Mutualism Breakdown by Amplification of Wolbachia Genes

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Abstract

Most insect species are associated with vertically transmitted endosymbionts. Because of the mode of transmission, the fitness of these symbionts is dependent on the fitness of the hosts. Therefore, these endosymbionts need to control their proliferation in order to minimize their cost for the host. The genetic bases and mechanisms of this regulation remain largely undetermined. The maternally inherited bacteria of the genus Wolbachia are the most common endosymbionts of insects, providing some of them with fitness benefits. In Drosophila melanogaster, Wolbachia wMelPop is a unique virulent variant that proliferates massively in the hosts and shortens their lifespan. The genetic bases of wMelPop virulence are unknown, and their identification would allow a better understanding of how Wolbachia levels are regulated. Here we show that amplification of a region containing eight Wolbachia genes, called Octomom, is responsible for wMelPop virulence. Using Drosophila lines selected for carrying Wolbachia with different Octomom copy numbers, we demonstrate that the number of Octomom copies determines Wolbachia titers and the strength of the lethal phenotype. Octomom amplification is unstable, and reversion of copy number to one reverts all the phenotypes. Our results provide a link between genotype and phenotype in Wolbachia and identify a genomic region regulating Wolbachia proliferation. We also prove that these bacteria can evolve rapidly. Rapid evolution by changes in gene copy number may be common in endosymbionts with a high number of mobile elements and other repeated regions. Understanding wMelPop pathogenicity and variability also allows researchers to better control and predict the outcome of releasing mosquitoes transinfected with this variant to block human vector-borne diseases. Our results show that transition from a mutualist to a pathogen may occur because of a single genomic change in the endosymbiont. This implies that there must be constant selection on endosymbionts to control their densities.

Author Summary

Insects frequently carry intracellular bacteria that are passed from generation to generation through their eggs. These intracellular symbionts can be beneficial or parasitic, but because of their mode of transmission, they are always dependent on the reproduction of their carriers. They therefore have to control their own growth in order to minimize
deleterious effects on the host. Bacteria of the genus *Wolbachia* are the most common maternally transmitted intracellular bacteria in insects. Most *Wolbachia* variants that are naturally associated with the fruit fly *Drosophila melanogaster* are benign to their hosts and provide them with protection against viruses. However, *w*MelPop is a virulent *Wolbachia* variant that over-replicates massively and shortens the lifespan of its fruit fly host. Here we show that amplification of a *Wolbachia* genomic region containing eight genes—called Octomom—is responsible for the pathogenic effects of *w*MelPop. Our results provide a link between genotype and phenotype in *Wolbachia* and show that virulence in symbionts can be simply caused by increases in gene copy number. These results also indicate that gene copy number variation may be a common mechanism underlying rapid evolution of intracellular symbionts.

**Introduction**

Vertically transmitted bacterial endosymbionts are widespread in arthropods, particularly in insects [1]. Many endosymbionts are mutualists and confer a fitness advantage to the host. The benefits may range from metabolic provisioning to protection against pathogens [2]. Other symbionts act as parasites and manipulate host reproductive biology in order to increase the relative fitness of their carriers [3]. In both cases, the density of endosymbionts within hosts is a crucial factor determining their prevalence in host populations [4,5].

Symbiont densities are determined by host and symbiont genetic diversity and environment [4,6–10]. These densities are under selection at the level of the host and the symbiont. Interestingly, there are conflicting selective forces at the level of the symbiont. Higher symbiont densities are associated with higher transmission fidelity and stronger phenotypes induced in the host [4,5,8,11–17]. Theoretically, this should lead to a selection for higher densities. On the other hand, high symbiont levels may have a negative impact on host fitness [8,9,17–19]. Since vertical transmission leads to dependence of the symbiont on the fitness of the host, it is advantageous for endosymbionts to limit their densities and consequently minimize the cost to their hosts. Thus, a key question in the field of host–microbe interactions is how symbionts regulate their replication and resulting densities to achieve an equilibrium between these opposing selective forces.

*Wolbachia* is conceivably the most prevalent bacterial endosymbiont of insects [20,21], and its interactions with hosts have been studied extensively. *Wolbachia* is maternally transmitted and exhibits a range of phenotypes. These include cytoplasmic incompatibility and other reproductive manipulations that potentiate *Wolbachia* spread in host populations [22]. Some *Wolbachia* strains have also been shown to be metabolic mutualists [23] or to protect insects from viral infections [24–27]. The *Wolbachia* strain infecting *Drosophila melanogaster*, *w*Mel, exerts only a weak cytoplasmic incompatibility in laboratory conditions [28], and this reproductive manipulation seems not to be expressed in field conditions [29]. Since cytoplasmic incompatibility cannot explain *Wolbachia* prevalence in *D. melanogaster* populations [28,30], it was suggested that *w*Mel exerts positive fitness effects on its hosts [29]. More recently, it was shown that *w*Mel provides *Drosophila* with strong resistance to systemic and oral infection with the natural pathogen *Drosophila* C virus (DCV) [24,25,31]. This protection extends to RNA viruses of different families [24,25,27,32], indicating that *w*Mel protects against a wide range of RNA viruses. Some of the fastest evolving genes in *D. melanogaster* are involved in antiviral RNA interference and are under strong positive selection [33]. Therefore, viruses seem to be a strong selective force in *D. melanogaster*. Moreover, several viruses, including DCV,
have been isolated from natural populations of *D. melanogaster* [34–36]. Although there are no data regarding *Wolbachia* antiviral protection in natural populations, the *D. melanogaster–wMel–DCV* interaction fulfills many of the criteria for defensive mutualism [37]. Therefore, antiviral protection may be the cause of wMel maintenance in *D. melanogaster* natural populations.

Natural wMel variants have a small effect on host longevity, yet they provide a strong antiviral protection [8]. This protection is positively correlated with *Wolbachia* density: the higher the titers of *Wolbachia*, the higher the antiviral protection [8,17,18,26,38–40]. On the other hand, high endosymbiont densities can have a cost in the absence of viral infection, and *Wolbachia* variants conferring strong protection often shorten the lifespan of the flies [8,18]. There is thus a fine balance between density, benefit, and cost to the host.

