

Research

Full-length messenger RNA sequences greatly improve genome annotation

Brian J Haas*, Natalia Volfovsky*, Christopher D Town*, Maxim Troukhan§, Nickolai Alexandrov§, Kenneth A Feldmann§, Richard B Flavell§, Owen White* and Steven L Salzberg*

Addresses: *The Institute for Genomic Research, 9712 Medical Center Drive, Rockville, MD 20850, USA. §Ceres Inc., 3007 Malibu Canyon Road, Malibu, CA 90265, USA.

Correspondence: Steven L Salzberg. E-mail: salzberg@tigr.org

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Abstract

Background: Annotation of eukaryotic genomes is a complex endeavor that requires the integration of evidence from multiple, often contradictory, sources. With the ever-increasing amount of genome sequence data now available, methods for accurate identification of large numbers of genes have become urgently needed. In an effort to create a set of very high-quality gene models, we used the sequence of 5,000 full-length gene transcripts from *Arabidopsis* to re-annotate its genome. We have mapped these transcripts to their exact chromosomal locations and, using alignment programs, have created gene models that provide a reference set for this organism.

Results: Approximately 35% of the transcripts indicated that previously annotated genes needed modification, and 5% of the transcripts represented newly discovered genes. We also discovered that multiple transcription initiation sites appear to be much more common than previously known, and we report numerous cases of alternative mRNA splicing. We include a comparison of different alignment software and an analysis of how the transcript data improved the previously published annotation.

Conclusions: Our results demonstrate that sequencing of large numbers of full-length transcripts followed by computational mapping greatly improves identification of the complete exon structures of eukaryotic genes. In addition, we are able to find numerous introns in the untranslated regions of the genes.

Background

The scientific community has recently witnessed the publication of several large eukaryotic genomes in various stages of completion, including the human genome [1,2], the nematode *Caenorhabditis elegans* [3], the fruit fly *Drosophila melanogaster* [4], and the model plant *Arabidopsis thaliana* [5,6]. Each of these genomes contains over 10,000 genes, and

as scientists attempt to study these genes more closely, the need for accurate gene models becomes increasingly apparent. For high-throughput genome sequencing projects, equally rapid high-throughput genome annotation is necessary, and bioinformaticists use a variety of computational methods to generate this annotation. Despite many improvements in recent years, these computational methods still fall

short of producing correct models for every gene. In order to improve the annotation and facilitate further research, it is essential that we develop methods to identify genes correctly.

Annotation of a gene model should include a precise description of where the genomic DNA sequence is transcribed into messenger RNA, the positions in the mRNA of any and all introns, and the translated protein sequence of the gene. If alternative splice variants are present, these should be annotated as well. Computational methods for genome annotation have several shortcomings that result in the following errors in annotation.

Gene prediction programs predict exon boundaries correctly only about 80% of the time, even for the most intensively studied organisms [7-9]. Thus a gene with five exons will be completely correct only $0.8^5 = 33\%$ of the time, and genes with more exons will be even less likely to be correct. Gene prediction programs also tend to merge and split gene models. Thus two real genes may be merged into one predicted transcript, or vice versa. In addition, programs to align genomic DNA to protein sequences often miss small exons, especially when the homologous proteins are not well conserved. Annotation protocols also tend to miss short genes. For example, recent work has shown that at least one large family of *Arabidopsis* genes encodes a short (80-120 amino acid) protein similar to a secreted polypeptide ligand for a receptor-like kinase that functions in meristems [10]. Most of these were missed in the original, automated annotation of the *Arabidopsis* genome. Alignment programs also make mistakes when genes occur in tandemly repeated copies. Finally, alignment of protein sequence to genomic DNA cannot predict untranslated regions (UTRs), and the leading *ab initio* gene prediction programs (Genscan [11], GlimmerM [12], Genemark.HMM [13]) have great difficulty predicting UTRs; most of them predict only the coding portion of a transcript.

The solution to many of these problems is to identify the complete sequence of the transcribed portions of the genome. Sequencing the mature transcripts (spliced mRNA) solves three major problems: first, it permits accurate identification of the 5' and 3' UTRs. Second, in conjunction with complete genomic sequence, it enables alignment software to identify the precise locations of all introns. Third, it aids in the discovery of new genes.

Results and discussion

We used sequences from 5,000 full-length transcripts sequenced by Ceres, Inc. and released to the public in March 2001 ([14], and at GenBank as accession numbers AY084215-AY089214). We merged this data with a small set of full-length transcripts created in a pilot study, yielding 5,016 complementary DNA (cDNA) sequences in total. As described in detail below, after comparing the cDNA alignments to the

previous annotation, we found that 62% of the matching gene models correctly predicted the exon-intron structure for the gene, 33% needed to be modified, and 5% represented previously undiscovered genes.

We mapped and aligned the cDNA sequences to the five chromosomes of *A. thaliana*. This step is not entirely straightforward, in part because several different programs are available for aligning cDNA to genomic sequence (see Materials and methods). Although the programs are similar, they are not identical, and we conducted a detailed comparison in order to determine which one would produce the best results for these data. Note that the conclusions from this comparison might change for alignments between cDNAs and genomic sequences from different species - for example, alignments between human cDNAs and the mouse genome - where the genome and the cDNAs match less closely than in this study. Because the cDNA sequences are derived from two *A. thaliana* ecotypes (Wassilewskija and Landsberg erecta; see Materials and methods) that differ from the genome's ecotype (Columbia), we expected to find some polymorphisms when we compared the cDNAs to the genome; on average, the three ecotypes are more than 99% identical.

