Mitochondrial fission is important for organelle transport, inheritance, and turnover, and alterations in fission are seen in neurological disease. In mammals, mitochondrial fission is executed by dynamin-related protein 1 (Drp1), a cytosolic guanosine triphosphatase that polymerizes and constricts the organelle. Recruitment of Drp1 to mitochondria involves receptors including Mff, MiD49, and MiD51. MiD49/51 form foci at mitochondrial constriction sites and coassemble with Drp1 to drive fission. Here, we solved the crystal structure of the cytosolic domain of human MiD51, which adopts a nucleotidyltransferase fold. Although MiD51 lacks catalytic residues for transferase activity, it specifically binds guanosine diphosphate and adenosine diphosphate. MiD51 mutants unable to bind nucleotides were still able to recruit Drp1. Disruption of an additional region in MiD51 that is not part of the nucleotidyltransferase fold blocked Drp1 recruitment and assembly of MiD51 into foci. MiD51 foci are also dependent on the presence of Drp1, and after scission they are distributed to daughter organelles, supporting the involvement of MiD51 in the fission apparatus.

**Introduction**

Mitochondria are highly dynamic organelles and undergo continuous fission and fusion events, a balance of which is vital for their function and cellular distribution (Westermann, 2010). Mutations in genes associated with mitochondrial dynamics have been linked with peripheral and optic neuropathies (Alexander et al., 2000; Delettre et al., 2000; Züchner et al., 2004; Davies et al., 2007) and fatal developmental abnormalities (Ishihara et al., 2009), whereas defects in mitochondrial dynamics have been implicated in Parkinson’s, Alzheimer’s, and Huntington’s disease (Wang et al., 2008; Arduíno et al., 2011; Reddy et al., 2011). The primary regulator of fission, controlled through both posttranslational modifications and interactions with mitochondrial adaptor proteins, is the dynamin family member Drp1 (Dnm1 in yeast; Bui and Shaw, 2013; Elgass et al., 2013). Cytosolic Drp1 is recruited to mitochondrial constrictions, often where the endoplasmic reticulum crosses (Friedman et al., 2011). There, it polymerizes into spirals and through GTP-dependent conformational changes it constricts the organelle leading to membrane scission (Lackner et al., 2009). The crystal structures of dynamin and Drp1 revealed interfaces involved in protein oligomerization along with a mechanism for polymer constriction (Faelber et al., 2011; Ford et al., 2011; Fröhlich et al., 2013). Unlike dynamin-1, Drp1 lacks a pleckstrin homology domain and is recruited to mitochondria through outer membrane receptors Fis1 (Mozdy et al., 2000; Yoon et al., 2003; Stojanovski et al., 2004), Mff (Gandre-Babbe and van der Bliek, 2008; Otera et al., 2010), along with chordate-specific MiD49 and MiD51 (Palmer et al., 2011; Zhao et al., 2011). Individually, Mff, MiD49, and MiD51 have been shown to be sufficient to recruit Drp1 to drive fission (Koirala et al., 2013; Losón et al., 2013; Palmer et al., 2013), whereas the role of Fis1 as a Drp1 receptor has been questioned (Otera et al., 2010). Recently, addition of stoichiometric amounts of MiD49 was found to increase Drp1-mediated constrictions.

© 2014 Richter et al. This article is distributed under the terms of an Attribution-Noncommercial-Share Alike-No Mirror Sites license for the first six months after the publication date (see http://www.rupress.org/terms). After six months it is available under a Creative Commons License (Attribution-Noncommercial-Share Alike 3.0 Unported license, as described at http://creativecommons.org/licenses/by-nc-sa/3.0/).
Results and discussion

