Expression of Yeast NDI1 Rescues a Drosophila Complex I Assembly Defect

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Abstract

Defects in mitochondrial electron transport chain (ETC) function have been implicated in a number of neurodegenerative disorders, cancer, and aging. Mitochondrial complex I (NADH dehydrogenase) is the largest and most complicated enzyme of the ETC with 45 subunits originating from two separate genomes. The biogenesis of complex I is an intricate process that requires multiple steps, subassemblies, and assembly factors. Here, we report the generation and characterization of a Drosophila model of complex I assembly factor deficiency. We show that CG7598 (dCIA30), the Drosophila homolog of human complex I assembly factor Ndufaf1, is necessary for proper complex I assembly. Reduced expression of dCIA30 results in the loss of the complex I holoenzyme band in blue-native polyacrylamide gel electrophoresis and loss of NADH:ubiquinone oxidoreductase activity in isolated mitochondria. The complex I assembly defect, caused by mutation or RNAi of dCIA30, has repercussions both during development and adulthood in Drosophila, including developmental arrest at the pupal stage and reduced stress resistance during adulthood. Expression of the single-subunit yeast alternative NADH dehydrogenase, Ndi1, can partially or wholly rescue phenotypes associated with the complex I assembly defect. Our work shows that CG7598/dCIA30 is a functional homolog of Ndufaf1 and adds to the accumulating evidence that transgenic NDI1 expression is a viable therapy for disorders arising from complex I deficiency.

Introduction

Alterations in mitochondrial energy metabolism have been implicated in aging and age-onset disease [1]. Mitochondrial complex I (NADH:ubiquinone oxidoreductase/NADH dehydrogenase), is one of the most complicated enzymes of the eukaryotic cell, with more than 40 subunits originating from two separate genomes [2] and a still growing list of accessory and assembly factors [3]. Functioning as the major entry site of electrons from the mitochondrial electron transport chain (ETC), the fully assembled enzyme is thought to be L-shaped, embedded in the mitochondrial inner membrane by a hydrophobic membrane arm with a hydrophilic peripheral arm protruding perpendicularly into the mitochondrial matrix [4]. A current model of biogenesis of human complex I supports the assembly of the membrane arm in a stepwise process with subsequent addition of a partially, independently assembled peripheral arm and additional nuclear encoded subunits to complete the holoenzyme [5], [6]. Moreover, recent studies have shown that complex I subassemblies can act as scaffolds for the assembly of respiratory supercomplexes consisting of multiple ETC complexes [7].

Proper assembly of complex I is intimately associated with the presence of a multitude of assembly factors and chaperones, and their loss results in diseases that mimic complex I subunit defects [3]. Complex I assembly factors were first isolated in studies of Neurospora crassa mutants that accumulated complex I subassemblies, with associated assembly factors, due to a mutation in a membrane arm subunit [8]. Of the two complex I intermediate associated (CIA) proteins identified in this study, only the 30 kDa protein (CIA30) has been shown to have a human homolog (NDUFAF1) [9] that functions as an assembly factor, interacting with mid-stage membrane arm subassemblies but not with a fully assembled holoenzyme or a late-stage subunit. A patient with mutations in Ndufaf1 has also been described who shows severely reduced levels of complex I holoenzyme and suffers from cardioencephalomyopathy [10]. Complete deletion of CIA30 in N. crassa result in complete loss of complex I and respiration exclusively via an alternative internal NADH:ubiquinone oxidoreductase [8].

