Conserved introns reveal novel transcripts in *Drosophila melanogaster*

Michael Hiller, Sven Findeiß, Sandro Lein, et al.

*Genome Res.* 2009 19: 1289-1300 originally published online May 20, 2009
Access the most recent version at doi:10.1101/gr.090050.108

**Supplemental Material**  
http://genome.cshlp.org/content/suppl/2009/05/21/gr.090050.108.DC1.html

**References**  
This article cites 69 articles, 35 of which can be accessed free at:  
http://genome.cshlp.org/content/19/7/1289.full.html#ref-list-1

Article cited in:  
http://genome.cshlp.org/content/19/7/1289.full.html#related-urls

**Email alerting service**  
Receive free email alerts when new articles cite this article - sign up in the box at the top right corner of the article or click here

To subscribe to *Genome Research* go to:  
http://genome.cshlp.org/subscriptions

Copyright © 2009 by Cold Spring Harbor Laboratory Press
Conserved introns reveal novel transcripts in *Drosophila melanogaster*

Michael Hiller,1,2,9 Sven Findeiß,3,4 Sandro Lein,5 Manja Marz,3 Claudia Nickel,5 Dominic Rose,3 Christine Schulz,6 Rolf Backofen,1 Sonja J. Prohaska,3,4,7 Gunter Reuter,5 and Peter F. Stadler3,4,6,7,8

1Bioinformatics Group, Albert-Ludwigs-University Freiburg, 79110 Freiburg, Germany; 2Department of Developmental Biology, Stanford University, Stanford, California 94305, USA; 3Bioinformatics Group, Department of Computer Science, University of Leipzig, D-04107 Leipzig, Germany; 4Interdisciplinary Center of Bioinformatics, University of Leipzig, D-04107 Leipzig, Germany; 5Institute of Genetics, Biologicum, Martin Luther University Halle-Wittenberg, D-06108 Halle, Germany; 6RNonics Group, Fraunhofer Institut für Zelltherapie und Immunologie–IZI, D-04103 Leipzig, Germany; 7Institut für Theoretische Chemie und Molekulare Strukturbiologie, Universität Wien, A-1090 Wien, Austria; 8Sante Fe Institute, Santa Fe, New Mexico 87501, USA

Noncoding RNAs that are—like mRNAs—spliced, capped, and polyadenylated have important functions in cellular processes. The inventory of these mRNA-like noncoding RNAs (mlncRNAs), however, is incomplete even in well-studied organisms, and so far, no computational methods exist to predict such RNAs from genomic sequences only. The subclass of these transcripts that is evolutionarily conserved usually has conserved intron positions. We demonstrate here that a genome-wide comparative genomics approach searching for short conserved introns is capable of identifying conserved transcripts with a high specificity. Our approach requires neither an open reading frame nor substantial sequence or secondary structure conservation in the surrounding exons. Thus it identifies spliced transcripts in an unbiased way. After applying our approach to insect genomes, we predict 369 introns outside annotated coding transcripts, of which 131 are confirmed by expressed sequence tags (ESTs) and/or noncoding FlyBase transcripts. Of the remaining 238 novel introns, about half are associated with protein-coding genes—either extending coding or untranslated regions or likely belonging to unannotated coding genes. The remaining 129 introns belong to novel mlncRNAs that are largely unstructured. Using RT-PCR, we verified seven of 12 tested introns in novel mlncRNAs and 11 of 17 introns in novel coding genes. The expression level of all verified mlncRNA transcripts is low but varies during development, which suggests regulation. As conserved introns indicate both purifying selection on the exon–intron structure and conserved expression of the transcript in related species, the novel mlncRNAs are good candidates for functional transcripts.

[Supplemental material is available online at www.genome.org. Partial sequences of experimentally confirmed novel transcripts have been deposited in GenBank (http://www.ncbi.nlm.nih.gov/Genbank/) under accession nos. FJ528666–FJ528673 and FJ845365–FJ845382.]

A large portion of the transcriptional output of eukaryotic genomes consists of "mRNA-like noncoding RNAs" (mlncRNAs) (Maeda et al. 2006; The ENCODE Project Consortium 2007). These transcripts are capped, polyadenylated, and often spliced (sometimes alternatively spliced) just like protein-coding mRNAs, but they lack discernible open reading frames. These mlncRNAs are typically much larger than the "housekeeping" RNAs such as transfer RNA (tRNA), small nuclear RNAs (snRNAs), small nuclear RNAs (snRNAs), and they do not seem to have well-conserved secondary structures.

To date, molecular functions have been described only for a small minority of mlncRNAs. They act at different levels of cellular regulatory networks, employing a series of different mechanisms. Some are precursors of small RNAs (Kapranov et al. 2007; Riccardo et al. 2007; Carlile et al. 2008). For example, exons as well as introns of mlncRNAs can be processed into micro RNAs (miRNAs) (Cai and Cullen 2007; He et al. 2008), or snoRNAs (de los Santos et al. 2000; Tycowski and Steitz 2001).

Other mlncRNAs exert their function as large RNAs (Nakamura et al. 1996; Kelley and Kuroda 2000; Ginger et al. 2006). For example, the Evf-2 transcript acts as a transcriptional enhancer for distal-less homeobox genes (Faedo et al. 2004; Feng et al. 2006), related transcripts appear to have a role in the epigenetic regulation of homeotic transcription factors (Dinger et al. 2008), the *hsrw* transcript in the gut is key to the heat shock response in *Drosophila* (Arya et al. 2007), the *NRON* (LOC641373) transcript suppresses the transcription factor NFAT (Willingham et al. 2005), and the HOTAIR transcript is involved in transcriptional repression of the mammalian HOX-D cluster (Rinn et al. 2007). Computational studies provide evidence that a sizeable fraction of mlncRNAs is subject to purifying selection (Ponjavic et al. 2007). Many mlncRNAs are specifically expressed in cell types or developmental stages and may even have a specific localization within cells (Inagaki et al. 2005; Tupy et al. 2005; Ravasi et al. 2006; Mercer et al. 2008), suggesting that their expression is regulated and not due to transcriptional background noise. Furthermore, mlncRNAs are up- or down-regulated in human cancer cells (Calin et al. 2007). Given that transcription of noncoding regions is abundant (Manak et al. 2006), this indicates that the few well-studied examples are only the tip of the iceberg of functionally important noncoding RNAs (ncRNAs) (Amaral et al. 2008).
As a consequence of specific expression and the generally lower expression levels of mlncRNAs compared with their protein-coding counterparts (Ravasi et al. 2006), it is likely that despite these large expressed sequence tag (EST) sequencing projects, we have not yet discovered the complete repertoire of ncRNAs even in the best-studied model organisms. Since a complete inventory of transcripts is a prerequisite for the comprehensive understanding of cellular regulation, a method to uncover mlncRNAs that does not depend on expression data is highly desirable.

