACC003, Alomone) and Src (sc-18, Santa Cruz). Secondary antibodies for Western blots were: Alexa Fluor® 680 goat anti-rabbit (A-21109, Invitrogen) and IRDye 800CW conjugated goat anti-mouse (926–32210, Li-COR); and for immunocytochemistry were: Atto 647 N goat anti-mouse (50185, Sigma), and Atto 647 N goat anti-rabbit (15048, Active Motif).

2.2. Animals

Protocols received institutional approval. Male 3 mo old C57BL/6NCrl mice were injected (i.p.) with heparin (200 IU/kg). After 20 min, animals were anesthetized with sodium pentobarbital (70 mg/kg, i.p.). The heart was surgically removed and rapidly arrested in ice-cold phosphate buffer saline (PBS) (in mM): 2.7 KCl, 10 HEPES-Na, 1 EDTA-Na2, pH 7.4. The sample was centrifuged in a fixed angle rotor at 50,000×g for 45 min. The homogenate was transferred into a 2.0 ml Eppendorf tube and centrifuged at 1,300×g for 3 min. The supernatant was carefully transferred into a clean 1.5 ml Eppendorf tube and centrifuged at 10,000×g for 10 min. The pellet containing crude mitochondria was resuspended in 55 μl of isolation buffer A.

The crude mitochondria preparation was carefully overlaid on 3 ml of 30% (v/v) Percoll (Graham, 2001) in buffer B (in mM): 250 sucrose, 10 HEPES-Na, 1 EDTA-Na2, 50 Tris–HCl, pH 7.4 using a Potter-Elvejem homogenizer (10 rapid strokes). The homogenate was transferred into a 2.0 ml Eppendorf tube and centrifuged at 1,300×g for 3 min. The supernatant was carefully transferred into a clean 1.5 ml Eppendorf tube and centrifuged at 10,000×g for 10 min. The pellet containing crude mitochondria was resuspended in buffer A for organelle immunochemistry and in buffer B for functional assays. Purified mitochondria were used within 2 h after isolation.

2.4. Western blot analysis

Isolated mitochondrial pellets from the different fractions were resuspended in lysis buffer (in mM): 50 Tris–HCl, 150 NaCl, 1 EDTA-Na2, 1 EGTA-Na4, 0.1% (w/v) Na-deoxycholate and 0.1% (w/v) SDS, pH 7.4 supplemented with protease inhibitors (1 tablet/50 ml, Roche), and incubated for 1 hour with shaking. Samples were centrifuged at 10,000×g for 5 min, the supernatants were collected, and the protein concentration was measured. Equal amounts of protein (50 μg) were loaded for 10% SDS-PAGE, and transferred to nitrocellulose membranes. Proper loading was corroborated with Ponceau S staining. Membranes were blocked with 5% (w/v) milk in TBS (150 mM NaCl, 20 mM Tris-HCl pH 7.4) for 60 min at room temperature. Membranes were washed three times with TBS (5 min at room temperature) and incubated overnight with primary antibodies at 4°C. Primary antibodies were used at the following concentrations: for plasma membrane (Caderherin, 19.4 μg/ml), endoplasmic reticulum (GRP78 BiP, 0.2 μg/ml), nuclear envelope (Lamin B1, 0.2 μg/ml), Golgi (GM130, 0.5 μg/ml), mitochondria (VDAC1, 0.2 μg/ml and Cox2, 0.1 μg/ml), T-tubules (α1C, 0.45 μg/ml) and for a signaling tyrosine kinase family (Src, 0.04 μg/ml). After washing three times with TBS (5 min, room temperature) membranes were incubated with 0.01 μg/ml secondary antibodies (anti-mouse/rabbit) for 1 h at room temperature, and washed again three times with TBS for 5 min.

