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# The transcription start site landscape of *C. elegans*

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More than half of *Caenorhabditis elegans* pre-mRNAs lose their original 5' ends in a process termed “trans-splicing” in which the RNA extending from the transcription start site (TSS) to the site of trans-splicing of the primary transcript, termed the “outtron,” is replaced with a 22-nt spliced leader. This complicates the mapping of TSSs, leading to a lack of available TSS mapping data for these genes. We used growth at low temperature and nuclear isolation to enrich for transcripts still containing outtrons, applying a modified SAGE capture procedure and high-throughput sequencing to characterize 5' termini in this transcript population. We report from this data both a landscape of 5'-end utilization for *C. elegans* and a representative collection of TSSs for 7351 trans-spliced genes. TSS distributions for individual genes were often dispersed, with a greater average number of TSSs for trans-spliced genes, suggesting that trans-splicing may remove selective pressure for a single TSS. Upstream of newly defined TSSs, we observed well-known motifs (including TATAA-box and SPI) as well as novel motifs. Several of these motifs showed association with tissue-specific expression and/or conservation among six worm species. Comparing TSS features between trans-spliced and non-trans-spliced genes, we found stronger signals among outtron TSSs for preferentially positioning of flanking nucleosomes and for downstream Pol II enrichment. Our data provide an enabling resource for both experimental and theoretical analysis of gene structure and function in *C. elegans*.

[Supplemental material is available for this article.]

In most organisms, the locations of transcription start sites (TSSs) can be determined by establishing the sequences of mRNA 5' ends. For a subset of single-cell eukaryotes and animals, however, a processing event known as “trans-splicing” obscures the locations of the TSSs (Sutton and Boothroyd 1986; Krause and Hirsh 1987; Lasda and Blumenthal 2011). Trans-splicing is an efficient process that results in removal of the 5' end of the pre-mRNA, replacing it with a short 5' leader that is then retained on the mature mRNA. The removed sequence (between the TSS and the first active 3' splice site in the newly transcribed mRNA precursor) is called the “outtron” (Conrad et al. 1991). Trans-splicing is a spliceosome-catalyzed process that can be thought of as a surrogate use of a mobile 5' splice site contained on a short (usually ~100 nt) RNA called an SL RNA. SL RNAs are present in the nucleus in the form of Sm protein-bound small nuclear ribonucleoprotein particles (snRNPs), where the 5' 22 nt forms the SL exon, with the 3' portion serving as “intron.”

*Caenorhabditis elegans* has been a valuable model system for studying a variety of aspects of gene expression (including trans-splicing) (Krause and Hirsh 1987; Lasda and Blumenthal 2011). Approximately 70% of *C. elegans* genes are subject to trans-splicing (Allen et al. 2011; Lasda and Blumenthal 2011). *C. elegans* has two types of trans-splicing, each with a distinct splice leader (SL1 and

SL2) (Spieth et al. 1993). SL1 is the more common splice leader, with 55% of *C. elegans* genes spliced to SL1 (Allen et al. 2011). SL1 trans-spliced genes are either singly transcribed genes or the first genes in cotranscribed operons. The SL1 trans-spliced genes are thus those for which the outtron 5' ends correspond to TSS positions. About 15% of all genes are trans-spliced by the spliced leader SL2 or one of its close relatives (Allen et al. 2011). SL2 trans-spliced genes are always downstream in operons, so they generally do not have TSSs present just 5' of the gene body. Due to the rapidity of trans-splicing and the overwhelming predominance of post-trans-splicing forms among stable cytoplasmic RNAs, the annotation of the worm genome has lacked a global description of TSSs, confounding analysis of cis-regulatory sequences, chromatin configurations around promoters, and identification of specialized features such as bidirectional promoters.

Here we perform a 5'-targeted analysis on nuclear RNA from *C. elegans* grown under conditions that slow the processing of outtrons, enabling the mapping of a large number of TSS for both SL1 trans-spliced and non-trans-spliced genes and a consequent picture of transcriptional initiation repertoire in this system.

## Results

### Capture of 5'-end RNA reads from nuclei

Animals grown at the low end of the normal temperature range were used to isolate potential pre-trans-splicing intermediates. The choice of lower growth temperatures to support identification of pre-spliced intermediates follows observations that (1) *C. elegans* strains partially defective in key trans-splicing RNP factors show a cold sensitivity in organismal phenotype (MacMorris et al. 2007), and (2) splicing can be mechanistically sensitive to cold

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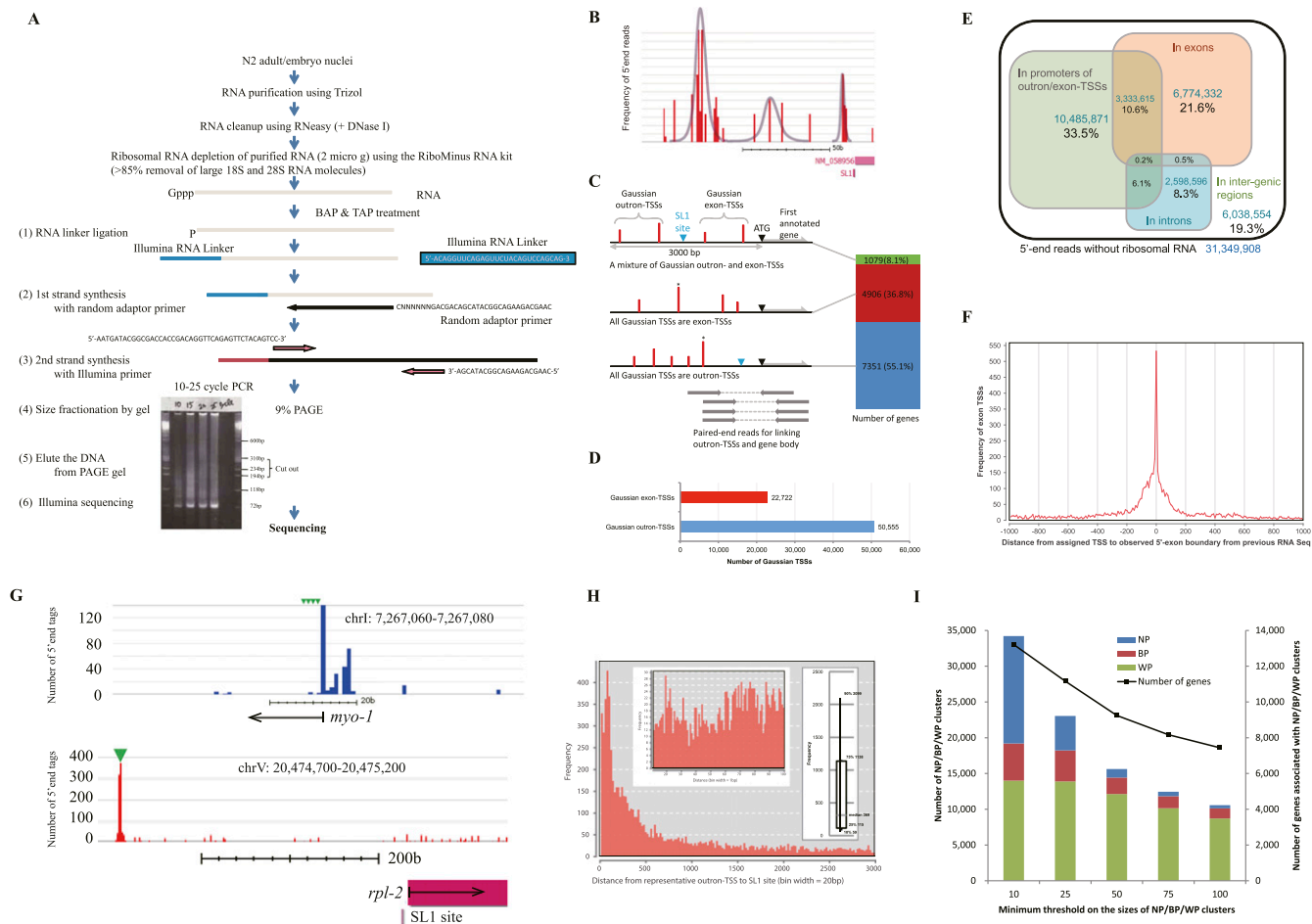
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temperatures as a result of numerous base-pairing interactions that must be remodeled during the process (Strauss and Guthrie 1991).

To identify TSSs that are processed before *trans*-splicing or are free of *trans*-spliced leaders, we grew populations of animals at 16°C, isolated nuclei from embryos and whole adult tissues, and extracted total nuclear RNA. To minimize contaminating ribosomal RNA (rRNA) molecules, we used hybrid subtraction-based enrichment of RNA with the RiboMinus RNA Kit before sequencing (Ruan et al. 2004; Chen and Duan 2011). We then used a 5'-SAGE method (Fig. 1A) to capture 5' ends, including DNase treatment and a phosphatase treatment preceding cap removal and

linker ligation that strongly enriches for RNA molecules where a 5' phosphate is only revealed following decapping (protocol described in Hashimoto et al. 2009). Illumina sequencing was then carried out, resulting in approximately 119 million and 76 million 76-bp single-end reads collected from the nuclei of embryo and adult samples, respectively (see Methods). These reads were aligned to the *C. elegans* genome WS220 (Hillier et al. 2009; Harris et al. 2010) using BWA (Li and Durbin 2009) (see Methods). From this analysis, approximately 55.6 million and 30.4 million reads, respectively, were anchored to unique positions (Table 1), with the approximately 55.6 million reads from embryo samples used as the



**Figure 1.** Identification of TSSs from alignments of 5'-end reads. (A) Process for 5'-end library construction for Illumina GA sequencing. See details in Methods. (B) Frequency distribution of reads aligned to genomic locations upstream of NM\_058956 (phosphatase). Three peaks were calculated as the best-fitting set of Gaussian distributions according to Bayesian information criteria. (C) Three classes of genes with different patterns of Gaussian TSS distributions in the regions of 3000 bp upstream of the translation initiation sites. (Top figure) Case in which both Gaussian outron/exon-TSSs occur because of the presence of an SL1 site between the two classes of Gaussian TSSs. (Middle and bottom figures) Cases in which all Gaussian TSSs are exon-TSSs and outron-TSSs, respectively. (Vertical bars with asterisk) Representative TSSs with the maximum number of aligned 5'-end tags. (Right bars) The numbers of genes in the three classes. As shown in the bottom, alignments of paired-end reads were useful in linking representative TSSs to their gene bodies over SL1 sites. (D) Frequency of Gaussian exon/outron-TSSs. (E) 5'-End reads excluding rRNA reads were categorized into the four groups of reads that were mapped onto promoters of outron/exon-TSSs, exons, introns, and intergenic regions, respectively. The Venn diagram illustrates the relationship among the four groups. (F) Frequency distribution of distances between 5'-exon boundaries from WS220 (Hillier et al. 2009; Harris et al. 2010) and proximal exon-TSSs identified in our data. (G) Examples of 5' capture data for genes that have been previous subjects of TSS mapping. (Above) Exon 5' end for the gene *myo-1* with original 5' region having four candidate 5' ends (from S1 nuclease mapping with accuracy to within a few bp) (Okkema et al. 1993) marked with the green arrows. (Below) Outron 5' region for *rpl-2* with original 5' end (from 5' RACE mapping with accuracy to within 1–2 bp) (Sleumer et al. 2012) highlighted by the green arrow. (H) Frequency distribution of the distances between SL1 sites and most abundant representative outron-TSSs supported by 10 or more single-end reads. (Left inset) This magnifies the region ranging from 10 to 100 bp; (right inset) the 90th, 75th, 25th, and 10th percentiles and the median of outron lengths. (I) Numbers of TSS clusters categorized into NP, BP, and WP types in terms of five values of the minimum threshold on the size of TSS clusters. Because multiple TSS clusters may be associated with each gene, the second vertical axis shows the numbers of genes involved. Observe that TSS clusters of WP type are dominant for threshold  $\geq 25$ .