While computational methods can identify protein-coding genes with high accuracy in genomic sequences, the identification of ncRNAs is generally more difficult (Eddy 2002). Protein-coding sequences exhibit a clear evolutionary signature (such as an excess of synonymous mutations and reading frame preserving insertions/deletions) that allows their detection by comparative genomics methods (Badger and Olsen 1999; Stark et al. 2007b). However, apart from transcriptional footprints (Glusman et al. 2006), ncRNAs generally lack such characteristic evolutionary patterns. An important subclass, which includes the housekeeping ncRNAs, has evolutionarily conserved secondary structures. These ncRNAs can be identified by methods such as RNAz (Washietl et al. 2005) and Evofold (Pedersen et al. 2006) that search for regions with an excess of mutations that maintain the secondary structure. A few specific classes of ncRNAs, in particular miRNAs and snoRNAs, can be predicted by searching for typical sequence and/or secondary structure motifs (Lai et al. 2003; Hertel and Stadler 2006; Stark et al. 2007b; Hertel et al. 2008). By design, however, all these approaches will miss ncRNAs that are unstructured or whose structure is not well conserved in evolution.

Here, we present a new approach to identify intron-containing mlncRNAs from genomic sequence data alone. Our method exploits characteristic evolutionary signatures of conserved introns. The rationale behind this approach is driven by the observation that intron positions are generally well conserved both in protein-coding RNA and ncRNA genes (Nesterova et al. 2001; Mouse Genome Sequencing Consortium 2002; Rodriguez-Trelles et al. 2006; Ponjavic et al. 2007). For example, the *Drosophila hsr6* ncRNA exhibits a conserved two-exon transcript structure with conserved splice sites but a high exonic sequence divergence (Garbe et al. 1989). Furthermore, changes in the exonic-intron structure are largely independent of the sequence evolution of neighboring exons (Yandell et al. 2006).

The assumption underlying our approach is that a functional pair of donor (5') and acceptor (3') splice sites will be retained over long evolutionary timescales only if (1) the locus is transcribed into a functional transcript, and (2) accurate intron removal is necessary to produce a functional transcript. Thus, conserved introns can be employed to determine the presence of a functional transcript directly from comparative genomics data. The advantage of this approach is that we do not need to make any assumption of the transcript itself. In particular, it is irrelevant whether there is an open reading frame or significant sequence conservation, or whether the functional product arises from the intronic or the exonic part of the transcript. We assume here that the sequence of the intron itself is not conserved; hence, we focus on transcripts whose exonic parts are likely to carry the function, even though we used the intronic part for their detection.

We applied this intron-based approach to 15 insect genomes and reliably predicted novel mlncRNAs. We show that these mlncRNAs are largely unstructured and often not associated with significant sequence conservation, implying that they cannot be predicted by existing methods. Our screen also identified unannotated protein-coding genes and provides a refinement of several gene structures by identifying introns in incomplete coding or untranslated regions (UTRs). Experimental verification succeeded for 18 of 29 tested predictions. Furthermore, we showed that conserved introns imply conserved expression of the surrounding transcript in other species.

### Results

#### Computational identification of spliced RNAs in *Drosophila* genomes

Our approach consists of three steps. First, we predicted introns in individual insect genomes. Second, we used genome-wide alignments to identify orthologous introns, defined here as introns that are independently predicted in at least two genomes and where both donor and acceptor sites are exactly aligned. Third, we compiled a set of evolutionary signatures that are characteristic for introns with conserved splice sites, and use machine learning to distinguish between real and false intron predictions. These steps are illustrated in Figure 1 and are detailed below.

We chose *Drosophila* as a model system to test our approach for several reasons: (1) There is a sufficient number of sequenced insect genomes, which allows comparative genomics methods to annotate features such as protein-coding genes, structured RNAs, and regulatory motifs with high accuracy (*Drosophila* 12 Genomes Consortium 2007; Kheradpour et al. 2007; Stark et al. 2007a, b). (2) The majority (54%) of introns in *Drosophila melanogaster* is not longer than 81 nucleotides (nt). This length was determined as a natural cutoff between long and short introns in the literature (Mount et al. 1992; Lim and Burge 2001) and is used here as well. This simplifies the initial identification of putative introns in the genomic sequence. (3) The short introns in *D. melanogaster* contain basically all the information needed to identify them in pre-mature transcripts (Lim and Burge 2001). In contrast, most mammalian introns are longer (Deutsch and Long 1999), and substantial information about intron positions is contained in exons, complicating their prediction (Dewey et al. 2006).

The intronscan tool predicts short introns in premature transcript sequences by utilizing scores for the donor and acceptor splice sites, the branch point, the intron length and the sequence composition (Lim and Burge 2001). Intronscan detects short (≤81 nt) introns in *D. melanogaster* with an accuracy of sensitivity and specificity of 94%. We benefit from this accuracy in the first step, where we applied intronscan on the entire genomic sequence. (3) The short introns in *D. melanogaster* contain basically all the information needed to identify them in pre-mature transcripts (Lim and Burge 2001). In contrast, most mammalian introns are longer (Deutsch and Long 1999), and substantial information about intron positions is contained in exons, complicating their prediction (Dewey et al. 2006).

The intronscan tool predicts short introns in premature transcript sequences by utilizing scores for the donor and acceptor splice sites, the branch point, the intron length and the sequence composition (Lim and Burge 2001). Intronscan detects short (≤81 nt) introns in *D. melanogaster* with an accuracy of sensitivity and specificity of 94%. We benefit from this accuracy in the first step, where we applied intronscan on the entire genomic sequence. (3) The short introns in *D. melanogaster* contain basically all the information needed to identify them in pre-mature transcripts (Lim and Burge 2001). In contrast, most mammalian introns are longer (Deutsch and Long 1999), and substantial information about intron positions is contained in exons, complicating their prediction (Dewey et al. 2006).

