A multi-systemic mitochondrial disorder due to a dominant p.Y955H disease variant in DNA polymerase gamma

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Abstract

Mutations in the mitochondrial DNA polymerase, POLG, are associated with a variety of clinical presentations, ranging from early onset fatal brain disease in Alpers syndrome to chronic progressive external ophthalmoplegia. The majority of mutations are linked with disturbances of mitochondrial DNA (mtDNA) integrity and maintenance. On a molecular level, depending on their location within the enzyme, mutations either lead to mtDNA depletion or the accumulation of multiple mtDNA deletions, and in some cases these molecular changes can be correlated to the clinical presentation. We identified a patient with a dominant p.Y955H mutation in POLG, presenting with a severe, early-onset multi-systemic mitochondrial disease with bilateral sensorineural hearing loss, cataract, myopathy, and liver failure. Using a combination of disease models of Drosophila melanogaster and in vitro biochemistry analysis, we compare the molecular consequences of the p.Y955H mutation to the well-documented p.Y955C mutation. We demonstrate that both mutations affect mtDNA replication and display a dominant negative effect, with the p.Y955H allele resulting in a more severe polymerase dysfunction.

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Introduction

The mitochondrial DNA polymerase γ (POLγ) is required for replication of the mitochondrial genome (mtDNA). The holoenzyme consists of the catalytic subunit POLγA, encoded by the POLG gene (MIM 174763), and by the dimeric processivity factor POLγB, encoded by the POLG2 gene (MIM 604983) (1). POLγA belongs to the family A polymerases, with an N-terminal 3’–5’ exonuclease domain, a central linker domain and a C-terminal polymerase domain (2). Replication of the mitochondrial genome is independent of the cell cycle with individual mtDNA molecules being randomly selected for replication, a phenomenon referred to as relaxed replication (3–6). The total mtDNA copy number, however, is maintained at a relatively constant level.

Defects in mtDNA replication or nucleotide metabolism can lead to rearrangements, deletions, point mutations, or depletion of mtDNA, often resulting in mitochondrial dysfunction and ultimately mitochondrial disease (7). Although several factors involved in mtDNA replication have been associated with mitochondrial diseases, mutations in POLG are most common with close to 230 different disease-causing mutations reported (8) [recently summarized in (9)]. The associated clinical symptoms can be quite variable, both with respect to disease onset and clinical presentation, and cause a number of different disease entities, such as Alpers syndrome (MIM 203700), mitochondrial neurogastrointestinal encephalopathy (MNGIE: MIM 613662), sensory ataxic neuropathy, dystarthritis and opthalmo-paresis (SANDO: MIM 607459), spino-cerebellar ataxia-epilepsy (SCAE: MIM 607459), and chronic progressive external ophthalmoplegia (CPEO: MIM 157640 and MIM 258450). On a molecular level, mutations in POLG lead to the accumulation of multiple mtDNA deletions or mtDNA depletion (5,9), which in turn cause reduced oxidative phosphorylation (OXPHOS).

CPEO is the most common mitochondrial myopathy, defined by a progressive bilateral ptosis and diffuse, symmetric reduction in ocular motility, often associated with additional symptoms, e.g. hearing loss and ataxia [reviewed in (10)]. About half of all CPEO cases are inherited and disease-causing mutations at seven different loci have so far been identified, including loci coding for the mitochondrial adenine nucleotide translocator 1 (SLC25A4: MIM 103220) (11), the mitochondrial DNA helicase, TWINKLE (TWNK: MIM 606075) (12) and POLγA (13). The p.Y955C mutation of human POLG results in adult onset autosomal dominant CPEO (13) as well as premature ovarian failure (14) and is associated with the accumulation of multiple mtDNA deletions in affected patients (13,15). Mutagenesis experiments identified Y955, together with residues R943, L947 and A957 of POLγA to be essential for nucleotide specificity, and processivity (16–21), with the Y955C mutation resulting in replicative stalling and formation of multiple mtDNA deletions (21).

