Role of Dynamin Related Protein 1 (Drp1)-Mediated Mitochondrial Fission in Oxygen-Sensing and Constriction of the Ductus Arteriosus

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ABSTRACT

**Rationale:** Closure of the ductus arteriosus (DA) is essential for the transition from fetal to neonatal patterns of circulation. Initial PO$_2$-dependent vasoconstriction causes functional DA closure within minutes. Within days a fibrogenic, proliferative mechanism causes anatomical closure. Though modulated by endothelial-derived vasodilators and constrictors, O$_2$-sensing is intrinsic to ductal smooth muscle cells (DASMC) and oxygen-induced DA constriction persists in the absence of endothelium, endothelin and cyclooxygenase mediators. O$_2$ increases mitochondrial-derived H$_2$O$_2$ (mitoROS), which constricts DASMC by raising intracellular calcium and activating rho kinase. However, the mechanism by which oxygen changes mitochondrial function is unknown.

**Objective:** Determine whether mitochondrial fission is crucial for O$_2$-induced DA constriction and closure.

**Methods and Results:** Using DA harvested from 30 term infants during correction of congenital heart disease, as well as DA from term rabbits, we demonstrate that mitochondrial fission is crucial for O$_2$-induced constriction and closure. O$_2$ rapidly (<5 minutes) causes mitochondrial fission by a cyclin-dependent kinase-mediated phosphorylation of dynamin-related protein 1 (Drp1) at serine 616. Fission triggers a metabolic shift in the DASMC that activates pyruvate dehydrogenase and increases mitochondrial H$_2$O$_2$ production. Subsequently fission increases complex I activity. Mitochondrial-targeted catalase overexpression eliminates PO$_2$-induced increases in mitoROS and cytosolic calcium. The small-molecule Drp1 inhibitor, Mdivi-1, and siDRP1 yield concordant results, inhibiting O$_2$-induced constriction (without altering the response to phenylephrine or KCl) and preventing O$_2$-induced increases in oxidative metabolism, cytosolic calcium and DASMC proliferation. Prolonged Drp1 inhibition reduces DA closure in a tissue culture model.

**Conclusions:** Mitochondrial fission is an obligatory, early step in mammalian O$_2$-sensing and offers a promising target for modulating DA patency.

**Keywords:** Oxygen sensing, persistent patent ductus arteriosus, mitochondrial division inhibitor-1 (Mdivi-1), mitochondrial-targeted photoactivatable green fluorescent protein, mitochondrial fission

**Non-standard Abbreviations:**

- DA: ductus arteriosus
- DASMC: ductus arteriosus smooth muscle cell
- Drp1: dynamin-related protein 1
- Drp1 p-Ser-616: dynamin-related protein 1 (phosphorylated at serine 616)
- ECAR: extracellular acidification rate
- H$_2$O$_2$: hydrogen peroxide
- Mdivi-1: mitochondrial division inhibitor 1
- MFC: mitochondrial fragmentation count
- mitoCAT: mitochondrial-targeted catalase
- mitoGFP: mitochondrial-targeted green fluorescent protein
- mito-PA-GFP: mitochondrial-targeted photoactivatable green fluorescent protein
- mito-DsRED: mitochondrial targeted red fluorescent protein
- mitoROS: mitochondrial-derived reactive oxygen species
- MNF: mitochondrial Networking Factor
- OCR: oxygen consumption rate
- PDH: pyruvate dehydrogenase

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INTRODUCTION

Closure of the ductus arteriosus (DA) at birth diverts blood from the right ventricle through the newly expanded lungs, an essential step in the transition from the fetal to neonatal circulation. Functional DA closure due to vasoconstriction begins within minutes of birth and is required for subsequent anatomical closure1-2. Anatomical closure occurs within days by a process that involves breakdown of the internal elastic lamina, proliferation of cells in the media3 and fibrosis. Persistent patency of the DA (PDA) is prevalent in low birth-weight, preterm infants4, resulting in pulmonary congestion and failure to thrive. Prostaglandin H synthase inhibitors such as ibuprofen5 can close the DA in ~70% of infants; however, morbidities (e.g. oliguria) are not uncommon6. Failure of medical therapy often leads to invasive closure7. Conversely, maintenance of DA patency, by prostaglandin E infusion8, is sometimes required as a bridge to corrective congenital heart surgery. This too can cause toxicity (notably apnea and morbidity related to central venous access).

Oxygen (O2) is the primary stimulus for DA constriction. O2-induced DA vasoconstriction is reinforced by endothelin9 and inhibited by vasodilator prostanoids10 and nitric oxide11. Coceani et al proposed that DA constriction to O2 results from the loss of the vasodilator prostaglandin E2 combined with an increase in levels/activity of the potent vasoconstrictor, endothelin-112. While endothelin can constrict the DA and endothelin levels gradually increase with oxygenation, endothelin is not essential for DA closure/constriction in animal models13, 14. Moreover, simultaneous inhibition of endothelin receptors and endothelin converting enzyme (at doses that inhibit constriction to exogenous endothelin and reduce endothelin levels) does not inhibit acute constriction of the human DA to oxygen15. Our group has proposed an alternative mechanism for the O2-induced DA16 in which the core mechanism of O2-induced constriction reflects changes in the function of a mitochondrial-based redox sensor that triggers a cascade of reinforcing constrictor mechanisms that are intrinsic to the DA smooth muscle cell (DASMC)15, 17-19. Within 5 minutes of increasing PO2, mitochondria-derived reactive oxygen species (mitoROS), notably H2O2, increase and act as diffusible signaling molecules that by inhibiting O2-sensitive potassium channels, lead to DASMC depolarization, activation of the L-type calcium channel, calcium influx and initiation of vasoconstriction. Subsequently, mitoROS activate rho kinase, which sustains the O2-constriction20-22. Some calcium also enters the DASMC via store-operated channels or transient receptor potential channels22. MitoROS originate from the electron transport chain (ETC) complexes I or III and inhibition of these complexes (by rotenone or antimycin A) selectively inhibits DA constriction to O2, without altering responses to other vasoconstrictors, such as phenylephrine or KCl20.

Anatomical closure of the DA follows, and may depend on, functional closure. There are several theories for the basis of the DA closure, including that hypoxic cell death occurs in cells in the wall of the full-term ductus due to reduced blood flow in vasa vasorum of the thick walled DA 23. This mechanism may not be active in all species and was found not to be crucial in mice24.