Unlike mitochondrial complex I, flavone-sensitive, rotenone-insensitive, single-subunit, alternative internal NADH dehydrogenase (Ndi1) genes are restricted to plant and fungal mitochondria where they function as NADH dehydrogenases without translocating protons across the inner mitochondrial membrane [11–13]. In fungus, NDI1 activity has been shown to be sufficient to complement complete loss of complex I holoenzyme [14], and...
some yeast, such as *Saccharomyces cerevisiae* lack a multi-subunit mitochondrial complex I entirely [15]. Due to its much simpler genetics and functionality in non-native systems [16], transgenic NDI1 has been shown to be highly effective as a therapeutic tool for complex I associated diseases in non-fungal systems, including nematodes [17], arthropods [18], and mammals [19–23]. Moreover, the contrasting sensitivity to inhibitors between NDI1 (flavone sensitive, rotenone insensitive) and endogenous complex I (rotenone sensitive, flavone insensitive) allows for accurate assessment of the contribution of NADH:ubiquinone oxidoreductase activity from the two sources.

Previous work from our lab and others has shown that expression of *Nadi1* in the fruit fly, *Drosophila melanogaster*, can increase metabolic activity and lifespan [24], [25]. Here, we have expanded upon this paradigm by examining the impact of exogenous *Nadi1* expression in flies with a complex I assembly defect. To do so, we first characterized the consequences of reduced expression of the *Drosophila* homolog of the complex I assembly factor *Ndufaf1*/*CG7598* (dCIA30). We demonstrate that flies carrying a mutation in *dCIA30* display a reduction of complex I holoenzyme in blue native polyacrylamide gel electrophoresis (BN-PAGE) to undetectable levels. Moreover, mitochondria isolated from larvae with *dCIA30* knock down show drastic reductions in endogenous rotenone sensitive, flavone insensitive NADH:ubiquinone oxidoreductase activity. In addition, RNAi knock down of *dCIA30* allows for development into adulthood and these flies show sensitivity to a range of extrinsic stressors. We show that expression of *Nadi1* partially rescues developmental arrest in mutants, and complex I deficiency-associated phenotypes in *dCIA30* RNAi knock down flies. These results support the idea that expression of *Nadi1* may be an effective therapeutic strategy to treat disorders resulting from defects in complex I assembly. Furthermore, our *dCIA30* knock down model provides a powerful tool to better understand the pathophysiological mechanisms of human disease arising from complex I deficiency.

**Materials and Methods**

**D. melanogaster Culture**

Flies were maintained on standard agar-cornmeal-yeast-sugar media [26] in humidified incubators at 25°C, on 12:12 hour light:dark cycles. Flies were switched to new media every 2–3 days. Fly lines *CG7598* (stock #16923) [27] and *UAS-Gfp-IR* (stock #9931) were obtained from the Bloomington Drosophila Stock Center, and the *UAS-CG7598-IR* (stock #14859) fly line was obtained from the Vienna Drosophila RNAi Center [28]. *UAS-Nadi1* flies were generated as previously described [24]. An imprecise excision of the *CG7598* (P element line, *dCIA30* exon 8) was generated as previously described [29], and verified by sequencing. A precise excision that restores the gene was also generated, verified by sequencing, and used as controls in all experiments. *UAS-dCIA30* flies were generated by transforming flies with pUAST plasmids containing *dCIA30* cDNA using standard procedures.

**qRT-PCR**

Total RNA was extracted using TRIzol reagent (Invitrogen, USA) following manufacturer protocols from 5 L3 larvae or 5 adult flies. Amplicons of *Actin5C* were used as a reference to normalize *dCIA30* amplicons. cDNA synthesis and qRT-PCR were performed in one step using Power SYBR Green RNA-to-CT 1-Step kit (Applied Biosystems, USA), and DNA amount was monitored during a 40-cycle PCR with an Applied Biosystems 7300 Thermal Cycler (Life Technologies, USA). Primer sequences: *Actin5C*, TTGTCTGGGCAAGAGGATCG and AC-CACCTGCACTTTGCACTTTC, *dCIA30*, TCACACAAAAGGATGCCATTA and GCATGTGGTACATGGCTCCAG.

**Mitochondrial Isolation**

Mitochondria were purified from larvae or adult flies by differential centrifugation as previously described [30]. Briefly, larvae or adults were homogenized in chilled mitochondrial isolation medium (MIM, 250 mM sucrose, 10 mM Tris (pH 7.4), 0.13 mM MgCl2) and debris was pelleted by centrifugation (500 xg, 5 min at 4°C). Mitochondria were pelleted from the supernatant by centrifugation (5,000 xg, 5 min at 4°C) and stored at −80°C.