**Table 1.** Statistics of sequenced 5'-end reads from nuclei of embryo and adult samples and their alignment to the worm genome (WS220)

	Number of reads	Number mapped	Number mapped uniquely	Number mapped redundantly
Embryo (single-end)	119,198,792	90,575,557	55,591,595	34,967,142
Adult (single-end)	76,731,798	49,634,285	30,453,251	19,181,034

primary data set for initial TSS identification. As expected from the mass of ribosomal RNA present even after (incomplete) hybrid subtraction, the pool of sequenced reads still included approximately 24.2 million 5'-end rRNA reads. We removed these after their alignment to the genome, leaving a total of 31.4 million reads from the embryonic library.

Selecting representative TSSs is not simple because the distribution of TSSs is not necessarily sharp and unique but is often broad and/or multi-modal (Suzuki et al. 2001; Schug et al. 2005; Carninci et al. 2006; Frith et al. 2006; Saxonov et al. 2006; Zhu et al. 2008; Rach et al. 2011), as illustrated in Figure 1B. We take several approaches in working with the multi-modal character of TSS distributions. These include both approaches based on calling one or a few most prominent starts for each gene as representative and approaches based on a base-by-base distribution of TSS frequencies covering an extensive upstream region of every gene.

In calling individual reference starts based on many noisy peaks present in the frequency distribution, one approach is to model start site regions as a mixture of Gaussian distributions; such an approach has been widely used in the recent analysis of TSS distributions (Boyle et al. 2008; Ni et al. 2010; Rach et al. 2011). We calculated a series of best-fitting Gaussian distributions using Bayesian information criteria (see Fig. 1B). We then quantified the expression level of the resulting Gaussian-based TSS models by the number of reads in the corresponding Gaussian window. To eliminate noise in genome-wide analysis, we used 73,277 Gaussian TSS peaks that were supported by 10 or more reads (see their positions in Supplemental Table S1). In addition, we used a noise-filtering method using feature density estimator F-Seq (Boyle et al. 2008) to exclude TSS peaks whose F-Seq values are less than 4 standard deviations above the mean of the background (F-Seq has been widely used in other studies on TSS distributions) (Ni et al. 2010; Rach et al. 2011).

The identified "peaks" from the above analysis ranged from cases showing a small number of starts in a tight cluster to those spread over hundreds of bases of DNA. While the unique-site peaks are readily recognizable as a defined entity, parsing of more disperse TSS regions into individual peaks is by nature dependent on the algorithm and parameters. With >80% of the observed Gaussian peaks showing a dispersion (standard deviation) of <20 bp, the most challenging cases are in the minority (for reference, Supplemental Table S2 shows a list of Gaussian peaks parsed from the data with a maximal standard deviation parameter of 20 bp, and Supplemental Fig. S1 presents a statistical analysis of Supplemental Table S2).

We defined outtron-TSSs as candidate TSSs by two criteria: (1) They had previously mapped downstream SL1 sites (Allen et al. 2011); and (2) the TSS occurred in a window from 10 bp to 3000 bp upstream of the translation initiation site of the first downstream

annotated gene (illustrated in Fig. 1C). The 10-bp lower limit was chosen to avoid putative outtrons that would likely be too short to allow *trans*-splicing (also see below) as well as confounding instances resulting from short upstream near-matches to SL1 sequence; the 3000-bp upper limit was chosen based on general characteristics of *C. elegans* promoters (Hunt-Newbury et al. 2007). The term "outtron-TSS" is derived from the term "outtron" (Blumenthal 2005) indicative of the sequence between the natural transcription start site and spliced leader attachment site of a relevant gene (Blumenthal 2005). We defined exon-TSSs as candidate TSSs with an annotated coding region with no SL1 *trans*-splice sites that were not also intron 3' splice sites. (A low level of *trans*-splicing sometimes occurs at intron 3' splice sites [Allen et al. 2011].) Figure 1D indicates the incidence of all Gaussian outtron/exon-TSSs.

Among 13,336 genes having Gaussian TSS peaks in the region of length 3000 bp upstream of their translation initiation sites, 7351 genes (55.1%) had only Gaussian outtron-TSSs and were therefore defined as *trans*-spliced genes, and 4906 genes (36.8%) having only Gaussian exon-TSSs were categorized as non-*trans*-spliced genes (Fig. 1C). For the other 1079 genes (8.1%) that had both types of Gaussian TSSs in our data, we place the genes in neither category, avoiding for this analysis any decision of whether they were *trans*-spliced. For genes that could be defined unambiguously as *trans*-spliced or non-*trans*-spliced, we assigned gene-unique representative outtron/exon-TSSs with the maximum number of aligned reads. This allowed us to pursue further analysis in parallel with (1) all Gaussian peaks as TSSs ("The Gaussian Peak Collection"), and (2) representative gene-unique TSSs (one for each gene).

To assess the specificity of our 5'-SAGE method in detecting TSSs, we examined the balance between ends present within known RNAs and those in regions upstream of annotated genes, assigning the approximately 31.4 million embryonic alignments to one or more of the categories: promoters of outtron/exon-TSSs, exons, introns, and intergenic regions. The former three categories are overlapping as illustrated by the Venn diagram in Figure 1E. We compared our results with those by the CAGE method, another widely used method of monitoring 5' ends of RNA. The ratio of 5'-end tags mapped to promoter regions is 50.2%, while the ratio of 5'-end tags by the CAGE method in the human and mouse genomes was smaller than 30% (Carninci et al. 2006), indicating that the background signal of our 5'-SAGE method is similar to that of CAGE. Although the modest number of 5' ends present in annotated transcribed regions (21.6% of the total) are under some suspicion as potentially representing capture of fragmented mRNA precursors at positions other than the 5' end, the fact that the vast majority (78.4%) are in regions with no known abundant transcripts suggests that the latter RNAs are almost certainly TSSs. In particular, we stress that a situation in which 5' ends were captured at random from fragmented RNA present in the samples would invariably yield a dramatic enrichment for tags within regions annotated in other RNA sequencing projects. That this was not the case argues strongly for the enrichment in this analysis of bona fide capped 5' ends.

An independent evaluation of 5'-end mapping was possible for known exon-TSSs that are free from *trans*-splicing. We first evaluated the correspondence of 5' mapping for several *C. elegans* genes in which single-gene studies (generally individual primer extension reactions either with a sequencing endpoint or with a PCR/sequencing endpoint, or nuclease protection mapping) had defined an exon 5' end within a few nucleotides. Although not

large in number, such specific examples served as valuable benchmarks. For the set of genes evaluated (*unc-54*, *myo-1*, *myo-2*, *sqt-1*, *col-12*), the major 5' ends defined in earlier studies were within a few base pairs of prevalent 5' capture sites from this work (Krause and Hirsh 1987; Dibb et al. 1989; Park and Kramer 1990, 1994; Okkema et al. 1993). In addition to this individual gene analysis, we examined 5268 presumed non-*trans*-spliced first-exon 5' boundaries that were derived from whole RNA-seq data in WS220 (Hillier et al. 2009; Harris et al. 2010) and that reside  $\geq 100$  bp upstream of ATG sites of 1078 genes, analyzing the frequency distribution of distances between the known 5' ends and their proximal exon-TSSs in our data. This data set would be expected to be highly enriched for exon TSS positions. As would be expected from concordance of the two data sets, we observed a remarkable peak at zero in the distance distribution, confirming the significant concordance between known 5' ends of mRNAs and our exon-TSSs (Fig. 1F).

Existing outtron-TSS mapping studies are much rarer in *C. elegans*, as noted above, due to the challenges inherent in capturing a transient 5'-end sequence. We know of five genes that have been mapped at high precision through primer extension (*rol-6* and *col-13*) (Park and Kramer 1990, 1994), 5'-anchored PCR and sequencing (*rpl-2* and *ace-1*) (Culetto et al. 1999; Sleumer et al. 2012), or S1 nuclease protection (*ubq-1*). In our 5'-end repertoires, we see support for the mapped outtron-TSSs for *rpl-2* (Fig. 1G), *col-13*, and for two of three 5' ends mapped for *ace-1* outtron sequences. *Rol-6* expression is very low in the samples used for this work, presumably due to the precise staging of transcription. Thus, little or no signal is seen in this study for any sequence upstream of *rol-6*. For *ubq-1*, we observe a different 5' end from that mapped in the original study by S1 nuclease protection. (The prominent 5' end inferred from the original study was at  $-455$  nt. We observed a strong end at  $-294$ , in a region of the gene that was electrophoresed out of the pictured gel in the S1 mapping study. That study was also carried out with total RNA from heat-shocked embryos, while we used nuclear RNA from embryos grown at the lower end of the *C. elegans*' viable temperature range.) These comparisons, as well as comparisons with the regional outtron mapping of Morton and Blumenthal (2011), serve both to support the utility of the present data and to provide caution in any universal interpretation of data from a limited set of developmental timing and growth conditions. Even if stable mRNA is present in the sample, these assays will only provide a positive outtron signal based on recent transcription, specifically addressing the (likely transient) presence of the pre-*trans*-splicing nuclear precursor under the growth conditions of the assay.