We here identify an autosomal dominant p.Y955H mutation in POLγA, leading to a severe multi-systemic mitochondrial disease with bilateral sensorineural hearing loss, cataract, myopathy, liver failure and feeding difficulties that required a percutaneous endoscopic gastrostomy. He had a normal intellectual capacity and development. Measurements of mitochondrial OXPHOS activity in a skeletal muscle biopsy was unremarkable at 10 months of age, but a reduced ATP production rate was determined in a second biopsy around 1 year later (Fig. 1D and E). Morphological analysis confirmed a mitochondrial dysfunction, exhibiting a high number of COX negative muscle fibres (Fig. 1F) and ragged-red fibres (Fig. 1G).

The p.Y955H mutation in POLγA is associated with early onset multi-systemic mitochondrial disease

Sequencing of the POLG locus in subject 1 revealed that the patient was heterozygous for the previously reported c.2864A>G, p.Y955C, mutation (13) (Fig. 1H). Southern blot analysis showed the accumulation of multiple mtDNA deletions (Supplementary Material, Fig. S1B), while mtDNA sequencing was unremarkable. Sequencing of POLG in subject 2 detected a previously unreported mutation, c.2863T>C, p.Y955H, additionally revealing a heterozygous p.Q1236H mutation, previously shown to be benign (22). Detailed allele-specific analysis revealed that the p.Y955H mutation was in cis with p.Q1236H on the paternal POLG allele but absent in the father (Table 1). The p.Y955H mutation had thus occurred de novo. Multiplex ligation-dependent probe amplification (MLPA) and cDNA analysis of POLG did not detect any other disease-causing mutation in the gene. Whole exome sequencing on DNA samples from subject 2, was performed as described previously (23,24), but revealed no other potentially disease-causing mutation involved in mitochondrial or metabolic diseases.

Previous reports suggested that mutations affecting position Y955 in human POLγA can lead to an increase in incorrect nucleotide incorporation, but cloning and sequencing of mtDNA from human muscle samples showed no difference in point mutation load, when compared with control samples (Supplementary Material, Table S1).

Mutations at p.Y873 are recessive lethal in Drosophila melanogaster

In order to understand the molecular and physiological consequences of the p.Y955H mutation we generated Drosophila melanogaster (Dm) models for both the p.Y955C and p.Y955H mutations (Y955 in human POLγA corresponds to Y873 in DmPOLγA). To this end we targeted the Dm yamas locus, coding for DmPOLγA, following a previously described procedure (25) (for details see materials and methods) (Supplementary Material, Fig. S2).
Flies heterozygous for the p.Y873C and p.Y873H mutations did not show any obvious phenotypic abnormalities. Eclosure rates were comparable to control flies (Fig. 2A), although flies heterozygous for p.Y873C mutation were developmentally delayed (Fig. 2B). Lifespans (Fig. 2C) were normal even after 15 generations of intercrossing (Supplementary Material, Fig. S3), whereas Dm lines homozygous for either of the two mutations were larval lethal at the third instar larval stage (Fig. 2D).

The p.Y873H and p.Y873C mutations lead to mtDNA depletion in Drosophila melanogaster

The p.Y955C mutation causes multiple mtDNA deletions in humans (13,15) as observed in muscle samples from subject 1 (Supplementary Material, Fig. S1B). In contrast, no multiple deletions were observed in muscle samples from subject 2, carrying the p.Y955H mutation (Supplementary Material, Fig. S1B). Interestingly, flies carrying the p.Y873H or p.Y873C mutation did not present multiple deletions; neither as heterozygous flies (Fig. 3A) nor as homozygous larvae (Fig. 3B). However, qRT-PCR and Southern blot analysis revealed severe mtDNA depletion in L3 larvae homozygous for either of the two mutants, with a

| Nucleotide | Protein |
|------------|---------|
| Subject 2  | c.[2863T>C; 3708G>T];[=] p.[Y955H; Q1236H];[=] |
| Father     | c.[3708G>T];[=] p.[Q1236H];[=] |
| Mother     | c.[=][=] p.[=][=] |

