Hypothesis

Computational and molecular characterization of multiple isoforms of \(lfe-2\) gene in nematode \(C.\) \textit{elegans}

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Abstract:
\(C.\) \textit{elegans} C46H11.4 gene encodes a Let-23 fertility effector/regulator protein of the EGF-receptor class of the tyrosine kinase family. Alternative splicing is a major mechanism of generating protein diversity in higher eukaryotes. \(C.\) \textit{elegans} genome sequencing consortium has reported three alternatively spliced transcripts of C46H11.4 gene which encodes for three hypothetical proteins namely, C46H11.4a, C46H11.4b and C46H11.4c. Using a combination of various bioinformatics tools like gene or exon finding programmes, blast searches, alignment tools etc followed by experimental validation, we report the presence of three more alternatively spliced transcripts which encode for novel hypothetical proteins C46H11.4d, C46H11.4e and C46H11.4f. These isoforms arise as a result of alternative splicing in the pre-mRNA encoded by gene C46H11.4. These novel un-reported spliced variants not only point towards the extent of alternative splicing in \(C.\) \textit{elegans} genes but also hint towards the complex nature of alternative splicing.

Keywords: \(lfe-2\); Let-23 fertility effector/regulator protein; exon prediction; alternative splicing; \(C.\) \textit{elegans}

Background:
Protein tyrosine kinases (PTKs) are important molecules for intra- and inter-cellular communication as well as for survival in eukaryotes, playing a major role in signal transduction processes. [1] There are two main classes of PTKs: Receptor Tyrosine Kinases (RTKs) or cellular, and Non-Receptor Tyrosine Kinases (NRTKs) or non cellular. [2] RTKs are transmembrane proteins that are involved in the control and regulation of several key cellular processes. To date, two groups of RTKs have been described in \(C.\) \textit{elegans}: orthologs of RTKs known in mammals (e.g. DAF-2, EGL-15, KIN-8, LET-23 AND VAB-1) and RTKs identified only in \(C.\) \textit{elegans} (e.g. KIN-15 and KIN-16). [3] \(C.\) \textit{elegans} C46H11.4 gene encodes \(lfe-2\), a Let-23 fertility effector/regulator protein of the gene class LET-23, a member of the EGF-receptor tyrosine kinases family. \(lfe-2\) activity is required for normal levels of hermaphrodite fertility and for tissue-specific negative regulation of the LET-23 signaling pathway. [4] LET-23 is seen to play a pivotal role in free living nematode \(C.\) \textit{elegans}. It is required for vulval induction as it helps the VPCs (vulval precursor cells) to respond to the signals that induces vulval differentiation. [5] It also signals through the RAS/MAPK pathway to specific cell fates in the vulval cord, vulva, male tail and excretory system and through a PLC/IP3 pathway to regulate ovulation. [6] Large numbers of spliced transcripts have been reported from various groups of RTKs and their associated families in almost all higher eukaryotes like humans, mouse and several other organisms including \(C.\) \textit{elegans}. Alternative splicing is a powerful means of regulating gene expression and enhancing protein diversity. In fact, the majority of metazoan genes encode pre-mRNAs that are alternatively spliced to produce anywhere from two to tens of thousands of mRNA isoforms. Alternatively spliced exons are typically identified by aligning EST clusters to reference mRNAs or genomic DNA. [7] This approach has several limitations; firstly they may underestimate alternative splicing because of their incomplete coverage and lack of information regarding combinations of exons that are utilized. Secondly, it is hard to detect highly specific as well as low copy number splice variants by EST-based approaches [8] or microarray technologies. [9, 10] Most of the other approaches used work on the genomic level by predicting splice sites and introns/exons using sequence signals and composition differences. [11, 12] These methods are more devoted to gene prediction and usually yield only one optimal gene structure. Thus, methods that facilitate the identification of alternative exons would be quite useful to assist in genome annotation. Although several approaches for the ab initio prediction of gene structure have been developed, the ab initio prediction of alternative splicing
using a combination of gene/exon finding programmes has not been considered. Genefinder prediction by the *C. elegans* sequencing consortium of genomic sequence of C46H11.4 has reported three spliced variants C46H11.4a, C46H11.4b and C46H11.4c. In spliced transcript C46H11.4a, a new exon originates from the large intronic region between exon 5 and exon 6 of C46H11.4c (the biggest isoform of C46H11.4 gene) and splices with the 6th exon of the parent transcript C46H11.4c to form a new spliced variant (Figure 1). Similarly, a new exon originates from the intronic gap between exon 4 and exon 5 of C46H11.4c and splices with the 5th exon of the parent transcript C46H11.4c to form its new spliced transcript C46H11.4b (Figure 1). Here, we report the detailed analysis data of the large intronic and 5' and 3' untranslated regions (5'-UTR and 3'-UTR) of C46H11.4 gene using various gene/exon finding programmes and several other bioinformatics tools. Computational analysis predicted the existence of three new spliced variants C46H11.4d, C46H11.4e and C46H11.4f which were subsequently confirmed by the presence of different corresponding transcripts by RT-PCR using gene specific primers and RNA isolated from mixed population of *C. elegans*.