The intronscan tool predicts short introns in premature transcript sequences by utilizing scores for the donor and acceptor splice sites, the branch point, the intron length and the sequence composition (Lim and Burge 2001). Intronscan detects short (≤81 nt) introns in *D. melanogaster* with an accuracy of sensitivity and specificity of 94%. We benefit from this accuracy in the first step, where we applied intronscan on the entire genomic sequence. (3) The short introns in *D. melanogaster* contain basically all the information needed to identify them in pre-mature transcripts (Lim and Burge 2001). In contrast, most mammalian introns are longer (Deutsch and Long 1999), and substantial information about intron positions is contained in exons, complicating their prediction (Dewey et al. 2006).

The intronscan tool predicts short introns in premature transcript sequences by utilizing scores for the donor and acceptor splice sites, the branch point, the intron length and the sequence composition (Lim and Burge 2001). Intronscan detects short (≤81 nt) introns in *D. melanogaster* with an accuracy of sensitivity and specificity of 94%. We benefit from this accuracy in the first step, where we applied intronscan on the entire genomic sequence. (3) The short introns in *D. melanogaster* contain basically all the information needed to identify them in pre-mature transcripts (Lim and Burge 2001). In contrast, most mammalian introns are longer (Deutsch and Long 1999), and substantial information about intron positions is contained in exons, complicating their prediction (Dewey et al. 2006).
First, real splice sites often have substitutions that are consistent with the splice site consensus. For example, the nucleotide position −3 of acceptor splice sites is usually C or T, with G being extremely rare. Consistent with this preference, positive samples show more frequently C/T substitutions than negative samples. Likewise, positive and negative samples differ in the frequency of specific substitutions at positions of the donor splice site and the polyadenylate tract. To capture this information, we determined position-specific scores for all possible substitutions for all species pairs *D. melanogaster* with another species. Pairwise substitution scores are more accurate because they account for species-specific nucleotide preferences at certain splice site positions, which are often correlated to the genomic A+T content (Fig. 2). Summing these scores for the donor (positions +2...+6) and acceptor splice site (positions −7...−3) gives a strong separation of positive and negative samples (Fig. 3). This approach is motivated by the codon substitution frequency model used by Lin et al. (2007) to confirm and reject protein-coding genes.

Furthermore, we observed that most positive samples exhibit a poor sequence conservation in the middle of the intron, while numerous negatives show an atypical high conservation (Fig. 3). Averaging phastCons conservation scores (Siepel et al. 2005) for the intron region +8...+20 and −20...−8 (which occurs in all introns between 40 and 81 nt) gives a highly significant difference between positives and negatives (0.056 vs. 0.425, Wilcoxon P-value < 2.2 × 10−16). This pattern is expected because the middle part of an intron usually contains unconstrained sequence (Halligan and Keightley 2006). Moreover, positive samples show some length variation between species (Parsch 2003), while negatives rarely do. Finally, degenerated positions in the splice site consensus leads to some variation in the sequence-based splice site scores (computed by intronscan) for positive samples, but usually not for negative samples that often have identical splice sites between species.

To combine these features into a single decision (real intron vs. false prediction), we trained a support vector machine (SVM) using randomly selected ~95% of our set for training (22,278 positives and 111,530 negatives). On the remaining unseen 5% of the set, the SVM distinguishes between positive and negative samples with a very high accuracy (area under the receiver operating curve, 0.983) (Fig. 1C). Classifying the unseen part of the set with a probability ≥0.95 as real introns, we obtained a true positive rate of 80% (977 of 1221) at a false-positive rate of 0.12% (seven of 5936). At a probability cutoff of 0.99, we still have a true-positive rate of 71.8% (877 of 1221) at a false-positive rate of 0.07% (four of 5936). We manually examined these four false-positives and provide evidence that three of them are real introns (Supplemental Fig. S1), likely reducing the number of

### Figure 1.
Overview of the computational intron prediction procedure. (A) Introns are predicted using intronscan on both strands of the *D. melanogaster* genome, yielding a total of ~1.4 million predictions. Independent intronscan predictions in the other insect genomes were made. (B) Only those *D. melanogaster* intron predictions are retained that have an orthologous prediction in at least one additional genome. (C) A support vector machine (SVM) classifier based on five features is used to distinguish positive (real introns) and negative training samples (false predictions). These features measure characteristic splice site substitutions, sequence conservation in the middle part of introns, and variation of the intron length, donor, and acceptor score between species. As indicated by the distributions, these features are highly discriminative for positive and negative samples. By using this classifier, we predict 369 conserved introns.
false-positives to just one case. The performance of the SVM is better for intron predictions located in the coding sequence (CDS) compared with introns in UTRs (Supplemental Fig. S2A). The performance difference on predicted introns that are longer or shorter than 60 nt is only marginal (Supplemental Fig. S2B). Overall, this shows that a machine learning approach based on features that capture characteristic intron evolution patterns allows a highly accurate detection of real introns among a set containing only 17% (1221 of 7157) real introns. This high accuracy is advantageous for detecting introns in the rest of the genome, where real introns are likely to be rarer than in this set.

Novel spliced transcripts

We used the SVM to evaluate the 342,785 predictions without an overlap to annotated protein-coding transcripts on the same strand to uncover novel introns and therefore novel transcripts. Using a stringent probability threshold of 0.95, we predict 369 introns (Supplemental Table S1). We searched ESTs and noncoding FlyBase transcripts and found 131 (35.5%) introns where both splice sites are transcript-confirmed, with the rest (238 cases, 64.5%) being currently unconfirmed (six of these 238 have support for either the donor or acceptor site). Of these 238 unconfirmed introns, 44 (18%) are supported by ESTs in other Drosophila species. This indicates that our approach is successful in uncovering novel transcripts. Figure 4 shows examples of confirmed introns belonging to the 5′ UTR of a gene, to an intronic antisense transcript, to a potentially tissue-specific ncRNA, and to a structured ncRNA that represents a precursor for short interfering RNAs (siRNAs) (Okamura et al. 2008).