The wMel variant wMelPop breaks this balance and is clearly pathogenic: it over-proliferates and dramatically shortens the lifespan of infected flies [8,19,41,42]. wMelPop is, hence, an exceptional vertically transmitted symbiont. Its uniqueness was immediately recognized as providing a tool to better understand regulation of vertically transmitted symbionts and the biology of *Wolbachia* [43].

Understanding the cause of the wMelPop phenotype and regulation of *Wolbachia* densities is also important from an applied perspective. Several *Wolbachia* strains, including wMelPop, have been transinfected into mosquito vectors of human diseases, where they can interfere with arboviruses or other pathogens [44–52]. The purpose of this research is to release *Wolbachia*-carrying mosquitoes refractory to dengue virus and are already being tested in the field [44,50,53–55]. Different variants of wMel transinfected into *A. aegypti* show a trade-off between host fitness and resistance to dengue virus. A wMelPop-derived strain gives higher resistance to dengue but has a high fitness cost, which may prevent it from stably infecting natural mosquito populations [56,57]. On the other hand, a wMel-derived strain confers lower protection to dengue virus but is able to stably invade *A. aegypti* populations [50,53–55]. Ideally, a further understanding of the system would allow researchers to use *Wolbachia* strains with a better ratio of antiviral protection to cost. Moreover, since wMelPop has been transinfected into mosquitoes [44–48], understanding the pathogenicity of this *Wolbachia* variant is crucial for predicting wMelPop dynamics in the released mosquito populations.

Finding the genetic basis of wMelPop pathogenicity is essential to understanding its phenotype. Difficulty in the functional analysis of *Wolbachia* lies in its refractoriness to genetic manipulation. Nonetheless, genomic analyses have provided insight into the cause of wMelPop pathogenicity. The first genomic map of wMelPop was published in 2003 [58], while the full genome of the similar wMel was published in 2004 [59]. Analyses of polymorphic genomic markers and whole genome assemblies have shown that wMelPop is closely related to wMelCS variants [8,60–62]. We have recently identified genetic differences between wMelPop and the closely related non-pathogenic wMelCS_b [8]. The wMelPop genome contains an amplification of a ~21-kb region, named Octomom, that includes eight *Wolbachia* genes (WD0507 to WD0514) flanked by direct repeats. This amplification in wMelPop was also described by Woolfit and colleagues [62]. Apart from this amplification, we found only one synonymous SNP unique to wMelPop (position 943,443, G>A) [8]. Therefore, we hypothesized that Octomom region amplification underlies wMelPop virulence. Gene amplification has previously been reported to change the pathogenicity of other bacteria and viruses [63–67].

Here we show strong evidence that, in support of our original hypothesis, Octomom region amplification is the cause of the wMelPop phenotypes of over-replication and pathogenicity.
Results

Currently, *Wolbachia* cannot be genetically manipulated, which hinders functional studies on *Wolbachia* genes. However, bacterial amplified DNA sequences have been described before as unstable [64], leading us to test the hypothesis that natural variation in Octomom copy number exists and causes distinct phenotypes. To detect Octomom copy number variation, we tested several single *Drosophila* females for the copy number of the Octomom gene *WD0513* in their *Wolbachia* bacteria (Fig. 1A). The copy number of *WD0513* was determined by quantitative PCR on genomic DNA samples from single flies carrying *Wolbachia*, using *Wolbachia wsp* (*Wolbachia* surface protein) as a reference gene. *wMelCS_b* samples were used as reference samples for one *WD0513* copy, based on the coverage analysis of our previous *Wolbachia* sequencing data [8]. We analyzed two fly stocks infected with *wMelPop: w^{118}* iso, derived from the original stock in the Benzer lab [19], and a DrosDel isogenic *w^{118}* iso stock into which we introgressed *wMelPop* from the *w^{118}* stock [8]. All *wMelPop* samples analyzed had at least a duplication of the Octomom region, with high variation in *WD0513* copy number between individual females, ranging from two to ten copies (Fig. 1A). This copy number corresponds to the average *WD0513* copy number in the *Wolbachia* of each individual female (thus, differences in Octomom copy number between *Wolbachia* cells within each female may exist).

To check whether the *Wolbachia* Octomom region is amplified as a unit, we tested *WD0507* and *WD0513* copy number simultaneously in individual flies. The copy numbers of the two genes are the same in each fly (Figs. 1B and S1), suggesting integrity of the Octomom region. A common mechanism of gene amplification in bacteria leads to tandem duplications and the formation of new junctions between units [64]. We detected the presence of this new predicted *WD0514–WD0507* junction by PCR and Sanger sequencing (Fig. 1C; S1 Text). These data show that Octomom copy number is highly variable and that the amplification is consistent with a tandem duplication.

**Fig 1. Individual wMelPop flies differ in Octomom copy numbers.** (A) *WD0513* copy number variability in single females from two wMelPop stocks with w^{118} and iso genetic backgrounds, relative to wsp. We tested two replicates of w^{118} stock and five replicates of iso stock. wMelCS_b iso flies were used for copy number normalization (control [ctr]). Lines are medians of the replicates. Supporting data can be found in S1 Data. (B) Relation between *WD0507* and *WD0513* abundance in single wMelPop females. Each dot represents a female, and the regression line is shown. The estimates for the fitted regression line are slope = 1.036 ± 0.041, intercept = 0.182 ± 0.204, $R^2 = 0.92$. Supporting data can be found in S2 Data. (C) PCR of the predicted *WD0514–WD0507* junction in wMelPop flies. wMelCS_b was used as a negative control. Three samples of each Wolbachia variant were used. PCR for wsp gene was used as a DNA quality control.

