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Filipe Carvalho, Anna Spier, Thibault Chaze, Mariette Matondo, Pascale Cossart, et al.. *Listeria monocytogenes exploits the MICOS complex subunit Mic10 to promote mitochondrial fragmentation and cellular infection*. 2019. pasteur-02389987v1

HAL Id: pasteur-02389987

<https://pasteur.hal.science/pasteur-02389987v1>

Preprint submitted on 2 Dec 2019 (v1), last revised 3 Jan 2021 (v2)

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***Listeria monocytogenes* exploits the MICOS complex subunit Mic10 to promote mitochondrial fragmentation and cellular infection**

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Running title: *L. monocytogenes* targets Mic10 to fission mitochondria

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Abstract: 172 words; Text: 4847 words

23 Abstract

24 Mitochondrial function adapts to cellular demands and is affected by the ability of the organelle to
 25 undergo fusion and fission in response to physiological and non-physiological cues. We previously
 26 showed that infection with the human bacterial pathogen *Listeria monocytogenes* elicits transient
 27 mitochondrial fission and a drop in mitochondrial-dependent energy production through a
 28 mechanism requiring the bacterial pore-forming toxin listeriolysin O (LLO). Here, we performed
 29 quantitative mitochondrial proteomics to search for host factors involved in *L. monocytogenes*-
 30 induced mitochondrial fission. We found that Mic10, a critical component of the mitochondrial
 31 contact site and cristae organizing system (MICOS) complex, is significantly enriched in
 32 mitochondria isolated from cells infected with wild-type but not with LLO-deficient
 33 *L. monocytogenes*. Increased mitochondrial Mic10 levels did not correlate with upregulated
 34 transcription, suggesting a post-transcriptional regulation. We showed that Mic10 is necessary for
 35 *L. monocytogenes*-induced mitochondrial network fragmentation, and that it contributes to
 36 *L. monocytogenes* cellular infection independently of MICOS proteins Mic13, Mic26 and Mic27.
 37 Together, *L. monocytogenes* infection allowed us to uncover a role for Mic10 in mitochondrial
 38 fission.

39 **Importance.** Pathogenic bacteria can target host cell organelles to take control of key cellular
 40 processes and promote their intracellular survival, growth, and persistence. Mitochondria are
 41 essential, highly dynamic organelles with pivotal roles in a wide variety of cell functions.
 42 Mitochondrial dynamics and function are intimately linked. Our previous research showed that
 43 *Listeria monocytogenes* infection impairs mitochondrial function and triggers fission of the
 44 mitochondrial network at an early infection stage, in a process that is independent of the main
 45 mitochondrial fission protein Drp1. Here, we analyzed how mitochondrial proteins change in
 46 response to *L. monocytogenes* infection and found that infection raises the levels of Mic10, a
 47 mitochondrial inner membrane protein involved in formation of cristae. We show that Mic10 is

48 important for *L. monocytogenes*-dependent mitochondrial fission and infection of host cells. Our
49 findings thus offer new insight into the mechanisms used by *L. monocytogenes* to hijack
50 mitochondria to optimize host infection.

Introduction

Mitochondria constitute one of the most important eukaryotic organelles due to their role in several essential cellular processes, such as energy production, biosynthesis of metabolic intermediates, calcium storage and signaling, autophagy, apoptosis, as well as redox and innate immune signaling (1–3). The overall morphology and cellular distribution of the mitochondrial network are controlled by a succession of fusion and fission events referred to as “mitochondrial dynamics”. This dynamic equilibrium is fundamental to meet cellular energetic and metabolic demands and respond to stress-inducing conditions (4).

Mitochondrial dynamics are governed by a family of large GTPases with membrane-shaping properties necessary to drive fusion or fission of mitochondria (5, 6). Mitochondrial fusion requires the sequential merging of the outer (OMM) and inner mitochondrial membranes (IMM) by the action of the OMM-anchored mitofusin 1 (Mfn1) and 2 (Mfn2) proteins, and the IMM-bound optic atrophy 1 (Opa1) protein. Mitochondrial fission is mainly mediated by the cytosolic dynamin-related protein 1 (Drp1), although the endoplasmic reticulum, actin and septins also play significant roles (6, 7). The importance of mitochondrial dynamics in cell physiology is attested by numerous neuromuscular pathologies associated with genetic defects affecting the expression or activity of proteins involved in mitochondrial fusion and fission (8).

Due to their involvement in essential cellular processes, mitochondria are an attractive target for viral and bacterial pathogens (9–12). Indeed, many pathogenic bacteria were shown to modulate mitochondrial dynamics to create the ideal conditions for intracellular replication, immune evasion and persistence (11, 13–17). We previously explored how mitochondrial function and dynamics are impacted by infection with *Listeria monocytogenes* (16, 18), a facultative intracellular bacterial pathogen responsible for listeriosis, a life-threatening disease in immunocompromised individuals (19). We showed that *L. monocytogenes* causes fragmentation of the host mitochondrial network early in infection. This event requires the bacterial pore-forming toxin listeriolysin O (LLO), which

76 promotes calcium influx into the host cell (16), causing a drop of the mitochondrial membrane
 77 potential and Drp1-independent mitochondrial fission (18). *L. monocytogenes* infection has thus
 78 revealed an unconventional mechanism of mitochondrial fission, but the mechanistic details and
 79 molecular players involved in modulation of mitochondrial dynamics and function upon
 80 *L. monocytogenes* infection still remain unclear.

81 Here, we set out to increase our understanding of the impact of *L. monocytogenes* infection on host
 82 cell mitochondria and identify novel factors involved in *L. monocytogenes*-induced mitochondrial
 83 fission by performing a quantitative characterization of the mitochondrial proteome upon infection.
 84 We report that *L. monocytogenes* infection significantly upregulates the mitochondrial levels of
 85 Mic10, a core subunit of the mitochondrial contact site and cristae organizing system (MICOS)
 86 complex (20). We show that this increase in Mic10 abundance requires LLO but is not correlated
 87 with increased transcription. Finally, we demonstrate that Mic10 is necessary for *L. monocytogenes*-
 88 induced mitochondrial fragmentation, and contributes to bacterial infection.

Results

Quantitative proteomic analysis of the human mitochondrial response to *L. monocytogenes*

infection. To understand how the human mitochondrial proteome is affected by *L. monocytogenes* infection, we performed quantitative, label-free proteomic analysis of mitochondria isolated from human cells infected with *L. monocytogenes* (Figure 1A). We used HCT116 cells, a human intestinal epithelial cell line rich in mitochondria and efficiently infected by *L. monocytogenes*. Since *L. monocytogenes*-induced mitochondrial fission occurs early in infection and requires LLO (16), we performed short infections (2 h) with wild-type *L. monocytogenes* or with an LLO-deficient strain to focus on LLO-dependent processes. Mitochondria were isolated from infected and uninfected cell lysates by magnetic immunoaffinity purification and mitochondria-associated proteins were processed for LC-MS/MS analysis (Figure 1A). A total of 2,370 unique proteins were identified, with 2,039 (86%) proteins detected in every condition (Figure 1B). Among all identified proteins, 862 (36.4%) were annotated as mitochondrial (Figure 1B), which represents a good mitochondrial enrichment degree in our samples (compared to 7–8% of mitochondrial proteins in the human proteome) and a high coverage of the annotated mitochondrial proteome (53% of 1626 proteins; IMPI version Q2, June 2018). This overrepresentation of mitochondrial proteins is reflected in the results of a Gene Ontology (GO) term enrichment analysis, showing eight mitochondrial terms among the ten most enriched GO biological processes (Figure 1C).

