CRISPR/Cas9-Based Edition of Frataxin Gene in Dictyostelium discoideum for 1 Friedreich's Ataxia Disease Modeling 2 3 4 Authors: 5 Gentili, Hernan G.<sup>1</sup>; Pignataro, María Florencia<sup>1</sup>; Olmos, Justo<sup>1</sup>; Pavan, Florencia<sup>2</sup>; 6 Ibáñez, Itati<sup>2</sup>; Santos, Javier<sup>1,3\*</sup> Velázquez Duarte, Francisco<sup>1,1</sup> 7 8 <sup>1</sup> Instituto de Biociencias, Biotecnología y Biología Traslacional (iB3). Departamento de 9 Fisiología v Biología Molecular v Celular, Facultad de Ciencias Exactas v Naturales, 10 Universidad de Buenos Aires. Intendente Güiraldes 2160, Ciudad Universitaria, 11 C1428EGA, Buenos Aires, Argentina. 12 13 <sup>2</sup> Instituto de Química Física de los Materiales, Medio Ambiente y Energía (INQUI-14 MAE), CONICET, FCEN, UBA, Intendente Güiraldes 2160, Ciudad Universitaria, 15 C1428EGA, Buenos Aires, Argentina. 16 <sup>3</sup> Departamento de Química Biológica, Facultad de Ciencias Exactas y Naturales, 17 18 Universidad de Buenos Aires. Intendente Güiraldes 2160, Ciudad Universitaria, 19 C1428EGA, Buenos Aires, Argentina. 20 21 \*Corresponding Authors: Francisco Velazquez Duarte at fvelazquez.ib3 @gmail.com and Javier Santos at javiersantosw@gmail.com 22 23 24 25 Running Title: Modeling Friedreich Ataxia in Dictyostelium 26 27 28 Keywords: Iron-Sulfur Cluster Assembly/ Frataxin/ Friedreich Ataxia/Rare Disease Abbreviations: ACP, acyl carrier protein; Cas9, CRISPR associated protein 9; CD, 29 30 circular dichroism; CTR, C-terminal region; CRISPR, clustered regularly interspaced 31 short palindromic repeats; DLS, dynamic light scattering; Fe-S, iron-sulfur; FA, 32 Friedreich's Ataxia; FXN, frataxin; DdFXN, D. discoideum frataxin; DTT, 33 dithiothreitol; HPLC, high-performance liquid chromatography; ISCU, iron-sulfur cluster assembly enzyme; ISD11, NFS1 interacting protein; NFS1, mitochondrial L-34 35 cysteine desulfurase enzyme; NMR, nuclear magnetic resonance; PAGE, 36 polyacrylamide gel electrophoresis; PDB, Protein Data Bank; SDS, sodium dodecyl sulfate; SEC, size exclusion chromatography. 37

- 38
- 39
- 40 Abstract

41 In this paper we describe the development of a new model system for Friedreich's Ataxia (FA) using *Dictyostelium discoideum*. We investigated the conservation of function 42 between humans and D. discoideum and showed that DdFXN can substitute the human 43 version in the interaction and activation of the Fe-S assembly supercomplex. We edited 44 the  $f_{xn}$  locus and isolated a defective mutant, clone 8, which presents landmarks of 45 frataxin deficiency, such as a decrease in Fe-S cluster-dependent enzymatic functions, 46 growth rate reduction, and increased sensitivity to oxidative stress. In addition multicel-47 lular development is affected as well as grow on bacterial lawn. 48

49 We also assessed the rescuing capacity of DdFXN-G122V, a version that mimics a hu-50 man variant present in some FA patients. While the expression of DdFXN-G122V res-51 cues growth and enzymatic activity defects, as DdFXN does, multicellular development 52 defects were only partially rescued

53 The results of the study suggest that this new model system offers a wide range of pos-54 sibilities to easily explore diverse phenotypes in FA and develop drug or treatment 55 screenings for designing and evaluating therapeutic strategies.

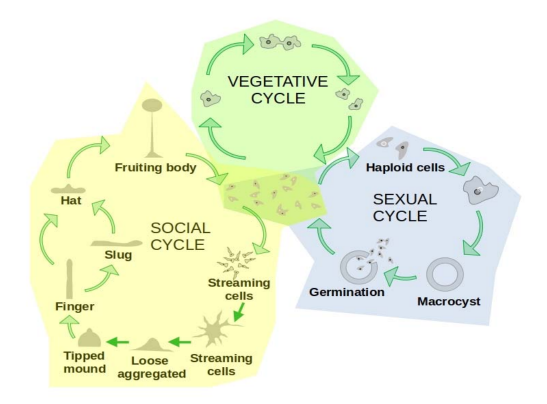
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#### 57 Introduction

Dictyostelium discoideum is a social amoeba that lives in the soil and feeds on bacteria 58 and other microbes. Dictyostelids belong to a separate branch of eukaryotic organisms, 59 distinct from plants, fungi and animals. Its cells lack a cell wall resembling animal cells 60 in organization (1). D. discoideum has become a very attractive eukaryotic non-61 mammalian model organism to study molecular mechanisms, cell physiology and 62 63 human pathology (2, 3). The studies carried out with D. discoideum have provided great 64 insights into diverse areas such as bacterial infection, immune cell chemotaxis, and 65 autophagy/phagocytosis as well as mitochondrial and neurological disorders (4).

amoeba has been widely used to study human diseases, including 66 This neurodegenerative illnesses such as Alzheimer's, Parkinson's, Huntington's, as a result 67 of which important discoveries of the pathophysiology of these pathologies have been 68 obtained (5). Furthermore, D. discoideum has been used as a model to identify drug 69 targets and discover new compounds with therapeutic potential, and these advances 70 have even served as a platform for new clinical trials (6, 7). This organism allows 71 72 studying the effect of very specific molecular alterations, from different viewpoints, combining biochemistry, structural biology, and cell biology to generate a 73 comprehensive picture concerning the outcomes of the alteration at different levels of 74 75 complexity. D. discoideum has a unique developmental life cycle among eukaryotes, 76 presenting both unicellular and multicellular phases. This allows the study of different levels of cell organization and cell-cell interaction and communication (Figure 1). 77





79 Figure 1. D. discoideum Life Cycle. D. discoideum grows as unicellular amoebae as 80 long as there are nutrients available (Vegetative Cycle). During this growth phase, 81 haploid amoeboid cells feed on bacteria and divide by a binary fission process. In 82 response to starvation, D. discoideum triggers a multicellular developmental program 83 that leads to the formation of a fruiting body bearing a mass of spores for dispersal 84 (Social Cycle). During this program, starved cells gather by chemotaxis and form an 85 aggregate. This aggregate suffers several morphogenetic changes going through 86 different stages. First, cells form a mound, enclosed within an extracellular matrix. After 87 that, the mound develops into a standing structure (finger). The finger may develop into 88 a migrating slug, or it may directly progress all the way to the culmination stages. 89 Finally, the fruiting body is formed. It will contain spores that will be released and 90 eventually germinate to produce growing amoeboid cells (8). Alternatively, upon 91 starvation, D. discoideum can go through a sexual cycle if sexually compatible cells are 92 present and other environmental conditions are met (9). 93

Recently, CRISPR/Cas9-mediated technology has been implemented to *D*. *discoideum*, allowing rapid genome editing by transiently expressing single guide RNA
(sgRNA) and Cas9 using an all-in-one vector and the generation of genomic mutants
(10–12). This possibility may accelerate gaining knowledge of specific components of
the molecular machinery underlying the complex life cycle in *D. discoideum*.

