

The genomic basis of Red Queen dynamics during rapid reciprocal host–pathogen coevolution

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Red Queen dynamics, involving coevolutionary interactions between species, are ubiquitous, shaping the evolution of diverse biological systems. To date, information on the underlying selection dynamics and the involved genome regions is mainly available for bacteria–phage systems or only one of the antagonists of a eukaryotic host–pathogen interaction. We add to our understanding of these important coevolutionary interactions using an experimental host–pathogen model, which includes the nematode *Caenorhabditis elegans* and its pathogen *Bacillus thuringiensis*. We combined experimental evolution with time-shift experiments, in which a focal host or pathogen is tested against a coevolved antagonist from the past, present, or future, followed by genomic analysis. We show that (i) coevolution occurs rapidly within few generations, (ii) temporal coadaptation at the phenotypic level is found in parallel across replicate populations, consistent with antagonistic frequency-dependent selection, (iii) genomic changes in the pathogen match the phenotypic pattern and include copy number variations of a toxin-encoding plasmid, and (iv) host genomic changes do not match the phenotypic pattern and likely involve selective responses at more than one locus. By exploring the dynamics of coevolution at the phenotypic and genomic level for both host and pathogen simultaneously, our findings demonstrate a more complex model of the Red Queen, consisting of distinct selective processes acting on the two antagonists during rapid and reciprocal coadaptation.

host–pathogen coevolution | Red Queen hypothesis | population genomics | copy number variation | *Caenorhabditis elegans*

More than 40 y ago, Van Valen (1) proposed the Red Queen hypothesis stating that evolutionary lineages persist only if they continuously change and adapt to ongoing selective pressures. A later refinement of the hypothesis put the spotlight on host–pathogen interactions (2, 3): Because these interactions are antagonistic and many pathogens depend on their hosts for survival, they are likely shaped by repeated cycles of adaptation and counteradaptation from both partners. The consequences of these dynamics are potentially far-reaching. For example, they may account for the extremely high rates of molecular evolution in host immunity and pathogen virulence genes (4, 5), they might favor the evolution and maintenance of outcrossing and sexual reproduction (6, 7), they might generally promote genetic diversity (3), and they can also affect a diversity of life history traits (8). Despite the enormous number of studies on host–pathogen coevolution, two core aspects are as yet only poorly understood: the genome regions targeted by reciprocal selection and the resulting allele frequency dynamics in these regions.

Two main dynamics are generally proposed as alternative mechanisms underlying host–pathogen coevolution: recurrent selective sweeps (RSS) and antagonistic frequency-dependent selection (aFDS) (2). Under RSS, the two antagonists sequentially accumulate adaptive mutations, each one sweeping to fixation through the host or pathogen population. Under aFDS, genetic diversity is maintained, and adaptive alleles oscillate in frequency

as a result of selection against the most common host or pathogen type. The relevance of these two selection mechanisms can be assessed through time-shift experiments, where fitness-related traits for one of the antagonists is evaluated in the presence of the other antagonist from the evolutionary past, present, and future (9, 10). The expectation for RSS is that the focal organism produces high fitness toward its past and low fitness toward its future antagonist. Under aFDS the faster-evolving partner (usually the pathogen) is expected to have higher fitness in the presence of the contemporaneous antagonist than those from the evolutionary past or future (11). To date, such time-shift experiments have primarily been applied to bacteria–phage models, revealing that coevolution leads to RSS dynamics initially (12, 13) and possibly aFDS dynamics at later time points depending on nutrient availability, spatial structure, and fluctuating environments (11, 14, 15). Time-shift experiments have occasionally been used for eukaryotic unicellular hosts, for example, unicellular algae and coevolving viruses, revealing transient RSS dynamics followed by the dominance of a generally resistant host (16). Time-shift experiments with multicellular hosts were, for example, performed for the water flea *Daphnia magna* and the flax *Linum marginale* host systems, in both cases suggesting aFDS as the main determinant of coevolution (17, 18).

