The mitochondrial acyl-carrier protein interaction network highlights important roles for LYRM family members in complex I and mitoribosome assembly

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**Running Title:** LYRM proteins in mitochondrial function

**Abbreviations**  
4'PP: pantetheine-4'-phosphate  
ACN: Acetonitrile  
AE-MS: Affinity-enrichment mass spectrometry  
CCCP: Carbonyl cyanide m-chlorophenyl hydrazone  
LFQ: Label free quantitation  
LYRM: Leucine-Tyrosine-Arginine motif  
OXPHOS: oxidative phosphorylation  
SILAC: Stable isotope labelling with amino acids in cell culture
Abstract

NDUFAB1 is the mitochondrial acyl carrier protein (ACP) essential for cell viability. Through its pantetheine-4’-phosphate post-translational modification, NDUFAB1 interacts with members of the leucine-tyrosine-arginine motif (LYRM) protein family. Although a number of LYRM proteins have been described to participate in a variety of defined processes, the functions of others remain either partially or entirely unknown. We profiled the interaction network of NDUFAB1 to reveal associations with 9 known LYRM proteins as well as more than 20 other proteins involved in mitochondrial respiratory chain complex and mitochondrial ribosome assembly. Subsequent knockout and interaction network studies in human cells revealed the LYRM member AltMiD51 to be important for optimal assembly of the large mitoribosome subunit, consistent with recent structural studies. In addition, we used proteomics coupled with topographical heat-mapping to reveal that knockout of LYRM2 impairs assembly of the NADH-dehydrogenase module of complex I, leading to defects in cellular respiration. Together, this work adds to the catalogue of functions executed by LYRM family of proteins in building mitochondrial complexes and emphasises the common and essential role of NDUFAB1 as a protagonist in mitochondrial metabolism.
Introduction

The acyl carrier protein (ACP) is a component of the type II fatty acid synthase and polyketide synthase systems found in bacteria, chloroplasts and mitochondria [1, 2]. ACP can be found as a matrix soluble protein [3], where it interacts with the leucine-tyrosine-arginine motif (LYRM) containing family of proteins. This family is comprised of small (<15kDa), basic proteins, which harbour the LYRM towards the N-terminus followed by a highly conserved downstream phenylalanine, typically flanked by arginine residues. ACP is essential for cell viability in both yeast and human cells where it participates with individual LYRM members in diverse processes including respiratory chain assembly and Fe-S cluster biogenesis [4-6]. A pantetheine-4’-phosphate (4’-PP) prosthetic group, covalently bound at an invariant serine residue in ACP, serves as the attachment-point for acyl chains as thioesters [7]. The 4’-PP is essential for LYRM protein interactions, while its acylation increases binding affinity [8]. This suggests that the 4’-PP may act as a docking site for the interaction with the LYR motif, while the extending acyl chain acts like a ‘key in a lock’ to help stabilise interactions and facilitate its activity. It was recently shown in yeast that acylation of ACP acts as an allosteric activator of LYRM family members in respiratory chain assembly through the availability of acetyl-CoA, linking free levels of the substrate with regulation of respiration [8].

In metazoans and some fungi, mitochondrial ACP (NDUFAB1 in humans), is also integrated into mitochondrial complex I (NADH-ubiquinone oxidoreductase) where it acts as an accessory/supernumerary subunit critical for assembly [4, 7]. Uniquely, NDUFAB1 is found at two sites – on the distal end of the membrane arm and as a protrusion from the matrix arm [9]. At these sites, NDUFAB1 directly interacts with complex I subunits NDUBF9 (membrane arm) and NDUA6 (matrix arm) [10], both of
which are also members of the LYRM family. In complex I, the 4′-PP conjugated acyl-chain of NDUFA1 inserts into a hydrophobic channel present in both NDUFA6 and NDUFB9, while salt-bridge contacts are made between two conserved glutamates in NDUFA1 and the positively-charged residues of the LYRM5s [10].

The full breadth of the mammalian NDUFA1-LYRM interaction network is not yet fully known, however it is clear that NDUFA1 interacts with LYRM4 during the biogenesis of mitochondrial Fe-S clusters in a similar fashion to that seen with NDUFB9/NDUFA6 in complex I [11]. Interestingly, Fe-S biogenesis has been coupled to fatty acid synthesis via mutual interdependence on the 4′-PP [12]. Similarly, LYRM5 was recently found to interact with NDUFA1 where it is involved in deflavinating the electron transferring flavoprotein [5]. Additional LYRM members involved in the assembly of other complexes of the oxidative phosphorylation (OXPHOS) machinery including complex II [13], complex III [14] and complex V [15] have also been shown to interact with NDUFA1 [5]. The functions of a number of other LYRM family members are yet to be characterised, and new members are also being discovered.

For example, an open reading frame in the 5′ region of the transcript in the mitochondrial fission adapter MiD51 was found to encode the LYRM member AltMiD51 [16]. While AltMiD51 (also termed MIEF1-microprotein and L0R8F8) was suggested to influence mitochondrial dynamics [17], it was reported to associate with mitochondrial ribosomal subunits where it directly regulates translation [18]. However, rather than direct regulation of ribosome translation activity, AltMiD51 may instead function in ribosome assembly. Structural studies of an assembly intermediate of the large mitoribosome subunit was found to contain the known assembly factor MALSU1 bound to AltMiD51 along with NDUFA1, none of which are present in the mature ribosome [19]. The importance of AltMiD51 in ribosome function therefore requires
clarification. In this study, we dissect the interaction network of NDUFAB1 and investigate the interplay with partially or entirely uncharacterised LYRM family members. In particular, we establish roles for LYRM2 in complex I biogenesis, and AltMiD51 in MALSU1 stabilisation and ribosome assembly.
EXPERIMENTAL PROCEDURES

Tissue Culture and SILAC labelling
HEK293T cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal calf serum, 1% penicillin/streptomycin, 50 µg/ml uridine and 1 X Glutamax (ThermoFisher Scientific). Inhibition of mitochondrial translation was performed by the addition of 50 µg/ml chloramphenicol (Sigma) for up to 96 hours as indicated. For stable isotope labelling with amino acids in cell culture (SILAC), cells were cultured as previously described [4, 20]. In short, SILAC DMEM lacking lysine and arginine (ThermoFisher Scientific) was supplemented with 600 mg/ml L-proline and 10 kDa dialysed fetal calf serum (Sigma-Aldrich). Media additionally contained either ‘light’ lysine and arginine (146 µg/ml L-lysine-HCl and 42 µg/ml L-arginine-HCl) or ‘heavy’ (180 µg/ml $^{13}$C$_6^{15}$N$_2$-L-lysine-HCl and 44 µg/ml $^{13}$C$_6^{15}$N$_4$-L-arginine-HCl) (Cambridge Isotope Laboratories and Silantes respectively). Cells were incubated at 37 °C supplemented with 5% CO$_2$.