MiD51 belongs to the nucleotidyltransferase fold superfamily of proteins

To search for a construct amenable to crystallization, we performed limited proteolysis on mouse MiD49 lacking its transmembrane domain. We identified a large (≥40 kD) trypsin-resistant fragment of MiD49 that lacks a predicted disordered region of the protein (Fig. S1 A). When human MiD51 lacking a homologous disordered region (residues 50–123) but still containing its

of liposomes from 31 to 15 nm, a point at which membrane scission is possible (Koirala et al., 2013). This indicates that MiD49/51 may be actively involved in facilitating Drp1 scission similar to the unrelated yeast adaptors Mdv1/Caf4 that are absent in higher eukaryotes (Bui and Shaw, 2013). MiD49/51 lack significant sequence similarity to any other proteins and they do not possess characteristic sequence domains/motifs. Here we determine the crystal structure of the cytosolic domain of MiD51 and identify a critical region in MiD51 important for Drp1 recruitment and assembly of the fission apparatus.

Figure 1. Crystal structure of a MiD51 cytosolic domain sufficient for Drp1 recruitment. (A) MiD51-GFP and MiD51-DKR-GFP were expressed in wild-type MEFs and MEFs lacking endogenous MiD51 (MiD51 TALEN). Cells were subsequently immunostained for Drp1 and the mitochondrial marker protein Tom20 and visualized by fluorescence microscopy. (right) Line scans demonstrating colocalization of Drp1 with MiD51-GFP and MiD51-DKR-GFP. Bars, 20 µm. (B) Structure of MiD51 N118 displaying the nucleotidyltransferase domain in blue and DRR domain in green. (C) Surface representation of MiD51 N118; view and coloring as in B. (D) Topology of MiD51 and illustration of mutant constructs used in this study. TM, transmembrane domain; DR, disordered region; NT, nucleotidyltransferase domain.
N-terminal transmembrane anchor (termed MiD51\textsubscript{ATM}) was transiently expressed in mouse embryonic fibroblasts (MEFs), it was still able to recruit cytosolic Drp1 to mitochondria like transiently expressed in mouse embryonic fibroblasts (MEFs), consistent with a previous knockdown study (Palmer et al., 2011).

We crystallized recombinant, monomeric human MiD51\textsubscript{ATM} (lacking its N-terminal transmembrane anchor and disordered region [Fig. S1 C]) and refined the native structure to R\textsubscript{work} and R\textsubscript{free} values of 0.223 and 0.263, respectively (Table 1). We observed clear and continuous electron density for residues 133–461 in all four copies of MiD51\textsubscript{ATM} present in the asymmetric unit, with the remaining residues at the N terminus presumed to be disordered. Overall MiD51\textsubscript{ATM} serves clear and continuous electron density for residues 133–461 in all four copies of MiD51\textsubscript{ATM} (Fig. 1 A and Fig. S1 B). These MEFs lacking MiD51 have more elongated mitochondria than control MEFs, consistent with a previous knockdown study (Palmer et al., 2011).
MiD51 specifically binds ADP and GDP

The region in MiD51 structurally homologous to the nucleotide binding site in cGAS and other structural homologues revealed additional electron density not associated with the polypeptide chain, which we were unable to assign (Fig. S2 A). Isothermal titration calorimetry (ITC) studies of MiD51 with various nucleotides showed specific binding to only ADP and GDP (Fig. 2, A and B) with affinities of 2.9 and 3.8 μM, respectively. We determined the crystal structures of MiD51 bound to either GDP or ADP (Fig. 2, C and D; and Table 1). Both structures revealed clear densities for a single nucleotide diphosphate (Fig. S2, B and C). Superimposition of the Cα backbone of MiD51 with MiD51–GDP or MiD51–ADP revealed no significant conformational changes, with RMSD variations ranging from 0.18 to 0.63 Å (Fig. S2 D). In the MiD51–ADP complex, Ser-187 and Ser-340 form hydrogen bonds with the purine, and Ser-189, His-201, Arg-342, and Lys-368 contact the phosphate groups (Fig. 2, C and D). In the MiD51–GDP complex, an additional hydrogen bond is formed by GDP with Gln-203 because of the purine group in GDP pointing inwards toward the back of the nucleotide binding pocket. It should be noted that the purine group of ADP is rotated away at a 180° angle compared with GDP, leading to an additional conformational shift in the ribose ring (Fig. 2 E).