99 In addition to what was stated above, D. discoideum helps to implement the 100 "Three Rs principle" in animal research (Replacement, Reduction, and Refinement); so 101 development of *Dictyostelium*-based disease models will be highly beneficial in preliminary drug screenings (13). D. discoideum has been identified as an exceptional 102 103 model organism to study rare diseases (14, 15). More than 8000 different pathologies, 104 affecting more than 8% of the world human population, have been classified to date as 105 rare diseases; they are generally underrepresented in scientific agendas and suffer from 106 a lack of tools, model organisms and therapeutics.

107 In this context, our laboratory is interested in the development of new tools to 108 facilitate the search for therapeutic solutions for Friedreich's Ataxia (FA), a 109 neurodegenerative disease that affects 1:50000 of the population worldwide. The main

110 cause of this disease is a decrease in the expression or in the functionality of a 111 mitochondrial protein named frataxin, which is encoded in the nuclear genome.

112 Frataxin is involved in the biosynthesis of the iron-sulfur cluster (Fe-S) in the mitochondrial matrix. The deficiency of frataxin affects several enzymatic reactions that 113 depend on Fe-S clusters, which are essential cofactors involved in several enzymatic 114 115 functions. The Krebs cycle enzymes, aconitase and succinate dehydrogenase (16), and 116 the respiratory chain complexes (17, 18); critical processes such as DNA repair (19) or chemical modification of transfer RNAs (20); lipoic acid synthase, which catalyzes the 117 118 final step in the novo pathway for the biosynthesis of lipoic acid (a key coenzyme of 119 pyruvate dehydrogenase and the  $\alpha$ -ketoglutarate dehydrogenase enzymes) (21), all depend on Fe-S clusters. 120

The Fe-S assembly reaction depends on a supercomplex formed by at least five different subunits: i) the L-Cys desulfurase NFS1, a pyridoxal-phosphate (PLP)– dependent enzyme, which catalyzes the desulfurization of L-cysteine, generating as products the precursor sulfide attached as a persulfide group to a Cys residue (Cys-S-SH) and L-alanine; ii) the ISD11 protein, which is only present in eukaryotic organisms (*16*, *22*); iii) the mitochondrial acyl carrier protein (ACP), which stabilizes ISD11(*23*, *24*); iv) the scaffolding protein (ISCU); and v) frataxin.

The NFS1 dimer is stabilized by the ACP-ISD11 heterodimer (25). The Fe-S cluster assembly site is situated on the scaffolding protein ISCU and not only is frataxin the kinetic activator of the reaction, but also its surface is part of the Fe-S cluster assembly. The stoichiometry of the supercomplex is (NFS1-ACP-ISD11-FXN-ISCU)<sub>2</sub>. Its architecture is intricate as each heterodimer ACP-ISD11 also interacts with both NFS1 subunits as a bridge and two assembly sites formed by ISCU/FXN and both NFS1 chains. That is, frataxin simultaneously interacts with both NFS1 subunits and 135 ISCU (26). Even though it has been previously demonstrated that frataxin binds iron, it 136 is not clear whether it works as a chaperon of this metal ion in the context of the 137 supercomplex. It has also been shown that ISCU is able to bind this metal and that this 138 activity is crucial for Fe-S cluster assembly (27).

139 In addition to having Friedreich's Ataxia caused by mutations in frataxin gene, 140 the alteration of transcription, aberrant splicing or the presence of point mutations in 141 other proteins involved in the cluster assembly and affecting their expression or 142 functionality all result in severe human diseases. Thus, the mutation of NFS1 results in 143 an autosomal recessive mitochondrial disease characterized by a respiratory chain 144 complex II and III deficiency and multisystem organ failure (28), and the mutation of 145 ISCU results in ISCU myopathy (29), whereas the mutation of ISD11 (R68L) is 146 associated with the development of a mitochondrial genetic disorder, i.e., an autosomal recessive disease, known as Combined Oxidative Phosphorylation Deficiency 19 147 148 (COXPD19) (17).

149 The mutations of frataxin that result in FA affect the protein at different levels: 150 conformational stability (e.g., L198R, G137V, G130V) (30-32), the mitochondrial import and processing pathway (e.g., W168R and W173G) (33), iron binding affinity 151 152 (e.g., D122Y) (32, 34) or alterations at the assembly site architecture (e.g., W155R and 153 N146K and Q148R) (35, 36). In addition, the truncation of frataxin at position 193 154 (deletion of the last stretch of residues named the C-terminal region, which conforms a 155 non-periodic structure) results in a pathogenic variant (37). A very similar variant, 156 truncated at position 195, exhibits strong alterations in its internal motions and also 157 reduced conformational stability, besides a decrease in its iron binding capability (30).

Frataxin has been described as an essential protein in eukaryotic organisms as the deletion of this protein is lethal in yeast and mammalian cells (*38*). Moreover, in 160 multicellular eukaryotic organisms (plants and mice), the complete deletion of frataxin leads to early embryonic lethality (39, 40) or to the arrest of the larval stage L2 / L3 in 161 Caenorhabditis elegans (41) and reduced larval viability along with metamorphosis 162 failure in Drosophila melanogaster (42, 43). Regarding the use of cell cultures, different 163 models have been described. Since the patient's cell line (fibroblasts or lymphocytes) 164 165 does not consistently express the biochemical phenotypes associated with FA under 166 basal culture conditions, RNA interference (RNAi) strategies have been developed to reproduce partial frataxin deficiency in human and murine cell lines (44, 45). 167 168 Furthermore, "humanized" murine cell models have been developed to eliminate 169 endogenous frataxin and express frataxin with pathogenic mutations (46). A murine 170fibroblast cellular model, with the ability to deactivate frataxin transcription, was also 171 generated using the Cre /loxP recombination system. On the other hand, stem cells from 172 patients have been used, whereas the use of inducible pluripotent stem cells that can 173 mimic tissues affected by FA is under development (47, 48).

174 In a previous paper we examined the *D. discoideum* genome and found the 175 complete dotation of proteins involved in the Fe-S cluster assembly (49). Furthermore, 176 by analyzing the sequences and structure models of these proteins, we found that 177 residues located in the protein-protein interaction surfaces are highly conserved between 178 the amoeba and the human. In fact, the frataxin residues involved in FA are fully 179 conserved, with the exception of a core residue His183 of the human FXN that interacts 180 with residues located in the CTR (the last stretch of the protein), whereas in DdFXN, 181 this residue is an Arg fully exposed to the solvent, and a Tyr residue (in DdFXN) is 182 located at the position corresponding to the Trp168 in the human variant, as inferred by 183 means of a structure model of DdFXN.

In this paper, we investigated the effect of the functional deficiency in frataxin on the metabolism and physiology of the amoeba *D. discoideum*. The short duplication time, simple genetic manipulation, the simplicity of creating knockout cell lines by CRISPR/Cas9 due to its haploid genome and the simplicity of the characterization of phenotypes were conceived as key advantages of this model.