To provide valid outtron-TSS candidates, we sought where possible to associate each with the body of the corresponding gene. Exon-TSS assignments have the advantage of potential validation through a continuous set of RNA-seq reads that go into the relevant mRNA. To further evaluate outtron-TSS assignments, we performed paired-end sequencing of the embryonic library, generating 42.2 million paired-end reads of 5'-end RNA fragments; 23.1 million pairs mapped to unique positions in the genome (Table 2). We then applied an analysis that checked whether the SL1 site in a gene was sandwiched by the aligned paired-end reads, providing support for the assignment of an outtron-TSS to a gene body associated with an SL1 site as shown in Figure 1C. One might be concerned with the difficulty of linking outtron-TSSs to SL1 sites because the distance between paired-end reads ( $\sim 250$  bp being the cDNA insert size in our experiment) could be smaller than the length of typical outtrons. Examining the frequency distribution of

**Table 2.** Statistics of sequenced paired-end reads for TSSs from embryo nuclei and their alignment to the worm genome

	Number of paired-end reads	Number mapped	Number mapped uniquely	Number mapped redundantly
Embryo (paired-end)	42,232,142	37,066,515	23,145,796	10,029,550

the distances between the SL1 sites and the most abundant representative outtron-TSSs (defined by single-end sequencing) shows that, although many TSSs were located at  $>250$  bp upstream of corresponding SL1 sites, 40.3% of 7351 TSSs were seen within 250 bp of SL1 sites (Fig. 1H), indicating that our paired-end sequencing data will be useful in linking many TSSs to associated gene bodies.

In Figure 1H, we find that lengths of outtrons, the region between the TSS and the *trans*-splice site, vary greatly, from a 90th percentile of 2099 to a 10th percentile of 59, with a median length of 369. Whereas the long outtrons are similar in length to long introns in worms, the putative outtrons defined by these data were in some cases much shorter than introns can be, consistent with the idea that the constraint on the lower limit of intron length results from a minimum distance between the two splice sites needed for spliceosome assembly. While the possibility of short outtrons is intriguing, we note that the type of data acquired in this work (a static picture of 5'-end structures) does not provide information about precursor-product relationships. Apparent short outtrons, based on the distance from the predominant TSS to the *trans*-splice site, may lead to misinterpretation. The true outtrons may derive from initiation events upstream. Indeed, we observed upstream TSSs in 118 (76%) among 154 genes for which the predominant TSS was within 20 bp upstream of the SL1 site (Supplemental Table S2). In the remaining 36 genes, because such outtrons may be processed efficiently, these upstream TSSs might either not appear in our data or appear as rarely used sites. Conversely, the outtron sites that appear in nuclear RNA would have been enriched for slower 5' processing, potentially enriching the signal for any short outtrons that are spliced inefficiently.

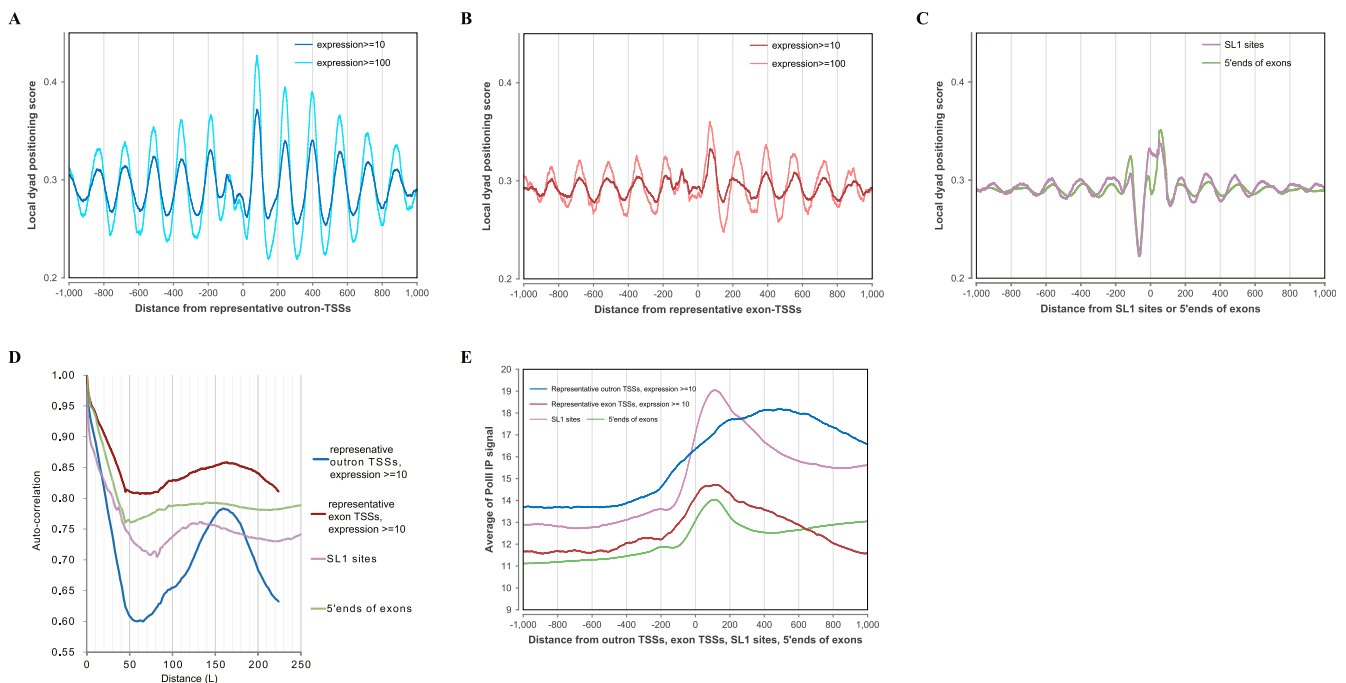
An intriguing diversity among initiation elements comes in the degree to which initiation is specified at a single position in the DNA sequence versus a range of positions. Classifiers for peak breadth have been applied extensively in such analyses, including previous studies on human, mouse, and fruit fly genomes: In each case, once a set of reference TSSs was determined, shapes of TSS distributions in promoters of individual reference TSSs would be assigned (Carninci et al. 2006; Frith et al. 2006; Ni et al. 2010; Rach et al. 2011). We classified *C. elegans* TSS distributions into three types: NP (Narrow with Peak), BP (Broad with Peak), and WP (Weak Peak) according to the definitions proposed by Ni et al. (2010). The precise category fractions varied somewhat depending on the minimum threshold on the size of the TSS cluster (e.g., 10, 25, 50, 75, and 100). For start sites with modest-to-high expression (thresholds of more than 25 reads observed) started, WP-type shapes were dominant (Fig. 1I). This observation is consistent with WP dominance in other studies of human, mouse, and fruit fly genomes (Carninci et al. 2006; Frith et al. 2006; Ni et al. 2010; Rach et al. 2011). TSS distribution of WP type frequently ranged over a long region (typically of length  $>500$  bp) and had multimodal peaks; for some analyses, we thus decompose each WP TSS distribution into Gaussian peaks to examine characteristics as noted below.

### Nucleosome and Pol II positioning around TSSs and SL1 sites

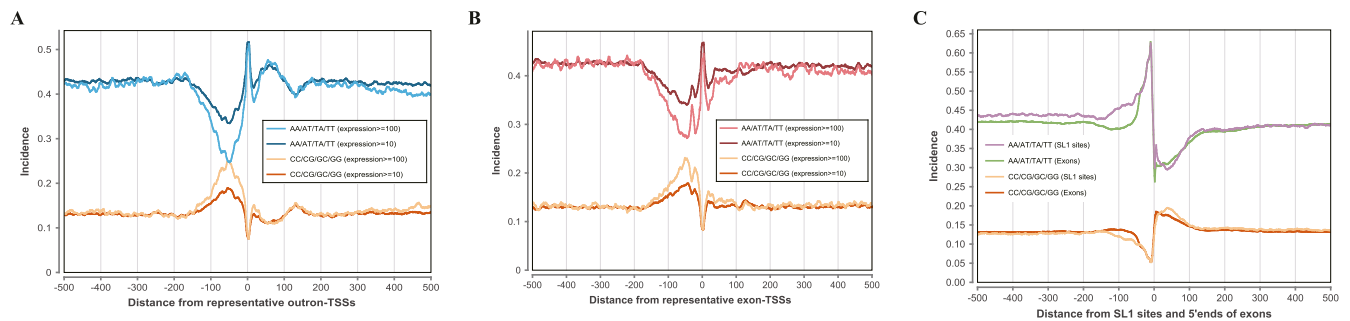
Some types of modified histones and RNA Pol II are known to accumulate at promoter regions (Whittle et al. 2008; Baugh et al. 2009), with information related to this accumulation having been used to approximate the locations of transcription initiation sites. To check whether the TSS collection defined in this study shares these characteristics with other species, we examined the publicly available data of nucleosome positioning estimated from MNase digestion of whole chromatin (Valouev et al. 2008) or of chromatin pre-selected for its association with the histone modification mark H3K4me2/3 (Gu and Fire 2010), also examining Pol II positioning (Gerstein et al. 2010) surrounding representative TSSs. Using either representative TSSs (Fig. 2A,B) or the full Gaussian peak collection (Supplemental Fig. S2A,B), we observed arrays of positioned and phased nucleosomes, with an apparent nucleosome-depleted region immediately upstream of TSSs. The upstream nucleosome depletion is a conserved feature that has also been reported in mammals and other eukaryotes (Mavrich et al. 2008; Kaplan et al. 2009; Sasaki et al. 2009; Weiner et al. 2010; Valouev et al. 2011). The characteristic features of nucleosome positioning relative to representative TSSs were also reconfirmed by comparison with H3K4me2/3 data (Supplemental Fig. S2D). The period of the nucleosome positioning around representative TSSs was estimated as  $\sim 160$  bp, as evaluated using an autocorrelation measure (Fig. 2D). The  $\sim 160$ -bp spacing around representative TSSs was smaller than the  $\sim 175$ -bp spacing in the entire genome (Valouev et al. 2008). Nucleosome positioning scores were more pronounced for

representative TSSs of higher expression (Fig. 2A,B). These observations indicate that transcription correlates with more consistent packing of nucleosomes. This observation is consistent with barrier nucleosome models (Mavrich et al. 2008) in which Pol II stalled at the TSSs potentially serves as a barrier that can facilitate packing of nucleosomes downstream from TSSs. Indeed, Figure 2E exhibits Pol II enrichment at  $\sim 170$  bp and  $\sim 100$  bp downstream from representative outtron/exon-TSSs, respectively. Interestingly, the distance between TSS and maximum Pol II enrichment differed between outtron-TSS sites ( $\sim 170$  bp) and exon-TSSs ( $\sim 100$  bp), respectively. Although the mechanistic basis for this difference was not clear, the existence of a difference suggested the possibility that the biological system would have the opportunity, if needed, to distinguish functionally between the two types of 5' end.