The precise mechanism by which O2 increases mitoROS and the relevance of mitochondrial O2-sensing to functional and anatomical DA closure were the subject of this study. We observed that physiologic increases in PO2 rapidly fragment the DASMC’s mitochondrial network, suggesting that structural changes in mitochondria are a key early step in O2-sensing. O2 induces fission by stimulating phosphorylation of the guanosine triphosphatase (GTPase), dynamin related protein 1 (Drp1), at serine 616. Fission activates pyruvate dehydrogenase (PDH), increases oxygen consumption rate (OCR) and initiates redox based O2-sensing. Drp1 inhibition selectively blocks oxygen-mediated DA constriction and, if sustained, prevents DA closure in a tissue culture model. Thus, mitochondrial fission is an obligatory early step in O2-sensing that is crucial to DA constriction and closure.
METHODS

Human ductus.
DA were obtained from infants with various forms of congenital heart disease at the time of surgical correction either at the University of Nebraska Children’s Hospital or the University of Chicago under protocols approved by the local IRB of both institutions. Demographics are provided in the supplemental table I. None of the patients had a persistently patent DA, defined as continued patency in infants older than 3 months. The 2 oldest patients in this cohort had associated congenital heart disease (tetralogy of Fallot and coarctation of the aorta).

Animal studies.
The University of Chicago Animal Care Committee approved all protocols.

Statistics.
Values are stated as mean ± SEM. When applicable, normality was confirmed with a Kolmogorov-Smirnov test. Inter-group differences were assessed by Student’s t-tests (unpaired or paired) or ANOVA (simple or repeated-measures), as appropriate. Post hoc analysis was performed using Tukey’s test. A p<0.05 was considered statistically significant.

Term human DA ring physiology.
The DA from 28 infants were immediately placed in sterile, iced, hypoxic M231 culture media and shipped overnight by FedEx to the University of Chicago. DA from two infants originated at the University of Chicago. Of 30 infants, 25 (83%) had been on prostaglandin prior to removal of DA at surgery. The DA was cut into rings and suspended in hypoxic Earle’s Balanced Salt Solution (PO2~40mmHg) at an experimentally determined optimal resting tension of 1g, as previously described. 10μm meclofenamate and 100μm L-NAME were present throughout all experiments. Mdivi-1 (20μM) was given 30 minutes before normoxic challenges (PO2~120mmHg).

Term rabbit DA ring physiology.
Pregnant New Zealand White rabbits at 30-31 days of gestation (term =31days) were anesthetized with ketamine 75 mg and xylazine 20 mg, intramuscularly and 50 mg pentobarbital, intravenously. The pups were delivered by cesarean section and a midline sternotomy was performed, before initiation of respiration. The heart, lungs, and great vessels were excised en bloc and placed in deoxygenated Earle’s solution. The DA was carefully dissected free from adventitia under a dissecting microscope and severed distal to the takeoff of the left pulmonary artery and proximal to the insertion into the descending thoracic aorta. DA physiology was studied as previously described. 10μm meclofenamate and 100μm L-NAME were present throughout all experiments.

DASMC culture (human and rabbit).
Human DASMCs from 14 infants (10 Male, 4 Female) were used in this part of the study. 4 term fetal pups from 4 different pregnant New Zealand White rabbits were used to isolate and culture rabbit DASMCs. The identity of DASMCs was confirmed morphologically and by immunohistochemical demonstration of the presence of SM α-actin and the absence of von Willebrand’s factor. A primary culture of DASMC was established and then the cells were harvested with trypsin, frozen in Freezing Media (10% DMSO, 20% FBS and 70% M231) and stored in Liquid Nitrogen for later use. DASMC were used within the first 5 passages in culture. DASMC were scrupulously maintained in hypoxia (PO2 40 mmHg, pH 7.35-7.45, PCO2 30-40mmHg) using an environmentally controlled, Tri-Gas CO2 incubator (Thermo Fisher Scientific), until the protocol called for exposure to normoxia.
**DA tissue culture (human and rabbit).**
The DA rings were cultured in M231 media with 5% smooth muscle growth media in normoxic conditions (PO$_2$ 120 mmHg, pH 7.35-7.45, PCO$_2$ 30-40 mmHg). The DAs were allowed to float in the culture medium. Once Mdivi-1 or vehicle (DMSO) was added to the media it was left in place for the duration of the experiment.

**Exposure of hypoxic human DASMC to acute normoxia.**
The experiment was performed in a chamber flushed with nitrogen. Inside the hypoxic chamber, two small glass containers (500 ml) were filled with Earle’s solution (37°C). They were bubbled either with a hypoxic (0% O$_2$, 5% CO$_2$; PO$_2$=28±2 mmHg) or normoxic (20% O$_2$, 5% CO$_2$; PO$_2$=134±2 mmHg) gas mixture. DASMCs were immersed in the hypoxic solution for 30 minutes and then either maintained in hypoxia for 5 and 20 minutes longer or quickly switched to the normoxia solution for 5 or 20 minutes. At the end of the experiment, cells were flash frozen in liquid nitrogen and later lysed at 4°C under hypoxic conditions using the PhosphoSafe™ Extraction Reagent (Calbiochem) and a Phosphatase Inhibitor Cocktail (Calbiochem, 1:100).

**Imaging Mitochondrial Network Fragmentation and Connectivity.**

**Mitochondrial Fragmentation Counts (MFC).**
Mitochondrial network fragmentation was assessed as previously described\textsuperscript{27, 28}. The MFC quantifies discrete mitochondrial particles and greater fission results in higher MFC values. To calculate MFC, cells were loaded with tetramethylrhodamine (TMRM) (50 nM, 20 min at 37°C, Molecular Probes, Eugene, OR) and imaged with the Zeiss 510 META confocal laser scanning microscope using an alpha-Plan Apo 100x/1.46NA objective with 3x digital zoom (excitation at 561 nm, emission recorded above 575 nm). Acquired images were background subtracted, filtered (median), thresholded and binarized to identify mitochondrial segments using ImageJ (NIH, Bethesda, MD). Continuous mitochondrial structures were counted with a particle counting subroutine and the number was normalized to the total mitochondrial area (in pixels) to obtain the MFC for each imaged cell (n>25 randomly selected cells/group).