The four programs used for the alignment comparison were sim4 [15], dds/gap2 [16], est_genome [17] and GeneSeqer [18]. Three of these are general-purpose alignment algorithms designed to map expressed sequence tag (EST) or cDNA sequences to a genome. One program, GeneSeqer, is more specialized, in that it has particular subroutines designed to recognize splice sites in *A. thaliana*, and it is able to find AT-AC introns. The other three programs require only that an intron begin with the dinucleotide GT and end with AG. Note that none of these programs is tuned to find AT-AC introns [19] or any other introns with non-consensus splice sites.

The first question we asked in the comparison was how often the four programs produced identical alignments between the cDNA sequences and the genome. Because the programs differed in how they treated the first and last nucleotides of the cDNA, we ignored those nucleotides in deciding if two alignments were identical. The number of identical alignments is shown in Table 1. The table shows that the two programs in greatest agreement with each other were gap2 and GeneSeqer, which agreed on 4,839 out of 5,016 alignments. The total number of cDNAs for which all four programs agreed was 4,124. This initial comparison does not provide a true picture of the extent of agreement, though, because most of the differences were in the alignment at the 5' or 3' ends of the cDNAs.

To determine how well the programs agreed at the ends of the alignments, we compared the lengths of the alignments. Because the cDNAs derive from a different ecotype of *Arabidopsis* than the genomic sequence, there are a few

Table 1

Number of identical cDNA alignments produced by four different programs on a set of 5,016 cDNA sequences

Program	gap2	est_genome	GeneSeqer
sim4	4,819	4,342	4,784
gap2		4,274	4,839
est_genome			4,257

polymorphisms in the UTRs, but even these regions are usually > 98% identical. Differences between ecotypes sometimes interrupt an open reading frame (ORF). The result is that a gene in one ecotype may be a pseudogene, or a severely truncated gene, in the other [20]. We addressed this problem by first aligning cDNAs to the genome, and then always using the genomic sequence to create the predicted protein. Therefore the alignments should span the entire length of the cDNA. Our analysis indicates that gap2 and sim4 performed the best at matching the entire cDNA to the genome, as shown in Table 2.

Two immediate observations from Table 2 are that gap2 produced longer alignments than any of the other programs, and that est_genome produced shorter alignments than all the others. Sim4 and gap2 are closest to each other, disagreeing on alignment length in only 34 cases, with gap2 generating the longer alignment on 32 of those. GeneSeqer has a few more differences than gap2, but is clearly much closer to gap2 than est_genome. Even though est_genome often produces shorter alignments, the unaligned sequence is almost always restricted to a small number of terminal nucleotides.

A more critical question is how many times the programs agreed on the precise locations of all the splice sites for a particular transcript. This question is answered for all pairwise comparisons between the programs in Table 3. Overall, there were 4,918 cDNAs that yielded identical splice sites

Table 2

Comparison of lengths of cDNA alignments for the 5,016 cDNA sequences

	gap2	est_genome	GeneSeqer
sim4	32/34	6/652	31/67
gap2		2/708	3/50
est_genome	706/708		706/733

Entries show the number of sequences for which the program listed along the top produced a longer alignment than the program listed on the left; for example, the entry 32/34 indicates that gap2 produced a longer alignment in 32 cases out of the total 34 for which sim4 and gap2 had alignments of different lengths.

(and therefore identical introns) regardless of which program was used to generate the alignment. As Table 3 shows, the programs disagreed on fewer than 100 alignments, less than 2% of the total. This still leaves open the question of whether one program is clearly superior on these problematic alignments. We therefore evaluated the 98 cDNAs (5,016 - 4,918) for which there was disagreement among the programs to determine the cause of the differences. In 64 of these cases, three programs agree and only one disagrees, while in the remaining 32 cases, there was no alignment shared by a majority of the programs. We looked individually at all 64 cases in which a majority agreed to evaluate whether or not a 'majority wins' rule would always produce the correct result, and came to the following conclusions.

Originally, there were 140 cDNAs for which the programs did not all agree on the locations of the introns. Of these, 106 were cases in which only one program disagreed with the three others. These differences were communicated to the authors of the GeneSeqer program (V. Brendel) and the sim4 program (L. Florea). Both authors identified bugs in their systems, fixed the bugs, and issued new releases, which were then re-run on all the data used in this study. The result was that these two programs are in much closer agreement with gap2 and with each other than previously.

Gap2 disagreed with the other three programs in 24 alignments. The most common reason for the disagreement was an erroneous alignment to non-consensus splice sites (other than GT and AG). Gap2's alignment appeared to be incorrect for all 24 cases. Sim4 disagreed with the other three programs 16 times. Like gap2, in most cases these were due to erroneous non-consensus splice sites. However, in one case, sim4 seems to have found a correct alignment missed by the other three programs. GeneSeqer disagreed with the majority 14 times, sometimes as the result of a tendency to create additional short exons. On the other hand, GeneSeqer is excellent at identifying potential short exons, especially at the termini: in four cases out of 14, GeneSeqer's alignment contained an additional exon that results in a greater percent identity in the overall alignment to the genome. Est_genome disagreed with the other three programs on 10 alignments; in all 10 cases the majority was correct. The mistake in eight of these alignments was that a short exon was missed.