2.5. Mitochondrial H2O2 measurement

Mitochondrial reactive oxygen species (ROS) generation was measured spectrophotometrically (560 nm excitation and 590 nm emission) by incubating 250 μg of mitochondrial protein in a solution containing (in mM): 20 Tris–HCl, 250 sucrose, 1 EDTA-Na4, 1 EDTA-Na2, and 0.15% (w/v) bovine serum albumin, pH 7.4 at 25 °C with 10μM Amplex Red (a H2O2 sensitive-dye, Invitrogen) and continuous stirring. Sodium succinate (3 mM) was used to activate complex II.

2.6. Labeling of mitochondrial proteins

Purified mitochondria were incubated with 500 nM Mitotracker® Red CMXRos (Invitrogen) for 60 min with shaking, plated drop-wise onto coverslips and incubated at 4 °C for 1 h. Coverslips (0.17 mm thickness; Warner Instruments, Cat no. 64–0712) were coated with 0.1% Poly-γ-Lysine (Sigma-Aldrich) for 2 h at room temperature and washed with PBS before use. Attached mitochondria were then washed with PBS once and fixed with 4% (w/v) paraformaldehyde in PBS for 10 min at room temperature. After three washes with PBS, mitochondria were permeabilized with 0.5% (v/v) Triton-X 100 in PBS for 10 min at room temperature. Mitochondria were further colabeled with specific antibodies (in 1% (w/v) normal goat serum (NGS), 0.2% (v/v) Triton X-100 in PBS) for mitochondrial (0.2 μg/ml anti-VDAC1 and 0.2 μg/ml anti-Cox2 monoclonal antibodies) and nuclear envelope (0.2 μg/ml anti-Lamin B1 polyclonal antibody) markers. ATTO 647 N-labelled secondary antibodies (1 μg/ml anti-rabbit and anti-rabbit) in 1% (w/v) NGS, 0.2% (v/v) Triton X-100 in PBS were used. Some mitochondria were only incubated with secondary antibodies as controls. Preparations were sequentially treated with 10, 25, 50 and 97% (v/v) 2,2′-Thiodiethanol (TDE, Sigma-Aldrich) solutions in PBS (pH 7.5) for 5 min each at room temperature (Staudt et al., 2007). TDE solutions were added and aspirated gently to prevent uplifting of mitochondria from the coverslips. Coverslips were mounted onto slides and sealed with regular nail polish. Samples used only for regular confocal microscopy were mounted using ProLong®Gold (P36934, Invitrogen) instead of TDE. Confocal images were acquired with an Olympus confocal microscope using a 60× oil immersion objective with 1.42 NA (PlanApo N) at a scanning resolution of 0.0575 μm per pixel. STED images were collected as described below.

2.7. STED microscopy

STED images were acquired with a custom-made STED system using an oil immersion objective (HCX PL APO CS 100x/1.40–0.70 OIL, Leica Germany). A 635 nm pulsed diode laser (LDH-D–C–635, PicoQuant GmbH) was used for excitation. The pulses for STED deple- tion were delivered by a tunable Ti:sapphire laser (Mai Tai HP, Spectra Physics) set at 780 nm. Fluorescence emission from ATTO 647 N-labeled secondary antibodies was collected through a Semrock BrightLine FF01-685/40–25 nm band pass filter in front of a photomultiplier (H7422PA-40, Hamamatsu Photonics K.K.). Images (955 × 960 pixels) were acquired with a 16 kHz line frequency (reso- nant mirror of 8 kHz) and summed 256 times. Pixel size was ~9.575 nm × 9.575 nm. For a fair comparison between conventional confocal images and STED images, all imaging parameters were kept identical except for the number of summations which was 64 when recording confocal images. For comparison, confocal images were acquired for the same field prior STED imaging.
2.8. Image analysis