**Blue-native Polyacrylamide Gel Electrophoresis (BN-PAGE)**

BN-PAGE was performed using a Novex Native PAGE Bis-Tris Gel System (Invitrogen) following manufacturer protocols. Briefly, mitochondria were purified from 20 L3 larvae or 20 adult flies, resuspended in 25 μl of 1×Native PAGE Sample buffer (Invitrogen) with 1% digitonin and protease inhibitors (Roche, USA), and incubated on ice for 15 min. After centrifugation at 16,100 xg for 30 min, 25 μl of the supernatant was resuspended with 1.25 μl of 5% G250 sample additive and 10 μl of 4×Native PAGE Sample Buffer (Invitrogen). Samples were loaded on 3–12% Bis-Tris Native PAGE gels and electrophoresed using 1×Native PAGE Running buffer system (Invitrogen). The cathode buffer included 1×Cathode Buffer Additive (Invitrogen). NativeMark Protein standard (Invitrogen) was used as the molecular weight marker. Protein concentrations of adult fly mitochondrial preps were determined with a Micro BCA Protein Assay Kit (Thermo Scientific, USA) following manufacturer instructions.

**Eclosion**

Twenty wandering L3 larvae were collected in vials and maintained as described above. For approximately 10 days, the number of flies that eclosed from each vial was recorded daily.

**NADH:ubiquinone Oxidoreductase Activity Assay**

Mitochondria were purified from 5 male L3 larvae, resuspended in 100 μl MIM, and 3 μl were added to 150 μl of a previously prepared colorimetric complex I activity assay buffer (1× PBS, 3.5 g/1 BSA, 0.2 mM NADH, 0.24 mM KCN, 60 μM DCIP, 70μM decylubiquinone, 25 mM antimony A). NADH:ubiquinone oxidoreductase activity was monitored as a drop in DCIP absorbance at 600 nm using an Epoch microplate spectrophotometer (BioTek, USA). Flavone or rotenone insensitive activity was measured as the difference in DCIP reduction in the presence of flavone (0.4 μM) or rotenone (2 μM) in the assay buffer, and baseline activity in an assay buffer that contained both inhibitors. All reported activities are normalized to citrate synthase activity.

**Citrate Synthase Activity Assay**

Mitochondrial preps used in the NADH:ubiquinone oxidoreductase assay were diluted 10 fold in MIM, and 3 μl were added to 150 μl of a previously prepared colorimetric citrate synthase activity assay buffer (50 mM Tris (pH 8.0), 0.1 mM 5,5′-dithiobis-(2-nitrobenzoic acid) (DTNB), 0.3 mM acetyl-CoA, 1 mM oxaloacetic acid). Citrate synthase activity was measured as an increase in DTNB absorbance at 412 nm using a microplate spectrophotometer.
Electron Micrographs (EM)

EMs were acquired as previously described [29] from male flies 2 or 10 days post eclosion. Briefly, dorsal indirect flight muscle was dissected from decapitated adult flies at 4°C in 2% paraformaldehyde with 1% glutaraldehyde and fixed in the same solution overnight. After postfixation in 1% osmium tetroxide at room temperature, samples were dehydrated in an ethanol series and embedded in Epon 812. Ultrathin sections (80 nm) were examined with a Philips 420 electron microscope (Philips, Netherlands) at 100 kV at a magnification of 4900X.

Weight

Flies were anesthetized under light N2 gas and weighed in groups of 5 in pre-weighed microcentrifuge tubes using an analytical scale (Torbal, USA).

