Featured Organism

Caenorhabditis elegans — applications to nematode genomics

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Abstract

The complete genome sequence of the free-living nematode Caenorhabditis elegans was published 4 years ago. Since then, we have seen great strides in technologies that seek to exploit this data. Here we describe the application of some of these techniques and other advances that are helping us to understand about not only the biology of this important model organism but also the entire phylum Nematoda. Copyright © 2003 John Wiley & Sons, Ltd.

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Background

Caenorhabditis elegans is a free-living soil nematode which has been developed as a model metazoan for over three decades (Brenner, 1974), work that recently resulted in the award of a Nobel Prize. C. elegans is a highly amenable experimental organism: it is relatively small (~1 mm in length), transparent, easy to culture and has a short generation time (~3 days). Furthermore, with the availability of its full genome sequence and with a host of well-developed experimental techniques C. elegans provides an ideal model for exploring the molecular aspects of a range of cell and developmental processes. Hence, as a model, it has proved invaluable in helping understand a number of biologically and medically important processes, including development, cancer, ageing and neurobiology (Boulton et al., 2002; Coates and de Bono, 2002; de Bono et al., 2002; Liu et al., 2002; Partridge and Gems, 2002; Wodarz, 2002).

In addition to its role as a model metazoan, C. elegans is also the best characterized member of the phylum Nematoda. Phylogenetic analyses reveal that nematodes can be split into five major taxonomic groups (or clades) with C. elegans being a clade V organism (Blaxter, 1998; Dorris et al., 1999). They are a highly diverse group of organisms occupying a diverse set of habitats, from obligate parasites of many forms of life, including plants, arthropods, fish and mammals to free-living organisms occupying niches such as soil and marine environments. It is interesting to note that parasitism appears to have evolved at multiple times during the evolution of the phylum (Dorris et al., 1999).

The C. elegans genome

Although over 97% of the 100 Mb C. elegans genome was available since the first publication in 1998, the full sequence was only finally finished at the end of 2002, making it the first truly complete animal genome. Analysis and annotation of the genome has been an ongoing process and will continue for the foreseeable future as new methods and data become available. Central to these analyses has been the WormBase consortium, a network of researchers involved in developing a web-based resource for the C. elegans genome (Harris et al., 2003; Stein et al., 2001). WormBase (available at http://www.wormbase.org, current release WS93b, 16 January 2003) is a continually evolving entity, incorporating not only the full genome sequence and associated gene models but also a
whole wealth of other functional annotation (e.g. gene expression data, and double-stranded RNA knockdown experiments). WormBase is regularly updated and is being continually expanded as new data becomes available.

The *C. elegans* genome, split into five autosomes and one sex chromosome, is currently predicted to encode 21,197 proteins, of which 2927 are splice variants. Protein coding genes and RNA genes are initially predicted *ab initio* using the GeneFinder algorithm (Wilson *et al*., unpublished; available at http://ftp.genome.washington.edu/cgi-bin/Genefinder). Genes are then further annotated in a manual process involving data from a range of different sources, including expressed sequence tag (EST)/mRNA sequence information, published papers and direct contact with the worm community. Interestingly, only ∼12,000 genes have been associated with ESTs despite the generation of over 191,524 ESTs (K. Bradnam, personal communication). This is probably due to the limited range of life cycle stages sampled by the cDNA libraries used to generate these sequences. Many genes may only be expressed at transient points in the life cycle or under specific conditions.

Protein-coding genes may be classified into three categories:

- **Confirmed (3633 genes):** there is transcript evidence for every base in every exon.
- **Partially confirmed (8743 genes):** there is some transcript evidence; however, the whole gene is not fully supported.
- **Predicted (8821 genes):** those without any transcript support.

Although gene prediction software is being continually developed, gene models associated with the last two categories may be inaccurate due to the lack of supporting data. The presence of pseudogenes (recently suggested to be as many as 20% of all annotated *C. elegans* genes; Mounsey *et al*., 2002) further complicates the process of gene annotation. In a large-scale attempt to aid annotation, Reboul *et al*. (2001) are utilizing a systematic PCR-based approach to verify the gene model and expression of the open reading frames (ORFs) for each predicted gene. To date this approach has confirmed over 12,000 genes (Vaglio *et al*., 2003) and current estimates argue for the existence of at least 17,300 genes. Furthermore, sequencing of the ORF-PCR products has helped correct the intron–exon structure for over a quarter of the predicted genes. There are also numerous non-protein-coding RNA genes: in addition to the easily recognizable rRNA, snRNA and tRNA genes, a number of small non-coding regulatory RNAs (termed microRNAs) have recently been discovered which appear to play an important role in regulating gene expression. Due to their small size (typically ∼22 bp), they are very difficult to recognize in the raw sequence and will therefore require sophisticated methods to identify such as the approach based on bioinformatics, comparative genomics and cDNA cloning used by Lee and Ambros (2001).