Microscopy sample preparation
Cells were seeded two days prior to experimental manipulation on HistoGrip (ThermoFisher Scientific) coated glass coverslips. Coverslips were coated according to manufacturer’s instructions. Cells were fixed with 4% paraformaldehyde in PBS for 10 min. After permeabilization with 0.5% Triton X-100 in PBS, cells were incubated with primary antibodies against Tom20 (rabbit polyclonal, 1:500, Santa Cruz, SC11415) and Flag (mouse monoclonal, 1:100, Sigma-Aldrich, F1804) for 90 min in 3% BSA, 0.02% Tween-20 in PBS at room temperature. Primary antibodies were labelled for 60 min with Alexa-Fluor-488 conjugated anti-mouse-IgG or Alexa-Fluor-568 conjugated anti-rabbit-IgG (ThermoFisher Scientific, A-11001 and A-11011,
respectively) diluted in 3% BSA, 0.02% Tween-20 in PBS at room temperature. Hoechst 33258 (1 µg/ml; Sigma) was used to stain nuclei.

**Microscopy**

Confocal microscopy was performed on a Leica TCS SP8 confocal microscope (405 nm, 488 nm, 552 nm, 647 nm; Leica Microsystems) equipped with HyD detectors using a 63×/1.40 NA oil immersion objective (HC PLAPO, CS2, Leica Microsystems). Microscopy data was recorded using the Leica LAS X Life software. Images in all experimental groups were obtained using the same settings. Z-sectioning was performed using 300 nm slices. Leica .lif files were converted to multi-colour .tiff composite stacks using custom-written Fiji/ImageJ macros.

**Image and statistical analysis**

Images were analysed using Fiji and custom-written macros [21]. Correlation coefficient R, measuring the correlation of Tom20 (mitochondria) and Flag antibody staining was measured using the Colocalization Analyzer function in Fiji [21]. All graphical representations and statistical analyses were carried out on Prism (v8.0.2, GraphPad) using two-way ANOVA with multiple comparisons.

**CRISPR-mediated knockout and verification**

Knockout cell lines were generated using the pSpCas9(BB)-2A-GFP (PX458) CRISPR/Cas9 construct (a gift from F. Zhang; Addgene, plasmid 4813 [22]) as described previously [4]. In brief, CRISPR/Cas9 gRNAs were designed for gene-disruption using CHOPCHOP software [23]. Transfections were performed using Lipofectamine LTX (ThermoFisher Scientific). Genomic DNA was isolated from each
cell line and the targeted region amplified. PCR products were ligated into pGEM4Z expression vectors [24]. Single clones were analysed by Sanger sequencing. Indels for individual alleles are summarised in Supplementary Table 1.

Mitochondrial isolation

Mitochondria were isolated as previously described [25]. Protein concentration was estimated by bicinchoninic acid assay (BCA) Pierce™ Protein Assay Kit (ThermoFisher Scientific).

Affinity enrichment

Affinity enrichment assays were performed on knockout cell lines complemented with Flag-tagged cDNA. Briefly, WT and Flag-tagged whole cell pellets were harvested in triplicate. Pellets, representing 2.5 mg total protein, were re-suspended in Solubilisation buffer (20 mM Tris-Cl pH 7.4, NaCl 50 mM, 10% glycerol, 0.1 mM EDTA) containing 1% digitonin (Merck). Pierce™ Spin Columns (ThermoFisher Scientific) were loaded with 30 µl anti-Flag M2 affinity gel (Sigma) and equilibrated with Wash buffer (20 mM Tris-Cl pH 7.4, 60 mM NaCl, 10% glycerol, 0.5 mM EDTA) and 0.1% digitonin. Following clarification, solubilised proteins and complexes were added to the columns and incubated on a rotating mixer for 2 hours at 4 °C. Columns were attached to a vacuum manifold and washed 20 times with 500 µl of Wash buffer. The columns were then incubated with 100 µl of elution buffer (Solubilisation buffer plus 150 µg/ml Flag peptide) for 15 minutes at 4 °C prior to low speed centrifugation and collection into a microcentrifuge tube. This was repeated twice and the eluates were pooled. These were combined with 5X volumes (1 ml) of ice-cold acetone and the samples centrifuged at 14000 X g for 10 minutes at 4 °C, following which the precipitated pellet
was solubilised in 8 M urea and 50 mM ammonium bicarbonate. This was sonicated in a water bath at room temperature for 15 min prior to addition of Tris(2-carboxyethyl)phosphine hydrochloride (TCEP) to a final concentration of 5 mM and chloroacetamide (Sigma Aldrich) to a final concentration of 50 mM and the sample was incubated at 37 °C while shaking. The sample was then treated with ammonium bicarbonate to dilute the urea to a final concentration of 2 M. Trypsin (1 μg; Promega) was added to the sample prior to incubation overnight at 37 °C. After this the sample was acidified with 10% trifluoroacetic acid (TFA) to a final concentration of 1%. Stage tips were generated with two plugs of 3M™ Empore™ SDB-XC Extraction Disks (Fisher Scientific) which were activated with 100% acetonitrile (ACN) via centrifugation [26]. All spins were performed at 1800 X g. The tips were washed with 0.1% TFA, 2% ACN three times. The sample was added to the stage tip and eluted with 80% ACN and 0.1% TFA. The eluates were subsequently dried down using a SpeedVac.

