

1 **Two sides to every coin: complementary introgression line populations in *Caenorhabditis***
2 ***elegans***

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22 gene expression, genetic interaction

23 Abstract

24 Quantitative genetics seeks to understand the role of allelic variation in trait differences.
 25 Introgression lines (ILs) contain a single genetic locus introgressed into another genetic
 26 background, and are one of the most powerful quantitative trait locus (QTL) mapping designs.
 27 However, albeit useful for QTL discovery, this homogenous background confounds genetic
 28 interactions. Here, we created a novel IL population, complementary to a previously created
 29 population, which enables identification of genetic interactions in ILs. The novel IL_{CB4856}
 30 panel was made by crossing divergent strains of the model nematode *Caenorhabditis elegans*
 31 (N2 and CB4856). The IL_{CB4856} panel comprises a population of 145 strains with sequencing
 32 confirmed N2 introgressions in a CB4856 background from which a core set of 87 strains
 33 covering the entire genome was selected. We present three experiments demonstrating the
 34 power of the complementary IL panels. First, we performed QTL mapping identifying new
 35 regions associated with lifespan. Second, the existence of opposite-effect loci regulating heat-
 36 stress survival is demonstrated. Third, by combining IL_{N2} and IL_{CB4856} strains, an interacting
 37 expression QTL was uncovered. In conclusion, the complementary IL panels are a unique and
 38 ready-to-use resource to identify, resolve, and refine complex trait architectures in *C. elegans*.

39 **Introduction**

40 In the last decade, many advances have been made in quantitative genetics using
 41 *Caenorhabditis elegans*, putting it at the forefront of the study of natural genetic variation (1).
 42 The most frequently used strains for exploring natural genetic variation in *C. elegans* are
 43 called N2 (Bristol) and CB4856 (Hawaii). These two strains differ in 176,543 single
 44 nucleotide variants 256,747 insertions-deletions(2-4). These polymorphisms also translate to
 45 differences in phenotypic traits, many of which were identified using genetic crosses (1).
 46 Many of these genes were identified using a combination of recombinant inbred lines (RILs)
 47 and introgression lines (ILs). These extensive discoveries illustrate how ILs are excellent tools
 48 for dissection of complex traits, but only a single genome-wide IL panel exists for *C. elegans*
 49 at this time.

50 Many quantitative genetics studies in *C. elegans* that focus on gene identification start
 51 with a two-step approach: first recombinant inbred lines (RILs) are used, followed by
 52 introgression lines (ILs) (5). RILs are lines that form a genetic mosaic of both parental
 53 genomes, whereas ILs consist of small segments of one parental strain introgressed into the
 54 genetic background of the other parental strain. In the RIL-IL approach, first a quantitative
 55 trait locus (QTL) is identified using RILs, followed by confirmation and fine mapping using
 56 ILs (for examples, see *e.g.*: (5-9)). Although this paradigm is used in many QTL studies
 57 where such populations can be established, different approaches do exist such as
 58 chromosome-substitution populations or whole-genome IL populations (10-14). In studies
 59 using whole-genome IL populations, it has generally been noted that ILs are more sensitive
 60 for small-effect QTL compared to RILs (11,15-19). The reason ILs are more sensitive lies in
 61 the reduction of genetic diversity which enables direct comparison of isolated genomic
 62 regions against the parent background; genetic variation is isolated to the introgression.

63 However, whole-genome ILs panels are generally less informative on the exact location of the
64 QTL, because they contain fewer genetic cross-overs than RIL panels.

65 One of the major challenges in quantitative genetics is the detection (or the inference)
66 of the contributions of multiple loci to trait differences. The existence of pervasive epistasis
67 for complex traits has been hinted upon in many studies (for a review see (20)). Still, it
68 remains prohibitively challenging to map and quantify genetically interacting loci (1,21-23).
69 Interestingly, ILs are potentially suitable for uncovering such genetic interactions. However,
70 in a single-background introgression line population, the locus-background interactions are by
71 definition entangled. Therefore, the current genetic panels present limited options to detect
72 and verify genetic interactions. Complementary IL populations (containing introgressions
73 from parent A in genetic background B and vice versa) would be of great value here, because
74 these can detect loci-background interactions. In a combined, complementary IL population,
75 loci-background interactions can be mapped when adjacent ILs overlap and be distinguished
76 from closely linked additive effects or interactions. Thus, adding a population with the
77 complementary genetic background can further elucidate the genetic architecture of traits, as
78 has already been demonstrated for specific QTL (*e.g.* (9,24,25)).

79 Here, we constructed a new IL population with N2 segments introgressed into a
80 CB4856 genetic background (IL_{CB4856}). This population complements a previously
81 constructed IL_{N2} population and consists of 145 strains with sequencing-confirmed
82 introgressions and a core set of 87 strains covering the entire genome. This makes *C. elegans*
83 the first organism for which a set of complementary whole-genome IL panels is available. We
84 present power simulations for QTL detection and show experimentally that this population
85 can identify multiple QTL for lifespan (a notoriously complex trait). Furthermore, we
86 demonstrate that the combination of the IL_{CB4856} and IL_{N2} panels can uncover genetic
87 interactions with distant loci for heat-stress survival and the expression of the gene *clec-62*.

88 Together, our research provides a toolset for uncovering and mapping the existence of
89 complex genetic architectures in *C. elegans*.

90 **Material and methods**

91

92 *Strains and maintenance*

93 The starting strains for introgression line construction were recombinant inbred lines (RILs)
94 with N2 and CB4856 parents, namely: WN001, WN007, WN025, WN068, WN071, and
95 WN110 (26). Strains were kept on NGM plates seeded with *Escherichia coli* OP50 and
96 culturing temperatures used during the crosses were 12°C, 16°C, or 20°C, depending on the
97 desired speed of population growth (27).

98

99 *Crossing scheme*

100 To generate the IL_{CB4856}, we divided crosses into two stages (**Supplementary table 1**). The
101 first stage was used for most loci, where a RIL male was back-crossed to a CB4856
102 hermaphrodite to ensure the presence of CB4856 mitochondria in the F1. This step was
103 followed by a second stage cross with CB4856 males to enable the integration of homozygous
104 CB4856 genotypes at the *peel-1 zeel-1* incompatibility locus on chromosome I in the F2 (28).
105 The exception were strains with WN001 as a parent, where we wanted to obtain coverage of
106 the *peel-1 zeel-1* locus. For WN001 crosses, the second cross was initially omitted.
107 Subsequently, selected genotyping was conducted in the F3 (4-9 markers; **Supplementary**
108 **table 2**), screening for strains with most CB4856 loci and absence of the N2 genotype at the
109 *peel-1 zeel-1* locus.

110 Next, selected strains were inbred and selected further to obtain as many homozygous
111 CB4856 loci as possible (for 5-12 generations). If a desired strain contained more than one N2
112 locus, further back-crosses with CB4856 males were conducted until only one detectable N2
113 locus remained in an otherwise CB4856 genotypic background. Finally, the strains were
114 inbred by transferring single hermaphrodites to new NGM plates for at least 10 generations.

In this way, 154 ILs were created initially. These ILs have been cryopreserved along with the parental strains. Ultimately, after genotyping by low-coverage whole-genome sequencing we could verify the introgressions in 145 IL_{CB4856} strains and selected a core set of 87 unique introgression lines covering the entire genome (**Supplementary table 3**).

Genotyping by fragment length polymorphisms

Initial genotyping (during the crossing of the strains) was PCR-based using primer pairs that detect insertions-deletion variants between the CB4856 and N2 strains (4). In total, 41 primer pairs were optimized with a bias for covering loci with a high-recombination frequency (**Supplementary table 2**) (29). The selection criteria for generating the primer pairs were: (i) the deletion occurred in the CB4856 strain, (ii) the deletion is larger than 25 bp and shorter than 150 bp, and (iii) it is not located in a repetitive region. All primers have been developed with Primer3 (primer3-win-bin-2.3.6) on the 1000 bp up- and downstream of the deletion (30). Primer3 was used with standard settings, selecting three primers in the size ranges of: 100-150 bp, 200-250 bp, 300-350 bp, 400-450 bp, 500-550 bp, 600-650 bp, 700-750 bp, and 800-850 bp. The annealing temperature was selected between 58°C and 60°C. The specificities of the primers were tested using BLAST (ncbi-blast 2.2.28 win64) against WS230 (settings: blastn -word_size 7 -reward 1 -penalty -3) (31). Only primers with fewer than five hits were considered for further selection. Final selection of the primers was based on the presence of a visible amplicon of the expected size after PCR of the N2 and CB4856 strains (**Supplementary figure 1**) and detection of heterozygous strains.

For genotyping during crossing, DNA was isolated from single adults that had generated offspring. Nematodes were lysed at 65°C for 30 minutes using a custom lysis buffer (32), followed by 5 minutes at 99°C. Genotyping PCRs were performed with GoTaq using the manufacturers recommendations (Promega, Catalogue No. M3008; Madison, USA). The

annealing temperature was 58°C (30 seconds), with an extension time of 1 minute for 40 cycles. All samples were run on 1.5% agarose gels stained with ethidium bromide. Ultimately, a low-resolution genetic map was constructed for 154 ILs based on the insertion-deletion markers.

Sequencing: DNA isolation, library construction, and sequencing

DNA was isolated from all the 154 initially generated IL_{CB4856}. Furthermore, DNA was isolated from the six parental N2xCB4856 RILs: WN001, WN007, WN025, WN068, WN071, and WN110. We also sequenced an additional 29 IL_{N2} that were generated previously (14): WN203, WN204, WN206, WN208, WN210, WN213, WN214, WN218, WN222, WN224, WN231, WN233, WN236, WN237, WN238, WN240, WN247, WN249, WN253, WN256, WN259, WN261, WN262, WN265, WN269, WN272, WN282, WN283, and WN285. Furthermore, we sequenced the two parental strains N2 and CB4856 as reference. A subset of 15 of the IL_{CB4856} population – covering parts of chromosomes I and IV – has been published previously (**Supplementary table 3**) (33,34).

The DNA isolation and library construction have been reported on previously (33). Shortly, genomic DNA was obtained from populations grown on 9 cm NGM plates. The Qiagen DNeasy Blood & Tissue Kit (Catalogue No. 69506; Hilden, Germany) was used for DNA isolation. DNA was quantified by Qubit. For each sample, a total of 0.75 ng of DNA was taken as input for the library construction. The libraries were sequenced on an Illumina MiSeq using a 300 cycle kit. Data have been deposited under accession number SRP154243 in the NCBI Sequence Read Archive (SRA; <https://www.ncbi.nlm.nih.gov/sra>).

Genotype calling using hidden Markov model and construction of the genetic map

The low-coverage sequencing data were used for variant calling using a hidden Markov model, as described previously (35,36). Subsequently, the hidden Markov model genotype calls were filtered for low-coverage areas (<100 supporting calls) and for introgressions less than 1×10^5 base pairs, which are unlikely to occur often given the crossing scheme. From the 154 initially created ILs, we could confirm the presence of an introgression in 145 ILs and a core set of 87 ILs was selected. The core set covered as many loci as possible with introgressions with as few multi-introgression ILs as possible. Ultimately, the genetic map of the new population and the additionally sequenced strains were integrated with the previously constructed and sequenced RILs (4,26) IL_{N2} strains (4,14) from the Laboratory of Nematology, Wageningen University. This genetic map with 1152 markers covers the recombination events in the whole set of strains (**Supplementary table 4**).

Lifespan experiment

We measured lifespan of the 87 IL_{CB4856} strains in the core set. Each experiment was started by transferring 10 L4 animals to three independent plates per genotype (30 animals per strain in total). The two parental strains were tested on nine plates (90 animals in total). Plates were screened daily until reproduction ended. Animals that bagged and those that were lost from the assay (i.e. that climbed to the sides of the plate) were censored from the data. For the IL_{CB4856} strains WN322, WN342, and WN351, all three replicates were not successful. For QTL mapping, we summarized the data to the mean, median, minimum, maximum, and variance in lifespan. These summarized traits were used in further analyses (**Supplementary table 5**).

Heat-shock survival experiment

We analysed previously published data to identify heat-stress survival QTL in *C. elegans* (37). The goal of the analysis was to test whether loci on chromosome IV could be associated with heat-stress survival. The data from Jovic *et al.* consist of survival measurements in 33 CB4856 x N2 RILs and 71 IL_{N2} strains (**Supplementary table 7**). For each strain, 32 animals were measured together on one plate on average. These animals were heat-shocked for four hours at 35°C, 48 hours after age-synchronization of the population by bleaching (see (37)). We took the data for survival at 72, 96, and 216 hours for mapping QTL.