Some of the alterations observed in mammalian cells are also present in *D*. *discoideum*, such as reduced aconitase and succinate dehydrogenase or a higher sensitivity to oxidative stress. In addition, the amoeba cultures deficient in frataxin grew at a lower rate and the life cycle exhibited alterations. In addition, we have explored the rescue capacity of the constitutive frataxin expression.

These cell lines obtained by CRISPR/Cas9 will allow us to carry out therapeutic screenings of different compounds and drugs that can be used in the treatment of FA.

197 Material and Methods

# 198 Strains, Cell Culture, Plasmid Constructions, CRISPR/Cas9 Guide Design and D. 199 discoideum Transformation

200The D. discoideum strain AX2 was cultured at 22 °C on axenic culture on 201 dishes, in bacterial medium SM agar plates with Klebsiella aerogenes lawn, or in 202 Erlenmeyer flasks, in HL5 medium (http://dictybase.org/techniques/index.html). The parental plasmid used for CRISPR/Cas9 genome editing (pTM1285) (50) was kindly 203 204 provided by Dr. Kamimura. It was used to insert the DNA fragment codifying for the 205 RNA guide to target the *D. discoideum fxn* gene. For the guide preparation, a pair of 206 oligonucleotides (FwGuide553/RevGuide553) designed using CRISPOR 207 (http://crispor.tefor.net) and Breaking-Cas (https://bioinfogp.cnb.csic.es/tools/breakingcas/index.php) were adequately annealed, 208

209 and for DNA ligation, the Golden Gate strategy was used (**Table 1**). Construct checking 210 was performed by colony PCR of *E. coli* DH5 $\alpha$ , using the RevGuide553 and 211 tRNA\_seq\_3 as primers.

212 For the transformation of *D. discoideum*, AX2 cells were harvested during the 213 exponential phase of growth, washed twice in ice-cold H50 buffer (20 mM HEPES, pH 214 7.0, 50 mM KCl, 10.0 mM NaCl, 1.0 mM MgSO<sub>4</sub>, 5.0 mM NaHCO<sub>3</sub> and 1.0 mM NaH<sub>2</sub>PO<sub>4</sub>), and re-suspended in H50 buffer at a concentration of  $5 \times 10^7$  cells/ml. A 215 volume of 100  $\mu$ l of cell suspension was electroporated using 10-20  $\mu$ g of plasmid (0.75 216 217 kV, 25  $\mu$ F, twice). Cells were transferred to culture dishes containing HL5 culture 218 medium and incubated for 16 h. For clone selection, G418 antibiotic (geneticin), which 219 blocks polypeptide synthesis, was added (20  $\mu$ l/mL) and the cells were incubated for 220 48h; the resistance to G418 is conferred by the neo gene located in the pTM1285 vector. 221 After that, HL5 medium containing G418 was removed from the plastic dishes 222 and the cells were re-suspended in a volume of 400  $\mu$ L of HL5 medium; a volume of 75 223  $\mu$ L was mixed with 250  $\mu$ L of bacteria *Klebsiella aerogenes* liquid culture and plated in 224 SM agar plates. After an incubation of 3-4 days at 22 °C, plaques were observed 225 (absence of bacterial lawn).

226

## 227 D. discoideum Clone Isolation and DNA Sequencing

To identify sgRNA/Cas9 editing, the isolated genomic DNA was directly carried out by picking up cell material from the corresponding *D. discoideum* plaque using a sterilized tip. The material was re-suspended in a volume of 50  $\mu$ L of Lysis Buffer (50 mM KCl, 10 mM Tris-HCl pH 8.3, 2.5 mM MgCl<sub>2</sub>, 0.45% NP40, 0.45% Tween-20 and Proteinase K (40  $\mu$ g). The cell suspension was incubated at 20-24 °C for 20 min and then heated at 95 °C for 3 min to inactivate the protease. The cell lysate was used as template for PCR. The DNA corresponding to the *fxn* gene was amplified using KOD hot start master mix (Cat. N° 71842 Millipore) and FwDdFXNOE and RevDdFXNOE as primers (**Table 1**). The PCR cycle comprised denaturation at 95 °C for 30 s, annealing at 55 °C for 30 s, an extension at 68 °C for 30 s, and then 40 PCR cycles were programed. The amplified DNA fragments were purified using Wizard SV Gel and the PCR Clean-Up System (Cat. N° A9281 Promega), then the DNA was sequenced using the Macrogen facility (https://dna.macrogen.com/).

241

#### 242 Evaluations of the D. discoideum Growth Rate and Life Cycle Alterations

To evaluate the generation time (g), *D. discoideum* growth curves (22 °C and 180 rpm) were made using flasks (250 mL) containing HL5 culture medium (75 mL) and starting from a  $1 \times 10^5$  amoeba cells/mL inoculum. The cell cultures were grown during a week, and small aliquots (0.1 mL) were regularly taken for cell counting using a Neubauer chamber. The growth curves were analyzed, and the maximum ( $\mu$ ) was obtained from the slope when the culture grows exponentially. The generation time was calculated according to Equation 1.

$$g = \frac{\ln 2}{\mu}(1)$$

For the study of the *D. discoideum* life cycle and development, a 6-cm petri dish was prepared with 10 mL of 1.8% Oxoid L28 agar in KK2 medium (16.5 mM KH<sub>2</sub>PO<sub>4</sub>, 3.9 mM K<sub>2</sub>HPO<sub>4</sub>, pH: 6.2) and 2.0 mM MgSO<sub>4</sub>, 0.1 mM CaCl<sub>2</sub> (complete KK2).

A volume of 5-15 mL of cell culture in the log phase  $(2-5 \times 10^6 \text{ cells/mL})$  was centrifuged at 1400 rpm for 3 min under and then two washes with KK2 were carried out. Next, the cells were resuspended in complete KK2 to give  $2.5 \times 10^7$  cell/mL. After that, a volume of 1.6 mL containing  $4 \times 10^7$  cells was added to each plate, and the plates were left in a levelled table for 15 min for cell adhesion, after which the medium was 258 carefully removed by tilting the plates. The plates were incubated in humidity at 22 °C259 for 18-22 h.

260

#### 261 Aconitase and Succinate Dehydrogenase Activities Measurements

Aconitase (ACO) and succinate dehydrogenase (SDH) enzymatic activity measurements were carried out on the soluble fraction of the cellular extracts of *D*. *discoideum* AX2 and **clone 8**.

For the SDH activity assay,  $2 \times 10^5$  cells were used per reaction (in multi well plates) and cell lysis was performed by three freeze-thaw cycles. The assay was carried out according to the supplier's instructions (Abcam: ab 228560).

To measure ACO activity,  $1.5 \times 10^6$  cells were used per reaction. In this case, cell lysis was performed by means of a detergent solution included in the commercial kit. The protocol was according to the supplier's instructions (Abcam: ab 109712).

#### 271 Frataxin Variants Molecular Dynamics Simulation

272 Molecular dynamics simulations and the analysis of production runs were 273 carried out using the YASARA Structure (51) on the following hardware: Processor 274 Intel CORE i7 10,700 10th generation; SOCKET 1200 2.9 GHZ (Max 4.8 GHZ) 16 M 275 cores/threads 8/16, 2; Memory Kingston HX426C16FB3/8G HyperX2666 MHz; Disc 276 SSD Kingston A400 240GB SATA 7 mm; Linux Ubuntu 20.04 LTS 64 bit. The 277 molecular models corresponding to wild-type DdFXN and a DdFXN fragment from 278 clone 8 strain were modeled using AlphaFold2 (52) and with SwissModel at 279 https://swissmodel.expasy.org/interactive.

