The canine hookworm genome: analysis and classification of

*Ancylostoma caninum* survey sequences


**Abstract**

Hookworms infect nearly a billion people. The *Ancylostoma caninum* hookworm of canids is a model for studying human infections and information from its genome coupled with functional genomics and proteomics can accelerate progress towards hookworm control. As a step towards a full-scale *A. caninum* genome project, we generated 104,000 genome survey sequences (GSSs) and determined the genome size of the canine hookworm. GSSs assembled into 57.6 Mb of unique sequence from a genome that we estimate by flow cytometry of isolated nuclei to be 347±1.2 Mb, substantially larger than other Rhabditina species. Gene finding identified 5,538 genes in the GSS assembly, for a total of 9,113 non-redundant *A. caninum* genes when EST sequences are also considered. Functional classifications of many of the 70% of genes with homology to genes in other species are provided based on Gene Ontology and KEGG associations and secreted and membrane-bound proteins are also identified.

**Keywords**

hookworm; *Ancylostoma caninum*; genome survey sequences; expressed sequence tags; genome; comparative genomics

Hookworms are parasitic nematodes that live in the host small intestine and affect mammals including humans, dogs, and cats. *Ancylostoma duodenale* and *Necator americanus* infect close to a billion people [1] causing anemia and malnutrition, and also diminished physical and cognitive development in children. *A. caninum* is a hookworm species that infects canids and is commonly used as a model for studying human infections. Hookworm infections are usually
treated with anthelmintic drugs [2]. However, recurrence of infections and lack of immunity has bolstered the need for vaccines to reduce infections.

Tremendous advances towards a hookworm vaccine have been made in the last decade. Ancylostoma Secreted Proteins, ASP-1 and ASP-2 [3,4] and several hookworm digestive proteases like aspartic haemoglobinase [5] are being studied as potential vaccines against hookworm infections. In addition, hookworm gene products may have application against other diseases. For instance, recombinant hookworm rNAPC2 has anti-coagulant properties and is being studied for the treatment of acute coronary syndrome in the U.S. [6]. A recent study has shown that burns treated with this protein heal more rapidly with reduced scar contracture [7].

Genomic information from hookworms can enhance development of new, safer and sustainable control strategies. Our earlier effort to explore the A. caninum genome involved an EST-based approach [8]. By sampling 9,331 ESTs from three stage-specific cDNA libraries we identified and analyzed 3,840 genes. In this study we sampled the A. caninum genome by whole-genome shotgun. While one would expect that the A. caninum genome size would be in the 53-59 MB range based on other clade V Strongyliida species including Haemonchus contortus [9], our recent genome size estimate using flow sorted nuclei surprisingly revealed a genome more than six times this size. Genome size was estimated by proidium iodide staining and flow cytometry of isolated nuclei following methods described in [10]. To prepare the nuclei, the A. caninum L3s were washed in cold Galbraith buffer pH 7.2 and then pipetted into a plastic petri dish along with 30-50 μl cold Galbraith buffer and chopped (50 times) with a fresh single edge razor blade. The chopped material was washed into one edge of the (tilted) petri dish using an additional 1 ml of cold Galbraith buffer (per liter: 4.26g MgCl2, 8.84 g sodium citrate, 4.2 g [N-morpholino] propane sulfonic acid, 1ml Triton X-100, 1 mg boiled ribonuclease A, pH 7.2-7.4 with 1M KOH) and this solution was pipetted into a 1.5 ml Dounce homogenizer. Standards (similarly chopped C. elegans, the head of a D. melanogaster (Iso-1) and/or D. virilis female) were added to the Dounce at this time and the final amount in the Dounce was adjusted to 1ml adding cold Galbraith buffer as necessary. Each sample, standard, and co-prepared sample was ground using 15 strokes of the “A” pestle at a rate of 3 strokes per two seconds. The ground solution was poured through 20 μm nylon filter into a microfuge tube, brought to a final volume of 1 ml using additional cold Galbraith buffer as needed and then maintained on ice. Propidium Iodide (50 μl of 1 mg/ml H2O) was added to a final concentration of 0.075 mM, the tube capped and inverted several times to mix, and then stored in the dark for 2-8 hours prior to running. Samples were run in a Beckman-Coulter Epics Elite Cytometer using 25 mW of 488 nm (blue) excitation. PI fluorescence was measured after passing the collected fluorescent output across a 610 nm long pass filter. Counting was activated by PI fluorescence (gating: Debris, partial nuclei and nuclei with adhering cytoplasmic “tags” were excluded from the final analysis on the basis of forward and side scatter parameters. Only those nuclei with the lowest level of scatter were counted, as these have been shown by sorting to be the intact, untagged nuclei). A total of five replicates with at least 6,000 scored nuclei in each replicate were used to estimate genome size. C. elegans (1C = 100 Mb), D. melanogaster (1C = 175 Mb) and D. virilis (1C = 333.5 Mb) were used as standards. In every replicate, A. caninum ran just above the Drosophila virilis 2C peak (Fig. 1). The number of nuclei isolated from individual A. caninum was modest (maximum 200). However, the relative mean position of these nuclei compared to the standard was very consistent (Fig 1). The average (± standard error) genome size of A. caninum was 347.2±1.2 Mb. According to our estimate we sampled the A. caninum genome to an anticipated ~17% coverage.