As the previously published dataset indicated chromosome IV might be involved in heat-stress survival, we designed a replicated experiment. We tested 17 IL_{N2}, 20 IL_{CB4856}, and the N2 and CB4856 parental strains for heat-shock survival (**Supplementary table 8**). Each experiment was started by transferring a food-deprived population to a new 9 cm NGM dish, where the population was allowed to develop for ~60h at 20°C. After that period, the population consisted of egg-laying adults, from which eggs were isolated by bleaching for developmental synchronization (day 0) (38). Isolated eggs were grown for 48 h at 20°C. At this time, 20-40 nematodes were transferred to 6 cm NGM plates containing 100 µl FUDR, which inhibits reproduction (39). Two plates per strains were generated, one as a control (remaining on 20°C) and one receiving a four hour 35°C heat-shock immediately after transfer (40). At 72h, 96h and day 216h of the experiment, the number of surviving and dead nematodes were counted. After gathering the data, we only included samples with greater than 10 animals found at the 72 hour observation (upon heat-shock, animals tend to crawl to the top of the plates where they desiccate). Each strain was assessed at least three times per treatment, parental lines were tested more frequently (n = 10 for N2 and n = 12 for CB4856, post filtering).

Experiment to measure gene expression of clec-62

We used the IL_{CB4856} population to test the QTL for the expression of *clec-62*, for which multiple QTL were uncovered in a previous study in the IL_{N2} panel (41). Expression of *clec-62* was measured in a similar setup as in two previous eQTL experiments (41,42). In short, we collected 48-hour old L4 nematodes of N2 (6 replicates), CB4856 (6 replicates) and a single replicate for 46 IL_{CB4856} strains, 50 IL_{N2} strains, and 52 N2xCB4856 RILs grown at 20°C (Supplementary table 9). The strains were tested in three experimental batches and each batch contained a randomized selection of strains and both parental lines in duplicate.

The RNA was isolated from these samples using a Maxwell16 LEV Plant RNA kit using the recommended protocol with one modification, namely adding 20 µL proteinase K to the lysis step after which the samples were incubated for 10 minutes at 65°C whilst shaking at 900 rpm (43). Some samples were removed from the analysis afterwards because of low RNA concentrations and/or RNA degradation (confirmed by gel electrophoresis). Subsequently, cDNA was constructed using a Promega GoScript reverse transcriptase kit and *clec-62* expression was measured using RT-qPCR. For the RT-qPCR, two primer pairs were designed for amplification of the two isoforms (A and B) of *clec-62*: P_MS_CLEC62_A_F (5' CGACACTTCATTCCCCGAGC 3'), P_MS_CLEC62_A_R (5' TTAAGCTGGAACGGCACCAAC 3'), P_MS_CLEC62_B_F (5' CGCGTTGGTGCCGCTTAAC 3') and P_MS_CLEC62_B_R (5' GATTGCTGATTGAGGACGGCG 3'). Gene expression of *clec-62* was normalized against the reference genes *rpl-6* and *Y37E3.8* as previously described by (43). After the experiment and quality control of the qPCR data, 5 N2 replicates, 6 CB4856 replicates, 41 IL_{CB4856}, 42 IL_{N2}, and 47 RIL samples were analysed further.

QTL mapping in the IL populations

We used three approaches for QTL mapping in the IL population(s): (i) to an individual IL, (ii) to a single genetic background using bin mapping (14), and (iii) to two backgrounds using bin mapping with an interaction term. The mapping to an individual IL uses a linear model where each IL is compared to the parent genetic background, correlating trait differences to the introgression region covered by the particular IL. To correlate trait values to the introgression, the linear model

$$y = I + e$$

where y is the trait and I is the presence of the introgression was used. Each IL was tested with this model versus the genetic-background parent (*e.g.*, for an IL_{CB4856} strain versus the CB4856 strain). The obtained p-values were corrected for multiple testing by using the p.adjust function in R with the Benjamini-Hochberg method (44).

The single-background bin mapping assumes that a single QTL exists per overlapping set of introgressions (14). Here, the ILs with an introgression at a specific marker were tested versus the genetic background strain. Similar to the individual introgression, a linear model

$$y = x_I + e$$

where y is the trait and x_I is the introgression genotype at a given marker is used. For each marker, only the ILs with an introgression are compared to the genetic background parent (*e.g.*, for an IL_{CB4856} strain versus the CB4856 strain). The significance threshold was calculated using a permutation approach with 1,000 permutations to determine whether a significance fell below a pre-set false-discovery rate ($q = 0.05$).

The two-background bin mapping also assumes that a single QTL exists per overlapping set of introgressions. However, because both parental backgrounds were considered, the interaction between the two backgrounds can also be solved

$$y = x_I + x_{BG} + x_I \times x_{BG} + e$$

Where y is the trait and x_I is the introgression genotype at a given marker, and x_{BG} is the genetic background of the IL. Also here, for each marker, only the ILs with an introgression are compared to the genetic background parent (*e.g.*, for an IL_{CB4856} strain versus the CB4856 strain). The significance threshold was calculated using a permutation approach with 1,000 permutations to determine whether a significance fell below a pre-set false-discovery rate ($q = 0.05$).

Using the outcome of the bin mapping models QTL peaks were called using a 1.5 LOD-drop and under the condition that two peaks should be at least 1×10^6 base pairs apart.

QTL mapping in the RIL population

Mapping in the RIL population was conducted as described previously (33,34). In short, we fitted the trait data to the linear model

$$y_i = x_{i,j} + e_{i,j}$$

where y is the trait as measured in RIL i , x is the marker of RIL i at location j , and e is the residual variance. The 1152 marker set was used for mapping, and the significance was determined by a permutation approach using 1,000 permutations ($q = 0.05$).

Power analysis in the IL populations

To determine the statistical power of bin mapping with the IL_{CB4856} and both IL panels, we used a simulation approach. QTL mapping was simulated for two scenarios: (i) using only the IL_{CB4856} panel (87 strains), or (ii) using the combined IL_{CB4856} and IL_{N2} panels. To test the power for QTL detection, simulated data was used as input for the analyses. First, we simulated scenarios of a single QTL, varying the amount of variance explained (0.2, 0.25, ..., 0.8) and various levels of replication in the RILs and ILs (2, 3, ..., 15). Second, we mapped the data using a bin mapping approach (as in (14)). Third, we bin mapped using both the IL_{N2}

and IL_{CB4856} populations enabling to detect additive QTL and local interacting QTL. QTL were mapped using the methods described above and compared to the simulated data.

Heritability analysis

The broad-sense heritability (H^2) was calculated as in (41,45,46), where

$$H^2 = \frac{\sigma_F^2 - \sigma_P^2}{\sigma_F^2}$$

here H^2 was the broad-sense heritability, σ^2 is the variance of either the population F (RIL, IL_{N2}, or IL_{CB4856}) or the parental strains P (N2 and CB4856). Where the variance of the parental populations was the pooled variance and used as an estimate of the measurement error. The significance threshold was determined using a permutation approach with 1,000 permutations per trait. A significance of $q = 0.01$ was taken to compensate for the upper-bound estimation this approach gives.

The narrow-sense heritability was calculated using a REML approach as provided by the “heritability” package (47-49). The significance threshold was determined by 1,000 permutations per trait.

Software, scripts, and data

Data were analysed using R (version 3.4.2, windows x64) in RStudio (version 1.1.383) with custom written scripts (50,51). The tidyverse packages (version 1.2.1) were used for organizing and plotting data (52). All scripts and data are available at https://git.wur.nl/published_papers/sterken_2022_cb4856-ils. In particular, the scripts include a set of functions for QTL mapping and power simulations in the IL populations.

Results

Genetic characteristics of the IL_{CB4856} population

A genome-wide population of introgression lines containing an N2 segment in a CB4856 background was constructed (IL_{CB4856}). This set was created by backcrossing a set of six recombinant inbred lines (26) with the CB4856 strain (**Supplementary table 1; Figure 1A**). During the crosses, the genotypes were monitored using 41 amplification fragment length polymorphism (FLP) markers (**Supplementary table 2; Supplementary figure 1**) and afterwards all 154 selected IL strains were whole-genome sequenced (**Supplementary table 3**). A total of 145 strains with sequence-confirmed N2 introgressions into the CB4856 strain was obtained, of which 99 contained a single N2 introgressed region in an otherwise CB4856 genetic background, 37 ILs contained two N2 introgressed regions, and nine strains contained multiple N2 introgressed regions (**Supplementary table 3**). From this set, a core set of 87 strains was assembled, covering the entire *C. elegans* genome with N2 segments in an otherwise CB4856 genetic background (**Figure 1B**). The median introgression size of this population was 3.13 Mb, with the smallest region spanning less than 0.02 Mb and the largest region spanning 16.09 Mb. Each chromosome was covered by regions found in a minimum of seven ILs (chromosome III) and a maximum of 20 ILs (chromosome IV). In conclusion, this set of strains can be used as resource for QTL exploration in *C. elegans*.

To facilitate integrated analyses using IL populations, we created and tested the power of a combined genetic map of the new population with our previously constructed N2-background IL population (IL_{N2}) (14). To complete the IL_{N2} genetic map, we sequenced 29 of the IL_{N2} strains, to supplement the 57 IL_{N2} strains that were sequenced previously (4). By integrating the IL_{CB4856}, IL_{N2} with the genetic map of the Wageningen N2xCB4856 RIL population (4,26), we constructed a map with 1152 informative markers, spanning 389 strains

(**Supplementary table 4**). We used this map for power analyses of the core sets of the two IL populations. With the establishment of an IL_{CB4856} population, we could simulate its ability to detect QTL when combined with the IL_{N2} population (**Supplementary figure 2; Figure 1C**). Using simulations, we found that three replicates are sufficient to detect 50% of the interacting QTL explaining at least 35% of variance. Therefore, this analytical design can effectively uncover local genetic interactions.

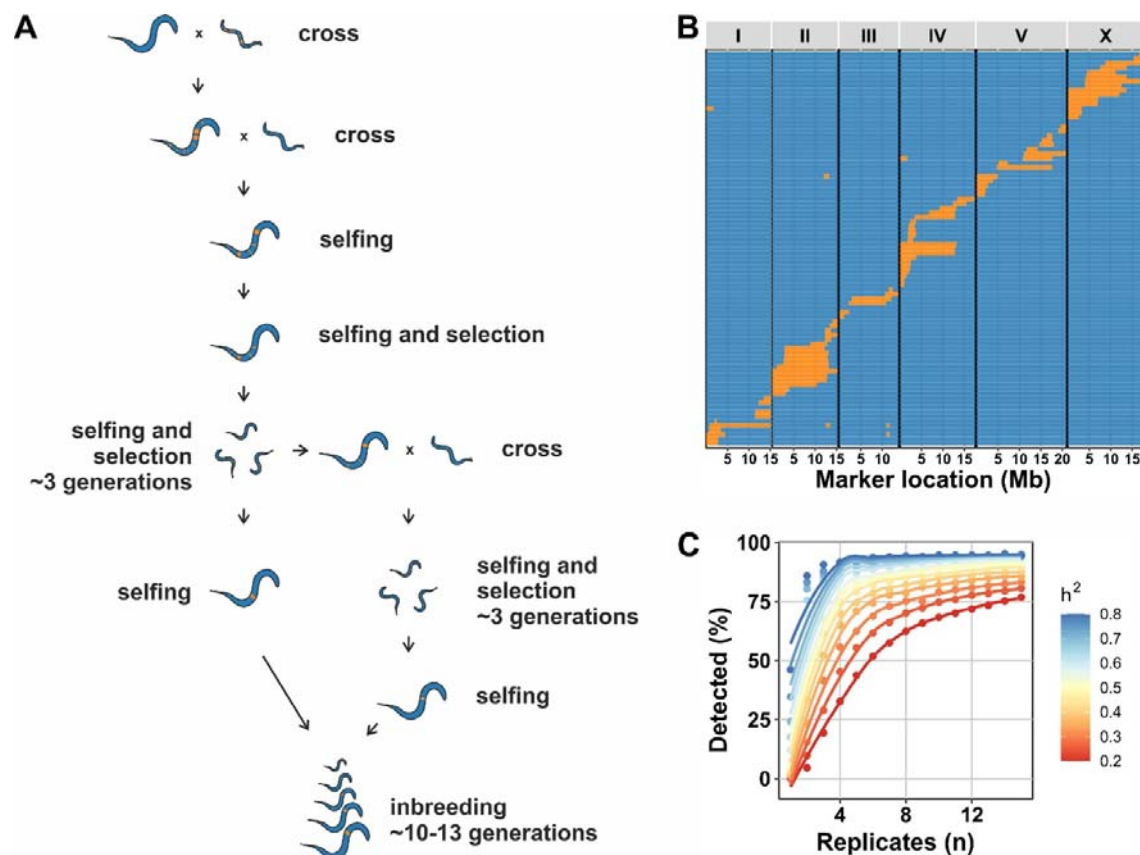


Figure 1: A novel N2>CB4856 introgression line panel and its mapping properties. (A) Schematic overview of the crossing scheme to create IL_{CB4856} strains. N2xCB4856 RIL strains were (back)crossed with CB4856 over several generations to obtain IL_{CB4856} strains with single introgressions. (B) The panel consisting of 87 strains covering the entire genome with N2 introgressions in a CB4856 genetic background. On the x-axis, the physical location is shown in million bases (Mb) split-out per chromosome. On the y-axis, the strains are shown (no label, each row represents a single strain). The blue colours indicate the CB4856 genotype, and the orange colours indicate the N2 genotype. (C) Power analysis of the

detection of genetic interactions when combining the IL_{CB4856} with the previously constructed IL_{N2} panel. On the x-axis, the number of replicates per IL is plotted against the percentage of the simulated QTL detected on the y-axis. The colours indicate the amount of variance explained (h^2) per simulated interaction QTL.