Mass spectrometry of SILAC treated cells
Mitochondria were isolated as above from LYRM1KO, LYRM9KO and AltMiD51KO cell lines cultured in SILAC DMEM and prepared as described previously [4]. Analysis was performed on SILAC labelled whole cell lysates of the LYRM2KO cell line. Mitochondrial protein yield and whole cell pellet protein amounts were assayed by BCA. For each triplicate set, two replicates containing 50 μg of protein from heavy labelled cells/mitochondria were mixed with the same amount of protein from light KO cells/mitochondria. The third replicate consisted of a label switch. Both whole cell and mitochondrial samples were treated as follows: samples were solubilised in 1% sodium deoxycholate, 100 mM Tris-Cl pH 8.1, 40 mM chloroacetamide and 10 mM TCEP prior to vortexing and heating for 5 min at 99 °C with 1500 rpm shaking.
Samples were then sonicated for 15 min in a room temperature water bath prior to the addition of 1 μg trypsin (Promega) and incubation overnight at 37 °C. The supernatant was then transferred to 3x 14G 3M™Empore™ SDB-RPS stage tips [26]. Ethyl acetate (99%) and 1% TFA was added to the tip before centrifugation at 3000 X g at room temperature as described [26]. Stage tips were washed first with 99% ethyl acetate and 1% TFA and then with ethyl acetate supplemented with 0.2% TFA. Samples were eluted in 80% ACN and 1% NH₄OH, and acidified to a final concentration of 1% TFA prior to drying in a SpeedVac.

Mass-spectrometry and data analysis

Peptides were reconstituted in 2% ACN, 0.1% TFA and transferred to autosampler vials for analysis by online nano-HPLC/electrospray ionization-MS/MS on either a ThermoFisher Scientific Orbitrap Q Exactive Plus, Orbitrap Fusion Lumos Trivid or Orbitrap Elite Hybrid Ion-Trap instrument as follows:

NDUFAB1Flag and yACP1Flag AE-MS were acquired on an Orbitrap Elite Hybrid Ion-Trap instrument connected to an Ultimate 3000 HPLC (ThermoFisher Scientific). Peptides were loaded onto a trap column (PepMap C18 trap column 75 μm x 2 cm, 3 μm, particle size, 100 Å pore size; ThermoFisher Scientific) at 5 μl/min for 3 min before switching the pre-column in line with the analytical column (PepMap C18 analytical column 75 μm x 50 cm, 2 μm particle size, 100 Å pore size; ThermoFisher Scientific). Peptide separation was performed at 300 nl/min using a 90 min non-linear ACN gradient of buffer A [0.1% formic acid, 2% ACN, 5% DMSO] and buffer B [0.1% formic acid in ACN, 5% DMSO]. Data was collected in Data Dependent Acquisition (DDA) mode using m/z 300-1650 as MS scan range, rCID for MS/MS of the 20 most intense ions. Other instrument parameters were: MS scan at 120,000 resolution, maximum
injection time 150 ms, AGC target 1E6, CID at 30% energy for a maximum injection time of 150 ms with AGC target of 5,000.

For LYRM2\textsuperscript{KO} whole cell SILAC measurements and LYRM2\textsuperscript{Flag} affinity enrichment, data was acquired on an Orbitrap Q Exactive Plus connected to an Ultimate 3000 HPLC. Peptides were loaded onto a trap column (Acclaim C18 PepMap nano Trap x 2 cm, 100 \(\mu\)m I.D, 5 \(\mu\)m particle size and 300 Å pore size; ThermoFisher Scientific) at 15 \(\mu\)l/min for 3 min before switching the pre-column in line with the analytical column (Acclaim RSLC C18 PepMap Acclaim RSLC nanocolumn 75 \(\mu\)m x 50 cm, PepMap100 C18, 3 \(\mu\)m particle size 100 Å pore size; ThermoFisher Scientific). The separation of peptides was performed at 250 nl/min using a non-linear ACN gradient of buffer A (0.1% formic acid, 2% ACN) and buffer B (0.1% formic acid, 80% ACN) over 280 min (LYRM2\textsuperscript{KO} whole cell) or 70 min (LYRM2\textsuperscript{Flag}). For LYRM2\textsuperscript{KO} analysis, data were collected in positive mode using Data Dependent Acquisition using m/z 375-1575 as MS scan range, HCD for MS/MS of the 12 most intense ions with charge \(\geq 2\). Other instrument parameters were: MS1 scan at 70,000 resolution (at 200 m/z), MS maximum injection time 54 ms, AGC target 3E6, normalized collision energy was at 27% energy, isolation window of 1.8 Da, MS/MS resolution 17,500, MS/MS AGC target of 2E5, MS/MS maximum injection time 54 ms, minimum intensity was set at 2E3 and dynamic exclusion was set to 15 sec. For LYRM2\textsuperscript{Flag}, data were collected in positive mode using Data Dependent Acquisition using m/z 375-2000 as MS scan range, HCD for MS/MS of the 10 most intense ions with charge \(\geq 2\). Other instrument parameters were: MS1 scan at 70,000 resolution (at 200 m/z), MS maximum injection time 118 ms, AGC target 3E6, normalized collision energy was at 27% energy, isolation window of 1.8 Da, MS/MS resolution 35,000, MS/MS AGC target of 5E5, MS/MS maximum
injection time 118 ms, minimum intensity was set at 5E3 and dynamic exclusion was set to 10 sec.

For LYRM1KO, AltMiD51KO, and LYRM9KO mitochondria, data were acquired on an Orbitrap Fusion Lumos instrument connected to an Ultimate 3000 HPLC. Peptides were loaded onto a trap column (PepMap C18 trap column 75 μm x 2 cm, 3 μm, particle size, 100 Å pore size; ThermoFisher Scientific) at 5 μl/min for 3 min before switching the pre-column in line with the analytical column (PepMap C18 analytical column 75 μm x 50 cm, 2 μm particle size, 100 Å pore size; ThermoFisher Scientific). The separation of peptides for was performed at 300 nl/min using a 90 min non-linear ACN gradient of buffer A [0.1% formic acid, 2% ACN, 5% DMSO] and buffer B [0.1% formic acid in ACN, 5% DMSO]. The mass spectrometer was operated in positive-ionization mode with spray voltage set at 1.9 kV and source temperature at 275°C. Lockmass of 401.92272 from DMSO was used. Data were collected using the Data Dependent Acquisition using m/z 350-1550 at 120000 resolution with AGC target of 5e5. The “top speed” acquisition method mode (3 sec cycle time) on the most intense precursor was used whereby peptide ions with charge states ≥ 2-5 were isolated with isolation window of 1.6 m/z and fragmented with high energy collision (HCD) mode with stepped collision energy of 30 ± 5%. Fragment ion spectra were acquired in Orbitrap at 15000 resolution. Dynamic exclusion was activated for 30 s.