Genomic DNA from adult A. caninum derived from experimental infection (strain Baltimore was isolated by G.A. Schad [11] from natural infection in the Baltimore area in the 1960s) was randomly sheared, end-repaired and size fractioned to enrich for 2-4 kb fragments. A total of
104,000 genome survey sequences (GSSs) were generated (72.6 Mb) (http://genome.wustl.edu/platforms_index.cgi), 95% passed quality screening and were submitted to dbGSS division of GenBank. Eighty nine per cent of clones had sequences from both ends. The 94,602 GSSs were assembled using PCAP [12]. Thirty five percent of the sequences assembled in 14,208 contigs and subsequently in 12,430 supercontigs (largest supercontig 9,358 bp). Total length of the contigs was 14.5 Mb and the 60,172 singletons contributed to 43.1 Mb of additional unique sequence. The low redundancy in the GSSs is consistent with the larger genome size estimate and rules out the possibility of a ~60 Mb genome like the Strongylid *H. contortus*.

As a first step to explore the available *A. caninum* genome sequence we identified the repetitive elements and masked them prior to gene identification. We evaluated repeat content in the assembly and singletons by masking simple repeats, low-complexity repeats and repeats identified by generating a custom library of repeat sequences. The custom library was built using RECON [13] and default parameters. This library was screened for non-coding RNA and protein-coding genes using Rfam [14] and Non-Redundant GenBank (built 03/23/2005) respectively, yielding 1,141 repeat families (Suppl. File S1). RepeatMasker (RepeatMasker Open-3.0.) was used to estimate the percentage of repeats. The total repeat content is estimated to be around 26.9 % (*C. elegans* repeat content is 16.5%) out of which 0.4% are simple repeats, 0.2% are low complexity repeats, and 26.3% are repeats identified by the custom library (The list of the ten most abundant repeats is available as on line Suppl. Table S1a). The GC content of the repeats was 44.6%, similar to the genome GC (43.2%) but lower than the protein coding exons GC (47.3%). Twenty-six repeat families had hits to repeats from various organisms in RepBase [15] (1e-05 cut-off; Suppl. Table S1b). Further analysis, especially on its resident mobile genetic elements, identified a novel mariner-like element [16], a non-long terminal repeat (LTR) retrotransposons [17] and transib transposon [18].