To determine proteins displaying significantly altered levels with infection, we performed a statistical test on all identified proteins to select those whose abundance changed at least two-fold. We obtained a list of 167 proteins, of which 32 (19.1%) were detected in every condition (Figure 1D). To find proteins that could play a role in mitochondrial fission, we focused on *bona fide* mitochondrial proteins or predicted to be functionally associated and/or interact with mitochondria. According to the IMPI, 35 of the 167 differentially abundant proteins (21%) were annotated as mitochondrial (Figure 1D, Table S1). Among these are proteins involved in the mitochondrial

114 electron transport chain, such as the NADH:ubiquinone oxidoreductase subunit B2 (NDUFB2) and
 115 assembly factor DMAC2, the ubiquinol-cytochrome c reductase assembly factors 1 and 3 (UQCC1
 116 and UQCC3), the cytochrome c oxidase subunits 6C and 7B (COX6C and COX7B), and the
 117 F₁F₀ ATP synthase subunit 6.8PL (ATP5MPL). Four of these proteins become significantly more
 118 abundant in response to infection, suggesting an increased activity of the respiratory chain. Other
 119 differentially abundant proteins identified in our analysis are associated with mitochondrial
 120 translation (cysteiny- and methionyl-tRNA synthetases 2, CARS2 and MARS2; tRNA
 121 isopentenyltransferase 1, TRIT1; and mitochondrial ribosomal proteins CHCHD1 and MRPL42),
 122 metabolism of sterols (lanosterol synthase, LSS), fatty acids (hydroxyacyl-thioester dehydratase
 123 type 2, HTD2) and branched-chain amino acids (branched-chain ketoacid dehydrogenase kinase,
 124 BCKDK), regulation of mitophagy (Bcl2-associated athanogene 5, BAG5 (21, 22); FUN14 domain-
 125 containing 1, FUNDC1 (23); peroxiredoxin 6, PRDX6 (24)) and apoptosis (Bcl2-like protein 1,
 126 BCL2L1), cristae formation (MICOS complex subunit Mic10, MICOS10), and organelle transport
 127 (myosin 19, MYO19 (25)). **GO enrichment analysis of the 35 differentially abundant mitochondrial**
 128 **proteins did not reveal any statistically significant overrepresented functional pathways.** The
 129 mitochondrial abundance of 14 of the 35 proteins (40%) was significantly increased upon infection
 130 with wild-type *L. monocytogenes*, whereas it decreased for 8 proteins (23%) (Table S1).
 131 Interestingly, 7 of the 14 upregulated proteins and 4 of the 8 downregulated proteins did not display
 132 these changes upon infection with LLO-deficient bacteria (Table S1), indicating that LLO triggers
 133 such alterations in their mitochondrial abundance.

134 We further focused our attention on proteins predicted to participate in mitochondrial dynamics or
 135 membrane-remodeling processes. Interestingly, we found Mic10 among the seven proteins enriched
 136 in mitochondria in response to infection by wild-type but not LLO-deficient *L. monocytogenes*
 137 (Figure 1E). Mic10 is a core subunit of the mitochondrial contact site and cristae organizing system
 138 (MICOS) complex, a conserved IMM complex responsible for the formation of crista junctions (sites

where the IMM invaginates to form cristae) (20). Importantly, Mic10 is a small transmembrane protein, whose V-shaped membrane topology and ability to oligomerize provide it with membrane-bending properties that are fundamental for driving the formation of crista junctions (26, 27).

Mic10 is required for *L. monocytogenes*-induced mitochondrial network fragmentation. To investigate the role of Mic10 in *L. monocytogenes* infection-induced mitochondrial fission, we either lowered or raised Mic10 levels in host cells before infection and analyzed how the mitochondrial network morphology was affected. To assess the effect of Mic10 depletion, we used U2OS cells because its mitochondrial morphology is better suited for microscopy analysis compared to HCT116 cells. U2OS were transfected with non-targeting control (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs (Figure 2A), after which they were left uninfected or infected with wild-type or LLO-deficient bacteria. Cells were fixed and mitochondria immunolabeled for confocal microscopy analysis. In agreement with our previous results, si-Ctrl cells showed a typical tubular mitochondrial network, which became fragmented upon infection with wild-type, but not LLO-deficient bacteria (Figure 2B). In si-Mic10 cells, the mitochondrial morphology was similar to that of si-Ctrl cells, indicating that Mic10 knockdown does not affect mitochondrial network shape. However, unlike si-Ctrl cells, mitochondrial fragmentation was not detected in si-Mic10 cells in response to wild-type *L. monocytogenes* (Figure 2B). To obtain an unbiased, quantitative representation of these observations, we used a semi-automated morphometric tool to analyze the mitochondrial network morphology from a large number of cells, which allowed us to calculate a mitochondrial fragmentation degree per cell. Results confirmed that the mitochondrial network of si-Mic10 cells does not undergo fragmentation in the presence of *L. monocytogenes*, in contrast to si-Ctrl cells (Figure 2C). We also performed this experiment in HCT116 cells and obtained similar results (Figure S1), thus corroborating the crucial role of Mic10 in mediating mitochondrial fission in response to *L. monocytogenes* infection.

Next, we examined the effect of increased Mic10 levels in *Listeria*-dependent mitochondrial fission. We followed the experimental approach described earlier but instead of siRNA, cells were transfected with a plasmid driving the constitutive expression of a C-terminal FLAG fusion of the human Mic10 protein (Mic10-FLAG), or the empty parental plasmid as a control. We observed that constitutive expression of Mic10-FLAG resulted in a concomitant reduction of the endogenous Mic10 levels (Figure 3A). This effect was previously reported for both Mic10 and Mic60 and suggests a tight regulation of the total levels of MICOS proteins (28). Whereas control plasmid-transfected cells displayed a fragmented mitochondrial network upon infection with wild-type but not LLO-deficient bacteria (Figure 3B,C), cells expressing high levels of exogenous Mic10-FLAG showed a highly vesiculated mitochondrial network, even in the absence of infection (Figure 3B,C). Unlike Mic10 depletion, which does not affect the mitochondrial network morphology (Figures 2B,C), excessive mitochondrial levels of Mic10 cause a collapse of the mitochondrial network.

These results indicate that *L. monocytogenes* requires basal Mic10 levels to trigger mitochondrial fission in an LLO-dependent manner. Moreover, together with our proteomic data, they suggest that this mitochondrial network breakdown could be a result of increased Mic10 levels in mitochondria.

Mic10 contributes to an efficient *L. monocytogenes* cellular infection. The dynamic state of the mitochondrial network was reported to play a role in the early steps of *L. monocytogenes* cellular infection, as cells with fragmented mitochondria were less susceptible to infection, whereas cells with hyperfused mitochondria showed improved infection levels (16). Considering our results regarding the effect of Mic10 levels on the morphological status of the mitochondrial network, we wondered whether and how Mic10 levels affect *L. monocytogenes* infection. We performed gentamicin protection assays in control cells and in cells either depleted of Mic10 or overexpressing Mic10, and after infection with wild-type bacteria, we quantified the intracellular bacterial load.

189 Compared to control cells, Mic10-depleted cells were 30% less infected (Figure 4A), whereas cells
 190 overexpressing Mic10 showed a 20% increase in infection (Figure 4B). To determine if the reduced
 191 infection levels observed under Mic10 depletion were due to alterations in cellular bioenergetics
 192 elicited by defects in mitochondrial function and energy metabolism, we analyzed the mitochondrial
 193 respiratory and ATP production capacity of these cells. The oxygen consumption rate and ATP
 194 levels in si-Mic10 cells were **similar** to those in si-Ctrl cells (Figure 4C,D), in agreement with other
 195 studies (29, 30). These results suggest that the effect of Mic10 on *L. monocytogenes* infection is not
 196 caused by changes in mitochondrial energy production.

197 Depletion of Mic10 in yeast and mammalian cells results in reduced levels of MICOS proteins
 198 Mic13, Mic26 and Mic27 (26, 28, 31, 32), which interact closely with Mic10 to form the Mic10
 199 subcomplex (20). We thus assessed if the decreased *Listeria* infection associated with Mic10
 200 knockdown was a consequence of the lower abundance of Mic13, Mic26 and/or Mic27 by
 201 performing siRNA-mediated silencing of the corresponding genes before infection with wild-type
 202 *L. monocytogenes*. We confirmed that Mic10 depletion results in partial downregulation of the other
 203 three members of the Mic10 subcomplex (Figure 4E). In turn, Mic13 and, to a lesser degree, Mic27,
 204 are also necessary to sustain basal Mic10 levels (Figure 4E), in agreement with previous reports (31,
 205 32). Quantification of intracellular bacteria showed again impaired infection of Mic10-depleted cells,
 206 but revealed no difference between control cells and cells depleted for either Mic13, Mic26 or Mic27
 207 (Figure 4F). This surprising result demonstrates that none of these MICOS subunits is individually
 208 required for *Listeria* cellular infection, and therefore supports a unique role of Mic10 in this process.
 209 In agreement with this finding, besides Mic10, none of the other six subunits of the metazoan
 210 MICOS complex (Mic13, Mic19, Mic25, Mic26, Mic27, and Mic60) showed significantly changed
 211 levels with *L. monocytogenes* infection in our proteomic analysis. We investigated if increased
 212 Mic10 levels resulted from infection-driven transcriptional upregulation. Quantitative real-time PCR
 213 analysis of RNAs from cells infected or not with wild-type bacteria showed that the relative level of

214 transcripts coding for Mic10 or any other MICOS subunits remained unchanged upon
 215 *L. monocytogenes* infection (Figure S2). This result suggests that instead of upregulating Mic10
 216 transcription to elevate its protein levels, *L. monocytogenes* infection promotes an accumulation of
 217 Mic10 in mitochondria through an unknown post-transcriptional mechanism.