In terms of chromosomal organization, gene distribution between the chromosomes is non-uniform (see Table 1; *C. elegans* Sequencing Consortium, 1998). Genes appear to be denser towards the centre of the chromosomes, whilst tandem repeats and pseudogenes are more prevalent at the ends of chromosome arms. These findings suggest that these regions are evolving at a higher rate than the centre of the chromosomes (*C. elegans* Sequencing Consortium, 1998). In terms of gene organization, *C. elegans* and other nematodes appear unique among metazoans in having genes organized in operons (Blumenthal, 1998; Evans *et al*., 1997; Guiliano *et al*., 2002). These operons are resolved by trans-splicing: the first gene in the operon receives the spliced leader sequence (SL1), while downstream genes receive either SL1 or one of a family of alternative spliced leader sequences (SL2s). Whereas SL1 is added to many transcripts that are not part of operons, it seems that SL2s are only found on downstream genes in operons. A recent analysis using the occurrence of SL2 to define operons suggests that there are ∼1000 operons containing two to eight genes (Blumenthal *et al*., 2002).

**The *C. briggsae* genome**

This year saw the release of the first draft of the *Caenorhabditis briggsae* genome. *C. briggsae* is another free-living soil nematode thought to have diverged from *C. elegans* ∼50–150 million years ago (Coghlan and Wolfe, 2002). The current *C. briggsae* genome assembly is 102 MB, comprising 142 pieces, and is essentially 98% complete (http://www.sanger.ac.uk/Projects/C_briggsae/).
Table 1. Distribution of the 21,197 predicted genes in *C. elegans* across the six chromosomes indicating their relative gene density, the proportion of genes giving an RNAi phenotype, and their homology to two different sequence databases

| Chromosome | No. of predicted genes | Length (Mb) | Gene density (genes/Mb) | RNAi phenotypes (% of genes) | Proteins with similarity to SWISS-TrEMBL<sup>1a</sup> (% of genes) | Proteins with similarity to *C. briggsae*<sup>1b</sup> (% of genes) |
|------------|------------------------|-------------|-------------------------|------------------------------|----------------------------------------------------------------|------------------------------------------------------------------|
| I          | 3,068                  | 15.08       | 203.4                   | 13.7                         | 1,949 (64)                                                        | 2,788 (91)                                                        |
| II         | 3,692                  | 15.26       | 241.9                   | 11.9                         | 1,955 (53)                                                        | 3,237 (88)                                                        |
| III        | 2,957                  | 13.78       | 214.6                   | 18.5                         | 1,853 (63)                                                        | 2,665 (90)                                                        |
| IV         | 3,409                  | 17.49       | 194.9                   | 10.9                         | 1,958 (57)                                                        | 3,085 (90)                                                        |
| V          | 5,082                  | 20.92       | 242.9                   | 5.2                          | 2,252 (44)                                                        | 4,603 (91)                                                        |
| X          | 2,989                  | 17.72       | 168.7                   | 5.6                          | 1,635 (55)                                                        | 2,567 (86)                                                        |

<sup>1</sup> Predicted proteins from the *C. elegans* genome were searched for sequence similarity using BLAST against two different databases: (a) BLASTP vs. SWISS-TrEMBL — a comprehensive protein sequence database which combines SWISS-PROT with the translation of all protein-coding sequences from the EMBL nucleotide sequence database. Nematode proteins were removed from the database before analysis; (b) TBLASTN vs. the *C. briggsae* draft genome. The numbers given are those proteins which had a raw BLAST score of $>$50 against the respective databases.