Stress Resistance

All stress assays were performed with male flies 6–8 days post eclosion in groups of 25–30 flies. For hypoxia resistance, flies were exposed to anoxic conditions in a 100% N2 chamber. Flies were maintained in the chamber for one hour, moved back into normoxia, and monitored for recovery (ability to stand) every 3 minutes for approximately 2 hours. For hyperoxia resistance, flies were maintained in a humidified chamber maintained at 85% O2 and survival was assayed at least once per day. For wet starvation, flies were maintained on water only medium (1% agar in ddH2O) and maintained in a 25°C incubator with 12 hour light:dark cycles. Survival was scored multiple times per day. For hyperthermia resistance, flies were maintained at 37°C and survival was scored every 2 hours.

Statistical Analysis

Unless indicated otherwise, significance was determined with a two-tailed, unpaired t test from at least three independent experiments and expressed as p values. All error bars reflect standard error of the mean.

Results

CG7598 (dCIA30), the Drosophila Homolog of Human Complex I Assembly Factor, Ndufa1, is Necessary for Complex I Assembly

A Drosophila melanogaster homolog of the Neurospora crassa CIA30 protein, CG7598 (dCIA30), was previously identified in a homology search of amino acid sequences [31]. The dCIA30 protein shares high homology with human NDUFAF1 (69% similarity, 44% identity) in a ClustalW amino acid sequence alignment [32], in particular, in the C-terminal half where a conserved domain search turns up a conserved CIA30 domain (Figure S1A). As part of the Drosophila Gene Disruption Project [27], an insertion mutation of dCIA30 (insertion EY09101, dCIA30EY09101) that contains an approximately 11 kb transposable P element (P{EPgy2}), in the 5' UTR is available and was acquired through the Bloomington stock center (stock #16925). In order to proceed with a line that disrupts the gene without the possible confounding presence of additional promoters, enhancers, and other genes, we generated additional lines in which the P{EPgy2} element was precisely and imprecisely excised by crossing dCIA30EY09101 to flies harboring a transposase. One imprecise excision resulted in removal of the bulk of the P element (P[Epay2]), in the 5' UTR and was acquired through the Bloomington stock center (stock #16925). In order to proceed with a line that disrupts the gene without the possible confounding presence of additional promoters, enhancers, and other genes, we generated additional lines in which the P[Epay2] element was precisely and imprecisely excised by crossing dCIA30EY09101 to flies harboring a transposase. One imprecise excision resulted in removal of the bulk of the P[Epay2] element, leaving only a 517 bp fragment that does not carry any identifiable genetic elements, but does contain start and stop codons in all three forward frames (dCIA30ex80). In addition, a precise excision line was generated and used as a control throughout this study (Figure S1B).

We checked the effects of the insertion mutation on dCIA30 gene activity by quantifying dCIA30 transcript levels during the third-instar larval stage of development (L3). Measurement of relative dCIA30 mRNA levels by quantitative reverse transcriptase

Figure 1. dCIA30 mutation results in loss of complex I holoenzyme. (A) dCIA30 transcript levels were measured by qRT-PCR in male third instar (L3) larvae (n = 5 male L3 larvae, 3 replicates). The dCIA30EY09101 and dCIA30EY09101 insertion mutants show a ca. 55–75% loss of transcript relative to a precise excision control (* p<0.05). (B) The dCIA30EY09101 and dCIA30EY09101 mutations result in a specific loss of the band that corresponds to complex I holoenzyme in blue native polyacrylamide gel electrophoresis (BN-PAGE) of L3 larvae. Precise excision controls (+) and larvae with expression of a dCIA30 cDNA construct (UAS-dCIA30/; da-GAL4, dCIA30EY09101, cDNA rescue) show the presence of the complex I holoenzyme band. (M = molecular size marker, CV2 = complex V dimer, CI = complex I, CV1 = complex V monomer, CIII = complex III, CV = complex IV, CII = complex II, mitochondria from 2.5, 5, and 10 larvae equivalents in successive lanes for each genotype). doi:10.1371/journal.pone.0050644.g001
Figure 2. dCIA30 mutation results in developmental arrest and defects in mitochondrial function and ultrastructure. (A) Homozygous dCIA30[EY09101] or dCIA30[ex80] mutants fail to develop past the pupal stage (***p<0.001). Expression of a cDNA rescue construct (cDNA rescue) is sufficient to rescue eclosion frequencies back to those of controls (+). (n = 20 L3 larvae, 8 replicates). (B) Isolated mitochondria from homozygous dCIA30[EY09101] or dCIA30[ex80] mutant male L3 larvae show negligible endogenous (rotenone sensitive, flavone insensitive) NADH:ubiquinone oxidoreductase activity relative to controls (+, *p<0.05). Expression of dCIA30 cDNA restores endogenous NADH:ubiquinone oxidoreductase activity back to control levels. (n = 5 male L3 larvae, 5 replicates). (C) Homozygous dCIA30[EY09101] or dCIA30[ex80] pupae are structurally similar but visibly smaller relative to precise excision controls. Expression of dCIA30 cDNA is sufficient to rescue the size deficit. Tick marks = 1 mm. (D) Electron micrographs (EMs) of flight muscle tissues show degeneration of myofibrils and interspersed mitochondria in homozygous dCIA30[ex80] male flies 2 days post eclosion. Expression of a dCIA30 cDNA construct restores mitochondrial and myofibrillar integrity and organization. Scale bar = 1 μm.