218 Overall, these results indicate that *Listeria* cellular infection efficiency is specifically and positively
 219 correlated with increased mitochondrial levels of Mic10.

220 Discussion

221 Bacterial pathogens like *L. monocytogenes*, *Legionella pneumophila*, *Shigella flexneri*, *Chlamydia*
 222 *trachomatis* and *Mycobacterium tuberculosis* interfere with mitochondrial dynamics to create an
 223 intracellular environment suited for survival and persistence (14–16, 18, 33–35). In particular,
 224 *L. monocytogenes* was shown to induce mitochondrial fission early in infection and cause a
 225 metabolic slowdown (16, 18), delaying mitochondria-dependent cellular responses, such as type III
 226 interferon signaling (36). Here, we employed quantitative proteomics to characterize the
 227 mitochondrial response to *L. monocytogenes* infection and search for host factors involved in
 228 *L. monocytogenes*-induced mitochondrial fission. We revealed the MICOS complex protein Mic10
 229 as a new player in this process. We showed that *L. monocytogenes* infection increases Mic10 levels
 230 in mitochondria in an LLO-dependent manner, and that Mic10 abundance is positively correlated
 231 with *L. monocytogenes*-induced mitochondrial fragmentation and host cell infection. This supports a
 232 model whereby *L. monocytogenes* infection promotes elevated mitochondrial Mic10 levels to trigger
 233 organellar fission and favor cellular infection.

234 Our proteomic approach yielded a degree of mitochondrial enrichment (36%) and mitochondrial
 235 proteome coverage (53%) comparable to those reported in other studies using varied mitochondrial
 236 isolation and mass spectrometry protocols (37–39). One of these studies explored the host
 237 mitochondrial response to *M. tuberculosis* infection, showing that virulent strains increased
 238 mitochondrial energy production and protected host cells from apoptosis, as opposed to avirulent
 239 bacteria (37). These changes were partially supported at the protein level, with upregulation of
 240 proteins involved in respiration and anti-apoptotic mechanisms, and reduced levels of proteins linked
 241 to anti-microbial response. Our proteomic data also hint that mitochondrial translation and
 242 respiration are enhanced in response to *L. monocytogenes* infection, possibly to compensate for the
 243 drop in mitochondrial membrane potential (16).

244 We chose to explore Mic10 because it represents a *bona fide* IMM protein with well-characterized
 245 membrane-shaping properties (26, 27, 40–42), and its mitochondrial levels showed an LLO-
 246 dependent upsurge with infection. We hypothesized that elevated Mic10 levels could drive
 247 deregulated IMM remodeling, resulting in mitochondrial fission. In support of this assumption, we
 248 showed that *L. monocytogenes* cannot fragment mitochondria in Mic10-depleted cells. In contrast,
 249 we observed clear mitochondrial fragmentation in cells overexpressing Mic10, even in the absence
 250 of bacteria. Others have reported that excessive Mic10 levels disrupt cristae structure (26), and that
 251 Mic10 knockdown or knockout also result in absent cristae junctions and unattached cristae stacked
 252 in the matrix (26–30, 40, 43). These phenotypes showcase the importance of Mic10 in IMM
 253 structure maintenance and suggest that *L. monocytogenes* could target Mic10 to induce IMM
 254 remodeling and trigger mitochondrial fission. We did not observe changes in the mitochondrial
 255 morphology of Mic10-depleted cells, implying that the ultrastructural defects caused by Mic10
 256 knockdown are not sufficient to elicit mitochondrial fragmentation, in contrast to the disruptive
 257 effect of excessive Mic10 levels. Consistently, knockdown of other MICOS subunits, such as Mic60
 258 (28), Mic19 and Mic25 (44), did not cause mitochondrial fragmentation, although Mic60- and
 259 Mic19-depleted mitochondria showed bulb-like enlargements (28, 44). Similar features were
 260 reported in Mic10-null yeast mitochondria (45), but we did not observe them in our si-Mic10 cells.

261 Surprisingly, the enrichment of Mic10 in mitochondria upon *L. monocytogenes* infection was not due
 262 to increased Mic10 transcription, which suggests that Mic10 accumulation in mitochondria occurs at
 263 the protein level. This could be caused by increased import or reduced turnover of Mic10 in
 264 mitochondria. As a nuclear gene-encoded protein, Mic10 is imported from the cytosol via the
 265 mitochondrial protein import machinery (46, 47). However, as other MICOS proteins are similarly
 266 imported (47), and our proteomics data showed no significant changes in their mitochondrial levels,
 267 it seems unlikely that increased Mic10 levels are caused by enhanced mitochondria import. Protein
 268 turnover in mitochondria is carried out by multiple proteases residing in the different mitochondrial

compartments (48). Interestingly, two of these proteases, Yme1L and Oma1, were reported to participate in the processing of Mic60 and Mic19, respectively (28, 44), suggesting that they may participate in Mic10 proteolysis. Future knockdown or loss-of-function experiments should clarify the involvement of Yme1L and/or Oma1 in Mic10 turnover.

Cells with fragmented mitochondria were shown to be less infected by *L. monocytogenes*, raising the hypothesis that pre-fragmented mitochondria are more resistant to the *L. monocytogenes*-induced bioenergetic slowdown (16). Here, we demonstrate that *L. monocytogenes* infection is partially impaired in cells with reduced Mic10 abundance, suggesting that Mic10-dependent mitochondrial fission induced by *L. monocytogenes* is important for subsequent cellular infection. In contrast, bacterial infection was improved by 20% in cells transfected with DNA driving Mic10 overexpression. Since not every cell overexpressed Mic10, it is possible that this margin may be higher. Further experiments using stable clones of Mic10-overexpressing cells will be helpful to confirm whether *Listeria* infection is enhanced due to increased mitochondrial Mic10 levels.

An important question is how *L. monocytogenes* manipulates events taking place inside mitochondria, even at early steps of infection when it is entering host cells or possibly still in the extracellular medium. The obvious trigger is LLO secreted by *L. monocytogenes*, which mediates Ca^{2+} influx into the host cytoplasm (16, 49, 50). Mitochondria take up Ca^{2+} from the cytosol via the mitochondrial calcium uniporter (MCU) complex (3), which includes the mitochondrial calcium uptake protein 1 (MICU1) that controls the Ca^{2+} concentration crossing the MCU channel (51). Interestingly, MICU1 was identified in our proteomic analysis, showing an apparent enrichment with *L. monocytogenes* infection in an LLO-dependent manner (Table S1). Mitochondrial Ca^{2+} efflux is mediated, among others, by the sodium/calcium exchanger NCLX (52), which can be activated by protein kinase A (PKA)-mediated phosphorylation (53). The catalytic subunit alpha of PKA (PRKACA) is one of four mitochondria-related proteins that are less abundant with infection in an LLO-dependent manner, suggesting that PKA-mediated NCLX activation is impaired during

294 *L. monocytogenes* infection. MICU1 upregulation and NCLX inhibition could result in increased
 295 mitochondrial Ca^{2+} concentration and, among other effects, a generalized collapse of the
 296 mitochondrial network (54). An investigation on the contribution of these mitochondrial proteins
 297 could clarify a role for mitochondrial Ca^{2+} uptake in *L. monocytogenes*- and possibly also Mic10-
 298 dependent mitochondrial fragmentation.

299 In conclusion, this work represents the first proteomic analysis of the mitochondrial response to
 300 *L. monocytogenes* infection and allowed us to reveal a novel actor in mitochondrial dynamics, which
 301 is specifically manipulated by *L. monocytogenes* to create the ideal setting for host cell infection.