Confocal double-labeled images were median filtered and analyzed with custom-made software to determine their protein proximity index as previously described (Li et al., 2010; Wu et al., 2010). For display, STED images were deconvolved by 20 iterations of the Richardson-Lucy deconvolution algorithm and median filtered with a radius of 2 pixels using the plug-in DeconvolutionLab developed for ImageJ by Cédric Vonesch, Raquel Terrés Cristofani and Guillaume Schmit at the Biomedical Image Group (BIG), EPFL, Switzerland. Analysis of cluster size was done using the raw images without any modifications by performing line scans across the clusters to measure pixel intensity as a function of distance and manually estimating the full width at half maximum (FWHM) of the light intensity peaks above the baseline noise. Cluster sizes were measured from 500 clusters each from three independent experiments. Histograms were manually fitted using IGOR Pro (WaveMetrics Inc.).

2.9. Statistics

Error bars were plotted as standard errors of the mean (±SEM) for a minimum of three independent experiments. Two-tailed Student’s t-test was performed and $p<0.05$ was considered significantly different.

3. Results

To image VDAC1 and Cox2 in purified murine cardiac mitochondria at the nanoscopic level, our strategy was to first evaluate the
purity and viability of our mitochondrial preparation using biochemical and immunochemical approaches followed by STED fluorescence nanoscopy analysis.

### 3.1. Mitochondrial fractions: purity and functional evaluation

Purified mitochondria were obtained using a 30% self-generated Percoll density gradient containing 250 mM sucrose (Graham, 2001; Zhang et al., 2008). Fig. 1A shows three distinct fractions obtained after Percoll separation, named M1, M2 and M3. The purity and viability of the mitochondrial fractions was assessed using three complementary parameters: 

- Production of reactive oxygen species (ROS),
- Western blot analysis of organelle protein markers; and
- Colocalization with mitotracker which labels mitochondria with intact membrane potential.

The viability of M1, M2 and M3 fractions was first tested by measuring ROS production using Amplex Red, a H$_2$O$_2$-sensitive dye, and succinate (3 mM), a substrate of complex II. As shown in Fig. 1B, mitochondria from M3 fraction maintained the same efficiency to produce ROS as crude mitochondria; however, M1 and M2 had only 25% and 50% of crude mitochondria capacity to produce ROS, respectively. These results indicated to us that M3 fraction was the most viable of the three purified fractions.

Next, M1, M2 and M3 fractions along with crude mitochondria (CM) and heart ventricular lysates (HVL) were examined for the presence of classical protein markers of different organelles. We used antibodies against VDAC1 and Cox2 for mitochondria, Cadherin for plasma membrane, GM130 for Golgi, Lamin B1 for nuclear envelope, and against GRP78 BIP for endoplasmic reticulum. In addition, we used antibodies against the L-type Ca$^{2+}$ channel α1C subunit, a T-tubule marker, and for Src, a family of signaling tyrosine kinases (SFKs). SFKs, known to be tethered to the plasma-membrane or with cytoplasmic localization, have been observed in mitochondria from brain by immunogold electron microscopy (Salvi et al., 2002; Tibaldi et al., 2008) and by immunocytochemistry in primary mouse pre-leptotene spermatocytes GC2 cells (Livigni et al., 2006); while in the heart, the presence of SFKs in mitochondria is supported by pharmacology and site- and phosphorylation state-directed antibodies showing increased Src-dependent phosphorylation of the mitochondrial protein, adenine nucleotide translocator 1 in isoflurane-preconditioned heart (Feng et al., 2010). Western blots in Fig. 1C show that M3 fraction produced robust signals for VDAC1 and Cox2 but no significant signals for Cadherin, GM130, Lamin B1, GRP78 BIP, α1C, and Src. Quantification of normalized signals (to Ponceau S signals) demonstrates that M3 fraction is significantly enriched in VDAC1 and Cox2 with respect to heart ventricular lysates (HVL) and crude mitochondria (CM), and that it is practically free of non-mitochondrial protein markers with the largest contaminant being from the nuclear envelope (Lamin B1, 0.06±0.03 in a scale 0–1; n = 3), and the least being from the T-tubules (α1C, 0.001±0.0003; n = 3). Interestingly, SFKs signals were also low in M3 (0.02±0.004; n = 3).