The coordinates were solvated, and standard minimization protocols were applied to remove steric clashes. The simulation cell was prepared by maintaining a 20 Å water-filled space around the protein with a density of 0.997 g/mL. The system (cubic

cell, periodic boundaries, and an 8.0 Å cut-off for long-range coulomb electrostatics 283 forces) was neutralized with 0.9% NaCl, and the temperature was maintained at 298 K 284 285 with a pH of 7.4. After the initial steepest descent minimization, unrestrained replicas of 286100 ns MD simulations using an ff14SB Amber force field were carried out with 2.50 fs 287 time steps (53). Snapshots were saved every 0.1 ns. The root-mean square deviation 288 (RMSD), root mean square fluctuation (RMSF), and secondary structure content were 289 calculated. Molecular dynamics simulations were performed using the Yasara Structure 290 program (54).

291

## 292 Western Blotting Analysis

For frataxin detection in cell lysates of *D. discoideum*, we used a set of nanobodies prepared in our laboratory using recombinant DdFXN as the target and phage display technology. Three panning rounds with increasing washing steps (10, 15 and 25 for rounds 1, 2 and 3, respectively) were carried out to select specific nanobodies. Since the nanobodies carry a 6xHis tag, we used anti RGS HIS6 HRP(QIAGEN) Cat.n°/ID 34450.

For the detection of DdFXN, wild-type and the G122V variant, an anti-FLAG monoclonal antibody (Cell Signaling Technology, [9A3] 8146S) was used followed by a secondary HRP-conjugated anti-mouse (ThermoFisher Cat. #31430) to specifically detect frataxin expressed from plasmidic DNA. For this detection, the FLAG of sequence DYKDDDDK was included in the C-terminal stretch of DdFXN. The HRP signal was detected using Clarity<sup>TM</sup> Western ECL subtrate (Bio-Rad #1705060).

#### 305 Table 1. Oligonucleotides Used in this Research

Name	Oligonucleotide Sequence <sup>1</sup>
FwDdFXNOE <sup>1</sup>	AAagatctATGATTTTCAACTTTTTAAACAAAGC

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RevDdFXNOE <sup>1</sup>	GTactagtAATTTCCATATCATATTTACAAAGTG
FwGuide 553 <sup>2</sup>	agcaTGGTGGTCATCACCATTGAG
RevGuide 553 <sup>2</sup>	aaacCTCAATGGTGATGACCACCA
tRNA_seq_3 <sup>3</sup>	GCTCGATTAGCTCAGTCGGCAG

306

<sup>1</sup> Upper-case letter indicates match and lower-case mismatch with target template.
 308

## 309 Oxidative Stress Sensitivity Assay

310 To assess sensitivity to oxidative stress, cell viability was determined upon  $H_2O_2$ 311 treatment using a crystal violet assay described by Feoktistova et al. (55). In summary, 312 100000 cells were seeded in a 24-well plate and allowed to adhere for 1h. Then the 313 medium was removed and fresh HL5 medium, or supplemented with 2 mM H<sub>2</sub>O<sub>2</sub>, was 314 added to each well. Since adherent cells detach from cell culture plates during cell 315 death, the wells were gently washed to remove the dying cells after the corresponding 316 time of treatment. Remaining cells were fixed and stained with crystal violet. Culture 317 plates were dried, and crystal violet was measured after solubilization with acetic acid 318 (A570 nm) as an estimate of the remaining cells. Viability was calculated referenced to 319 the crystal violet at time 0 for each strain. The experiment was performed at least 3 320 times for each strain and contained 4 replicates of each time point.

# 322 *Results*323

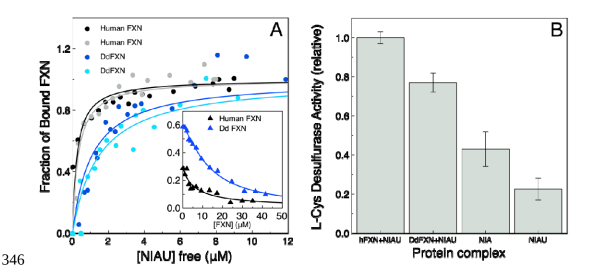
# 324 DdFXN Can Substitute Human Orthologue in the Supercomplex for Fe-S cluster 325 Assembly

Since we aimed to establish *D. discoideum* as a model organism for FA using DdFXN to explore its roles within the cell and to extrapolate it to the human context, we first tested the ability of DdFXN to substitute the human orthologue in the context of the supercomplex.

As proof of functional conservation, we assessed the ability of DdFXN to interact with the human supercomplex NIAU (NFS1/ACP-ISD11/ISCU)<sub>2</sub>. We carried out fluorescence anisotropy assays using a variant of DdFXN (DdFXN\_C35A) or human FXN S202C labelled with Texas-red.

The labelled frataxin variant, in the presence of saturating ISCU concentration, was incubated with increasing NIA concentrations. As it can be seen in **Figure 2A**, the DdFXN variant was able to bind to the human NIAU in a similar fashion as the mature form of the human FXN does, although with a lower apparent affinity. It is worthy of note that the wild-type DdFXN was able to compete with the labeled-human-frataxin variant (**Figure 2A**, **inset**), suggesting that the same binding site is involved for both proteins.

We then tested whether this binding is functional and if DdFXN is also able to act as the kinetic activator of L-Cys-desulfurase activity in a human supercomplex context. **Figure 2B** shows that the interaction of DdFXN with the human supercomplex was productive and Cys-desulfurase activity increased upon the addition of human FXN or DdFXN.



347 Figure 2. In Vitro Substitution of Human FXN by DdFXN. (A) DdFXN interaction 348 with the human supercomplex L-Cys desulfurase. Texas red-Labeled DdFXN (blue and cyan correspond to two independent experiments) or human FXN (black and gray 349 350 correspond to two independent experiments) were incubated with increasing 351 concentrations of the NIAU subcomplex and fluorescence anisotropy was monitored. A 352 fraction of bound frataxin was plotted as a function of the free NIAU complex. The 353 inset shows the competition with non-labeled proteins of a preformed supercomplex in 354 which human FXN was labeled with Texas red followed by fluorescence anisotropy. 355 The competition was carried out using non-labeled human FXN (black triangles) or 356 non-labeled DdFXN (blue triangles); the fraction of bound frataxin was plotted as a 357 function of the added FXN variant. (B) In vitro activity of the human supercomplex 358 activated by human FXN or DdFXN. The activity corresponding to NIA or the NIAU 359 complexes without frataxin are shown for reference. 360

#### 361 Fxn Locus Can Be Easily Edited in D. discoideum with CRISPR/Cas9 Technology

362 As noted above, we aimed to generate a new experimental model to study the 363 biochemical consequences of alterations in the functionality of frataxin in the cell. We 364 intended to get mutant strains that were highly deficient in frataxin so we would be able 365 to explore strategies for reestablishing homeostasis and to evaluate the rescue capacity 366 of wild-type and disease-associate variants. With this aim in mind, we carried out 367 CRISPR/Cas9 editing of D. discoideum endogenous locus, using a guide RNA targeted at Trp146. This residue, corresponding to the human Trp155, is extremely conserved in 368 frataxin along the tree of life because of a functional role (49). When looking at the 369 370 structure of the human supercomplex (NFS1/ACP-ISD11/ISCU/FXN)<sub>2</sub>, Trp155 is 371 located near the docking surface of ISCU and at Van der Waals distance of its [2Fe-2S]

372 assembly site, and from Leu386 of NFS1, which hints at its importance. Trp155 belongs

373 to a highly conserved region that contains the Motif 1 described by Gibson et al. (56).