Initial analysis suggests that the coding regions are more similar to the *C. elegans* genome than the non-coding regions. Thus, in addition to *C. elegans* providing a springboard for annotation of the *C. briggsae* genome, identification of homologous sequences within their respective genomes offers a new method of gene identification for *C. elegans* (Lee and Ambros, 2001). Furthermore, Webb *et al.* (2002) have found that those intergenic regions that also share homology between the two genomes may also be involved in gene regulatory functions. BLAST analysis of the *C. elegans* protein dataset suggests that ~10% of *C. elegans* genes do not share significant similarity with a *C. briggsae* gene and that these genes are spread equally between the six *C. elegans* chromosomes (our unpublished data; see Table 1). However, a lower proportion of predicted proteins from chromosome V appear to have a significant match to a sequence in the SWISS-PROT/TrEMBL database than those from other chromosomes. This may indicate that chromosome V contains a higher proportion of nematode-specific genes.

Global comparisons of sections of the *C. elegans* and *C. briggsae* genomes suggest that *Caenorhabditis* spp. have a rearrangement rate of 0.4–1.0 breakages/Mb/million years (Coghlan and Wolfe, 2002), which is at least four times faster than the rate in *Drosophila melanogaster*. These comparisons are an ongoing exercise and are expected to reveal many interesting insights into their evolution.

Other nematode genomes

Many species of nematode pose a significant risk to human health and agriculture worldwide. This has led to the initiation of a number of sequencing projects aimed at elucidating some of the molecular aspects of parasitism. At present 33 different nematode species are the subject of EST sequencing projects (Parkinson *et al.*, 2001). In an attempt to place this sequence data in a genomic context, ESTs from each species are clustered on the basis of sequence similarity to form a non-redundant set of putative gene transcripts. Initial comparisons of the ‘partial genomes’ from a few species reveals that the degree of similarity between parasitic nematodes and *C. elegans* reflects their underlying phylogeny (unpublished data). For example, of the predicted genes from the mouse whipworm *Trichuris muris* (a clade I nematode) only ~40% have significant similarity to a *C. elegans* gene, whilst of the predicted genes from the sheep hookworm *Haemonchus contortus* (a clade V nematode) ~65% appear to share significant similarity with *C. elegans*. Although these figures for similarity may be artificially low due to the small number of cDNA libraries sampled, they do suggest that whilst *C. elegans* may serve as a useful model of clade V nematodes, transfer of genomic knowledge to nematodes from less related clades may prove problematic. In addition to these EST initiatives, last year the Institute for Genome Research (TIGR) received funding for a three-fold shotgun coverage of the...
110 Mb genome of the filarial parasite Brugia malayi (http://www.tigr.org/tdb/e2k1/bma1).

To date there is little information on the amount of synteny between C. elegans and related parasites. However, last year one study published a comparison of a genomic fragment from B. malayi with the syntenic region from C. elegans. Synteny was observed, with the order, composition and configuration of genes being conserved for putative orthologous genes (Guiliano et al., 2002).

Functional genomics

Less than 5% of the 21 197 predicted proteins encoded by the C. elegans genome have so far been characterized by conventional genetics or biochemistry. To redress this deficit of biological information over sequence information, several laboratories have initiated high-throughput functional genomic approaches based on generation of loss-of-function phenotypes, analysis of expression profiles and protein–protein interactions.

RNA interference

In the 6 years since it was first discovered in C. elegans (Fire et al., 1998), double-stranded RNA-mediated interference (RNAi) has become an integral part of many C. elegans laboratories and is often used as a first approach when investigating gene function. When compared with classical mutagenesis screens, inactivation of specific gene products provides a more rapid link between sequence data and biological function. Knockdown of gene expression can be provoked by injection of the dsRNA of interest into the gonad (Fire et al., 1998), by soaking in dsRNA (Tabara et al., 1998) or by feeding on dsRNA-expressing bacteria (Timmons et al., 2001). Regardless of the route of administration, the RNAi effect is transported across cellular boundaries, so that its effect is seen throughout most of the worm and is transmitted to the offspring of treated worms. These effects are due to distinct mechanisms (Grishok et al., 2000; Ketting et al., 1999). Bacterial feeding, in particular, is an easy and efficient technique capable of being performed in most laboratories (Kamath et al., 2001).