302

303 **Materials and methods**

304 **Bacterial strains, cell lines and growth conditions.** The following *Listeria monocytogenes* strains
 305 were used in this study: wild type EGD (BUG 600), its isogenic LLO mutant EGD Δ hly (BUG 3650),
 306 and the corresponding GFP-expressing derivatives EGD-cGFP (BUG 2539) and EGD Δ hly-cGFP
 307 (BUG 2786). Bacteria were grown at 37 °C in brain heart infusion (BHI) media (Difco, BD),
 308 supplemented with chloramphenicol (7 µg/mL), when required. The following tissue culture cell
 309 lines were used in this study: HCT116 (human colorectal adenocarcinoma; ATCC CCL-247) and
 310 U2OS (human osteosarcoma; ATCC HTB-96). Cells were maintained in McCoy's 5A GlutaMAX
 311 medium (Gibco), supplemented with 1 mM non-essential amino acids (Gibco) and 10% (v/v) fetal
 312 bovine serum (FBS) (BioWest), and grown at 37 °C in a humidified 10% CO₂ atmosphere.

313

314 **Cell transfection.** For transient gene knockdown, cells were reverse transfected with siRNAs in 24-
 315 well plates, using Lipofectamine RNAiMax (Invitrogen) according to the manufacturer's
 316 instructions, except that McCoy's 5A was used as dilution medium. The medium was changed the
 317 following day and cells were assayed 48 h post-transfection. siRNA duplexes were used at the
 318 following concentrations: siRNA Universal Negative Control #1 (Sigma-Aldrich) and Mic10 (5'-
 319 CGGAUGCGGUCGUGAAGAUA^{tt}-3'; Eurofins Genomics) at 100 nM; Mic13 (Ambion, Silencer
 320 Select #s195661), Mic26 (Ambion, Silencer Select #s35601), and Mic27 (Ambion, Silencer Select
 321 #s225655) at 20 nM. For transient overexpression of Mic10, cells were seeded in 24-well plates one
 322 day before transfection with 0.5 µg of Mic10-FLAG plasmid DNA (pcDNA3.1(+)-MINOS1-DYK;
 323 GenScript, ORF cDNA clone ID OHu15514), using jetPRIME (Polyplus Transfection) according to
 324 the manufacturer's instructions. Control cells were transfected with empty plasmid DNA
 325 (pcDNA3.1(+); Invitrogen). Cells were assayed 24 h post-transfection. For immunofluorescence,
 326 cells were seeded in wells containing glass coverslips.

327

328 **Cell infections.** For cell infection assays, confluent monolayers were incubated for 1 h at 37 °C in
 329 FBS-free cell culture medium alone (non-infected cells) or inoculated with logarithmic phase
 330 bacteria (OD_{600nm} 0.6–1.0) at a multiplicity of infection (MOI, bacteria/cell) of 20 (for HCT116
 331 cells) or 50 (for U2OS cells). Medium was removed and cells were incubated for another hour (total
 332 infection time: 2 h) at 37 °C in FBS-containing culture medium supplemented with 20 µg/mL
 333 gentamicin sulfate (Sigma-Aldrich), to kill extracellular bacteria. Cells were then washed with
 334 Dulbecco's phosphate-buffered saline (DPBS; Gibco) before processing for further analyses. For
 335 immunofluorescence, cells were infected with GFP-expressing *L. monocytogenes* strains. To
 336 quantify intracellular bacterial levels, infected cells were lysed in ice-cold 0.2% (v/v) Triton X-100
 337 in DPBS, serially diluted in DPBS, and plated on BHI agar plates. Colony-forming units (CFUs)
 338 were counted after 24 h of incubation at 37 °C and bacterial numbers were normalized to the
 339 inoculum concentration.

340

341 **Mitochondrial isolation and LC-MS/MS sample preparation.** For label-free quantitative
 342 proteomic analysis of mitochondria, HCT116 cells (~5×10⁷) cultivated in 150-mm dishes were
 343 treated as outlined in Figure 1A. Cells were left uninfected (NI) or infected either with wild type (*Lm*
 344 WT) or LLO-deficient EGD (*Lm* Δ*hly*), as described above. Three independent biological replicates
 345 were prepared and analyzed for each condition. After infection, mitochondria were isolated from
 346 cells by magnetic immunoaffinity separation, using the Mitochondria Isolation Kit (human; Miltenyi
 347 Biotec). For this, cells were washed with ice-cold DPBS and scraped in ice-cold kit lysis buffer
 348 (1 mL/10⁷ cells) containing a cocktail of protease inhibitors (cOmplete, EDTA-free; Roche). Cells
 349 were then lysed in a Potter-Elvehjem homogenizer (~50 strokes), with lysis monitored by trypan blue
 350 staining. Lysates were centrifuged for 5 min at 800×g (4 °C) to pellet unbroken cells, and the
 351 supernatant was recovered for magnetic labeling and separation of mitochondria as detailed in the kit

instructions. Purified mitochondria were resuspended in urea lysis buffer (20 mM HEPES pH 8.0, 8 M urea) and protein concentration was measured with BCA Protein Assay kit (Pierce). Proteins were reduced for 30 min at 55 °C in the presence of 25 mM DTT, and then alkylated for 15 min in the dark in the presence of 50 mM iodoacetamide. Samples were diluted two-fold with 20 mM HEPES pH 8.0 and proteins digested with Lys-C (Promega) at a protease/protein ratio of 1:100 (w/w) for 4 h at 37 °C. Samples were diluted two-fold again and incubated overnight at 37 °C with trypsin (sequencing grade modified, Promega) at a 1:50 (w/w) ratio. Formic acid (FA) was added at 1% (v/v), and after 10 min on ice, samples were centrifuged for 10 min at 10,000×g to pellet any insoluble material. Peptides in the supernatant were purified in Sep-Pak C18 cartridges (100 mg; Waters), lyophilized, dissolved in solvent A [0.1% (v/v) FA in water/acetonitrile (ACN) (98:2, v/v)] and quantified by absorbance at 280 nm (NanoDrop, Thermo Fisher Scientific). Samples were analyzed by LC-MS/MS as described in Text S1.

Immunofluorescence. Cells grown on glass coverslips were fixed for 15 min at room temperature in 4% (v/v) paraformaldehyde in PBS, permeabilized for 5 min in 0.5% (v/v) Triton X-100 in PBS, and blocked for 20 min in blocking buffer [1% (w/v) BSA, 10% (v/v) goat serum in PBS]. Labelling with primary and fluorophore-conjugated secondary antibodies or dyes was performed in blocking buffer for 1 h at room temperature. Cells were washed three times in PBS between every step after fixation, except after blocking. Coverslips were mounted onto microscope slides with FluoroMount-G mounting medium (Interchim), and imaged the next day or stored in the dark at 4 °C. Primary antibodies were used as follows: rabbit polyclonal anti-C1ORF151/Mic10 (1:200; Abcam, ab84969), mouse monoclonal anti-Tom20 clone 29 (1:200; BD Transduction Laboratories), rabbit polyclonal anti-Tom20 clone F-10 (1:200; Santa Cruz Biotechnology), and mouse monoclonal anti-FLAG clone M2 (1:100; Sigma-Aldrich). Anti-rabbit and anti-mouse antibodies conjugated to Alexa Fluor 568 and 647 dyes (1:500; Molecular Probes) were used as secondary antibodies; Hoescht 33342

(Molecular Probes) was used to stain DNA. Cells were analyzed in a ZEISS AxioObserver.Z1 inverted microscope (Carl Zeiss AG) equipped with a high-speed CSU-X1 spinning-disk confocal system (Yokogawa) and an Evolve EM-CCD camera (Photometrics). Single focal plane images were acquired through a Plan-Apochromat 63×/1.4 Ph3 oil objective across multiple wavelength channels, using MetaMorph software (version 7.7.9.0). Fiji was used for image processing, including channel color selection, brightness and contrast adjustment, addition of scale bars and generation of composite images.

Mitochondrial morphology analysis. Confocal images of cells were taken from various fields of view randomly selected across the entire coverslip area and their mitochondrial morphology was analyzed using the semi-automated morphometric tool MiNA within Fiji (55). Mitochondrial networks (labeled with anti-Tom20) from individual cells were selected and digitally isolated before batch analysis. From the output data, a ratio of the values listed under the “Individuals” (number of unbranched mitochondrial particles, e.g. puncta and rods) and “Mitochondrial footprint” (mitochondrial area) parameters was calculated to determine the degree of mitochondrial fragmentation per analyzed cell. A minimum of 50 cells were analyzed per condition, in a total of three independent experiments.