**Methodology:**

**Animals**

The Bristol N2 wild-type strain of *C. elegans*, grown on NGM plates and maintained on *Escherichia coli* (OP50) essentially as described in [13] were used in all experiments.

**Reagents**

RevertAid™ M-MuLV Reverse Transcriptase and oligo (dT)$_{18}$, and single-stranded cDNA was synthesized using Revert Aid™ Reverse Transcriptase at 42°C for 1 h (total vol. 20 micro liter). Subsequently, 1 micro liter of the single-stranded cDNA preparation was added to PCR system buffer for PCR using appropriate sets of primers (total vol. 20 micro liter). PCR was performed for 30 cycles (denaturation, 94°C, 5 min, 1 cycle; denaturation, 94°C, 45 sec; annealing, 58°C, 1 min; extension, 72°C, 1 min; finally, 72°C for 7 min). RT-PCR product (5 micro liter) was subjected to electrophoresis on a 2% (w/v) agarose gel, stained with ethidium bromide and photographed on a UV-trans illuminator.

**Bioinformatics tools**

Genomic sequence of cosmid C46H11 (Accession No.U88314) was downloaded from the NCBI nucleotide database (http://www.ncbi.nlm.nih.gov/entrez). Various other databases used for searches were those at the *C. elegans* Sequencing Project Center at (http://sanger.ac.uk/Projects/Celegans/blasthserver.shtml) and Yuji Khoara’s *C. elegans* expressed sequence tag (EST) database (http://dmbj.nig.ac.jp/htmls/c-elegans/html/CE_INDEX.html). Homology and similarity searches of the polypeptide and nucleotide sequences were performed using the BLASTN and BLASTP non-redundant database (http://www.ncbi.nlm.nih.gov/BLAST). Gene/Exon finding tools used were HMMGene (http://www.ncbi.nlm.nih.gov/BLAST). Genescan (http://genes.mit.edu/GENSCAN.html), GeneSplicer (http://chcb.umd.edu/software/GeneSplicer/), FGENESH (http://www.saffery.com/ bery.phtml), ORF findings tools used were located at (http://www.bioinformatics.vg/sms/orf_find.html) and (http://www.prl.msu.edu/clab/dnajava/orf_find.html). Alignment analysis was carried out using the Gene stream Align tool (http://www2.igh.cnrs.fr/bin/align-guess.cgi) and ClustalW tool available at (www.ebi.ac.uk/clustalw). Primers were designed manually or using the tool Primer3 (http://biotools.umassmed.edu/bioapps/ primer3/www.cgi). Various other tools and bioinformatics programmes used were same as described in our previous study/earlier. [15]

### Hypothesis

**Preparation of RNA**

Total RNA was isolated from mixed-stage nematodes using the method described earlier. [14] Finally, total RNA was dissolved in diethyl pyrocarbonate-treated distilled water. Purity of RNA was checked using denaturing agarose gel electrophoresis.

**Reverse transcriptase (RT)-PCR**

Computationally predicted spliced variants were validated using RT-PCR. *C. elegans* total RNA (3 micro gram) was primed with oligo (dT)$_{18}$, and single-stranded cDNA was synthesized using Revert Aid™ Reverse Transcriptase at 42°C for 1 h (total vol. 20 micro liter). Subsequently, 1 micro liter of the single-stranded cDNA preparation was added to PCR system buffer for PCR using appropriate sets of primers (total vol. 20 micro liter). PCR was performed for 30 cycles (denaturation, 94°C, 5 min, 1 cycle; denaturation, 94°C, 45 sec; annealing, 58°C, 1 min; extension, 72°C, 1 min; finally, 72°C for 7 min). RT-PCR product (5 micro liter) was subjected to electrophoresis on a 2% (w/v) agarose gel, stained with ethidium bromide and photographed on a UV-trans illuminator.