The knockdown of 16 757 genes (around 80% of the predicted gene complement) by RNAi has recently been carried out, representing the first near-complete targeting of a metazoan genome (Kamath et al., 2003). RNAi phenotypes were assigned to 1528 genes, more than 1000 of which had not previously been annotated with a biological function. Overall, 63% of loci already known by mutation were detected by RNAi and gave similar phenotypes. Significant differences were detected in the types of genes resulting in 'non-viable' phenotypes (lethal and sterile phenotypes) and 'viable phenotypes' (post-embryonic phenotypes). C. elegans genes having clear orthologues in other eukaryotes were far more likely to produce a non-viable RNAi phenotype than non-orthologous genes, probably due to the fact that conserved genes tend to be essential. Viable phenotypes, on the other hand, were more likely to be related to genes of unknown function, underlining our current lack of understanding of metazoan development. Table 1 shows the distribution of phenotypes obtained across the six chromosomes. Significantly, genes targeted on chromosomes V and X are less likely to show a phenotype than those on the other chromosomes. In the case of chromosome V this may be associated with the greater number of genes that are nematode-specific (and hence less likely to perform essential functions). For chromosome X, many more genes are associated with worm behaviour or morphology and are thus less likely to show non-viable phenotypes.

A key advantage of this impressive study is the construction of a re-useable library of dsRNA-expressing bacteria, which is made available to other researchers via the HGMP Resource Centre (http://www.hgmp.ac.uk). The library has already been used to screen for genes involved in ageing (Lee et al., 2003) and fat regulation (Ashrafi et al., 2003).

In addition to RNAi, the C. elegans Gene Knockout Consortium has generated several hundred mutant lines using a PCR-based method to detect deletions in genes specifically requested from the C. elegans community (http://elegans.bcgsc.bc.ca/knockout.shtml). Practically, these mutants offer the advantage of generating stable mutant lines that can be maintained indefinitely.

Through these gene inactivation and deletion studies, we now know that the vast majority
of genes yield no obvious phenotype. A further observation in RNAi screens is that only about two-thirds of genes already identified by classical mutational genetics give RNAi phenotypes. Although RNAi and deletion mutants are undoubtedly powerful techniques, they can only generate loss-of-function phenotypes. Classical forward genetics, on the other hand, has the ability to generate different mutant phenotypes depending on how it alters genetic loci, e.g. single nucleotide changes can result in more subtle phenotypes than deletions, and gain-of-function alleles can reveal the function of genes with redundant function.

Expression profiling

Gene expression profiling using microarray technology is now a standard method for gaining functional information for unknown genes and to help understand aspects of gene regulation. One of the most widely used microarray tools for *C. elegans* is the resource provided by Stuart Kim’s lab in UC Stanford, USA. RNA samples may be sent for analysis against microarrays produced in-house and the results are stored on their local database (http://genome-www5.stanford.edu/MicroArray/SMD/). Initial studies were performed on a microarray comprising 11 990 genes (155 experiments). More recently, a ‘whole genome’ microarray has been produced containing ~1 kb fragments from 17 871 genes (Jiang et al., 2001). To date, more than 1000 experiments involving many growth conditions, developmental stages and varieties of mutants have been performed. The availability of a central database resource enables global analyses of all the experiments, revealing an expression landscape in which there are mountains of genes with related expression profiles (Kim et al., 2001).

An alternative commercial *C. elegans* microarray is available from Affymetrix (http://www.affymetrix.com/products/index.affx). Many other companies also provide the option to construct custom arrays from either PCR products or synthesized oligonucleotides. Given the almost weekly changes in annotation to *C. elegans* genes and the low relative cost of manufacturing new microarrays, projects wishing to use large numbers of microarrays may wish to consider this latter option.

Recently, the filarial genome project has begun to construct a microarray containing 4000 *B. malayi* transcripts selected from EST datasets (Steve Williams, Smith College, MA, personal communication). Microarrays are expected to be available from March this year and will provide an exciting source of new data on gene expression in this medically important organism.

Aside from microarray technology, two groups are analysing gene expression with reporter constructs and *in situ* hybridization. Ian Hope’s group has used a promoter trapping approach to assess gene expression (Hope et al., 1998). Profiles of 342 genes are available on their website (http://bgypc086.leeds.ac.uk/). Yuji Kohara and colleagues are hybridizing a non-redundant set of cDNA clones from their EST sequencing project. Images are available on their website (http://nematode.lab.nig.ac.jp/db/).