Table 3

Number of cDNA alignments, out of 5,016 total, for which all splice sites are identical

Program	gap2	est_genome	GeneSeqer
sim4	4,946	4,965	4,960
gap2		4,954	4,955
est_genome		4,961	

Overall, it does indeed matter which program is used to align cDNA sequences to genome sequences. Three of the four programs do an excellent job of extending the alignment to cover the full length of the cDNA; only *est_genome* consistently failed to extend the alignments. When the programs disagree, this sometimes indicates that an exon was missed by one or more programs, and manual inspection is necessary to determine the best alignment. Finally, there is a substantial difference in computational speed. *Sim4* is more than 200 times faster than any of the other programs, which can have a significant impact in efforts such as this to align large numbers of sequences. For the searches in this study, the average computation time per cDNA (on an 850 MHz Pentium III computer) for *sim4* was 0.026 second; for *gap2*, 6.4 seconds; for *est_genome*, 9.2 seconds; and for *GeneSeqer*, 12.2 seconds. *GeneSeqer*'s speed per search increases dramatically, approximating the speed of *sim4*, if the cDNAs are submitted as a batch rather than one at a time. In addition, memory requirements make it impossible to run some programs on a standard desktop (*dds/gap2* runs out of memory if one attempts to align a cDNA to a whole chromosome, for example).

Re-annotation of the *Arabidopsis* genome

The alignments generated from the cDNA sequences were used to create new gene models for the corresponding genes in the *A. thaliana* genome. Many of the genes have been manually curated, but many others were created by automated scripts [5,6]. Manual curation is still ongoing.

We used the cDNA alignments to create new gene models automatically according to the following criteria. As described above, there were 4,918 cDNAs for which all alignment programs agreed on the positions of all introns. Using a majority voting scheme for the remaining 98 cDNAs did not always give a correct answer, as discussed above, therefore we used these only after manual inspection. We assume the protein-coding region is the longest ORF on the forward strand, and required it to span at least 40% of the cDNA length. This allowed us to create 4,809 gene models automatically, leaving 109 cDNAs that were inspected manually to determine if they represent RNA genes, pseudogenes or other types of sequence. In one case, cDNA Ceres:104289, the protein-coding region was actually on the opposite strand, corresponding to expressed protein At2g23670, and Ceres:20125 matched the correct strand, supporting the gene annotation. (This could be explained in several ways: as an example of antisense-mediated translational control, as two separate proteins on opposite strands, perhaps expressed in different cell types, or simply erroneous data.) In most of the other cases, the problematic cDNA is either an RNA gene or a likely pseudogene.

Using the alignments from the 4,809 gene models, we updated the annotation of the genome, and evaluated how this had changed the previous annotation. For the vast

majority of genes, 5' and 3' UTRs had not been annotated previously, and these were added with the incorporation of the cDNA data. More interesting is how the protein-coding regions changed. Of the gene models, 2,978 contained identical protein-coding regions to what had already been annotated and required only UTR refinements, but 1,591 were adjusted, yielding more accurate protein sequences. Some of these contain very short 'micro-exons' that are usually missed by *ab initio* gene prediction programs. Perhaps most significant was the addition of 240 completely novel genes not previously included in the *Arabidopsis* genome annotation. Of the 240 novel genes, 92 have significant homology to known proteins, and the rest do not match any previously described proteins. In summary, we found that 62% of the matching gene models further validated the existing exon-intron structure for the gene, 33% needed to be corrected, and 5% represented previously undiscovered genes.

Micro-exons

We also used the cDNA alignments to detect 'micro-exons', very short exons that are typically missed by both gene-finding programs and alignment algorithms. Using new software protocols we developed, we found 47 micro-exons, ranging from 3 to 25 base pairs (bp) in length, distributed evenly across all five chromosomes.

To find micro-exons, we analyzed the results of *sim4* alignments using all 5,016 Ceres cDNAs. *Sim4* identified 36 cDNAs encoding exons of 25 bp or less. In an effort to identify additional micro-exons, *sim4* alignments containing imperfect intron-exon boundaries were examined. We selected only those cases with near-perfect alignments, requiring that all but one or two exons have 100% identity. We then checked to see if the 1-2 exons with slightly lower identity were misaligned as the result of the presence of a small, undetected, exon. We used the 5 bp segments at the boundaries of the exon as probes. If these 5 bp probes mismatched in the original alignment, we searched the adjacent intron (that is, the intron identified by the initial alignment) for short exons that would produce a perfect match with the cDNA. We also required that any new exon would generate introns with a standard GT-AG consensus on either end. This procedure therefore yielded valid exon-intron structures that always improved the identity of the alignment between cDNA and genomic DNA. Figure 1 shows an example of the cDNA alignments before and after inserting a micro-exon.