Significance

Pathogens are omnipresent and by definition detrimental to their hosts. Pathogens thus exert high selection on their hosts, which, if adapting, can exert similar levels of selection on the pathogen, resulting in ongoing cycles of reciprocal adaptation between the antagonists. Such coevolutionary interactions have a central influence on the evolution of organisms. Surprisingly, we still know little about the exact selection dynamics and the genome regions involved. Our study uses a controlled experimental approach with an animal host to dissect coevolutionary selection. We find that distinct selective processes underlie rapid coadaptation in the two antagonists, including antagonistic frequency-dependent selection on toxin gene copy number in the pathogen, while the host response is likely influenced by changes in multiple genome regions.

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Conclusive information on the involved selection dynamics further requires a combination of the time-shift experiments with data on allele frequency changes, using, for example, whole-genome sequence analysis. Several previous studies applied these two approaches (see summary in *SI Appendix, Table S1*). However, many of these studies had a different focus and thus did not quantify allele frequency changes across time. For example, previous genome analysis of host–phage coevolution focused on sequencing phage genomes only at the end point of the experiment (4) or only for a single phage gene across several time points (14). In general, longitudinal whole-genome analysis of coevolving populations remains rare and usually includes only one antagonist (*SI Appendix, Table S1*). In fact, whole-genome sequence analysis for both antagonists has been performed only for a bacteria–phage interaction, revealing RSS, as predicted by time-shift experiments, and an influence of phage diversity on selective sweeps dynamics (19, 20).

The objectives of the current study were to enhance our understanding of the selection dynamics and the genomic basis of host–pathogen coevolution by using (i) a multicellular animal host system, which can be easily manipulated and which allows cryopreservation for later time-shift experiments, (ii) laboratory-based coevolution experiments with control over interaction type (i.e., coevolution or not), population size, generation time (the latter only for the host), and high replication, (iii) time-shift experiments for the two antagonists, using cryopreserved material from the evolution experiment for both host and pathogen, and (iv) whole-genome sequence analysis for both antagonists, taking advantage of available high-quality reference genomes (Fig. 1). In detail, we used the nematode host *Caenorhabditis elegans* and its pathogen *Bacillus thuringiensis* as a model (21, 22). We implemented methods for nematode cultivation and worm–pathogen purification (*Methods*) to allow performance of evolution experiments at high throughput. To initiate coevolution in our experiments, we used genetically diverse host populations (with 3,000 individuals each, including >370,000 segregating sites per population) and infected them with a single *B. thuringiensis* strain, MYBT18247 (*SI Appendix, Figs. S1–S3*). In the coevolution treatment, host and pathogen populations were continuously paired to enforce reciprocal adaptation, while the control treatments were identical but without exposure to the respective antagonists (*Methods*, 16 biological replicates and 23 transfers per treatment). After the evolution experiment, we used the regularly frozen evolved hosts and pathogens for measuring changes in fitness by their exposure to the respective ancestral antagonists. We then characterized the selection dynamics using reciprocal time-shift experiments (all replicate populations, three time points) in combination with population genomic analysis of both coevolved hosts and pathogens (six replicate populations and three time points for each antagonist and treatment).

Results and Discussion

Coevolution Caused a General Increase in Host and Pathogen Fitness.

For our phenotypic analysis of the experimentally evolved hosts and pathogens, we focused on the fitness traits that were targeted by selection in our experimental protocol: host egg number and pathogen cell number inside a host body at exactly the transfer time point (*Methods*). We found that in the presence of the ancestral pathogen, the coevolved hosts had significantly more eggs than the control and ancestor, both at the end of the evolution experiment (Fig. 2A and *SI Appendix, Table S2*) and across the 23 transfers (*SI Appendix, Fig. S4* and *Tables S3* and *S4*). The host populations from the control treatment did not differ significantly from the ancestor (Fig. 2A and *SI Appendix, Fig. S4* and *Table S2*). These changes occurred independently of changes in resistance (*SI Appendix, Fig. S5A* and *Tables S5* and *S6*), a trait not under direct selection in our experiment. To estimate fitness of *B. thuringiensis*, the evolved pathogen lines were competed against a labeled ancestral clone for one infection cycle within the host. Bacterial fitness here is mainly determined by replication inside the host because the experimental medium