Raw files were analysed using the MaxQuant platform [27] version 1.6.3.4 searching against the UniProt human database containing 20,399 reviewed, canonical entries (January 2019) and a database containing 246 common contaminants. For label-free (LFQ) AE-MS experiments, default search parameters were used with “Label free quantitation” set to “LFQ” and “Match between runs” enabled. In all cases, Trypsin/P cleavage specificity (cleaves after lysine or arginine, even when proline is present)
was used with a maximum of 2 missed cleavages. Oxidation of methionine and N-terminal acetylation were specified as variable modifications. Carbamidomethylation of cysteine was set as a fixed modification. A search tolerance of 4.5 ppm was used for MS1 and 20 ppm for MS2 matching. False discovery rates (FDR) were determined through the target-decoy approach set to 1% for both peptides and proteins. Using the Perseus platform [28] version 1.6.5.0, proteins group LFQ intensities were \( \log_2 \) transformed. Values listed as being ‘Only identified by site’, ‘Reverse’ or ‘Contaminants’ were removed from the dataset. Mitochondrial annotations were imported by matching with the Mitocarta2 dataset [29] by gene name and/or ENSG identifier. Experimental groups were assigned to each set of triplicates and the number of valid values for row group calculated. For each experiment (containing a control and an enrichment group) rows having less than 3 valid values in the enrichment group were removed and the missing values in the relevant control group imputed to values consistent with the limit of detection. Proteins identified from less than 2 unique peptides were excluded from further analysis. A modified two-sided t-test based on permutation-based FDR statistics [28] was performed between the two groups. The negative logarithmic \( p \)-values were plotted against the differences between the \( \log_2 \) means for the two groups. The significance threshold used for these experiments is noted in the relevant figure legend and supplementary tables. Non-mitochondrial proteins were removed based on their absence from MitoCarta 2.0, prior to plotting of the data. For SILAC experiments, default search parameters were used with multiplicity set to 2 (Lys8, Arg10) and “Match between runs” enabled. Using the Perseus platform [28] version 1.6.5.0, proteins group normalized H/L ratios were \( \log_2 \) transformed. Label switched samples (L/H) were inverted to KO/Control orientation prior to this step. Values listed as being ‘Only identified by site’, ‘Reverse’ or
‘Contaminants’ were removed from the dataset. Mitochondrial annotations were imported by matching with the Mitocarta2 dataset [29] by gene name and/or ENSG identifier. Experimental groups were assigned to each set of triplicates rows with < 2 valid values for each group removed. A representative MS/MS spectra for each of the 17 proteins identified from a single peptide has been included in Supplemental Figure 4. Non-mitochondrial proteins were removed based on the absence of a Mitocarta 2.0 annotation, and a one sample Student’s two-sided t-test was performed within each group. The negative logarithmic p-values were plotted against the differences between the mean ratios for each group. A significance threshold (p<0.05) was used for all experiments. log₂-transformed median SILAC ratios were mapped on homologous subunits of the respiratory chain complexes, mitoribosome assembly intermediate and mitoribosome. Applicable Protein Data Bank accession codes provided in text.

Experimental Design and Statistical Rationale

SILAC labelled whole cell and mitochondrial analyses were performed in triplicate, with a label switch as we have done previously [4, 30, 31]. The statistical approaches used to analyse the data was consistent with published quantitative SILAC analyses from our [4, 31] and other labs employing similar instrumentation and methods. The log₂ ratio values of proteins in replicates were normally distributed. The fold change threshold we used for significance was > 1.5 and a p < 0.05 following a single sample two-sided Student’s t-test. AE-MS analyses were performed in triplicate using label free quantitation and compared to untagged control cells as we [32] and others have done previously using similar instrumentation and methods. Imputation was performed only on values in controls and random values were drawn from a narrow distribution equivalent to the limit of detection of the instrument. The log₂ intensity values of the
proteins in replicate were normally distributed. Significantly-enriched proteins were
determined through a modified two-sided t-test based on permutation-based FDR
statistics [28] between the two groups, with an FDR of 1% and the s0 value determined
iteratively to exclude any enriched proteins specific to the control group.

Protein electrophoresis, in-gel activity assays and western blot analysis
SDS-PAGE samples were solubilised in LDS sample buffer (Bio-Rad) with 10 mM DTT
and separated on Tris-Tricine 10-16% gradient gel as previously described [30, 33,
34]. BN-PAGE and semi-dry western transfer was performed as described previously
[35, 36]. Blots were incubated with horseradish peroxidase coupled secondary
antibodies and ECL chemiluminescent substrate (Bio-Rad) and immunoreactive
bands detected using a Chemidoc XRS system (Bio-Rad). Bands were subsequently
quantified using Image Lab (version 5.2.1; Bio-Rad). Antibodies were purchased for
Core 1 (ThermoFisher, 16D10AD9AH5), NDUFV1 (Proteintech 11238-1-AP), MrpL45
(Sigma, HPA023373); Cox4 (Abcam, ab110261), Flag M2 Clone (Sigma-Aldrich,
F3165), ATP5A (Abcam, ab14748) and SDHA (Abcam, ab14715). Polyclonal rabbit
antibodies were previously raised in-house for NDUFAF2 and NDUFA9 [37]. Tom40
antibodies were a kind gift from M. Mori (Kumamoto). In-gel activity was performed as
previously described [38].

Pulse labelling of mtDNA-encoded subunits
Labelling of mtDNA-encoded translation products with $^{35}$S-Methionine/cysteine was
performed as previously described [39]. Isolated mitochondria were subjected to SDS-
PAGE and transferred to PVDF membranes. Radiolabelled proteins were analysed
with an Amersham Typhoon Gel and Blot Imaging System (GE Healthcare Life Sciences).

Retroviral transduction

Knockout cell lines were stably complemented via retroviral transduction as described previously [4, 40]. Briefly, cDNA fragments were amplified from HEK293T RNA and sub-cloned into pBMN-Z (Addgene, 1764) in place of the LacZ cassette. HEK293T cells were transfected with retroviral constructs using Lipofectamine LTX (ThermoFisher Scientific). Resulting virus was harvested 72 hours after infection. Viral particles were sterile filtered with 0.45 µm, PDVF membrane (Millipore) and supplied with 1 µg/µl polybrene addition to relevant knockout cell lines. Target cells were infected 24 h before the media was replaced. Stable cells were selected via addition of puromycin (2 µg/ml) for up to 72 h. Genomic DNA was isolated from candidate cell lines and the cDNA region amplified and sequence verified.