Using this method, we were able to identify 11 additional micro-exons, all shorter than 12 bp. An extraordinarily short exon of only 3 bp was identified, corresponding to exon 2 of disease-resistance gene *RAR1* (At5g51700). A listing of these micro-exons from all chromosomes is shown in Table 4. Note that in some cases the length of the micro-exons is not a multiple of three; for these, one of the preceding or following exons had its boundary realigned to maintain the

Table 4**Micro-exons from each of the five chromosomes, listed in order of increasing length**

Locus	Gene name	cDNA accession	Exon number	Exon length (nucleotides)	Micro-exon sequence
At5g51700	RAR1	Ceres:99615	2 of 6	3	AG<GGA>GT
At1g63290	D-ribose-5-phosphate-3-epimerase	Ceres:37843	2 of 8	5	AG<GACGG>GT
At3g01850	D-ribose-5-phosphate-3-epimerase	Ceres:2398	2 of 9	5	AG<GACGG>GT
At4g01610	Cysteine protease	Ceres:20761	5 of 11	5	AG<ATCAG>GT
At5g14030	Expressed protein	Ceres:16313.	5 of 6	6	AG<GCCAAG>GT
At2g38880	Putative CCAAT-binding transcription factor subunit	Ceres:7805.	4 of 7	6	AG<TTGGAG>GT
At2g07340	Expressed protein	Ceres:34060.	4 of 6	7	AG<GAAGAAC>GT
At2g41710	AP2 domain transcription factor	Ceres:41462	3 of 9	9	AG<TTTATCTAG>GT
At2g36190	Beta-fructofuranosidase	Ceres:118038	2 of 6	9	AG<ATCCAAATG>GT
At4g13720	Auxin-regulated protein	Ceres:8361	4 of 8	10	AG<GGCCATACAT>GT
At4g29510	Arginine methyltransferase (pam1)	Ceres:38601	2 of 9	11	AG<GAATCCATGAA>GT
At3g55260	Beta-N-acetylhexosaminidase	Ceres:118286	7 of 15	17	AG<GTTTGCCAAAATGAGAG>GT
At1g80380	Auxin-regulated protein	Ceres:117698	3 of 7	18	AG<GTACCTAGGTACAATAAG>GT
At3g55630	Tetrahydrofolylpolyglutamate synthase	Ceres:230791.	6 of 15	18	AG<GAGAAAACCAGCAATGAG>GT
At5g61530	Auxin-regulated protein	Ceres:152557	5 of 10	19	AG<GGAGTTGCCAGCTCAGATG>GT
At5g03880	Auxin-regulated protein	Ceres:37668	4 of 12	19	AG<CTGTCCCTTCTGCCGGAAG>GT
*At4g23470	Expressed protein	Ceres:25694.	7 of 8	19	AG<GGTTTGCATGTATGCAG>GT
At1g67320	Expressed protein	Ceres:116252.	7 of 18	20	AG<TTGAAAACATTTACTACAAG>GT
At3g50210	Flavonol synthase	Ceres:25787.	8 of 12	20	AG<TGGAGCTCACACTGACTATG>GT
At4g37680	Expressed protein	Ceres:262351.	2 of 5	21	AG<GGTTTGTCTTTTCGAAATTCAG>GT
At3g60340	Palmitoyl-protein thioesterase	Ceres:38539.	5 of 12	21	AG<ACATCAGTTGTTTGTGAGAAG>GT
At3g23600	Expressed protein	Ceres:11339.	2 of 7	22	AG<GTTTTGAAGCTCCAAACTTAAG>GT
At5g46030	Expressed protein	Ceres:15222.	4 of 5	22	AG<CTTTGACTGAAGCTATAGACAT>GT
At4g33925	Expressed protein	Ceres:24360.	2 of 5	22	AG<TAACCGAAGAACAGCTCTCAAT>GT
*At4g23470	Expressed protein	Ceres:25694.	2 of 8	22	AG<ATTGTTGCTTCGCGTTGTGGTG>GT
At3g13860	Chaperonin, putative	Ceres:38045.	2 of 17	22	AG<CTCGTCTACTTCCAGGAACTG>GT
At1g73180	Expressed protein	Ceres:108165.	13 of 14	23	AG<TTACTTGAATAAGCACACAGG>GT
At5g09830	Expressed protein	Ceres:37422.	2 of 3	23	AG<GAAGTCATTGACATATCTGGAGG>GT
At2g23930	Small nuclear ribonucleoprotein E	Ceres:4850.	2 of 4	23	AG<GTACATGGATAAGAAGCTCCAAA>GT
At1g66940	Expressed protein	Ceres:110066.	3 of 5	24	AG<AATCTAATATTAGATGGATAATAG>GT
At5g51100	Expressed protein	Ceres:126592.	8 of 9	24	AG<CACGCTTACTATCTGGATTTTGAG>GT
At1g05070	Expressed protein	Ceres:13725.	2 of 3	24	AG<AGCTCAGTAATGCTTCTTTTGTCTG>GT
At2g32580	Expressed protein	Ceres:16625.	2 of 3	24	AG<GACTCAGCAATGGTTCATTCCTG>GT
At2g29960	Cyclophilin	Ceres:19211.	4 of 6	24	AG<AAAAC TTCAGAGCTTTGTGCACAG>GT

Table 4 (continued from the previous page)