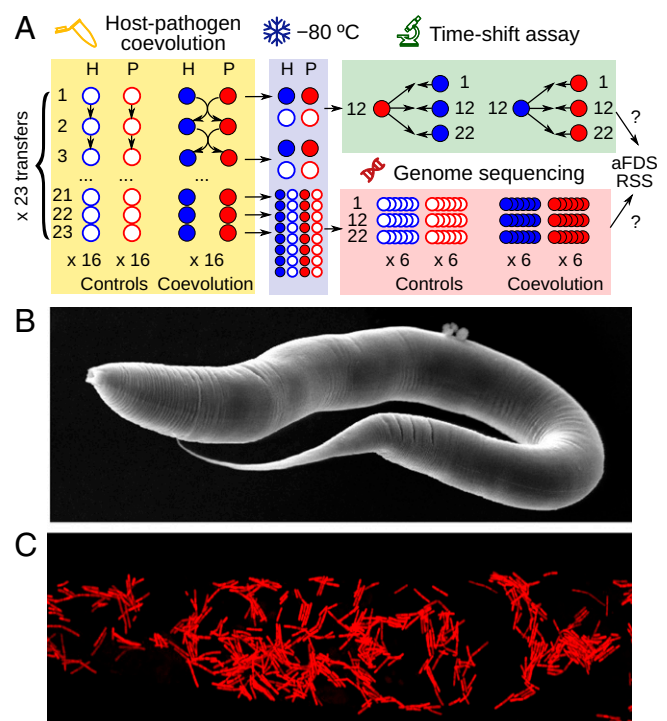


Fig. 1. A *Caenorhabditis elegans*–*Bacillus thuringiensis* model for experimental host–pathogen coevolution. (A) Overview of the general design of the evolution experiment, the subsequent phenotyping and time-shift experiment, and focus of the genomic analysis. H, host; P, pathogen. (B) Scanning electron micrograph of a *C. elegans* hermaphrodite. (C) Illustration of *B. thuringiensis*, labeled by red fluorescence, inside an infected and killed *C. elegans*.

did not support *B. thuringiensis* growth (*SI Appendix, Fig. S7*). In these competitions, coevolved pathogens produced significantly more cells than ancestral and control-evolved bacteria (Fig. 2B and *SI Appendix, Tables S7* and *S8*). The control lines did not differ from the ancestral clone (however, see relative fitness in broth, *SI Appendix, Fig. S7* and *Tables S9* and *S10*). Pathogen fitness evolved independent of virulence (*SI Appendix, Fig. S5B* and *C* and *Tables S11* and *S12*), which, like host resistance, was not directly targeted by selection in our experiments. We conclude that both host and pathogen adapted to the presence of their coevolving antagonist.

Time-Shift Experiments Indicated Antagonistic Frequency-Dependent Selection Dynamics for Both Antagonists.

We performed time-shift experiments to find out whether the antagonists adapted reciprocally to each other and, if so, whether adaptation was determined by RSS or aFDS. We ensured comparability of results for the two antagonists by assaying the same trait for both, namely, the effect on host egg production as one of the properties directly under coevolutionary selection. All 16 host populations from transfer 10 (H10, the focal time point) were infected with the corresponding pathogen populations from three different time points, including one from the past (P1, transfer 1), the present (P10, transfer 10), and the future (P23, transfer 23). In 10 out of 16 coevolved lines, pathogen P10 had the strongest negative effect on host fertility (Fig. 2C and *SI Appendix, Table S13*). This result is consistent with aFDS dynamics, where the pathogen is best adapted to the contemporaneous host. Similarly, when the pathogen from transfer 10 (P10) was used to infect the corresponding host populations from the past (H1), the present (H10), and the future (H23), the contemporaneous host H10 had the lowest fitness (Fig. 2D and *SI Appendix, Table S13*), again in line with aFDS.