As real transcripts usually have more than one intron, we expect that unconfirmed introns have a tendency to occur in “clusters” (Fig. 4E), defined here as a 1000-nt genomic region having at least two introns on the same strand. We found 15 clusters with two, two clusters with three, and one cluster with four unconfirmed introns. A simulation (see Methods) shows that this is significantly more than expected (empirical P-value < 0.0001), providing evidence for the existence of multi-exon transcripts.

Introns in novel coding genes and extensions of annotated genes

We observed that predicted introns have a tendency to be located closely upstream of the annotated start of a protein-coding gene. Twenty of the 238 (8.4%) unconfirmed introns are located within 100 nt upstream of a coding gene, but only 0.8% of the predictions classified as false, a 10-fold difference. Conspicuously, genes that have an unconfirmed intron closely upstream tend to have no annotated 5′ UTR (14 of 20 [70%] genes with vs. 77 of 218 [36%] genes without an intron prediction within 100 nt upstream; Fisher's exact test, P = 0.005). As shown in Figure 4A and Supplemental Figure S3, these introns often indicate an incomplete annotation of the 5′ UTR or CDS and help to narrow down the true transcription start site. We observed no comparable enrichment for the 3′ end of genes. This indicates that the annotation of the 3′ end is more complete, which is expected given that large-scale EST projects use poly-A tails to prime the reverse transcription reaction.

Next, we compared all 238 unconfirmed introns with coding gene predictions by CONTRAST (Gross et al. 2007), NSCAN (Gross and Brent 2006), and novel exons predicted by Lin et al. (2007). In addition, we used BLASTX to search for protein homology of the regions flanking these introns. All hits were manually examined. We found that 23 introns likely belong to coding region extensions of annotated genes (Table 1). Another 30 introns overlap gene predictions or BLASTX hits that have no overlap with annotated genes, indicating that these introns likely belong to novel coding genes that are missing in the current D. melanogaster genome annotation (Fig. 5). We conclude that predicted introns close to annotated genes likely belong to incomplete CDS or 5′ UTR. Our predicted introns, together with other annotation tools, thus also extend the catalog of Drosophila coding genes.

Novel spliced ncRNAs

In Table 1, we listed only introns that belong to coding genes with high confidence. To get a set of unconfirmed introns in likely noncoding regions, we used BLASTX again with a low threshold (E ≤ 10^{-5}) and excluded 94 introns (including the 53 introns in Table 1). For the subsequent analysis, we also excluded nine unconfirmed introns that overlap repeats and six introns that are located on chrU, chrUextra, and chr*Het. We note that these predictions can be real, but repeats and these gap-rich genome parts are generally difficult to analyze. The remaining 129 introns are considered to be bona fide mlncRNAs (Supplemental Table S1). Twenty-nine of these 129 (22.48%) have predicted orthologous introns in species outside the Sophophora subgenus (Drosophila virilis, Drosophila mojavensis, Drosophila grimshawi), which indicates
Conserved introns predict novel transcripts

Figure 3. Evaluating characteristic intron evolution. (A) Two predicted introns with orthologous intronscan predictions in other species are shown. The prediction on top exhibits several substitutions in the splice site regions that are characteristic for real introns (e.g., C-to-T substitutions at acceptor position –3). Furthermore, this prediction has a low sequence conservation within the intron (average phastCons score for the region +8...+20 and −20...−8 is only 0.002). This prediction gets a high probability for being a real intron (0.999). In contrast, the prediction at the bottom has substitutions that are inconsistent with intron evolution (e.g., A-to-G substitution at acceptor position −3), and it exhibits conservation throughout the intron (average phastCons score is 0.92). The SVM probability for being a real intron is consequently low (0.001). Positive substitution scores are shown in shades of green; negatives in shades of red. Substitution scores are only considered for the donor (positions +2...+6) and acceptor splice site (positions −7...−3). Note that the substitution scores are specific for each pair D. melanogaster with another species; thus, the same substitution with respect to different species can get different scores. (B) The distribution of the summed substitution scores (left) and the average conservation scores (right) show a substantial difference between our positive and negative samples. The position of the values of the introns from panel A are indicated. For a better visualization, the y-axis for positive and negative samples has a different scale.

exon–intron structure conservation over 63 million years (Myr) of evolution (Tamura et al. 2004).

In contrast to the ncRNAs identified by Tupy et al. (2005), our 129 introns are flanked by regions of rather low sequence conservation (average phastCons scores for the 100 nt up- and downstream flanks are 0.25). Note that this is no indication that the predictions are not real. Indeed, the seven unconfirmed introns that we experimentally verified (see below) show an even lower flank conservation (average 0.21). A large fraction of these 129 introns overlap coding genes in antisense direction (41 of 129; 32%); however, this is not surprising given that almost half of the D. melanogaster genome is covered by exons and introns of coding genes and the fact that many genes overlap each other on opposite strands (Misra et al. 2002).

Novel mlncRNAs are mostly unstructured

Our screen identified two introns located in known mlncRNAs with extensive secondary structures (pncr009:3L, CR32205; Fig. 4D) that function as siRNA precursors (Okamura et al. 2008). To test if our predictions are associated with conserved secondary structures, we applied RNAz to the regions flanking the 129 introns. We obtained two (1.6%) predictions of conserved secondary structures. Since RNAz has a certain false-positive rate, we used two control sets to test for enrichment or depletion of conserved structures. Five thousand randomly selected genome regions and their shuffled versions show a highly similar percentage of RNAz hits. Together with the observation that >98% of these mlncRNAs are not associated with conserved secondary structures, this indicates that our method mostly predicts unstructured mlncRNAs, which cannot be identified by RNAz and related methods.

Experimental verification of predicted mlncRNAs

We used RT-PCR with primers designed to flank the predicted intron to validate expression of the corresponding transcripts in five different developmental stages of D. melanogaster: embryo, larva, pupa, male, and female. We counted as a positive verification only those introns where the transcript is spliced and sequencing confirms the correctness of both splice sites.