335

336 *QTL mapping in the IL_{CB4856} panel uncovered nine lifespan QTL*

337 To investigate QTL detection using real data in the newly generated IL_{CB4856} strains, we
 338 performed a QTL mapping experiment for lifespan (**Supplementary table 5**). We picked this
 339 trait because it was tested previously in the IL_{N2} population and would allow us to compare
 340 the QTL architectures (14). In the previous panel, six QTL for mean lifespan were detected.
 341 When we measured lifespan, we found that the N2 strain lived longer than the CB4856 strain
 342 on average, which was similar to previous results (**Figure 2A**) (14). After bin mapping, we
 343 detected 42 QTL for the various summary statistics mapped, including nine QTL related to
 344 mean lifespan (**Supplementary figure 3; Supplementary table 6; Figure 2B**). Of these nine
 345 mean-lifespan QTL, three (1, 2, and 7) were also detected in the IL_{N2} population, including a
 346 QTL on chromosome IV. Interestingly, the effect direction (shorter mean lifespan related to
 347 CB4856) was recapitulated in the IL_{CB4856} population (**Figure 2C**), indicating that we
 348 uncovered an additive QTL on the right side of chromosome IV.

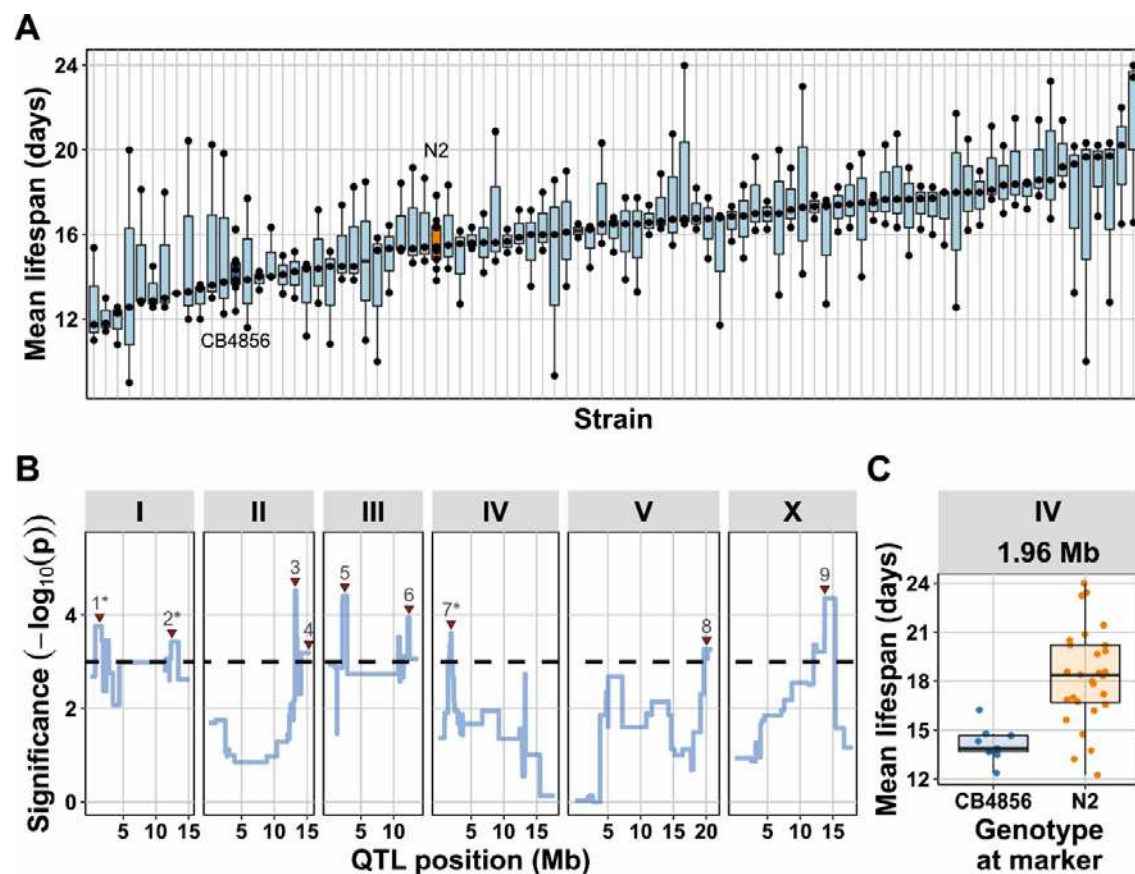


Figure 2: Lifespan analysis in the IL_{CB4856} panel. (A) Overview of the mean lifespan over all 87 IL_{CB4856} strains and the N2 and CB4856 parental lines. Each dot represents a plate (n = 10 animals per plate), the box plots are used as visual aids. Most of the ILs had longer lifespan than the two parental lines. (B) The QTL profile for the mean lifespan mapping. On the x-axis, the physical position in million bases (Mb) is shown. On the y-axis, the statistical significance of the association. The dashed horizontal line indicates the FDR = 0.05 threshold (based on 1,000 permutations). The red triangles indicate the peaks that were called, the asterisk indicates that this peak was mapped previously in the IL_{N2} panel (C) The trait values under the chromosome IV peak (7), where we found that the ILs with an introgression on the site (N2 genotype) had an increased lifespan compared to the CB4856 parental strain.

349

350 *QTL fine mapping using the IL_{CB4856} panel identified 18 heat-stress survival QTL on*
 351 *chromosome IV*

352 The most common usage of ILs is in fine mapping or experimental validation of previously
 353 identified QTL. Typically, once a QTL is detected in a RIL panel, a selected set of ILs can be
 354 used to validate a QTL detected in a mapping experiment (for examples, see *e.g.*: (5-9)). We
 355 used a set of ILs that could validate the detected QTL on chromosome IV in the heat-stress

response. Chromosome IV was previously implicated in fitness (reproduction) after heat shock, and a heat-shock expression QTL hotspot was found on chromosome IV (40-42). To further verify the QTL, we performed a QTL analysis on previously published data from a heat-shock survival experiment in the IL_{N2} panel (**Supplementary table 7**) (37). In this experiment, we found only one chromosome IV QTL associated with heat-stress survival (**Supplementary figure 4; Supplementary table 6**). We set out to investigate (i) whether this QTL was robust and (ii) whether it was additive or implicated in a genetic interaction with the genetic background.

We performed a heat-stress survival experiment on ILs that together covered chromosome IV: 17 IL_{N2} strains and 20 IL_{CB4856} strains. We measured survival of four hour exposure to 35°C at 24, 48, and 168 hours after the start of the exposure (**Supplementary table 8; Figure 3A**). We observed that the ILs typically showed a decreased survival when compared with the parental lines (**Figure 3B**). When we used these data for bin mapping, 18 heat-stress survival QTL were uncovered on chromosome IV (**Supplementary table 6; Figure 3C**). One locus, a QTL around 12.6 Mb was associated with a decrease in survival in both N2 and CB4856 ILs, indicating the existence of opposite-effect QTL in that region (**Figure 3D and E**). Therefore, we concluded that the use of strains from both IL panels is especially useful when mapping traits with complex genetic architectures.

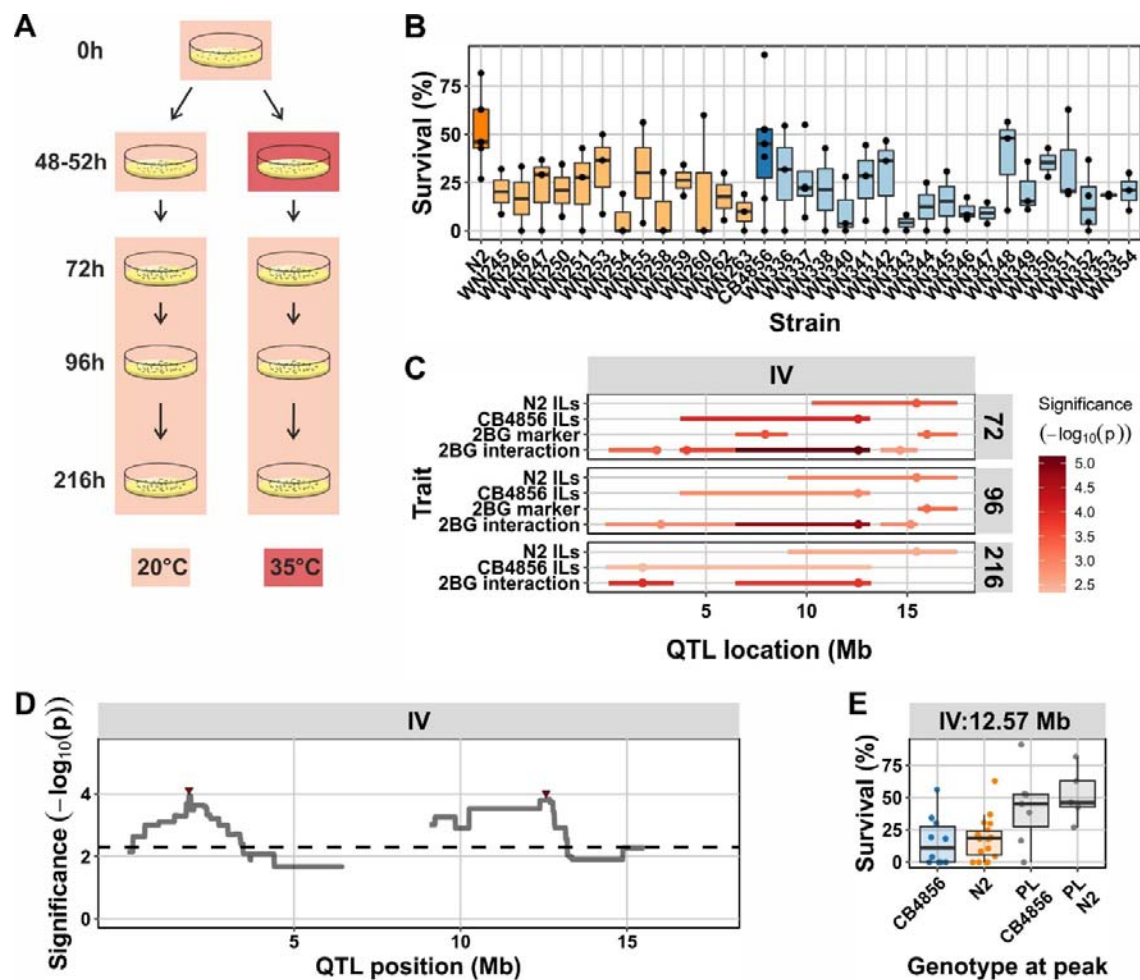


Figure 3: Heat-stress survival in ILs covering chromosome IV. **(A)** Experimental setup of the heat-stress survival experiment. **(B)** The survival one week after 4 hour exposure to a 35°C heat-shock. Each dot represents a plate (n = 20 – 40 animals per plate) and the box-plots are added as visual aid. **(C)** Overview of all QTL mapped in the IL populations using various types of models (N2 ILs: IL_{N2} bin mapping; CB4856 ILs: IL_{CB4856} bin mapping; 2BG marker: 2-background mapping model variance captured by marker; 2BG interaction: 2-background mapping model variance captured by the interaction marker-background). On the x-axis, the physical position in million bases (Mb) is shown. The colour scale indicates the significance of the association. Only significant associations are shown (FDR = 0.05; based on 1,000 permutations). **(D)** The QTL profile of the 2BG interaction term, the two red triangles indicate the locations of the peaks. On the x-axis, the physical position in million bases (Mb) is shown, on the y-axis the significance. The dashed horizontal line indicates the threshold (FDR = 0.05; based on 1,000 permutations). **(E)** The split-out of the right peak from panel (D). Where both parental strains have a similar survival percentage, the IL_{N2} (CB4856 introgression at the locus) and the IL_{CB4856} (N2 introgression at the locus) display a lower survival percentage.