The pattern of temporal genotype-specific coadaptation was additionally confirmed by considering all possible host–pathogen

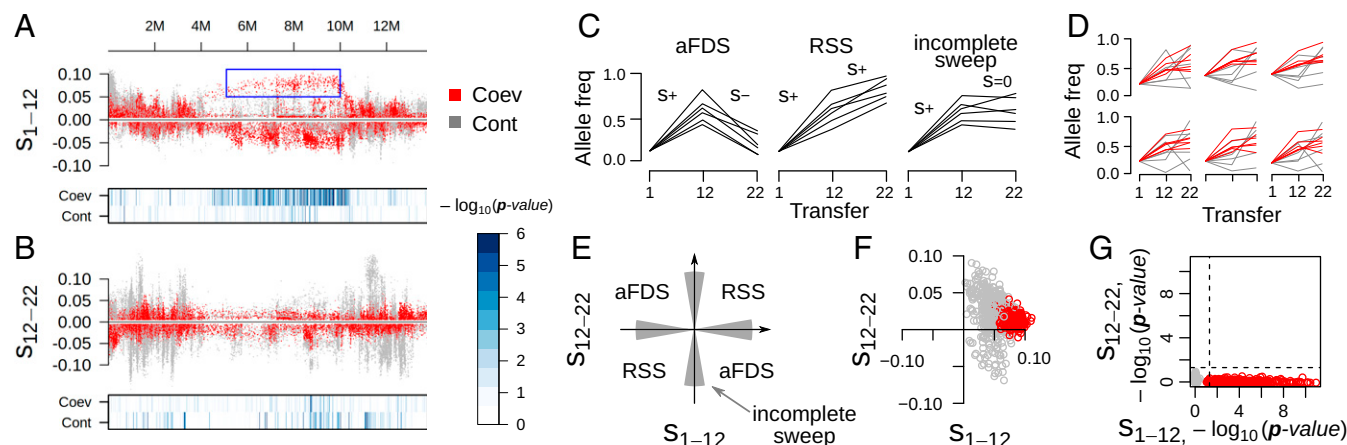


Fig. 3. Selective sweep dynamics of specific genome regions in coevolving *C. elegans* populations. (A) The distribution of the selection coefficient s along chromosome III during the first half of the experiment (transfers 1–12) estimated for coevolving (Coev; red) and control (Cont; gray) populations. The estimation of s is based on allele frequency changes of biallelic SNPs in six replicate populations. The heat maps below the main panel indicate genome regions with a significant change in allele frequency (more intense color corresponds to higher significance). The blue box highlights the genome region supported by many significant SNPs and high values of s (>0.05). (B) The estimated selection coefficients along chromosome III for the second half of the experiment (transfers 12–22). (C) Hypothetical allele trajectories expected for aFDS (Left), RSS (Middle), and incomplete sweeps (Right). (D) The actual allele trajectories for six representative SNPs from the highlighted region on A. (E) Hypothetical change in selection for three types of coevolutionary dynamics. If the SNPs observed are in complete linkage with the loci under selection, then a change in sign of the selection coefficient is expected for aFDS but not RSS dynamics. The shaded areas close to the axes correspond to incomplete sweeps. (F) The change in s for SNPs within the candidate region on chromosome III from A ($n = 489$). (G) The corresponding P values for the same SNPs, showing that allele frequency changes were significant from transfer 1 to 12 but not from transfer 12 to 22.

many loci are under selection but could result from high linkage due to the low recombination rates typical for the central regions of *C. elegans* chromosomes (26). These regions nevertheless include genes with functions related to the observed phenotypic variations, including fertility [e.g., a coevolution-specific duplication containing *trd-1* (27); *SI Appendix, Figs. S18 and S19*] and immunity-related genes like C-type lectin-like genes (ref. 28; for a complete list of change-of-function mutations see *Dataset S2*).

We conclude that none of the identified genome regions with significant changes under coevolution showed allele trajectories that match the expected patterns of the time-shift experiments. The lack of support for aFDS in the genomic data are unlikely due to a lack of statistical power. For example, many significant SNPs from the first half of coevolution could have lost significance during the second half due to increased population divergence (i.e., lack of parallel evolution). However, if we follow frequency changes of these particular SNPs in their replicate populations, then almost none of them changed the sign of selection, as required for aFDS. Consequently, the underlying genomic dynamics are likely more complex. It is conceivable that the phenotypic pattern is produced by nonparallel allele oscillations at different loci across the independently evolving replicate populations or, as a more parsimonious explanation consistent with the patterns identified in our genome data, by parallel-occurring incomplete sweeps, which occur at different loci during different phases of coevolutionary adaptation (Fig. 3C). Our findings are thus different from those of previous work, which, for example, suggested canonical allele oscillations at a single locus underlying aFDS in the *D. magna* model (29) or showed a series of complete allele fixations underlying RSS in (cyano)bacteria-phage experiments (12, 19).