We tested 12 introns that likely belong to mlncRNAs, and could verify seven (58%) of them (Fig. 6). We named these seven mlncRNAs according to their genomic location (cytogenic band). The expression level of all transcripts is low, consistent with previous findings of low expression levels of mlncRNAs (Ravasi et al. 2006). Only two of the seven mlncRNAs can be found in all five tested conditions (mlncRNA102B1 and mlncRNA42E5-2). The other five show variation in the expression pattern during development, which suggests that their expression is controlled. For example, mlncRNA36C10 is found only in embryos, mlncRNA6E83 shows only a weak expression at the pupal stage, and mlncRNA66A2 is expressed in all five conditions except for the embryo stage. MlncRNA69E3 shows two bands on the gel due to usage of an alternative acceptor splice site (Supplemental Fig. S4), and our predicted intron corresponds to the longer transcript. ESTs downstream of mlncRNA102B1 indicate that this mlncRNA
belongs to a longer *D. melanogaster* transcript and the orthologous *Drosophila yakuba* intron is confirmed by a four-exon transcript (Supplemental Fig. S5). Next, we tested 17 unconfirmed introns located in putative coding transcripts, and verified 11 (65%) of them (Supplemental Table S2). This shows that novel mlncRNA and coding transcripts are roughly equally likely to be verified in our experimental setup.

Overall, we verified 62% (18 of 29) of our tested predictions. As in all transcriptomic studies, this percentage represents a lower bound as we miss transcripts expressed in other conditions or at expression levels below our sensitivity.

**Conserved introns are predictive of conserved transcript expression in other *Drosophila* species**

Our approach is based on the assumption that conserved introns should be indicative of conserved transcripts, which should therefore be expressed in other *Drosophila* species. To test this, we determined whether the regions orthologous to the seven novel mlncRNAs are expressed in *Drosophila simulans*, *Drosophila erecta*, and *Drosophila pseudoobscura*, and whether the orthologous introns are correctly spliced. We used RT-PCR with primers specific for these three species and confirmed splicing of the transcript and correctness of both predicted splice sites by sequencing.

We found that five of the seven transcripts are expressed in *D. erecta* and *D. pseudoobscura* (Fig. 6; Supplemental Table S3). All seven transcripts are expressed in the closely related *D. simulans*. Expression of orthologous transcripts was detected in *D. pseudoobscura* despite the observation that the sequence up- and downstream of the intron has little similarity to *D. melanogaster* (Supplemental Fig. S6). It is noteworthy that the alternative acceptor of mlncRNA69E2 is used in *D. simulans* and *D. erecta* as well. Consistent with the absence of this alternative splice site in *D. pseudoobscura* (Supplemental Fig. S4), we detect only one *D. pseudoobscura* transcript. These results show that novel noncoding
Table 1. Predicted unconfirmed introns that likely belong to coding transcripts

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Locus and strand</th>
<th>SVM probability</th>
<th>Intron supported by</th>
<th>Supported splice site(s)</th>
<th>EST confirmed in*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3R</td>
<td>16418216–16418278</td>
<td>– 1</td>
<td>CONTRAST</td>
<td>Both</td>
<td>D. erecta, D. grimshawi</td>
</tr>
<tr>
<td>X</td>
<td>7223271–7223350</td>
<td>+ 1</td>
<td>CONTRAST</td>
<td>Both</td>
<td>D. simulans, D. yakuba, D. erecta</td>
</tr>
<tr>
<td>3L</td>
<td>4473080–4473137</td>
<td>+ 1</td>
<td>CONTRAST, NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>14754827–14754883</td>
<td>– 1</td>
<td>CONTRAST, NSCAN</td>
<td>Both</td>
<td>D. grimshawi</td>
</tr>
<tr>
<td>3L</td>
<td>18673369–18673426</td>
<td>+ 1</td>
<td>EDW46531, EDX10977b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>15234878–15234933</td>
<td>+ 1</td>
<td>EDW49469, EDX12334b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>4596738–4596798</td>
<td>– 1</td>
<td>EDX06217, EDV95486b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>21742889–21742942</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>9465182–9465246</td>
<td>– 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>18194675–18194735</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>16207852–16207912</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td>D. sechellia</td>
</tr>
<tr>
<td>3L</td>
<td>13129956–13130018</td>
<td>+ 1</td>
<td>EDW90731b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>3042603–3042664</td>
<td>+ 1</td>
<td>EDW81806b</td>
<td>Both</td>
<td>D. ananassae, D. viriliis</td>
</tr>
<tr>
<td>2L</td>
<td>17982366–17982435</td>
<td>– 1</td>
<td>CONTRAST, NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>16193374–16193427</td>
<td>+ 1</td>
<td>EDV52216b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>15663802–15663855</td>
<td>+ 0.99</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2L</td>
<td>19966361–19966419</td>
<td>+ 0.97</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>4972107–4972166</td>
<td>+ 0.96</td>
<td>CONTRAST</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>3803685–3803739</td>
<td>– 0.95</td>
<td>EDV55288b</td>
<td>Both</td>
<td>D. simulans</td>
</tr>
<tr>
<td>3L</td>
<td>11951423–11951472</td>
<td>+ 1</td>
<td>Acceptor</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>5959006–5959065</td>
<td>+ 1</td>
<td>Acceptor</td>
<td>Both</td>
<td>D. pseudoobscura, D. willistoni</td>
</tr>
<tr>
<td>X</td>
<td>16532209–16532265</td>
<td>+ 0.96</td>
<td>NSCAN</td>
<td>Both</td>
<td>Acceptor</td>
</tr>
<tr>
<td>2L</td>
<td>8973537–8973596</td>
<td>– 0.96</td>
<td>CONTRAST, NSCAN</td>
<td>Both</td>
<td></td>
</tr>
</tbody>
</table>