- Hence, it would be more likely to obtain deleterious mutations in this region. Using pTM1285, an all-in-one vector, to express the corresponding guide RNA, we
- 376 obtained hundreds of clones and selected 14 presumably edited clones (Figure S1).

After the isolation of these clones, the genomic DNA corresponding to the *fxn* gene was successfully amplified and sequenced for 13 clones. The analysis of the genomic DNA sequences showed a variety of mutations in the protein as the result of CRISPR/Cas9 editing and subsequent DNA repair (**Table 2, Figure S1**).

381

# 382 Table 2. Summary of CRISPR/Cas9 Edition of *fxn* Locus in *D. discoideum* in 383 Sequenced Clones.

384

Mutation	Number of
	Clones
Premature STOP codon	$9(7)^{1}$
Small Insertion	2
Point Mutation	1
Silence Mutation	1

385

386<sup>1</sup> Seven different editions.

387

# 388 A D. discoideum Clone Lacking Frataxin is Viable

389 The amino acid sequence analysis of D. discoideum clones obtained by CRISPR/Cas9

390 genome editing (e.g., clone 8) indicated that fxn gene functionality of DdFXN can be

391 obliterated without being lethal for the amoeba, at least in the experimental conditions

392 assayed in this paper.

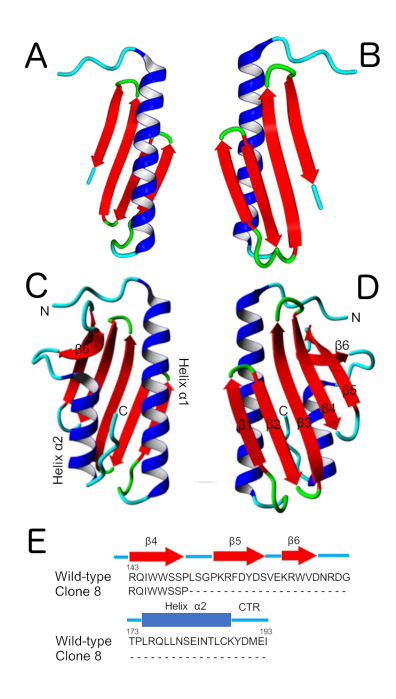
Edition in **clone 8** produces a large truncation of the protein chain because of one base pair deletion generating a frame shift, and premature stop codon (**Figure S2**). This truncation eliminates a stretch that includes more than 40% of the sequence, which may completely destabilize the structure of the remaining protein fragment, comprising residues 81-150 from DdFXN, using the numbering of the DdFXN precursor (**Figures 34 and B**). The truncation eliminated part of strand beta 4, strands beta 5 and 6, alpha helix 2 and the CTR.

We studied the molecular motions of this fragment by molecular dynamics simulations. Even though the global conformation of the fragment persisted during 100-200ns MDs simulations, significantly higher RMSD values and atomic fluctuations were observed, compared to the wild-type (**Figures 4A and B**). Alpha helix 1 presented high distortions, establishing non-native contacts with the beta sheet (**Figures 4E and F**) and higher internal motions compared to the wild-type frataxin, as judged by the fluctuations.

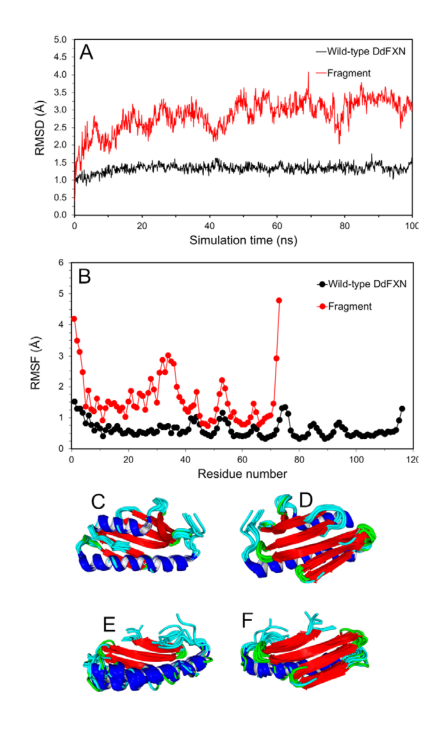
407 All this suggests that the fragment should be highly dynamic and more likely 408 easily degraded in a cellular environment. However, we cannot rule out that the 409 fragment might acquire a frataxin-like secondary structure and even some kind of 410 packing, although residual activity is highly unlikely.

AlphaFold2 predictions suggested a conformation for a putative dimeric fold of the fragment, in which the predicted accessible apolar surface for each monomer is highly reduced by the interaction, and the remanent Cys residue (Cys112) might establish an intermolecular disulfide bond stabilizing the hypothetical dimeric structure (**Figure S3**). Experiments will be done to evaluate this possibility.

416



418 **Figure 3. The Alterations of the Frataxin Structure.** (A) and (B) Two views of the 419 truncated frataxin DdFXN\_81-150 (clone 8). The topology shown is only for 420 visualization purposes. (C) and (D) Two views of the wild-type DdFXN model. DdFXN 421 models were constructed using AlphaFold2 (52). AlphaFold formed a disulfide bond 422 between Cys112 and Cys187, whereas other programs like the Swiss-model did not 423 establish the -S-S- bond. (E) Amino acid sequence truncation predicted from the DNA 424 sequences for **clone 8 (Figure S2** shows the complete sequence alignment).



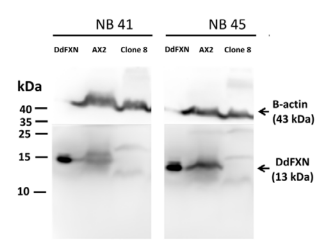
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- 428

429 Figure 4. Molecular Dynamics Simulations of the DdFXN Variant. (A) Root-mean-430 square deviation (RMSD) along the simulations (calculated for the alpha carbon atoms). 431 (B) Root-mean-square fluctuations (RMSF) of the alpha carbon atoms. Snapshots of the 432 wild-type variant (C and D) and the fragment (E and F). Wild-type and clone 8 433 (fragment) FXN were constructed using AlphaFold2 (52). A disulfide bond between 434 Cys112 and Cys187 is formed. For the simulations, three non-native-extra residues in 435 the N-terminal stretch were included: Met78-Gly79-Ser80-Pro81-Ile82-Ser83, as for 436 the recombinant proteins produced in our laboratory. 437

# 439 Edited Clone 8 Presents Undetectable Frataxin Expression Levels

First, we analyzed the expression levels of frataxin in the wild-type AX2 and the edited clone 8 strains. Two different nanobodies that detected the recombinant DdFXN in Western blotting were used to study the frataxin expression in total *D. discoideum* cell lysates (**Figure 5**).