Next, M1, M2 and M3 fractions were further tested by measuring membrane potential using the ratio of mitotracker and mitochondrial marker. The Viability of M1, M2 and M3 fractions was assessed using three complementary parameters:

- Production of reactive oxygen species (ROS),
- Western blot analysis of organelle protein markers; and
- Colocalization with mitotracker which labels mitochondria with intact membrane potential.

Western blots in Fig. 1C show that M3 fraction produced robust signals for VDAC1 and Cox2 but no significant signals for Cadherin, GM130, Lamin B1, GRP78 BIP, α1C, and Src. Quantification of normalized signals (to Ponceau S signals) demonstrates that M3 fraction is significantly enriched in VDAC1 and Cox2 with respect to heart ventricular lysates (HVL) and crude mitochondria (CM), and that it is practically free of non-mitochondrial protein markers with the largest contaminant being from the nuclear envelope (Lamin B1, 0.06±0.03 in a scale 0–1; n = 3), and the least being from the T-tubules (α1C, 0.001±0.0003; n = 3). Interestingly, SFKs signals were also low in M3 (0.02±0.004; n = 3).

### Visualization of mitochondrial proteins by confocal microscopy

Purified M3 fraction mitochondria were loaded with mitotracker and counter-stained for Cox2, VDAC1 and Lamin B1 proteins. A–B. Anti-Cox2 monoclonal antibody and mitotracker signals. C. Overlay of A and B shows high correlation between mitotracker and Cox2 signals. D–E. Anti-VDAC1 monoclonal antibody and mitotracker signals. F. Overlay of D and E shows high signal correlation of mitotracker with VDAC1. A–F. Squares in A–F are shown at higher magnification for better examination. G–H. Lamin B1 (nuclear envelope marker, negative control) and mitotracker signals. I. Overlay of G and H shows high signal correlation of mitotracker with Lamin B1. J–K. Mitochondria were loaded with mitotracker and labeled with secondary antibody alone. Secondary antibody (anti-mouse) showed no specific signals (J), while mitotracker labeling was robust (K) confirming the presence of mitochondria. L. Protein Proximity Index (Wu et al., 2010) of mitotracker towards Cox2, VDAC1, Lamin B1, or secondary antibody alone (Sec Ab) (black bars) and vice versa (open bars) (n = 5 mitochondria preparations). Values are given in the text. X is the protein being measured with respect to its mitotracker proximity. Confocal images were acquired at 0.0575 μm/pixel.

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**Fig. 2.** Visualization of mitochondrial proteins by confocal microscopy. Purified M3 fraction mitochondria were loaded with mitotracker and counter-stained for Cox2, VDAC1 and Lamin B1 proteins. A–B. Anti-Cox2 monoclonal antibody and mitotracker signals. C. Overlay of A and B shows high correlation between mitotracker and Cox2 signals. D–E. Anti-VDAC1 monoclonal antibody and mitotracker signals. F. Overlay of D and E shows high signal correlation of mitotracker with VDAC1. A–F. Squares in A–F are shown at higher magnification for better examination. G–H. Lamin B1 (nuclear envelope marker, negative control) and mitotracker signals. I. Overlay of G and H shows high signal correlation of mitotracker with Lamin B1. J–K. Mitochondria were loaded with mitotracker and labeled with secondary antibody alone. Secondary antibody (anti-mouse) showed no specific signals (J), while mitotracker labeling was robust (K) confirming the presence of mitochondria. L. Protein Proximity Index (Wu et al., 2010) of mitotracker towards Cox2, VDAC1, Lamin B1, or secondary antibody alone (Sec Ab) (black bars) and vice versa (open bars) (n = 5 mitochondria preparations). Values are given in the text. X is the protein being measured with respect to its mitotracker proximity. Confocal images were acquired at 0.0575 μm/pixel.
Although M1 and M2 fractions contain similar amounts of VDAC1 and Cox2 than M3, indicating the presence of mitochondria in these fractions, the signals from other organelle markers in M1 and M2 were still significant. Electron microscopy of M1 and M2 fractions showed very few intact mitochondria when compared to M3 (n = 3) suggesting that the strong signals of VDAC1 and Cox2 in M1 and M2 originate from damaged mitochondria. In summary, the degree of purity as reflected by the absence of extramitochondrial markers was optimal in M3 fraction. These results are consistent with the functional data in Fig. 1B where M3 had the best performance.