Introns in potentially novel coding genes

<table>
<thead>
<tr>
<th>Chromosome</th>
<th>Locus and strand</th>
<th>SVM probability</th>
<th>Intron supported by</th>
<th>Supported splice site(s)</th>
<th>EST confirmed in*</th>
</tr>
</thead>
<tbody>
<tr>
<td>3L</td>
<td>16099003–16099075</td>
<td>+ 1</td>
<td>EDW41673, EDX10642b</td>
<td>Both</td>
<td>D. sechellia, A. gambiae</td>
</tr>
<tr>
<td>2L</td>
<td>14782479–14782524</td>
<td>+ 1</td>
<td>EDW51777, EDX05070b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>18672140–18672192</td>
<td>+ 1</td>
<td>EDW94594, EDV1848b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>14499932–14499987</td>
<td>– 1</td>
<td>EDW48482, EDX07699b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2L</td>
<td>14782831–14782890</td>
<td>– 1</td>
<td>EDV57971b</td>
<td>Both</td>
<td>D. mojavensis</td>
</tr>
<tr>
<td>2L</td>
<td>13119784–13119844</td>
<td>– 1</td>
<td>EDV58744b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>13119936–13119996</td>
<td>+ 1</td>
<td>EDV58744b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>7751812–7751876</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td>D. sechellia, D. yakuba</td>
</tr>
<tr>
<td>2R</td>
<td>12268491–12268550</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td>D. erecta</td>
</tr>
<tr>
<td>3L</td>
<td>15710127–15710178</td>
<td>+ 1</td>
<td>DAA03006, EDW94927b</td>
<td>Both</td>
<td>D. simulans</td>
</tr>
<tr>
<td>2L</td>
<td>17486655–17486729</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>16360252–16360331</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>15710335–15710391</td>
<td>+ 1</td>
<td>DAA03006, EDW94927b</td>
<td>Both</td>
<td>D. simulans</td>
</tr>
<tr>
<td>2R</td>
<td>19040486–19040544</td>
<td>– 1</td>
<td>NSCAN</td>
<td>Both</td>
<td>D. yakuba</td>
</tr>
<tr>
<td>2R</td>
<td>7732172–7732229</td>
<td>+ 1</td>
<td>NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>15709741–15709795</td>
<td>+ 0.99</td>
<td>DAA03006, EDW94927b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3L</td>
<td>1164863–1164930</td>
<td>+ 0.99</td>
<td>EDX1011b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>15212035–15212094</td>
<td>+ 0.99</td>
<td>3R152111832, 3R15212093</td>
<td>Both</td>
<td>D. grimshawi, D. mojavensis</td>
</tr>
<tr>
<td>3L</td>
<td>18671860–18671916</td>
<td>+ 0.99</td>
<td>EDV51848b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2L</td>
<td>1789294–1789351</td>
<td>+ 0.99</td>
<td>CONTRAST, NSCAN</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>16579653–16579707</td>
<td>+ 0.98</td>
<td>DAA02671b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>18595958–18596014</td>
<td>– 0.98</td>
<td>DAA04274b</td>
<td>Both</td>
<td>D. simulans</td>
</tr>
<tr>
<td>3L</td>
<td>15709941–15709994</td>
<td>+ 0.98</td>
<td>DAA03006, EDW94927b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>14269140–14269201</td>
<td>– 0.98</td>
<td>CONTRAST</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>3R</td>
<td>6496222–6496293</td>
<td>+ 0.97</td>
<td>DAA03590b</td>
<td>Both</td>
<td>D. sechellia</td>
</tr>
<tr>
<td>3L</td>
<td>18061971–18062023</td>
<td>+ 0.97</td>
<td>EDV51895, EDV46481b</td>
<td>Both</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>15211777–15211831</td>
<td>+ 1</td>
<td>3R15211832</td>
<td>Accceptor</td>
<td>D. simulans, D. grimshawi</td>
</tr>
<tr>
<td>2R</td>
<td>8318195–8318255</td>
<td>+ 0.99</td>
<td>2R7946658b</td>
<td>Accceptor</td>
<td></td>
</tr>
<tr>
<td>2R</td>
<td>1150369–1150320</td>
<td>– 0.99</td>
<td>CONTRAST</td>
<td>Donor</td>
<td></td>
</tr>
<tr>
<td>2L</td>
<td>2287620–2287681</td>
<td>– 0.97</td>
<td>EDV53978, EDX03445b</td>
<td>Donor</td>
<td>D. sechellia, D. yakuba</td>
</tr>
</tbody>
</table>

*The orthologous intron is EST confirmed in other insects.
**BLASTX hit that flanks exactly the intron; NCBI protein identifier is given.
***congochr* identifier of novel exons from Lin et al. (2007).
transcripts that are expressed in several Drosophila species can be predicted from genomic sequence alone by searching for conserved introns.

Discussion

Eukaryotic cells produce a diverse class of intron-containing transcripts including protein-coding and noncoding ones. Protein-coding genes and particular classes of ncRNAs such as tRNAs, miRNAs, and snoRNAs can be identified by comparative genomics methods. However, the detection of mlncRNAs currently relies solely on experiments (Inagaki et al. 2005; Tupy et al. 2005) since no characteristic features are known that allow their prediction.

We describe here a method that predicts intron-containing transcripts by making use of evolutionary characteristics of conserved introns and the observation that introns rarely turnover or shift with respect to the exons. It is important to note that we solely use intron information for predictions. This allows us to identify (1) protein-coding transcripts (including UTRs thereof) as well as mlncRNAs; (2) transcripts without conserved secondary structures; and (3) transcripts without evolutionarily conserved sequences (see Fig. 4). The latter property is important as functional ncRNAs do not necessarily have significantly conserved sequences (Panga et al. 2006). For example, the XIST ncRNA has a clear function in X chromosome inactivation in mammals (Kelley and Kuroda 2000); however, a comparison of human and mouse XIST reveals a low overall sequence identity (Nesterova et al. 2001).

While our approach is unbiased with respect to these characteristics, it has limitations. Our method predicts only a partial transcript structure, which in general will have to be completed by experimental approaches such as full-length cDNA sequencing. However, gene prediction algorithms that predict only the CDS and high-throughput transcriptomic techniques suffer from the same problem. Here, we focused on short introns in D. melanogaster; consequently, transcripts containing exclusively longer introns cannot be predicted. It remains unclear whether longer introns and whether short introns in other species are predictable in a similar way.

Furthermore, we currently classify introns with a conserved intron body as false, because the great majority of real introns shows no sequence conservation in the middle. Thus, introns overlapping other functional elements such as putative promoter elements (Supplemental Fig. S7), introns that are miRNA precursors (Berezikov et al. 2007; Okamura et al. 2007), or retained introns that encode a protein domain (Hiller et al. 2005) are unlikely to be predicted.