Flies heterozygous for the p.Y873C and p.Y873H mutations did not show any obvious phenotypic abnormalities. Eclosure rates were comparable to control flies (Fig. 2A), although flies heterozygous for p.Y873C mutation were developmentally delayed (Fig. 2B). Lifespans (Fig. 2C) were normal even after 15 generations of intercrossing (Supplementary Material, Fig. S3), whereas Dm lines homozygous for either of the two mutations were larval lethal at the third instar larval stage (Fig. 2D).
milder reduction in the heterozygous state (Fig. 3C and D). Heterozygous flies (p.Y873H or p.Y873C) showed an mtDNA depletion, which was somewhat more pronounced after 15 generations intercrossing (Fig. 3C).

POLγ:Y955H has lower affinity to DNA than WT POLγ or POLγ:Y955C

We next investigated the biochemical consequences of the p.Y955C mutation and for comparison we also analysed the previously characterized p.Y955C mutation (20,21). To this end, wild-type POLγA protein and mutant derivatives (POLγA:Y955C and POLγA:Y955H) were expressed and purified. DNA binding properties of all three POLγA versions were first assessed in the absence or presence of recombinant POLβ, using an electrophoretic mobility shift assay (EMSA). Both POLγA:Y955C and POLγA:Y955H bound to a primed DNA template independently of POLβ (Fig. 4A). However, the $K_d$ (equilibrium dissociation constant) for binding to the template, was higher for POLγA:Y955H than for WT POLγA and POLγA:Y955C (Supplementary Material, Fig. S4).

Modifications at position Y955 are associated with difficulties of incorporating dATP

We previously demonstrated that the p.Y955C mutation leads to stalling at low dNTP concentrations. POLγA:Y955C is especially sensitive to low dATP concentrations, and the enzyme enters a polymerase/exonuclease idling mode at dATP insertion sites (21). To investigate if this is a general phenotype for mutations affecting Y955, we performed primer extension experiments, using two different primed DNA templates (Fig. 4B). In the absence of dNTPs, WT POLγA will use its 3′–5′ exonuclease activity to digest the labelled primer, but in the presence of dNTPs, the polymerase will initiate primer elongation. In this assay, POLγA:Y955H behaved as previously demonstrated for POLγA:Y955C and required higher dNTP concentrations to produce full-length products (Fig. 4B, upper panel). Increasing the number of thymines on the template strand, promoted stalling of both mutant polymerases (Fig. 4B, lower panel), consistent with preferred stalling at dATP insertion sites (Fig. 4B, compare upper and lower panel) (21).

POLγA:Y955H has a dominant negative effect on the replisome at low dNTP concentrations

In the presence of mtSSB and the TWINKLE helicase, WT POLγA in complex with POLβ is able to replicate a circular dsDNA template containing a preformed replication fork (Fig. 5A, lanes 1–4 and 13–16) (26). Neither POLγA:Y955H nor POLγA:Y955C could support DNA synthesis (Fig. 5A, lanes 5–8 and 17–20, respectively). In agreement with previous reports, POLγA:Y955C displayed a dominant negative effect on DNA synthesis in the presence of WT POLγA (Fig. 5A, lanes 21–24) (21). The dominant negative effect was less pronounced with POLγA:Y955H even when the mutant was added at a 3:1 molar ratio relative to WT POLγA (compare lanes 9–12 with lanes 21–24 in Fig. 5A). Postmitotic tissues contain lower dNTP concentrations than the 10 μM used here (27,28). We therefore repeated the experiment using 1 μM dNTP. At this concentration, POLγA:Y955H also displayed a dominant negative effect on WT POLγA activity (Fig. 5B, lanes 9–12). Furthermore, we also observed an accumulation
of shorter-than-input length DNA products (Fig. 5B, lane 12), suggesting that POL\textsubscript{c}A:Y955H causes template degradation at lower dNTP concentrations.