The masked assembly was used to call genes. In total, 5,538 genes were identified through a 6-tier gene-calling pipeline [19]. In addition, of the 3,840 genes that were identified by our previous *A. caninum* EST approach [8] 3,589 (represented by 4,816 contig consensus sequences) were not incorporated in calling genes, and therefore added to the list of identified unique genes. Similarly, 258 out of 498 contigs from external EST projects ([20] and Datu B., Loukas A., and Gasser, R., personal communication), were also non-overlapping genes and added to the non-redundant list of genes, making the total number of identified genes 9,385. The 9,385 genes are likely an overestimate of gene discovery, as one gene could be represented by multiple non-overlapping EST clusters or gene fragments identified from the GSS assembly. Such ‘fragmentation’ was estimated at 20% using *C. elegans* as a reference genome. As expected the fragmentation of the GSS derived genes was higher than the EST derived genes (21% vs 4%), and the overall fragmentation of 20% partially explains the low overlap between our EST and GSS gene collections. While the gene calling pipeline provides us with translations, we used prot4EST, a translational prediction pipeline optimized for EST datasets [21], to generate protein prediction for the EST clusters. The GFF and fasta files for the final set of 9,113 confident translations used in our analysis can be found at http://www.nematode.net [22]. Our approach indicates that the moderate number of ESTs and low-coverage obtained by GSS approach is a powerful combination which could allow an initial cost-effective identification of genes in neglected genomes. By this approach we identified 62 out of 77 *A. caninum* proteins previously deposited in the non-redundant GenBank (cut-off 1e-30) and increased the number of available genes from this species greater than 100-fold. The data contained 2.5 Mb of sequence from predicted genes, of which 1.9 Mb was exonic sequence. Direct comparison of the *A. caninum* genes to the homologous *C. elegans* counterparts indicated a smaller median exon size in *A. caninum* than *C. elegans* (128 bp vs. 174bp), and very similar median intron size of 72bp vs. 76bp (Suppl. Table S2). However, exons per genes and length of genes cannot be adequately estimated due to the fragmented...
nature of the assembly. At this coverage of the genome, it is premature to determine whether *A. caninum*’s larger genome will also contain an increase in the number or size of protein-coding genes compared to *C. elegans*. The GC content for *A. caninum* differed from *C. elegans* for the genome as a whole (43.2% for *A. caninum* versus 35.4% for *C. elegans*) and for protein coding exons (47.3% versus 42.7%).

Homology search of the 9,113 *A. caninum* genes with WU-BLAST (www://blast.wustl.edu) versus three specific phylogenetic databases (Fig. 2) revealed that 70% (6,335/9,113) had homologs among known and predicted proteins from other species. The majority of those with homology (63% or 3,996/6,335) matched all three databases. Of all proteins with homology, 92.5% shared similarity to *Caenorhabditis elegans* and/or *C. briggsae*, a higher percentage than the 80% match rate obtained using EST clusters alone [8]. Reasons for the increase in observed homology include the addition of non-abundant genes sampled by the GSS method, increased sequence length and quality contributing towards more significant hits, and not the effect of increased database size on score cut-off. This was confirmed by BLAST searching the EST clusters from Mitreva et al., [8] against the newly built databases used for this study (data not shown). The 7% of genes with homology only to *Caenorhabditis* species genes are potential nematode Clade V lineage-specific genes. Furthermore, 59% (3,330/5,641) of the hits to *C. elegans* were best reciprocal hits, therefore putative orthologs between *A. caninum* and *C. elegans*. RNA interference (RNAi) has not been reported in hookworms, therefore extrapolation from the *C. elegans* orthologs/homologs with observed phenotype by RNAi can be very informative for functional analysis of the orthologous counterparts in *A. caninum*. Of all orthologous *C. elegans* genes, 98% (3,253/3,330) had available RNAi information (RNAi data used from Wormpep v. 156), and of these 46% have observable phenotypes by RNAi knockdown. Of the genes with observable phenotypes, 60% (897/1,494) had severe phenotypes including embryonic, larval, or adult lethal, sterile, sterile progeny, and larval or adult growth arrest. A list of all RNAi information for *C. elegans* genes with *A. caninum* homolog is available as online Suppl. Table S3.

To functionally classify and categorize the genes, homologies to Kyoto Encyclopedia of Genes and Genomes (KEGG; [23]) and Interpro [24] database members were analyzed. The KEGG database contains information on metabolic pathways and interactions. *A. caninum* genes were mapped to the metabolic pathways using the highest-scoring WU-BLAST match and corresponding enzyme commission (EC) numbers of the homologous KEGG representative. Twenty-four percent of the *A. caninum* genes were mapped to 124 KEGG metabolic pathways classified into 11 major metabolic categories. Relative to mappings from the complete *C. elegans* genome, 70% of potential mappings were represented by *A. caninum* sequences. Biosynthesis of Secondary Metabolites and Energy metabolism were among the better represented categories using *C. elegans* as a reference (Table 1a). The complete listing of all mapping is available as online Suppl. Table S4. Metabolic pathways reconstruction is a productive approach for identifying novel anthelmintic drug targets. To facilitate this research, we have made the *A. caninum* KEGG associations to the 132 represented metabolic pathways graphically viewable on our web site (www.nematode.net; [22]). The viewer provides associations to specific enzyme commission (EC) numbers, strength of the match and associated KO identifiers. The *A. caninum* KEGG viewer coupled with our viewer of *C. elegans* metabolic mappings with incorporated RNAi information (http://www.nematode.net/KEGScan/cgi-bin/KEGScan_hit_distribution.cgi?species_selection=Caenorhabditis%20elegans) enables identification of potential ‘chokepoints’ in biochemical pathways and corresponding phenotypes. The combination of information from multiple sources can bolster the case for selection of new anti parasitic drug targets.