**Mitochondrial Networking Factor (MNF).**
To quantify functional networking of the mitochondria, cells were doubly transfected with mito-PA-GFP\textsuperscript{29} and mito-Ds-Red\textsuperscript{30} using Fugene HD transfection reagent (Roche, Indianapolis, IN), as previously described\textsuperscript{27, 28}. Imaging experiments were performed 24-96 hours following the transfection of the cells using a Zeiss 510 META confocal laser scanning microscope equipped with an environmental chamber, to ensure that experiments were conducted at 37°C and at physiologic pH, PCO$_2$ and either normoxic or hypoxic PO$_2$. A 100X alpha Plan-Apochromat (1.46NA) objective and 2X zoom were used to image single cells. Sequential images of fluorescent mitochondria were collected every 17.6 seconds. Mito-Ds-Red was excited at 561 nm and the emission was recorded using 565 nm dichroic and 575 nm long pass filters. After acquiring 3 control images, localized photoactivation of mito-PA-GFP was achieved by 10 passes of the 2-photon laser (at 700 nm) within a 23 µm$^2$ activation area. Mito-PA-GFP was excited at 488 nm and the images were acquired at 500-550 nm. Mito-Ds-Red permits visualization of the entire mitochondrial network, while the mito-PA-GFP enables real-time imaging of GFP diffusion from the discrete photoactivation region through the mitochondrial reticulum. MNF quantifies the spread of mito-PA-GFP after photoactivation and thus is proportional to the extent of network fusion, resulting in low values in states of fission\textsuperscript{27, 28}.

**Transmission Electron Microscopy.**
Two human ductus were used for this experiment. Each ductus was cut into 2 to 4 rings and treated with either hypoxia (20 minutes) or normoxia (20 minutes). These DA were then immediately fixed and processed for electron microscopy. After osmium tetroxide treatment, samples were stained with 1% uranyl acetate and embedded in SPURR (Electron Microscopy Sciences, Hatfield, PA). Images were
collected using a scanning transmission electron microscope at 300 KV (Tecnai F30; FEI, Hillsboro, Oregon) with a high-performance Gatan CCD camera (Pleasanton, CA).

**O2-consumption and metabolic measurements.**

Simultaneous measurements of oxygen consumption rate (OCR) and extracellular acidification rate (ECAR, a measure of lactate production) were performed on a XF24 extracellular flux analyzer (Seahorse Bioscience, North Billerica, MA). DASMC were plated on a XF24 cell culture microplate. Measurements were made the next day, after equilibration in XF assay medium supplemented with 4.5g/l glucose. To compare metabolic function in hypoxia versus normoxia, the entire XF24 extracellular flux analyzer was sealed in a plastic container which was flushed with either nitrogen (hypoxia) or room air (normoxia). The oxygen level within the chamber was monitored by an oxygen sensor and by the XF24’s PO2 sensor. OCR was measured at baseline and upon sequential challenge with 1 μM oligomycin (to inhibit ATP synthase), 2 μM FCCP (to uncouple the mitochondria and demonstrate maximal OCR) and 5 μM antimycin A (to inhibit the mitochondrial component of respiration).

**qRT-PCR and immunofluorescence.**

These techniques were performed as previously described27.

**Immunoblotting and immunofluorescence.**

Immunoblotting was performed on 30 μg of protein from human DASMC. The primary antibodies used were anti-Drp1 (Santa Cruz Biotechnology, Santa Cruz, CA), anti-phospho Ser616 Drp1, anti-phospho Ser637 Drp1 (Cell Signaling Technology, Boston, MA), anti-Actin (Millipore, Billerica, MA) and anti-translocase of outer mitochondrial membrane 20 (TOM20, Santa Cruz Biotechnology, Santa Cruz, CA). The mitochondria-enriched fraction was isolated using a mitochondria/cytosol fractionation kit (Abcam, Cambridge, MA).

**Measurement of cytosolic calcium concentrations ([Ca2+]cyt).**

DASMC were plated on coverslips and loaded with Fura-2 AM (3 μmol/L for 30 minutes at 37°C, Invitrogen, Carlsbad, CA). The ratio of 510 nm fluorescence signals elicited at 380 and 340nm stimulation was calculated to quantify [Ca2+]cyt, as previously described27. Compartmental calcium stores were assessed by measuring changes in [Ca2+]cyt when the sarcoplasmic reticulum pool was released by cyclopiazonic acid (CPA), or the mitochondrial pool was released by carbonyl cyanide p-trifluoromethoxyphenylhydrazone (FCCP).

**Small interfering RNA (siRNA).**

To achieve specific molecular knockdown of Drp1 or the putative Drp1 phosphorylating kinase, PKCδ, we used siRNA, targeting Drp127 or PKCδ (IDT, San Jose, CA). Human DASMC were grown to ~80% confluence and then transfected using Lipofectamine RNAiMAX (Invitrogen, Carlsbad, CA) and 10 nM of a validated Silencer Select siRNAs or a scrambled siRNA control (IDT, San Jose, CA). After 6 hours, normal culture medium was applied and gene knockdown was assessed using qRT-PCR (48 hours) and immunoblot (72 hours).

**PDH enzyme activity.**

PDH enzyme activity assay was performed on freshly isolated DA proteins using a PDH Enzyme Activity Dipstick Assay Kit following the manufacturer’s instruction (MitoSciences-Abcam, ab109882, Eugene Oregon). This technique uses anti-PDH antibody immobilized on a capture line section of the dipstick. The native (active) PDH is immunocaptured and activity is measured by coupling PDH-dependent production of NADH to the reduction of nitroblue tetrazolium. PDH activity results in an insoluble blue precipitate at the capture line, the intensity of which is proportional to PDH activity.

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ETC complex I activity.
Protein isolated from DASMC analyzed using the Mitoscience Complex I Enzyme Activity Dipstick Assay Kit (ab109720). Complex I from the DASMC is captured in its native (active) form on the dipstick which is then immersed in a Complex I activity buffer solution containing substrate (NADH) and an electron recipient (nitroblue tetrazolium). The immunocaptured Complex I oxidizes NADH and the electron transfer reduces nitroblue tetrazolium forming a blue precipitate at the capture line that is proportional to complex activity.

Mitochondrial targeted increases in catalase activity.
Human DASMC in culture (50% confluence) were infected with serotype 5 adenovirus carrying mitochondrial-targeted catalase (Ad-mitoCAT, 100 MOI) or an empty virus (100 MOI). Catalase activity was quantified after 3-5 days using a colorimetric activity assay, following the manufacturer's instructions (Biovision, Milpitas, California).

DASMC proliferation and apoptosis assays.
Proliferation was quantified using the EdU kit according the manufacturer's instructions (Invitrogen), as previously described27. Apoptosis was measured using an FITC annexin V apoptosis detection kit, as previously described (BD Pharmingen, San Diego, CA)27.