**POL\textsubscript{c}A:Y955C and POL\textsubscript{c}A:Y955H display reduced ligation efficiency**

At the end of replication, DNA ends are ligated to produce a closed circular mtDNA molecule. We have previously shown that the 3’–5’ exonuclease and 5’–3’ polymerase activities of POL\textsubscript{c}A must be correctly balanced in order to support ligation (29). Since mutations affecting position Y955 shift the balance towards the exonuclease activity, we decided to monitor effects on ligation. To this end we performed coupled polymerase-ligation assays, using a template consisting of a radioactively 5’-labelled 60-mer hybridized to a circular 100-mer (Fig. 6A). On this template, DNA polymerase needs to extend the 3’ end to full circle and then terminate at the downstream 5’ end, in order to produce a ligateable nick. WT POL\textsubscript{c}A extended the 60-mer and provided ligateable ends already at 1 μM dNTP (Fig. 6B, lanes 2–4). The POL\textsubscript{c}A:Y955C mutant was also capable of generating ligateable full-length products, but it needed 100 times more dNTP, which reasons well with its lower dNTP affinity (Fig. 6B, lanes 5–7). The POL\textsubscript{c}A:Y955H mutant on the other hand had problems to fully extend the 60-mer even in the presence of 100 μM dNTP and were therefore unable to create ligateable ends (Fig. 6B, lanes 8–10). Here again, we observed that both mutants have increased exonuclease activity at low dNTP concentrations (see Fig. 6B, lanes 5 and 8). Next, we asked whether the presence of the mutants could affect the ability of the WT to create ligateable ends, which would reflect a typical situation in vivo. To answer this question, we performed time-dependent ligation assays at 1 μM dNTPs, where WT POL\textsubscript{c}A was mixed with either of the two mutants (Fig. 6C). At this dNTP concentration only the WT protein is able to extend the primer. Addition of POL\textsubscript{c}A:Y955C or POL\textsubscript{c}A:Y955H severely inhibited polymerization and the ability of WT POL\textsubscript{c}A to create ligateable nicks (Fig. 6C and D).

**Discussion**

Mitochondrial diseases form a highly diverse group, with a range of clinical symptoms and different ages of onset. They are predominantly monogenic disorders and recent advances in

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**Figure 3.** Mutations at p.Y873 lead to mtDNA depletion in flies. (A) Southern blot analysis of mtDNA of heterozygous mutant flies. (B) Long-range PCR of homozygous mutant and control larvae. Primers were situated as indicated. (C) Southern blot analysis of mtDNA from homozygous mutant larvae. (D) Relative mtDNA levels determined by qPCR in L3 larvae (L0), or flies intercrossed for 1 (F1) or 15 (F15) generations. TaqMan probes used as indicated. Error bars are ±SD (*P<0.05, **P<0.01, ***P<0.001, n=5).
In vitro DNA binding and polymerization properties of POL\textsubscript{c} Y955C and POL\textsubscript{c} Y955H. (A) Electrophoretic mobility shift assays showing that POL\textsubscript{c} Y955C and POL\textsubscript{c} Y955H bind DNA both in absence (lanes 5 and 8) and presence (lanes 6 and 9) of POL\textsubscript{c} B. Lanes 1, 4 and 7 contained no protein. Lower panel: schematic representation of the DNA template. The asterisk indicates the \textsuperscript{32}P label on the 5' end of the 20-mer. (B) Coupled 3'–5' exonuclease/polymerase assays show that both POL\textsubscript{c} Y955C and POL\textsubscript{c} Y955H required higher dNTP concentrations than WT POL\textsubscript{c} A to synthesize DNA (all experiments were performed in the presence of POL\textsubscript{c} B). Reactions were run on denaturing 15% PAGE. Below: scheme of the primer-extension reaction starting from the 20-meric primer to produce a 35-mer product.