Whole-genome QTL mapping uncovering an N2-background-dependent QTL for *clec-62* gene-expression

One final way in which the IL_{CB4856} panel can be used, is by combining data from different genetic panels, especially the IL_{N2} and N2xCB4856 RIL populations. For this case, we investigated an expression QTL for the gene *clec-62*, which we previously picked up in the IL_{N2} panel (41). Previously, no expression QTL in N2xCB4856 mapping populations had been identified (42,53-56). We set out to verify previous QTL detection in the IL_{N2} panel (41), as well as explore the existence of eQTL for *clec-62* in the IL_{CB4856} population.

We measured the expression of the two isoforms of *clec-62* in 47 RILs, 42 IL_{N2} strains, and 41 IL_{CB4856} strains by RT-qPCR (**Figure 4A; Supplementary table 9**) and found that *clec-62A* expression was 3.5-fold higher than *clec-62B*. The expression levels of the two isoforms were highly correlated (**Supplementary figure 5A**; Pearson $R = 0.94$; $p < 10^{-15}$) and therefore the analysis was performed using the higher expressed isoform A. We also found a high correlation between the expression in the previous microarray-based experiment on the IL_{N2} population, but not on the RIL population (**Supplementary figure 5B; Supplementary table 10**). Analysis of broad-sense and narrow-sense heritabilities in the three populations showed there was only significant broad-sense heritability for *clec-62* expression in the IL_{N2} panel (H^2 estimates 0.86 – 0.98; $q < 0.01$; **Supplementary table 11**). Significant narrow-sense heritability was only found in the microarray data of the IL_{N2} panel ($h^2 = 0.56$; $q < 0.01$), which is probably due to the higher number of strains included in that experiment. Altogether, we conclude that genetic variation in *clec-62* expression can only be found in an N2 genetic background and no QTL should exist in the RIL and the IL_{CB4856} population. Therefore, the variation in *clec-62* expression must be explained by genetic interactions.

QTL mapping of *clec-62* expression indeed identified QTL interacting with the genetic background. As predicted, no QTL were detected in the RIL and the IL_{CB4856}

populations (**Supplementary figure 5C**). However, when mapping using the IL_{N2} panel, a highly similar QTL profile for *clec-62* expression as measured by RT-qPCR and microarray was found (**Supplementary figure 5C; Supplementary table 6**). When using the combined IL_{N2} and IL_{CB4856} panels, we identified two loci with significant interaction effects: one on chromosome IV and one on chromosome X (**Supplementary table 6; Figure 4B**). These loci were both associated with a low expression when a CB4856 introgression is present (**Figure 4C and D**). Overall, we conclude that the interaction of CB4856 introgressions with the N2 genetic background drives the *clec-62* expression QTL.

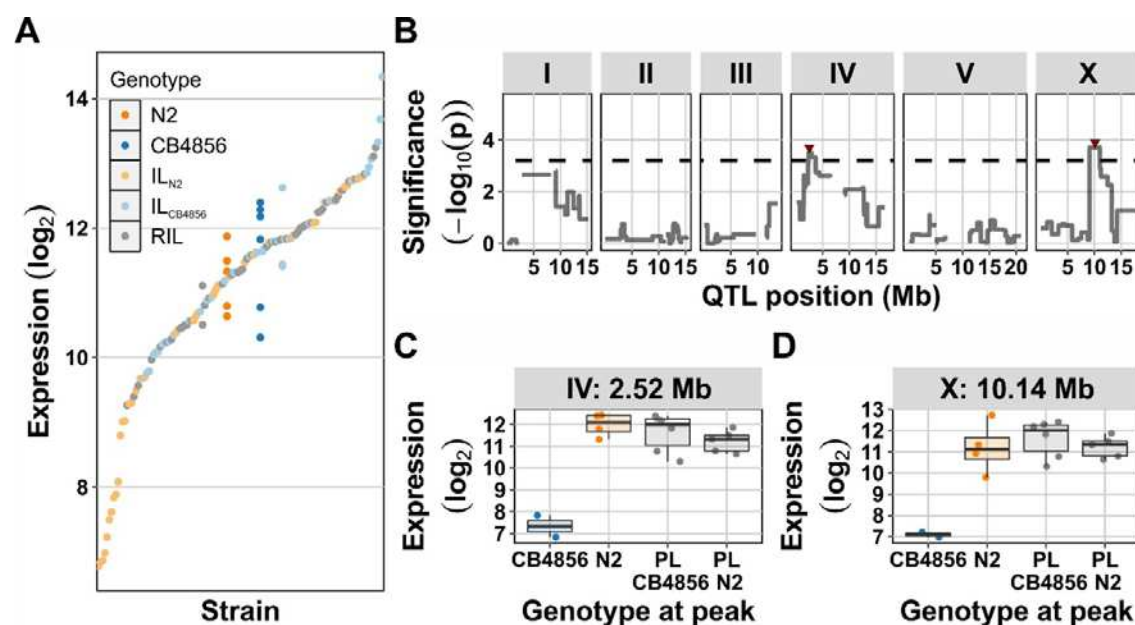


Figure 4: Expression of *clec-62* is due to QTL interacting with the N2 genetic background (A) The expression of *clec-62* as measured by qPCR in N2 (n = 5), CB4856 (n = 6), and three inbred panels: RIL (n = 47), IL_{N2} (n = 42), and IL_{CB4856} (n = 41). The log₂-normalized expression is shown per strain. (B) Profile of the significance of the interaction with the genetic background as mapped in the two IL panels. On the x-axis the physical position is shown in million bases (Mb). On the y-axis the significance of the association is shown (-log₁₀(p)). The dashed horizontal line indicates the FDR = 0.05 threshold. The red triangles indicate QTL-peaks. (C) The split-out of the chromosome IV peak from panel (B). Where both parental strains and the IL_{CB4856} population have a similar *clec-62* expression, the IL_{N2} (CB4856 introgression at the locus) has a lower *clec-62* expression. (D) As in (C), but for the

chromosome X peak.

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Discussion

Here we presented a novel whole-genome N2>CB4856 IL population in *C. elegans*. This new IL_{CB4856} population is complementary to the previous IL_{N2} population (14). The construction of the novel panel was facilitated by the availability of the first high-quality CB4856 genome (4), allowing for the selection of insertions/deletions as genetic markers which could be used during the crossing process. Genotyping by low-coverage whole-genome sequencing provided a detailed genetic map that allows the use of the novel population in various QTL approaches. We present three such approaches in this paper: (i) mapping lifespan using the panel on its own, (ii) further dissection of a previously implicated region in heat-stress resistance, and (iii) exploration of *clec-62* expression regulation using all three available N2xCB4856 populations. These cases show that the IL_{CB4856} panel empowers understanding the role of natural genetic variation in trait regulation, especially the role of genetic interactions.

The IL_{CB4856} panel for use in QTL mapping

The IL_{CB4856} panel is mostly free from large-effect laboratory-derived alleles that segregate in the IL_{N2} panel. The crossing scheme used to obtain the IL population considered results obtained from quantitative genetics studies in *C. elegans*. The scheme accounted for the *peel-1/zeel-1* locus (28) leading to marker distribution distortions in N2xCB4856 RIL populations (4,26,29). A double back-cross with CB4856 was used to remove the N2 *peel-1/zeel-1* allele, which failed to segregate otherwise (data not shown). As the background of the developed IL population is CB4856, it lacks the N2 laboratory derived alleles, such as *nath-10* (57), *glb-5* (9), and *npr-1* (9,25). Therefore, the new IL_{CB4856} panel may be especially useful if the studied trait might be affected by these pleiotropic and large-effect alleles (58).

One of the main strengths of IL populations in comparison to RIL populations lies in the detection of small phenotypic effects. RIL populations are hampered by the residual variance induced by the segregation of multiple QTL (11). In strains with a homogeneous background all QTL are fixed, except the introgressed locus, reducing the residual variance (the experimental variation) per QTL to a minimum. Many studies observed ILs resolve more QTL than RILs (11,15-19) and because the IL_{CB4856} population contains novel breakpoints compared to the IL_{N2} population QTL in these ILs can now also be pinpointed to a narrower locus. Therefore, the IL_{CB4856} population will likely be useful for fine mapping complex traits. Furthermore, the IL_{CB4856} population can serve as a resource for the generation of ILs with even smaller introgressions (19).

The compounding benefit of additional mapping populations

We show that complementary IL populations can easily identify genetic interaction dependent on background-effects. As examples, we identified interacting loci for both heat-stress resistance and *clec-62* expression. The reason that genetic interactions can more easily be identified when compared to RIL population (21,22) is that complementary IL populations can place the QTL effect in the context of the genetic background. There are many genome-wide IL populations, including for barley (12), *Arabidopsis* (11,13), tomato (59), maize (60), rice (61), and mice (62). However, to our knowledge there are no genome-wide complementary IL populations and until now observations in complementary ILs have only been made on a small scale. One difficulty of mapping interactions in complementary IL panels is that the partner locus (hiding in the genetic background) cannot readily be pinpointed. Still, QTL mapping in the combined IL_{N2} and IL_{CB4856} panels provides a starting point for investigations into genetic interactions.

From the experimental observations we made, we propose that genetic background effects are often overlooked in an IL panel with a single background. In such IL panels, the QTL-background interactions are confounded by definition. In other words: if an interaction with the genetic background exists and determines the trait levels in the individual introgression line, it cannot be distinguished from additive QTL. The homogenous background can lead to different estimations of QTL effect sizes compared to RILs (as reviewed (20)). The cause of this effect is due to the frequency of the genotype at the interacting locus. In an ideal RIL population, the loci are unlinked and therefore both genotypes affect the main effect at the QTL. However, in ILs, the loci are linked and the QTL main effects are therefore affected by the background interaction as exemplified by ILs generated for specific loci (9,24,63). The availability of two complementary IL populations therefore enables QTL dissection on a genome-wide scale.

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Author contributions

Conceived and designed the experiments: MGS, LBS, JEK. Constructed the panel: MGS, JWvC, JAGR. Sequencing and genetic map construction: DC. Performed experiments: MS, JS, KJ, YAW, JJS, SCH. Analysed the data: MGS, LvS. Wrote the paper: MGS, LvS. All authors edited and commented on the paper.

Supplementary figures and tables

Supplementary figure 1: Map of the amplification fragment length polymorphisms. For each primer-pair (indicated by the chromosome number and the start location of the deletion in CB4856) the amplicons in N2, CB4856 and the six RILs used for constructing the IL_{CB4856} panel are shown. The 100, 500, and 1000 bp bands of the 1kb+ marker are indicated. Photographs of the gels were stretched to cover the same area.

Supplementary figure 2: Power analysis in the IL_{CB4856} and the combined IL_{N2} and IL_{CB4856} panels. (A) Power analysis of bin mapping using the IL_{CB4856} panel core set of 87 ILs. The percentage of simulated QTL detected is plotted versus the number of replicates per IL. The colours indicate the amount of heritable variation simulated in the QTL. (B) Power analysis of bin mapping using both the 90 IL_{N2} and 87 IL_{CB4856} core sets. The simulated QTL contained only a marker effect. The percentage detected versus the number of replicates is for the marker-effect in the two-background model.

Supplementary figure 3: A figure of all lifespan QTL mapped. On the y-axis the various summary statistics (maximum, mean, median, and variance) for lifespan are shown. On the x-axis the position of the QTL in million bases (Mb) is shown. The dots indicate the peak of the QTL and the horizontal lines the confidence interval of the peak location. The colours indicate the significance.