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polymerase chain reaction (qRT-PCR) showed that both the dCIA30EY09101 and dCIA30ex80 homozygotes have a severe reduction of dCIA30 mRNA levels relative to the control line (Figure 1A). We checked the effect of the reduced dCIA30 mRNA levels on complex I assembly by BN-PAGE of mitochondria isolated from L3 larvae (Figure 1B). As expected, mitochondria from control larvae clearly show the presence of a band corresponding to complex I holoenzyme, identified by molecular mass. In contrast, the diminished levels of dCIA30 in both the dCIA30EY09101 and dCIA30ex80 homozygotes are insufficient to support assembly and accumulation of complex I to detectable levels. Importantly, disruption of dCIA30 did not noticeably affect the assembly or relative abundance of the other ETC complexes. To confirm that the complex I defect was due to loss of dCIA30, we expressed a wild-type dCIA30 cDNA [33] via a ubiquitous daughterless promoter (da-GAL4), in a dCIA300000 homozygous mutant background (UAS-dCIA30+/+; da-GAL4, dCIA30ex80/dCIA30ex80). These ‘cDNA rescue’ flies clearly showed the presence of a complex I holoenzyme band (Figure 1B), demonstrating that the presence of exogenous dCIA30 is sufficient to rescue complex I assembly.

Loss of dCIA30 Confers Developmental Arrest and Defects in Mitochondrial Function and Ultrastructure