Bacterial Coevolutionary Adaptation Is Driven by Copy Number Variation of a Toxin-Encoding Plasmid. In parallel to the host, we assessed genome sequence variation in six coevolved and six control *B. thuringiensis* populations. These populations were from the same time points and replicates as those of the host, analyzed above. For the pathogen, all replicate populations were founded by a single clone, and thus, adaptation should rely on only de novo mutations. As the phenotypic response was consistent across replicates from a specific evolution treatment, we again expected

to find patterns of parallel genomic evolution. Our analysis identified a total of 24 point mutations and small insertions and deletions (indels) during coevolution and 49 mutations in the control treatments (*SI Appendix, Tables S18–S20*). Eighteen mutations are nonsilent and are predicted to have functional consequences. Three mutations in the coevolving populations affected genes involved in sporulation initiation (*dnaA*, *kinB*, *spo0B*; *SI Appendix, Figs. S20 and S21*). For the control treatments, we also found mutations affecting sporulation, although in a different set of genes (e.g., *spo0B*, *abrB*, *kinE*, *ynzD*). These results suggest that changes in the process of sporulation are an adaptation to the general experimental evolution protocol. It cannot be excluded that the specific genes affected under coevolution contribute, at least to some extent, to the observed pattern of genotype-specific temporal coadaptation (*SI Appendix, Figs. S2D and S9*). Apart from the sporulation initiation genes, no other mutations showed parallelism in the coevolving populations.

As structural genome changes may also underlie evolutionary adaptation, we next sought the presence of copy number variations (Fig. 4). We found that a specific plasmid differed in copy number dynamics between control and coevolved populations (Fig. 4B and *SI Appendix, Table S21*). More precisely, a 113-kb plasmid carrying the *cry6B* toxin gene decreased in copy number across time in the control populations. In contrast, in the coevolving populations, the same plasmid first increased in copy number at transfer 12 and then returned to initial frequencies by transfer 22 (Fig. 4B and *SI Appendix, Table S21*). The encoded Cry6B belongs to the *cry6* family of pore-forming crystal toxins that cause *C. elegans* gut damage and mortality and reduced reproduction (30, 31). The general decrease of this plasmid in the control treatment is thus consistent with the observed reduced virulence (*SI Appendix, Fig. S5*), whereas oscillating copy number changes during coevolution indicates aFDS. Importantly, the change in *cry6B* copy number correlates significantly with the pathogen's effect on host fitness in the time-shift experiment (Fig. 4C, Spearman correlation $\rho_s = -0.58$, $P = 0.0053$). Thus, high copy numbers are associated with lowered host fertility, while low copy numbers lead to higher fertility in a host genotype-specific form during the time-shift experiments. This correlation thereby supports the involvement of copy number variation in the pathogen's temporal adaptation to the

coevolving host. Complete sequencing of the avirulent pathogen line C3_15 (Fig. 4D) revealed that the 113-kb plasmid was present at comparatively high frequency, yet it contained a 9.5-kb deletion encompassing the *cry6B* gene and five other ORFs (Fig. 4A), suggesting that the deletion caused the loss of virulence. Based on these results, we conclude that *cry6B* gene copy number variation largely determines the pathogen's response during coevolutionary adaptation.

Our findings point to the general importance of plasmid copy number variation as a mechanism underlying rapid adaptation of pathogenic bacteria. Here, we used an initially clonal pathogen and genetically diverse host. Such asymmetric levels of genetic diversity are rarely considered in coevolution experiments (SI Appendix, Table S1). However, under natural conditions, pathogens are often clonal and subjected to bottlenecks during the infection process and/or more generally during transmission, while the host population is usually genetically diverse. For the pathogen, this leads to reduced standing genetic variation and a limited supply of de novo mutation (24). Under such commonly

occurring conditions of low genetic diversity, it is unclear how pathogens acquire the relevant changes to quickly adapt to their host. Our findings now suggest that plasmid copy number variation could provide the means to respond fast, even when pathogens are initially clonal. Our results further demonstrate that copy number variations can cause aFDS-like dynamics during coevolution. Interestingly, our previous work already indicated the importance of copy number variation of a toxin-encoding plasmid in *B. thuringiensis* adapting to *C. elegans*, although for a genomically very distinct strain (MYBT18679) with a distinct set of Cry toxin genes and without any evidence for temporal coadaptation or aFDS dynamics (21). It is conceivable that such copy number changes also determine rapid adaptation in the large variety of other pathogens that similarly possess plasmid-encoded virulence factors (32).