RESULTS

Mitochondrial fission is required for O₂-constriction (Fig 1 and 2).

The small molecule inhibitor of Drp1, mitochondrial division inhibitor (Mdivi-1)31, prevents and reverses O₂-induced constriction in rabbit and human DA. Mdivi-1 does this selectively, without altering constriction to agents that act by depolarizing the DASMC (KCl) or releasing intracellular calcium (phenylephrine). This suggests that Mdivi-1 interferes with a mitochondrial oxygen-sensor, rather than a downstream constrictor mechanism.

Oxygenation rapidly causes mitochondrial fission in DASMC.

Exposure of hypoxic DASMC to O₂ at tensions similar to those experienced at birth (increasing PO₂ from 40 to 120 mmHg), fragments the DASMC’s mitochondrial network. This was quantified as an increase in the MFC and a reduction in MNF that occurred within 10 minutes of increasing PO₂ (Fig 3A-C). Mdivi-1, prevents this O₂-induced fission (Fig 3C), offering pharmacologic evidence that it is mediated by Drp1. To ensure that fission was not unique to DASMC in culture, we divided surgically resected human DAs into two pieces, one maintained in hypoxia (40 mmHg) and the other exposed to a PO₂ of 120 mmHg for 15 minutes. Transmission electron microscopy (TEM) confirms that O₂ causes mitochondrial fragmentation in DASMC within the media of freshly isolated DA rings (Fig 3D).

Expression profiling of fission and fusion mediators (Fig 4).

Mitochondrial fragmentation could reflect impaired fusion, enhanced fission or both. We constructed an mRNA expression profile for mediators of fusion (mitofusin-1, mitofusin-2 and optic atrophy protein) and fission (Drp1 and fission protein 1) at time points from 0-72 hours after exposure to normoxia (Fig 4A). Drp1 expression began to increase within 1-hour of O₂ exposure and by 24-hours Drp1 was the only fission or fusion mediator whose expression had changed (Fig 4A). After 72 hours of O₂, Drp1 expression increased ~8-fold and mitofusin-2 expression also increased.
Changes in Drp1 phosphorylation (Fig 4).

DA constriction begins within 5 minutes of an increase in PO$_2$, which is too rapid to reflect transcriptional regulation of Drp1. Consequently we assessed whether O$_2$ changes Drp1 phosphorylation at serine 616, a post-translational modification known to cause Drp1 to translocate to the mitochondria and form multimers that circumferentially constrict the mitochondrion and initiate fission. As expected, brief O$_2$-exposure (<10 minutes) did not change total Drp1 expression; however, it increased expression of activated DRP1 (Drp1-p-Ser-616, Fig 4C) in the mitochondrial protein fraction. O$_2$ also promoted DRP-1 activation by decreasing expression of the inhibitory phosphorylation form, p-Ser-637 (supplemental Fig IA).

Identification of kinases that phosphorylate Drp1 (Fig 4C).

Next we examined the identity of the kinase(s) that activate/phosphorylate Drp1. Several kinases can cause Drp1 phosphorylation, including cyclin-dependent kinases, calcium/calmodulin-dependent protein kinase, and protein kinase C. To identify the kinase(s) that mediate O$_2$-induced increase in Drp1 p-Ser-616, DASMC were exposed to oxygen in the presence of roscovitine, an inhibitor of cyclin-dependent kinase-1, -2 and -5, or KN93, a calcium/calmodulin-dependent protein kinase inhibitor or siPKC$_\delta$. Roscovitine and KN93 reduced Drp1 p-Ser-616 (Fig 4D); whereas, siPKC$_\delta$ had no effect (supplemental Fig I). All siRNA species were confirmed to achieve >80% knockdown of their targets. Thus cyclin-dependent kinase and calcium/calmodulin-dependent protein kinase (linked to cell proliferation and [Ca$^{2+}$]cyt, respectively) contribute to O$_2$-induced activation of Drp1 in DASMC.

Fission increases O$_2$-consumption and oxidative metabolism (Fig 5).

We next explored how this Drp1-mediated fission elicits vasoconstriction. Because increases in mitochondrial respiration are thought to mediate O$_2$-dependent increases in mitoROS in the DA, we investigated the effects of fission on mitochondrial metabolism. To do this we measured OCR and glycolysis (ECAR) in primary cultures of DASMC. To modulate PO$_2$ the Seahorse analyzer itself was housed in a hypoxic chamber (Fig 5). DASMC maintained in hypoxia were glycolytic, with low OCR and high ECAR (Fig 5A-B). Elevating PO$_2$ increased OCR and reduced ECAR. This metabolic change resulted from rapid, PO$_2$-dependent activation of PDH, the key regulator of mitochondrial glucose oxidation (Fig 5B). The obligatory role for fission in this oxidative metabolic shift is demonstrated by the finding that Drp1 inhibition (whether achieved by Mdivi-1 or siDrp1) inhibits O$_2$-induced increases in both OCR and PDH activity. As would be predicted, Mdivi-1 inhibits the rise in H$_2$O$_2$ production in response to increases in PO$_2$ (Fig 5A). The finding that PO$_2$-induced PDH activation is inhibited by Mdivi-1 demonstrates that fission is required for the burst of oxidative metabolism that serves as the redox signal of “normoxia” in the DA. This metabolic shift increases mitoROS, presumably by increasing electron flux through the ETC. Consistent with this, more sustained incubation in O$_2$ (2 days) also increases ETC mega-Complex I activity (Fig 5B).

Inhibition of Drp1 lowers cytosolic calcium (Fig 6).

Elevation of [Ca$^{2+}$]cyt is the accepted final common pathway for activation of the contractile apparatus in DASMC. Consequently, we examined the effects of Drp1 inhibition on O$_2$-induced elevation of [Ca$^{2+}$]cyt. Mdivi-1 impaired O$_2$-induced increases in cytosolic calcium in rabbit DASMC (Fig 6A). Mdivi-1 reduced the calcium released from the sarcoplasmic reticulum and mitochondria by CPA and FCCP, respectively (Fig 6B and 6C). Mdivi-1 is thought to be a selective Drp1 inhibitor; however, to confirm the putative mechanism, we used a highly specific molecular strategy (siDrp1) and observed
concordant results. Like Mdivi-1, siDrp1 prevented O₂-induced increases in [Ca²⁺]cyt without altering the response to KCl (Fig 6D). The trigger for the rise in calcium in DASMC is thought to be increasing H₂O₂. Supporting this, exogenous H₂O₂ increases [Ca²⁺]cyt in DASMC (Fig 6E).