Fly development proceeds through four distinct, easily identifiable stages, embryo, larva, pupa, and adult. Larvae homozygous for dCIA300000 or dCIA300000 were viable and survived to late...
Figure 4. *dCIA30* knockdown flies display developmental and physiological phenotypes that are rescued by NDI1. (A) RNAi knockdown of *dCIA30* (*dCIA30-RNAi*) results in a significant drop in the frequency of pupae that eclose to adult flies relative to controls (*da-GAL4/*+, ***p < 0.001). Co-expression of NDI1 in *dCIA30-RNAi* flies (Ndi1 rescue) restores eclosion back to levels found in control pupae. The expression of NDI1 from double copies of both UAS-Ndi1 and da-GAL4 in *dCIA30* insertion mutants (UAS-Ndi1;da-GAL4,dCIA30ex80) restores eclosion to approximately 70% of pupae (***p < 0.001). (n = 20 L3 larvae, 8 replicates). (B) Mitochondria isolated from male L3 larvae with RNAi knock down of *dCIA30* (*dCIA30-RNAi*) show negligible endogenous (rotenone sensitive, flavone insensitive) NADH:ubiquinone oxidoreductase activity relative to controls (*da-GAL4/*+), regardless of UAS-Ndi1 co-expression (Ndi1 rescue, **p < 0.01). Expression of UAS-Ndi1 causes a dramatic increase in flavone sensitive, rotenone insensitive NADH:ubiquinone oxidoreductase activity in a *dCIA30* knock down (Ndi1 rescue) or mutant background (UAS-Ndi1;da-GAL4,dCIA30ex80, ***p < 0.001). (n = 5 male L3 larvae, 5 replicates). (C) RNAi knock down of *dCIA30* produces pupae that are smaller relative to controls. The expression...
of *UAS-Ndi1* does not result in rescue of the pupal phenotype in RNAi knock down flies. However, expression of ND1 with double copies of both transgene and driver (*UAS-Ndi1; da-GAL4; dCIA30IR*) increases pupal size in *dCIA30* mutants (See figure 2C for comparison). Tick marks = 1 mm. (D) Adult male *dCIA30* knock down flies, 6 days post eclosion have significantly lower body weights compared to controls (**p < 0.001). Co-expression of *ND1* in these knock down flies partially restores body weight (**p < 0.001). (n = 5 male flies, 6 replicates). (E) EMs of thoracic flight muscles from male *dCIA30* knock down flies show degeneration of mitochondria and myofibrils at 10 days post eclosion. Co-expression of ND1 largely restores mitochondrial and myofibril integrity and organization. Scale bar = 1 μm.

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Yeast Ndi1 can Partially or Wholly Rescue Phenotypes Associated with *dCIA30* Deficiency

Next, we set out to determine whether the alternative NADH dehydrogenase, *Ndi1*, could complement the loss of *dCIA30*. Expression of ND1 (*UAS-Nd1; da-GAL4; UAS-dCIA30IR; UAS-Ndi1*, referred to as “*Ndi1 rescue*” in this report) in flies with RNAi knock down of *dCIA30* completely reverted the developmental arrest phenotype to control levels (Figure 4A) and expression of ND1 from two copies each of *UAS-Ndi1* and *da-GAL4* transgenes (*UAS-Ndi1; da-GAL4; dCIA30IR*) resulted in a significant rescue of the mutant phenotypes (Figure 4A–C). Adult males that eclosed from these pupae were able to fertilize wild type females whereas adult females failed to produce offspring in crosses with wild type male flies.

This rescue was not a result of a cryptic increase in complex I assembly by ND1 or by dilution of the *UAS-GAL4* system. NADH:ubiquinone oxidoreductase activity of isolated mitochondria from L3 larvae showed negligible endogenous rotenone sensitive, flavone insensitive activity in the knock down lines, relative to controls, even in the presence of *UAS-Ndi1* transgenes (Figure 4B). Instead, expression from *UAS-Ndi1* transgenes resulted in large increases in flavone sensitive, rotenone insensitive activity that corresponded to ND1 activity (Figure 4B). Moreover, the expression of two copies of *UAS-Ndi1* construct in a homozygous *dCIA30IR* background (*UAS-Ndi1; da-GAL4; dCIA30IR*), did not show any effect on the missing complex I band (Figure S3). Similarly, the co-expression of a *UAS-Ndi1* transgene in *dCIA30* RNAi knockdown lines using the same *UAS-GAL4* system had little effect on *dCIA30* mRNA knockdown (Figure 5A and 5C) or in the resulting defect in complex I holoenzyme assembly (Figure 5B and 5D).

RNAi knockdown of *dCIA30* led to an approximately 30% reduction in body weight relative to driver only controls (*da-GAL4* +). Co-expression of ND1 largely restored the adult body weight of flies with *dCIA30* knockdown (Figure 4D). RNAi of *dCIA30* also resulted in detrimental effects on mitochondrial and myofibril structure in adult flies. EM analysis of thoracic muscle revealed severe degeneration of mitochondria and myofibrils, similar to those seen in *dCIA30* mutant flies (Figure 4E). Co-expression of ND1 was sufficient to fully rescue the myofibril defect and partially rescue the mitochondrial ultrastructure defect.