It has been shown that nucleosomes have a tendency to associate with DNA sequences exhibiting high G/C nucleotide and dinucleotide content, while being excluded from DNA sequences exhibiting high A/T nucleotide and dinucleotide content (Bernstein et al. 2004; Field et al. 2008; Hughes and Rando 2009; Segal and Widom 2009; Tillo et al. 2010). We therefore examined the correlation of this underlying nucleotide signal to nucleosome positioning around newly identified representative TSSs (Fig. 3A,B) and around the full collection of Gaussian peaks (Supplemental Fig. S2E,F); in both analyses, we observed an unexpected phenomenon, a significant peak of C/G rate and a remarkable valley of A/T rate around  $\sim 100$  bp upstream of outtron- and exon-TSSs, corresponding to a site where observed nucleosomes were depleted.



**Figure 2.** (A,B) The distributions of nucleosome positioning (local dyad positioning score) around representative outtron-TSSs (A) and exon-TSSs (B) of expression levels  $\geq 10$  and  $\geq 100$  (see the distribution around Gaussian TSSs in Supplemental Fig. S2A,B). From these comparisons, note that the periodic pattern of nucleosome positioning was of substantially greater amplitude for TSSs of greater expression. (C) Nucleosome positioning (local dyad positioning score) around SL1 sites and 5' ends of exons. We used all available SL1 sites without considering their expression levels. (D) Autocorrelation analysis: Let  $s(x)$  denote the local dyad positioning score. The autocorrelation defined by  $R(L) = \int_{-\infty}^{\infty} s(x+L)s(x)dx$  was calculated around representative outtron/exon-TSSs, SL1 sites, and 5' ends of exons, where  $L$  is displayed in the x-axis. Sharp peaks at  $\sim 160$  bp for outtron-TSSs and broad peaks surrounding  $\sim 160$  bp for exon-TSSs highlight that arrays of positioned and phased nucleosomes are more prominent around outtron-TSSs than around exon-TSSs. Broad peaks at  $\sim 140$  bp are also seen for SL1 sites and 5' ends of exons. (E) Pol II occupancy around representative outtron/exon-TSSs of expression level  $\geq 10$ , SL1 sites, and 5' ends of exons.



**Figure 3.** (A,B) A/T and C/G dinucleotide incidence rates around representative outtron-TSSs (A) and exon-TSSs (B) of expression levels  $\geq 10$  and  $\geq 100$  (see the distribution for Gaussian TSSs in Supplemental Fig. S2E,F). AT richness immediately upstream of outtron-TSSs is due to the requirement for SL1 splicing motif "TTTTAG." In both types of TSSs, a significant valley in the A/T rate and a significant peak in the C/G rate were observed  $\sim 100$  bp upstream of TSSs. (C) A/T and C/G dinucleotide incidence rates around SL1 sites and 5' ends of exons. In all graphs, a running average over a 20-bp window is shown for smoothing lines.

Nucleosome positioning became more prominent around outtron-TSSs than around exon-TSSs (Supplemental Fig. S2A,B), suggesting that a higher Pol II signal and a more pronounced array of positioned nucleosomes were associated with *trans*-splicing activity. We therefore examined nucleosome positioning around SL1 sites and found that nucleosome cores were enriched  $\sim 40$  bp downstream from SL1 sites (Fig. 2C). One possibility was that this enrichment might reflect a combination of the distance distribution between TSSs and SL1 sites and the phased array of nucleosomes downstream from TSSs; however, this hypothesis was not supported by our statistical analysis (Supplemental Fig. S2G,H).

As an alternative and more direct link, we considered the possibility that genomic signals for splicing machinery and nucleosome constraint might be associated. Such an association has been observed at *cis*-spliced sites (Kolasinska-Zwierz et al. 2009), hence an expectation that a disambiguation of TSSs and SL1 spliced sites might show such an effect. Consistent with such an expectation, nucleosome positioning around 5' ends of exons was quite close to that around SL1 sites (Fig. 2C). Furthermore, we also observed RNA Pol II enrichment at  $\sim 100$  bp downstream from SL1 sites and 5' ends of exons (Fig. 2E), and a strong correlation of A/T and C/G dinucleotide rates to the nucleosome enrichment downstream from SL1 sites and 5' ends of exons (Fig. 3C). These observations suggest an available confluence in biological regulation in which shared genomic signals and underlying dependencies on the underlying sequence can be used to coordinate nucleosome positioning, splicing, and polymerase initiation.

### Evolutionarily conserved sequence motifs around TSSs

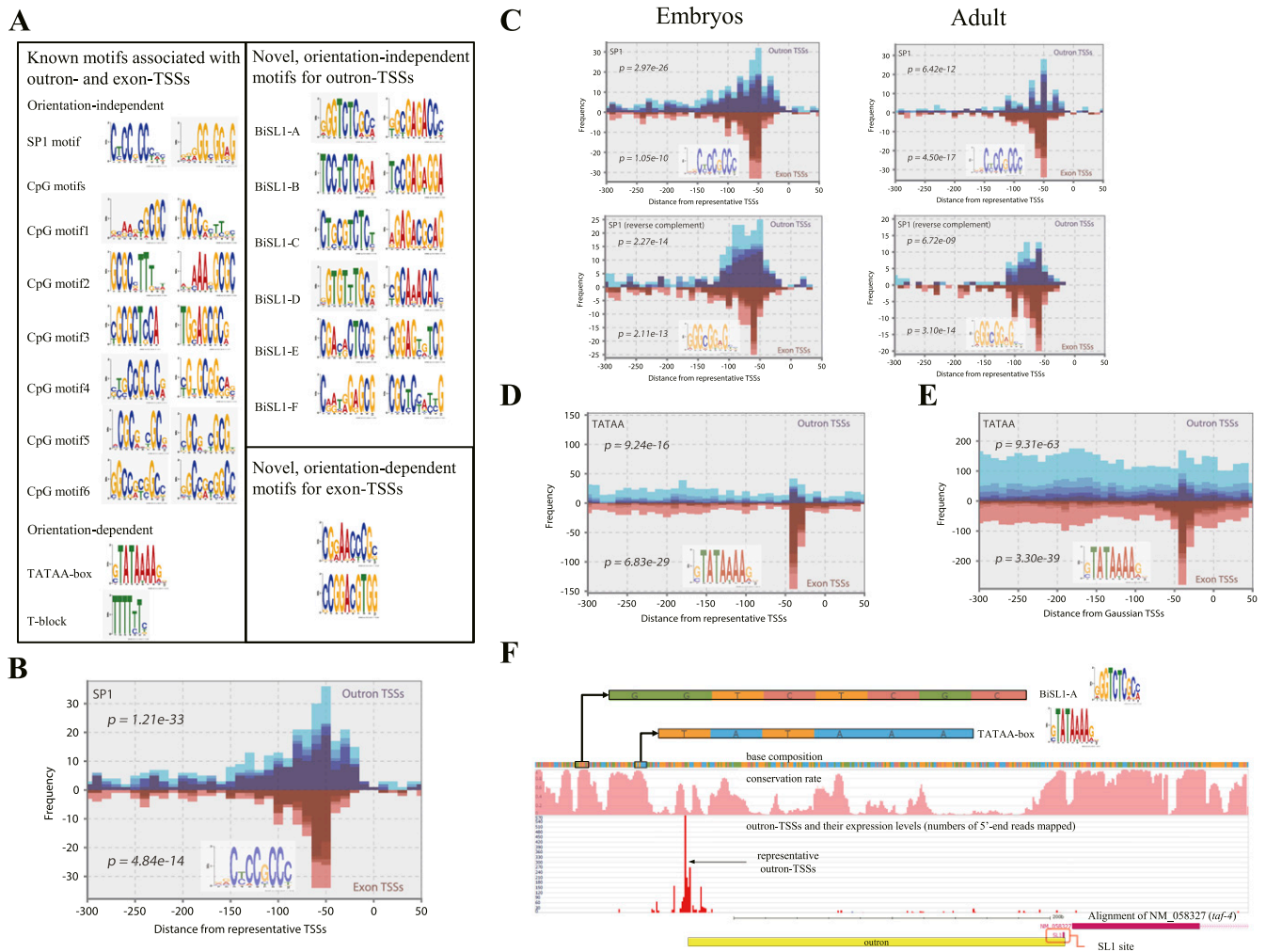
The large number of newly identified TSSs allowed us to search regions surrounding the TSSs for sequence motifs that might serve as *cis*-regulatory elements, in particular, transcription factor binding sites. Genomic regions ranging from  $-300$  bp to  $+50$  bp of representative TSSs (and in a separate analysis around the full collection of Gaussian TSS peaks) were selected as putative promoters. Both analyses were of use, although we note with the full Gaussian-TSS collection that multiple closely spaced TSSs in this collection would allow some single *cis*-elements to be multiply counted, providing some impetus to mine the representative TSS collection preferentially. Candidate promoter regions were analyzed using MEME (Bailey and Elkan 1994), a widely used program that identifies over-represented sequence motifs within a set of input sequences (see Methods). We used MEME to search for

statistically significant motifs of  $\leq 10$  bp ( $E$ -value  $< 10^{-20}$ ), noting that these computationally predicted motifs may not necessarily be functional despite their high statistical significance.

The evolutionary conservation of a motif is one of the strongest lines of evidence for its functional importance. We therefore assessed the conservation of each motif occurrence with respect to the average conservation rate among the genomes of six related nematode species for all positions in the occurrence (see Methods). Seventeen evolutionarily conserved motifs were detected in this analysis; of these, nine were previously known, and the remaining eight were novel (Fig. 4A). SP1 is a family of transcription factors involved in transcriptional control in several different systems. Figure 4B illustrates the frequency distribution of SP1 motif (Huber et al. 1998; Xi et al. 2007) occurrences around representative TSSs. Highly conserved SP1 motif occurrences were significantly enriched in promoter regions of representative outtron-TSSs ( $P = 1.21 \times 10^{-33}$ ) and exon-TSSs ( $P = 4.18 \times 10^{-14}$ ) according to the Wilcoxon rank-sum test (see Methods). Since all motifs were of length  $\leq 10$ , the number of all possible  $\leq 10$ -mers approximates  $(4/3) \times 10^6$  [ $\sim (4/3) \times 4^{10}$ ]. To reduce the possibility of selecting false-positive motifs from  $\leq 10$ -mers, we performed multiple hypothesis testing using the Bonferroni correction to check each motif at a significance level of 5% divided by  $(4/3) \times 10^6$  ( $=3.75 \times 10^{-8}$ ).