Statistics. Statistical analyses were performed in Prism 8 (GraphPad Software). Unpaired two-tailed Student’s t-test was used to compare the means of two groups; one-way ANOVA was used with Tukey’s post-hoc test for pairwise comparison of means from more than two groups, or with Dunnett’s post-hoc test for comparison of means relative to the mean of a control group. Difference between group means were considered statistically significant at p-value < 0.05. Significance levels

400 are indicated as: ns, not significant ($p > 0.05$); *, $p < 0.05$; **, $p < 0.01$; ***, $p < 0.001$, ****,
401 $p < 0.0001$.

402

403 **Data availability.** Mass spectrometry proteomics data have been deposited to the ProteomeXchange
404 Consortium via the PRIDE (56) partner repository with the dataset identifier PXD014667.

Acknowledgements

We thank current and past lab members for helpful discussions; Francis Impens and Evy Timmerman (VIB Proteomics Core, University of Ghent, Belgium) for training and assistance with proteomic analyses; and Alessandro Pagliuso for critical reading of the manuscript.

This study was supported by grants to P.C. from the European Research Council (H2020-ERC-2014-ADG 670823-BacCellEpi), the Agence Nationale de la Recherche (ANR) and the French Government's "Investissements d'Avenir" program Laboratoires d'Excellence "Integrative Biology of Emerging Infectious Diseases" (LabEx IBEID, ANR-10-LABX-62-IBEID). A.S. was supported by a BioSPC doctoral fellowship from the Université Paris Diderot. P.C. is a Senior International Research Scholar of the Howard Hughes Medical Institute. F.S. is a CNRS permanent researcher.

416 **Figure legends**

417 **Figure 1 – Analysis of changes in the human mitochondrial proteome elicited by** 418 ***L. monocytogenes* infection.**

419 A) Schematic diagram of the experimental procedure used for proteomic analysis of human
420 mitochondria isolated from cells infected or not with *L. monocytogenes*.

421 B) Venn diagram of unique and common proteins identified in mitochondria isolated from
422 uninfected and *L. monocytogenes*-infected cells. Pie chart shows distribution of proteins classified as
423 mitochondrial and non-mitochondrial according to MitoMiner's IMPI database (Q2, 2018).

424 C) Top ten most prevalent Gene Ontology (GO) biological processes obtained from functional
425 enrichment analysis (PANTHER gene analysis tool) of all proteins identified in mitochondria
426 isolated from uninfected and *L. monocytogenes*-infected cells.

427 D) Venn diagram of unique and common proteins identified in mitochondria isolated from
428 uninfected and *L. monocytogenes*-infected cells with statistically significant changes in their
429 abundance (fold change>2, FDR=0.05). Pie chart shows distribution of differentially abundant
430 proteins classified as mitochondrial and non-mitochondrial according to MitoMiner's IMPI database
431 (Q2, 2018).

432 E) Heat map showing relative changes in abundance of the 35 proteins annotated as mitochondrial in
433 D. For each protein, the intensity levels in every condition (each box represents a triplicate) were
434 normalized to the mean value using Z-score. Intensity levels higher and lower than the mean are
435 indicated in green and red shades, respectively. Proteins are indicated by their gene symbol.

436

437 **Figure 2 – Mic10 knockdown blocks *L. monocytogenes*-induced mitochondrial fragmentation.**

A) Immunoblot analysis of Mic10 levels in U2OS osteosarcoma cells transfected with control negative (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs. Beta-actin protein was used as loading control.

B) Immunofluorescence analysis of U2OS cells transfected with control negative (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs, which were left uninfected (NI) or infected (MOI 50, 2h) with GFP-expressing wild type (*Lm* WT) or LLO-deficient (*Lm* Δ hly) *L. monocytogenes*. Mic10 is shown in green, mitochondria (anti-Tom20) in red, nuclei (Hoescht 33342) in blue, and bacteria (*Lm*) in yellow. White box indicates a region of the mitochondrial network magnified (2x) in the inset shown below (mitochondrial labeling only). Scale bar (top right): 10 μ m.

C) Quantitative analysis of the mitochondrial fragmentation degree in U2OS cells analyzed in B. Mitochondrial network morphology was analyzed using the morphometric ImageJ plugin tool MiNA on mitochondria-labeled images. Fragmentation degree per analyzed cell was determined by the ratio between number of individual mitochondrial particles and total mitochondrial area. Scatter plot graph shows mitochondrial fragmentation degree values for each analyzed cell (dots, n>95) and the mean (horizontal bar). Statistically significant differences were determined by one-way ANOVA with Tukey's post-hoc test: *** p<0.001; **** p<0.0001.

Figure 3 – Mic10 overexpression triggers mitochondrial fragmentation regardless of *L. monocytogenes* infection.

A) Immunoblot analysis of Mic10 levels in HCT116 cells transiently transfected with control plasmid (Ctrl) or a plasmid constitutively expressing C-terminal FLAG-tagged Mic10 (Mic10-FLAG). Both endogenous Mic10 and exogenous Mic10-FLAG were detected with an anti-Mic10 antibody. Beta-actin protein was used as loading control.

B) Immunofluorescence analysis of HCT116 cells transiently transfected with control plasmid (Ctrl) or a plasmid constitutively expressing C-terminal FLAG-tagged Mic10 (Mic10-FLAG), which were left uninfected (NI) or infected (MOI 20, 2h) with GFP-expressing wild type (*Lm* WT) or LLO-deficient (*Lm* Δ hly) *L. monocytogenes*. Mitochondria (anti-Tom20) is shown in red, Mic10-FLAG (anti-FLAG) in green, nuclei (Hoescht 33342) in blue, and bacteria (*Lm*) in yellow. White box indicates a region of the mitochondrial network magnified (2x) in the inset shown below (mitochondrial labeling only). Scale bar (top right): 10 μ m.

C) Quantitative analysis of the mitochondrial fragmentation degree in cells analyzed in B. Mitochondrial network morphology was analyzed using the morphometric ImageJ plugin tool MiNA on mitochondria-labeled images. Fragmentation degree per analyzed cell was determined by the ratio between number of individual mitochondrial particles and total mitochondrial area. Scatter plot graph shows mitochondrial fragmentation degree values for each analyzed cell (dots, n>70) and the mean (horizontal bar), and is representative of three independent experiments. Statistically significant differences were determined by one-way ANOVA with Tukey's post-hoc test: **** p<0.0001.

Figure 4 – Mic10 contributes to *L. monocytogenes* cellular infection.

A) Quantification of intracellular bacteria in HCT116 cells transfected with control negative (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs, following infection with wild type *L. monocytogenes* (MOI 20, 2h). Results are shown as mean $\pm$ SEM of three independent experiments, and represented as percentage of intracellular bacteria relative to those quantified in si-Ctrl cells. Statistically significant difference was determined by unpaired, two-tailed t-test: *** p<0.001.

B) Quantification of intracellular bacteria in HCT116 cells transiently transfected with control plasmid (Ctrl) or a plasmid constitutively expressing C-terminal FLAG-tagged Mic10 (Mic10-

485 FLAG), following infection with wild type *L. monocytogenes* (MOI 20, 2h). Results are expressed as
486 percentage of intracellular bacteria relative to those quantified in Ctrl cells, and shown as mean $\pm$
487 SEM of three independent experiments. Statistically significant difference was determined by
488 unpaired, two-tailed t-test: *** $p < 0.001$.

489 C) Oxygen consumption rate (OCR, pmol/min) of HCT116 cells transfected with control negative
490 (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs was measured in a Seahorse XF Analyzer. Electron
491 transport chain inhibitors (oligomycin, FCCP, and antimycin A/rotenone) were added at defined time
492 points to monitor specific components of cellular respiration. Results are expressed as fraction of the
493 first OCR value (basal respiration) of si-Ctrl cells, and shown as mean $\pm$ SEM of three independent
494 experiments.

495 D) Cellular ATP levels in HCT116 cells transfected with control negative (si-Ctrl) or Mic10-
496 targeting (si-Mic10) siRNAs were quantified by luminescence-based plate assay, using ATPlite
497 Luminescence Assay kit. Negative controls consist of cells treated with Triton X-100 (TX100).
498 Results are expressed as percentage of ATP levels relative to those quantified in si-Ctrl cells, and
499 shown as mean $\pm$ SEM of three independent experiments. Statistically significant differences were
500 determined by unpaired, two-tailed t-test: ns, not significant.