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To test Octomom amplification’s effect on wMelPop virulence, we established Drosophila lines carrying Wolbachia with different Octomom copy numbers. Individual females with the highest and the lowest Octomom copy number were selected throughout several generations in both w^{1118} and iso backgrounds (Figs. 2 and S2). Octomom copy number is heritable: Drosophila mothers carrying high-copy Wolbachia produce mostly offspring with high-copy Wolbachia, while the inverse is observed for mothers with low-copy Wolbachia. In the course of selection for low Octomom copy number in the w^{1118} background, we recovered a Drosophila line carrying wMelPop with only a single copy of Octomom (Fig. 2C). This single-copy Octomom line had also lost the WD0514–WD0507 junction detected in wMelPop with multiple Octomom copies (S3 Fig.). Therefore, from generation six onwards, we maintained three selection regimes: high, two, and one Octomom copy number. The wMelPop unique synonymous SNP is present in all three selection lines, including the line carrying Wolbachia with a single Octomom copy (S4 Fig.).

Taking advantage of the different selection lines, we compared the phenotypes of flies with wMelPop with different Octomom copy numbers. We predicted that the higher the copy number, the more severe the pathogenic phenotype, and that the one-copy Octomom line would be phenotypically identical to wMelCS_b. To perform these assays, we used the progeny of females individually tested for Octomom copy number. As Wolbachia wMelCS_b was associated with the iso fly genetic background and the one-copy Octomom line appeared only in the w^{1118} background, we used hybrids between iso and w^{1118} to directly compare the two (S1 and S2 Tables). All female hybrids resulting from these crosses have the same host genetic background.
Mutualism Breakdown by Amplification of Wolbachia Genes

Fig 3. Octomom copy number determines wMelPop phenotypes. (A) Lifespan of female flies with different wMelPop Octomom copy numbers, flies with wMelCS_b and Wolbachia-free controls at 29°C. Seventy females per line were analyzed; flies are the progeny from crosses between iso and w1118 lines. Bold letters on the right indicate groups with significantly different survival curves by Tukey’s test of all pairwise comparisons of Cox hazard ratios. Supporting data can be found in S4 Data. (B) Lifespan of female flies from the forward selection iso low-copy line two (two Octomom copies) and the matched reverse selection line (seven copies) at 25°C. Mixed effects Cox model fit, \( p < 0.001 \). Supporting data can be found in S5 Data. (C) Time-course analysis of Wolbachia densities in female flies with different wMelPop Octomom copy numbers, starting at eclosion (day zero). Each bar represents wsp genomic levels in 16–20 single females (progeny from crosses between iso and w1118 lines). The boxes extend from the 25th to 75th percentile, and the whiskers include all values. Statistical analysis was performed using a log-linear model, and the \( p \)-values refer to comparisons of slopes. Supporting data can be found in S6 Data. ns, non-significant. (D) Western blot with anti-WSP antibody of pools of ten 10-d-old iso female flies with three or ten Octomom copies. Drosophila tubulin was used as a loading control. (E) Survival of female flies with different wMelPop Octomom copy numbers upon viral infection at 18°C. Fifty females per line were analyzed; flies are the progeny from crosses between iso and w1118 lines. Bold letters on the right indicate groups with significantly different survival curves by Tukey’s test of all pairwise comparisons of Cox hazard ratios. Supporting data can be found in S7 Data. 

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(heterozygous between iso and w1118) and differ in the Wolbachia inherited from the mother. Two high-copy Octomom lines, one in each genetic background, were used to control for potential host-genotype-specific maternal effects. Survival data demonstrate that differences in Octomom copy number lead to differences in host longevity: the more Octomom copies, the earlier the flies die (Figs. 3A and S5A–G). The line with one Octomom copy derived from wMelPop is indistinguishable from wMelCS_b and Wolbachia-free control (Figs. 3A and S5E–G). Even a single duplication of this region is enough to significantly shorten the host lifespan (median time to death is reduced by 39%) (Figs. 3A and S5E–G). The lifespan of flies from the two high-copy Octomom lines is further reduced, and there is no difference between these two lines (Figs. 3A and S5E–G). To further test the dependence of the phenotype on Octomom copy number, we reversed the direction of the selection in selected iso lines (choosing females with wMelPop with the highest Octomom copy number from the low-copy lines and with the lowest copy number from the high-copy lines, from generation 17 onwards) (S6A Fig.), simultaneously maintaining the forward selection regime as controls (S2 Fig.). Comparison of the lifespans of females from the forward and reverse selections confirmed that Wolbachia Octomom copy number determines wMelPop pathogenicity (Figs. 3B and S6B–D). Overall, Octomom copy number negatively correlates with longevity (S7 Fig.), and by manipulating copy number we can control Wolbachia virulence.
We next asked whether \textit{Wolbachia} growth is associated with Octomom copy number. We tested \textit{Wolbachia} levels in flies carrying \textit{Wolbachia} with different Octomom copy numbers over time by real-time quantitative PCR (qPCR) (Fig. 3C). The higher the Octomom copy number, the higher the density of \textit{Wolbachia}. The levels are different at eclosion, and the growth of \textit{Wolbachia} is faster in flies with higher Octomom copy number. Both high-copy lines have the same \textit{Wolbachia} growth rate, which is higher than the \textit{Wolbachia} growth rate of the two-copy line. This growth rate, in turn, is higher than that of one-copy \textit{wMelPop} and \textit{wMelCS}\_b, which have the same \textit{Wolbachia} growth rate (Fig. 3C). We confirmed this effect of Octomom copy number on \textit{Wolbachia} densities by comparing \textit{Wolbachia} WSP protein abundance between flies harboring \textit{wMelPop} with three versus ten Octomom copies (Fig. 3D). Flies carrying \textit{Wolbachia} with ten Octomom copies had more WSP protein than flies harboring \textit{Wolbachia} with three copies.

The density of \textit{Wolbachia} is known to be related with \textit{Wolbachia}-conferred antiviral protection, and \textit{wMelPop} provides very strong protection [8,17,18,26,38–40]. This protective effect is best analyzed when flies are kept at 18°C, the temperature at which \textit{wMelPop} is not pathogenic [41]. In flies that are raised from egg to adult at 25°C, \textit{Wolbachia} levels at the time of infection are still related to Octomom copy number (see Fig. 3C). The survival of virus-infected flies confirmed that the higher the Octomom copy number, the stronger the antiviral protection (Figs. 3E and S5H). As with pathogenicity and growth rate, \textit{wMelPop} with one Octomom copy is phenotypically identical to \textit{wMelCS}\_b in terms of antiviral protection.