To quantify the physiological effects of complex I loss, we tested the ability of *dCIA30* knockdown flies to cope with various forms of extrinsic stress. To explore the role of complex I/ND1 in the ability to withstand oxidative stress, we examined recovery from hypoxia and survivorship under hyperoxia. RNAi knockdown of *dCIA30* conferred a drastically reduced ability to recover from exposure to hypoxia, which was completely rescued by ND1 expression (Figure 5A). Flies with reduced expression of *dCIA30* also showed increased sensitivity to hyperoxia, which was also rescued by ND1 expression (Figure 5B). Loss of complex I is expected to have drastic effects on metabolism, and accordingly, flies without detectable complex I were much more susceptible to...
wet starvation (Figure 5C). Providing an alternative source of mitochondrial NADH dehydrogenase activity by expressing NDI1 partially restored survival under starvation conditions. Similarly, loss of complex I drastically reduced resistance to elevated temperature that was largely restored by NDI1 expression (Figure 5D).

Discussion

In the current report, we have demonstrated the role of CG7598/dCIA30, the Drosophila homolog of human Ndufaf1, as a vital factor in the assembly of mitochondrial complex I. We show that loss of dCIA30 is sufficient to reduce complex I holoenzyme to levels that are undetectable in BN-PAGE. Reduction of complex I holoenzyme levels can be reverted by precise excision of the inserted element in the dCIA30 mutants or by expression of a wild type dCIA30 cDNA construct. These findings confirm that dCIA30 is a necessary factor for complex I assembly in Drosophila. dCIA30 mutation almost completely abrogated development during the pupa stage, and reduced complex I holoenzyme levels in dCIA30 RNAi knockdown flies caused an approximately 70% drop in eclosion frequency. In flies that did reach adulthood, dCIA30 knock down further reduced complex I holoenzyme to undetectable levels in BN-PAGE. In addition to being smaller, these flies showed dramatically increased sensitivity to a variety of extrinsic stressors including hypoxia, hyperoxia, hyperthermia, and starvation. Interestingly, while sensitivity to hypoxia, hyperoxia and high temperature (conferred by loss of complex I) can be completely or almost completely rescued by Ndi1, sensitivity to starvation conditions was only partially rescued by Ndi1. Therefore, transkingdom gene therapy using Ndi1 may have limitations with respect to complementing defects of energy metabolism.

Among other detriments, loss of complex I may result in reduced ETC function, increased oxidative stress, skewing of the NAD:NADH ratio, and reduced ability to form supercomplexes [7]. Of these, loss of ETC activity, increased oxidative stress, and skewing of the NAD:NADH ratio may be improved by supplementation with NDI1 [18], [24], [34]. In some fungi, such as S. cerevisiae, NDI1 functions as the sole matrix facing NADH:ubiquinone oxidoreductase [15], coupling NADH oxidation to ubiquinone reduction like mitochondrial complex I, but without translocating protons across the inner mitochondrial membrane. As Ndi1 in the present study was cloned from S. cerevisiae [24], it is unlikely that it could act as a scaffold for supercomplex formation or otherwise participate specifically and directly in other endogenous pathways when expressed in Drosophila, which does not have an endogenous Ndi1 homolog. The near complete rescue of sensitivity to hypoxia, hyperoxia, and hyperthermia in complex I deficient flies by NDI1 supplementation suggests that sensitivity to hypoxia, hyperoxia, and hyperthermia may be linked to loss of ETC activity, increased oxidative stress, and/or skewing of the NAD:NADH ratio. Conversely, the major detriment in wet starvation may be a result of a complex I function that is not supplemented or insufficiently supplemented.