Supplementary figure 4: The heat-stress survival QTL mapped in the data obtained from (37). (A) the QTL profiles as mapped in the IL_{N2} and the N2xCB4856 RIL population. On the x-axis the physical position is shown in million bases (Mb). On the y-axis the significance of the association is shown (-log₁₀(p)). The dashed horizontal line indicates the FDR = 0.05 threshold. The red triangles indicate QTL-peaks. (B) The split-out per genotype at the Chromosome IV QTL at 13.2 Mb. The ILs with the CB4856 introgression at the marker are shown versus the measurements in the N2 strain.

Supplementary figure 5: The *clec-62* expression QTL. (A) The expression of *clec-62* isoform A versus isoform B as measured by RT-qPCR in the same strains. The expression has been normalized and log₂-transformed. The Pearson correlation coefficient is shown as well as the significance of correlation. The colours indicate the various strains/panels tested. (B) the correlation between the expression of *clec-62* as measured by microarray (x-axis) versus RT-qPCR measured *clec-62* isoform A expression. Each dot represents the expression measured in a strain of either the IL_{N2} panel (orange) or RIL panel (grey). The Pearson

correlation coefficient is shown, as well as the significance of the correlation. (C) The QTL profiles of *clec-62* expression. On the x-axis the physical position is shown in million bases (Mb). On the y-axis the significance of the association is shown ($-\log_{10}(p)$). The dashed horizontal line indicates the FDR = 0.05 threshold. The red triangles indicate QTL-peaks. Where qPCR data is shown, *clec-62* isoform A was used for analysis.

Supplementary table 1: Description of the crossing setup and generations underlying each IL.

Supplementary table 2: List with the fragment length polymorphism primers used for initial tracking of segments.

Supplementary table 3: Complete list of genotypes of all the Wageningen Nematology strains of the N2xCB4856 RIL, IL_{N2} and IL_{CB4856} panels.

Supplementary table 4: Integrated genetic map for all the Wageningen Nematology strains of the N2xCB4856 RIL, IL_{N2} and IL_{CB4856} panels.

Supplementary table 5: Data of the lifespan experiment in the IL_{CB4856} panel.

Supplementary table 6: Locations and intervals of all QTL mapped for lifespan, heat-stress survival and *clec-62* expression.

Supplementary table 7: Heat stress survival data from (37) that was analysed.

Supplementary table 8: Heat stress survival data from chromosome IV IL_{N2} and IL_{CB4856} strains.

Supplementary table 9: Expression data of the *clec-62* gene as measured by RT-qPCR.

551 **Supplementary table 10:** Expression data of the *clec-62* gene as measured by microarray,
552 obtained from (41,42).

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554 **Supplementary table 11:** Heritability analysis of *clec-62* expression.

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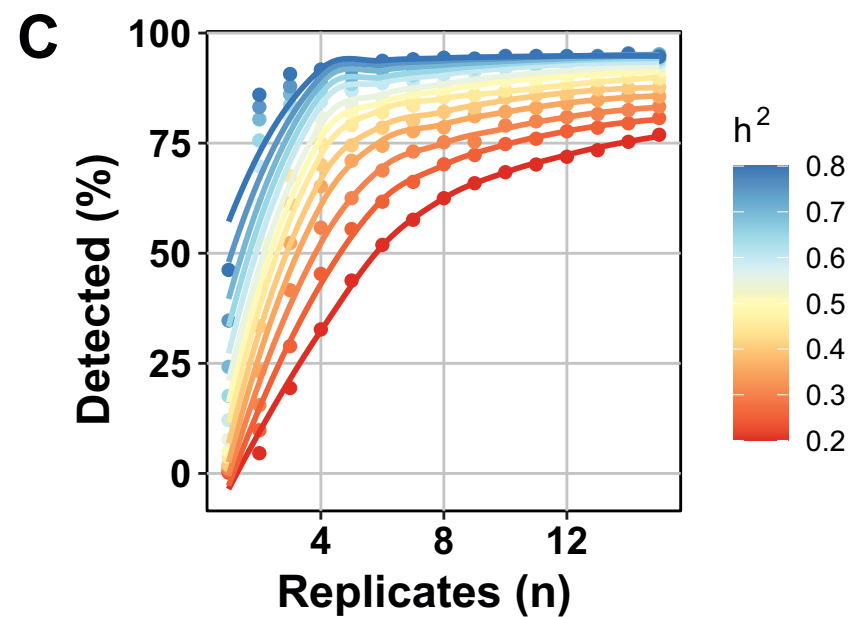
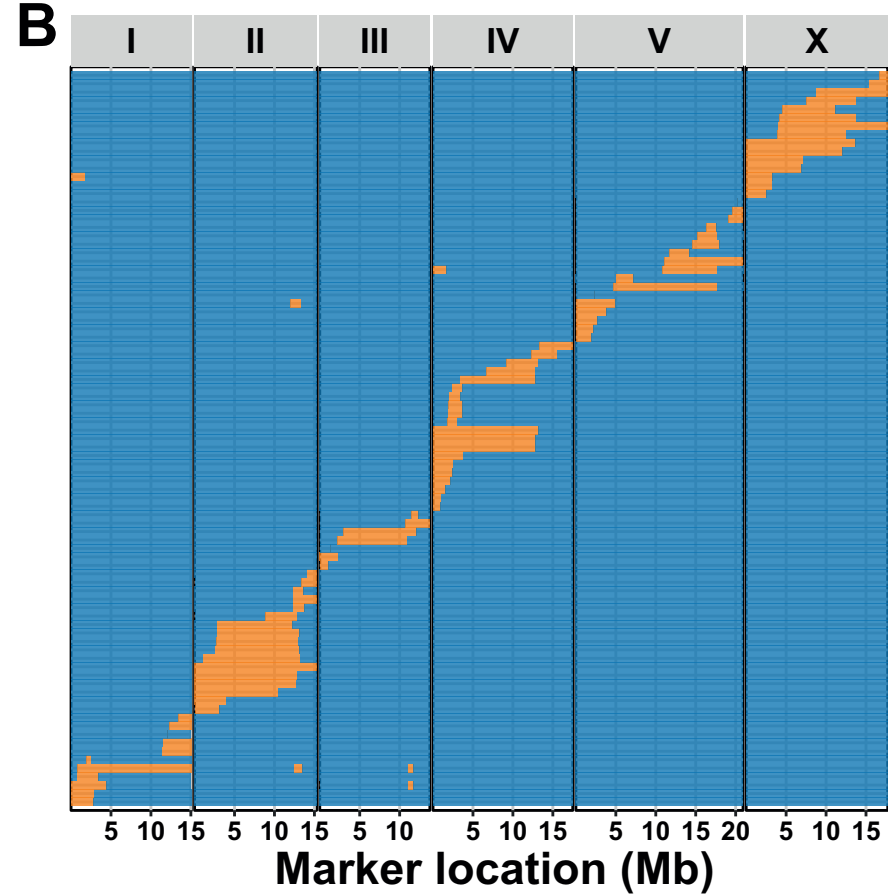
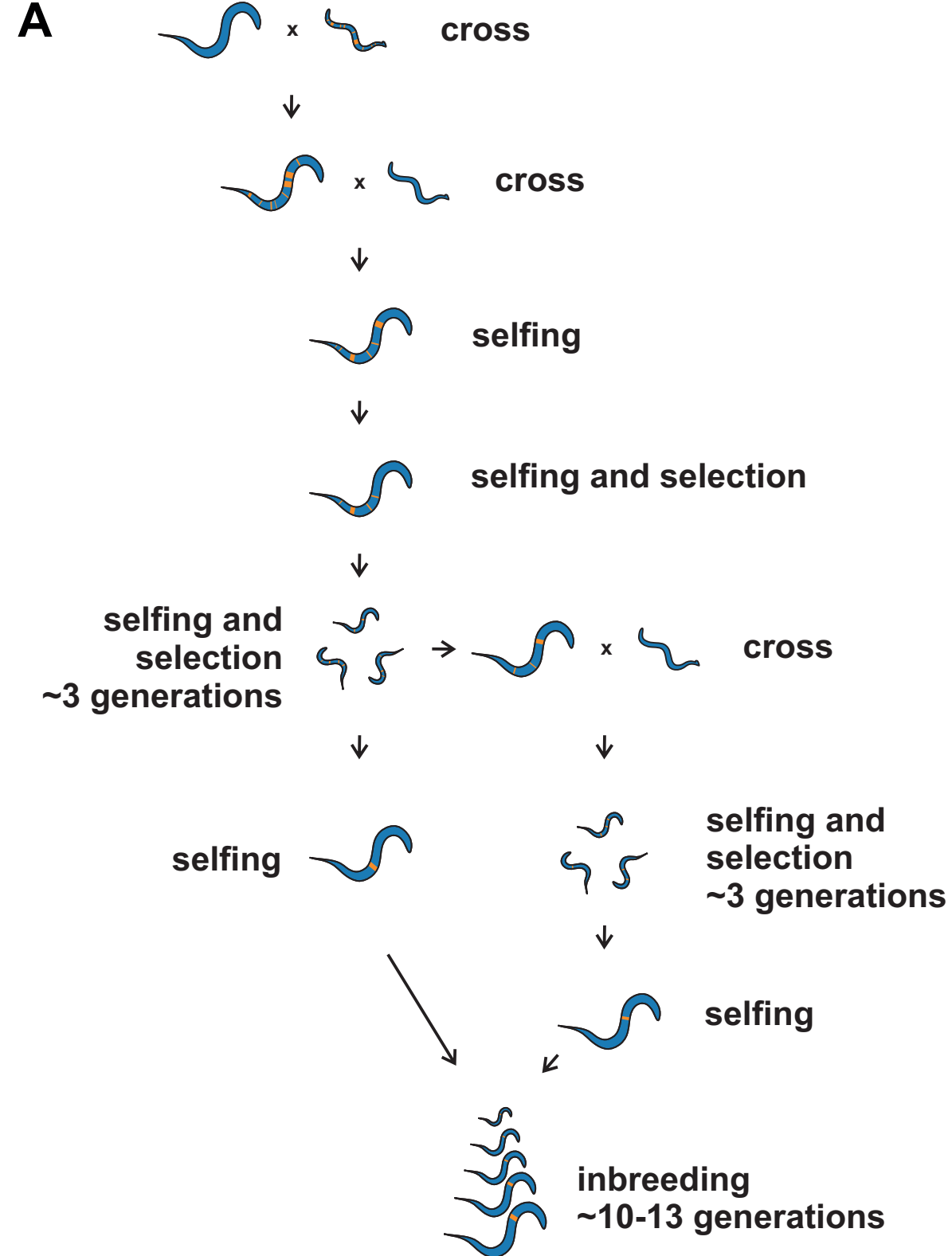
References