We showed that Octomom copy number can change rapidly under direct selection (Figs. 2 and S2). Next we questioned whether Octomom copy number would be stable if this selection were relaxed. We observed that releasing our lines from copy number selection and maintaining them at 25°C in crowded vials for five generations caused a decrease in copy number in three out of four lines tested (S8 Fig.). The only line where the copy number did not change over the five generations started with two Octomom copies. Also, examination of another \textit{wMelPop} stock did not show the expected life-shortening phenotype and, accordingly, Octomom amplification (S9 Fig.). Presumably, Octomom copy number reverted to one copy, and the phenotype was lost in this stock. All these results demonstrate that \textit{wMelPop} \textit{Wolbachia} is genetically and, consequently, phenotypically unstable.

Octomom amplification could promote \textit{wMelPop} virulence in several ways, including via local or overall gene expression deregulation. The most parsimonious explanation, however, is that Octomom genes are overexpressed and that this causes the phenotype. Thus, we checked the expression of Octomom genes, immediately adjacent genes, and genes distant from the region by reverse transcription real-time qPCR. All Octomom genes, except WD0514, had a statistically significant higher expression in \textit{wMelPop} than in \textit{wMelCS}\_b, but immediately adjacent genes did not (S10 Fig.). Moreover, analysis of one Octomom gene (WD0511) showed that expression level was dependent on \textit{wMelPop} Octomom copy number (Fig. 4).

**Discussion**

Here we identify the genetic basis of \textit{Wolbachia} \textit{wMelPop} virulence. By selecting for \textit{Wolbachia} with different Octomom copy numbers, we show a functional link between copy number and \textit{wMelPop} phenotypes. The more copies of Octomom, the higher the densities of \textit{Wolbachia}, and the faster the hosts die, but the stronger the antiviral protection. The evidence we provide is stronger than a simple correlation because we are controlling Octomom copy number and determining its effect. Furthermore, all \textit{wMelPop} phenotypes are reverted in the line selected for one Octomom copy, establishing that Octomom copy number drives these phenotypes. There is evidence that \textit{Wolbachia} levels determine the strength of the \textit{Wolbachia}-associated
phenotypes [8,17,18,26,38–40]. Therefore, the different replication capacities of wMelPop variants with distinct Octomom copy numbers are the likely cause of the differences in the other phenotypes.

Woolfit and colleagues also identified Octomom amplification in the D. melanogaster wMelPop genome and a deletion of the Octomom region in a mosquito-adapted wMelPop variant, wMelPop-PGYP [62]. As wMelPop-PGYP retained a strong life-shortening effect in A. aegypti, while an A. aegypti–adapted wMel variant was benign, the authors dismissed Octomom as responsible for the high virulence of wMelPop also in D. melanogaster. We argue that the difference between wMelPop-PGYP and wMel phenotypes in mosquitoes may be due to other genetic changes accumulated during their adaptation to a new host, some already described for wMelPop-PGYP [62]. This phenotypic difference may also exist because wMel and wMelPop belong to the two different monophyletic groups of Wolbachia from D. melanogaster: wMel group and wMelCS group [8,68–70]. wMelCS-like variants replicate faster than wMel-like variants and sometimes shorten the lifespan of their natural host [8], and this difference may be exacerbated in mosquitoes. Relatedly, some Wolbachia bacteria transinfected into a new host species induce new pathogenic phenotypes [18,71–73].

Amplification of Octomom is in agreement with common gene amplification by nonequal recombination in bacteria [64]: (i) Octomom is flanked by direct repeats (see [8,62]), (ii) it seems to amplify as a unit, since different Octomom genes are equally amplified in the same fly (Figs. 1B and S1A), (iii) we confirmed the predicted novel joint point (Figs. 1C and S3; S1 Text), and (iv) the amplification is unstable.

The degree of Octomom amplification, and the associated strength of the phenotypes, can rapidly change and is fully reversible. This shows that Wolbachia can evolve rapidly, and adds...
to the understanding of genome evolution of endosymbionts. Many endosymbionts have evolutionarily dynamic genomes [1,74]. Genomes of Wolbachia and other endosymbionts (including Hamiltonella defensa, Serratia symbiotica, Sarocladium oryzae) are rich in mobile elements, prophages or phage-derived regions, and other repetitive DNA sequences [1,59,74–79]. These DNA elements may mobilize, amplify, or reduce in numbers, leading to genomic changes, but they can also mediate recombination and other genomic rearrangements. Comparative genomics of some closely related endosymbionts show extensive genomic rearrangements [75,76,78,80–85]. The same repetitive DNA elements may serve as a basis for gene amplification, as observed for Octomom. Consequently, gene copy number variation may be a common feature in these endosymbionts and may promote fast but reversible evolutionary changes. Accordingly, gene amplifications in other wMel variants [8], other Wolbachia strains [80,81], and the whitefly endosymbiont Portiera [78] have previously been reported, although without any associated phenotypes.