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As an alternative method for categorizing predicted proteins we used Interpro [25], an integrated database of known protein domains from well characterized data sources, to identify signatures/domains in our sequences. InterproScan [26] was used to locate homology to the Interpro database. 73% of the A. caninum genes mapped to one or more of 1,826 unique Interpro domains. The protein kinase (IPR000719), Protein kinase-like (IPR011009), Transposase, type 1 (IPR001888) and allergen V5/Tpx-1 related domains (IPR001283) were among the most frequently identified (Table 1b). Intriguingly, there were no C. elegans genes mapped to the IPR001888. Blasting of the 65 A. caninum genes against the C. elegans proteome did not yield any hits (cutoff 1e-10). We also looked at the Pfam entries for PF01359 (seed and full alignments)[27] and the corresponding Interpro id IPR001888 and we found 4 C. elegans proteins included in this entry. However all of these proteins were superseded or retired from Wormbase. Among the 65 A. caninum genes mapped to this IPR was the novel mariner-like element, bandit-transposon, that we characterized from the genome of A. caninum [16].

The phylogenetic analysis of Tc1/mariner superfamily of transposons indicated that the closest relative to bandit was the human HSmar1 rather than nematodes or arthropods originated transposons.

Furthermore, based on A. caninum InterPro protein domain matches, 32% of the gene products mapped to one or more organizing principles of the Gene Ontology (GO; [28]), representing 789 unique GO identifiers. Among the most common GO categories were ATP binding (361 genes mapped to GO:0005524), membrane (355 to GO:0016020), and intracellular (213 to GO:0005622). The GO associations are available on line through the Amigo viewer - (http://www.nematode.net/cgi-bin/amigo/go_AC/go.cgi).

Most of the nematode vaccine targets studied to date are excretory/secretory (ES) or intestinal antigens that are secreted or membrane-bound. Many chemotherapeutic targets are receptors or channels also found in membranes. We processed the A. caninum sequences through the Phobius server [29] which predicts secreted proteins (SP) and transmembranes containing domains (TM) by comparing the Hidden Markov Models of different sequence regions. There were 490 genes with predicted SPs for secretion and 1,437 with predicted transmembrane domains. Of the SP, 17% had associations to 48 unique IPR entries and 25% of the TM to 206 unique IPR entries (on line Suppl. Table S5).

The generation and analysis presented here is a valuable addition to resources for the study of hookworms. The A. caninum sequencing data have been submitted to public databases and the functional classifications and sequence characterizations are available on line, and are therefore accessible to researchers working on hookworms and other parasites. The generated data can also serve as a resource for more complete microarrays, RT-PCR, RNA interference and proteomic experiments and analysis. Such studies will aid in the identification of genes involved in host recognition, infection, migration and immune evasion as well as the characterization of targets for vaccines and anthelmintic drugs.