Conclusions

Based on a controlled experimental approach, we demonstrated that rapid coevolutionary adaptations between *C. elegans* and its pathogen are consistent with aFDS at the phenotypic level. This finding highlights the potential importance of antagonistic frequency dependence for shaping coevolutionary dynamics in eukaryotic host systems. Most surprisingly, our genome sequencing results did not conform to the classic aFDS model with several alleles cycling at a single locus. Instead, we found that the phenotypic pattern of aFDS is determined by more complex changes at the genomic level and that these changes are clearly distinct in the two antagonists. In the case of *B. thuringiensis*, rapid adaptation of the initial clone was mediated by copy number variation of the *cry6B* toxin gene rather than oscillations of different alleles. Such copy number variations may generally enhance fast evolutionary responses (33) and may explain why many virulence factors are associated with mobile genetic elements (32). In the case of *C. elegans*, the phenotypic pattern of aFDS is likely produced by changes in different genome regions. These may occur independently across the replicate populations, possibly each still mirroring aFDS. Alternatively, they may occur in parallel, but for each genome region at different time periods of coevolution, thus yielding the identified pattern of multiple incomplete sweeps. Interestingly, the primary trait under selection in the experiment, host fertility, is known to have a quantitative genetic basis in this nematode, controlled by multiple loci of small effect (26). The quantitative nature of this trait may explain the involvement of several genome regions during coevolutionary adaptation. It is a clearly different response than that previously observed for aFDS dynamics in *Daphnia* water fleas, in which the interaction with the coevolving pathogen was found to be controlled by one or a few host loci (29, 34). In fact, to date, it is unclear what type of selection dynamics emerge if coevolution is mediated by changes in a quantitative trait (35) and/or by copy number variation. Our results highlight that both can simultaneously shape coevolution and cause rapid coadaptation of the two antagonists.

Methods

All methods are described in more detail in the SI Appendix. The evolution experiment was started with a previously established, genetically diverse *C. elegans* population (36) and the nematocidal *B. thuringiensis* strain MYBT18247 (21, 22, 37). Experimental evolution was performed in a viscous medium which allowed us to maintain *C. elegans* in microtiter plates in a homogeneous environment, where they can mate and reproduce at rates comparable to those on agar plates. All protocols for *C. elegans* maintenance and manipulation (e.g., synchronization, feeding, worm transfer, cryopreservation, infection protocol) were specifically optimized for this medium (SI Appendix, Material and Methods). The nematode populations were preadapted to the experimental evolution protocol over 22 wk. The evolution experiment was run over 23 wk with weekly transfers and included three treatments (SI Appendix, Fig. S3; 16 biological replicates per treatment): a coevolution treatment, in which evolving hosts and pathogens were forced to coadapt to each other, and two control treatments, in which either *C. elegans* or *B. thuringiensis* was subjected to the passaging in the absence of the antagonist. Host populations were maintained at a constant population size of 3,000 individuals, while *B. thuringiensis* was always added