Catalytically active members of the nucleotidyltransferase family harbor a highly conserved set of acidic residues in their active sites (DE[h][DE][h] and h[DE][h]), which coordinate divalent ions and activate hydroxyl acceptor groups in a substrate (Kuchta et al., 2009). In MiD51, only a single aspartic acid residue (Asp-200) is present in this region, indicating that MiD51 is unlikely to support nucleotide hydrolysis and transfer. Furthermore, no evidence for a catalytic Mg2+ ion site was discernible from the electron density, and cocrySTALLizing MiD51 with nucleotide and MnCl2 did not reveal bound Mn2+ ions in an anomalous difference Fourier map (Fig. S2, E and F). Thus MiD51 resembles a set of proteins, including NF45 and NF90, that do not have enzymatic activity.

We undertook extensive mutagenesis of residues involved in nucleotide binding (Fig. 2 F) and transiently expressed constructs in cells. In all cases, we found that Drp1 was still recruited to mitochondria (unpublished data). Simultaneous mutation of residues involved in contacting the phosphate backbone of ADP or GDP (K368E, K372E, R342E, and H201D) was found to disrupt nucleotide binding according to ITC measurements (Fig. 2 B), yet expression of this construct (terming nucleotide binding disruption, MiD51NBD) in MiD51TALEN MEFs still induced Drp1 recruitment to mitochondria (see Fig. 3 B). In addition, the mitochondrial network in MiD51TALEN MEFs shifted from an extended to a reticular phenotype after 4-hydroxytamoxifen (4-OHT) induced expression of either MiD51 or MiD51NBD (Fig. 2 G). Whether the nucleotide binding groove is physiologically relevant or whether mitochondria have simply adapted the fold for a novel purpose remains to be fully clarified.

Identification of the Drp1 recruitment domain in MiD51

MiD51 contains an additional domain inserted into the nucleotidyltransferase fold (Fig. 1, B and D; and Fig. 3 A). To investigate if this region is involved in mediating mitochondrial fission, MiD51 mutants were expressed in MiD51TALEN MEFs. A MiD51-GFP fusion construct lacking a defined loop region corresponding to residues 238–242 (MiD51APPEYP) was still targeted to mitochondria, yet, unlike MiD51-GFP, the active recruitment of Drp1 to the mitochondrial surface was blocked (Fig. 3 B). Similarly, a MiD51 mutant (MiD51R235A) that disrupts a salt bridge (R235-D249) located below this loop, was also unable to recruit Drp1 to mitochondria. Because of leaky GFP expression in this cell line, we repeated the expression in COS7 cells where such leakage was not observed (unpublished data). Again, Drp1 recruitment defects for MiD51APPEYP and MiD51R235A were observed with these mutants resembling expression of the negative control protein Miro1 (Fig. 3 C). In contrast, an adjacent nonsalt bridge mutant (MiD51R234A) did not block Drp1 binding to mitochondria (Fig. 3 C). Similar findings were also observed when we induced the expression of untagged mouse MiD51 constructs in MiD51TALEN MEFs (Fig. S3, A and B).

We recently found that redirecting MiD51 to lysosomes by using a heterodimerizer assay led to the recruitment of Drp1 to that organelle (Palmer et al., 2013). This and other studies (Koirala et al., 2013; Losón et al., 2013) have shown that expression of MiD49 or MiD51 is sufficient for Drp1 recruitment and subsequent fission. We repeated this assay using MiD51APPEYP and found that although this construct could be directed to lysosomes, Drp1 was not recruited (Fig. 3 D). In contrast, MiD51 constructs lacking only the transmembrane anchor (MiD51ATM) or additionally lacking the disordered region (MiD51AN118) were able to redirect Drp1 to lysosomes, hence underscoring the critical role of the β4–α4 loop in Drp1 recruitment.