3.2. Confocal immunofluorescence of purified mitochondria

To further validate the suitability of M3 fraction for nanoscopy studies, we colabeled this mitochondria fraction with antibodies against Cox2 (Fig. 2A, A'), VDAC1 (Fig. 2D, D'), and Lamin B1 (Fig. 2G, G') together with mitotracker, which was efficiently uptaken by purified mitochondria (Fig. 2B, B'; E, E', H, H'). Both Cox2 and VDAC1 immunofluorescence highly coincided with mitotracker signals (Fig. 2C, C' and Fig. 2F, F') supporting the mitochondrial nature of both types of signals. As expected from the Western blot analysis in Fig. 1C, D the immunofluorescence signals of the nuclear envelope marker Lamin B1 were scarce and showed no colocalization with mitotracker (Fig. 2I, I'). Thus, organelle labeling by specific antibodies represents a further purification step as it can readily discern between scarce contaminant proteins and mitochondria-associated proteins. Controls with secondary antibody in the absence of primary antibody showed no labeling (Fig. 2J vs. mitotracker control in Fig. 2K) and VDAC1 antibody signal could be blocked with recombinant VDAC1 protein (Fig. S1).

We next quantified the colocalization index of Cox2, VDAC1, Lamin B1 and secondary antibodies with mitotracker using the protein proximity index (PPI) method (Wu et al., 2010) (Fig. 2L). This method isolates specific colocalization signals from those produced by image heterogeneity and non-specific colocalization as those arising from the antibody background signal. The PPI values were: mitotracker towards Cox2 = 0.83 ± 0.05; Cox2 towards mitotracker = 0.66 ± 0.04; mitotracker towards VDAC1 = 0.79 ± 0.03; and VDAC1 towards mitotracker = 0.61 ± 0.03 (n = 5). These data indicate that about 80% of mitochondria labeled by mitotracker are also labeled by Cox2 and VDAC1; whereas, only about 65% of mitochondria labeled with Cox2 and VDAC1 were also labeled by mitotracker. The latter suggests that ~35% of mitochondria could not uptake or maintain mitotracker although they could be labeled for Cox2 and VDAC1. In contrast to Cox2 and VDAC1, PPI of Lamin B1 and secondary antibody alone (Sec Ab) with mitotracker were very low and similar in both directions. Values were: mitotracker towards Lamin B1 = 0.14 ± 0.01; Lamin B1 towards mitotracker = 0.15 ± 0.01; mitotracker towards Sec Ab = 0.12 ± 0.02; Sec Ab towards mitotracker = 0.14 ± 0.03 (n = 5). These results demonstrate as a reference that Cox2 and VDAC1 but not Lamin B1 are located in mitochondria from M3 fraction. Because mitotracker is concentrated by functional mitochondria with an intact membrane potential, the results also confirm that M3 fraction contains viable mitochondria.

We have also tested the presence of α1C and SKFs proteins in purified mitochondria from M3 fraction by colabeling with polyclonal antibodies against these proteins and with mitotracker (Fig. S2). As expected for M3 fraction, mitochondria were readily labeled with mitotracker; however, α1C produced scarce signals that did not coincide with mitotracker, while Src signals were undetectable and similar to secondary antibody alone. To validate the labeling properties of both antibodies, we labeled isolated murine cardiomyocytes under identical conditions and obtained clear labeling with the same antibodies. These results are consistent with the Western blot analysis showing that α1C and Src proteins are negligible in the purified M3 mitochondrial fraction isolated from murine heart.