Locus	Gene name	cDNA accession	Exon number	Exon length (nucleotides)	Micro-exon sequence
At1g65220	Expressed protein	Ceres:21223.	2 of 8	24	AG<CTCAAAGGAGAAGCCCACTCTCGG>GT
At5g23310	Iron superoxide dismutase 3	Ceres:26637.	7 of 8	24	AG<CACTCTTATTATCTGGACTACAAG>GT
At4g25100	Superoxide dismutase	Ceres:32935.	6 of 7	24	AG<CATGCTTACTACCTTGACTTCCAG>GT
At3g55920	Cyclophilin-like protein	Ceres:94608.	5 of 8	24	AG<AGAACTTTCGGTCACTTTGCACGG>GT
At1g77060	Carboxyphosphoenolpyruvate mutase, putative	Ceres:12293.	4 of 6	25	AG<GACCAAGCATGGCCAAAGAAGTGTG>GT
At4g15900	PRL1 protein	Ceres:123113.	2 of 17	25	AG<CAAGCAGATTCGTCTCAGCCATAAG>GT
At2g47640	Putative small nuclear ribonucleoprotein D2	Ceres:26123.	3 of 6	25	AG<CAAGCCAATGGAAGAGGATACCAAT>GT
At2g41630	Transcription factor IIB (TFIIB)	Ceres:2657.	2 of 7	25	AG<GTTGGGACTTGTTCGCAACTATCAAG>GT
At3g62840	Small nuclear ribonucleoprotein	Ceres:32457.	3 of 5	25	AG<TAAACCAATGGAAGAGGATACCAAC>GT
At2g21270	Putative ubiquitin fusion-degradation protein	Ceres:34470.	4 of 10	25	AG<CCACAAC TTGAAAGTGGTGACAAGA>GT
At3g10330	Transcription initiation factor IIB (TFIIB)	Ceres:38950.	2 of 7	25	AG<GTTGGGACTTGTTCGACCATCAAG>GT
At1g42480	Expressed protein	Ceres:42677.	7 of 9	25	AG<ATTGCTGGAGGAACTGAAGATGAG>GT
At2g23985	Expressed protein	Ceres:252843.	2 of 4	25	AG<TGTCTTGTTCAGGTGAACAAAAAAG>GT

*At4g23470 contains two micro-exons.

available on-line [26] and is also provided as Additional data with this paper online. Figure 2 highlights several interesting examples. In Figure 2a, the alternative 3' splice site on the second intron leads to a shift in the reading frame, producing a different protein sequence. In Figure 2b, alignments of several cDNAs indicate that the last intron is unspliced. Figure 2c shows that different 5' ends lead to differing 5' introns and exons, while not changing the protein sequence in this particular example. Figure 2d shows a centrally located exon that is spliced out along with the surrounding introns. Figure 2e contains three different 5' transcription start sites, three different 3' termination sites, and two unspliced introns in the middle transcript. The unspliced introns occur within exon 2 of GI:14335057, which corresponds to three exons and two introns in both the other transcripts. Note that some of the alternative splicing events occur within the same ecotype.

Neither collection of cDNAs can be considered a random sample of transcripts, and therefore the number of examples of alternative splicing discovered in this data (approximately 10% of the overlapping transcripts) should not be used to extrapolate to the entire genome. The discovery of transcripts with different introns spliced out raises the question of whether the different spliced products are translated and whether the splicing differences reflect

programmed developmental variation or simply splicing errors. It is not possible to answer these questions now, but incomplete splicing and consequential variants in plants have been noted previously to be associated with gene silencing and were postulated to reflect the regulated production of aberrant RNA products not destined to be translated [27]. One clear conclusion is that alternative splicing can be discovered via analysis of cDNAs and genomic sequence, and that a fuller collection of cDNAs will provide a valuable resource for more discoveries about splicing and gene regulation.

Are the sequences full-length?

An independent project to sequence complete *Arabidopsis* cDNAs is ongoing by the SPP consortium [28], using clones created by K. Shinozaki at RIKEN in Japan. These sequences are publicly available from GenBank (search for "RIKEN cDNA Arabidopsis"). These data provided the opportunity to compare the two sets of cDNAs and measure independently how many of them appear to cover the entire length of the predicted mRNA transcript. The sequencing of the RIKEN cDNAs generated 2,996 sequences as of October 2001; we compared these to the 5,016 cDNAs from Ceres and found 1,129 sequences that are contained in both data sets. Of the 1,129 sequences, 941 alignments yield the same exon-intron structure for the underlying gene. We then asked, for each of

Table 5**Alternative acceptor and donor splice sites, alternative 5' exons, and exon skipping examples based on cDNA alignments**