Consistent with the signaling role of H₂O₂ in DASMC, prior studies had shown that augmenting intracellular catalase attenuated O₂-sensing in DASMC, we established the compartmental specificity of the signaling H₂O₂ pool using Ad-mitoCAT. Mitochondria-specific overexpression of catalase inhibited O₂-induced elevation of cytosolic calcium (Fig 6E-F and supplemental Fig III). Thus, fission triggers O₂-sensitive H₂O₂ generation by the DASMC’s mitochondria.

**Prolonged Drp1 inhibition prevents DASMC proliferation and DA closure (Fig 7 & 8).**

We next assessed whether prolonged inhibition of fission would prevent the anatomical closure of the DA. In a tissue culture model, we demonstrated that a PO₂ of 140 mmHg caused DA occlusion after 3 days (rabbit) and 6-14 days (human) (Fig 7 A-B). Mdivi-1, but not vehicle, prevented closure of the DA. Because DA closure involves cellular proliferation and because prior experience shows Mdivi-1 can prevent SMC proliferation, we established the compartmental specificity of the signaling H₂O₂ pool using Ad-mitoCAT. Mitochondria-specific overexpression of catalase inhibited O₂-induced elevation of cytosolic calcium (Fig 6E-F and supplemental Fig III). Thus, fission triggers O₂-sensitive H₂O₂ generation by the DASMC’s mitochondria.

The antiproliferative effects of Mdivi-1 also occurred in DA rings in tissue culture, evident as a reduction in the number of PCNA-positive DASMC in rings cultured in the presence of Mdivi-1 (Fig 8A). Mdivi-1 significantly prevented fibrotic remodeling of the DA, which normally occurs with closure (Fig 8B-C). These findings, together with the time dependent increase in Drp1 expression suggest that fission is important to DA closure, as well as to the rapid initiation of O₂-induced constriction.

**DISCUSSION**

This study identified a critical role for Drp1-mediated mitochondrial fission in acute constriction of the DA to O₂ and also strongly suggests this mechanism participates in the DA’s subsequent anatomical closure. Mitochondrial fission increases oxidative metabolism in DASMC. The resulting changes in redox signaling (particularly increases in mitochondrial-derived H₂O₂) elevate [Ca²⁺]cyt and initiate vasoconstriction. Drp1 activation is rapid, occurring with onset within 5 minutes, and reflects posttranslational modification. With normoxia cyclin-dependent kinase and calcium/calmodulin-dependent protein kinase increase Drp1 phosphorylation, increasing the ratio of p-Ser-616/p-Ser-637 and consequently favoring fission. These kinases link mitochondrial morphology to the regulation of cell proliferation (cyclin-dependent kinase) and cytosolic calcium (calcium/calmodulin-dependent protein kinase). The concordant findings in rabbits and humans speak to the conservation of this mechanism. The basis for the fission-induced redox signal is demonstrated to lie in an alteration in mitochondrial metabolism. Fission in DASMC rapidly activates PDH (within 10 minutes) and subsequently increases activity of Complex 1 in the mitochondrial electron transport chain (Fig 5B). The demonstration of the importance of Drp1-mediated fission to DA closure in DAs from human infants strengthens the translational relevance of this discovery (Supplemental Table I). The physiological normality of the term DAs was evident in their robust and reversible O₂ constrictor responses (Figure 2A-B). None of these patients had persistent DA; rather the DA had been maintained patent by infusion of prostaglandin E1 where medically necessary.
The DA is a prototypic member of the specialized oxygen homeostatic system, which also includes the type 1 cell in the carotid body, the neuroepithelial body, and resistance pulmonary artery SMC\textsuperscript{16}. Prior studies have showed that increasing PO\textsubscript{2} caused a proportionate increase in the generation of H\textsubscript{2}O\textsubscript{2} by DASMC\textsuperscript{16}. H\textsubscript{2}O\textsubscript{2} is produced by dismutation of the superoxide that results from the increased ETC activity that accompanies oxidative metabolism \textsuperscript{20, 26, 37, 38}. H\textsubscript{2}O\textsubscript{2} serves as a diffusible oxidant and second messenger, signaling the PO\textsubscript{2} to voltage-gated ion channels (e.g. Kv1.5 and the L-type calcium channel) \textsuperscript{20, 26} and enzymes (e.g. rho kinase) \textsuperscript{16}. Prior studies in mammals (including humans) and birds have demonstrated that the superoxide in DA originates from ETC Complexes I and III\textsuperscript{37, 39}. However, it was assumed that oxygen increased mitoROS by changing mitochondrial function, rather than structure. It is now evident that the metabolic and redox changes reflect earlier dynamic changes in morphology.

Multiple lines of evidence indicate that fission in the DASMC results from the increased activity and expression of a single GTPase, Drp1. Drp1 exists as a cytosolic homo-tetramer; however, when activated, it is recruited to the mitochondria. Drp1’s activity is determined by GTP hydrolysis, which is regulated by its C-terminal GTPase effector domain. Upon activation Drp1 assembles in the outer mitochondrial membrane creating a multimeric collar that mediates mitochondrial division \textsuperscript{40,41}. Mdivi-1 inhibits Drp1 multimerization\textsuperscript{31} and in DASMC this prevents oxygen-induced fission (Fig 3A). Fission results in more numerous, smaller mitochondria, which we demonstrate in DASMC (increased MFC) and in human DA rings (decreased mitochondrial size on TEM, Fig 3D).

Mdivi-1 is the most selective and effective of the antifission molecules identified from a screen of 23,000 candidates\textsuperscript{31, 42}. While Mdivi-1 can inhibit cytochrome-c release in cell-free assays, where mitochondrial division does not occur\textsuperscript{42}, the concordant results with Mdivi-1 and siDrp1 in this study provide mechanistic specificity. As an additional method of ensuring mechanistic certainty, we carefully quantified fission using two complementary techniques, MFC, a particle counting technique, and MNF, which measures the rapidity of diffusion of mito-PA-GFP. These assays yielded concordant results; physiological increases in PO\textsubscript{2} fragmented the mitochondrial network and impaired the diffusion of mitochondrial matrix proteins. Finally, we replicated key findings with Mdivi-1 on metabolism, calcium levels and proliferation/apoptosis using a specific molecular approach-siDrp1 (Fig 5B, 6D and 7C).