Oxygen consumption stress test

Oxygen consumption and extracellular acidification rates were measured in live cells using a Seahorse Bioscience XFe-96 Analyzer. Briefly, 25,000 cells were plated per well in culture plates treated with 50 µg/ml poly-D-lysine. For each assay cycle, there were 4 measurements of 2 minutes mix, 2 minutes wait and 5 minutes measure. The following inhibitor concentrations were used: 2 µM oligomycin, 0.5 µM carbonyl cyanide m-chlorophenyl hydrazone (CCCP), 0.5 µM rotenone and 0.3 µM antimycin A. Data were normalised using the CyQUANT assay kit (Invitrogen) and analysed using Wave2.6.0 and Prism (v8.0.2, GraphPad) software packages.
Results

The NDUFAB1 interactome

Previously, we generated a HEK293T cell line expressing a doxycycline inducible, Flag-tagged copy of NDUFAB1 (NDUFAB1Flag) harbouring CRISPR-mediated disruptions in the genomic wild-type NDUFAB1 alleles [4]. Cell viability was dependent on the addition of doxycycline (and thus NDUFAB1 expression), confirming the essential nature of the protein. To examine the role of NDUFAB1 in complex I, we also generated NDUFAB1 knockout cells rescued via the expression of the Flag-tagged yeast ACP (yACP1). Viability in this cell line was also dependent on addition of doxycycline, however complex I assembly and respiration were not restored [4]. Based on these results, we concluded that the essential role of NDUFAB1 is separate to its role as a subunit of complex I. Affinity-enrichment quantitative mass spectrometry (AE-MS) on the NDUFAB1Flag cell line, revealed a suite of mitochondrial proteins, including nine LYRM family members, to be enriched (Fig. 1A,B; Supplemental Table 1). Many of these proteins were also detected in an independent study [41]. In order to help determine which of these binding partners and associated mitochondrial processes are essential, AE-MS analysis was similarly performed on yACP1Flag cells. This revealed the enrichment of five LYRM-family proteins (Fig. 1C,D; Supplemental Table 1). Common to both was the association with LYRM4, which is involved in mitochondrial Fe-S cluster biogenesis [11, 33, 42]. Other proteins involved in Fe-S cluster biogenesis (ISCU, NFS1) were also enriched, and this process is essential in eukaryotic cells including single-celled yeasts [43, 44]. Taken together, it is likely that yACP1 complements the function of NDUFAB1 in this fundamental process, thereby leading to the survival of NDUFAB1 knockout HEK293T cells.
Analysis of changes to the proteome in LYRM\textsuperscript{KO} cell lines

LYRM2, a LYRM family member of unknown function, was also highly enriched in both the NDUFA1 and yACP1 pull-downs and thus presented itself as an obvious candidate for further characterisation. The LYRM member AltMiD51 was also highly enriched and we sought to clarify its functional importance in binding to the mitoribosome intermediate assembly [19]. We also chose to interrogate the functions of the other uncharacterised LYRM-family members LYRM1 and LYRM9, one of which (LYRM9), was enriched in the NDUFA1 pull-down. We generated and validated HEK293T KO lines for LYRM1, LYRM2, LYRM9 and AltMiD51 via CRISPR/Cas9 mediated gene-editing (Supplemental Table 2). AltMiD51 is located on a bicistronic transcript upstream of the mitochondrial fission receptor MiD51 [17] and has been found to be the main gene product [45]. Western blot analysis verified that the mutations we generated did not interfere with downstream translation of MiD51 (Supplemental Fig. 1). The validated clonal cell lines were subsequently subjected to SILAC [20] and quantitative MS analysis in order to determine the effect of each knockout on the proteome. While loss of LYRM1 or LYRM9 appeared to have a minimal impact on the overall mitochondrial proteome, significant changes were observed following disruption of LYRM2 or AltMiD51 (Fig. 2A-D; Supplemental Table 3). LYRM2\textsuperscript{KO} whole cell proteomic analysis showed reduced levels of a subset of complex I subunits and assembly factors (Fig. 2A) along with a small number of proteins involved in one-carbon metabolism, nucleotide synthesis and protein import (Supplemental Table 4). In the AltMiD51\textsuperscript{KO} cell line, subunits belonging to complex I, complex IV, the large subunit of the mitoribosome, as well as mitoribosome assembly factor MALSU1 were all reduced (Fig. 2B).
We used blue-native gel electrophoresis (BN-PAGE) and western blot analysis to validate our findings. Mitochondria were isolated from control and each of the four KO cell lines, solubilised in digitonin and separated by BN-PAGE. Western blot analysis was performed using antibodies against each of the OXPHOS complexes and the large subunit of the mitoribosome (Fig. 2E-l). Consistent with the proteomic analysis, a specific defect in the migration of the respiratory chain CI/CI\textsubscript{II}/CIV supercomplex (SC) was observed in LYRM2\textsuperscript{KO} mitochondria, resulting in a faster migrating form (indicated by SC\textsuperscript{+} in Fig. 2E). When blots were incubated with antibodies specific to complex III (Fig. 2F) and complex IV (Fig. 2G), a similar defect was present in the supercomplex. Importantly, no changes in migration were present in the free complex III (CI\textsubscript{I}I\textsubscript{I}) and complex IV holoenzymes, nor the CI\textsubscript{I}I\textsubscript{I}/CIV complex, indicating that the defect was specific to complex I. Both complex V (Fig. 2H), and a mitoribosome complex containing large subunit protein MRPL45, appeared to migrate normally in LYRM2\textsuperscript{KO} mitochondria (Fig. 2I). Quantification of the levels of supercomplex and non-assembled free complexes from digitonin-solubilised samples, as well as the total levels of individual holo-complexes following Triton X-100 solubilisation (which dissociates supercomplexes; supplemental Fig. 3), confirmed the complex I defect in LYRM2\textsuperscript{KO} mitochondria (Fig. 2J). An upregulation in complex IV was also observed and this has previously been observed in other HEK293T cells with complex I deficiency [4].