Locus	Gene name	cDNA accessions
Alternative acceptor splice sites		
At3g58710	WRKY DNA-binding protein	Ceres:100465, gi:15991735
At5g35680	Expressed protein	Ceres:11304, gi:14596002
At2g38860	Expressed protein	Ceres:114031, gi:13122287
At4g31550	DNA-binding protein	Ceres:11953, gi:15384214
At1g22700	Expressed protein	Ceres:120133, gi:15294175
At4g30480	Expressed protein	Ceres:12573, gi:14423435
At1g52870	Expressed protein	Ceres:126586, gi:14326544
At5g41810	Expressed protein	Ceres:126660, gi:14532565
At1g63970	Expressed protein	Ceres:15758, gi:11386014
At2g33830	Auxin-regulated protein	Ceres:1711, gi:1127600
At5g20040	IPP transferase	Ceres:19250, gi:14279069
At3g55330	Oxygen-evolving complex subunit	Ceres:21674, Ceres:3747
At1g60850	RNA polymerase subunit	Ceres:21961, gi:514321
At1g76405	Expressed protein	Ceres:23773, gi:13358245
At1g22630	Expressed protein	Ceres:37537, gi:15010607
At1g02500	S-adenosylmethionine synthase	Ceres:37800, gi:15450420
At4g20380	Zinc finger protein Lsd1	Ceres:38456, gi:1872520
At3g54380	Expressed protein	Ceres:38778, gi:14423485
At3g04830	Expressed protein	Ceres:38917, gi:15293268
At1g11840	Lactoylglutathione lyase	Ceres:39107, gi:11094298
At1g02090	COP9 complex subunit	Ceres:40042, gi:3288822
At1g79650	DNA repair protein RAD23	Ceres:40579, gi:14334441
At2g25625	Expressed protein	Ceres:465, gi:14334615
At4g02640	Expressed protein	Ceres:6568, gi:10954094
At3g11930	Ethylene-responsive protein	Ceres:7474, gi:13926249
At2g20820	Expressed protein	Ceres:91872, gi:14190456
At3g09150	Expressed protein	Ceres:98026, gi:13359272
Alternative donor splice sites		
At1g16460	Mercaptopyruvate sulfurtransferase	Ceres:111646, gi:6009982
At5g16540	Zinc finger protein 3	Ceres:113763, gi:4689375
At2g41070	bZIP family transcription factor	Ceres:114632, gi:13346156
At2g36000	Expressed protein	Ceres:123727, gi:14532493
At3g21175	Expressed protein	Ceres:12996, gi:14596058
At5g61880	Expressed protein	Ceres:146274, gi:14517497
At3g14230	RAP2 family protein	Ceres:158240, gi:15450917
At3g03890	Expressed protein	Ceres:18355, gi:14190432
At1g67700	Expressed protein	Ceres:19973, gi:15215605
At3g55630	Tetrahydrofolylpolyglutamate synthase	Ceres:230791, gi:15292866
At2g21620	Expressed protein	Ceres:31655, gi:15320407
At4g10100	Expressed protein	Ceres:35962, gi:6635742
At1g23950	Expressed protein	Ceres:41387, gi:15146261
At1g24260	Floral homeotic protein	Ceres:5055, gi:2345157

Table 5 (continued from the previous page)

Locus	Gene name	cDNA accessions
At2g39730	Expressed protein	Ceres:7114, gi:15450670
At3g07760	Expressed protein	Ceres:7246, gi:15451021
At3g06720	Importin alpha	Ceres:9351, gi:4191743
Alternative 5' exons		
At3g57810	Expressed protein	Ceres:101256, Ceres:29384
At3g03780	Methionine synthase	Ceres:111720, gi:14532771
At5g59890	Actin depolymerizing factor 4	Ceres:11691, gi:15215858
At3g49010	60S ribosomal protein L13	Ceres:12182, gi:15292840
At5g52210	GTP-binding protein	Ceres:16621, gi:1184980
At1g01100	Acidic ribosomal protein	Ceres:24367, gi:15293082
At2g41430	ERD15	Ceres:31388, gi:13926319
At3g08580	Adenylate translocator	Ceres:36818, gi:1433
At5g05000	GTP-binding protein	Ceres:6734, gi:1151243
At3g48880	Expressed protein	Ceres:99337, gi:15028346
Exon skipping		
At5g54940	Translation initiation factor	Ceres:103464, Ceres:32071
At2g46800	Zinc transporter	Ceres:207558, gi:4206639
At5g53860	Expressed protein	Ceres:22860, gi:15215799
At5g27840	TOPP8 Ser/Thr protein phosphatase type-I	Ceres:38656, gi:14596132
At3g23280	Expressed protein	Ceres:41648, gi:15010671
At1g77080	Expressed protein	Ceres:92459, gi:11545544, gi:11545546, gi:13649968

A full list with illustrations and supporting alignment data is available at [26].