501 E) Immunoblot analysis of the levels of Mic10 subcomplex members (Mic10, Mic13, Mic26 and
502 Mic27) in HCT116 cells transfected with control negative siRNA (si-Ctrl), or siRNA targeting
503 Mic10 (si-Mic10), Mic13 (si-Mic13), Mic26 (si-Mic26) or Mic27 (si-Mic27). Beta-actin protein was
504 used as loading control.

505 F) Quantification of intracellular bacteria in cells treated as in E, after infection with wild type
506 *L. monocytogenes* (MOI 20, 2h). Results are shown as mean $\pm$ SEM of three independent
507 experiments, and represented as percentage of intracellular bacteria relative to those quantified in si-

Ctrl cells. Statistical significance was determined by one-way ANOVA with Dunnett's post-hoc test: ns, not significant; ** $p < 0.01$.

Figure S1 – Confirmation of Mic10-dependent *L. monocytogenes*-induced mitochondrial fragmentation in HCT116 cells.

A) Immunoblot analysis of Mic10 levels in HCT116 cells transfected with control negative (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs. Beta-actin protein was used as loading control.

B) Immunofluorescence analysis of HCT116 cells transfected with control negative (si-Ctrl) or Mic10-targeting (si-Mic10) siRNAs, which were left uninfected (NI) or infected (MOI 20, 2h) with GFP-expressing wild type (*Lm* WT) or LLO-deficient (*Lm* Δ hly) *L. monocytogenes*. Mic10 is shown in green, mitochondria (anti-Tom20) in red, nuclei (Hoescht 33342) in blue, and bacteria (*Lm*) in yellow. White box indicates a region of the mitochondrial network magnified (2x) in the inset shown below (mitochondrial labeling only). Scale bar (top right): 10 μ m.

C) Quantitative analysis of the mitochondrial fragmentation degree in cells analyzed in B. Mitochondrial network morphology was analyzed using the morphometric ImageJ plugin tool MiNA on mitochondria-labeled images. Fragmentation degree per analyzed cell was determined by the ratio between number of individual mitochondrial particles and total mitochondrial area. Scatter plot graph shows mitochondrial fragmentation degree values for each analyzed cell (dots, $n > 50$) and the mean (horizontal bar), and is representative of three independent experiments. Statistically significant differences were determined by one-way ANOVA with Tukey's post-hoc test: * $p < 0.05$; ** $p < 0.01$, *** $p < 0.001$.

Figure S2 – Transcription of Mic10 or any other MICOS complex genes is not upregulated upon *L. monocytogenes* infection.

532 Analysis of gene expression of human MICOS complex subunits Mic10, Mic13, Mic19, Mic25,
 533 Mic26, Mic27, and Mic60 in response to *L. monocytogenes* infection. HCT116 cells were left
 534 uninfected (NI) or infected (MOI 20, 2h) with wild type (*Lm* WT) or LLO-deficient (*Lm* Δhly)
 535 *L. monocytogenes*, and total cellular RNAs were isolated and used for RT-qPCR analysis. Results are
 536 shown as mean $\pm$ SEM of three independent experiments, and represented as fold change in
 537 transcript levels relative to those in NI cells.

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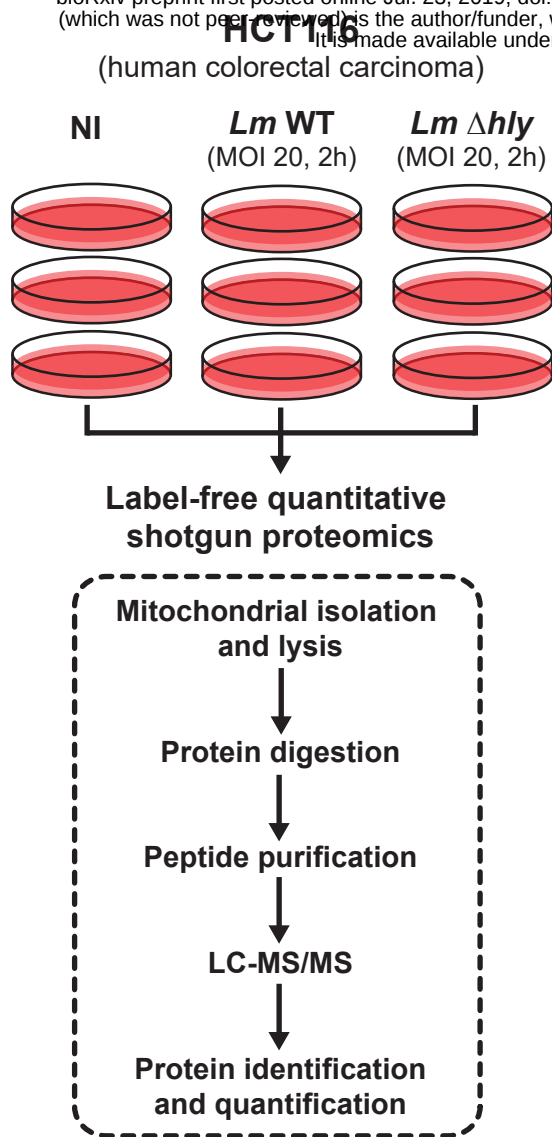
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696