Furthermore, the importance of generating as complete a genome sequence as possible from both Necator and Ancylostoma species is recognized. Complete genome sequences from these species will serve as ongoing references for improved methods of parasite control, not only for drug and vaccine target identification, but also for development of diagnostic tools, drug resistance monitoring, vaccine response tracking, and parasite population surveys. Therefore, among several Strongylida species proposed for genome sequencing in 2006, N. americanus was considered because it is the most prevalent hookworm species infecting humans, and therefore most important from a public health standpoint. A. caninum was recommended as the Ancylostoma species to be sequenced based on its use as a model and the preliminary sequencing data (ESTs and GSSs generated here and in [8]). This genus is made up of very closely related species, which should make comparisons between them relatively easy. In
October 2006, the National Human Genome Research Institute announced that the Large-Scale Sequencing Research Network received financial support for sequencing the two hookworm genomes (http://www.genome.gov/10002154). We anticipate that complete annotated genomes for these species are approximately 3-4 years away. Based on the size estimate of the *A. caninum* genome reported here, nearly 2.7 million ABI3730 reads (average length of 800 bp) would be required for 6X coverage of the genome. The greater genome size increases the need for a physical map to enable sequence assembly. The larger than expected *A. caninum* genome size together with changing sequencing technology will also lead to a reconsideration of sequencing strategy. For example, one such ‘massively parallel’ sequencing platform is the FLX sequencer from 454 Life Sciences. This is a sequencing system that offers a 100-fold increase in throughput over the current state-of-the-art Sanger sequencing technology on capillary electrophoresis instruments [30]. The apparatus uses a novel fiber-optic slide (PicoTiterPlates) of individual 40 μm wells in which beads containing individual DNA fragments amplified by an emulsion PCR step are subjected to sequencing by synthesis using a pyrosequencing protocol optimized for solid support and picoliter-scale volumes. The FLX Genome Sequencer is capable of generating up to 100 Mb of data in a single run (7 hours) as 200-210 bp reads. No cloning is involved in the sample preparation; therefore no cloning bias can be introduced.

The extended genomic studies can offer a growing and fundamental base of information, which when coupled with downstream functional genomics and proteomics, could shorten the route towards developing more efficient and sustainable control programs.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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**References**


10. Bennett MD, Leitch JJ, Price HJ, Johnston JS. Comparisons with Caenorhabditis (~100 Mb) and Drosophila (~175 Mb) using flow cytometry show genome size in Arabidopsis to be ~157 Mb and thus ~25 % larger than the Arabidopsis initiative estimate of ~125 Mb. Ann Botany 2003;547–557. [PubMed: 12646499]


**Abbreviations**

**BLAST**  
basic local alignment search tool

**EST**  
expressed sequence tags

**GSS**  
genome survey sequence

**GO**  
gene ontology

**KEGG**  
Kyoto encyclopedia of genes and genomes
Figure 1.
The number of nuclei scored at differing levels of red fluorescence. The red fluorescence corresponds to binding of propidium iodide to the DNA of 2C nuclei in: A. Co-prepared heads of female *D. melanogaster* Iso-1 (1C = 175 Mb) and female *D. virilis* (1C = 333.5 Mb); B. Co-prepared heads of female *D. melanogaster* and *D. virilis* with *A. caninum*. C. A single female *A. caninum* (1C = 347.2 Mb). Small differences in PMT voltage shift means to slightly higher or lower channels in the three histograms. The DNA estimate is based on the ratio of co-prepared sample and standard as in 1B and 1C.
Figure 2.
Distribution of *Ancylostoma caninum* BLAST matches by database. *Caenorhabditis* spp., represents *C. elegans*, *C. briggsae* and *C. remanei*; Other nematodes, all nematode beyond *Caenorhabditis* and *Ancylostoma*; and Non-nematoda, all species beyond nematodes.
Databases were built 06/09/06, e-value scores $\leq 1e^{-05}$ are considered.
### Table 1a

KEGG biochemical pathway category mappings\(^a\) for *A. caninum* and *C. elegans* genes

<table>
<thead>
<tr>
<th>KEGG PATHWAY</th>
<th><em>A. caninum</em></th>
<th></th>
<th><em>C. elegans</em></th>
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<td>Genes</td>
<td>Enzymes</td>
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<td>Glycan Biosynthesis and Metabolism</td>
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<tr>
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<td>89</td>
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<tr>
<td>Xenobiotics Biodegradation and Metabolism</td>
<td>154</td>
<td>37</td>
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\(^a\) WU-BLAST (www://blast.wustl.edu) with top hits of each gene; KEGG v. 40 was used.
### Table 1b

The top 10 most abundant Interpro identifiers of *A. caninum* in *C. elegans*

<table>
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<th>Interpro</th>
<th>Description</th>
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<th>C. elegans</th>
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<td>IPR000719</td>
<td>Protein kinase</td>
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<tr>
<td>IPR011009</td>
<td>Protein kinase-like</td>
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<td>Transposase, type 1</td>
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