Interaction screens

A genome-wide yeast two-hybrid analysis aims to eventually provide a complete catalogue of protein–protein interactions (http://vidal.dfci.harvard.edu/interactome.htm). Initially the feasibility of such a high-throughput technique was tested by fishing for interactions within a set of 27 genes involved in vulval development (Walhout et al., 2000). Eleven interactions between these genes had already been reported in the literature. By pair-wise matching each protein, six of these 11 interactions were detected. The failure to detect the remaining interactions is probably due to the inherent limitation of the two-hybrid assay (weak interactions may not be detected) or the physiology of the yeast cell (important post-translational modifications may not be performed correctly). Further interactions were sought by using the 27 proteins to screen a worm cDNA library, identifying a total of 148 interaction partners, of which only 15 had previously been described. Further work successfully identified the full complement of interacting partners in the proteasome (Davy et al., 2001). The large amount of genome-wide expression, RNAi and interaction data is now beginning to be correlated (Walhout et al., 2002). Finding correlates between these datasets helps validate the interaction data, e.g., in the yeast *Saccharomyces cerevisiae* potential interaction partners are more likely
to be co-expressed, and their knockdown phenotypes are often the same (Ge et al., 2001). Conversely, clustering of genes by their phenotype (Piano et al., 2002) or expression data (Roy et al., 2002) can be used to hypothesize on the function of novel proteins within clusters where many genes have already been ascribed biological functions. These studies have also shown that clustered genes are found within groups along chromosomes (Roy et al., 2002).

Transgenesis in parasites

Much has been learned about basic nematode biology, and of some drug targets that have close counterparts in C. elegans. To understand the function of the many parasite-specific gene products, techniques developed by the C. elegans community are starting to be adapted in parasite studies (Hashmi et al., 2001)

Several attempts at transgenesis in parasites have met with variable success (Davis et al., 1999; Hashmi et al., 1995; Jackstadt et al., 1999; Lok and Massey, 2002). Microinjection and ballistic DNA transfer into several species have demonstrated promoter activity and protein production in injected parasites, but rarely in their progeny. Often transgene activity can only be seen around the site where the DNA was introduced, e.g. in Strongyloides stercoralis, a parasite of the human gut. injection of an actin–green fluorescent protein (GFP) construct gave rise to fluorescence only in gonadal tissues and released embryos (Lok and Massey, 2002). All of the GFP-positive embryos failed to hatch. Although many nematodes will prove refractory to microinjection techniques as developed for C. elegans, these techniques will continue to be developed and could provide valuable tools for parasite research in the future.

Parasite genes in C. elegans

Promoter regions of parasitic genes have been shown to be active in C. elegans, but correct timing or location is not always seen (Britton et al., 1999; Gomez-Escobar et al., 2002). Orthology between Haemonchus contortus and C. elegans cysteine protease genes was suggested in studies that showed that the Haemonchus gene, under control of the C. elegans promoter, could functionally rescue a C. elegans mutant line (Britton and Murray, 2002). Expression of Onchocerca volvulus glutathione S-transferase (GST) in C. elegans demonstrated correct processing of the mRNA as well as signal peptide cleavage and glyclosylation of the mature protein (Krause et al., 2001), suggesting that this may be a useful system for producing parasite proteins.

RNAi in parasites

Although tremendously successful in C. elegans, the use of RNAi is only just starting to be addressed in parasites. The short period of protein knockdown may well limit its use in parasites, especially those with extended developmental cycles. To date, RNAi by soaking has proved successful in the gut nematode Nippostrongylus brasiliensis (Hussein et al., 2002), the filarial nematode B. malayi (Aboobaker and Blaxter, in preparation) and two plant parasitic nematodes Heterodera glycines and Globodera pallida (Urwin et al., 2002). Suppression in N. brasiliensis lasted for at least 6 days after exposure to dsRNA for 16 h. This length of knockdown is sufficient to assess the survival of treated worms once they are reintroduced into their animal host. A similar method was used to suppress a gene required to form the protective sheath around the first larval stage of B. malayi; this resulted in a drastic reduction in the number of released larvae (Aboobaker and Blaxter, in preparation).

Conclusions

This review summarizes the techniques currently used to assess gene function in C. elegans and other species of nematodes. Many of the techniques that have been developed are high-throughput, but can easily be adapted to analyse single genes, or groups of genes, in more detail. The data and reagents already available represent an advanced starting point for many projects. The C. elegans model is also a foundation for studying protein function in many medically and economically important nematode species, and has already helped define what genes may encode species-specific proteins in other nematodes. In the future, molecular and
structural studies will help define the function of many nematode proteins and their interactions within large biological networks.