Rolling circle replication assay reveals dominant negative effect of POL\textsubscript{c} Y955C and POL\textsubscript{c} Y955H. (A) In vitro replication reactions performed at 10 \textmu M dNTP and indicated POL\textsubscript{c} versions (all experiments were performed in the presence of POL\textsubscript{c} B). Lanes 1–4, WT POL\textsubscript{c} A; lanes 5–8; POL\textsubscript{c} A:Y955H; lanes 9–12, WT POL\textsubscript{c} A and POL\textsubscript{c} A:Y955H; lanes 13–16, WT POL\textsubscript{c} A; lanes 17–20, POL\textsubscript{c} A:Y955C; lanes 21–24, WT and POL\textsubscript{c} A:Y955C. (B) As in (A) but at 1 \textmu M dNTP.
sequencing technologies have revolutionized the diagnosis of patients with mitochondrial diseases. Despite these advances, the correlation between genotype and phenotype remains difficult to predict and combined with the large heterogeneity of the human genome makes functional validation of novel disease variants essential. A complicating factor is that a biochemical defect is not always observed in patient samples (30–32). To date, almost 230 mutations have been identified in the human POLG gene alone (8), with both dominant and recessive inheritance patterns. A substantial number of mutations are also reported to only prompt clinical symptoms, when inherited in trans with other mutations as compound heterozygous, and it is thus important to fully characterise variants. We here used a combination of in vitro biochemistry, model organisms and
patient investigations to demonstrate pathogenicity of a mutation in POLG, which causes a tyrosine to histidine change at position 955 in the human POLA protein.

The tyrosine residue at position 955 of POLA is essential for nucleotide incorporation into the growing DNA strand. An interaction between Y955 and E895 is required for the binding of the incoming nucleotide (Fig. 6f). The hydroxyl group of Y955 interacts with the carboxyl group of E895, which is located adjacent to the catalytic helix (residues 943–955). In T7 DNA polymerase this glutamate selectively stabilizes the incoming nucleotide. The polar contacts, rotamer conformation and steric influence of E895 likely contribute to the formation of a tight binding pocket. In addition, Y955 may aid catalysis via E895-mediated alignment of the catalytic helix towards the nucleotide.

The severity of the p.Y955C and p.Y955H mutations is due, in part, to the steric influence of Y955 in the active site (33). The large phenyl ring excludes solvent from the active site, while providing a tight binding pocket for the nucleotide. Removal of this phenyl ring results in a loss of favourable π–π stacking interactions between the incoming base and Y955 (negatively affecting nucleotide binding), which likely destabilises the active site. Differences in Kc were also observed between the p.Y955C and p.Y955H mutated versions of POLA, with the latter having 5-fold weaker binding to a primed DNA template. This is not surprising given that the introduction of a polar histidine into an apolar buried cavity can negatively affect both protein stability and ligand binding (34–36). Replacement of Y955 with histidine is thus likely responsible for (i) loss of steric packing, (ii) loss of a stabilizing hydrogen bond with E895, (iii) loss of π-stacking interactions with Y951 and (iv) electrostatic repulsion within the binding cavity (Fig. 6e, lower left panel).

The p.Y955C mutation leads to reduced polymerase activity and replication stalling (16–21). Previous data also suggested that the p.Y955C mutation could cause increased mtDNA mutation loads (37–40), but this conclusion was not supported by data obtained here from human muscle.

