A genome-wide library of CB4856/N2 introgression lines of Caenorhabditis elegans

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ABSTRACT

Recombinant inbred lines (RILs) derived from Caenorhabditis elegans wild-type N2 and CB4856 are increasingly being used for mapping genes underlying complex traits. To speed up mapping and gene discovery, introgression lines (ILs) offer a powerful tool for more efficient QTL identification. We constructed a library of 90 ILs, each carrying a single homozygous CB4856 genomic segment introgressed into the genetic background of N2. The ILs were genotyped by 123 single-nucleotide polymorphism (SNP) markers. The proportion of the CB4856 segments in most lines does not exceed 3%, and together the introgressions cover 96% of the CB4856 genome. The value of the IL library was demonstrated by identifying novel loci underlying natural variation in two ageing-related traits, i.e. lifespan and pharyngeal pumping rate. Bin mapping of lifespan resulted in six QTLs, which all have a lifespan-shortening effect on the CB4856 allele. We found five QTLs for the decrease in pumping rate, of which four colocated with QTLs found for average lifespan. This suggests pleiotropic or closely linked QTL associated with lifespan and pumping rate. Overall, the presented IL library provides a versatile resource toward easier and efficient fine mapping and functional analyses of loci and genes underlying complex traits in C. elegans.

INTRODUCTION

Over the last decades, Caenorhabditis elegans has become a versatile model organism for unravelling genetic mechanisms underlying many diverse phenotypes such as stress response, ageing, host–pathogen interaction, behaviour and even learning and memory. Gene discovery in C. elegans is based on the wealth of advanced genetic methods and is facilitated by the ease of forward and reverse genetics. Although standard forward and reverse genetic analyses have led to extensive knowledge about individual genes and their specific molecular functions in pathways, we still know little about their effects on natural phenotypic variation and the overall genetic architecture of complex traits including the total number of genes, typical effect sizes for alleles or the interactions between them (1,2).

Mapping natural genetic variation in C. elegans has recently received increased attention using approaches based on quantitative trait loci (QTL) analysis using recombinant inbred lines (RIL). In QTL mapping, phenotypic traits are compared with the molecular marker genotypes of a segregating population to search for genomic regions (also called QTL) associated with the trait variation (3). For instance in C. elegans, not only QTLs have been mapped to study genetic determinants of natural variation in sensitivity to volatile anesthetics (4) and oxygen levels (5), but variations in other complex traits have also been mapped such as longevity, reproduction, body growth rate and gene expression (6–10). A QTL approach was used also to investigate phenotypic plasticity, pleiotropy, genotype–environment interactions and epistasis in C. elegans life histories (6–9,11–12).

However, mapping analyses in RILs have their own drawbacks because of the masking effects of major QTL and epistatic interactions of multiple QTL (13). In addition to RILs, introgression lines (ILs) are a more powerful resource facilitating high-resolution QTL mapping (14,15). In principle, the genome of an IL (which can also be referred to as a congenic strain or near isogenic line) is composed of a recipient genome contributed by one of the parental strains and a short, homozygous segment of the donor genome contributed by another, genetically distinct, parental strain. In this way, the difference in the phenotype of the recipient (background) strain and the IL can be precisely attributed to the introgressed locus. Without the confounding effects of other segregating QTLs that occur in RIL populations, the effect of the locus of interest can be determined with high accuracy. The additional value of IL libraries is that the selected ILs can be further backcrossed to the recipient parent and the resulting set of lines with shorter introgressions...
(sub-ILs) can be used to provide high-resolution information on the QTL architecture. ILs were successfully applied in plants to dissect QTLs of large effects into multiple, tightly linked loci (16,17) and facilitate subsequent positional cloning (18,19) providing clues for the molecular basis of gene functions.

Here, we report the first genome-wide library of *C. elegans* ILs developed from N2 Bristol (recipient) and CB4856 (donor) strains. The N2 Bristol strain is the standard strain for genetic and developmental studies. Its genome was fully sequenced in 1998 and provided the first genomic platform to study metazoan gene function (20). We chose the Hawaiian CB4856 wild type as a donor strain because of its highest genetic divergence from N2 compared to other wild isolates and a rapidly growing interest in mapping natural genetic polymorphisms underlying many diverse phenotypes. CB4856 and N2 differ by one single-nucleotide polymorphism (SNP) per 840 bp (21,22) and show a high overall difference in gene content as indicated by the comparative genomic hybridization (23). Numerous phenotypic differences between these two strains include clumping and bordering behaviour (24), ethanol responses (25), RNAi sensitivity (26), pathogen susceptibility (27) and some life-history traits (7).