Strikingly, loss of AltMiD51 led to a decrease in the total abundance of the CI/CI\textsubscript{I}I\textsubscript{I}/CIV supercomplex and the CIV holoenzyme (Fig. 2E-G). Defects were also seen for complex V, with the apparent accumulation of the F\textsubscript{i} module (Fig. 2H) [46]. Consistent with broader defects in the OXPHOS machinery, loss of AltMiD51 led to reduced levels, but not a complete block in assembly of the mitoribosome (Fig. 2I).
Quantification of complexes (Fig. 2J) showed the strong reduction in complexes I and IV. Interestingly the levels of free CIII\textsubscript{2} was increased in AltMiD51\textsuperscript{KO} cells, possibly reflecting the reduced formation of the supercomplex. A residual complex III dimeric pool has also been observed in mitochondria depleted of the general insertase OXA1L, even when other respiratory complexes were lost [47], suggesting that this dimeric form is a more stable entity. Quantification of complex III holoenzyme levels following Triton X-100 solubilisation (supplemental Fig. 3) revealed a slight overall reduction in total levels (Fig. 2J). Total complex V levels were also similar to control. These results indicate that even though loss of AltMiD51 results in reduced levels of the mitoribosome, the individual OXPHOS complexes are differentially affected.

Given that LYRM1\textsuperscript{KO} and LYRM9\textsuperscript{KO} cells showed no strong defects at the proteomic level, nor defects in the assembly of major mitochondrial complexes responsible for OXPHOS (except for the apparent increase in complex III levels; Fig 2J), we sought to verify that the LYRM members are resident mitochondrial proteins. Flag-tagged AltMiD51, LYRM1, LYRM2 and LYRM9 were re-expressed in their respective KO lines and subjected to immunofluorescence microscopy. In all cases, mitochondrial localisation was clearly observed (Supplemental Fig. 2A and B). We conclude that LYRM1 and LYRM9 may have redundant or condition-specific roles in mitochondria. We therefore chose to focus on AltMiD51 and LYRM2 for further characterisation.

**Loss of AltMiD51 leads to mitoribosome and translation defects**

Using the proteomic data from AltMiD51\textsuperscript{KO} cells, a topographical proteomic heat-map was fitted onto the mature mitoribosome structure. This analysis clearly showed that the large subunit was specifically affected in AltMiD51\textsuperscript{KO} cells (Fig. 3A). Recently, the structure of a large subunit intermediate containing the assembly factor MALSU1, with
AltMiD51 bound to NDUFAB1, was reported [19]. When the relative abundance of individual proteins in AltMiD51KO cells were mapped onto this structure, a reduction in all proteins found within the assembly intermediate was observed, as well as a particularly strong reduction in MALSU1 (Fig. 3B). This suggests that AltMiD51 is required for stabilisation of MALSU1 during its role in mitoribosome assembly.

Topographical proteomic heat maps were also fitted onto the structures of each of the five OXPHOS complexes (Fig. 3C) [48-52]. Consistent with the BN-PAGE results (Fig. 2), complexes I and IV were most affected by loss of AltMiD51, pointing to the importance of the hydrophobic mtDNA-encoded subunits for stability and structural integrity of the assembly modules [4, 53]. Indeed, in our proteomics dataset we could only detect three mtDNA-encoded subunits from complex I (ND5) and complex IV (CO2 and CO3). Each of these were the most reduced subunits in their respective complex upon loss of AltMiD51 (Fig. 2B). While not significantly affected, subunits from complex V also trended toward protein turnover in the knockout, while subunits from complex II, the only respiratory chain enzyme not containing mtDNA-encoded subunits, were unaffected (Fig. 3C). To assess the translation of all mtDNA-encoded subunits, we performed pulse-labelling with 35S-methionine. The AltMiD51KO cells showed clear defects in the translation of subunits belonging to complex I (ND1, ND2, ND3, ND4L, ND5 and ND6), complex III (Cyt b) and complex IV (CO1, CO2 and CO3) (Fig. 3D). Interestingly, translation of complex V subunits ATP6 and ATP8 were not reduced (see discussion). Pulse labelling of cells re-expressing Flag-tagged AltMiD51 led to an increase in the translation of most subunits in comparison to AltMiD51KO cells and a concomitant reduction in ATP6 and ATP8. Partial rescue was only observed consistent with the incomplete restoration in the mitoribosome assembly (Fig. 3E), and may be due to insufficient expression levels of AltMiD51 or due to the Flag tag.
hindering activity. However, this rescue was sufficient to restore apparent steady-state supercomplex levels (Fig. 3F).

**LYRM2 is required for complete assembly of complex I**

Using the proteomic profile of LYRM2\(^{KO}\) mitochondria, we again prepared a topographical heat map of relative subunit ratios for complex I (Fig. 4A) [48]. As can be seen, the levels of subunits belonging to the NADH-dehydrogenase (N-) module were strongly reduced while subunits belonging to the membrane arm were generally unaffected (Fig. 4A and B).

In our previous study, we generated KO cell lines specifically lacking the N-module of complex I [4]. These cell lines had a faster migrating subcomplex consistent with that observed in LYRM2\(^{KO}\) mitochondria shown here. To analyse this in more detail, we investigated the assembly of the complex I holoenzyme using the stronger non-ionic detergent, Triton X-100, which dissociates supercomplexes into their holoenzyme forms [54]. Complexes from control and LYRM2\(^{KO}\) mitochondria were subjected to BN-PAGE and western blotting. Incubation with antibodies specific for the Q-module subunit NDUFA9 revealed the presence of a faster migrating species, consistent with loss of the N-module (Fig 4C, CI\(^{ΔN}\)). As it was unclear if the absence of the N-module was due to a defect in assembly, or a defect resulting in destabilisation, we tested for the presence of NDUFAF2 within this complex. NDUFAF2 is a late-stage assembly factor which acts to stabilise the nascent complex I assembly intermediate, but dissociates from the complex once the N-module is attached [37, 55]. In this way, the presence of NDUFAF2 acts as a proxy for a complex I assembly intermediate devoid of the N-module. BN-PAGE analysis revealed that NDUFAF2 was stably associated with the CI\(^{ΔN}\) assembly in LYRM2\(^{KO}\) mitochondria while absent from control
mitochondria (Fig. 4D). These results still do not discount the possibility that the Cl\textsuperscript{AN} assembly present in LYRM2\textsuperscript{KO} cells represents a breakdown product that is subsequently stabilised by NDUFAF2. We therefore sought to test if this could be observed over a time course analysis where complex I was turned over. Control and LYRM2\textsuperscript{KO} cells were cultured in the presence of chloramphenicol (CAP), which arrests mitochondrial translation. As can be seen, the complex I-containing supercomplex in control cells was depleted over time (Fig. 4E left panel) but with no detection of a disassembled complex containing NDUFAF2 (right panel). In LYRM2\textsuperscript{KO} cells, the supercomplex lacking the N-module and containing NDUFAF2 was also depleted over time, showing the same kinetics as the fully assembled supercomplex in control cells (Fig 4E,F).