The p.Y955C mutation is consistently associated with multiple deletions in mtDNA, but we could not reproduce this phenotype in flies harbouring the p.Y955C mutation. Neither did the p.Y955H mutation lead to multiple deletions in either human skeletal muscle samples or in the corresponding fly model. However, this is not surprising, since it has been shown that the number of deletion-molecules increases with age. The lack of deletion in the flies and in the p.Y955H patient could simply reflect the short lifespan of flies and the very young age of the patient. In humans, multiple mtDNA deletions do not become visible in Southern blots until ≈20 years of age (41). Nevertheless, we observed mtDNA depletion with both p.Y873C and p.Y873H, consistent with observations made in murine and yeast models of the p.Y955C mutation (18,37–40). Depletion is and p.Y873H, consistent with observations made in murine and

**Figure 7.** A schematic model for the effects of mutant POLA/Y955C and POLA/Y955H on replication fork progression. The mitochondrial replication machinery at the fork is dynamic and POLA is regularly coming on and off the 3’ end of the newly synthesized DNA. When WT POLA (purple) is present at the replication fork of a heterozygous patient, it prevents mutant POLA (orange) from binding and mtDNA synthesis can therefore progress (left panel). Once mutant POLA binds the primed template, it enters an idling mode (repeated cycles of incorporation and degradation at the primer terminus) and simultaneously blocks the WT protein from accessing the 3’ end (right panel). mtSSB is green and TWINKLE is blue.

are subtle, they do lead to fundamentally different clinical presentations. Our data therefore further demonstrates the complexity of mitochondrial diseases and how difficult it is to predict clinical phenotypes based on molecular defects in these patients.

**Materials and Methods**

The Regional Ethics Committee at Karolinska Institutet approved the use of patient material in this study.

**DNA and RNA isolation from patients**

Genomic DNA from the patients was isolated from skeletal muscle, blood or fibroblasts, using QIAamp DNA Mini Kit (Qiagen). For RNA analysis, total RNA was isolated from fibroblasts using RNeasy Mini Kit (Qiagen).

**Molecular analysis in patients**

The entire mtDNA sequence from the patients was determined from total DNA isolated from skeletal muscle, while the POLG gene was sequenced from total DNA isolated from the patient’s blood. MtDNA was amplified by PCR in 28 overlapping M13-tailed fragments as described before (42). The coding exons 2–23 of POLG were amplified by PCR using M13-tagged intronic primers. Subsequent sequencing of all PCR products was carried out with M13 primers, using the BigDye version 3.1 sequencing kit (Applied Biosystems) on a 3130xl Genetic Analyser (Applied Biosystems). MtDNA sequence data were compared with the revised Cambridge reference sequence for human mtDNA (GenBank NC_012920.1) and for POLG to the reference sequence NM_002693.2. MLPA analysis, using kit P010 (MRC Holland, Amsterdam), for detection of deletions or duplications of each exon of the POLG gene was performed according to the manufacturer’s protocol as described earlier (43). RT-PCR was performed on isolated RNA, using GeneAmp RNA PCR kit (Applied Biosystems) and exons 2–23 of POLG cDNA were amplified by PCR in 10 overlapping fragments. The size and amount of the PCR products was compared with a control sample in 2% agarose gel. Sequencing of the PCR products from cDNA was performed as described for genomic DNA above. Southern blot analysis was done using aliquots of 0.2 μg total DNA isolated from skeletal muscle, which were digested with the restriction enzyme DdeI.
enzyme PvuII (New England Biolabs) and fractionated by electrophoresis in 0.5% agarose gels. The DNA was then transferred to Hybond XL nylon filter (GE Healthcare) by capillary blotting under standard procedures. The filter was hybridized with an equimolar mix of radiolabelled mtDNA probes corresponding to nucleotides 1–12,640, as described previously (44).

Expression and purification of recombinant human proteins

Mutated versions of POLγA were constructed using the QuikChange Lightning Site-directed mutagenesis kit according to the provided protocol (Agilent Technologies) and confirmed by sequencing (Eurofins MWG Operon). Recombinant baculoviruses coding for TWINKLE, mitochondrial single-stranded DNA-binding protein (mtSSB), WT POLγA, POLγA:Y955C, POLγA:Y955H and POLγB were expressed in Sf9 cells and purified as previously described (26).