**Primer3:**

1. F1 (7457-7478; 22 mer):
   - 5' CGTCGGGTACGGGAAGCAGCTG 3'
2. F2 (8919-8943; 25 mer):
   - 5' CCTAGTCAGGAGCCTGTTGTAC 3'
3. F3 (7981-8002; 22 mer):
   - 5' GAACTACGCTCATTCTGCTAGTGAG 3'
4. F4 (4138-4164; 27 mer):
   - 5' CCTAGTCAGGAGCCTGTTGTAC 3'
5. R1 (16984-17005; 22 mer):
   - 5' CGTCGGGTACGGGAAGCAGCTG 3'

The direction and relative position of the primers are indicated in figure 1.

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Results and discussion:
Computational analysis and prediction of new alternatively spliced transcripts of \(lfe-2\) encoding gene \(C46H11.4\)

Computational analysis of genomic DNA sequences using gene finders to predict spliced transcripts and subsequent confirmation of the predicted spliced variants have been successfully demonstrated in \(C.\) \(elegans\). \[14\] Several studies aimed at detecting new spliced variants arising due to alternative splicing have been done in various RTKs and related families in Humans \[16, 17, 18\], mouse \[19, 20\] etc. In light of these previous studies, we started with the detailed analysis of the 5’ and 3’ untranslated (UTR) and large intronic regions of \(C46H11.4\) gene. Analysis comprised of a series of bioinformatics tools like gene/exon finding programmes, ORF finders and several other bioinformatics tools (as mentioned in materials and methods). These tools predicted several exons possibly capable of replacing the existing exon(s) and thus creating alternatively spliced variant of the gene. From the several exons predicted above, we selected only the “common exons” capable of replacing the existing exon(s) and thus creating spliced transcript of the gene without affecting the reading frame of the protein. Further, the possibility of occurrence of that spliced exon/transcript was analyzed by a comparative analysis between the original protein and the new protein formed by addition/exclusion of alternative exons using various alignment tools. Lastly, several other parameters like percent-amino acid replacement, codon usage, sense nature i.e. whether from positive or negative strand, the probability score of occurrence of that exon etc. were also checked to ensure accuracy of predicted spliced transcript of the gene. Thus, Gene/Exon finder analysis of about 1.6 kb of intronic gap region between exon 1 and exon 2 of existing \(C46H11.4\) and about 4 kb of 5’ untranslated region (between the predicted stop codon of the upstream gene \(C46H11.6\) and the initiation codon of \(C46H11.4\)) predicted three new exons. These new exons 1d, 1e and 1f encoded 16, 22 and 24 amino acids residues and have methionine as the initiation codon (Table 1 in supplementary material). These newly identified exons were potentially capable of splicing with exon 2 or exon 3 of the parent gene \(C46H11.4\) (Figure 1, Table 1 in supplementary material) without affecting the reading frame of the protein.

Thus computational analysis of the large intronic and 5’ and 3’ untranslated (UTR) region predicted three new spliced transcripts of the gene \(C46H11.4\), which we have named \(C46H11.4d\), \(C46H11.4e\) and \(C46H11.4f\). In all the three new spliced transcripts a new N-terminal exon in each splices with the already existing exon of the parent isoform in the manner as depicted (Figure 1). In \(C46H11.4d\), a new potential exon originates from the 5’UTR region of \(C46H11.4c\) and splices with the 3rd exon of the parent transcript \(C46H11.4c\). (Figure 1). Whereas, in spliced variant \(C46H11.4e\), a new potential exon originates from the large intronic gap between exon 1 and exon 2 of \(C46H11.4c\) and splices with the 2nd exon of \(C46H11.4c\) (Figure 1). Similarly, a new potential exon originates from the large intronic gap between exon 1 and exon 2 of \(C46H11.4c\) and splices with the 3rd exon of \(C46H11.4c\) to form the new transcript \(C46H11.4f\). The three new spliced transcripts \(C46H11.4d\), \(C46H11.4e\) and \(C46H11.4f\) have a 16, 22 and 24 amino acids long first exon and have their new 5’ start 3322 bp upstream, 473 bp downstream and 1438 bp downstream respectively with respect to the parent gene \(C46H11.4c\) (Table 1 in supplementary material). In all spliced transcripts, the last exons remain the same only the N-terminal (5’) encoding exon varies depending upon the splicing pattern. Following the computational predictions of new spliced isoforms, Yuji Kohara’s \(C.\) \(elegans\) EST database was searched for putative EST/cDNA support for possible occurrence of these new exons/transcripts. A search of Yuji Kohara’s \(C.\) \(elegans\) EST database didn't yield any EST match for these new exons, the most probable reason for this being that the current available EST database for \(C.\) \(elegans\) is not adequately represented. Secondly, \(C.\) \(elegans\) sequencing consortium has also made prediction \(C46H11.4a\) which has no conclusive EST/cDNA support for its unique 1st exon present at the 5’end of the transcript. This also supports our hypothesis that only EST data is not good enough for detection of all possible alternatively spliced variants of a gene. NCBI BLAST search was accomplished for finding out homology of these new spliced variants; however, no significant information was available about the prospective similarity with other polypeptides. Due to non-availability of required information for supporting EST/cDNA matches for the newly predicted exons, presence of transcripts were validated by performing RT-PCR using total RNA isolated from the mixed population of \(C.\) \(elegans\).