The physiologic importance of fission is evident from the observation that Mdivi-1 selectively prevents constriction of the DA to O\textsubscript{2}, without altering the response to vasoconstrictor stimuli, such as KCl or phenylephrine (Fig 1). Prior to Mdivi-1, the only agents known to selectivity inhibit O\textsubscript{2}-induced DA constriction were redox agents that mimic hypoxia (e.g. the reducing agent, dithiothreitol\textsuperscript{26, 38}) or ETC Complex I and III inhibitors, rotenone and antimycin A\textsuperscript{20}, which in DASMC reduce mitoROS production. The commonality amongst these inhibitors of DA constriction is that each creates a more reduced redox environment in the DASMC. The central role for the oxidant stimulus, mitochondrial-derived H\textsubscript{2}O\textsubscript{2}, is confirmed by the demonstration that mitochondrial-targeted catalase prevents oxygen-induced elevation of [Ca\textsuperscript{2+}]\textsubscript{cyt}, an accepted surrogate for DA constriction (Fig 6E and F).

To account for O\textsubscript{2}-sensing, fission must occur rapidly before the onset of vasoconstriction. Indeed, Drp1 activation occurs within 3-5-minutes of elevating PO\textsubscript{2} (prior to elevation of tone). This rapidity reflects O\textsubscript{2}-induced, post-translational activation of Drp1 mediated by an increased ratio of active/inhibitory phospho forms (i.e. an increased p-Drp1-Ser 616/637 ratio) (Fig 4 & supplemental Fig I). Activated Drp1 rapidly translocates to the mitochondria (Fig 4C and D). Phosphorylation allows fission to occur in minutes, critical to the very rapid fission we observed. In this regard we identify two relevant kinases that link fission to mitosis (cyclin-dependent kinase) and elevation of intracellular calcium (calcium/calmodulin-dependent protein kinase). The importance of an increased ratio of phosphor-Drp1 Serine-616/637 is supported by the observation that inhibitors of Drp1 phosphorylation reduce O\textsubscript{2}-induced DASMC proliferation, while enhancing apoptosis (Fig 7C), similar to observations in
Inhibition of fission (by Mdivi-1 or siDrp1) largely eliminates the PO2-dependent rise in [Ca2+]cyt without altering the response to KCl (Fig 6A and D), consistent with the O2-specific effects of Drp1 inhibitors on DA constriction. Mdivi-1 appears to affect [Ca2+]cyt primarily by inhibition of mitochondrial H2O2 generation (Fig 6E-F). However, fission may alter [Ca2+]cyt by changing the connection between the mitochondria and organelles, such as the sarcoplasmic reticulum43-45. Supporting this, Mdivi-1 does alter compartmentalization of calcium, reducing calcium release from the sarcoplasmic reticulum and mitochondria during hypoxia (Fig 6 B-C). The changes in calcium may also be linked to metabolism. Lowering mitochondrial calcium, which is known to inhibit PDH46, 47, might contribute to Mdivi-1’s ability to inhibit oxidative metabolism (Fig 6C). A recent study investigating glucose-induced activation of mitochondrial fission and metabolism in the hypothalamus found that siDrp1 decreased mitoROS production and impaired substrate-driven respiration48, consistent with our findings (Fig 6D).

The consequence of fission that triggers vasoconstriction is a rapid increase in oxidative metabolism, evident as an increased OCR (with unchanged ECAR) in DASMC (Fig 5). That this shift results from fission is demonstrated by the observation that pharmacologic (Mdivi-1) or molecular Drp1 inhibition (with siDrp1) prevents the normoxic metabolic shift (Fig 5). We show that O2-induced PDH activation is inhibited by Mdivi-1 (Fig 5B).

There are several mechanisms by which inhibiting fission might inhibit respiration. The role of fission as a stimulant for oxidative metabolism has recently been identified in a heritable form of Parkinsonism that is caused by a mutation of a mitochondrial Ser/Thr kinase, PTEN-induced kinase 1 (PINK1)49. The PINK1 mutation impairs mitochondrial fission and depresses mitochondrial respiration in neurons by decreasing the expression and activity of ETC complex I and IV. Drosophila lacking PINK1 also display impaired mitochondrial fission and metabolism, both of which are rescued by Drp1 overexpression49. In these flies, reduced ETC complex activity reflects defective ETC complex assembly and is rescued by increasing mitochondrial fission. Consistent with this, fission in DASMC increases the activity of ETC Complex I (Fig 4B). We interpret this as indicating that fission changes the architecture of the mitochondria in a manner, which favors assembly and activation of Complex I, a major source of the ROS that signal PO2.

We acknowledge that the role of PINK1/Parkin in familial Parkinsonism is thought to lie in mitochondrial quality control and may not be directly relevant to the more dynamic role of fission in increasing ETC activity in the DA.

There are likely important temporal considerations in metabolic effects of fission. Oxygen’s effects initial effects are on Drp1 phosphorylation (evident in <5 minutes) and this is sufficient to activate PDH activity (evident in <20 minutes). These early changes are critical for initial redox signaling. Inhibiting mitochondrial H2O2 production prevents the rise in cytosolic calcium that triggers vasoconstriction (Fig 5E-F). We show that fission is an important early upstream mediator of calcium signaling through its effects on mitochondrial metabolism (OCR) and redox signaling (H2O2 production). Our prior observations the normoxic redox signaling by H2O2 derives from the mitochondria is supported by the observation that oxygen-induced elevation of cytosolic calcium is prevented by eliminating mitochondrial-derived hydrogen peroxide, using Ad-Mito-Catalase (Fig 6E-F). Fission increases mitochondrial respiration and mitochondrial hydrogen peroxyde generation. Mitochondria-derived hydrogen peroxide serves as a redox signaling molecule that triggers calcium entry and organelar calcium release thereby elevating cytosolic calcium and promoting calcium sensitization in DASMC17, 21, 22, 50-54. Inhibiting fission lowers calcium in all compartments (Fig 6A-C) by changing mitochondrial redox signaling (Fig 6E). Consequently blocking fission (using Mdivi-1) not only prevents constriction.
but also, inhibits PDH activity, a calcium-sensitive mitochondrial enzyme. Likewise, Mdivi-1 and siDRP1, which both inhibit respiration, abrogate O$_2$-dependent increases in cytosolic calcium (Fig 6A and 6D). Thus there is a positive feedback between mitochondrial respiration, H$_2$O$_2$ generation and calcium levels, such that fission-induced increases in respiration increase the magnitude of redox-induced calcium levels.