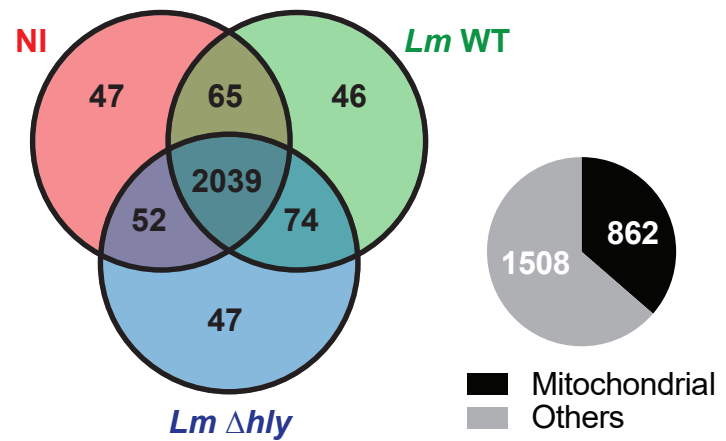
697

A



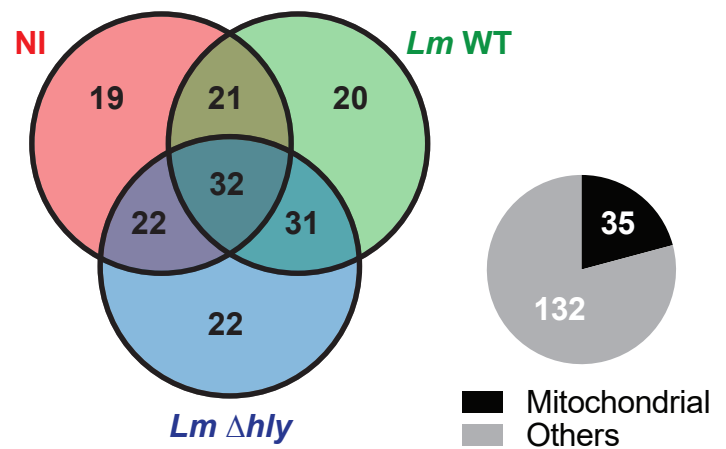
B

Total identified proteins: 2370



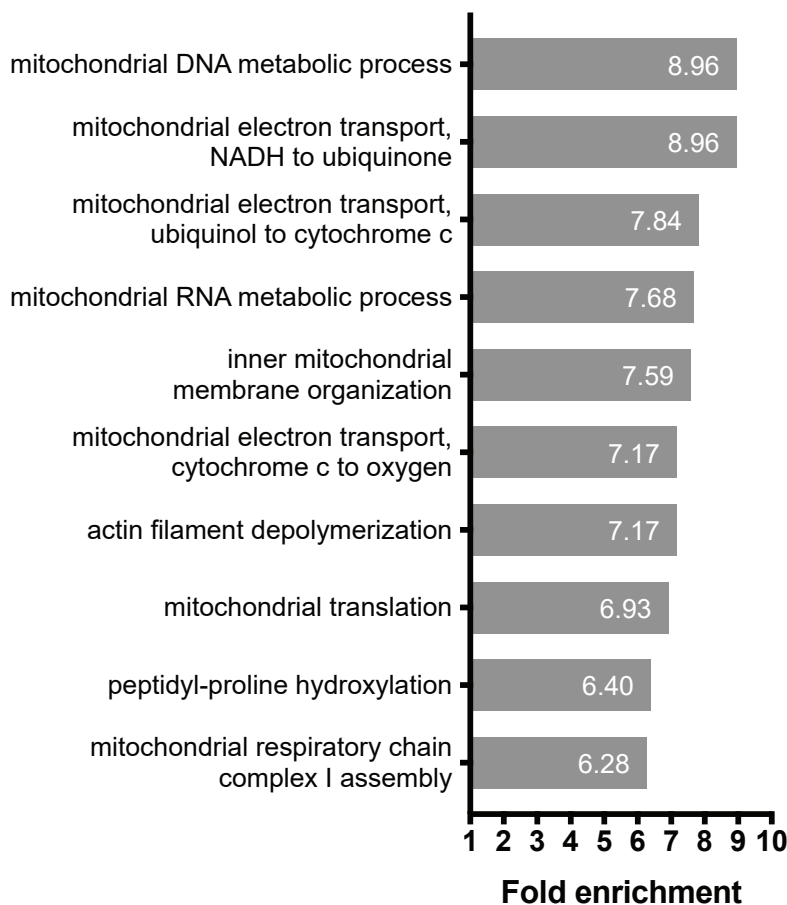
D

Differentially abundant proteins: 167
(fold change > 2; FDR = 0.05)