Genetic and mutant analyses in N2 led to extensive knowledge about many complex human disease pathways. On the other hand, we have become more aware of the profound role of genetic and environmental interactions behind complex disease phenotypes (28). It is, therefore, of fundamental importance to understand the exact mechanisms underlying background modifiers. In mutation mapping studies using SNPs of the CB4856 strain, it has been frequently observed that the penetrance of mutations can be greatly modified by the CB4856 background indicating a high potential of the N2/CB4856 IL library for the mechanistic studies of the modifier genes. Several strategies for such studies have already been proposed and can be readily implemented (1,29).

Overall, the genome-wide *C. elegans* IL library is a unique and powerful resource facilitating the development of the new direction of complex trait mapping in *C. elegans*. Here, we genetically characterize the constructed CB4856/N2 IL library and complement this description with phenotypic analyses of lifespan (LSP) and pharyngeal pumping rate and subsequent mapping of these traits. Both traits are correlated in N2 in the sense that pharyngeal pumping rate declines cause a decline in survival probability suggesting a shared regulatory system (30).

**MATERIALS AND METHODS**

**Construction of ILs**

Three hermaphrodites of each selected (parental) RIL (WN17, WN19, WN28, WN29, WN57, WN64, WN77, WN80; Supplementary Figure 1) were backcrossed to N2 males (six males per hermaphrodite). Depending on the proportion of the CB4856 segments in the parental RIL, three to five rounds of backcrossing were performed and followed by 10 generations of selfing to obtain homozygous strains. This procedure resulted in the creation of ~500 lines (85 derived from each RIL) with a strongly reduced, but unknown contribution of donor segments. All developed lines were initially subjected to a rough genotype screening with 7–12 SNP markers. The markers were distributed evenly over all CB4856 segments previously identified in their parental RIL. Only the lines with CB4856 alleles detected on no more than one chromosome were used for detailed genotyping. These lines (~15 lines per parental RIL) were fully genotyped with 123 evenly spaced SNPs.

This procedure resulted in the identification of ~20 ILs homozygous for a single CB4856 segment. Some of these ILs and several other lines with few homozygous donor segments were further backcrossed to N2 and subsequently selfed (the protocol as mentioned above) to construct ILs with either shorter or uniquely located CB4856 segments. The lines with more than one introgression are not included in the presented IL library. All lines are stored as frozen stocks at −80°C (31).

**SNP markers**

The SNP markers and the genotyping method were the same as previously applied for creating the RIL library (10). Briefly, all markers were selected on the *C. elegans* SNP data website (http://www.genome.wustl.edu/genome/celegans/celegans_snp.cgi). For chromosomes I, II and III, we selected 20 evenly spaced markers, for chromosomes IV, V and X we selected 22, 21 and 20 markers, respectively. We selected easily detectable (i.e. with a common restriction enzyme) SNP markers with high $P_{SNP}$ values ($P_{SNP} \geq 0.7$). Suspected SNP mistypings were checked for a second time.

**Nematode culturing**

During the IL construction, the nematodes were grown, backcrossed and selfed at 20°C in six-well plates filled with the nematode growth medium, NGM, containing 2% agar and seeded with *Escherichia coli* OP50 (31). The derived lines were subsequently cultured at 15°C on 6 cm petri dishes with NGM and *E. coli*.

Prior to the LSP and pumping rate assays, the ILs, N2 and CB4856 strains were grown on 9-cm Petri dishes at 20°C for 2 weeks and synchronized by bleaching (32). Synchronization was performed during 1 day in three batches with the lines randomly assigned to each batch. After ~40 h, 12 hermaphrodites in fourth larval stage (L4) per IL were randomly picked and placed in a 6 cm petri dish for phenotyping. All experiments were carried out in Elbanton climate chambers (Elbanton, NL, USA). Temperature was monitored with a Tinytag Transit temperature logger (Gemini Data Loggers, UK). All measurements were conducted using a dissecting microscope.

**Pumping rate**

Pumping rate was defined as the number of contractions in the terminal bulb in a 15 s period and recalculated to the number of contractions per minute. This was repeated three times for each nematode, and 12 nematodes from
each plate per IL were measured by two observers. Pumping rate was measured on Days 4, 6, 8 and 10 after hatching.

**Lifespan**

The LSP measurement started with 12 nematodes per IL (in the fourth larval stage). In the first week, the number of surviving individuals was counted every day and after that, every second day. The nematodes were assumed dead if no reaction was observed after repeated touching with a hair. The nematodes were transferred daily to a new plate until the reproduction was finished. If they were observed to have crawled up the wall of a plate and have died there, they were removed from the measurement. The same was done when a nematode was observed to have died due to unnatural causes, such as developing a ‘bagging’ phenotype, where eggs hatch inside the adult.