In normal, term human infants, most DAs close by vasoconstriction within hours-days and by 2 months only 4.5% of DAs are patent (and most of these are clinically silent)\textsuperscript{55}. However, in premature infants, immaturity of the mitochondrial O$_2$-sensing system and the downstream ion channel pathways\textsuperscript{18, 20} is associated with a 21% incidence of persistent DA patency\textsuperscript{56}. The demonstration that DA patency is maintained by inhibiting Drp1 is remarkable. Prolonged O$_2$-exposure selectively increased Drp1 expression without altering expression of the other mediators of fission (e.g. Fis1, Fig 4A) or fusion mediators (e.g. mitofusin-land -2 or optic atrophy protein-1, Fig 4A). Conversely, inhibiting Drp1 in tissue culture preserved DA patency (Fig 7B), at least in part by inhibiting oxygen-induced DASMC proliferation (Fig 7C) and preventing DA fibrosis (Fig 8B-C).

At least part of the effect of inhibiting Drp1 may relate to inhibition of proliferation. Prior work in hyperproliferative, pulmonary hypertensive pulmonary artery SMCs and lung cancer cells indicates that cyclin B- cyclin-dependent kinase 1, a key regulator of mitotic entry coordinates the cell cycle with mitotic mitochondrial division, ensuring equal distribution of mitochondria to daughter cells\textsuperscript{27, 28}.

Because of the clinical efficacy of cyclooxygenase inhibitors (such as indomethacin) in achieving DA closure it might be questioned why other strategies are needed. Beyond the fundamental scientific importance of understanding the molecular basis for oxygen sensing, there is a practical imperative. First, cyclooxygenase inhibitors fail to close the ductus in 30% of neonates\textsuperscript{57}, This lack of efficacy is even more common in extremely low gestational age infants. Second, indomethacin has adverse effects, including significant reductions in renal function (oliguria and reduced creatinine clearance), and reductions in mesenteric and cerebral blood flow\textsuperscript{58}. The combination of indomethacin and steroids also increases the incidence and morbidity of necrotizing enterocolitis\textsuperscript{59}.

LIMITATIONS

While Mdivi-1’s effects were largely mimicked by siDRP1, and while Mdivi-1 had no effect on phenylephrine or KCL constriction or calcium homeostasis, we cannot exclude the possibility that Mdivi-1 has unknown properties, such as acting as an antioxidant’’

Here we show that DASMC proliferation is also reduced by inhibiting kinases that activate Drp1. Pharmacological strategies to activate Drp1 could be developed as a means to close the DA, whilst inhibition of Drp1, by congeners of Mdivi-1, could be used to maintain DA patency in infants awaiting congenital heart surgery.

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DISCLOSURES
None

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DOI: 10.1161/CIRCRESAHA.111.300285

**FIGURE LEGENDS**

**Figure 1:** The Drp1 inhibitor Mdivi-1 selectively inhibits oxygen-induced constriction in rabbit ductus. **A-B:** Representative images showing that Mdivi-1 prevents and rapidly reverses O₂ constriction in rabbit ductus. Mean (±SEM) in panel B show that Mdivi-1 inhibits O₂-constriction without altering phenylephrine- or KCl-mediated constriction, measured under hypoxic condition (***p<0.01 compared to control, n=8, 4, 4).**

**Figure 2:** The Drp1 inhibitor Mdivi-1 selectively inhibits oxygen-induced constriction in human. **A-B:** Representative image and mean data show that Mdivi-1 inhibits O₂-constriction without altering KCl-constriction. n=6.

**Figure 3:** Increased PO₂ causes mitochondrial fission in rabbit and human DASMC. **A:** Serial observations of a rabbit DASMC transfected with mitochondrial targeted Ds-Red shows that 10-15 minutes of increased PO₂ (from 40 to 120 mmHg) fragments the mitochondrial network, thus increasing the MFC. Scale bar=10 µM. n=18-19 cells. **B:** There is increased fission after exposure of rabbit DASMC to increased PO₂. Representative images and mean ± SEM show impaired spread of mito-PA-GFP in oxygenated DASMC, evident as reduced MNF (n=18-19). **C:** The mitochondrial network in human DASMC fragments rapidly with brief (20 minutes) increases in PO₂, evident as an increasing the MFC and reduction in MNF. This is corrected by exposure to Mdivi-1 (20 µM for 20 minutes). **D:** Transmission electron microscopy data confirming that in DASMC within freshly isolated human DA rings, mitochondrial length is shorter after 20 min exposure to normoxia vs. hypoxia (n=62 and 56 mitochondria, respectively).
**Figure 4:** Increased PO$_2$ leads to rapid phosphorylation and activation of Drp1 in human DASMC.  

A: Incubation at PO$_2$ 120mmHg preferentially increases the expression of Drp1 mRNA in human DASMC. N=8 to 10.  

B: Immunoblot of whole cell lysate shows that brief O$_2$-exposure causes phosphorylation (activation) of Drp1 at serine 616 within 3-5 minutes (n=12).  

C-D: Immunoblot of the mitochondrial fraction shows that brief O$_2$-exposure (20 minutes) causes phosphorylation of Drp1 at serine 616, and causes Drp1 translocation to the mitochondria (n=4). Phosphorylation of Drp1 is prevented by inhibitors of its regulatory kinases (KN93- calcium/calmodulin-dependent protein kinase and roscovitine- cyclin-dependent kinase, n=3 to 5). Forskolin, an activator of protein kinase A which can phosphorylate Drp1 at serine 637, had no effect.

**Figure 5:** Normoxia increases oxidative metabolism and H$_2$O$_2$ production in rabbit and human DASMC.  

**A:** Rabbit DASMC: Normoxia increases OCR (left, n=17), decreases ECAR (middle, n=17) and H$_2$O$_2$ (right, n=6). Mdivi-1 (20 μM) inhibits these changes.  

**B:** Human DASMC.  

Top row: Normoxia (20 minutes) increases OCR (left, n=11) and PDH activity (third panel, n=9). Mdivi-1 (20 μM) inhibits the changes. Long-term normoxia (2 day) increases ETC complex 1 activity (right, n=10).  

Bottom row: Mdivi-1 (20 μM) inhibits the basal and FCCP induced maximal OCR and increases basal ECAR in human DASMC during normoxia condition (n=10). Oligomycin, used to show coupling of respiration to ATP synthesis, had similar effects with or without Mdivi-1. Antimycin is an inhibitor of ETC complex 3 and demonstrates that the OCR in DASMC is almost exclusively based on mitochondrial respiration. Antimycin’s effects were unaltered by Mdivi-1. Like Mdivi-1, SiDrp1 inhibits the basal OCR in human DASMC during normoxia condition while scrambled siRNA had no effect (n=11).