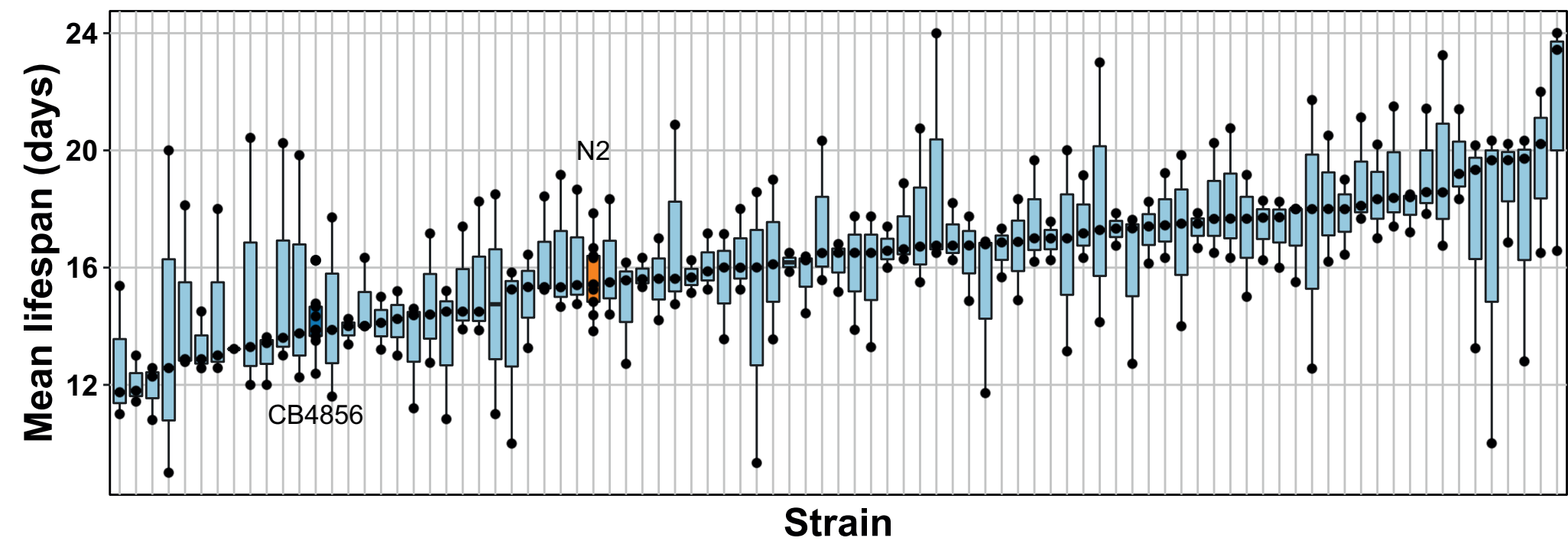
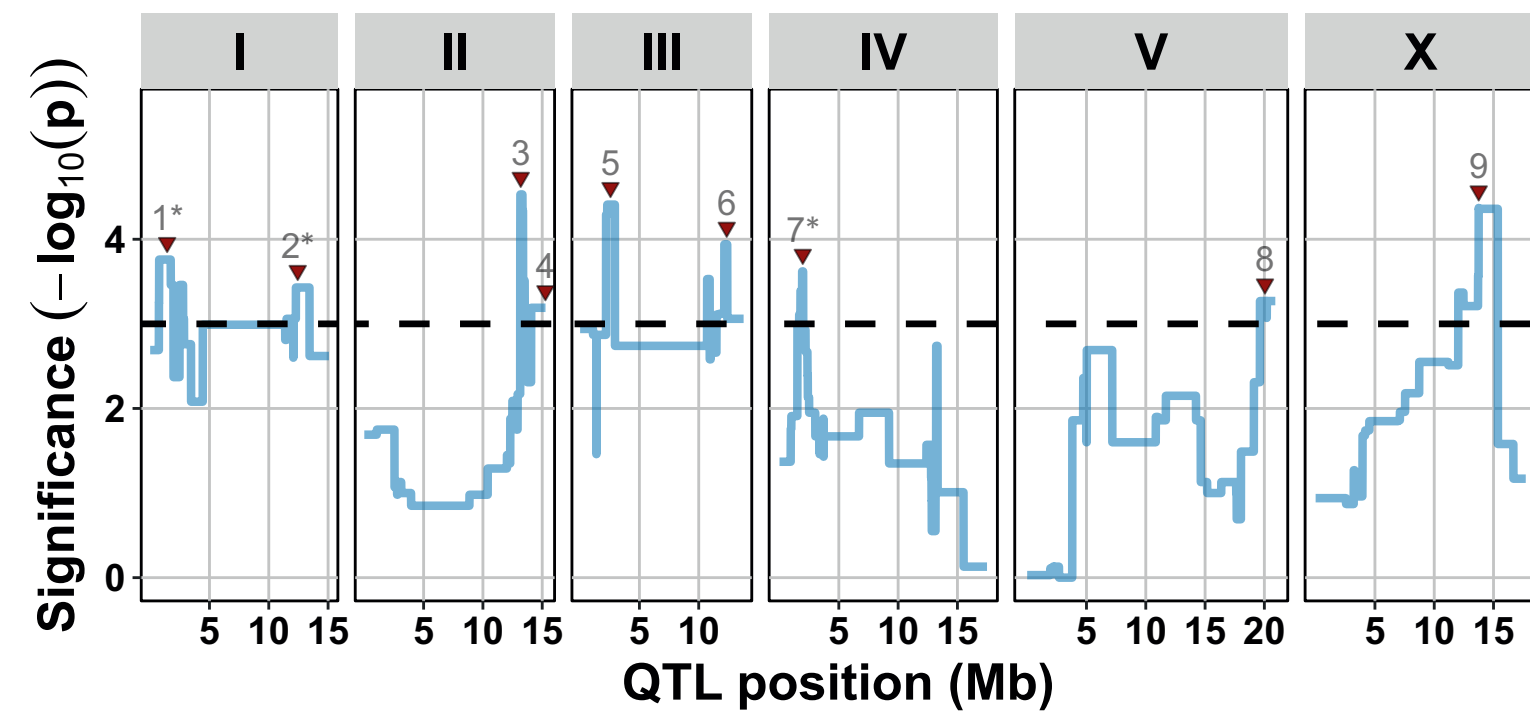
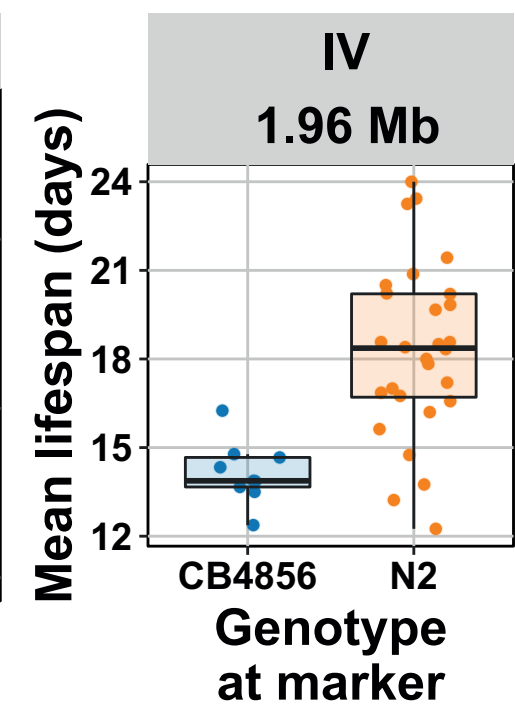
1. Evans, K.S., van Wijk, M.H., McGrath, P.T., Andersen, E.C. and Sterken, M.G. (2021) From QTL to gene: *C. elegans* facilitates discoveries of the genetic mechanisms underlying natural variation. *Trends Genet*, **37**, 933-947.
2. Kim, C., Kim, J., Kim, S., Cook, D.E., Evans, K.S., Andersen, E.C. and Lee, J. (2019) Long-read sequencing reveals intra-species tolerance of substantial structural variations and new subtelomere formation in *C. elegans*. *Genome Res*, **29**, 1023-1035.
3. Lee, D., Zdraljevic, S., Stevens, L., Wang, Y., Tanny, R.E., Crombie, T.A., Cook, D.E., Webster, A.K., Chirakar, R., Baugh, L.R. *et al.* (2021) Balancing selection maintains hyper-divergent haplotypes in *Caenorhabditis elegans*. *Nat Ecol Evol*, **5**, 794-807.
4. Thompson, O.A., Snoek, L.B., Nijveen, H., Sterken, M.G., Volkers, R.J., Brenchley, R., Van't Hof, A., Bevers, R.P., Cossins, A.R., Yanai, I. *et al.* (2015) Remarkably Divergent Regions Punctuate the Genome Assembly of the *Caenorhabditis elegans* Hawaiian Strain CB4856. *Genetics*, **200**, 975-989.
5. Andersen, E.C. and Rockman, M.V. (2022) Natural genetic variation as a tool for discovery in *Caenorhabditis* nematodes. *Genetics*, **220**.
6. Kammenga, J.E., Doroszuk, A., Riksen, J.A., Hazendonk, E., Spiridon, L., Petrescu, A.J., Tijsterman, M., Plasterk, R.H. and Bakker, J. (2007) A *Caenorhabditis elegans* wild type defies the temperature-size rule owing to a single nucleotide polymorphism in *tra-3*. *PLoS Genet*, **3**, e34.
7. Bendesky, A., Tsunozaki, M., Rockman, M.V., Kruglyak, L. and Bargmann, C.I. (2011) Catecholamine receptor polymorphisms affect decision-making in *C. elegans*. *Nature*, **472**, 313-318.
8. Frezal, L., Demoinet, E., Braendle, C., Miska, E. and Felix, M.A. (2018) Natural Genetic Variation in a Multigenerational Phenotype in *C. elegans*. *Curr Biol*, **28**, 2588-2596 e2588.
9. McGrath, P.T., Rockman, M.V., Zimmer, M., Jang, H., Macosko, E.Z., Kruglyak, L. and Bargmann, C.I. (2009) Quantitative mapping of a digenic behavioral trait implicates globin variation in *C. elegans* sensory behaviors. *Neuron*, **61**, 692-699.
10. Glauser, D.A., Chen, W.C., Agin, R., Macinnis, B.L., Hellman, A.B., Garrity, P.A., Tan, M.W. and Goodman, M.B. (2011) Heat avoidance is regulated by transient receptor potential (TRP) channels and a neuropeptide signaling pathway in *Caenorhabditis elegans*. *Genetics*, **188**, 91-103.
11. Keurentjes, J.J., Bentsink, L., Alonso-Blanco, C., Hanhart, C.J., Blankestijn-De Vries, H., Effgen, S., Vreugdenhil, D. and Koornneef, M. (2007) Development of a near-isogenic line population of *Arabidopsis thaliana* and comparison of mapping power with a recombinant inbred line population. *Genetics*, **175**, 891-905.
12. Schmalenbach, I., Korber, N. and Pillen, K. (2008) Selecting a set of wild barley introgression lines and verification of QTL effects for resistance to powdery mildew and leaf rust. *Theor Appl Genet*, **117**, 1093-1106.
13. Fletcher, R.S., Mullen, J.L., Yoder, S., Bauerle, W.L., Reuning, G., Sen, S., Meyer, E., Juenger, T.E. and McKay, J.K. (2013) Development of a next-generation NIL library in *Arabidopsis thaliana* for dissecting complex traits. *BMC Genomics*, **14**, 655.
14. Doroszuk, A., Snoek, L.B., Fradin, E., Riksen, J. and Kammenga, J. (2009) A genome-wide library of CB4856/N2 introgression lines of *Caenorhabditis elegans*. *Nucleic Acids Res*, **37**, e110.