Genotype–phenotype links are very rarely established in endosymbionts, as many of them cannot be cultured in vitro. Previous examples include a point mutation in Buchnera aphidicola that affects thermal tolerance provided to the aphid Acyrthosiphon pisum [86] and the loss of a prophage in Hamiltonella defensa, abrogating induced protection to parasitoids in the same aphid [87]. The involvement of Octomom genes in Wolbachia virulence provides a unique point of entry into understanding Wolbachia–host interactions at the molecular level. As Octomom genes are overexpressed and may cause the phenotype, functional analysis of Octomom-encoded proteins is required to better understand the Wolbachia–host interaction. The Octomom region is part of the Wolbachia accessory genome since it is not present in all Wolbachia strains and shows signs of horizontal gene transfer [8,88–91]. There are genes putatively encoding mobile elements in the flanking region (WD0506 and WD0515, in the direct repeats) and inside Octomom (WD0511). Because of its structure and associated phenotype, the Octomom region resembles bacterial pathogenicity islands [92,93]. However, the pathogenicity seems to be expressed only when the region is amplified. The functions of Octomom genes are unknown and can only be speculated about based on the sequence of predicted proteins. Proteins encoded by three genes (WD0512, WD0513, and WD0514) have eukaryotic protein domains or homologs in arthropods (mosquitoes and Daphnia) and therefore may be effector proteins that interact with the host [8,88–91]. When highly expressed, these proteins could suppress host control over the symbiont. Other genes (WD0506–WD0511 and WD0515) encode proteins that may be involved in transposition, DNA replication and repair, or transcriptional regulation [8]. Overexpression of these proteins may increase Wolbachia’s replication rate. It is crucial to determine which of these genes are involved in the regulation of Wolbachia density and which structural characteristics of the Octomom region are important.

Octomom copy number instability may confound past and future analyses of wMelPop phenotypes. For instance, Octomom copy number variation may have contributed to changes in wMelPop pathogenicity over time or associated with different host species or host genetic backgrounds [41,42,94]. This instability has to be taken into consideration in future applications of wMelPop-transinfected mosquitoes to prevent transmission of human pathogens. In the dengue vector A. aegypti transinfected with wMelPop-PGYP, currently being tested in the field [44,95], Octomom copy number instability is not a factor since this region is deleted [62]. However, wMelPop is also being transinfected to other vectors of human diseases, such as the malaria-transmitting Anopheles gambiae [96] and the dengue and chikungunya vector Aedes albopictus [97].

We have shown variation in Wolbachia Octomom copy number between individual hosts within a population and across time. Genetic heterogeneity within individual hosts has been previously shown at the nucleotide level in wCer1 and wCer2 [98]. The instability of Octomom
copy number suggests that there is also a high level of heterogeneity between Wolbachia bacteria within individual insects. Analysis of the dynamics and consequences of heterogeneity in gene copy numbers in somatic or germline tissues may be important to understand host–endosymbiont interactions.

Vertically transmitted endosymbionts are subjected to different levels of selection. An increase in replication may confer a fitness advantage to the bacteria in intra-host competition but a disadvantage at the inter-host level, as it can have a high cost to the host and reduce symbiont transmission. A Drosophila line harboring wMelPop was most probably isolated in the laboratory because husbandry conditions buffered the cost to flies of pathogenic bacteria and because low host population numbers increased drift. Our results demonstrate that a single mutation (a duplication) can profoundly alter endosymbiont replication. This conversion of a mutualist into a pathogen by a single genomic event suggests that virulent mutations in microbial symbionts may be frequent and constantly counter-selected. Therefore, symbiont titers may be at a labile equilibrium achieved in the course of co-evolution and to a large extent selected at the level of the symbiont.

**Materials and Methods**

**Fly Strains**

*D. melanogaster* w1118 stock with Wolbachia wMelPop was kindly provided by Markus Riegler and Scott O’Neill. wMelPop OPL stock was kindly provided by William Sullivan and Laura Serbus. Both wMelPop stocks are derived from Min and Benzer original stock [19]. DrosDel isogenic background (iso) flies with no Wolbachia and with wMelCS_b or wMelPop were described before [8,24,99]. The wMelPop and mitochondria of this DrosDel isogenic background line derive from the w1118 stock [8].

**DNA Extractions**

DNA was extracted from individual flies (wMelPop) or pools of ten flies (wMelCS_b controls in the selection experiments). Each fly or pool of flies was squashed in 250 μl of 0.1 M Tris HCl, 0.1 M EDTA, and 1% SDS (pH 9.0) and incubated 30 min at 70°C. Next, 35 μl of 8 M CH3CO2K was added, and samples were mixed by shaking and incubated for 30 min on ice. Samples were then centrifuged for 15 min at 13,000 rpm at 4°C, and the supernatant was diluted 100× for qPCR.

**RNA Extractions and cDNA Synthesis**

For each sample, ten 3- to 6-d-old flies were pooled and homogenized with a plastic pestle in 1 ml of Trizol Reagent (Invitrogen). RNA was extracted according to manufacturer’s protocol and resuspended in 50 μl of DEPC-treated water (Ambion). RNA concentrations were determined using a NanoDrop ND-1000 Spectrophotometer. cDNA was prepared from 1 μg of total DNAse-treated RNA using Random Primers and M-MLV Reverse Transcriptase (all Promega). Primers were pre-incubated with template RNA for 5 min at 70°C. Next, the enzyme was added, and reactions were placed at 25°C for 10 min, 37°C for 60 min, and 80°C for 10 min. cDNA was diluted 100× for qPCR.

**Real-Time Quantitative PCR**

The real-time qPCR reactions were carried out in the CFX384 Real-Time PCR Detection System (Bio-Rad) as described before [8]. Briefly, each of the reactions was performed with 6 μl of iQ SYBR Green Supermix (Bio-Rad), 0.5 μl of each primer (3.6 mM), and 5 μl of diluted DNA.
We performed at least two technical replicates per biological sample for each set of primers. Primer sequences were described before [8]. The following thermal cycling protocol was applied: initial 2 min at 50°C, denaturation for 10 min at 95°C, followed by 40 cycles of 30 s at 95°C, 1 min at 59°C, and 30 s at 72°C. Melting curves were examined to confirm the specificity of amplified products. Ct values were obtained using Bio-Rad CFX Manager software with default threshold settings. Ct values were subjected to a quality check—samples with standard deviation between technical replicates exceeding one were discarded. Relative amounts of transcripts and genes were calculated by the Pfaffl method [100]. To apply the method, the efficiency of each of the primer pairs was predetermined in a separate experiment. For the Octomom expression data, values were normalized to gmk expression. For the determination of the number of genomic Octomom copies, values were normalized to the single-copy wsp gene. For Wolbachia quantification, wsp levels were normalized to Drosophila Rpl32.