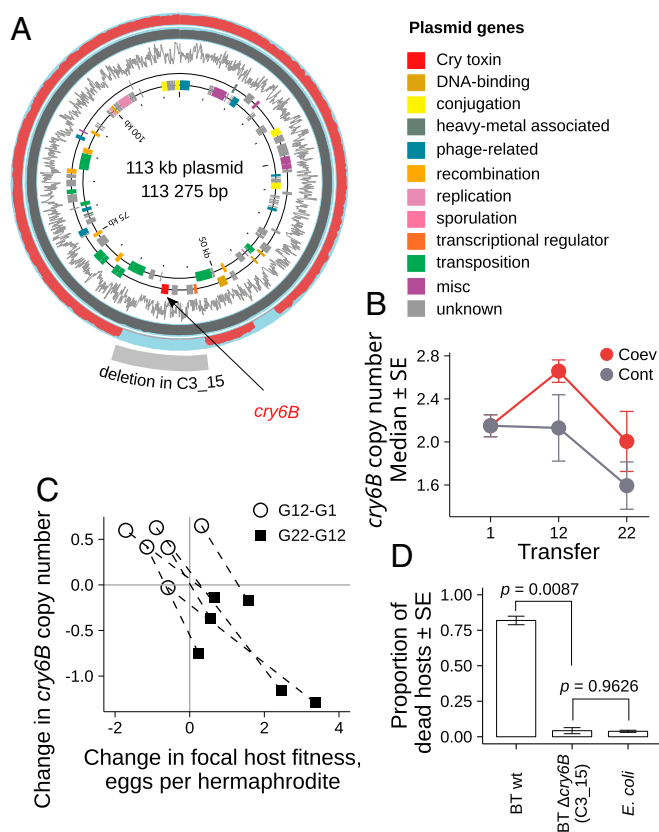


Fig. 4. Coevolving *Bacillus thuringiensis* show copy number changes in a toxin-encoding plasmid. (A) The 113-kb plasmid encoding Cry-toxin gene *cry6B*. Circles starting from the inside: coding DNA sequences on positive and negative strands, with colors indicating gene functions, GC content, sequence coverage for the virulent ancestor (gray on blue background), coverage for evolved avirulent pathogen line C3_15 (red on blue background), and deleted region in C3_15. Pathogen line C3_15 is a replicate population from the coevolution treatment that completely lost virulence after 23 transfers. (B) Change in *cry6B*-encoding plasmid copy number (relative to the chromosome) under coevolution and control conditions (six lines per treatment). Avirulent line C3_15 is not included. (C) The *cry6B* copy number differences covary with the effect of the pathogen on the fitness of hosts from transfer 12 in the time-shift experiment (see the results of the full time-shift experiment in SI Appendix, Fig. S8). Dashed lines connect two observations for the same lines. (D) Virulence of pathogen line C3_15 lacking *cry6B* relative to the ancestor (Wilcoxon rank sum test; *P* values adjusted with the Holm-Bonferroni method for performing three non-independent comparisons). BT wt, *B. thuringiensis* wild-type strain MYBT18247.

at a defined concentration of 2×10^8 spores/mL at each transfer step. Host and pathogen could be effectively separated by either a bleaching protocol (only survived by host eggs) or a pasteurization step (only survived by pathogen spores), allowing full experimental control over the (co)evolutionary process. Host and pathogen populations were cryopreserved at almost every second transfer. After the evolution experiment, the cryopreserved material was subjected to phenotypic analysis in the presence of either the ancestral antagonist or the respective coadapted antagonists from different time points. We characterized changes in the following traits: (i) host fertility (or the pathogen's effect on host fertility), (ii) pathogen competitive ability inside the host (using the red fluorescent evolved bacteria in combination with a green fluorescent ancestral bacterium), and (iii) host resistance (or, analogously, pathogen virulence).

Complete genome sequences were generated with the Illumina HiSeq2500 technology for entire host or pathogen populations at three time points across the evolution experiment (transfers 1, 12, and 22; always six biological replicates per treatment and time point). The paired-end sequence reads were processed using a custom pipeline, including mapping to reference genomes for *B. thuringiensis* (37) and *C. elegans* (version WS235) and calling SNPs and indels (SI Appendix, Material and Methods). Plasmid copy number estimation was based on read coverage of the plasmid relative to the average of the corresponding chromosome. The selection coefficient s was estimated for two time intervals (transfers 1–12 and 12–22) based on the

change in allele frequencies of biallelic SNPs using a tailored statistical modeling approach. This approach allowed us to simultaneously estimate s for a given time interval and the associated P value based on information from all replicate populations. Moreover, we evaluated genetic differentiation of the evolved populations from the ancestor with F_{ST} statistics using a sliding window approach as implemented in PoPoolation2 (38).

The data are supplied as Datasets S1 and S2. Sequence data are available from the National Center for Biotechnology Information (NCBI) BioProject PRJNA475030 (39).

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