Apart from the motivation to identify novel mlncRNAs, we aimed at predicting putatively functional mlncRNAs as opposed to transcriptional noise. Despite the observation that our predictions are generally not associated with strong sequence conservation, the detection of a conserved intron indicates that the exon–intron structure is under purifying selection and that the failure to correctly excise the intron likely affects the function of the transcript. Consistent with this, mlncRNA sequences, their splice sites, and promoters show reduced substitution, insertion, and deletion rates indicative of purifying selection (Ponjavic et al. 2007). Furthermore, we showed that conserved introns imply that the respective transcripts are expressed in other flies. While conserved exon–intron structure and conserved expression indicate function, the specific functional aspects of these mlncRNAs have to be addressed in future studies.

Methods

Genomes, alignments, and data

We downloaded the following genome assemblies, alignments, and annotation tracks from the UCSC genome browser (Karolchik et al. 2008): D. melanogaster (DM3), D. simulans (droSim1), Drosophila sechellia (droSec1), D. yakuba (droYak2), D. erecta (droEre2), Drosophila ananassae (droAna3), D. pseudoobscura (DP4), Drosophila persimilis (droPer1), Drosophila willistoni (droWil1), D. virilis (droVir3), D. mojavensis (droMoj3), D. grimshawi (droGrt1), A. gambiae (anoGam1), T. castaneum (triCas2), A. mellifera (apiMel3), genome-wide pairwise alignments of D. melanogaster with another insect produced by BLASTZ (Schwartz et al. 2003), FlyBase and RefSeq coding transcripts (24,090 transcripts), and FlyBase noncoding transcripts (1108 transcripts). The sequence conservation was quantified by the phastCons scores (Siepel et al. 2005) for a multiple alignment of these 15 insects. ESTs and mRNAs were downloaded from the UCSC genome browser or from GenBank.
Orthologous introns

To determine a set of *D. melanogaster* introns that have orthologous intronscan predictions, we obtained from the pairwise alignments for each intron the genomic locus (chr:start-end) and the strand in another species. Then, we checked if exactly this locus and strand correspond to an intronscan prediction in the respective species. Note that both splice sites have to be exactly aligned to get an orthologous intron, which makes this step sensitive to alignment errors. We discarded 900,708 predicted *D. melanogaster* introns without orthologous predictions, leaving 498,231 predictions that have between one and 14 orthologous predictions.

SVM training

To get a training set, we obtained the subset of 155,446 predictions that overlap a protein-coding FlyBase and/or RefSeq transcript. We compared these predicted introns to the exon–intron annotation of the FlyBase/RefSeq transcripts and found 23,499 predictions where both splice sites are annotated (real introns, positive samples), 14,481 predictions where only one splice site (donor or acceptor site) is annotated (these were omitted in the following), and 117,466 predictions where no splice site are annotated (negative samples). All positive and negative samples were combined into our test set. To get an unseen part of the test set, we randomly extracted ~5% (1221 positive and 5936 negative samples). The remaining 22,278 positives and 111,530 negatives were used as the SVM training set and for substitution score calculations.

We used the following five features: (1) the sum of all pairwise substitution scores for the region +2...+6 (positions are relative to the donor site) and −7...−3 (positions are relative to the acceptor site), (2) the average phastCons score of the region +8...+20 and −20...−8, (3) the median of all intron length differences, (4) the median of all differences of donor splice sites scores (computed by intronscan), and (5) the median of all differences of acceptor sites scores.

For feature 1, we first computed species-specific log-odds scores for each of the 10 splice site positions (+2...+6 and −7...−3) and each possible nucleotide pair (x, y) by computing \( \log[f_{\text{pos}}(x, y)]/f_{\text{neg}}(x, y) \). Here, nucleotide x occurs in *D. melanogaster* and nucleotide y occurs in one of the other 14 species (11 Drosophila species, A. gambiae, T. castaneum, An. mellifera). We write \( f_{\text{pos}} \) and \( f_{\text{neg}} \) for the frequencies of an \((x, y)\) pair in positive and negative samples, respectively. These log-odds scores were summed up for each of the 10 splice site positions (+2...+6 and −7...−3), considering only substitutions, i.e., disregarding unchanged positions. The more substitutions are consistent with splice site evolution, the higher is the total substitution score. We do not divide this sum by the number of orthologous introns since this implicitly captures the fact that positive samples have more orthologous predictions than negatives (medians 9 vs. 2).

Feature 2 captures that positive samples usually have poor sequence conservation in the middle part, leading to lower phastCons scores compared to negatives. We restrict this average to the region +8...+20 and −20...−8 since this region represents the middle part of the smallest introns in our set, which are 40 nt long. The differences in features 3 to 5 were always computed *D. melanogaster* vs. the orthologous intron of another species. These three features capture sequence variation at the splice sites and variation of the intron length in evolution.

We used libsvm 2.85 (http://www.csie.ntu.edu.tw/~cjliu/libsvm/) to train a radial basis kernel on input data consisting of the five features. For training and testing, all input features were

**Figure 6.** Experimentally verified introns in mlncRNA transcripts. The expression of the spliced transcript was tested in embryo (E), larva (L), pupa (P), male (♂), and female (♀) stages. Ethidium bromide–stained agarose gels show the RT-PCR results for *D. melanogaster* (D.mel) data. Genomic DNA (gen.) was used as a PCR control, and size standards (L) were run on a 100-bp Ladder (M). PCR products were visualized by sequencing.

**Intron prediction with intronscan**

We downloaded the intronscan program (Lim and Burge 2001) from the investigators’ website ftp://hollywood.mit.edu/pub/intronscan.tar.gz and applied it to the forward and reverse strand of the entire genome of all species. We set the minimum and maximum intron length to 40 and 81 nt, respectively; used a cutoff for the overall score of 11; and used parameters for intron length (-intlen), intron composition (-intcomp), and the branch site model with a branch site probability of 0.5 (-br-pB 0.5).
scaled to the interval \([-1,1]\) using svm-scale. Since training the SVM kernel on the entire training set is computationally very expensive due the large size (133,808 introns), we randomly extracted 5000 predictions and trained the kernel on them.

We extensively tested other features (such as average and position-specific phastCons scores of the donor and acceptor splice sites, standard error/deviation of donor/acceptor site scores and intron length, extending the substitution scores to positions further away from the splice sites) but found no significant improvement by adding more features or other feature combinations. It should be noted that extracting a different unseen 5% part of the test set and training on the remaining 95% leads to a similar performance.