**Figure 6:** Drp1 inhibition impairs oxygen-induced increases in cytosolic calcium DASMC.  

A-C: Mdivi-1 lowers calcium in all subcellular pools in DASMC and prevents oxygen-induced elevations in cytosolic calcium. A; n=44-46; B; n=8-13; C; n=38-57 cells.  

D: SiDrp1 inhibits increases in calcium caused by oxygen (n=90-109 cells) without altering the response to KCl (n=15-30 cells) in human DASMC.  

E-F: Representative images and mean data showing that elimination of mitochondrial H$_2$O$_2$ using Ad-mitoCAT prevents O$_2$-induced elevation of cytosolic calcium (n=46 to 51) without altering the response to KCl (n=19 to 37 cells) in human DASMC. E right panel: t-butyl hydroperoxide (1 μm) increases cytosolic calcium (n=17 cells).

**Figure 7:** Prolonged Drp1 inhibition prevents DASMC proliferation and prevents DA closure.  

A-B: In a tissue culture model, occlusion of the lumen occurs in response to incubation in oxygen after 3 days in rabbits (n=10 to 13 DA rings) and 6-14 days in humans (n=7 DA rings). Incubation with Mdivi-1 prevents DA closure in this model. C: Mdivi-1 prevents the proliferation of human DASMC, measured using EdU flow cytometry (n=6 to 9). Roscovitine, KN93 and siDrp1 inhibit the proliferation of human DASMC during normoxia, whereas siPKCδ has no effect (middle panel, n=4 cultures of DASMC). SiDrp1 induces apoptosis in human DASMC during normoxia condition (right panel, n=4 cultures of DASMC).

**Figure 8:** Mdivi-1 significantly prevented fibrotic remodeling of the rabbit DA.  

8A: Time course of channels in apoptosis (TUNEL) and proliferation (PCNA) in rabbit DAs maintained in tissue culture with or without mdivi-1 (n=5 DA rings/group).  

8B-C: Immunostaining shows that Mdivi-1 inhibits elastin production in vivo, impairing an important step in DA closure. The dramatic O$_2$-induced reduction in DA lumen and overall DA size over time is also attenuated by Mdivi-1 (n=5-8 DA rings at D6).  

8D: Proposed mechanism.
Novelty and Significance

What Is Known?

- Closure of the ductus arteriosus (DA) is essential for the transition from fetal to neonatal circulation.
- PO$_2$-dependent vasoconstriction causes functional DA closure within minutes.
- PO$_2$-dependent vasoconstriction occurs by a mechanism that is intrinsic to the DA smooth muscle cells and reflects the regulation of oxygen-sensitive potassium and calcium channels and rho kinase, by a mitochondrial redox sensor.
- Within days at elevated PO$_2$ a fibrogenic, proliferative mechanism causes anatomical closure.

What New Information Does This Article Contribute?

- Mitochondrial fission is crucial for O$_2$-induced constriction and closure of human and rabbit ducti.
- O$_2$ rapidly (<5 minutes) causes mitochondrial fission by post-translational modification and activation of dynamin-related protein 1.
- Fission triggers an oxidative metabolic shift in the DA that increases production of the redox signaling molecule, hydrogen peroxide.
- Dynamin-related protein 1 can be targeted to modulate ductal patency.

Closure of the ductus arteriosus (DA) is essential for the transition from the fetal to neonatal circulation. Functional closure, due to O$_2$-induced vasoconstriction, begins within minutes of birth and is required for subsequent anatomical closure. Within minutes of increasing PO$_2$ there is an increase in mitochondria-derived reactive oxygen species (mitoROS). The mitoROS inhibit O$_2$-sensitive potassium channels and activate calcium channels and rho kinase, initiating vasoconstriction. The precise mechanism by which O$_2$ increased mitoROS and the relevance to anatomical DA closure are unknown. We observed that physiologic increases in PO$_2$ fragment the DASMC’s mitochondrial network, suggesting that structural changes in mitochondria are a key early step in O$_2$-sensing. O$_2$ induces fission by phosphorylating dynamin related protein 1 (Drp1), at serine 616. Fission activates pyruvate dehydrogenase, increases oxygen consumption and mitoROS and initiates redox-based O$_2$-sensing. Drp1 inhibition selectively blocks oxygen-mediated DA constriction and, if sustained, prevents DA closure. These findings suggest that mitochondrial fission is an obligatory early step in O$_2$-sensing that is crucial to DA constriction and closure. Drp1 and its regulatory kinases are appealing therapeutic targets for modulating DA patency. The translational relevance of these finding is enhanced by the proof of this concept in ducti from 30 human patients with congenital heart diseases.
Figure 2 human DA

A

B

Δ tension O₂ (mg)

Control  Mdivi-1

Δ tension KCl (mg)

Control  Mdivi-1

P<0.01 **
Role of Dynamin Related Protein 1 (Drp1)-Mediated Mitochondrial Fission in Oxygen-Sensing and Constriction of the Ductus Arteriosus

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Supplemental Material

TABLE I: Patient demographics

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<th>Gender</th>
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**Abbreviations:** HLHS=Hypoplastic left heart syndrome, D-TGA=D-transposition of great arteries, VSD=Ventricular septal defect, COA=coarctation of the aorta, TOF=Tetralogy of Fallot, DILV=Double inlet left ventricle, PS=Pulmonic stenosis, ASD=Atrial septal defect, TAPVC=Total anomalous pulmonary venous connection
Supplemental Figure Legends

Supplemental Fig. I: A: Immunoblot of mitochondrial-enriched fraction of human DASMC exposed to 20 minutes of normoxia shows that SiPKCδ had no effect on the Drp1 phosphorylation at Ser 616 nor did it alter dephosphorylation of Drp1 at Ser 637, n=3 DA culture plates. B: qRT-PCR demonstrates that SiRNA decreases PKCδ mRNA, n=6 DA plates.

Supplemental Fig. II: SiDrp1 decreases Drp1 mRNA (A) and protein level (B) in human DASMC. n=3 DA cell culture each.

Supplemental Fig. III: Transfection of Ad-mitoCAT increases the catalase activity in human DASMC, n=3 DA culture plates each.
Supplemental Figure II

A

qRT-PCR

Relative Intensity compared to B2M

Control Scramble SiDrp1

** P<0.01

B

western blot

Relative Intensity compared to actin

Control Scramble SiDrp1

Drp1

actin