Figure 5. Loss of dCIA30 results in multiple stress sensitivity phenotypes that are ameliorated by NDI1. (A) Knock-down of dCIA30 (dCIA30-RNAi) in adult flies increased recovery time after exposure to hypoxia (1 hr). Loss of dCIA30 more than doubled mean recovery time compared to controls (da-GAL4/+). Co-expression of NDI1 (Ndi1 rescue) restored recovery time to control levels. (n = 120 male flies). (B) Knock down of dCIA30 in adult flies increased survival time under hyperoxia (84% O2). On average, dCIA30-RNAi flies lived approximately half as long as controls. Co-expression of NDI1 restored mean survival duration to approximately 95% of controls. (n = 120 male flies). (C) Knock down of dCIA30 in adult flies decreased survival duration under starvation with water. On average, flies with dCIA30 knock down survived less than half as long as control flies. Co-expression of NDI1 increased survival times to approximately 70% of controls. (n > 120 male flies). (D) Knock down of dCIA30 in adult flies decreased survival duration under high temperature (37°C). Mean survival times of flies with reduced dCIA30 were approximately half those of control flies. Co-expression of NDI1 increased mean survival time to approximately 90% of controls. (n > 135 male flies). doi:10.1371/journal.pone.0050644.g005
by NDI1 function. Although beyond the scope of this work, a further analysis of the contribution of the different harmful effects of complex I deficiency to various stresses may yield further insights into the etiology of complex I deficiencies. NDI1 has previously been shown to be effective as a therapeutic for complex I disorders, including fly [18], mouse [19], and rat [20] models of Parkinson’s disease, and in the fungus *Podospora anserina*, overexpression of endogenous *Ndi1* was shown to be able to rescue complex I holoenzyme deficiency [14]. To our knowledge, however, this is the first demonstration that exogenous NDI1 is able to restore development to a metazoon that has no detectable complex I holoenzyme in BN-PAGE or NADH:ubiquinone oxidoreductase activity in a colorimetric assay. Moreover, the generation and characterization of a fly model of complex I assembly factor deficiency will facilitate future work into the underlying pathophysiology of neurodegeneration and aging.

**Supporting Information**

**Figure S1** CG7598 is the *D. melanogaster* homolog of Ndufaf1/CG7598 identified by homology. Amino acid sequence alignment shows 69% similarity (44% identity), concentrated near the C-terminal half, which contains the CIA30 domain. (B) CG7598 maps to chromosome 3R at 99B9 and spans approximately 1.1 kb. The coding sequence consists of two exons separated by a 58 bp intron. A fly line with an insertion of an approximately 11 kb long P-element (P{EPgy2}) into the 5′UTR, 96 bp downstream from the transcriptional start site (*dCLA30*3′UTR1069) was used for initial mutant studies. A line that contains a smaller insertion of approximately 500 bp was generated by imprecise excision of the transposable element (*dCLA30*ex80). A hairpin RNAi construct targeting the 325 bp of the first exon with no reported off target knock downs (UAS-*dCLA30*IR) was used for RNAi knock down studies. (TIF)

**Figure S2** Complex I loss due to *dCLA30* mutation persists in adult flies. The *dCLA30*′mut* mutation results in specific loss of complex I holoenzyme band in BN-PAGE of adult male flies, 2 days post eclosion. In contrast, controls (+) and cDNA rescue flies show the presence of the complex I holoenzyme band. (M = molecular size marker, CV2 = complex V dimer, CI = complex I, CV1 = complex V monomer, CII = complex III, CIV = complex IV, CIII = complex II, 10, 25, 50 μg of total mitochondrial protein in successive lanes for each genotype). (TIF)

**Figure S3** Expression of ND11 does not affect complex I assembly. Expression of a UAS-ND11 construct does not affect the absence of the complex I holoenzyme band in a *dCLA30*′mut* mutant background. (M = molecular size marker, CV2 = complex V dimer, CI = complex I, CV1 = complex V monomer, CII = complex III, CIV = complex IV, CVI = complex II, mitochondria from 2.5, 5, and 10 larvae equivalents in successive lanes). (TIF)

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**Author Contributions**

Conceived and designed the experiments: JC JHH SB DWW. Performed the experiments: JC JHH JG. Analyzed the data: JC JHH JG DWW. Wrote the paper: JC JHH DWW.

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