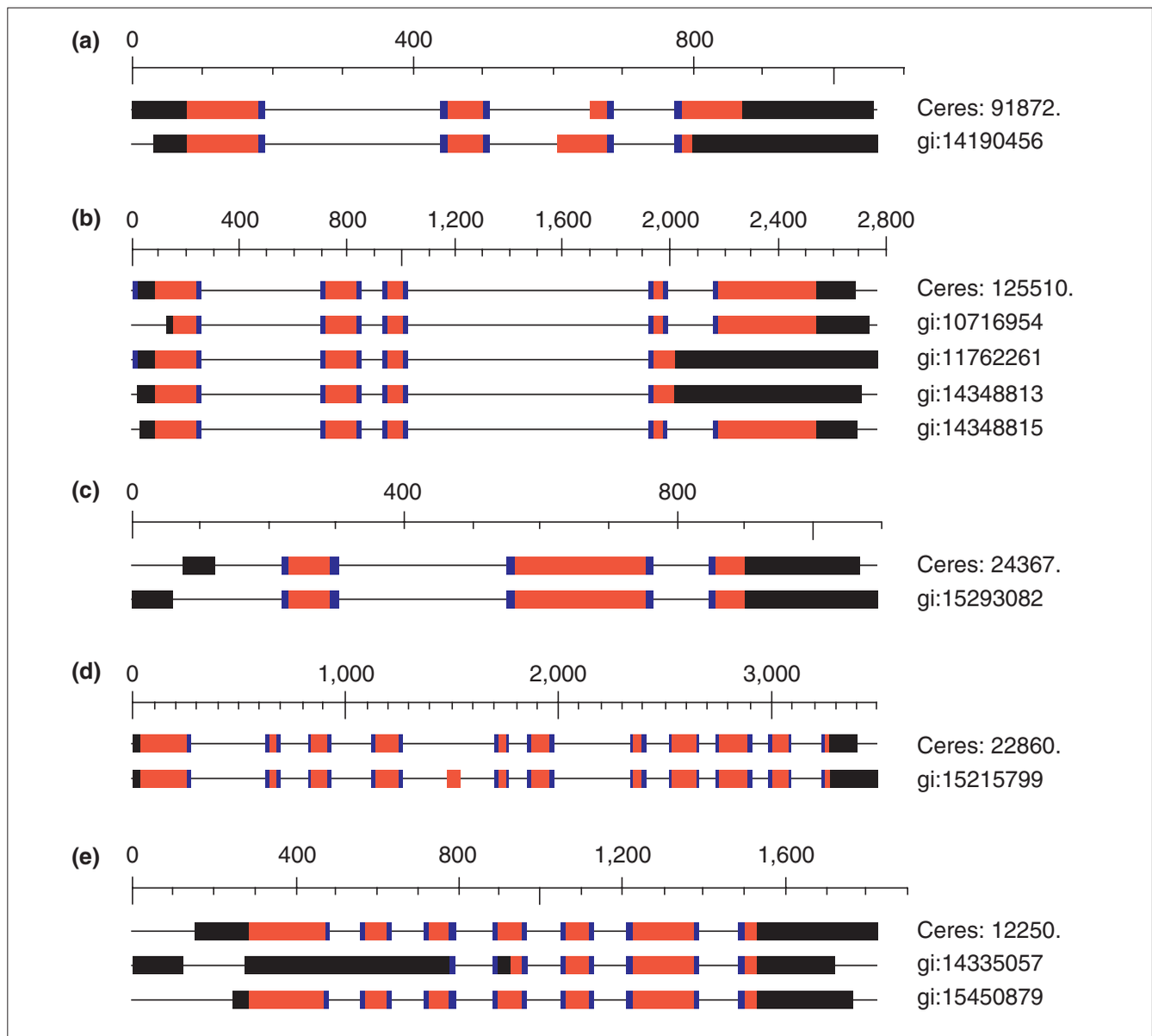
the sequences containing identical introns, do the 5' and 3' ends match, and if not, how large is the difference? The results are illustrated in Figure 3.

Several observations can be made about these results. First, it is important to note that the Ceres clones were selected for full-length sequencing from among a large number of clustered 5' sequences (see Materials and methods), whereas the RIKEN clones were sequenced on the 3' end followed by clustering and selection of a clone for sequencing [29]. The methods for creating the full-length cDNA sequences at both centers involve multiple sequencing runs, followed by assembly of the overlapping sequences. Second, we observed that in the Ceres data, many mRNAs appeared to have two or more putative alternative transcription start sites. This became apparent when different cDNA assemblies were found to overlap exactly except for an extension on the 5' end on one or more clones. It is interesting to note that when the RIKEN clones were longer or shorter on the 5' end, clones of the equivalent length could often be found in the Ceres collection. Multiple clones with the same 5' end provided strong validation that these were truly representative of alternative transcription initiation sites or repeatable artifacts of the

cloning process. Overall, there were 397 Ceres clones that were > 10 bp longer on the 5' end, and 136 RIKEN clones that were longer on the 5' end. If alternative transcription initiation is the correct explanation, then it is relatively common. It is worth noting that in almost all cases, both alternative cDNAs contain complete ORFs.

On the 3' end, the Ceres and RIKEN databases each contained 316 sequences that were >10 bp longer than their match from the other set. If these represent alternative polyadenylation sites or stabilized ends of RNA that get polyadenylated, then these are quite common. Further investigation will be necessary to determine if the 3' end of transcripts truly varies at such a high frequency.

In summary, work described in this study on *Arabidopsis* illustrates the utility of full-length cDNAs for finding alternative splice variants, short exons, UTRs, short genes and alternative transcription start sites. The annotation of eukaryotic genomes is currently an inexact and developing science, and the results described here demonstrate the power of full-length cDNA sequences for improving the quality of multiple aspects of genome annotation.

**Figure 2**

Alternative splice variants discovered by cDNA alignments. Red bars indicate the protein-coding portion of each exon. Black bars indicate noncoding exons and the UTR portions of the initial and terminal exons. Exon boundaries that line up exactly between two or more cDNAs are highlighted in blue. Thin lines connecting the exons represent introns. The genes involved are: **(a)** auxin-regulated protein, At2g20820, chromosome (chr) 2; **(b)** SKP1-interacting partner 5 (SKIP5), At3g54480, chr 3; **(c)** acidic ribosomal protein, At1g01100, chr 1; **(d)** auxin-regulated protein, At5g53860, chr 5; **(e)** unknown expressed protein, At2g45740, chr 2.

Materials and methods

Preparation and sequencing of cDNA

Starting material for cDNA synthesis was polysomal RNA isolated from the top-most inflorescence tissues (ecotype Wassilewskija) and from roots (ecotype Landsberg erecta). RNA from roots of Landsberg erecta was used to construct the libraries because of the availability of high-quality RNA. Nine parts inflorescence to one part root, as measured by wet mass, was used to make three size-fractionated libraries.