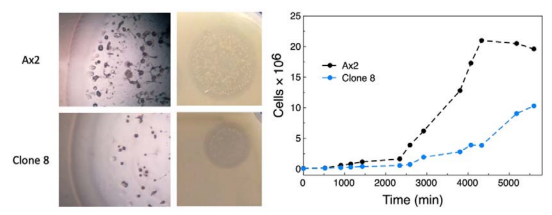
444

Figure 5. Analysis of Frataxin Expression by Western Blotting. Total cell lysates
corresponding to AX2 and clone 8 were analyzed by Western blotting using two
different nanobodies that carry a 6xHis-tag (NB41 and NB45) raised against DdFXN
and a HRP–conjugated secondary antibody antiHis tag (lower panel) and anti β-actin as
loading control (upper panel).

As it can be seen in **Figure 5**, the cell lysate corresponding to the wild-type AX2 undoubtedly showed detectable levels of frataxin. On the other hand, in clone 8 a band corresponding to frataxin or to a frataxin fragment was not detectable. The absence of a detectable signal in clone 8 was not due to the inability of nanobodies to recognize the mutated versions since when the truncated version was heterologous expressed in bacteria, it was readily detected by both nanobodies (**Figure S4**). This result indicates the virtual absence of frataxin in the CRISPR edited cells (**Figure 5**).

458 A D. discoideum Clone Lacking Frataxin Has Severe Defective Growth

As an initial global analysis on the effects of altering frataxin functionality in *D*. *discoideum* biology, we evaluated the growth of the frataxin-deficient clone in the HL5 rich medium and on bacteria. As it is shown in **Figure 6 and Table 3**, clone 8 presents a growth defect in both conditions. The wild-type strain AX2 exhibited a generation time of  $9.5 \pm 2.2$  h, which is in accordance with the literature (57). On the other hand, the edited clone 8 carrying the truncated frataxin grew at a lower rate (14.9 ± 2.2 h).



465

466 **Figure 6.** *D. discoideum* **Growth.** Ax2 and clone 8 were grown on bacterial lawn (left) 467 or in HL5 culture medium (right) starting from a  $\sim 1 \times 10^5$  amoeba cells mL<sup>-1</sup> inoculum. 468 The cell cultures were grown during a week, and small aliquots were regularly taken for 469 cell counting using a Neubauer chamber. Three independent experiments were carried 470 out; a representative experiment is shown. 471

472

# 473 Table 3. Generation Time Summary

Strain	Ax2	Clone 8	Ax2 (empty)	Ax2 (G122V)	Ax2 (FXN wt)	Clone 8 (empty)	Clone 8 (G122V)	Clone 8 (FXN wt)
Generation	$9.5 \pm 2.2$	$14.9 \pm 2.2$	$12.6\pm2.6$	$13.2 \pm 2.8$	$12.7 \pm 2.3$	$19.3 \pm 2.4$	$12.8\pm2.7$	$13.2 \pm 2.8$
Time (h)								

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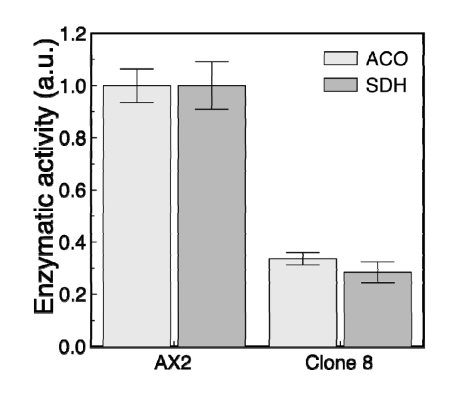
# 476 A D. discoideum Clone Lacking Frataxin Presents a Strong Decrease in Fe-S Cluster

# 477 Dependent Enzymatic Activities

Given that the typical phenotype of a reduction in frataxin functionality is a decrease in Fe-S cluster dependent enzymatic activities, and that this is a feature shared among the FA models, we studied the Krebs cycle enzymes aconitase (ACO) and 481 succinate dehydrogenase (SDH, the Complex II from the respiratory chain). We
482 compared the activity of these mitochondrial enzymes in cellular extracts of strains AX2
483 and clone 8 (Figure 7).

484 Clones 8 exhibited a significant reduction of the enzymatic activities by 485 comparison to the wild-type AX2. The activity decreased to 30-40% of the level 486 observed in the wild-type; these values are comparable to those obtained in other cell 487 models deficient in frataxin, where a  $\sim$ 50% reduction of ACO and SDH activities were 488 detected (*46*, *58*).

489



491 492

490

493 Figure 7. Effect of Frataxin Alteration on Fe-S Cluster Dependent Enzymatic
494 Activities. Aconitase (ACO) and succinate dehydrogenase (SDH) activities were
495 assayed on total lysate extract of *D. discoideum* cells from the AX2 strain and clone 8.
496 Three independent experiments were carried out. The activity measurements are relative
497 to the total protein quantified by triplicate in each extract by the Bradford method.
498

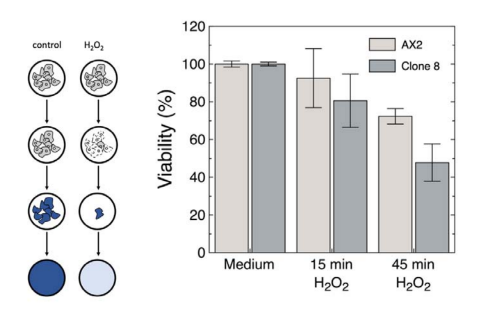
# 499 A D. discoideum Clone Lacking Frataxin Presents Higher Sensitivity to Oxidative

500 Stress

501 Another hallmark of FA disease is an increased sensitivity to oxidative stress. The 502 evidence provided by several organism models supports the idea that the deficiency in 503 frataxin function causes a deregulation in antioxidant response (42, 59–64). In a similar 504 fashion, one can think that clone 8, which exhibits a significant lower growth rate, may 505 show an inefficient handling of reactive oxygen species (ROS), with an ineffective 506 elimination of ROS by the antioxidant system. To study this issue, we carried out a 507 multi-well plate assay and the effect on cellular viability of a treatment with  $H_2O_2$  was 508 determined. We observed a higher impact of hydroperoxide on clone 8 cells compared 509 to the wild-type AX2 cells (Figure 8).

510

511



512 513

514 **Figure 8. Sensitivity to Oxidative Stress.** For determining the viability of cultured 515 cells, a Crystal Violet assay was carried out. The differential effects of  $H_2O_2$  on clone 8 516 cells compared to the wild-type AX2 cells was evaluated. A treatment of frataxin-517 deficient amoeba with hydroperoxide resulted in a significantly increase of the cell 518 detachment from the multi-well plate surface.