It should be noted that in the absence of LYRM2, some fully assembled complex I was still present (42 ± 5%; Fig. 2J). A complex I in-gel activity assay confirmed that this complex contained an active NADH-dehydrogenase, and hence N-module (Fig. 5A). Nevertheless, a significant defect in basal and maximal respiration was evident in LYRM2\textsuperscript{KO} cells indicating the importance of LYRM2 in normal mitochondrial respiration (Fig. 5B). Finally, the KO cell line was stably complemented with a Flag-tagged copy of LYRM2 (LYRM2\textsuperscript{Flag}). BN-PAGE analysis using antibodies against the N-module subunit NDUFV1 revealed rescue of the complex I assembly defect (Fig. 5C). We used the complemented cell line for immunoprecipitation and AE-MS analysis, revealing a strong interaction between LYRM2 and NDUFAB1 (Fig. 5D; Supplemental Table 3). The lack of complex I subunits or assembly factors points to a transient interaction between these proteins during N-module assembly or integration.
Discussion

LYRM-family proteins perform a variety of important functions within the mitochondrion [5, 11, 56, 57]. These functions are executed via interactions with the soluble form of NDUFAF1 [8]. As such, hypothesising a function for any novel or otherwise uncharacterised LYRM protein solely on the presence of the tripeptide motif is unrealistic. Unbiased quantitative proteomic approaches in human KO cell lines were employed to obtain insights into potential roles of poorly characterised LYRM proteins. The absence of any notable changes in the protein profiles in either LYRM1KO or LYRM9KO cell lines is surprising, especially given their relatively uniform expression across different cell and tissue types [58]. When our KO cell lines were complemented with Flag-tagged proteins and subjected to AE-MS analysis, there was very little co-enrichment of other mitochondrial proteins with the exception of NDUFAF1 (data not shown). The lack of a strong phenotype for LYRM9 is in agreement with data from the International Knockout Mouse Consortium, which reports no phenotypic changes in a Lyrm9−/− mouse [59]. No such data was available for LYRM1. We cannot rule out discrete and measurable functions for these proteins, such as in stress-induced, aged or environment-specific circumstances. For example, knockdown of LYRM1 in P19 mouse embryonic carcinoma cells have been reported to cause mitochondrial defects including reduced ATP levels and increased ROS production along with defects in differentiation and enhanced apoptosis [60]. In the HEK293T cells used here, mitochondria and cell growth defects were not apparent. However, both LYRM1KO and LYRM9KO cell lines exhibited an increase in complex III levels which may indicate that loss of these proteins may impact on mitochondrial function leading to a compensatory mechanism in increased complex III assembly/stability. Clearer functionality however, was evident from our studies of AltMiD51 and LYRM2.
AltMiD51 was earlier identified as an alternative transcript upstream of MiD51 [17], a protein involved in mitochondrial fission [61]. AltMiD51 was also suggested to be involved in fission, with mutations in the LYR motif leading to mitochondrial elongation [17]. However, it was more recently revealed that AltMiD51 indirectly bridges the connection between NDUFA1 with a stable mitoribosome assembly intermediate via the assembly factor MALSU1 [19]. Analysis of our AltMiD51KO cells highlights the importance of AltMiD51 in optimal mitoribosome assembly. We observed reduced levels of the large mitoribosome subunit and its constituent proteins while proteins belonging to the small ribosomal subunit were not significantly reduced. Of note was the strong reduction in MALSU1 (c7orf30), a protein previously identified to be involved in biogenesis of the large mitoribosomal subunit [62-64]. Our results are therefore consistent with AltMiD51 playing a role in mitoribosome assembly rather than acting in direct translational activation [18]. Like that seen for knockout of AltMiD51 shown here, along with previous knockdown studies [18], knockdown of MALSU1 did not completely arrest mitoribosome biogenesis, nor translation of mtDNA encoded subunits [62-64]. Nevertheless, AltMiD51KO cells showed reduced levels of most mtDNA-encoded subunits and concomitant defects in the levels of respiratory chain complexes containing those proteins. Interestingly, ATP6 and ATP8 showed no translation defects during a pulse-labelling experiment, and this may reflect increased levels of their bicistronic transcript or translational activation due to the accumulation of the F1-arm, as seen in yeast studies [65]. Additionally, previous studies, including those involving knockdown of MALSU1 [62], have suggested that while a defect in mitochondrial translation will typically result in decreased levels of mitochondrial peptides, it can instead result in increased levels of some mtDNA-encoded proteins.
via apparent preferential translation of specific mRNAs [62, 66]. Future targeted studies are required to investigate this possibility.

Topographical heat-map analysis of proteomic data also gave predictive insights into the importance of LYRM2 in OXPHOS and its function in assembly of the complex I N-module. This was confirmed by BN-PAGE western blotting and in-gel activity, and agrees with extensive data in the literature which suggests that the N-module is the final module to assemble [4, 67, 68]. Given that complexes III and IV can still assemble with the complex lacking the N-module, our results also question conclusions made that the N-module integrates onto complex I prior to supercomplex assembly [67]. Our results are also consistent with earlier findings that N-module assembly is not a prerequisite for supercomplex assembly [69]. AE-MS studies were unable to elucidate other binding partners besides NDUFAB1, meaning the precise function of LYRM2 remains unclear. However, it is well established that LYRM2 is not a subunit of complex I and thus may be interacting with the N-module in a transient manner. Alternatively, it is possible that LYRM2 may be involved in the maturation of individual subunits such as incorporation of flavin mononucleotide (FMN) or Fe-S clusters that are present within the N-module. Several attempted approaches to uncover this transient interaction were unsuccessful, but may depend on the metabolic status of the cell and acylation length of NDUFAB1 [8]. Complete functionalisation of LYRM2 in complex I assembly is an avenue for future research arising from our work.
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Data Availability

The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the dataset identifier PXD013706 [70].
Figures

Figure 1. Determination of NDUFAB1 and yACP1 interactomes.