Of note, the SP1 motif was "orientation independent" in the sense that occurrences of its reverse complement were enriched and conserved around both outtron-TSSs and exon-TSSs (Fig. 4C). Other examples of orientation-independent motifs were six variants of the CGCG motif. These motifs were frequently observed in CG-rich regions for both outtron-TSSs and exon-TSSs, were preferentially located 20–80 bp upstream of outtron-TSSs, and were significantly conserved around outtron-TSSs (Supplemental Fig. S3). The evolutionary conservation of a motif and its positional enrichment at a specific location upstream of TSSs are both potential indicators of the functional importance of these motifs. Although *C. elegans* has several SP1 homologs, it is not clear if the promoter-proximal motif represents binding of one of these factors or of another unrelated factor with similar sequence specificity.

Another well-known motif found to be enriched in putative promoters was the TATAA-box (Lifton et al. 1978; Goldberg 1979; Basehoar et al. 2004). TATAA motifs are highly conserved promoter elements shared by all eukaryotes, are typically found closely upstream of TSSs, and are correlated with low-CG-content promoters, narrow distributions of TSSs, and a larger variability in expression



**Figure 4.** Conserved sequence motifs upstream of TSSs. (A) List of known motifs identified for outtron- and exon-TSSs, and a list of novel motifs. (B) Frequency distribution of motif occurrences and their conservation around representative TSSs. (The upper histogram) The frequency distribution of motifs around outtron-TSSs with downstream SL1 sites; (the lower histogram) the motif distribution around exon-TSSs without SL1 sites. The conservation rate of a motif occurrence is the average conservation rate of all positions in a given instance, where the conservation rate in the worm genome was calculated using six allied species (see Methods). Conservation rates ranging from 0 to 1 were divided into five levels. Darker colors indicate higher conservation rate; for example, the darkest color denotes a range from 0.8 to 1. In the figure, an SP1 motif (Xi et al. 2007) was observed around outtron-TSSs and exon-TSSs. The SP1 motif was significantly conserved around a strong peak  $\sim 50$  bp upstream of the TSS, and  $P$ -values were calculated for representative TSSs from the Wilcoxon rank-sum test (Methods). (C) The SP1 motif is orientation independent in the sense that its reverse complement was observed and conserved around outtron/exon-TSSs. Motifs were examined upstream of TSSs collected from two samples, embryos and adults. The distributions from the two samples tend to be similar; thus, the graphs for embryos are displayed exclusively hereafter unless significant differences were observed between the distributions of the two samples. (D) Distribution of TATAA-box occurrences around representative TSSs. TATAA-box occurrences were enriched significantly upstream of both types of TSSs according to the rank-sum test (Methods). (E) Distribution of TATAA-box around Gaussian TSSs. (F) A TATAA-box occurs upstream of the representative outtron-TSS for NM\_058327 (*taf-4*) that is encoded in the plus strand. The outtron is displayed as a yellow box. The information is available at <http://wormtss.utgenome.org/> by making a query of *taf-4*.

(Blake et al. 2003; Raser and O'Shea 2004; Carninci et al. 2006; Landry et al. 2007; Yang et al. 2007). We found (Fig. 4D) that TATAA-box motifs showed high conservation across the six nematode species and were significantly enriched 30–40 bp upstream of both representative outtron-TSSs ( $P = 9.26 \times 10^{-16}$ ) and exon-TSSs ( $P = 6.83 \times 10^{-29}$ ). The degree of TATAA-box enrichment appears to differ between representative outtron-TSSs and representative exon-TSSs in Figure 4D, leading to a concern that TATAA-box occurrences around the full collection of Gaussian outtron-TSSs might have been overlooked. Indeed, more TATAA-box occurrences were seen around the full collection of Gaussian outtron-TSSs as well as around the full collection of Gaussian exon-TSSs

(Fig. 4E), although most of them were less conserved. Figure 4F presents the promoter region for the *taf-4* gene (TBP-associated transcription factor, refseq ID NM\_058327). *taf-4* features a 240-bp outtron, indicated by the region between the representative TSS and the SL1 site, and has a TATAA-box motif located 31 bp upstream of its TSS. The TSS data add to previous analysis indicating an upstream TATAA in *C. elegans* (Grishkevich et al. 2011), providing a demonstration of TATAA-box preference at a specific location upstream.

Supplemental Figure S4 shows several other previously described motifs that we identified upstream of *C. elegans* TSSs. T-blocks (stretches of TTT or more Ts) are known to occur

frequently at 50–100 bp upstream of a start codon, with longer T-block occurrences correlated with lower nucleosome occupancy and a higher expression level (Grishkevich et al. 2011). Using our TSS information, we found that T-blocks and A-blocks, the reverse complement of T-blocks, were significantly conserved around outtron-TSSs, while they were not conserved around exon-TSSs. The motif associated with Initiator (Itr) binding ([CT][CT]A.[AT][CT][CT]) (Smale and Baltimore 1989) occurred frequently and was enriched around outtron-TSSs. As expected, splice acceptor enrichment was observed proximal to outtron-TSS sites.

In contrast, Supplemental Figure S5 presents known motifs, most of which were identified in the human or fruit fly genomes but were less evident in the worm genome. BRE, a TFIIB recognition element, was reported to be located immediately upstream of TATAA elements in the human genome (Lagrange et al. 1998); however, this tendency was not evident in the worm genome. BRE occurrences were conserved significantly, but the number of BRE occurrences was fairly small. A DPE (downstream promoter element) was identified in the fruit fly genome (Burke and Kadonaga 1996) but was not enriched as a downstream promoter element in the worm genome (Supplemental Fig. S5). A DCE (downstream core element) (Ohler et al. 2002) was investigated by checking the appearance of one of the three short motifs CTTC, CTGT, and AGC, and was seen ubiquitously (Supplemental Fig. S5). An MTE (Motif Ten Element) that was reported to be conserved from fruit fly to humans (Lim et al. 2004) was not identified as enriched in the worm promoterome.

In addition to these previously identified sequence motifs, we identified eight novel motifs that were statistically enriched proximal to worm TSSs and also showed evidence for evolutionary conservation (Fig. 4A). Six of these motifs may have usage or roles specific to outtron-TSSs because they were conserved more significantly around outtron-TSSs than around exon-TSSs. The six motifs were each orientation independent and likely to occur in narrow regions, usually ~50 bp upstream of outtron-TSSs (Fig. 5). These motifs were modestly frequent around representative TSSs as well as around Gaussian TSSs. We also noticed that the novel motifs were prevalent in bidirectional promoters (discussed later), suggesting their functional relevance to transcription factor-binding sites. Two less frequent, orientation-dependent motifs were found upstream of exon-TSSs (Supplemental Fig. S6).

Sequence analysis alone cannot conclusively demonstrate that individual motifs are transcription-factor (TF) binding sites. To this end, the worm modENCODE project (Gerstein et al. 2010) provided an informative list of 12 candidate motifs that were observed in proximity to transcription-factor (TF) binding sites HLH-1, LIN-13, MEP-1, and PHA-4. Of note, the list includes sequences that are highly similar to four of our novel motifs (BiSL1-A, BiSL1-F, reverse complements of BiSL1-D and BiSL1-C in our notation), motivating the experimental validation of other novel motifs as TF-binding sites. We analyzed the frequency distribution of distances between binding sites of each TF and newly determined representative TSSs, observing that binding sites of each TF were enriched 50–100 bp upstream of representative TSSs (Fig. 6A). In addition, we examined the frequency distribution of distances between binding sites of each TF and positions of each sequence motif. We found that the positions of each of four motifs (BiSL1-A, BiSL1-C, BiSL1-D, and BiSL1-F) significantly accorded with those of each TF (Fig. 6B; Supplemental Fig. S7,  $P < 10^{-18}$ ). They would be good starting points for future functional analysis.

## Motifs associated with tissue-specific gene expression

Motifs identified upstream of TSSs might have a fundamental role as *cis*-elements that regulate expression of genes in particular tissue types. We therefore examined the association between each motif and the expression of its associated gene (defined as the nearest downstream gene). As a measure of tissue-specific expression, we used the publicly available LongSAGE data, a comprehensive collection of SAGE data derived from a variety of tissues (e.g., oocytes, embryos, gut, muscle, hypodermal, neurons, neural cells, pharynx cells, and gonad: British Columbia *C. elegans* Gene Expression Consortium, <http://elegans.bcgsc.bc.ca/>) (Jones et al. 2001; McKay et al. 2003; Blacque et al. 2005; Khattra et al. 2007; McGhee et al. 2007; Meissner et al. 2009; Wang et al. 2009). Two thousand three hundred and forty-two (2342) genes have SAGE data as well as at least one of our identified motifs upstream of their TSS. To test whether the presence of a particular motif tends to be associated with higher expression (represented by steady-state mRNA levels) in a given tissue compared with genes lacking the motif, we performed Wilcoxon rank-sum tests for assessing the null hypothesis that distributions of expression levels from the two sets of genes were equivalent. Figure 6C and Supplemental Figure S8 show the results of this analysis. We performed multiple hypothesis testing to check each motif at a significance level of  $3.75 \times 10^{-8}$  as we examined the significance of the conservation of a motif. *P*-values less than the significance level are highlighted in red in Figure 6C. Notably, the presence of the novel BiSL-A motif upstream of a gene's TSS was associated with significantly higher expression in oocytes, embryos, gut, muscle, hypodermal, pharynx, and gonad samples. Genes with CpG motif4 in their putative promoters showed elevated gene expression in neuronal cells. High expression in gonad cells was associated with the presence of several upstream motifs, including AAAA, BiSL1-A, CpG motif 1-2, and SL1. The statistical association of several of our identified motifs with tissue-specific patterns of mRNA expression indicates that these sequences may be functionally important in proper gene expression and are strong candidate transcription factor binding sites.