To determine whether a loss of Drp1 recruitment by MiD51APPEYP could be attributed to either misfolding or a conformational change in the Drp1 recruitment region (DRR), we determined the structures of MiD51APPEYP bound to nucleotides (Fig. 3 E and Table 1). Superimposition of mutant and wild-type structures showed a distinct lack of conformational changes, with RMSD variation ranging from 1.09 to 1.32 Å (Fig. S3 C). Small differences are observed in helix α6 of MiD51AN118APPEYP, which is located in a crystal contact, leading to the unwinding of two turns of helix. Loss of this secondary structure allows extension of α5 by five residues in MiD51AN118APPEYP (residues 259–275) compared with MiD51AN118 (residues 259–270). Despite the shortened β4–α4 loop, the DRR in the deletion mutant exhibited a largely native conformation and the structurally important R235-D249 salt bridge was maintained (Fig. 3 E). A comparison of the electrostatic surface between wild type and MiD51APPEYP revealed only small changes in the charge distribution between both structures (Fig. S3, D and E), suggesting that the topology of the β4–α4 loop is important for Drp1 recruitment. Notably, this loop is part of a region that is absent in
all other structures determined for members of the nucleotidyltransferase fold superfamily with the exception of the recently determined structure of cGAS (Civril et al., 2013; Gao et al., 2013a). However, the β4–α4 loop is shorter in cGAS and the residues equivalent to PEYFP are located on the side of the loop rather than on the apex as is the case for MiD51.
membrane like that of the control GFP-Miro1 (Fig. 4 A). We expressed MiD51-GFP in Drp1−/− MEFs (Wakabayashi et al., 2009) and found that it also did not assemble into foci and was instead distributed along the extended mitochondrial filaments (Fig. 4 B). We conclude that MiD49/51 assembly into foci is dependent on the presence of Drp1. Moreover, live-cell imaging of MiD51-GFP foci revealed that fission events indeed occurred at these sites and that a population of MiD51 in foci was inherited by each daughter organelle (Fig. 4 C and Videos 1 and 2).

Drp1 induces MiD51 assembly at foci that form scission sites
MiD51 and MiD49 can be found at foci at mitochondrial constriction sites, consistent with their role in fission (Palmer et al., 2011). Interestingly, MiD51 mutants still able to recruit Drp1 (e.g., MiD51NBD and MiD51R235A) also retained their ability to form foci at mitochondria, yet mutants MiD51NBD and MiD51R235A that are deficient in Drp1 recruitment ability did not form foci and were instead diffusely distributed along the outer membrane like that of the control GFP-Miro1 (Fig. 4 A). We expressed MiD51-GFP in Drp1−/− MEFs (Wakabayashi et al., 2009) and found that it also did not assemble into foci and was instead distributed along the extended mitochondrial filaments (Fig. 4 B). We conclude that MiD49/51 assembly into foci is dependent on the presence of Drp1. Moreover, live-cell imaging of MiD51-GFP foci revealed that fission events indeed occurred at these sites and that a population of MiD51 in foci was inherited by each daughter organelle (Fig. 4 C and Videos 1 and 2).
The distribution of MiD51 foci between organelles after a successful scission event also resembles that of yeast Drp1, Dnm1 (Bleazard et al., 1999). In yeast, Fis1 is a membrane receptor that helps assemble Mdv1p onto the mitochondrial surface where Mdv1p facilitates Dnm1 assembly into scission complexes (Tieu and Nunnari, 2000; Karren et al., 2005). Thus, MiD51 appears to act in a similar way to Mdv1p, yet it does not require an additional receptor because it is already anchored in the mitochondrial outer membrane. Both MiD51 and Mdv1p have different structural folds and hence the final scission complexes are likely to adopt different topologies. Consistent with this, it was recently proposed that variations in the sequence of the Insert B region of dynamin-related proteins accommodates diversity in adaptor binding (Ford et al., 2011; Bui and Shaw, 2013).

