A network of conserved co-occurring motifs for the regulation of alternative splicing

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ABSTRACT
Cis-acting short sequence motifs play important roles in alternative splicing. It is now possible to identify such sequence motifs as conserved sequence patterns in genome sequence alignments. Here, we report the systematic search for motifs in the neighboring introns of alternatively spliced exons by using comparative analysis of mammalian genome alignments. We identified 11 conserved sequence motifs that might be involved in the regulation of alternative splicing. These motifs are not only significantly overrepresented near alternatively spliced exons, but they also co-occur with each other, thus, forming a network of cis-elements, likely to be the basis for context-dependent regulation. Based on this finding, we applied the motif co-occurrence to predict alternatively skipped exons. We verified exon skipping in 29 cases out of 118 predictions (25%) by EST and mRNA sequences in the databases. For the predictions not verified by the database sequences, we confirmed exon skipping in 10 additional cases by using both RT-PCR experiments and the publicly available RNA-Seq data. These results indicate that even more alternative splicing events will be found with the progress of large-scale and high-throughput analyses for various tissue samples and developmental stages.

INTRODUCTION
Alternative splicing is a cellular process that provides distinct, context-dependent isoforms of genes to generate proteomic diversity (1), and is regulated in a tissue- and developmental stage-specific manner. The fraction of human genes that undergo alternative splicing had been first estimated as ~40% based on mRNA and EST sequence data (2,3). It has dramatically increased with the progress of microarray and high-throughput sequencing technologies, and the current estimate is as high as 74–95% of multiexon genes in human (4–6).

Deciphering the set of rules that govern the regulation of alternative splicing has been an important step toward understanding the mechanisms of alternative splicing (7–9). Besides the core splice signals in splice site selection, there are two major groups of cis-acting regulatory sequences for both constitutive and alternative splicing: exonic and intronic splicing regulators. The accumulation of cDNA and EST sequences, and the determination of the human genomic sequences have made it possible to computationally analyze exonic splicing regulators (10,11). However, until recently, it had been difficult to identify intronic splicing regulators by comparative genomic approaches because of the lack of sequence information of introns in other mammalian species than human.

The progress of genome sequencing projects of mammalian species has changed the situation, and now we can make use of genome sequence alignments to identify not only intronic splicing regulators but also various cis-
regulatory elements. The underlying idea is that, due to the purifying selection acting on functionally important sites, regulatory elements evolve much more slowly than the rest of non-functional sequences. Thus, highly conserved sequences in the genome alignments can be considered as candidate regulatory elements (12). Analyses in this direction successfully identified many cis-elements in promoter and 3'-untranslated regions (13) and insulator sites (14). Although it has been reported that many of the alternative splicing events are species-specific, i.e. they are not well conserved even among mammalian species (15–17), several studies have shown that genome alignments can be used to identify potential intronic splicing regulators (18–21).

In alternative splicing, not only a single motif can regulate spatio-temporal patterns of isoforms, but there must be combinatorial mechanisms to achieve complex regulation. To address this issue, we first identified highly significant potential intronic splicing regulators by using genome sequence alignments of mammalian species. Then we analyzed the relationships among these potential regulatory elements in terms of their co-occurrence in the intronic sequences.

MATERIALS AND METHODS

mRNA sequences and their coordinates on the human genome

To detect alternatively skipped exons, we used high-quality non-redundant mRNAs from human in the Reference Sequence (RefSeq) Database (22) (release 16, March 2006, http://www.ncbi.nlm.nih.gov/RefSeq). There are several categories in the RefSeq mRNAs, and we only used well-annotated mRNA sequences, i.e. ‘reviewed’, ‘provisional’, and ‘validated’. For our purposes, this high-confident set of exon skipping is more effective than using all possible event on the cost of false positives, e.g. by including EST sets. The final number of the mRNA sequences we obtained is 21 754. These mRNA sequences are mapped on the human chromosomes (hg18, March 2006, http://hgdownload.cse.ucsc.edu/goldenPath/hg18/bigZips/chromFa.zip) by using the BLAT program (23) to determine the coordinates on the human genomic sequences. For each gene, we compared the structure and coordinates of the transcripts to identify skipped exons. Here, we considered an exon as skipped if: (i) the exon is present in a transcript but completely absent in another one; and (ii) the 5'-end of the upstream exon and the 5'-end of the downstream exon have exactly the same coordinates. In total, we identified 1736 skipped exons in 1473 genes.