Experimental confirmation of computationally identified new spliced transcripts encoded by \(lfe-2\) gene \(C46H11.4\)

In order to confirm the possible existence of new transcripts of \(lfe-2\) fertility effector/regulator protein encoded by \(lfe-2\) gene as predicted by bioinformatics tools, RT-PCR amplification was performed. We selected the transcript encoding \(C46H11.4c\) as a control in our experiments. Total RNA, prepared from mixed-stage \(C.\) \(elegans\), was reverse transcribed and the resulting single-stranded cDNA was PCR amplified (as detailed in methodology). Forward and Reverse gene specific primers corresponding to each of the newly predicted exons were designed using a combination of bioinformatics tools and manual methods. These Forward and Reverse primers, specific for a particular exon were successfully able to confirm the occurrence of the spliced transcript by giving a band of anticipated size (Figure 2) when the primer pair specific for that exon was PCR amplified.
Our goal here was to use a novel bioinformatics method capable of delineating all possible spliced transcripts of a gene. Our results therefore, extend the findings of the previous studies that the conventional methods of detection of spliced variants of a gene are not good enough to detect all possible spliced isoforms. So for efficient delineation of all possible putative spliced variants of a gene and to minimize junk predictions (biological noise or false predictions), we propose to combine the ab initio prediction of alternative splicing using a combination of gene/exon finding programmes with experimental validation. These new coding sequences, not annotated or identified earlier will not only add to the available splice data base of *C. elegans* but will also enhance our knowledge about understanding of the genome structure and evolution of higher eukaryotes specially in context to humans. Moreover as now its well established that *C. elegans* are well characterized organisms that have a lot of biological functions that are similar to almost all higher organisms including humans, so based on our work, similar studies can be conducted in several other organisms specially humans with whom *C. elegans* share a close gene homology. Lastly, due to limited domain of our work, further studies using more
advanced techniques like the RNA interference (RNAi) could be taken up which would enhance our knowledge about the biological and functional significance of these spliced transcripts and their possible role in *C. elegans* gene working and regulation.

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**References:**


**Supplementary material**

<table>
<thead>
<tr>
<th>Exon</th>
<th>Amino acid sequence</th>
<th>N-terminal exon size</th>
<th>Splicing pattern</th>
</tr>
</thead>
<tbody>
<tr>
<td>1c</td>
<td>MLEAIYAKIMGVTSR</td>
<td>8</td>
<td>1c-2-3-4-5-6-7-8-Last(CN)</td>
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<tr>
<td>1d</td>
<td>MCLMSNVECKEVNCNGSSNSE</td>
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<td>1d-3-4-5-6-7-8-Last(PR-1)</td>
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<tr>
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<td>1e-2-3-4-5-6-7-8-Last(PR-2)</td>
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<tr>
<td>1f</td>
<td>MNYAHSASEKKSENKLKKKHJAGSSNSE</td>
<td>24</td>
<td>1f-3-4-5-6-7-8-Last(PR-3)</td>
</tr>
</tbody>
</table>

**Table 1:** Amino acid sequence encoded by the predicted alternatively spliced N-terminal exons

Deduced amino acid sequences encoded by alternative first exons (shown in regular font) and the first part of the amino acid sequence encoded by exon 2 or exon 3 (underlined) depending on the splicing pattern; number of amino acids encoded by the N-terminal exon is shown under N-terminal exon size. 1c, 1d, 1e and 1f are the first exons of C46H11.4c (control), C46H11.4d (predicted), C46H11.4e (predicted) and C46H11.4f (predicted) respectively.