Affinity-enrichment mass spectrometry (AE-MS) were performed on (A) NDUFAB1\textsuperscript{Flag} cells and (C) yACP1\textsuperscript{Flag} cells. Following imputation of missing values for the control experiments, log\(_2\) transformed LFQ Intensities were submitted to a modified two-sided two-sample t-test with significance determined by permutation-based FDR statistics [28]. The negative logarithmic \(p\)-values are plotted against the differences between the means for the two groups. A threshold of significance of \(< 1 \% \text{ FDR @ s0} = 5\) for panel A and 2 for panel C was determined as described in the methods. Schematic representation of LYRM-family proteins interacting with (B) NDUFAB1 and (D) yACP1. Enriched proteins known to interact with LYRM proteins and their associated pathways are shown in the outer ring.

Figure 2. Proteomic and BN-PAGE analysis of select LYRM\textsuperscript{KO} lines.

Volcano plots demonstrating relative levels of proteins in (A) LYRM2\textsuperscript{KO}, (B) AltMiD51\textsuperscript{KO}, (C) LYRM1\textsuperscript{KO} and (D) LYRM9\textsuperscript{KO} cells. The mean of log\(_2\) transformed SILAC ratios for mitochondrial proteins detected for each KO against the HEK293T Control was analysed by a single sample two-sided Student’s t-test and plotted against the negative log\(_{10}\) \(p\)-value. \(n = 3\) biological replicates. Dot colours as indicated. (E-I) Mitochondria were isolated from control and KO cell lines, solubilised in 1% digitonin, subjected to BN-PAGE and immunoblotting with antibodies for (E) NDUFA9 (complex I), (F) Core 1 (complex III), (G) COX4 (complex IV), (H) ATP5A (complex V) and (I) MRPL45 (mitoribosome large subunit, LSu). Blots were re-probed with complex II subunit SDHA as loading control. SC, supercomplex; SC\textsuperscript{†}, supercomplex assembly defect; CIII\textsubscript{2}, complex III dimer, CIV, complex IV; mtRibo, mitoribosome; F\textsubscript{1}, F\textsubscript{1}-ATPase
assemblies; asterisk, non-specific complex; #, non-assembled COX4. (J) Mitochondrial complexes were quantified and shown as a percentage of control levels. SC(Cl), SC(ClII), free ClII, CV and mitoribosomal complex (mitoRibo) represent signals from digitonin-solubilised samples while total Cl, total ClII and total CIV represent signals from Triton X-100 solubilised samples. Data shown as mean ± S.E.M., n=3.

**Figure 3. Analysis of mitoribosome and translation defects in AltMiD51KO cells.**

Topographical heat maps were fitted to the (A) mature mitoribosome (PDB: 3J9M [71]) and (B) stable assembly intermediate (PDB: 500L [19]). Blue is relatively less abundant, red is relatively more abundant, white is unchanged, dark grey is not detected and yellow is AltMiD51. rRNA is depicted in gold. (C) Topographical heat-maps of the mitochondrial respiratory chain enzymes (complex I PDB: 5XTH [48], complex II PDB: 1ZOY [49], complex III dimer PDB: 1BGY [50], complex IV PDB: 5Z62 [72] and complex V PDB: 5ARE [52]). (D) SDS-PAGE and phosphorimage analysis of 35S-Met pulse labelled mtDNA subunits from control, AltMiD51KO and AltMiD51KO cells expressing AltMiD51Flag. Immunoblotting using antibodies directed to the Flag epitope and SDHA were used as controls. (E) Mitochondria were isolated from control, knockout and stably complemented cell lines, separated by BN-PAGE and assayed by western blot with antibodies against mitochondrial ribosome (MRPL45). CII serves as a loading control. (F) As for E except probing for complex I (NDUFV1). LSu, large subunit; SSu, small subunit.
Figure 4. Defects in assembly of the complex I N-module in LYRM2KO cells.

(A) Topographical heatmap of proteomic analysis of LYRM2KO fitted to the complex I structure. (B) Heatmap showing relative decrease in abundance of each complex I subunit. ‘X’ denotes undetected protein. (C-D) Mitochondria from control and LYRM2KO cells were solubilised in Triton X-100, subjected to BN-PAGE and western blot analysis using antibodies against NDUFA9 and NDUFAF2 respectively. (E) Mitochondria were isolated from cells grown in the presence of chloramphenicol (CAP) for the time indicated, solubilised in digitonin and subjected to BN-PAGE and western blotting using antibodies directed to NDUFA9 or NDUFAF2. TOM40 (detecting the TOM complex) was used as a loading control. (F) Data in (E) was quantified and shown relative to ‘0 hrs’ for each sample. Data shown as mean ± S.D., n= 3.

Figure 5. Loss of LYRM2 leads to respiration defects.

(A) Mitochondrial proteins were separated by BN-PAGE prior to NADH-dehydrogenase in-gel activity assay; *, NADH dehydrogenase activity from dihydrolipoamide dehydrogenase [73]. (B) Oxygen consumption rates of control and LYRM2KO cell lines using treatments as indicated. Data reported as mean ± S.D. n= 8-9. (C) Mitochondria from control, LYRM2KO, and LYRM2KO cells stably complemented with Flag LYRM2Flag were separated by BN-PAGE prior to western blotting and incubation with antibodies against the complex I N-module subunit NDUFV1. (D) AE-MS was performed on LYRM2Flag whole cell lysates as in Figure 1A. Threshold of significance of <1 % FDR @ s0 = 1 was used, revealing specifically interacting proteins. Biological replicates n = 3. CI, complex I. CIAN, complex I missing N-module.
References


A) LYRM2KO

B) IB: NDUFA9

C) IB: NDUFA9

D) IB: NDUFAF2

E) IB: NDUFA9

F) Relative Abundance (%)

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**Figure 5**

**A** In-gel activity

**B** Oxygen Consumption Rate (pmol O₂/min/25,000 cells)

**C** IB: NDUFV1

**D** Log2 enrichment

**Legend:**
- CI/CIII₂
- CI
- 669 kDa
- 440 kDa
- Control
- LYRM2 KO
- LYRM2 Flag
- Oligomycin
- CCCP
- Rotenone/Antimycin
- Log2 (p-value)
- Log2 enrichment