Genome sequence alignment

The genome sequence alignments of vertebrate genomes generated by the TBA program (24) were downloaded from the UCSC Genome Browser Database (25) (http://hgdownload.cse.ucsc.edu/goldenPath/hg18/multiz17way). Although the genome alignment contains 17 species, we focused on mammalian genomes (12 species; release date and the assembly identifiers are in the parenthesis): human (March 2006, hg18), chimpanzee (November 2003, panTro1), macaque (January 2006, rheMac2), mouse (February 2006, mm8), rat (November 2004, rn4), rabbit (May 2005, oryCun1), dog (May 2005, canFam2), cow (March 2005, bosTau2), armadillo (May 2005, dasNov1), elephant (May 2005, loxAf1), tenrec (July 2005, echTel1) and opossum (January 2006, monDom4).

Counting conserved pentamers in the flanking introns of the skipped exons

We searched for intronic splicing regulators within introns 1000-bases upstream and downstream of the skipped exon (Figure 1). We adopted this length cutoff according to some related studies (17,26,27). The 60 residues at the upstream of 3' splice site of the skipped exon and the 5 residues at the downstream of the 5' splice site of the skipped exon are excluded from the analysis because these regions contains splice site signals, polypyrimidine tract and branch point sequences, which are rather highly conserved and might bias our analysis. We are aware that some motifs might exist in these regions, but they will again come at the cost of false positives. If the flanking intron of the skipped exon is shorter than 1000 bases, then only the region up to the neighboring exon is taken. In such cases, the 60 residues upstream of 3' splice site of the neighboring exon and the 5 residues downstream of 5' splice site of the neighboring exon are also excluded from the analysis to ensure that the region does not contain splice site signals, polypyrimidine tract, or branch point sequences.

Then we retrieved the multiple sequence alignments of genomic sequences corresponding to these regions, and searched for strictly conserved pentamers among at least 10 mammalian species. Here, we masked the regions that are highly conserved (> 85% identity) for more than 50-residues usually cis-elements for alternative splicing.

![Figure 1](image_url)

**Figure 1.** The regions analyzed to search for intronic splicing regulators. The skipped exon is shown in gray box. The regions analyzed are shown in gray arrows. If the flanking intron of the skipped exon is shorter than 1000 bases, then only the region up to the neighboring exon are taken. In such case, the 60 residues upstream of 3’ splice site of the neighboring exon and 5 residues downstream of 5’ splice site of the neighboring exon are also excluded from the analysis to ensure that the region does not contain splice site signals, polypyrimidine tract, or branch point sequences.
are much shorter and such a long stretch of conserved region might contain unannotated RNA genes embedded within introns or unannotated exons, and counting these conserved regions would bias the analysis. These cutoffs are taken from the average length and sequence identity of orthologous exons between human and mouse (28). In counting conserved pentamers, simple repeats, e.g. runs of Ts (TTTTTTTTTT), are counted only once to prevent over counting.

Statistical significance of conserved pentamers

To assess the statistical significance of the conserved pentamers, first we counted the number of pentamers that are strictly conserved in at least 10 mammalian genomes in the region defined above. We also counted the number of pentamers that are not conserved in the same regions in human. As a control, we counted the numbers of both conserved and unconserved pentamers in the intron between the constitutive exons. For this, we used the same RefSeq gene set (1473 genes), in which we identified exon skipping, and defined an exon as constitutive if there are at least three transcripts for a gene and all the transcripts have exactly the same exon. We set these criteria to minimize the effect of alternative splicing not observed in the current RefSeq database. Then we applied Fisher’s exact test to evaluate the specific enrichment of the conserved pentamers in the flanking introns of the skipped exons.