520

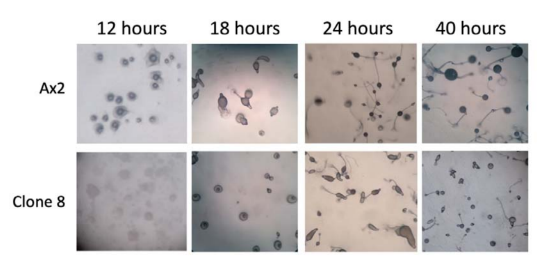
# 521 A D. discoideum Clone Lacking Frataxin Presents Defects in Multicellular

522 Development Progress

<sup>519</sup> 

523 To continue with the exploration of the frataxin deficiency effects on D. discoideum biology, we monitor the progression of the multicellular development 524 525 program in the frataxin-deficient clone 8 (Figure 9). As mentioned previously, D. discoideum grows as isolated ameboid cells that divide mitotically in a nutrient rich 526 culture medium, or in the presence of bacterial lawn. On the other hand, starvation 527 induces a developmental program, the result of which is the formation of a fruiting body 528 529 bearing spores that disseminate and germinate when nutrients become available. This 530 program involves chemotaxis, cell differentiation, morphogenetic movements, and the 531 integration of internal and external signals. In amoeba with mitochondrial deficiency, as 532 a compensatory mechanism for an energy deficit, there is a chronic increase in AMPK 533 activity (AMP-activated protein kinase) that maintains mitochondrial mass and ATP 534 levels at normal levels; however, it affects other processes that require energy, such as 535 growth, multicellular development, phototaxis and chemotaxis (65). Furthermore, 536 previous reports of *D. discoideum* with mitochondrial disease show that mitochondria 537 defective cells present poor growth both in fluid and in bacterial grasses, and also 538 alterations in multicellular development (66).

As it can be seen in **Figure 9**, clone 8 presents an altered development; after 18 h of starvation, whereas the wild-type AX2 already exhibited mature fruiting bodies, clone 8 presented immature stages (tipped mound). This significant delay seems to start from the onset of development as it was already evident after 12 h of starvation; at this point, the wild-type strain reached the mound stage while the clone was only at an early loose aggregate stage.



549 Figure 9. Starvation-Triggered Multicellular Development **Progression.** Exponentially growing cells of Ax2 (top) and clone 8 (bottom) were seeded in non-550 nutrient agar plates to induce multicellular development and then incubated at 22 °C. 551 Pictures were taken at different times (12, 18, 24 and 40 hours). A representative picture 552 553 of each timepoint is shown. 554

555

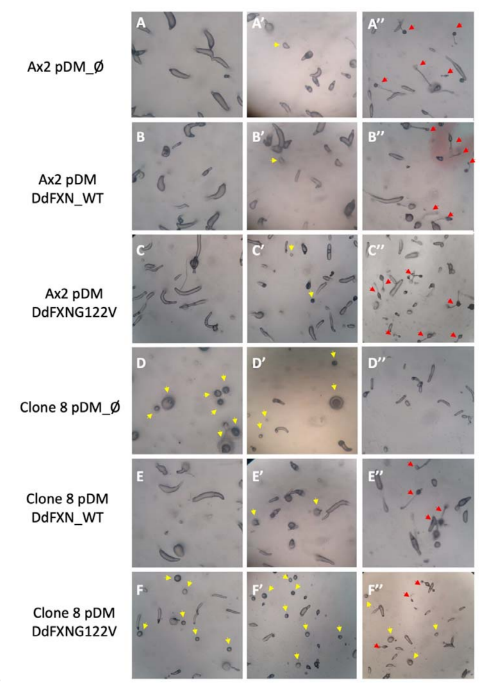
# 556 Constitutive Expression of Frataxin Wild-Type Rescues All Phenotypes in Clone 8

557 To undoubtedly establish that all defects reported in clone 8 were due to the lack 558 of functional frataxin, we expressed an intron-less, C- terminal flag-tagged wild-type 559 frataxin (DdFXNwt) version from a constitutive promoter.

560 We then evaluated the capacity of the wild-type DdFXN to rescue the 561 phenotypes reported above. As it can be seen, both growth (Table 3) and development 562 (Figure 10) of clone 8 expressing DdFXNwt were not significantly different to the wild-type strain AX2 transformed with an empty vector or over-expressing DdFXNwt. 563 564 On the other hand, clone 8 transformed with an empty vector recapitulates the reported alterations: an increased generation time (Table 3), decreased aconitase and succinate 565 dehydrogenase enzymatic activities (Figure 11B) and delayed development (Figure 566 10). 567

568 Interestingly, the rescue of the decrease in the iron-sulfur cluster dependent 569 enzymatic activities was not complete (**Figure 11B**). The expression of DdFXNwt from

- 570 the plasmid was corroborated by Western blotting analysis, using an anti-FLAG
- 571 monoclonal antibody (Figure 11C). As expected, no expression was detected when the
- 572 transformation was carried out with an empty vector.



**Figure 10.** Starvation-Triggered Multicellular Development Progression. Exponentially growing cells of each strain were seeded in non-nutrient agar plates to induce multicellular development and incubated at 22 °C. Representative pictures after 14 hours (left column), 16 hours (middle column) and 20 hours (right column) of

578 development are shown. Yellow arrows and red arrows mark immature mound/tipped 579 mound and early culminants/fruiting bodies.

580

### 581 Constitutive Expression of Frataxin G122V in Clone 8 Fully Rescues Growth Defect 582 but Produces Only Partial Recovery of Developmental Impairment

583

Finally, we wondered whether a high expression of a FA frataxin variant can rescue the wild-type features of *D. discoideum* cells. We transformed clone 8 cells with plasmid pDM326-DdFXN-G122V, which encodes the precursor form of G122V frataxin variant. This variant carries a mutation that produces a very specific and less aggressive FA phenotype in humans (G130V). It is worthy of note that clone 8 cells transformed with pDM326-DdFXN-G122V recovered the growth rate of wild-type AX2

590 (**Table 3**, **Figure 11A**).

We then evaluated whether the G122V frataxin might be able to rescue the defects in multicellular development observed for clone 8. Although a small improvement was detected when these cells were transformed with pDM326-DdFXN-G122V, compared with the same cells transformed with empty pDM326, development was not fully rescued.

At 20 h after starvation, we were able to still see immature structures in clone 8 overexpressing the G122V variant while none were observed when complemented with the wild-type frataxin (**Figure 10**).

We also evaluated the rescue capacity of DdFXN-G122V on the decrease of enzymatic activities (**Figure 11B**); although rescue was not total, the enzymatic levels reached were like the values obtained when DdFXNwt was expressed.

These results suggest that, because of the specific features of this variant, the expression of DdFXN-G122V cannot fully compensate all deficiencies lacking in the wild-type frataxin. This fact also suggests different activities, which are finely tuned by frataxin structure and dynamics, or different controls over the metabolic pathways.