3.3. Localization of mitochondrial proteins with STED

The diameter of mitochondria in M3 fraction is around 200–1000 nm, while the maximum resolution of conventional confocal microscopy (limited by the diffraction limit of the light) is ~200 nm in the focal plane and ~500 nm along the optic axis. Therefore, using traditional confocal microscopy, it is near impossible to differentiate protein clusters from the blurred mitochondrial image. However, with STED nanoscopy protein clusters can be readily identified. For comparison, we have acquired both confocal (Fig. 3A, H) and STED (Fig. 3B, E, I and L) images of the same preparation. The overlays (Fig. 3C, D, J and K) contrast the dramatic increase in resolution with STED nanoscopy, allowing the visualization of clusters of VDAC1 and Cox2 proteins. Cox2 showed a diffused cluster distribution (Fig. 3B and E), whereas VDAC1 had discreet clusters (Fig. 3I and L). Fig. 3F and M shows examples of resolution measurements using a line scan (corresponding lines in Fig. 3E and L) through clusters and measuring the FWHM of individual peaks (arrows). Similar measurements of 500 clusters from 50 isolated mitochondria (10 clusters per mitochondrion from three independent experiments) were used to construct the cluster size frequency histograms. Histograms were fitted to multiple Gaussian functions (red lines) revealing that in purified cardiac mitochondria Cox2 clusters are distributed mostly in clusters of ~33 nm (Fig. 3G), while VDAC1 clusters had in addition larger size populations of ~55 and 83 nm (Fig. 3N). As acquisition was at ~10 nm/pixel, peaks at ~20 nm in Fig. 3G and N may represent noise measurements or smaller cluster sizes that can not be resolved at the limit of the resolution.

4. Discussion and conclusions

We have been able to visualize protein clusters of VDAC1 and Cox2 in cardiac mitochondria using a custom-made STED fluorescence microscope and distinguish specific cluster populations of the proteins examined.

The visual power of this method (hundreds of mitochondria can be visualized and analyzed in a single experiment) together with the usage of specific antibodies should allow unambiguous demonstration about the presence of proteins in mitochondria. In the M3 purified mitochondria fraction, VDAC1 and Cox2 were readily detected by Western blot and super resolution fluorescence microscopy. In contrast, neither SKFs nor the α1C subunit of the L-type Ca2+ channel gave significant signals (Fig. 1 and Fig. S2). These results are consistent with the LC/MS/MS analysis of Percoll (30%/60% gradient) purified murine cardiac mitochondria, where neither α1C nor SKFs peptides were detected (Zhang et al., 2008). However, the existence of SKFs in rat heart mitochondria has been supported by pharmacological and biochemical studies where Western blots of Percoll (25%)-separated mitochondria showed robust signals for Src (Feng et al., 2008, 2010). Src signals were also robust in the M1 and M2 mitochondrial fractions but were basically absent in the purest and functional M3 fraction using both Western blot and confocal microscopy. It is possible that the concentration of Percoll influences the purification process or that species variation (rat vs. mouse) or perhaps mitochondria subpopulations may account for the differences.

Previous qualitative evaluation of Flag-tagged VDAC3 expressed in U2OS cells, gave similar large clusters (40 to 90 nm) (Neumann et al., 2010) in comparison to VDAC1 (~55 and 83 nm) in our studies. However, we identified additional smaller size clusters for VDAC1 (~30 nm). The usage of isolated mitochondria instead of cells may avoid out of focus fluorescence and contribute to increased resolution in our measurements.

4.1. Conclusions

Using STED microscopy and purified murine cardiac mitochondria, we have shown that native VDAC1 and Cox2 are distinctly organized in cardiac mitochondria forming clusters of different sizes.
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