**Web-based resources**

**WormBase**

http://www.wormbase.org/

The most comprehensive and up-to-date resource, covering many aspects of the *C. elegans* genome and its biology. The genome browser boasts a highly configurable interface, allowing the display of many features and annotations, including gene models, alignments to ESTs and the *C. briggsae* draft genome (and other sequences). In addition, WormBase includes a number of genome-wide data sets, including microarray expression data from a variety of life stages and conditions, RNAi experiments (16,757 genes), GFP expression data (covering 748 genes), single nucleotide polymorphisms (6,386 in total) and cell lineage information. WormBase is an ongoing project; new releases are published every ~2 weeks and will continue to develop and include new features as data becomes available.

The Wellcome Trust Sanger Institute and the Washington University Genome Sequencing Center *C. elegans* pages

http://www.sanger.ac.uk/Projects/C_elegans/

http://genome.wustl.edu/projects/c elegans/

These two centres were jointly responsible for generating the majority of the *C. elegans* genome sequence. The pages provide information on the *C. elegans* sequencing project, access to the sequence data and links to a number of related sites. They also both host a BLAST server allowing the user to search available *C. elegans* sequence data (http://www.sanger.ac.uk/Projects/C_elegans/blast_server.shtml and http://genome.wustl.edu/blast/briggsae-client.cgi).

**NEMBASE and NemaGene**

http://www.nematodes.org/nematodeESTs/nembase.html

http://www.nematode.net/NemaGene/index.php

These related databases are currently being developed as part of the parasitic nematode EST initiatives. For a number of species of nematodes, sequences are clustered (on the basis of sequence identity) into groups that putatively derive from the same gene, to form ‘partial genomes’. Web-based forms allow the identification of genes of interest via simple BLAST annotation or sequence similarity. Other features include the ability to define groups of genes on the basis of expression or similarity profiles (Parkinson and Blaxter, 2003).

**WorfDB**

http://worfdb.dfci.harvard.edu/

A database created to organize the data associated with the cloning of the complete set of predicted protein-encoding open reading frames (ORFs). It allows remote users to search for the availability and quality of cloned ORFs and displays the ORFs in context with the gene predictions.

**WormPD**

http://www.incyte.com/sequence/proteome/data bases/WormPD.shtml
A commercially produced venture which attempts to collate all currently published information on *C. elegans*, along with a number of automated analyses, (e.g. BLAST searches against a variety of protein databases) into one central resource.

NEXTDB: The Nematode Expression Pattern DataBase
http://nematode.lab.nig.ac.jp/

The Kohara lab has been constructing an expression pattern map of the *C. elegans* genome through EST analysis and systematic whole mount in situ hybridization. NEXTDB attempts to integrate all information from their expression pattern project.

The Interactome Project
http://vidal.dfci.harvard.edu/interactome.htm

These pages describe the initial findings from Dr Marc Vidal’s interactome project.

*C. elegans* transposon insertion project
http://www.cgmc.univ-lyon1.fr/

Dr Laurent Segalat’s lab is producing a transposon-based strategy to identify *C. elegans* mutants. These pages allow the user to search for a gene of interest and request any available mutants.

*C. elegans* gene knockout project
http://elegans.bcgsc.bc.ca/knockout.shtml

This is a worldwide consortium whose goal is to produce null alleles of all known genes in the *C. elegans* genome. You can submit requests of genes to target, and browse the lists of genes for which an allele has already been generated (currently ~600).

The Worm DNA Microarray Center
http://cmgm.stanford.edu/~kimlab/wmdirectorybig.html

The page provides information on Dr Stuart Kim’s large-scale microarray-based gene expression analysis project for *C. elegans*. It also contains information on how to take part and how to access the data (stored at the Stanford Microarray database).

Caenorhabditis Genetics Center
http://biosci.umn.edu/CGC/CGChomepage.htm

The center (situated in Minnesota) is responsible for collecting, maintaining and distributing stocks of *C. elegans*, maintaining a *C. elegans* bibliography, and publishing and distributing the *Worm Breeder’s Gazette*. A team in Oxford is responsible for coordinating genetic nomenclature and maintaining the *C. elegans* genetic map. A team in Dallas is responsible for maintaining the *C. elegans* web server (http://elegans.swmed.edu/).

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