In conclusion, we found that MiD51 adopts a nucleotidyltransferase fold and can bind ADP and GDP. Considering the low micromolar affinities for both GDP and ADP, the ability of MiD51 to bind both nucleotides is intriguing, particularly in light of the absence of catalytic activity or ATP/GTP binding. Potential roles include nucleotide sensing or as an effector of Drp1’s GTPase activity. However, we found that fission events still occurred at foci formed by the MiD51 nucleotide binding mutant (Video 3). A more prosaic function such as nucleotides being a cofactor to stabilize MiD51 is therefore possible. The nucleotidyltransferase fold acts as a protein recruitment platform to assemble oligomers of the key mitochondrial fission regulator Drp1 into foci on the mitochondrial membrane. This unexpected functionality is mediated, at least in part, by a region inserted into the nucleotidyltransferase lobe that harbors the β4–α4 loop crucial for MiD51–Drp1 interactions. Intriguingly, it is not only the loss of the PEYFP motif in the loop, but also destabilization of the loop, that impairs Drp1 recruitment because mutation of R235, which forms a salt bridge directly below the β4–α4 loop, also abrogates Drp1 recruitment. This suggests that the loop topology may be a critical factor in Drp1 recruitment because loss of the loop does not adversely affect the protein fold and only minor changes in the charge distribution are seen after loop removal. Furthermore, the PEYFP sequence is not fully conserved in MiD49, which instead harbors a LEFCP motif in the β4–α4 loop, supporting the notion that it is loop topology rather than the precise amino acid sequence that is important. In a previous study, it was found that deletion of residues 160–169 in MiD51 also resulted in loss of Drp1 recruitment (Zhao et al., 2011). However, this region is a central part of the α2 helix and the deletion would most likely disrupt a key secondary structure element causing misfolding. Despite the identification of this critical loop element in MiD51 for Drp1 recruitment, the precise topology of the putative MiD51–Drp1 oligomer complex or its stoichiometry remains elusive. A previous study has suggested that MiD51 and Drp1 copolymerize, yet at least in vitro MiD51 does not appear to self-assemble into oligomers (Koirala et al., 2013). Whether or not the β4–α4 loop is the sole region of contact between MiD51 and Drp1 remains to be determined. Considering the proposed copolymerization of MiD51 and Drp1 it seems likely that other regions in MiD51 make direct contact with a Drp1 oligomer. Indeed, such contact surfaces may be necessary to achieve the reduction in diameter of Drp1 polymers from ~31 to 15 nm to enable mitochondrial membrane scission. Additional contacts may also be mediated by other players of the fission machinery, including Mff, actin, and the endoplasmic reticulum (Friedman et al., 2011; Korobova et al., 2013). The interplay of these with MiD49/51 and Drp1 remains to be clarified.

Materials and methods

Accession numbers
Coordinates and structural factors have been deposited in the Protein Data Bank using accession numbers 4NXT, 4NXU, 4NXV, 4NXW, and 4NXX.
Expression and purification of MiD51

Mature MiD51 [UniProt accession no. Q9NNG6; aa 118–463] was cloned into the pGEX-4T1 expression vector (GE Healthcare). Plasmid DNA was transformed into chemically competent Escherichia coli (BL21(DE3)polyS) as described previously (Sambrook and Russell, 2001) with positive transformants being selected for by carbenicillin addition to culture medium. Recombinant protein was induced by addition of 0.5 mM IPTG at a culture density of OD600 = 0.6, followed by 24 h expression at 22°C.