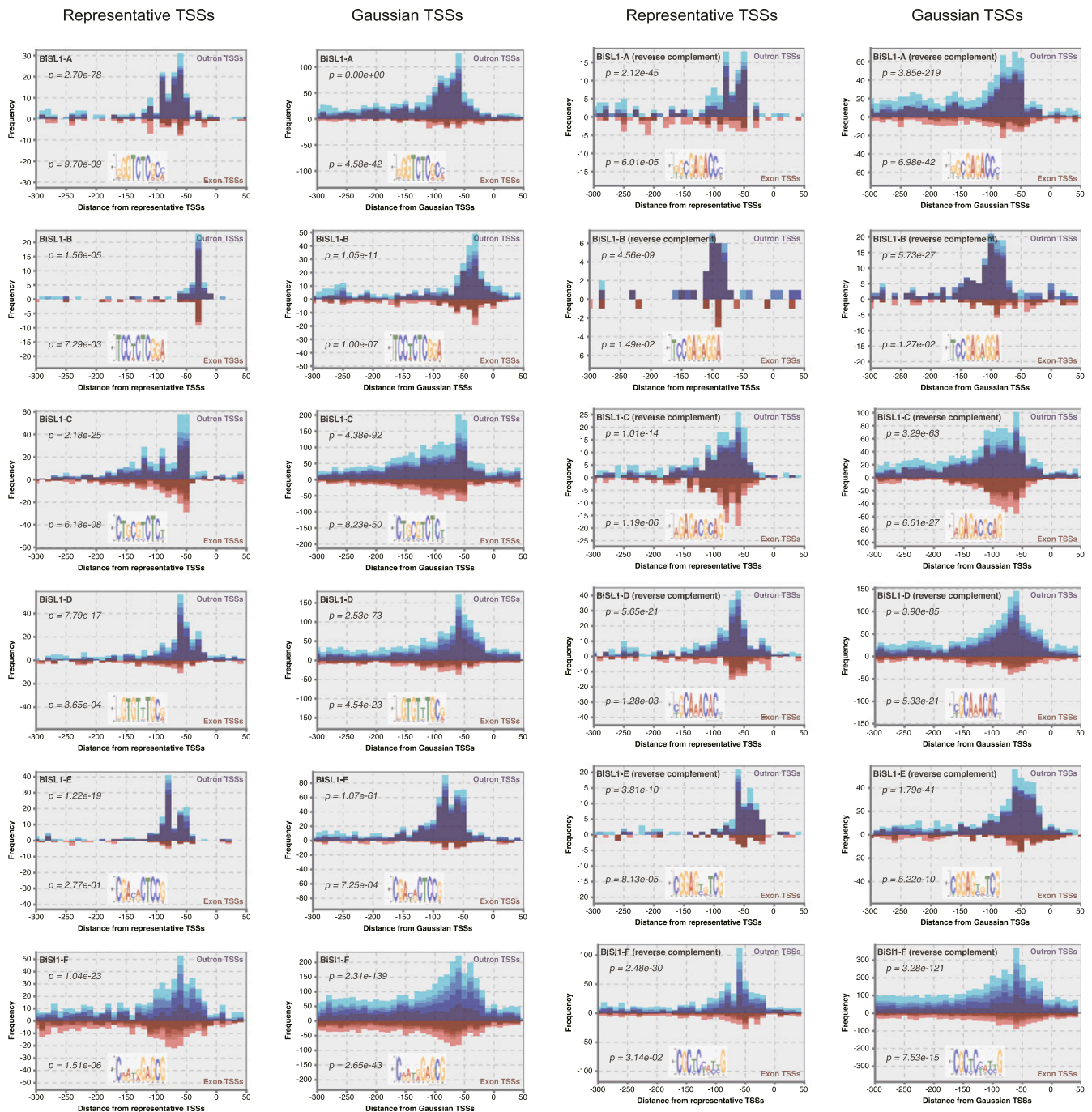
## Potential bidirectional promoters

Bidirectional promoters provide a compact mechanism for the joint transcriptional control of two coding regions. A comprehensive catalog of potential bidirectional promoters has not been available for *C. elegans*, due to limits in our knowledge of TSSs. Our comprehensive collection of TSSs together with information about gene expression levels provides a means of identifying candidate bidirectional promoters. Examining the candidate bidirectional promoters (those associated with TSSs 60–160 bp of each other) revealed that many of the identified motifs were frequently observed in these regions (Fig. 7; Supplemental Fig. S9). This class of *C. elegans* promoter should be of great interest for future experimental analysis.

## Discussion

The study provides a comprehensive collection of TSSs for 7351 *trans*-spliced genes in *C. elegans* such that their TSSs and gene bodies were linked by substantial paired-end reads, confirming the validity of the TSSs for the corresponding *trans*-spliced genes.

Transcription of many genes initiates predominantly at a single site. This could be due to selection for a single, regulatable promoter or for a single defined 5' UTR. If the selection is at the



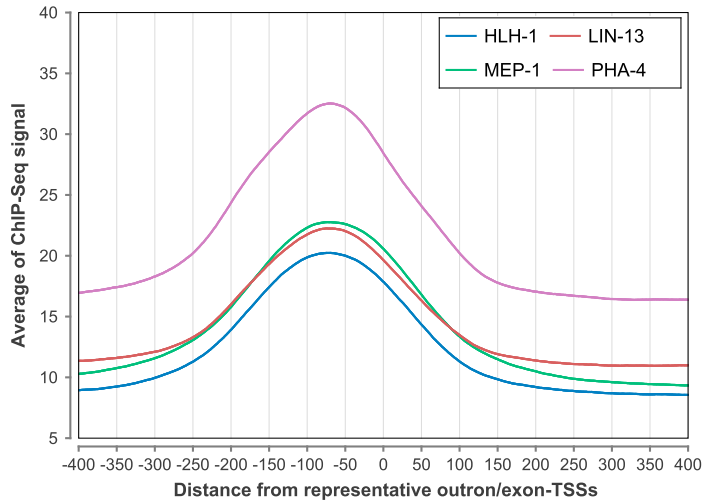
**Figure 5.** Frequency distributions of six novel orientation-independent motifs around representative and Gaussian outtron/exon-TSSs. Individual motifs are given new identifiers BiSL1-A–F. The distributions of the reverse complements of individual motifs are shown in the last two columns. *P*-values according to the Wilcoxon rank-sum test (see Methods) are shown for the distributions around representative outtron/exon-TSSs.

level of the UTR, we might expect that *trans*-splicing would eliminate that pressure, since the upstream region is replaced by a uniform 22-nt UTR. To address this question with the current data, we analyzed the average numbers of Gaussian outtron/exon-TSSs in the regions of the same length, 3000 bp, upstream of SL1 sites for outtron-TSSs and upstream of genes for exon-TSSs. We observed that the outtron-TSS average is significantly larger than the exon-TSS average (Supplemental Fig. 10A) according to the

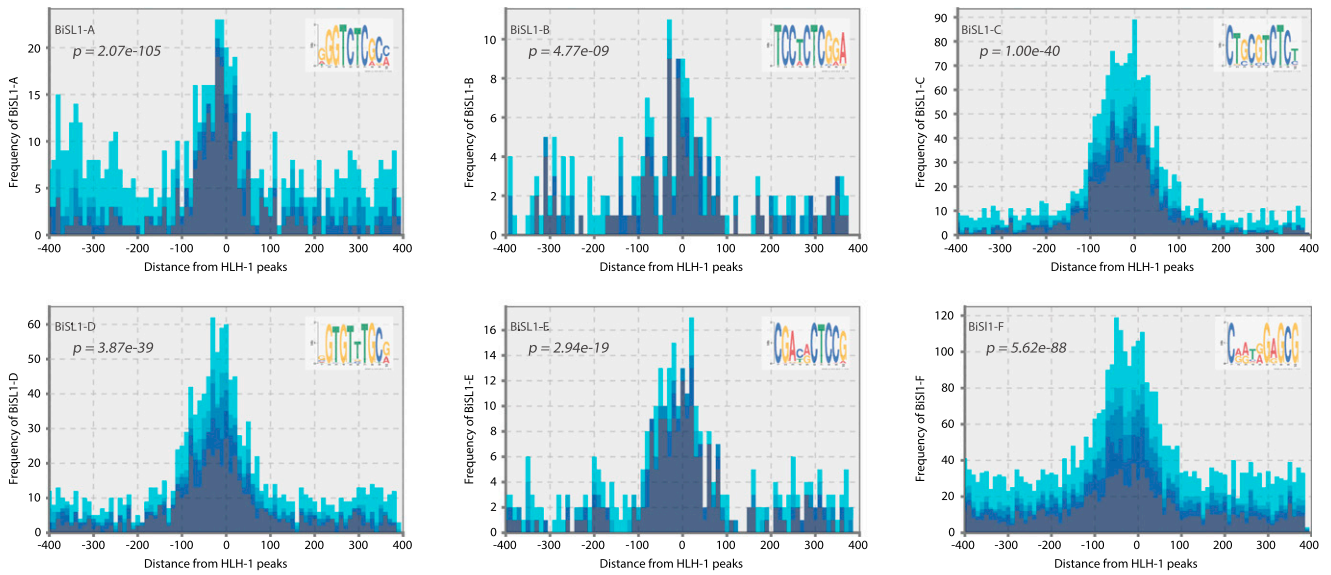
*z*-test ( $P$ -value  $< 10^{-116}$ ). In addition, the analysis of the distributions of the numbers of outtron- and exon-TSSs per gene also showed that the outtron-TSS distribution was significantly greater than the exon-TSS distribution (Supplemental Fig. 10B) according to the Wilcoxon rank-sum test ( $P$ -value  $< 10^{-117}$ ).