Sequencing of the WD0514–WD0507 Junction
The WD0514–WD0507 junction was amplified using specific primers (Link_seq_1 and Link_seq_2), and Sanger sequencing was performed with these primers and the primers annealing inside the junction (Link_seq_3–7) by Source Bioscience. Primer sequences are listed in S3 Table.

Selection Experiments
Selection for high- and low-copy Octomom wMelPop lines in w1118 and iso backgrounds was initiated with females from a single vial of each background. For each background, ten single females were separated into individual vials and allowed to lay eggs for 5 d before being sacrificed for determination of Wolbachia WD0513 copy number. The offspring of the female with the highest and the lowest Octomom copy number was used to start the next generation. This general procedure was repeated at every generation of selection. Three replicates of high- and low-copy Octomom selection lines for each background were established at generation two. From that point on, we selected one female/line/generation with the desired Octomom copy number (based on real-time qPCR). Female age for egg laying (0–2 d) and qPCR (5–7 d) was controlled from generation four and two for the iso and w1118 lines, respectively. At generation seven of the w1118 lines, we started to also select for one-copy Octomom wMelPop. At this point we selected the female with Wolbachia with WD0513 copy number closest to one for this selection regime, and the female with Wolbachia with WD0513 copy number closest to two for the low-copy Octomom lines.

From generation two to generation 13 of the w1118 selection lines and from generation two to generation 22 of the iso selection lines, we were selecting from among six to ten females. From these generations on, we selected from three females per line.

At generation 14 of the w1118 lines and generation 18 of the iso lines, the selection was not performed.

Preparation of Flies for Phenotypic Analyses
For phenotypic analyses of flies carrying wMelPop with different Octomom copy numbers, single females were placed in vials, allowed to lay eggs for 5 d, and sacrificed to determine WD0513 copy number. The progeny of females carrying Wolbachia with the specified Octomom copy numbers were selected for the phenotypic analyses. All lifespan assays were performed at 25°C and 29°C, the temperature regimes applied in the first report on wMelPop phenotypes [19].

In order to directly compare flies with wMelPop with the full range of Octomom copy numbers, flies with wMelCS_b, and flies without Wolbachia, we used hybrids between w^{1118} and iso genetic backgrounds (S1 and S2 Tables). Females with the desired Wolbachia status, which is transmitted to the next generation, were crossed with males from the other genetic background. Since females were used in the phenotypic analyses, their genetic backgrounds were all equal and heterozygous between w^{1118} and iso, irrespective of the direction of the crosses. The mitochondria from these two lines should be identical since they share a very recent common ancestor [8]. We used females with high Octomom copy number from both genetic backgrounds to control for the possible influence of the direction of the cross and maternal effects potentially associated with different backgrounds.

**Lifespan and Wolbachia Levels Experiments**

Females whose mothers’ Octomom copy number was assessed by qPCR were collected at eclosion (ten per tube), allowed to mate for 24 h (five males per tube), separated from males, and either checked for survival at 25°C or 29°C every day or kept at 25°C and sacrificed at the indicated time points for Wolbachia density quantification. Females were maintained on a standard cornmeal diet without live yeast and were passed to fresh vials every 3 d. The mothers of females used for phenotypic analyses were derived from selection lines at the generations indicated in S2 Table.

**Virus Production and Infection**

DCV was produced and titrated as described before [8,24]. Infections were performed by prick- ing 1- to 2-d-old female flies with virus at 10^9 TCID_{50} (median tissue culture infectious dose)/ml. After infection, flies were kept in vials without live yeast, ten flies per vial, at 18°C. It has been shown previously that wMelPop is not pathogenic to the flies at this temperature [41]. Flies were checked for survival daily and passed to fresh vials every 5 d.

**Statistical Analysis**

Survival data were analyzed by Cox proportional hazard mixed effects models. Octomom copy number was considered a fixed effect, and replicate tube (containing ten flies) within the same experiment was considered random. Model fitting was done using the coxme package in R [101]. Tukey’s test was applied for pairwise comparisons of Cox hazard ratios between flies with all wMelPop lines, flies with wMelCS_b, and flies without Wolbachia.

Analysis of growth curves of wMelPop lines with different Octomom copy number was performed with log-linear model fits (lm in R). The slopes of different fitted regression lines were compared and corrected for multiple comparisons (Bonferroni correction).

Spearman correlation between Octomom copy number and median time to death was performed in R (cor.test).

Comparison of the expression of several Wolbachia genes between wMelCS_b and wMelPop (S10 Fig.) was done with the t-test in R (t.test) and was corrected for multiple comparisons with the Bonferroni correction.

Comparison of wsp and WD0511 gene expression between fly lines carrying different Wolbachia (Fig. 4) was performed with a log-linear model fit (lm in R), and the different lines were compared pairwise with a Tukey’s test.
Western Blot

Ten mated females from high- and low-copy iso selection lines, whose mothers were individually tested for Octomom copy number, were aged for 10 d before protein extraction. Flies without Wolbachia were used as a negative control. Anti-WSP rabbit polyclonal antibody was kindly provided by Kostas Bourtzis [102,103] and pre-absorbed in fixed Wolbachia-free D. melanogaster embryos. Anti-beta-tubulin mouse monoclonal E7 antibody was acquired from the Developmental Studies Hybridoma Bank [104].