After harvesting and resuspension in GST binding buffer (0.1 M NaCl, 0.02 M Tris-Cl, pH 8.5, and 0.02 M EDTA), bacterial cells were lysed by freeze-thawing and sonication. Lysate was cleared by centrifugation (20,000 g for 30 min) and supernatant was applied to GST-resin (BD) loaded into a 20-mL gravity flow column. Resin was washed with 30 column volumes of wash buffer (PBS and 10% vol/vol glycerol) and eluted in 0.1 M reduced glutathione and 0.05 M Tris, pH 8.5. Removal of the GST affinity tag occurred by in-solution cleavage with thrombin, followed by dialysis against PBS and collection of the flow-through upon reaplication to GST resin. Recombinant MiD51 was subjected to gel filtration (Superdex200 10/300 GL; GE Healthcare) where it eluted as a single peak and concentrated using Amicon Ultra 30,000 MWCO centrifugal filters (EMD Millipore) to a final concentration of 6 mg/mL. The identity of MiD51 was confirmed by mass spectrometry and protein concentration was determined using a NanoDrop 2000 Spectrophotometer (Thermo Fisher Scientific).

Crystallization, derivatization, and structure determination of MiD51

Crystals of MiD51 and MiD51 complexes were obtained by hanging drop vapor diffusion in 16% (wt/vol) polyethylene glycol 8000, 0.10 M Hepes, pH 7.5, 10.0% (vol/vol) 2-propanol, and 0.2 M ammonium sulfate at 293 K by combining 1 µL of protein with 1 µL of crystallant over 500 µL of reservoir solution. Co-crystallization of nucleotides was performed by the addition of 10 mM nucleotide and 20 mM MnCl2, followed by flash-cooling in mother liquor supplemented with 15% glycerol, 0.02 M nucleotide, and 20 mM MnCl2. Xenon derivatization of native crystals was performed using a xenon chamber (Hampton) at the Australian Synchrotron. Native and derivative data were collected at the Australian Synchrotron, Beamline MX2, 0.005% (vol/vol) xenon in the data collection wavelength of 0.95 Å, and 1.4586 Å, respectively. Initial experimental phases of native MiD51 were obtained using Xe-SAD in PHENIX Autosol. Secondary structure elements were identified from the initial experimental map and modeled using polyalanine chains. Initial polyalanine models were used to improve xenon sites until the entire MiD51 chain could be traced. The polyalanine model was subsequently improved using automated building in Buccaneer (Cowtan, 2006, 2008), followed by manual rebuilding using Coot. The final model was refined to 2.0 Å with a final Rwork and Rfree of 0.218 and 0.266, respectively.

Improvement of this model was achieved using automated building in Buccaneer (Cowtan, 2006, 2008), where it eluted as a single peak in 300 mM NaCl, 0.05 M Tris-Cl, and 0.02 M EDTA. After dialysis against buffer (0.1 M NaCl, 0.05 M Tris-Cl, pH 8.0, and 0.02 M EDTA), bacterial cells were lysed by freeze-thawing and sonication. Lysate was cleared by centrifugation (20,000 g for 30 min) and supernatant was applied to GST-resin (BD) loaded into a 20-mL gravity flow column. Resin was washed with 30 column volumes of wash buffer (PBS and 10% vol/vol glycerol) and eluted in 0.1 M reduced glutathione and 0.05 M Tris, pH 8.5. Removal of the GST affinity tag occurred by in-solution cleavage with thrombin, followed by dialysis against PBS and collection of the flow-through upon reaplication to GST resin. Recombinant MiD51 was subjected to gel filtration (Superdex200 10/300 GL; GE Healthcare) where it eluted as a single peak and concentrated using Amicon Ultra 30,000 MWCO centrifugal filters (EMD Millipore) to a final concentration of 6 mg/mL. The identity of MiD51 was confirmed by mass spectrometry and protein concentration was determined using a NanoDrop 2000 Spectrophotometer (Thermo Fisher Scientific).