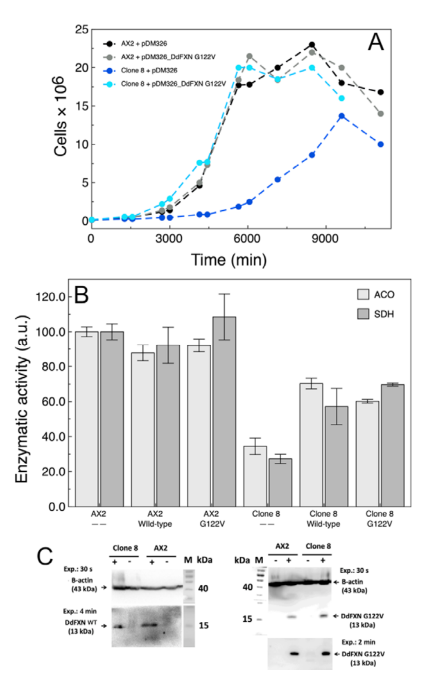




Figure 11. The Expression Frataxin Variant G122V Rescued Clone 8. (A) Growth 608 rate and (B) aconitase and succinate dehydrogenase enzymatic activities of D. 609 discoideum AX2 and clone 8 cells transformed with pDM326, pDM\_DdFXNwt or 610 pDM\_DdFXN G122V encoding the precursor variant of DdFXN G122V. Growth was 611 carried out at 22 °C and 180 rpm in HL5 culture medium, starting from a 1×10<sup>5</sup> amoeba 612 cells mL<sup>-1</sup> inoculum. The cell cultures were grown during a week, and small aliquots 613 614 were regularly taken for cell counting using a Neubauer chamber. The antibiotic 615 blasticidin was present in culture medium to maintain the pDM326 plasmid in D. discoideum cells. For activity assays, the activities were corrected by the total protein 616 mass, measured by Bradford. (C) Western blotting analysis of the expression of DdFXN 617

618 wild-type (WT) or the DdFXN G122V variant from plasmid pDM326. For G122V, 619 upper and lower panels correspond to 30 s and 2min exposition, respectively. The + and 620 – symbols indicate transformation of *D. discoideum* with the corresponding plasmid 621 without a frataxin gene insert (the wild-type or the DdFXN G122V variant). The 622 detection of the recombinant variant was carried out by using an anti-FLAG monoclonal 623 antibody that recognizes the FLAG sequence located in the C-terminal stretch of the 624 plasmidic frataxin.

625 626

#### 627 Discussion and Conclusions

628 Friedreich's Ataxia is a rare disease caused by a disfunction of frataxin, a mitochondrial protein, coded in the nucleus. Frataxin is a key component of the 629 630 mitochondrial supercomplex responsible for Fe-S cluster assembly. Fe-S cluster dependent activities are essential for the homeostasis of the cells. There is a high 631 632 diversity of cell functions that need the correct insertion of the Fe-S cluster into protein structure. The energetics of the cell, which is commanded by the mitochondrial 633 634 metabolism, directly depends on Fe-S clusters because clusters are crucial in electron 635 transport reactions. Thus, the alteration of Fe-S cluster biosynthesis affects Complex I, II and III. But Fe-S are also essential for other enzymatic activities like lipoic acid 636 637 synthesis, necessary for pyruvate dehydrogenase activity or substrate binding in mitochondrial aconitase (Krebs cycle); ultimately, given that pyruvate dehydrogenase 638 639 complex, Krebs cycle, and the electron transport chain, are all metabolic pathways highly coupled to ATP synthesis, the alteration of the Fe-S cluster assembly results in 640 641 the impairment of numerous ATP-dependent cellular processes.

In previous research we showed that *D. discoideum* has a complete pathway involved in Fe-S cluster assembly and transferring, as found in mammalian cells. Our inferences from protein sequence conservation and structure models suggested that some proteins (human and *D. discoideum*) may be exchangeable. In fact, the FXN residues that when mutated result in FA are practically fully conserved between human *D. discoideum*. In this study, we observed that DdFXN can bind to the human 648 supercomplex NIAU with a high affinity, similarly to the human FXN variant. 649 Additionally, it was able to activate L-Cys desulfurase function. This is indicative of the 650 role of protein-protein interaction surfaces and the high conservation score for the 651 residues involved in these interactions. This highlights the potential of *D. discoideum* as 652 a model system for FA.

653 We then showed that frataxin locus can be easily edited in D. discoideum. 654 Taking advantage of the haploid context of this amoeba, we constructed a completely 655 frataxin-deficient strain. In the edited D. discoideum clone 8, studied in this paper, a premature STOP codon occurred. The frataxin fragment encoded in the genomic DNA 656 657 seemed to be extremely unstable as judged by its absence in clone 8 amoeba cellular 658 lysates, a result that matches with molecular dynamics simulations, which suggests a 659 highly mobile protein backbone compared to the wild-type DdFXN. This strain presents 660 a significant decrease of the Fe-S cluster dependent enzymatic functions and an altered 661 phenotype. Reduced aconitase and succinate dehydrogenase activities, a decrease in 662 growth rate, and higher sensitivity to oxidative stress were consistently observed, being all these landmarks of frataxin deficiency in other Friedreich's Ataxia cellular and 663 organism models. Furthermore, we explored other processes that have been linked to 664 665 mitochondrial disfunction in D. discoideum and found defects during multicellular 666 development. Remarkably, we were able to rescue phenotype alterations with the constitutive expression of wild-type DdFXN. The expression of the wild-type protein 667 was able to fully rescue growth and multicellular behavior alterations while the 668 decreased enzymatic activities were only partially rescued. There are a few conceivable, 669 670 not mutually exclusive, explanations for this result. The frataxin expression levels 671 obtained when expressed from plasmids are different than endogenous. Furthermore, 672 given that frataxin is imported into the mitochondrial matrix, the higher expression 673 levels of the protein might also affect the importing and processing machinery, although 674 no effect was observed when a constitutive expression was carried out in the wild-type 675 AX2 strain. On the other hand, it is also worth noting that the DdFXN sequence used for protein expression from plasmids was engineered carrying a C-terminal FLAG tag 676 for protein identity assignment, which could affect some roles of frataxin but not others. 677 The fact that the FLAG sequence includes a highly negatively charged stretch of 678 679 residues, makes feasible that it alters in some degree frataxin interaction with the D. 680 discoideum NIAU supercomplex. Experiments with untagged versions would help to test this hypothesis. 681

682 We then used this set-up to assess the rescuing capacity of DdFXN G122V, a frataxin version that mimics a pathogenic variant found in Friedreich's Ataxia patients 683 684 (i.e., G122V in *D. discoideum* corresponds to G130V in humans). This variant, when it 685 is constitutively expressed, fully rescues the defects on growth but only marginally rescues the defects on multicellular development. This underlies the fact that DdFXN 686 G122V is not merely a less stable frataxin variant, because increased expression levels 687 should bypass the defect. So it may be that other features of the protein affected in 688 G122V are important for specific roles played by frataxin. Hence, clone 8 can help to 689 690 dissect the different functions of FXN within the cell. On the other hand, the defects of 691 enzymatic activities were rescued to the same levels as in the case of the expression of 692 the wild-type C-terminal tagged DdFXN.

In this context, this new biological model offers a wide range of options to easily explore diverse phenotypes occurring in FA; this makes *D. discoideum* a very attractive approach for studying, in a straightforward manner, the effect that diversity of FA variants has on the cellular metabolism. Moreover, this model may help to understand

697 whether frataxin works as a bottleneck for some processes, whereas in other processes it

698 has lower flux control, at least under the experimental conditions assayed.

In summary, we have generated a system where we can easily monitor the effects of a lack of frataxin activity; this opens the door to developing drug or treatment screenings that would help to design and/or evaluate therapeutical strategies. Moreover, this biological model offers a wide range of possibilities to easily explore diverse phenotypes in FA. To complete the list of desirable attributes for a disease model organism, it has been proved that frataxin locus can be easily edited, which enables the generation of specific strains that express the variant of interest from an endogenous locus.

707

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713

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