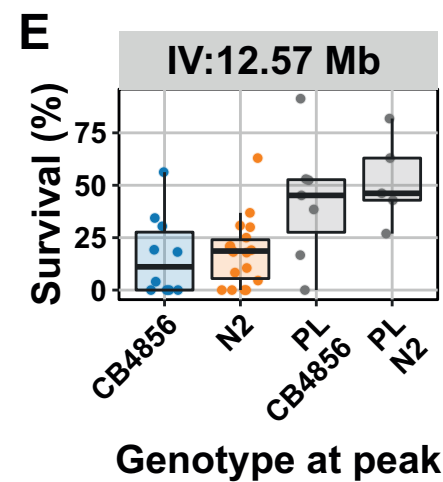
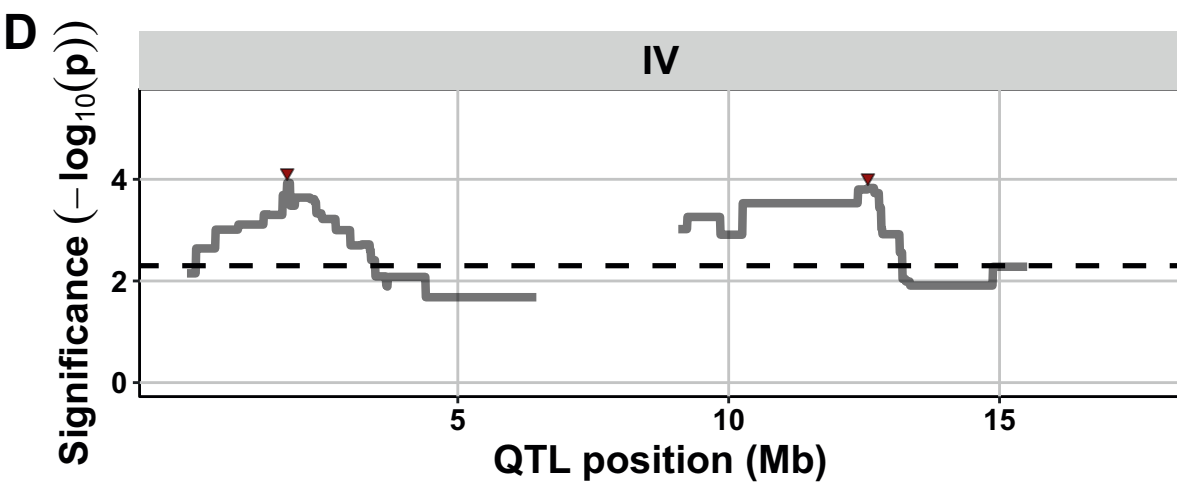
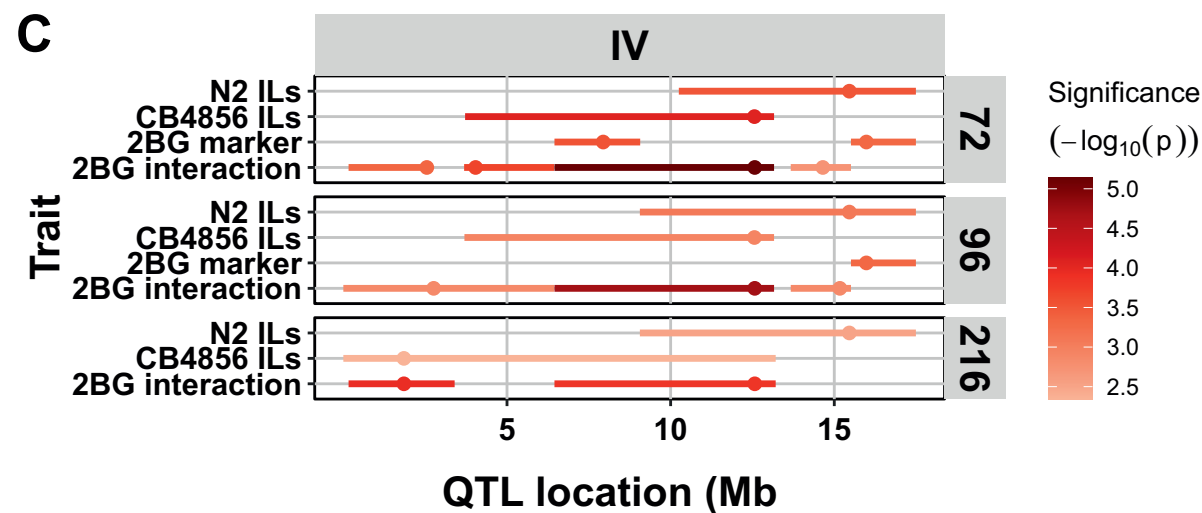
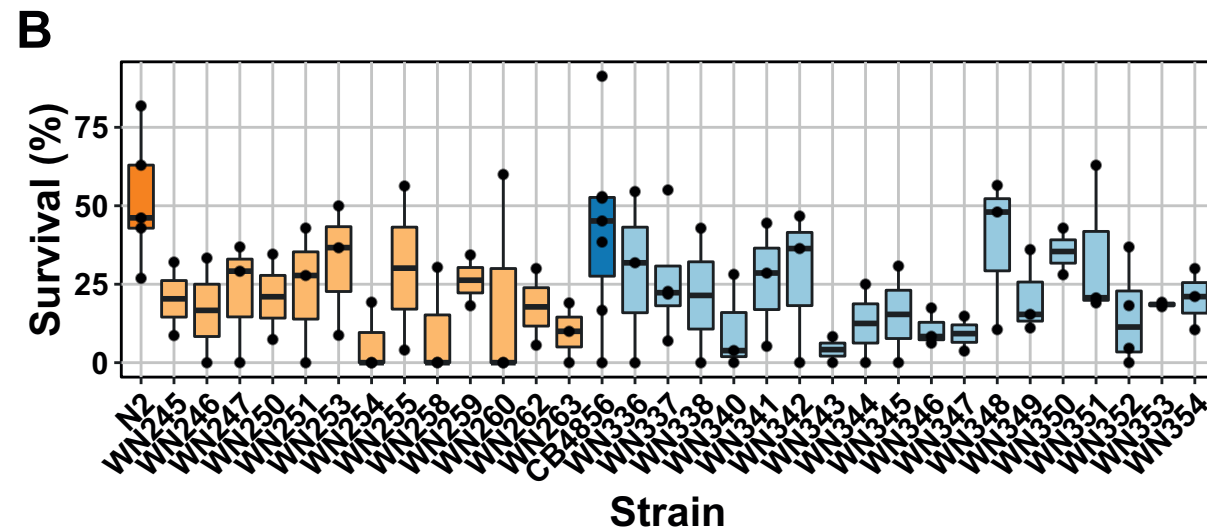
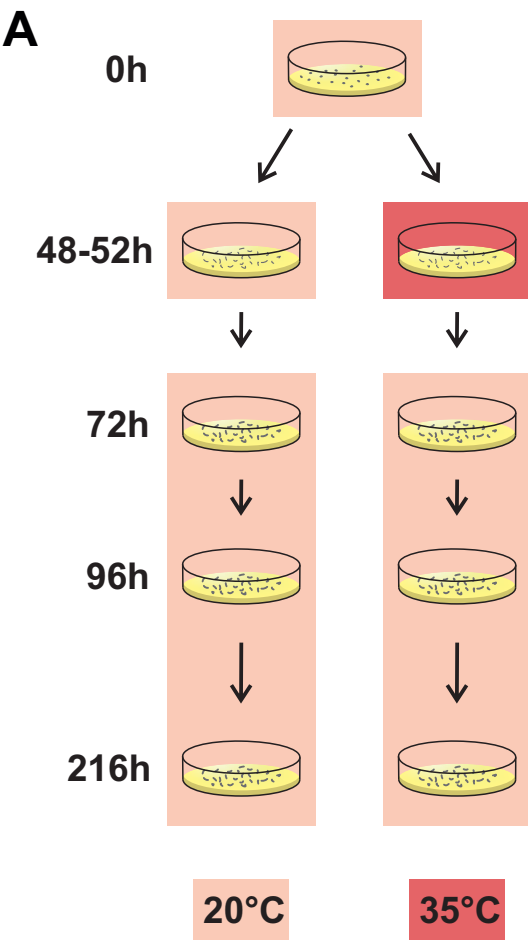
- 603 15. Glater, E.E., Rockman, M.V. and Bargmann, C.I. (2014) Multigenic natural variation
604 underlies *Caenorhabditis elegans* olfactory preference for the bacterial pathogen
605 *Serratia marcescens*. *G3 (Bethesda)*, **4**, 265-276.
- 606 16. Green, J.W., Snoek, L.B., Kammenga, J.E. and Harvey, S.C. (2013) Genetic mapping
607 of variation in dauer larvae development in growing populations of *Caenorhabditis*
608 *elegans*. *Heredity (Edinb)*, **111**, 306-313.
- 609 17. Evans, K.S. and Andersen, E.C. (2020) The Gene *scb-1* Underlies Variation in
610 *Caenorhabditis elegans* Chemotherapeutic Responses. *G3 (Bethesda)*, **10**, 2353-2364.
- 611 18. Evans, K.S., Zdraljjevic, S., Stevens, L., Collins, K., Tanny, R.E. and Andersen, E.C.
612 (2020) Natural variation in the sequestosome-related gene, *sqst-5*, underlies zinc
613 homeostasis in *Caenorhabditis elegans*. *PLoS Genet*, **16**, e1008986.
- 614 19. Bernstein, M.R. and Rockman, M.V. (2016) Fine-Scale Crossover Rate Variation on
615 the *Caenorhabditis elegans* X Chromosome. *G3 (Bethesda)*, **6**, 1767-1776.
- 616 20. Mackay, T.F. (2014) Epistasis and quantitative traits: using model organisms to study
617 gene-gene interactions. *Nat Rev Genet*, **15**, 22-33.
- 618 21. Bloom, J.S., Kotenko, I., Sadhu, M.J., Treusch, S., Albert, F.W. and Kruglyak, L.
619 (2015) Genetic interactions contribute less than additive effects to quantitative trait
620 variation in yeast. *Nat Commun*, **6**, 8712.
- 621 22. Forsberg, S.K., Bloom, J.S., Sadhu, M.J., Kruglyak, L. and Carlborg, O. (2017)
622 Accounting for genetic interactions improves modeling of individual quantitative trait
623 phenotypes in yeast. *Nat Genet*, **49**, 497-503.
- 624 23. Huang, W., Richards, S., Carbone, M.A., Zhu, D., Anholt, R.R., Ayroles, J.F.,
625 Duncan, L., Jordan, K.W., Lawrence, F., Magwire, M.M. *et al.* (2012) Epistasis
626 dominates the genetic architecture of *Drosophila* quantitative traits. *Proc Natl Acad*
627 *Sci U S A*, **109**, 15553-15559.
- 628 24. Gaertner, B.E., Parmenter, M.D., Rockman, M.V., Kruglyak, L. and Phillips, P.C.
629 (2012) More than the sum of its parts: a complex epistatic network underlies natural
630 variation in thermal preference behavior in *Caenorhabditis elegans*. *Genetics*, **192**,
631 1533-1542.
- 632 25. Andersen, E.C., Bloom, J.S., Gerke, J.P. and Kruglyak, L. (2014) A variant in the
633 neuropeptide receptor *npr-1* is a major determinant of *Caenorhabditis elegans* growth
634 and physiology. *PLoS Genet*, **10**, e1004156.
- 635 26. Li, Y., Alvarez, O.A., Gutteling, E.W., Tijsterman, M., Fu, J., Riksen, J.A.,
636 Hazendonk, E., Prins, P., Plasterk, R.H., Jansen, R.C. *et al.* (2006) Mapping
637 determinants of gene expression plasticity by genetical genomics in *C. elegans*. *PLoS*
638 *Genet*, **2**, e222.
- 639 27. Brenner, S. (1974) The genetics of *Caenorhabditis elegans*. *Genetics*, **77**, 71-94.
- 640 28. Seidel, H.S., Ailion, M., Li, J., van Oudenaarden, A., Rockman, M.V. and Kruglyak,
641 L. (2011) A novel sperm-delivered toxin causes late-stage embryo lethality and
642 transmission ratio distortion in *C. elegans*. *PLoS Biol*, **9**, e1001115.
- 643 29. Rockman, M.V. and Kruglyak, L. (2009) Recombinational landscape and population
644 genomics of *Caenorhabditis elegans*. *PLoS Genet*, **5**, e1000419.
- 645 30. Untergasser, A., Cutcutache, I., Koressaar, T., Ye, J., Faircloth, B.C., Remm, M. and
646 Rozen, S.G. (2012) Primer3--new capabilities and interfaces. *Nucleic Acids Res*, **40**,
647 e115.
- 648 31. Altschul, S.F., Gish, W., Miller, W., Myers, E.W. and Lipman, D.J. (1990) Basic local
649 alignment search tool. *J Mol Biol*, **215**, 403-410.
- 650 32. Vervoort, M.T., Vonk, J.A., Mooijman, P.J., Van den Elsen, S.J., Van Megen, H.H.,
651 Veenhuizen, P., Landeweert, R., Bakker, J., Mulder, C. and Helder, J. (2012) SSU