Cell culture and construction of MiD51-driven MiEFs

MiEFs, COS-7, and Hela cells were cultured as previously described (Palmer et al., 2011). Wild-type and Drp1−/− MEFs were previously obtained by Wakabayashi et al. (2009). Generation, selection, and induction of stable MiD51-inducible MEF cell lines were performed as previously described (Dunning et al., 2007). In brief, HEK293T cells were used to generate lentiviral particles and the resulting particle supernatants were obtained by transiently transfecting two plates of HEK293T cells using Lipofectamine 2000 (Invitrogen). Each plate was transiently transfected with pCMV6/iRX (packaging plasmid) and pGAG54 E20 (viral envelope). One plate was also transiently transduced with pU6G4 (transcriptional activator), whereas the other was cotransfected with either p5xUAS-MiTAL, p5xUAS-MiTAL, or p5xUAS-MiTAL, or p5xUAS-MiTAL. After 24 h incubation, viral supernatants were harvested and filtered through a 0.45-µm membrane filter and incubated with MiEF cells; MEFs were also transduced with pCMV6/iRX for 48 h. After this the viral media was replaced with 10% (vol/vol) FCS DMEM containing 5 µg/ml puromycin and 200 µg/ml hygromycin.

To generate an MEF clonal cell line lacking MiD51 using TALEN technology, the initiation start codon in the first coding exon of mouse Smc7/1 (Genbank accession no. NM_178719.5) encoding MiD51 was targeted using ZifIT (Sander et al., 2007, 2010), yielding the Tal effector repeat encoding DNA fragments (Bi, XbaI sites) of the FRB domain in the pC4-RHE vector, containing an in-frame GFP fusion (transcriptional activator), whereas the other was cotransfected with either p5xUAS-MiTAL, p5xUAS-MiTAL, or p5xUAS-MiTAL. After 24 h incubation, viral supernatants were harvested, filtered, and incubated with MEF cells; MEFs were also transduced with pCMV6/iRX for 48 h. After this the viral media was replaced with 10% (vol/vol) FCS DMEM containing 5 µg/ml puromycin and 200 µg/ml hygromycin.

To generate an MEF clonal cell line lacking MiD51 using TALEN technology, the initiation start codon in the first coding exon of mouse Smc7/1 (Genbank accession no. NM_178719.5) encoding MiD51 was targeted using ZifIT (Sander et al., 2007, 2010), yielding the Tal effector repeat encoding DNA fragments (Bi, XbaI sites) of the FRB domain in the pC4-RHE vector, containing an in-frame GFP fusion. FKBP-containing plasmids (Lazarou et al., 2005) were then co-transfected with miMiD51-RNA (NG NG HD HD HD NG HD NI NG NG HD HD NI NN NN NI NN NI NN NI) and mMiD51-R (NG NG HD HD HD HD HD HD HD NG HD HD HD HD NI NN NN NI NN NI) into wild-type MEFs. After 48 h incubation, viral supernatants were harvested, filtered, and incubated with MEF cells; MEFs were also transduced with pCMV6/iRX for 48 h.

Mutagenesis of MiD51

MiD51 lacking the N-terminal transmembrane anchor and the disordered region (MiD51NT19) and MiD51 lacking the Drp1 recruitment domain (MiD51PEYTP) were cloned downstream of the FKBp domain, following by GFP. Other point mutations and truncation constructs were created using either extension overlap PCR or the Quick Change Site Directed Mutagenesis kit (Agilent Technologies). All MiD51 mutants directed to mitochondria still contained the N-terminal transmembrane anchor. All mutations and deletion constructs were verified by nucleotide sequencing.