**Clustering**

To determine if real introns have a tendency to occur in clusters (defined here as a 1000-nt region having at least two introns on the same strand), we used a simulation test. We controlled for the different numbers of positive and negative introns since a higher number of introns will always result in more clusters. To this end, we randomly selected from the total set of negatively predicted introns as many as there are unconfirmed positive introns (total of 238) requiring that from each pair (chr, strand) the exact number of observed positive introns is selected. We repeated this 10,000 times and counted the number of clusters in each iteration. In those 10,000 iterations, we observed a maximum number of five clusters with two introns and no cluster with three or more introns. Compared with the 15 clusters with two, two clusters with three, and one cluster with four unconfirmed introns, this gives an empirical P-value of 0.0001.

**Experimental verification**

We did verification experiments for a set of 29 predicted introns. Primers were designed for the region 100 nt up- and downstream of each intron using FastPCR, version 5.2.118 (http://www.biocenter.helsinki.fi/bi/programs/fastpcr.htm), with the following design options: length of 20–24 nt, 55°C–62°C annealing temperature, 45–65 GC%, overlapping primers, primer quality control, primers low complexity control, and synchronizing melting temperature for primer pair ±4°C (default for all other options). To check nonspecific primer matches, we used a BLAST (E-value 1) against the whole genome, excluding the regions flanking the intron. Since the 3’ end of a primer has to fit nearly perfectly, we kept only BLAST hits having at least 16 matches together with an 18th primer–nucleotide match. Then, we discarded all primer pairs where the forward and reverse primer have a nonspecific match at the same chromosome arm. Supplemental Table S2 lists the primers for all tested introns.

Total RNA from *D. melanogaster* embryos, larvae, pupae, males, and females was extracted with TRIzol (Invitrogen). After treatment with RNase-free DNAse (Promega), total RNA was reverse transcribed with M-MLV reverse transcriptase (Promega). Standard fly miniprep was performed to isolate genomic DNA. Predicted primer pairs were used for PCR with PCR SuperMix (Invitrogen). PCR products were analyzed by agarose gel electrophoresis on 1.2% agarose gels, cloned in pGEM-T (Promega) and sequenced with the ABI PRISM dRhodamine Terminator Cycle Sequencing Kit (Applied Biosystems).

**Introns in protein-coding genes**

We downloaded the CONTRAST and NSCAN annotation from the UCSC genome browser. Annotations of predicted novel exons (Lin et al. 2007) were downloaded from http://www.broad.mit.edu/~mlin-fly_genome/ and converted to DM3 using the UCSC liftover tool (http://genome.ucsc.edu/cgi-bin/hgLiftOver). We compared the loci of our introns with these annotations to find cases where one or both splice sites are exactly consistent with these predictions.

We also used the NCBI BLASTX program (Altschul et al. 1990) to search the database of nonredundant protein sequences (nr) using an E-value threshold of $1 \times 10^{-5}$. As query sequences we used a 200-nt sequence consisting of 100 nt from the upstream and 100 nt from the downstream intron flank. In addition, we downloaded the CDS for *D. simulans*, *D. scelidilia*, *D. yakuba*, and *D. erecta* from http://tana.lnb.org/~venkyy/AAA/freeze_20061030/protein_coding_gene/GLEANR/sequence/cds and used BLAT (Kent 2002) to map these sequences to the *D. melanogaster* genome. All hits were manually inspected. While we did not generally exclude introns on chrU, chrUextra, and the heterochromatin regions (chr*Het) from our analysis, we excluded these introns from Table 1 since we are often not able to confirm that the protein annotation uniquely refers to this region.

**Discarding repeats**

We downloaded the RepeatMasker annotation of *D. melanogaster* from the UCSC genome browser excluding simple repeats and low complexity regions. We discarded introns that overlap a repeat with at least 10%.

**RNAz**

We retrieved sequence 60 nt upstream of the donor and 60 nt downstream of the acceptor site of predicted introns. After aligning these regions using ClustalW, RNAz (Washietl et al. 2005) was applied to analyze the secondary structure conservations of the resulting alignments. The concatenated 120-nt sequence fragments can be interpreted as potential splice products; 120 nt is the default input sequence length of RNAz. Several alignment preprocessing steps (control for appropriate alignment lengths [>50 nt], valid gap patterns [<25% gap characters], and a base composition inside the definition range of RNAz [GC-content >0.25 and <0.75]) were performed using the Perl script mazWindows.pl, which is part of the RNAz package (Washietl et al. 2005). By optimizing the latter features, this script maximally chooses six sequences out of an n-way alignment to match the RNAz training scope.

As a control, we used the alignments for 5000 randomly selected genome regions and obtained 118 (2.3%) RNAz hits. After randomizing these 5000 regions by column-wise shuffling, RNAz predicted 54 (1.1%) hits. Both percentages are not significantly different from the two hits for the 129 introns (Fisher’s exact test, $P = 0.77$ and $P = 0.65$, respectively).

**Data availability**

The genomic coordinates for the DM3 assembly, the strand, and the SVM probability of all 369 predicted introns, the 129 unconfirmed introns in putative mlncRNAs, and the 94 unconfirmed introns in putative novel coding transcripts are available in Supplemental Table S1. http://cs.stanford.edu/people/hillerm/Data/Introns_mlncRNA/ contains links to load these tables into the UCSC genome browser to view them in the context of other genome annotations. Partial sequences of experimentally confirmed novel transcripts have been deposited in GenBank under accession numbers FJ528666–FJ528673 and FJ845365–FJ845382.
Acknowledgments

We thank Aaron Wenger for critical reading of the manuscript. This work was supported by the German Research Foundation (STA 850/7-1 [under the auspices of SPP-1258 “Sensory and Regulatory RNAs in Prokaryotes”] and Hi 1423/2-1), the Graduiertenkolleg Wissensrepräsentation at the University of Leipzig, SF6/10, the European Network of Excellence “The Epigenome,” and the 6th Framework Programme of the European Union (project SYNET-LET, 043312).

References


Received December 8, 2008; accepted in revised form March 18, 2009.