Supporting Information

S1 Data. Relative WD0513 copy number in single females carrying wMelPop from different stocks (data for Fig. 1A).
(XLSX)
S2 Data. Relative WD0507 and WD0513 copy number from individual flies carrying wMelPop (data for Fig. 1B).
(XLS)
S3 Data. Relative WD0513 copy number in wMelPop in w1118 flies throughout selection (data for Fig. 2).
(XLSX)
S4 Data. Lifespan data for flies carrying wMelPop with different Octomom copy numbers (data for Fig. 3A).
(XLS)
S5 Data. Lifespan data for flies carrying wMelPop from forward and reverse selection (data for Fig. 3B).
(XLS)
S6 Data. Time-course analysis of relative levels of Wolbachia wMelPop with different Octomom copy numbers (data for Fig. 3C).
(XLS)
S7 Data. Survival data for DCV-infected flies carrying wMelPop with different Octomom copy numbers, wMelCS_b, or no Wolbachia (data for Fig. 3E).
(XLS)
S8 Data. Relative expression of wsp and WD0511 in flies carrying wMelPop with different Octomom copy numbers or wMelCS_b (data for Fig. 4).
(CSV)
S9 Data. Relative WD0507, WD0510, rpoD, and gmk copy numbers from individual flies carrying wMelPop (data for S1 Fig.).
(XLS)
S10 Data. Relative WD0513 copy number in wMelPop in iso flies throughout selection (data for S2 Fig.).
(XLSX)
S11 Data. Lifespan data for iso flies carrying wMelPop with different Octomom copy numbers (data for S5A Fig.).
(XLS)
S12 Data. Lifespan data for iso flies carrying wMelPop with different Octomom copy numbers (data for S5B Fig.).
(XLS)

S13 Data. Lifespan data for \textit{w}^{1118} flies carrying wMelPop with different Octomom copy numbers (data for S5C Fig.).
(XLS)

S14 Data. Lifespan data for \textit{w}^{1118} flies carrying wMelPop with different Octomom copy numbers (data for S5D Fig.).
(XLS)

S15 Data. Lifespan data for flies carrying wMelPop with different Octomom copy numbers, wMelCS\textsubscript{b}, or no Wolbachia (data for S5E Fig.).
(XLS)

S16 Data. Lifespan data for flies carrying wMelPop with different Octomom copy numbers, wMelCS\textsubscript{b}, or no Wolbachia (data for S5F Fig.).
(XLS)

S17 Data. Lifespan data for flies carrying wMelPop with different Octomom copy numbers, wMelCS\textsubscript{b}, or no Wolbachia (data for S5G Fig.).
(XLS)

S18 Data. Survival data of DCV-infected flies carrying wMelPop with different Octomom copy numbers, wMelCS\textsubscript{b}, or no Wolbachia (data for S5H Fig.).
(XLS)

S19 Data. Relative WD0513 copy number in iso flies carrying wMelPop throughout reverse selection (data for S6A Fig.).
(XLSX)

S20 Data. Lifespan data for flies carrying wMelPop from forward and reverse selection (data for S6B Fig.).
(XLS)

S21 Data. Lifespan data for flies carrying wMelPop from forward and reverse selection (data for S6C Fig.).
(XLS)

S22 Data. Lifespan data for flies carrying wMelPop from forward and reverse selection (data for S6D Fig.).
(XLS)

S23 Data. Median time to death and Octomom copy number in experiments shown in Figs. 3A and 5A–G (data for S7 Fig.).
(XLS)

S24 Data. Relative WD0513 copy number in flies carrying wMelPop in the absence of selection (data for S8 Fig.).
(XLSX)

S25 Data. Relative WD0513 copy number in single females carrying wMelPop from different stocks (data for S9A Fig.).
(XLSX)
S26 Data. Lifespan data for flies carrying wMelPop OPL, wMelCS_b, or no Wolbachia (data for S9B Fig.).

(XLS)

S27 Data. Relative expression of Octomom genes and other Wolbachia genes in flies carrying wMelCS_b or wMelPop (data for S10 Fig.).

(XLS)

S1 Fig. Different Octomom genes are amplified to the same extent in individual wMelPop flies. Octomom gene copy number variability relative to wsp between wMelPop iso flies. qPCR was performed on DNA from single females from iso line three (Fig. 1A) for WD0507, WD0510, and WD0513 (A) and cpoD and gmk (B). wMelCS_b flies were used for copy number normalization. Supporting data can be found in S9 Data.

(TIF)

S2 Fig. Selection for wMelPop with high and low Octomom copy number in iso flies. The bars for generation zero correspond to the data for iso line three from Fig. 1A. The female with the highest or lowest WD0513 copy number was always the founder of the next generation. After the first generation, three females with high and low copy number gave rise to three replicate lines that were maintained separately for the subsequent generations. The boxes extend from the 25th to 75th percentile, and the whiskers include all values. Dashed lines separate the generations. Gen, generation; Rep, replicate. Supporting data can be found in S10 Data.

(TIF)

S3 Fig. PCR of the predicted WD0514–WD0507 junction in flies harboring wMelPop with a single Octomom copy. wMelCS_b was used as a negative control, and wMelPop with two and ten Octomom copies were used as positive controls for the WD0514–WD0507 junction. Flies without Wolbachia (iso) were used as a negative control for wsp. Two samples of each Wolbachia variant were used.

(TIF)

S4 Fig. Alignment of the sequences containing the wMelPop unique SNP site from wMelCS_b and wMelPop selection lines with one, two and a high number of Octomom copies. CLUSTAL O (1.2.1) multiple sequence alignment [105–107] was used to align the sequences surrounding the wMelPop unique SNP at position 943,443 in the w1118 selection lines. Position 943,443 is highlighted in yellow.

(TIF)

S5 Fig. Octomom copy number determines wMelPop phenotypes. (A and B) One hundred iso females from high- and low-copy selection regimes were checked for survival at 25°C every day. Mixed effects Cox model fit, high versus low copy number for both replicates, \( p < 0.001 \). Supporting data can be found in S11 and S12 Data. (C and D) One hundred w1118 females from high- and low-copy selection regimes were checked for survival at 25°C (C) or 29°C (D) every day. Mixed effects Cox model fit, high versus low copy number at both temperatures, \( p < 0.001 \). Supporting data can be found in S13 and S14 Data. (E–G) Sixty–seventy females carrying wMelPop with different Octomom copy numbers were monitored daily for survival at 29°C (E) or at 25°C (F and G). Females are the progeny from crosses between iso and w1118 lines. Letters refer to groups with significantly different survival curves according to Tukey’s test of all pairwise comparisons of Cox hazard ratios. The experiment at 29°C is a replicate of the one presented in Fig. 3A. Supporting data can be found in S15–S17 Data. (H) One hundred females with different wMelPop Octomom copy numbers were pricked with DCV (10⁹ TCID₅₀/ml), and survival was followed daily. Females are the progeny from crosses between iso
and w^1118 lines. Letters refer to groups with significantly different survival curves according to Tukey’s test of all pairwise comparisons of Cox hazard ratios. This experiment is a replicate of the one shown in Fig. 3E. Supporting data can be found in S18 Data.