Mitochondrial isolation and Western blotting

Mitochondrial isolation from cultured cells has been described previously (Palmer et al., 2011). In brief, cells were resuspended (20 mM HEPES, 220 mM
mannitol, 70 mM sucrose, 1 mM EDTA, and 0.5 mM PMSE, pH 7.6) and homogenized using a glass homogenizer and a drill-fit pestle. The homogenate was then centrifuged at 800 g to pellet nuclear and cellular debris. The resulting supernatant was centrifuged at 10,000 g for 10 min at 4°C and the pellet was resuspended and centrifuged at 10,000 g for 10 min at 4°C to yield an enriched mitochondrial fraction. Proteins were separated by SDS-PAGE followed by Western blot analysis using primary antibodies and horseradish peroxidase-conjugated secondary antibodies. Signals were detected using ECL substrate (GE Healthcare) and images were obtained using a G:Box Chemi XT system (Syngene).

**Microscopy**

Cells were fixed, permeabilized, stained with the appropriate antibodies, mounted in mounting medium containing DABCO (Harlow and Lane, 1988), and then analyzed using a confocal microscope (LSM 510; Carl Zeiss) equipped with a ConfoCor 3 system containing an avalanche photodiode detector (40× oil objective with a 1.3 numerical aperture) and an avalanche photodiode detector using a 100× oil immersion objective with a 1.4 numerical aperture. Images were obtained with the same settings, except for detector gain adjustments that were performed to normalize saturation levels.

For 4-OHT-induced cells (Fig. S3 B), fixed cells were viewed using a fluorescence microscope (1X81; Olympus) using a 100× oil immersion objective with a 1.4 numerical aperture objective with a 1.4 numerical aperture. Images were acquired using an f-view 2 camera (Olympus) and processed using SoftSyst SIS (Olympus).

For live-cell imaging (Fig. 4 C and Videos 1 and 2), COS-7 cells expressing MiD51-GFP were stained with 25 nM MitoTracker deep red (Molecular Probes) and imaged with a fluorescence microscope (1X81; Olympus) using a 100× oil immersion objective with a 1.4 numerical aperture. Images were acquired using an f-view 2 camera (Olympus) and processed using SoftSyst SIS (Olympus).

**Online supplemental material**

Fig. S1 shows the generation of MiD51 soluble protein (A and B), a Western blot analysis of wildtype and MiD51^{ΔNHE} mitochondria (B), a structural sequence alignment (C), and an alignment with CGAS (D). Fig. S2 shows structural detail of the MiD51 nucleotide binding pocket and RMSD values between different MiD51 structures. Fig. S3 shows an analysis of Dmp1 recruitment in MiD51-inducible MEFs and the structural details of MiD51^{ΔNHE}. Videos 1 and 2 show time-lapse imaging of fission events at MiD51-GFP foci in COS-7 cells and Video 3 shows fission events occurring at MiD51^{ΔNHE}-GFP in MiD51^{ΔNHE}-GFP. Online supplemental material is available at http://www.jcb.org/cgi/content/full/jcb.20131014/DC1.

We thank Santosh Panikkar, Janet Newman, Australian Synchrotron MX beamline staff, Sofia Caria, Grant Mills, Matt Pengini, and the CSIRO C3 Centre for assistance, Megan Maher, Begona Heras, and Mark Hulet for helpful discussions, and Richard Voute and Michael Lazzaro for reagents.

This work was supported by grants from the Australian Research Council and National Health and Medical Research Council (to M.T. Ryan and M. Kvarnstru) and National Institutes of Health (to H. Sasaki; grant number GM089853).

The authors declare no competing financial interests.

Submitted: 4 November 2013
Accepted: 6 January 2014

References


Korobova, F., V. Ramabhadran, and H.N. Higgs. 2013. An actin-dependent step of Mff and Fis1 in Drp1 recruitment and are specific for mitochondrial fission machinery. J. Cell Biol. 201:155–170. http://dx.doi.org/10.1083/jcb.201206128


Reddy, D., J.M. McCaffery, and J.M. Shaw. 2013. Adaptor proteins MId49 and MId51 can act independently of Mff and Fis1 in Drp1 recruitment and are specific for mitochondrial fission. J. Biol. Chem. 288:27584–27593. http://dx.doi.org/10.1074/jbc.M113.479873