EMSA and $K_d$ determination

The affinity between POLγ (POLγA in complex with POLγB) and a primer template was monitored using an EMSA as previously described (25). Each experiment was repeated at least three times. Band intensities representing unbound and bound DNA were quantified using Fujifilm Multi Gauge V3.1 software. The fraction of DNA bond in each reaction was plotted versus the concentration of POLγ. Data were fitted with the binding equation (Fraction bound = (MaxB × [POLγ])/(MaxB + [POLγ])) using EXCELS Add-in ‘Solver’ to perform non-linear regression and obtain values for $K_d$ (as the value corresponding to the midpoint of MaxB) and using MaxB set to 1 (the fraction bound at which the data plateaus).

Coupled 3′–5′ exonuclease/polymerase assays

A 20-mer (5′-GGG TCG AGT CTA GAG GAG CC-3′) labelled at the 5′ end with [γ-32P] ATP was annealed to a 35-mer oligonucleotide containing a poly-dT stretch, 5′-TTT TTT TTT TAT CCT CCG TCC TCT AGA AGC GC-3′, or an oligonucleotide without a poly-dT stretch (5′-GAC AAC CAG CCG GCC TCC TCT AGA AGC GC-3′, as illustrated in Fig. 4B. These two templates were used as substrates to investigate DNA polymerization and 3′–5′ exonuclease activity as previously described (21).

In vitro rolling circle DNA replication

The reaction mixtures (20 μl) contained 10 fmol of rolling circle template and reactions were performed as described previously except that lower dNTP concentrations (1 μM or 10 μM) were used. POLγ (75 fmol WT POLγA in complex with 300 fmol POLγB, 225 fmol POLγA:Y955C in complex with 675 fmol POLγB, or 225 fmol POLγA:Y955H in complex with 675 fmol POLγB) were added as indicated in the figure legends (26).

Coupled DNA synthesis-ligation assay

The synthesis-ligation assay was adapted from (29). Here a circular DNA template was used instead of a linear (Fig. 4A). The closed circular template was constructed by circularizing the 100 nt oligonucleotide (5′-GAG GGG TAT GTG GCC ACA GGA CTT AAA CAC ATC TCT GCC AAA CCC AAA AAC AAA GAA CCC TAA CAC CAG CCT AAC CAG ATT TCA AAT ATC ACC GAA T-3′) with CircLigase TMssDNA ligase (Epiterror), followed by annealing to a 32P-labelled 60-mer (5′-TCT GGT TAG GCT GTT AGG GTT CTT GTG TTT TGG GTT CAG AGA TGT TTA GTG-3′). The reactions were performed in a volume of 20 μl containing 20 mM Tris–HCl (pH 7.5), 1 mM DTT, 0.1 mg/ml BSA, 10 mM MgCl2, 0.5 mM ATP, 4 units of T4 DNA ligase, 10 fmol circular DNA substrate, indicated POLγ version, and varying amounts of dNTPs as specified in the figure legends. In the mixing experiments, 75 fmol WT POLγA, POLγA:Y955H, or POLγA:Y955C in complex with 300 fmol POLγB were pre-incubated in the mixture on ice for 10 min before an additional 75 fmol of WT POLγA in complex with 300 fmol POLγB was added to each reaction. Reactions were incubated for 30 min if nothing else is indicated in the figure legends and terminated with 2× stop buffer (Formamide with 10 mM EDTA, 0.025% bromophenol blue, 0.025% xylene cyanol). Samples were run in 7 M urea/10% polyacrylamide gels and visualized with a PhosphorImager or autoradiography.

Drosophila stocks and maintenance

All genomically engineered fly strains were constantly back-crossed into a white Dahomey Wolbachia-free (wDahT) WT strain. All fly stocks were maintained at 25 °C on a 12:12 h light/dark cycle with 60% humidity and fed on a sugar/yeast/agar (SYA) diet (25).