**Bin mapping**

Per marker, the ILs containing a CB4856 allele were grouped into bins. Also, the ILs containing a CB4856 marker +1 or −1 of the marker under study were selected. The trait values of the grouped ILs were than tested against N2, by a two-sided t-test (assuming unequal variance) for average LSP and by linear regression for the decrease in pumping rate in time. The pumping rate coefficient (see Supplementary Table 1) of this regression equation was used for QTL mapping. A further decrease in QTL size was done by obtaining the locus that contained the most individual ILs significantly different from N2.

**RESULTS**

**Construction of a C. elegans IL population**

We constructed a population of 90 ILs, each containing a single CB4856 introgression in the N2 background. Together these ILs contain almost the whole CB4856 genome. The library was created by backcrossing a limited set of RILs of the CB4856 × N2 population (10) with N2 (Supplementary Figure 1). These RILs were selected based on a low proportion of CB4856 segments (~30%). After backcrossing and selfing, the genotypes of the obtained ILs were determined by the 123 markers used to genotype the CB4856 × N2 RILs (Supplementary Table 1).

Genetic constitution of the IL library is presented in Figure 1. The CB4856 segments in the different ILs predominantly overlapped; however, a minor fraction of adjacent introgressions did not overlap. We estimated that a maximum of ~3.8 Mb (megabase) was not covered by our set of ILs giving the set >96% coverage of the CB4856 genome. Furthermore, we obtained 32 ILs with two separate CB4856 segments (Supplementary Table 1).

The median segment length, defined as the distance between the two markers having N2 alleles flanking the CB4856 introgression, was ~3.5 Mb (average ~4.5 Mb) with a minimum of ~1.45 Mb and a maximum of ~16.3 Mb. Collectively, these 90 ILs have 79 distinguishable loci with a median length of ~1.7 Mb (average ~2.0 Mb; min.: ~0.6 Mb; max.: ~7.0 Mb).

**Loci affecting LSP and pumping rate**

To benchmark the use of our population of ILs in identifying QTLs, we measured LSP and pharyngeal pumping (PP) of 86 ILs giving a genome-wide coverage (Supplementary Table 1). A relation was found between LSP and PP, where pumping decreased in aging worms across the ILs (Figure 2A and B). This is in agreement with similar studies carried out in N2 by Huang et al. (30). The average PP at each time point was also comparable to the values reported by Huang et al. (30). The average PP across the IL population decreased and the variation increased as the worms got older (Figure 2A). We found variation among ILs in absolute PP (Figure 2B) as well as in the rate of decrease in time. CB4856 showed a sharper decrease in pumping than N2. Most ILs had a PP decrease rate in between the parental strains, mostly similar to N2 (Figure 2B). We also found a large difference for LSP across the ILs, varying from 10 days to 21 days, extremes larger and smaller than the parental strains (Figure 2C).

Obtaining the QTLs for both traits can give an estimate for the level of genetic intertwinement by colocation of QTLs. While mutant studies show a large decrease in LSP when a gene affecting PP is functionally compromised, such as reported by (30), not much is known about natural occurring alleles affecting this relation. Bin mapping of average LSP resulted in six QTLs, which all have a LSP-shortening effect on the CB4856 allele (Figure 3). Although CB4856 has a shorter average LSP than N2, the difference between CB4856 and N2 is less than the sum of effects of the loci detected with the ILs (6.8 days versus 26.3 days, respectively). This indicates that the effects of some LSP affecting loci are caused by the interaction between the CB4856 allele and the N2 background. The detection of QTL for LSP...
on chromosomes II, IV and X confirms the position of similar QTL detected by (6,9,11) although these studies were based on a mapping population derived from different parental strains. The QTLs on chromosomes I and III have not been previously reported and might harbour novel loci and candidate genes affecting LSP. We found five QTLs for the decrease in PP, of which four colocalated with QTLs found for average LSP (Figure 3). This suggests pleiotropic or closely linked QTL affecting LSP and PP.
DISCUSSION

We report the first construction and properties of a genome-wide mapping library consisting of 90 ILs in *C. elegans*, a powerful resource to study complex traits in this model species. The choice of the parental strains of our IL library is one of the major factors ensuring a high power of genetic dissection. While the overall level of polymorphism among wild isolates of *C. elegans* is relatively low, the genetic distance between the N2 and CB4856 is high. In fact, CB4856 is reported to show the highest genetic distance from N2 compared to other wild isolates (21). CB4856 exhibits genetic isolation from all other global *C. elegans* populations that were demonstrated to intercross and recombine (33). This divergence is also reflected by the profound differences in the traits like formation of the copulatory plug (34,35) and many life-history traits (7,8). The use of the N2 strain as the recipient for CB4856 introgressions is additionally justified by the common use of N2 in *C. elegans* research and the wealth of available genetic and genomic tools.