Because the ecotypes were mixed before library construction, we cannot determine the source ecotype for any individual cDNA. Polysomal RNA was isolated from a detergent-generated supernatant on a 2 M sucrose cushion. To capture full-length cDNAs, an oligonucleotide is first attached to intact 5' ends, taking advantage of the cap. After first- and second-strand synthesis, the full-length cDNAs were selected, size fractionated and cloned into pBluescript. The ligation mixture was transformed into bacteria, selected

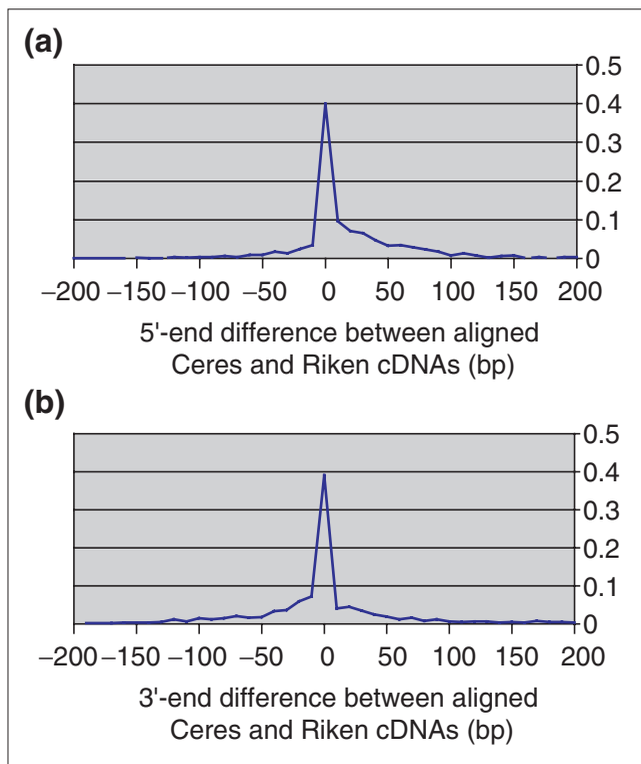


Figure 3
Comparison of the lengths of the 941 cDNAs from the clones that are contained in both the Ceres and RIKEN collections. **(a)** Comparison of the 5'-end difference between Ceres and RIKEN clones; **(b)** comparison of the 3'-end difference between Ceres and RIKEN clones. Peak height indicates the percentage of sequences with a length difference as indicated along the horizontal axis. Positive values on the horizontal axis correspond to longer Ceres clones, while negative values correspond to longer RIKEN clones.

on appropriate antibiotics and picked into 384-well microtiter plates. In repeated rounds of sequencing, several tens of thousands of clones from the three libraries were sequenced from the 5' end, the sequences clustered, and the clone with the longest 5' end in each cluster selected for complete sequencing.

The number of clones sequenced in each round depended on the percentage of new full-length clones that could be obtained from each of the size-fractionated libraries. As the clones reported in this study came from non-normalized libraries, only three rounds of 5' sequencing were employed; 42,000 in the first round, 59,000 in the second round and 22,000 in the final round. Following each round of 5' sequencing, all sequences were clustered using a clustering algorithm that forms separate clusters if there are more than 6 nucleotide differences in any 30-nucleotide window of the match. In this way, clones would not fall into separate clusters simply because of ecotypic differences, different putative transcription start sites or sequencing errors. However, they would cluster separately if alternative splicing

occurred in the first approximately 500 nucleotides and involved more than 6 nucleotides. Following clustering, the clone that was longest on the 5' end was selected for full-length sequencing. If clones were of comparable length on the 5' end, the clone to be sequenced was selected from the library with the highest percentage of full-length clones. Sequencing of 5' ends was performed on capillary sequencers (Molecular Dynamics); full-length sequencing was done on ABI377 sequencers using primer walking. The 5,016 clones analyzed in this study were selected from all full-length clones based on length (> 400 nucleotides), non-redundancy (eliminating alternatively spliced clones), and length of the putative ORF relative to overall clone length.

Alignment of cDNA sequences to the *A. thaliana* genome

Four programs were used to align all 5,016 Ceres cDNA sequences to the *A. thaliana* genome as follows. First, each program was used to align each cDNA sequence to the genome. Some programs cannot efficiently handle a search comparing a cDNA to a 30+ Mb eukaryotic chromosome, and to compensate for those programs, we created a modified procedure that first used BLASTN to identify and extract a region of 20,000 bp surrounding the gene. Each cDNA was aligned to the corresponding 20 kb genome sequence segment using all four programs with default parameter settings. The resulting alignments were then compared automatically to generate the comparison data appearing in the main text. The programs are sim4, available from [30]; dds/gap2, available from [31]; GeneSeqer, available from [32]; and est_genome, available from [33].

Gene models were constructed by first recreating the cDNA sequence using the *Arabidopsis* genome sequence, employing the longest alignment for which all programs predicted identical splice sites. The longest ORF was identified along the forward strand of the cDNA followed by a division of the ORF into protein-coding exon segments and untranslated regions of exons. These constructed gene models were then compared to the existing gene annotation at the mapped genomic region. Previously annotated gene structures that disagreed with the cDNAs were replaced by the cDNA alignment-based gene models, and new gene models were created where pre-existing gene annotations were lacking.

Additional data files

Additional data corresponding to anomalous splicing, including png image files and text-formatted multiple alignments, is available with the online version of this paper.

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32. **GeneSeqer** [<http://bioinformatics.iastate.edu/cgi-bin/gs.cgi>]
33. **est_genome** [http://www.hgmp.mrc.ac.uk/Registered/Option/est_genome.html]