Generation of genomically engineered DmPOLγA flies

Human POLG sequence (NP_002684.1) was aligned to the Drosophila melanogaster homolog (tamas; NP_476821.1) and the fly-equivalent position for p.Y955 was identified at position p.Y873. Genetically modified flies were generated essentially as described previously (25,45). We previously cloned the Tamas locus into the fly-specific targeting vector pGEattB-GMR (25), which was here modified by site-directed mutagenesis to introduce the p.Y873C and p.Y873H mutations. Mutant variants of the tamas gene were then injected into the embryos of DmPOLγA (Tamas) KO founder embryos (25), expressing ε31 integrase by the in-house Drosophila transgenic core facility. Positive flies were selected by eye colour and confirmed by Southern blot/sequencing.

Life-span determination

Newly eclosed adult flies were mated for 2 days before they were sorted for lifespan analyses. Two hundred females per genotype were individually picked and placed into vials with SYA food. The number of eclosed flies was scored every 12 h. At least five biological replicates were done for each genotype.

Hatching and eclosion rates

Fly development was assessed as follows. Flies were allowed to lay eggs on grape juice agar plates for 2 h. One hundred eggs per genotype were individually picked and placed into vials with SYA food. The number of eclosed flies was scored every 12 h. At least five biological replicates were done for each genotype.

DNA isolation and southern blot analysis from flies

For relative mtDNA copy number determination, total DNA extractions were prepared from L3 larvae using DNeasy Blood and
Tissue Kit (Qiagen). Five biological replicates, each with 10 larvae, were prepared for each genotype. Quantification of mtDNA levels was done using SYBR-Green qPCR analyses and primers targeting the mitochondrial encoded genes 12S, COX1 and CytB, and normalized to the nuclear encoded histone gene. All data were normalized against wild-type levels.

For Southern blot analysis, total DNA was extracted from 20 to 30 L3 larvae, either using the DNeasy Blood and Tissue kit (Qiagen), or by homogenizing samples with a tissue grinder in 400 μl of buffer A (100 mM Tris–HCl, pH 7.5; 100 mM EDTA; 100 mM NaCl and 0.5% SDS). After incubation at 65°C for 30 min, 800 μl of freshly prepared Buffer B (4 μl of 5 M KOAc and 10 μl of 6 M lithium chloride) was added and samples were left on ice for 120 min. After incubation, samples were centrifuged at 12 000g for 15 min and supernatant was transferred into a new tube. About 540 μl of isopropanol was added to the supernatant and samples were further centrifuged at 12 000g for 15 min. Pellets were washed with 70% ethanol, dried and resuspended in 100 μl nuclease-free water containing 20 mg/ml RNase A. After incubation at 37°C for 1 h, samples were stored at -4°C.

Approximately 1–3 μg of total DNA was cut using either XhoI or Syl restriction endonuclease (NEB). Digestions were run on 0.8% agarose gels, and blotted to Hybond-N membrane (Amer sham Bioscience). COXI or ND6-CytB (positions 10169–11169) were used as probes, and signals were visualized by autoradiography.32P-labelling of Southern probes was done according to manufacturer’s instructions (Prime-IT II Random Prime Labelling Kit, Agilent).

Long-Range PCR was performed using primers: Dmel_Long 1: 5’GCT CTA AAA TAT GTA CAC ATC GCC C3’, Dmel_Long 2: 5’AAT TAA GCT ACT GGG TTC ATA CCC C3’, Dmel_Long 3: tions (Prime-IT II Random Prime Labelling Kit, Agilent).

Long-Range PCR was performed using primers: Dmel_Long 1: 5’AAT TAA GCT ACT GGG TTC ATA CCC C3’, Dmel_Long 2: 5’GCT CTA AAA TAT GTA CAC ATC GCC C3’, Dmel_Long 3: 5’TTT GAA GCA GCT GCA GTA TAT TGA C3’ and Dmel_Long 4: 5’AGA GCT TGA CAT TGA AGA TGT TAT GGA C3’.

Supplementary Material

Supplementary Material is available at HMG online.

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Conflict of Interest statement. None declared.

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