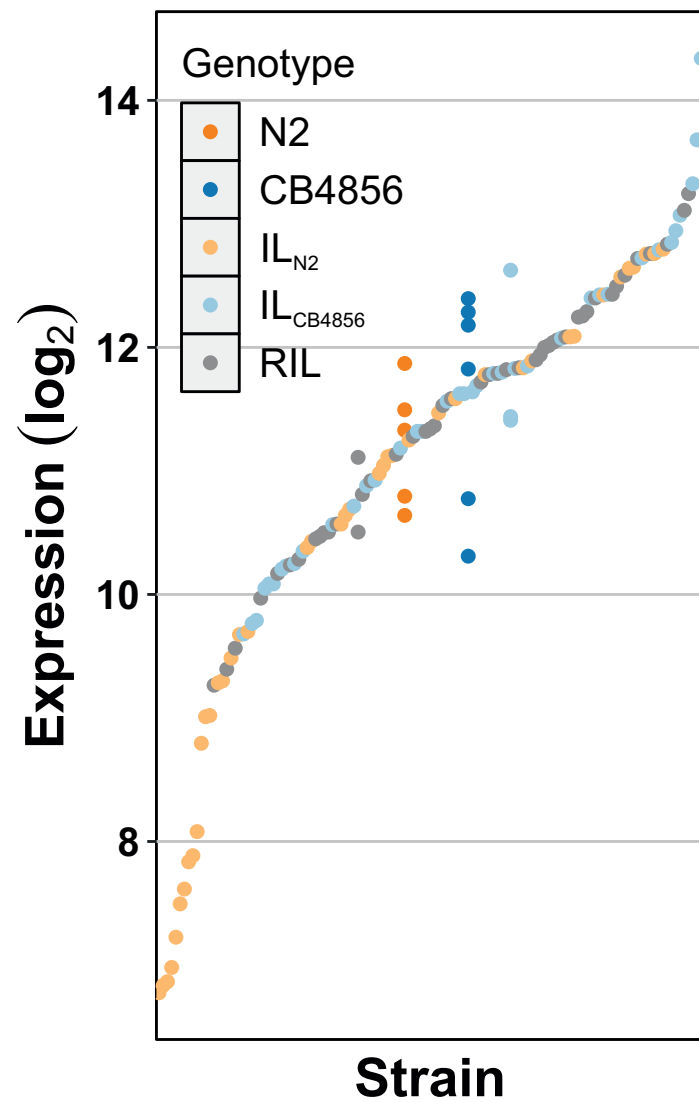
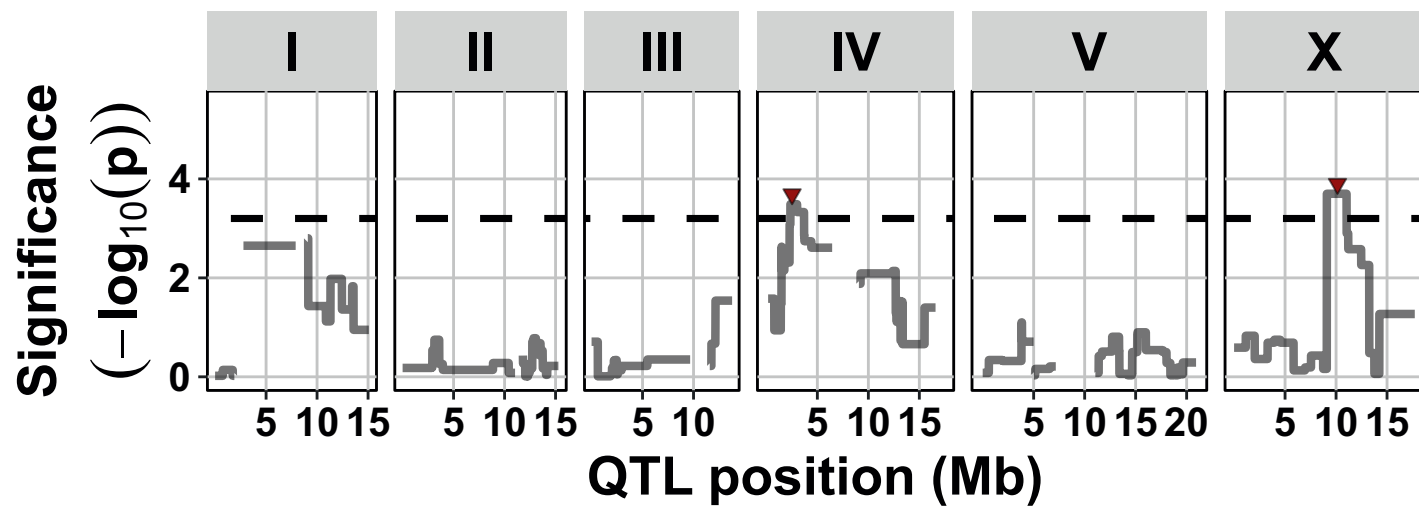
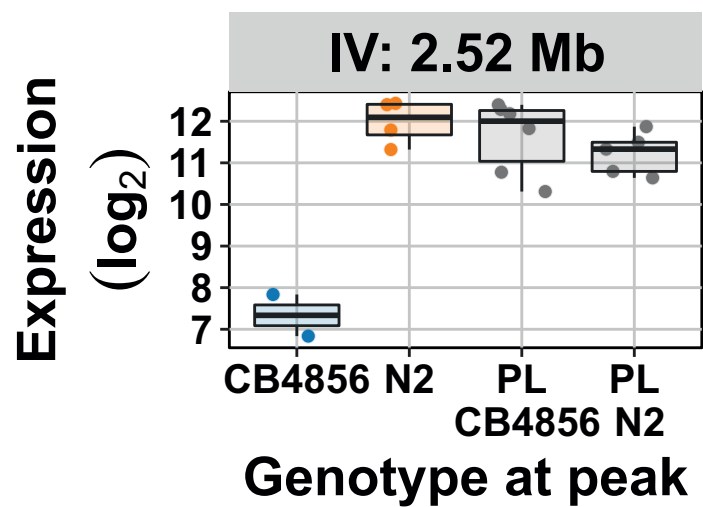
- 652 ribosomal DNA-based monitoring of nematode assemblages reveals distinct seasonal
653 fluctuations within evolutionary heterogeneous feeding guilds. *PLoS One*, **7**, e47555.
- 654 33. Gao, A.W., Sterken, M.G., Uit de Bos, J., van Creijl, J., Kamble, R., Snoek, B.L.,
655 Kammenga, J.E. and Houtkooper, R.H. (2018) Natural genetic variation in *C. elegans*
656 identified genomic loci controlling metabolite levels. *Genome Res*, **28**, 1296-1308.
- 657 34. Sterken, M.G., van Sluijs, L., Wang, Y.A., Ritmahan, W., Gultom, M.L., Riksen,
658 J.A.G., Volkers, R.J.M., Snoek, L.B., Pijlman, G.P. and Kammenga, J.E. (2021)
659 Punctuated Loci on Chromosome IV Determine Natural Variation in Orsay Virus
660 Susceptibility of *Caenorhabditis elegans* Strains Bristol N2 and Hawaiian CB4856. *J*
661 *Virol*, **95**.
- 662 35. Cook, D.E. and Andersen, E.C. (2017) VCF-kit: assorted utilities for the variant call
663 format. *Bioinformatics*, **33**, 1581-1582.
- 664 36. Cook, D.E., Zdravljivic, S., Tanny, R.E., Seo, B., Riccardi, D.D., Noble, L.M.,
665 Rockman, M.V., Alkema, M.J., Braendle, C., Kammenga, J.E. *et al.* (2016) The
666 Genetic Basis of Natural Variation in *Caenorhabditis elegans* Telomere Length.
667 *Genetics*, **204**, 371-383.
- 668 37. Jovic, K., Grilli, J., Sterken, M.G., Snoek, B.L., Riksen, J.A.G., Allesina, S. and
669 Kammenga, J.E. (2019) Transcriptome resilience predicts thermotolerance in
670 *Caenorhabditis elegans*. *BMC Biol*, **17**, 102.
- 671 38. Emmons, S.W., Klass, M.R. and Hirsh, D. (1979) Analysis of the constancy of DNA
672 sequences during development and evolution of the nematode *Caenorhabditis elegans*.
673 *Proc Natl Acad Sci U S A*, **76**, 1333-1337.
- 674 39. Hosono, R. (1978) Sterilization and growth inhibition of *Caenorhabditis elegans* by 5-
675 fluorodeoxyuridine. *Exp Gerontol*, **13**, 369-374.
- 676 40. Rodriguez, M., Snoek, L.B., Riksen, J.A., Bevers, R.P. and Kammenga, J.E. (2012)
677 Genetic variation for stress-response hormesis in *C. elegans* lifespan. *Exp Gerontol*,
678 **47**, 581-587.
- 679 41. Sterken, M.G., Bevers, R.P.J., Volkers, R.J.M., Riksen, J.A.G., Kammenga, J.E. and
680 Snoek, B.L. (2020) Dissecting the eQTL Micro-Architecture in *Caenorhabditis*
681 *elegans*. *Front Genet*, **11**, 501376.
- 682 42. Snoek, B.L., Sterken, M.G., Bevers, R.P.J., Volkers, R.J.M., Van't Hof, A., Brenchley,
683 R., Riksen, J.A.G., Cossins, A. and Kammenga, J.E. (2017) Contribution of trans
684 regulatory eQTL to cryptic genetic variation in *C. elegans*. *BMC Genomics*, **18**, 500.
- 685 43. Sterken, M.G., Snoek, L.B., Bosman, K.J., Daamen, J., Riksen, J.A., Bakker, J.,
686 Pijlman, G.P. and Kammenga, J.E. (2014) A heritable antiviral RNAi response limits
687 Orsay virus infection in *Caenorhabditis elegans* N2. *PLoS One*, **9**, e89760.
- 688 44. Benjamini, Y. and Hochberg, Y. (1995) Controlling the false discovery rate: a
689 practical and powerful approach to multiple testing. *Journal of the Royal statistical*
690 *society: series B (Methodological)*, **57**, 289-300.
- 691 45. Brem, R.B. and Kruglyak, L. (2005) The landscape of genetic complexity across 5,700
692 gene expression traits in yeast. *Proc Natl Acad Sci U S A*, **102**, 1572-1577.
- 693 46. Keurentjes, J.J., Fu, J., Terpstra, I.R., Garcia, J.M., van den Ackerveken, G., Snoek,
694 L.B., Peeters, A.J., Vreugdenhil, D., Koornneef, M. and Jansen, R.C. (2007)
695 Regulatory network construction in *Arabidopsis* by using genome-wide gene
696 expression quantitative trait loci. *Proc Natl Acad Sci U S A*, **104**, 1708-1713.
- 697 47. Kruijer, W., Boer, M.P., Malosetti, M., Flood, P.J., Engel, B., Kooke, R., Keurentjes,
698 J.J. and van Eeuwijk, F.A. (2015) Marker-based estimation of heritability in immortal
699 populations. *Genetics*, **199**, 379-398.
- 700 48. Speed, D., Hemani, G., Johnson, M.R. and Balding, D.J. (2012) Improved heritability
701 estimation from genome-wide SNPs. *Am J Hum Genet*, **91**, 1011-1021.

49. Kruijer, W. and Kruijer, M.W. (2014) Package ‘heritability’.
50. Team, R. (2015) RStudio: integrated development for R. *RStudio, Inc., Boston, MA*
URL <http://www.rstudio.com>, **42**, 84.
51. Team, R.C. (2013) R: A language and environment for statistical computing.
52. Wickham, H. (2019) *Advanced r*. CRC press.
53. Snoek, B.L., Sterken, M.G., Hartanto, M., van Zuilichem, A.J., Kammenga, J.E., de
Ridder, D. and Nijveen, H. (2020) WormQTL2: an interactive platform for systems
genetics in *Caenorhabditis elegans*. *Database (Oxford)*, **2020**.
54. Li, Y., Breitling, R., Snoek, L.B., van der Velde, K.J., Swertz, M.A., Riksen, J.,
Jansen, R.C. and Kammenga, J.E. (2010) Global genetic robustness of the alternative
splicing machinery in *Caenorhabditis elegans*. *Genetics*, **186**, 405-410.
55. Rockman, M.V., Skrovanek, S.S. and Kruglyak, L. (2010) Selection at linked sites
shapes heritable phenotypic variation in *C. elegans*. *Science*, **330**, 372-376.
56. Sterken, M.G., van Bemmelen van der Plaat, L., Riksen, J.A.G., Rodriguez, M.,
Schmid, T., Hajnal, A., Kammenga, J.E. and Snoek, B.L. (2017) Ras/MAPK Modifier
Loci Revealed by eQTL in *Caenorhabditis elegans*. *G3 (Bethesda)*, **7**, 3185-3193.
57. Duveau, F. and Felix, M.A. (2012) Role of pleiotropy in the evolution of a cryptic
developmental variation in *Caenorhabditis elegans*. *PLoS Biol*, **10**, e1001230.
58. Sterken, M.G., Snoek, L.B., Kammenga, J.E. and Andersen, E.C. (2015) The
laboratory domestication of *Caenorhabditis elegans*. *Trends Genet*, **31**, 224-231.
59. Monforte, A.J. and Tanksley, S.D. (2000) Development of a set of near isogenic and
backcross recombinant inbred lines containing most of the *Lycopersicon hirsutum*
genome in a *L. esculentum* genetic background: a tool for gene mapping and gene
discovery. *Genome*, **43**, 803-813.
60. Szalma, S.J., Hostert, B.M., Ledeaux, J.R., Stuber, C.W. and Holland, J.B. (2007)
QTL mapping with near-isogenic lines in maize. *Theor Appl Genet*, **114**, 1211-1228.
61. Li, Z.-K., Fu, B.-Y., Gao, Y.-M., Xu, J.-L., Ali, J., Lafitte, H., Jiang, Y.-Z., Rey, J.D.,
Vijayakumar, C. and Maghirang, R. (2005) Genome-wide introgression lines and their
use in genetic and molecular dissection of complex phenotypes in rice (*Oryza sativa*
L.). *Plant Molecular Biology*, **59**, 33-52.
62. Gale, G.D., Yazdi, R.D., Khan, A.H., Lusi, A.J., Davis, R.C. and Smith, D.J. (2009)
A genome-wide panel of congenic mice reveals widespread epistasis of behavior
quantitative trait loci. *Mol Psychiatry*, **14**, 631-645.
63. Kroymann, J. and Mitchell-Olds, T. (2005) Epistasis and balanced polymorphism
influencing complex trait variation. *Nature*, **435**, 95-98.



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