In both RILs and ILs, the mapping resolution depends on the number of lines used and the frequency of recombination. In the case of RILs, the recombination frequency in the lines is fixed and can be increased by either adding more RILs or by intercrossing lines before fixation by inbreeding. In ILs, the resolution can be increased by minimizing the introgressed segment of each IL. Consequently, to maintain genome-wide coverage more lines are required. Because RILs and ILs have different genetic makeups, they both require different mapping tools. Overall, the recombination frequency in RILs is higher compared to an equal number of ILs. Therefore, fewer lines are required for analysis to obtain the same resolution. Each RIL has more than one introgressed fragment, and, on average, each genomic region is represented by the same number of parental genotypes. For these reasons, replication of individual RILs is often not necessary. Moreover, the multiple introgressions per RIL allow the detection of genetic interactions between loci (epistasis). Additionally, QTL of large effect might hide the detection of QTL with small additive effects. In contrast, ILs increase the power to detect small-effect QTL, but the presence of a single introgressed fragment does not allow testing for genetic interactions.

The introgressed loci within the ILs are mostly much smaller than the QTL intervals obtained with the *C. elegans* RIL population (10) increasing the resolution, while reducing the number of candidate genes and effort of fine mapping. The current IL library provides a relatively high mapping resolution, as compared to other genome wide IL populations. For example, there are many more donor segments compared to recently developed ILs of mice. These are called consomic strains because they are chromosome substitution strains and represent the most recent addition to the mouse genetic resources aiming to genetically analyse QTL (36,37).

So far there are only a few other genome-wide IL libraries available in barley (38) and *Arabidopsis* (13). In *Arabidopsis*, 40 lines derived from *Ler* and *Cvi* parental accessions contained a single introgression, while 52 lines carried several *Cvi* fragments in a *Ler* background. The genetic length of the introgression fragments ranged from 31.7 to 5.2 cM. Lines with multiple *Cvi* fragments carried a main large introgression and several much smaller *Cvi* fragments. They selected a core set of 25 lines covering >90% of the genome.

In principle, there are two main approaches of QTL mapping using IL libraries. In the first one, a number of ILs are selected based on the location of QTLs previously identified with the use of another mapping population (e.g. RILs) and analysed to confirm and fine-map the putative QTLs. The second approach, applied in this paper, involves direct library-wide screens or bin mapping. The advantage of the first approach is that it might result in the initial detection of QTLs interacting with another locus, which might become undetectable when introgressed to a common background. These QTLs can be further analysed with the lines carrying combined donor segments constructed from multiple ILs. The second approach, in turn, allows the detection of the QTLs of small effects due to the reduced background noise and does not require primary QTL analysis with other mapping populations (39).

While the examination of 123 SNP markers indicated no undesired CB4856 segments in the presented ILs (except for a single marker in ewIR28 and ewIR75), the presence of additional short donor segments within a region flanked by two adjacent markers (or the regions between the ends of the chromosomes and the closest markers) cannot be entirely excluded. Genotyping using high-density marker would allow the detection of the additional CB4856 segments and taking them into account in QTL analysis. Nevertheless, the current average distance between markers is 2.38 cM and the probability of crossovers within such a small genomic region is low. Consequently, undetected CB4856 segments are expected to be rare.

Currently, one of the greatest challenges of QTL research is the development of strategies and resources that allow for the efficient and accurate dissection of QTLs of small effects with the aim to detect causal genes. The presented genomic library of ILs is the first step to achieve these goals and significantly improve the QTL detection process in *C. elegans*.

Although beyond the scope of this article, the LSP QTL on chromosome X may guide us toward gene identification underlying LSP. This QTL harbours *npr-1*. This gene is known to be polymorphic between CB4856 and N2 (24). The two alleles cause a difference in pathogen susceptibility (27) and social feeding (24). The difference in pathogen susceptibility was measured by a shorter LSP of lines carrying the CB4856 allele of *npr-1*. It is likely that LSP of *C. elegans* grown on *E. coli* (OP50) also depends on the allele type of *npr-1*. This could explain the reduced LSP of the IL ewIR81 carrying the CB4856 allele of *npr-1*. Moreover, this line shows significant more clumping behaviour than N2 confirming the functional difference between the N2 and CB4856 *npr-1* alleles. This IL library might therefore be useful to further focus on this candidate gene, and, for instance to search for
epistatic interactions with other loci using crosses between different ILs.

The present IL library was used to search for novel loci associated with LSP and pumping rate to illustrate the potential application of this resource. We believe that the library will accelerate the discovery of natural polymorphisms underlying complex traits and will lead to a better understanding of the mechanisms behind the observed phenotypic variation in *C. elegans*. The ILs are freely available on request by sending an email to the corresponding author J.K.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online.

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