These observations are consistent with models in which the presence of *trans*-splicing might remove selective pressure for having a single TSS. In addition to potential differences in selective

A



B



C

Motif	SW032	SWN22	SWEG1	SWEM1	SW031	SW030	SW028	SW023	SW033	SW034	SW035	SW037	SW038	SW039	METALONG	SW040	Number of genes associated with each motif
	purified oocytes	N2 Embryos (longSAGE)	sorted gut cells	FACS sorted muscle cells (replicate 1)	FACS sorted muscle cells (replicate 2)	FACS sorted hypodermal cells	FACS sorted pan-neuronal cells (replicate 1)	FACS sorted ciliated neurons	FACS sorted pharynx cells	FACS sorted AFD neurons	FACS sorted pharyngeal marginal cells	FACS sorted ASER neurons	FACS sorted punc-4::GFP cells	FACS sorted pharyngeal gland cells	All longSAGE data metlibrary	dissected gonad	
AAAA	1.39E-02	6.19E-01	8.20E-01	2.63E-01	7.62E-01	8.69E-01	7.50E-01	2.12E-02	4.69E-01	8.50E-02	2.41E-02	6.33E-02	2.01E-01	3.25E-01	5.87E-02	4.08E-07	2122
BISL1.A	2.06E-10	1.81E-08	2.61E-13	2.51E-08	2.69E-09	2.80E-08	1.98E-03	1.66E-02	2.83E-10	7.63E-07	4.14E-05	6.22E-04	7.68E-04	1.36E-04	4.06E-10	2.95E-21	137
BISL1.B	1.61E-01	7.80E-01	5.56E-01	2.02E-01	5.12E-01	8.36E-01	2.45E-02	9.46E-02	5.70E-01	5.61E-01	3.18E-02	5.15E-01	1.18E-01	5.97E-01	1.02E-01	1.89E-02	22
BISL1.C	8.33E-01	8.06E-01	3.14E-01	2.36E-01	4.88E-01	4.28E-01	2.49E-05	4.24E-04	4.89E-01	1.40E-02	7.70E-04	4.96E-03	3.18E-05	3.88E-01	2.86E-02	1.30E-01	207
BISL1.D	1.34E-02	4.52E-01	2.82E-01	9.51E-01	5.58E-01	5.51E-01	4.90E-02	5.88E-01	6.68E-01	7.50E-01	3.74E-01	2.11E-01	2.84E-01	6.52E-02	2.72E-01	1.29E-02	209
BISL1.E	2.79E-01	4.73E-01	1.55E-01	5.20E-01	3.92E-01	8.71E-01	3.94E-01	4.59E-01	8.09E-01	8.51E-02	1.35E-01	2.99E-02	1.40E-01	5.33E-01	4.28E-01	4.57E-05	81
BIS1.F	1.20E-02	1.55E-01	1.12E-01	5.63E-02	5.04E-02	4.38E-02	5.64E-03	1.29E-02	7.08E-02	1.06E-02	5.21E-03	4.47E-02	3.48E-02	5.70E-03	1.18E-02	2.68E-06	435
CpG_motif1	1.47E-01	4.12E-01	2.33E-01	8.54E-01	9.84E-01	6.13E-02	2.70E-02	4.78E-02	8.19E-01	1.14E-01	3.87E-02	5.30E-01	8.93E-03	1.08E-05	4.55E-01	2.24E-12	1206
CpG_motif2	1.17E-02	9.73E-01	1.77E-02	2.54E-01	1.30E-01	9.43E-01	3.36E-01	5.12E-01	1.74E-01	7.57E-01	4.38E-01	8.96E-01	4.55E-01	5.52E-02	2.97E-01	4.96E-16	366
CpG_motif3	2.52E-02	2.85E-01	8.54E-02	9.79E-01	1.24E-01	3.25E-01	7.84E-01	9.14E-01	1.06E-01	5.60E-01	9.00E-01	8.95E-01	8.58E-01	3.32E-01	6.23E-01	2.30E-06	61
CpG_motif4	6.24E-01	4.27E-03	3.88E-01	2.03E-05	1.61E-02	3.58E-02	2.50E-04	1.11E-06	6.40E-04	6.85E-09	6.15E-05	5.17E-09	2.68E-05	7.53E-05	5.56E-08	5.48E-01	737
CpG_motif5	3.55E-03	1.99E-02	3.05E-01	1.88E-03	2.77E-03	3.91E-02	1.16E-02	3.21E-02	6.70E-03	2.20E-01	6.84E-02	4.01E-02	1.09E-03	1.84E-02	3.03E-04	5.15E-04	551
CpG_motif6	4.13E-01	5.80E-01	8.45E-01	5.42E-02	9.64E-01	9.71E-01	5.06E-02	4.13E-01	2.08E-01	1.05E-02	1.02E-01	1.48E-02	1.68E-02	1.71E-02	7.69E-02	4.36E-02	291
Kozak	6.12E-01	1.52E-01	8.42E-01	3.30E-03	2.98E-01	9.86E-01	1.23E-01	4.61E-02	5.71E-01	2.16E-02	2.17E-01	1.32E-01	1.46E-01	6.88E-01	1.12E-01	1.11E-04	1507
ExUni.A	4.28E-01	7.22E-01	3.71E-01	7.85E-01	6.31E-01	9.46E-01	6.54E-01	1.85E-01	2.36E-01	5.59E-02	1.53E-01	1.17E-02	1.38E-01	4.65E-01	1.49E-01	8.96E-01	18
ExUni.B	1.29E-01	3.58E-01	4.51E-01	6.70E-01	9.94E-01	8.95E-01	9.71E-01	5.45E-01	1.94E-01	2.24E-02	1.12E-01	7.68E-02	7.72E-01	1.25E-01	1.57E-01	2.78E-01	2
SL1	4.76E-01	4.73E-02	1.17E-01	7.79E-01	7.91E-01	9.35E-01	3.98E-02	2.49E-01	6.80E-01	1.57E-01	2.15E-03	2.89E-03	3.16E-01	6.22E-01	2.97E-02	1.72E-09	473
SP1	7.05E-01	6.58E-03	4.74E-03	2.09E-04	3.34E-04	1.89E-04	4.15E-03	1.53E-04	7.86E-04	4.63E-07	1.48E-04	5.63E-07	1.43E-05	9.37E-02	6.37E-08	6.27E-03	183
T.block	7.13E-01	5.87E-02	7.21E-01	2.86E-04	1.51E-01	1.29E-02	5.72E-03	8.15E-03	6.56E-02	4.91E-03	9.82E-02	5.36E-03	8.72E-02	8.20E-02	7.84E-03	1.10E-04	2259
TATAA	7.97E-01	3.27E-01	9.88E-01	1.27E-01	4.89E-01	5.04E-01	2.58E-01	3.59E-01	2.33E-01	2.80E-01	6.21E-01	8.77E-02	5.47E-01	4.37E-01	3.61E-01	5.08E-01	445

Figure 6. (Legend on next page)

pressure, several conceivable mechanistic differences might serve as a basis for the observed difference in dispersion between exon- and outon-TSS distributions (e.g., subtle differences in sequence composition or chromatin structure between outon and upstream exon regions). The combination of mechanistic and selective differences in the two 5' regions should certainly become clearer as additional analytic and experimental tools are applied to *C. elegans* promoters. While sequence composition and technical influences cannot be ruled out, these observations are intriguingly consistent with the possibility that *trans*-splicing may afford an advantage to the organism in permitting flexibility of 5' gene structures (Blumenthal 2005).

Arrays of positioned and phased nucleosomes have been seen around TSSs in mammals and other eukaryotes (Mavrich et al. 2008; Kaplan et al. 2009; Sasaki et al. 2009; Weiner et al. 2010; Valouev et al. 2011), with this study confirming this characteristic in *C. elegans*. In addition, nucleosome positioning and Pol II enrichment downstream from SL1 sites suggested a distinctive association between signals for nucleosome positioning and signals for *trans*-splicing. In the regions upstream of newly identified TSSs, several evolutionarily conserved motifs were uncovered, including known motifs (e.g., TATAA-box and SP1) found in other species as well as six novel TSS-specific motifs with downstream SL1 sites. These discoveries led us to reveal 694 potential bidirectional promoters of ~100 bp in length that harbor orientation-independent motifs such as CpG, SP1, and novel motifs. We hope that this data set will serve as a valuable resource in functional genomics studies for *C. elegans*.

## Methods

### 5'-End library construction

Figure 1A illustrates the process of 5'-end library construction (Hashimoto et al. 2009). *C. elegans* nuclei were prepared by disruption of animals and sedimentation of lysates through a 2 M sucrose cushion (lysis and gradient reagents from Sigma-Aldrich, catalog no. NUC-101). We started with ~2 mL of worm grind (adult or embryo) for each nuclear preparation, with purified nuclei stored in 200  $\mu$ L of the NUC-101 storage buffer. We used TRIzol for the isolation of total RNA, with subsequent cleaning using the RNeasy Mini (QIAGEN). We then replaced the cap structure of mRNA with an Illumina RNA linker according to the oligo-cap method (Maruyama and Sugano 1994) as follows: The purified total RNA (starting with 5  $\mu$ g) was treated with bacterial alkaline phosphatase (TaKaRa). The RNA was extracted twice with phenol:chloroform (1:1), ethanol-precipitated, and treated with tobacco acid pyrophosphatase (TAP). Then a RNA linker was ligated using RNA ligase (TaKaRa): a 5'-oligo (5'-ACAGGUUCAGAGUUCUACAGUCCAGCAG-3') linker. After linker ligation, the RNA was depleted of ribosomal RNA using the RiboMinus Eukaryote Kit (for RNA-seq, Invitrogen, catalog no. 10837-08). After removing ribo-

somal RNA, a cDNA library was produced from 10 pmol of random hexamer primer (5'-CAAGCAGAAGACGGCATAACGACAGCAGN NNNNNC-3') using SuperScript III (Invitrogen) by incubating for 10 min at 26°C, 90 min at 50°C, and 15 min at 70°C. After first-strand synthesis, RNA was no longer needed and was degraded in 15 mM NaOH for 1 h at 65°C. cDNA was amplified in a volume of 100  $\mu$ L using PCR with 16 pmol of 5' (5'-AATGATACGGCGA CCACCGACAGGTTTCAGAGTCTACAGTCC-3') and 3' (5'-CAAG CAGAAGACGGCATAACGA-3') PCR primers. The cDNA was produced using 15–20 cycles of 1 min at 94°C, 1 min at 58°C, and 10 min at 72°C. The PCR fragments were size-fractionated by 8% polyacrylamide gel electrophoresis, and the fraction of 150–300 bp was recovered. The quality and quantity of the obtained single-stranded first-strand cDNAs were assessed using Bio-Analyzer (Agilent).