**S6 Fig. Phenotypic responses to reverse selection.** (A) At generation 17 of the selection for wMelPop iso lines with high and low WD0513 copy number (S2 Fig.), the selection was reversed. This reverse selection was performed in all three replicate lines from the high- and low-copy selection regimes by selecting the female with the highest WD0513 abundance from each low-copy line and the female with the lowest WD0513 abundance from each high-copy line (forward selection also continued, as shown in S2 Fig.). The boxes extend from the 25th to 75th percentile, and the whiskers include all values. Dashed lines separate the generations. Gen, generation; Rep, replicate. Supporting data can be found in S19 Data. (B and C) Lifespan of females of reversely selected high-copy lines was compared with that of high-copy females under forward selection at generation 22. Fifty females per line were used. (B) High-copy line one (nine Octomom copies) versus reverse high-copy line one (five copies) (C) High-copy line three (ten copies) versus reverse high-copy line three (six copies). Tukey’s test on the mixed effects Cox model fit, high versus low copy number, \( p < 0.001 \) and \( p = 0.0321 \) for lines one and three, respectively. Supporting data can be found in S20 and S21 Data. (D) Lifespan of females from forward selection low-copy line three (3.5 Octomom copies) and the corresponding reverse selection line (eight copies) at generation 22. Fifty females per line were used. Tukey’s test on the mixed effects Cox model fit, high versus low copy number, \( p < 0.001 \). Supporting data can be found in S22 Data.

**S7 Fig. Negative correlation between Octomom copy number and host longevity.** Median time to death (days) for lifespan experiments performed (Figs. 3A and S5A–G) is plotted as a function of Octomom copy number (relative WD0513 copy number). These data refer to flies with two different genetic backgrounds and experiments performed at two different temperatures. The two variables are negatively correlated (Spearman correlation rho = −0.701, \( p < 0.001 \)). Supporting data can be found in S23 Data.

**S8 Fig. Release of selection pressure leads to a change in Octomom copy number.** Selection was released in wMelPop iso flies at generation 26. The progeny of single females from generation 26 were kept without selection for Octomom copy number for five generations by passing all the flies to a new tube every 20 d. After these five generations, ten females per line were scored for WD0513 copy number in their Wolbachia bacteria. Plotted are the original selection lines at generation 26, the same selected lines at generation 31 (the high-copy-number line was selected for ten Octomom copies from generation 29 onwards), and released selection lines at generation 31. The mothers of selected lines are represented by triangular data points, the mothers of the released selection lines are represented by blue circular data points. Lines are medians of the points at each generation/treatment. Octomom copy number decreased in three out of four lines released from selection. The only line that did not show a decrease started with two copies of Octomom. Supporting data can be found in S24 Data.

**S9 Fig. Lack of Octomom amplification and virulent phenotype in a different wMelPop stock.** (A) Comparison of WD0513 copy number within different wMelPop iso and w^1118 stocks kept in the Teixeira lab (Fig. 1A) with wMelPop stock obtained from another lab (wMelPop OPL [original Popcorn line]). DNA from single females was extracted for qPCR.
wMelCS_b iso flies were used for copy number normalization, and wsp was used as a reference gene. Lines are medians of the replicates. Supporting data can be found in S25 Data. (B) Lifespan of females without Wolbachia, with wMelCS_b, and with wMelPop OPL. Females are the progeny from crosses between flies of the iso and the wMelPop OPL genetic backgrounds. One hundred females were collected at eclosion, allowed to mate for 24 h, separated from males, and scored daily for survival at 29°C. Letters refer to groups with significantly different survival curves according to Tukey’s test of all pairwise comparisons of Cox hazard ratios. Supporting data can be found in S26 Data.

S10 Fig. Octomom amplification leads to higher expression of Octomom genes. Expression of genes in the Octomom region (WD0507–WD0514), in the flanking repeated region (WD0506/WD0515), in the immediately adjacent region (WD0505 and WD0519), and in other locations of the chromosome (wsp and rpoD) in wMelCS_b (A) and wMelPop (B) (both in DrosDel isogenic background). The expression levels of WD0506–WD0513 are higher in wMelPop than in wMelCS_b (t-test, p < 0.001 for all). The expression levels of Octomom gene WD0514 and genes outside Octomom (wsp, rpoD, WD0505, and WD0519) are not significantly different between the two Wolbachia variants. Relative expression for each gene is calculated using gmk as a reference gene and is relative to that of wMelCS_b samples. RNA was extracted from eight samples of ten 3- to 6-d-old iso males, and real-time qPCR was performed on cDNA with specific primers. Lines are medians of the replicates. Cycle threshold values for the genes WD0507, WD0513, and WD0514 are high, indicating low gene expression levels for these genes. These cycle threshold values fall in a nonlinear section of the standard curve, making the quantification inaccurate. Moreover, cycle threshold values for some reactions were below the detection limit. Supporting data can be found in S27 Data.

S1 Table. Genetic background of females used in reciprocal crosses to generate w1118 × iso hybrids (Figs. 3A, 3C, 3E, 4, and S5E–H).

S2 Table. Selection generation number origin of mothers of the flies used for phenotypic analyses.

S3 Table. Oligonucleotide primers used for amplification and sequencing of the WD0514–WD0507 junction.

S1 Text. Sequence of the new WD0514–WD0507 junction. The sequencing of the PCR band (Fig. 1C) was performed with primers Link_seq_1–7 (S3 Table).

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**Author Contributions**
Conceived and designed the experiments: EC LT. Performed the experiments: EC LT. Analyzed the data: EC LT. Wrote the paper: EC LT.

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