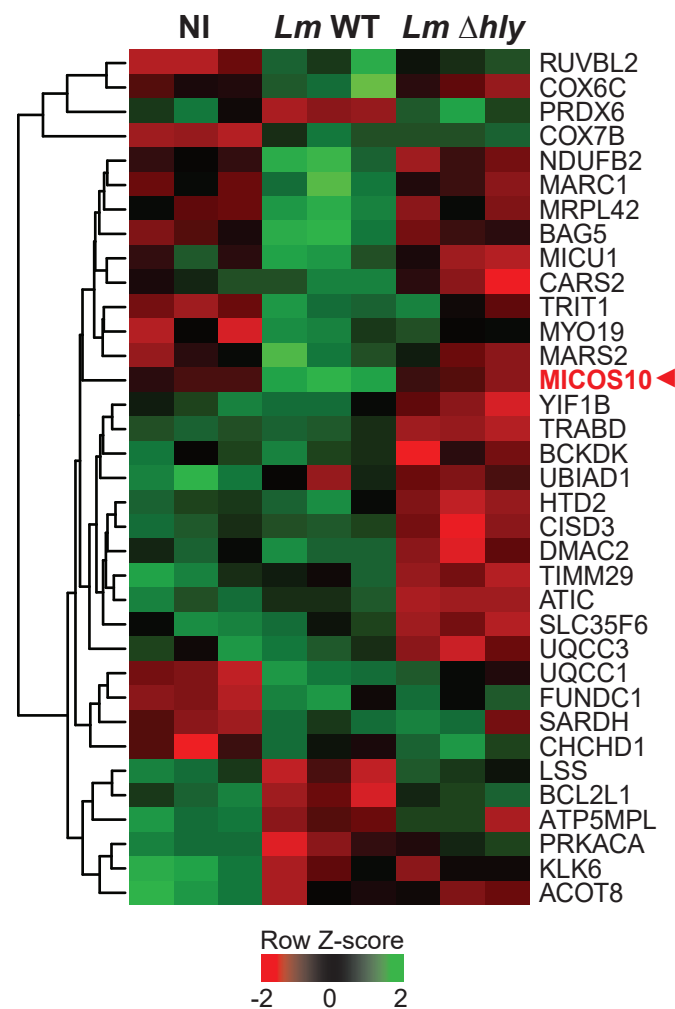


C

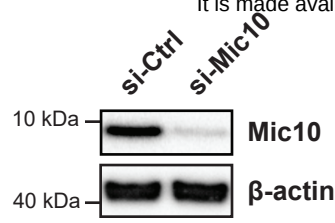
Top 10 enriched GO Biological Processes



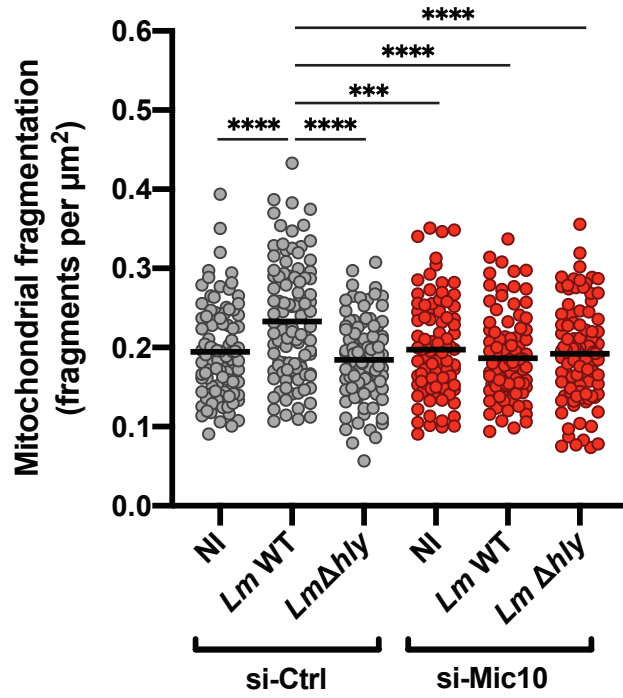
E



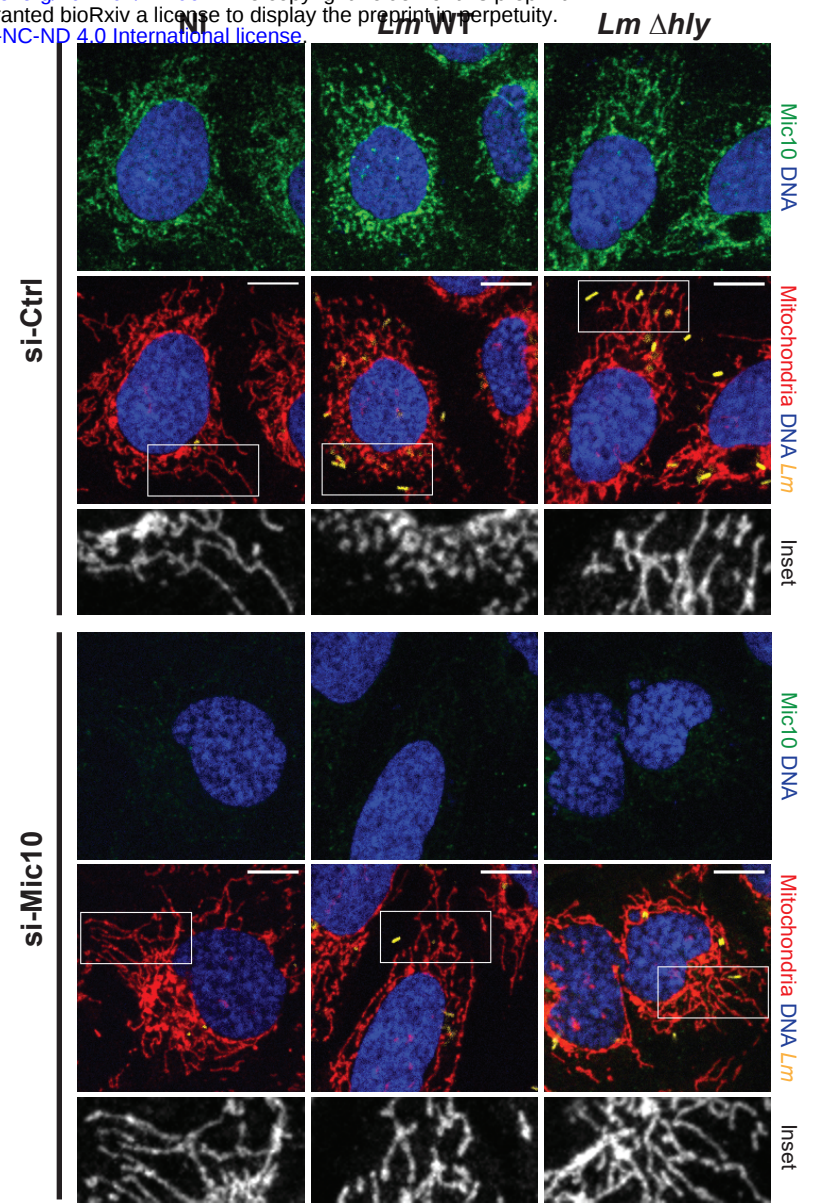
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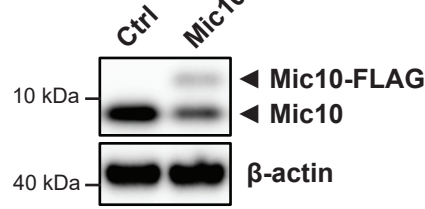
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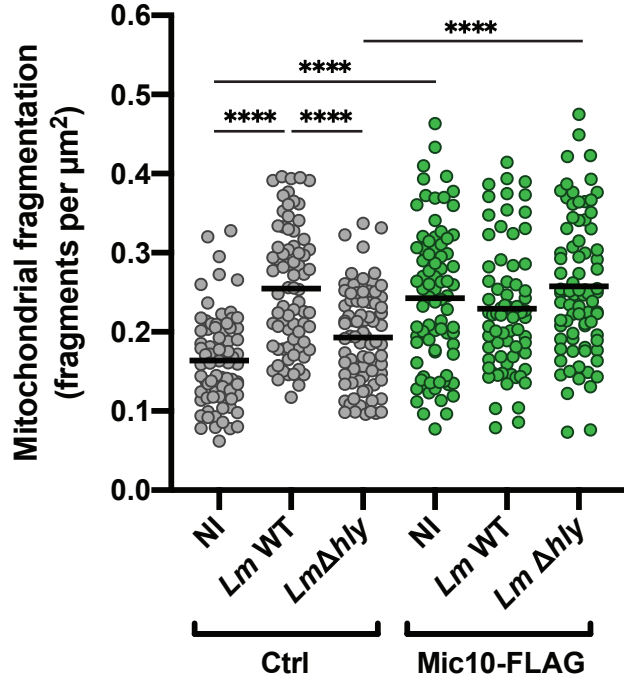
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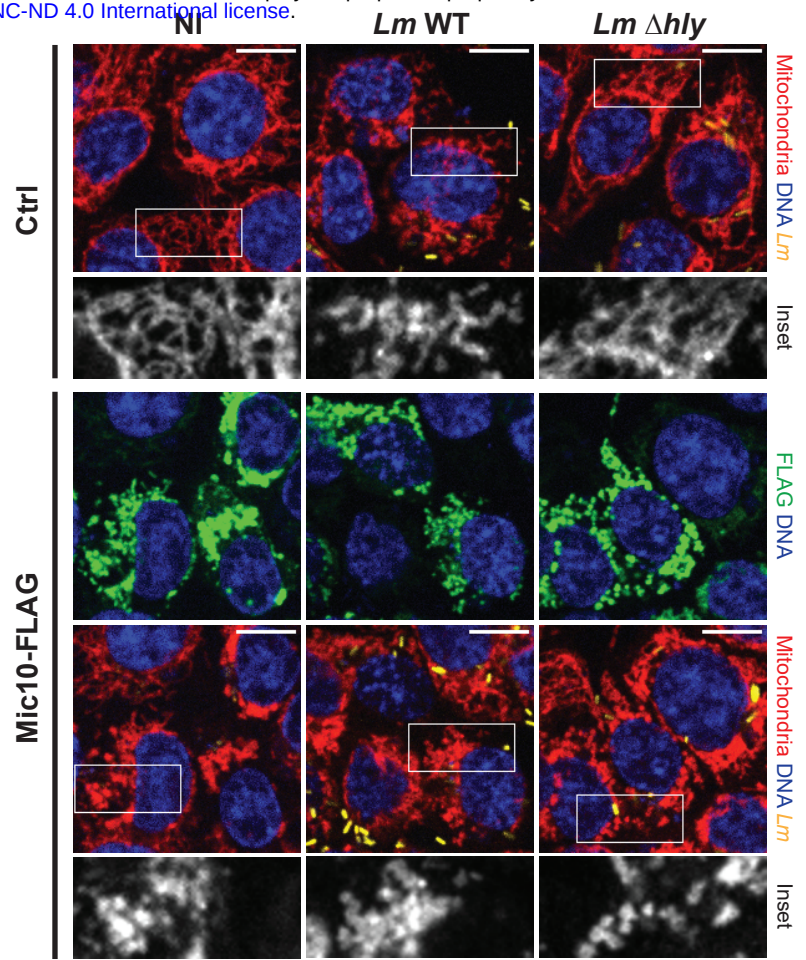
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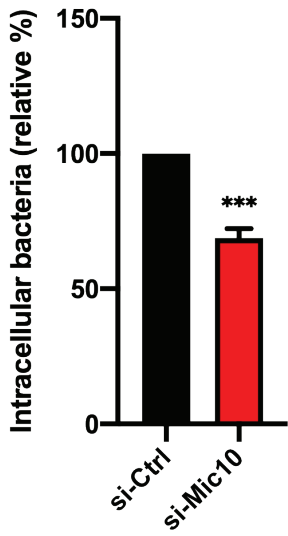
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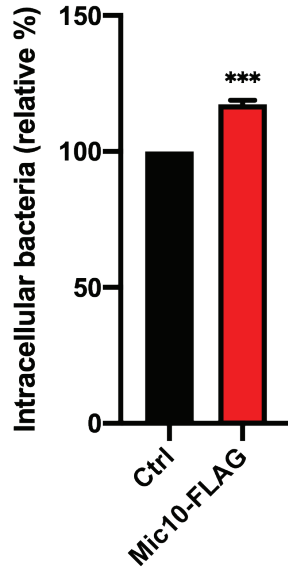
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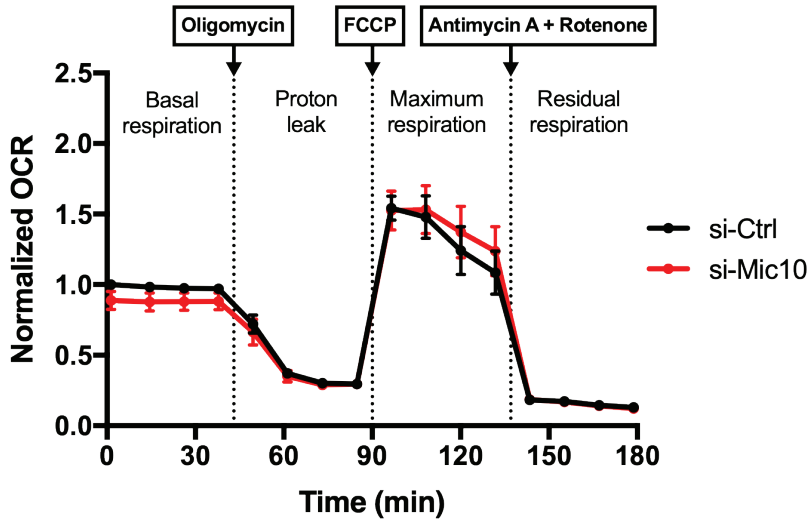
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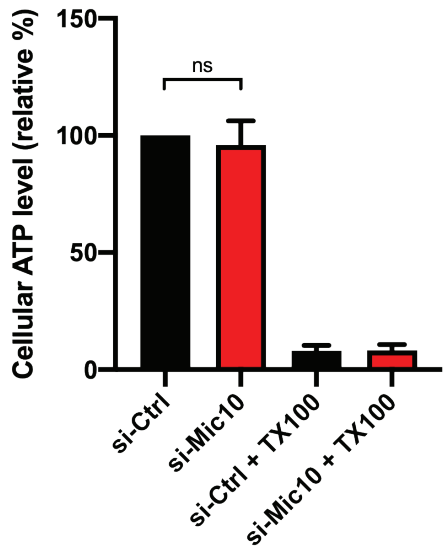
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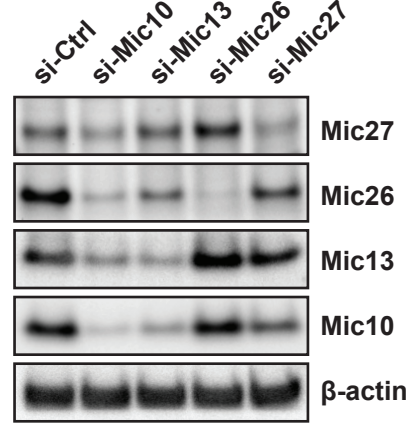
C



D



E



F