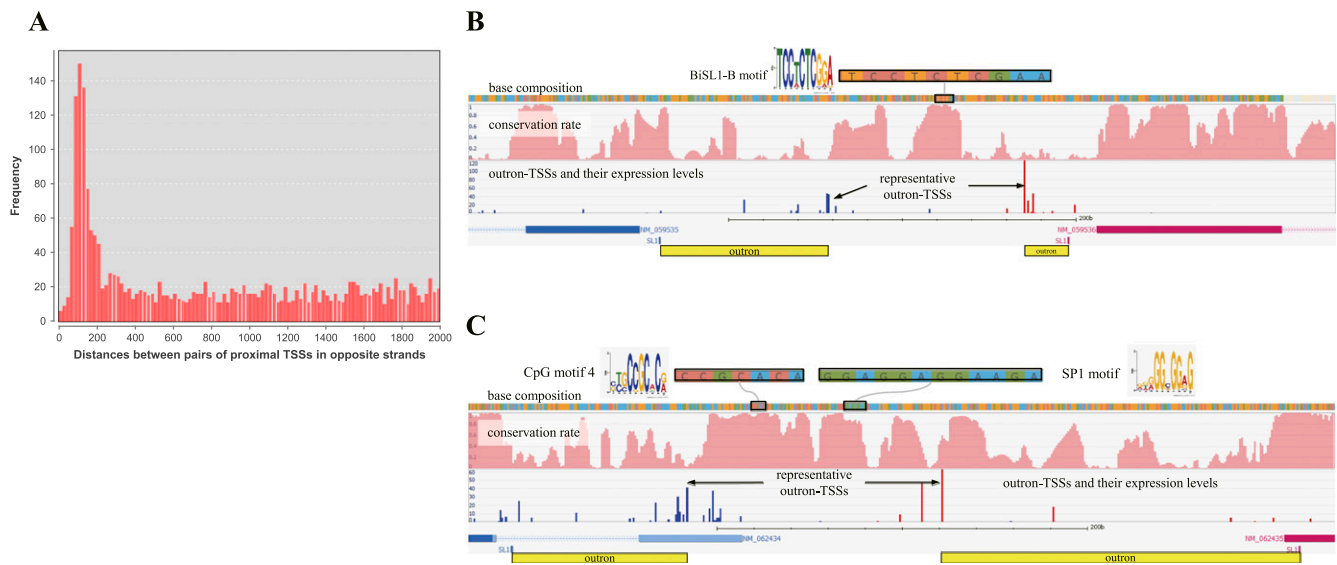
### Read alignment

To map the short reads of Illumina GALL to the reference genome of *C. elegans*, we used BWA with the default setting (4% of mismatches of read lengths were allowed). Many reads were eliminated unless they contributed to the TSS identification. For example, several *trans*-spliced reads failed to be mapped because of splice-reader sequences (SL1, SL2, etc.) attached to the head of the sequences after transcription. These unmapped reads were useless in identifying TSSs and were therefore discarded. Despite application of the RiboMinus Eukaryote Kit, we found that one-third of the reads were fragments of rRNA. These numerous rRNA reads simply reconfirmed rRNA regions that were well annotated in WormBase.

### TSS peak calling

After mapping of the short-read sequences to the reference genome, we sorted reads in the order of the genome coordinate and then computed the number of reads mapped to the same position to collect candidate TSS peaks. The distributions of the peaks showed various shapes; single-peak, multiple peaks, broadly distributed peaks, etc. To classify these peak types, we attempted to fit a Gaussian mixture model to our data. One issue was to determine the number of Gaussian distributions. We used X-means clustering (Pelleg and Moore 2000) of peaks that used Gaussian model and penalty score to avoid overfitting (Bayesian-information criteria, BIC), so that the result of X-means clustering is expected to have a nice fit to a Gaussian-mixture model representing the observed peaks. The observed number of clusters before genes ranged from 1 to 70. These clusters contain noisy peaks that may inadequately distort the Gaussian distributions. Thus, for each cluster, we trimmed marginal peaks (less than 5 tag counts) and applied the Gaussian distribution so as to compute  $N$  (gene expression level) and  $\sigma$  (distribution shape) values. TSSs within 10 bp from the SL1 sites were removed to avoid confusion with the *trans*-spliced tags. We selected clusters with the highest expression level within

**Figure 6.** Motifs associated with transcription factors and tissue-specific genes. (A) For transcription factors (TF), HLH-1, LIN-13, MEP-1, and PHA-4, we obtained TF-binding scores (ChIP-seq signals) from the modENCODE database (Gerstein et al. 2010), calculating the average TF-binding score around the representative TSSs. The TF-binding score is enriched 80–50 bp upstream of representative exon-TSSs as well as outon-TSSs, respectively. (B) Frequency distribution of motif occurrences around positions associated with locally maximum HLH-1 binding scores. The frequency distributions for LIN-13, MEP-1, and PHA-4 can be found in Supplemental Figure S7. Color shade represents conservation rates (see Fig. 4B; Methods). We checked the null hypothesis that two distributions of conservation rates of a motif in proximity to TF-binding sites and in the other nonpromoter genomic regions were equal, by using the Wilcoxon rank-sum test (see Methods). A lower  $P$ -value of a motif indicates a higher enrichment of the motif at the binding sites. (C) The rows present sequence motifs, and the columns show tissue types in the LongSAGE data, except for the last column that displays the number of genes associated with each sequence motif among 2342 genes. The table presents  $P$ -values of the Wilcoxon rank-sum test for assessing the null hypothesis that two distributions of expression levels of genes with the presence or the absence of the upstream motif were equal in each tissue. To highlight significantly low  $P$ -values according to the Bonferroni correction, we colored  $P$ -values  $\leq 3.75 \times 10^{-8}$  red, and  $P$ -values ranging from  $3.75 \times 10^{-8}$  to  $10^{-3}$  orange.



**Figure 7.** Potential bidirectional promoters. (A) Distribution of distances between paired proximal TSSs in opposite strands. The remarkable number of loci (total instances more than 600) with paired divergent TSSs at a distance of 60–160 bp suggests an abundance of potential bidirectional promoters. This range (60–160 bp) is consistent with the observation that motif sequences (candidate *cis*-elements) are typically enriched ~50 bp upstream of the TSSs. In reality, many motifs, including novel ones, were observed frequently in those potential regions. (B) Example of candidate bidirectional promoter region. Two expressed representative outtron-TSSs in the opposite strands were separated by 115 bp. A novel motif (TTCTCTCGGA), named “BiSL1-B” in Figure 5 (respectively, its reverse complement), was identified ~50 bp (~70 bp) upstream of the plus (minus) strand representative TSS. This is in accordance with the enrichment of the motif ~30–50 bp (70–100 bp) upstream of the plus (minus) strand TSS obtained from our genome-wide analysis (see BiSL1-B in Fig. 5). Of note, the motif was highly conserved among the six allied species. The two outtrons from the representative TSSs to their nearest downstream SL1 sites were different in length, demonstrating the difficulty of estimating outtrons and TSSs solely from SL1 sites and the genomic sequence. (C) Another example of a bidirectional promoter. Two common motifs (SP1 and CpG motif 4) were highly conserved between two expressed outtron-TSSs. Multiple highly expressed and isolated TSSs were seen in both strands, and the wide distribution of the occurrences of these motifs may account for the multiple outtron-TSSs (SP1 in Fig. 4B and CpG motif 4 in Supplemental Fig. S3). Many more examples of regions harboring conserved motifs between expressed outtron-TSSs were seen (Supplemental Fig. S9), which could function as potential bidirectional promoters (696 candidates in Supplemental Table S4). Expression levels between the pairs of genes have a very small similarity (Pearson’s correlation coefficient <0.066).

3000 bp before the gene start sites as representative TSSs. Supplemental Table S1 gives a gene-by-gene list of representative TSSs as well as Gaussian TSSs. In the table, we used the gene annotations in WormBase WS220. We merged gene models that overlapped in individual regions of the genome into unique groups (choosing the longest model in each case) so as to associate each TSS with a single neighboring entity.

### Finding motifs and their occurrences around TSSs

Finding motif sequences is a computationally expensive task because of numerous combinations of ACGT letters of various lengths. MEME is a widely used program for detecting motif sequences. To speed up the analysis, we used an MPI version of MEME, and it took us ~1 d to run the program using a cluster of machines equipped with 64 CPUs using the option “-p 64 -maxsize 10000000 -dna -nmotifs 30 -mod zoops -minw 3 -maxw 10.” Each letter in a motif is associated with bits that express a likelihood of the presence of the letter at the position in the motif. In searching all promoter regions (350 bp to –50 bp around TSSs) for a motif, we used letters associated with  $\geq 1$  bits in a motif for testing exact matches, while we replaced other letters by ?, a wild card matching any one letter. In this way, we transformed a motif into a regular expression. For example, we converted the SP1 motif in Figure 4B into the regular expression “C?CC?CC” and the TATAA-box in Figure 4D into “TATAAAA.” We then searched all promoter regions for the matches of the regular expression corresponding to a motif. We note that the use of regular expressions to search for many known motifs was preferable over weight-based methods, because

the relevant publications lacked bit-significance scores. To be consistent with MEME, for the motifs found by MEME, we also used regular expressions of characters whose bit-significance scores were >1.

### Measuring the significance of the conservation of a motif around TSSs

We calculated the conservation rates of motifs at each position from a comparison of the genomes of six related nematode species, *C. elegans* (The *C. elegans* Sequencing Consortium 1998), *Caenorhabditis brenneri* (Stein et al. 2003), *Caenorhabditis briggsae*, *Caenorhabditis remanei*, *Caenorhabditis japonica*, and *Pristionchus pacificus* (Thomas 2008), with alignments derived from the UCSC Genome Browser resource (Siepel et al. 2005; Fujita et al. 2011).

Subsequently, we tested the significance of the conservation of a motif in promoter regions that range from 300 bp to –50 bp of representative TSSs. If Gaussian TSSs are considered, the same motif occurrence can be counted more than once. To avoid this double counting, we focused on representative TSSs so as to associate any motif occurrence with its downstream representative TSS, uniquely. We checked the null hypothesis that two distributions of conservation rates of a motif in promoter regions and in the other nonpromoter genomic regions were equal, by using the Wilcoxon rank-sum test. The number of all occurrences of a focal motif in nonpromoter regions was huge, and using them all was likely to lead to incorrect calculation of *P*-values. We therefore selected 10,000 instances at random from all the occurrences,

and performed this trial 100 times to calculate *P*-values. Since some *P*-values happened to be extremely small or large, the median of the *P*-values was treated as the representative *P*-value. Because we considered motifs of  $\leq 10$ -mers in length, according to the Bonferroni correction, we tested each motif at a significance level of 5% divided by  $(4/3) \times 10^6$  ( $=3.75 \times 10^{-8}$ ), where  $(4/3) \times 10^6$  approximates the number of all possible  $\leq 10$ -mers.

## Data access

All sequence data are deposited at the NCBI Sequence Read Archive (SRA) (<http://www.ncbi.nlm.nih.gov/sra>) under accession number SRA060670. Our genome browser is available at <http://wormtss.utgenome.org/>. Both the primary data and analytical tables in BAM and WIG format are also available at <http://wormtss.utgenome.org/browser/download.jsp>, so that public databases such as WormBase and UCSC can incorporate our data easily into their sites.

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