Recent studies have shown that the RNA-Seq method, the transcriptome analysis using next-generation sequencing platforms, reveals more instances of alternative splicing events than the analyses by using mRNA and EST sequences (5,6). This means that some of the constitutive exons that we used as a control set might be alternatively skipped, and might affect our ability to detect motifs. To measure the amount of such cases, we analyzed the exons in our data sets with the RNA-Seq data (5,6) obtained from the GEO database (29). We classified an exon as skipped if there are at least three reads that cover five or more nucleotides of either the upstream and downstream exon. Using the RNA-Seq data from Wang et al. (6), only 68 exons out of 3086 exons that we classified as constitutive exons by RefSeq mRNAs are observed to be skipped. We also carried out the same analysis using the RNA-Seq data from Pan et al. (5), and detected only 37 exons out of the 3086 constitutive exons as skipped. By merging the results obtained from these two RNA-Seq data, total of 88 (2.9%) constitutive exons are detected as skipped, indicating that the quality of the classification for constitutive exons by using RefSeq mRNAs is high enough to be used as the control.

Since our analysis is based on the conserved pentamers, we may get two or more pentamers from a single highly conserved motif of the length longer than 5nt. We grouped these pentamers as clustered motifs according to the overlap in position with highly significant probability \(P < 1.0 \times 10^{-12}\) by the cumulative hypergeometric distribution function (30); see below.

Statistical significance of co-occurrence of motifs

The statistical significance of motif co-occurrence was calculated by the cumulative hypergeometric distribution function (30). The probability, \(P\), of obtaining the number of co-occurrence equal to or higher than \(c\) purely by chance is calculated by the following equation:

\[
P(c) = \sum_{i = c}^{\min(m_1, m_2)} \binom{m_1}{i} \binom{N - m_1}{m_2 - i} \binom{N}{m_2}
\]

where \(m_1\) and \(m_2\) are the number of motif1 and motif2, respectively, \(N\) the total number of introns examined and \(i\) the summation index. For example, if the numbers of motif1 and motif2 are 60 and 80, respectively, and 20 of them co-occur in the total of 800 introns, then the probability to have such co-occurrence purely by chance is calculated as \(1.8 \times 10^{-7}\).

Prediction of skipped exons in the human gene set

To evaluate whether the motif co-occurrence can be a strong indicator for alternative splicing, we predicted exon skipping in the Ensembl exon set (31) by using the motif co-occurrence and the conservation among the mammalian genomes. For this prediction, we used all Ensembl exons except for those identical with the exons identified in the initial RefSeq analysis. We considered a prediction to be correct if there is at least one for each transcript variant in ESTs from dbEST or mRNAs from GenBank. As a control, all internal exons of the Ensembl gene set are evaluated. The success rates between the predicted set and the control set are evaluated by \(\chi^2\) test.

Exon skipping supported by RNA-Seq data

To further confirm our predictions of exon skipping, we used RNA-Seq data (5). From the GEO database at NCBI (29), we downloaded RNA-Seq data (the GEO accession number: GSE13652) for the following six tissues: brain, cerebral cortex, heart, liver, lung and skeletal muscle. For each prediction of exon skipping, we created a 64-nt-long splicing junction sequence by using the bowtie program (32). To eliminate false positive hits, we did not allow any mismatches/indels. We also set a condition that at least five nucleotides of the read should overlap either the upstream or downstream exon.

To see if more tissue data can improve the detection rate of exon skipping events, we applied a re-sampling method to the existing data. Since we have data for six tissue samples, we counted the number of supported exon skipping for all possible combination from one to six samples, and the number of supported exon skipping is plotted along the number of tissue samples we used.
**Reverse transcription polymerase chain reaction and amplification of splicing products**

Poly (A)^+ RNAs from the following 12 human tissues (or brain regions) were purchased from Clontech Laboratories, Inc. (Mountain View, CA): fetal brain, hippocampus and amygdala of adult brain, skeletal muscle, bone marrow, prostate, thymus, pancreas, heart, liver, kidney and testis. We did not use any cancer cell lines, because it is known that there are many cancer-specific alternative splicing events, which might bias the analysis (33). Using the poly (A)^+ RNAs, single-stranded cDNAs were prepared as described previously (34). The polymerase chain reaction using the single-stranded cDNA as a template [Reverse transcription polymerase chain reaction (RT–PCR)] was carried out with an LA-Taq polymerase (Takara Bio, Inc., Shiga, JAPAN) in the presence of thermostable Rec A protein as described previously (35). Using thermostable Rec A, we could use the common PCR cycling conditions for all the primer sets given in Supplementary Table S1 as follows: denaturation step, 95°C for 1 min, 30 cycles of PCR consisting of 95°C for 15 s, annealing at 52°C for 30 s and extension at 72°C for 45 s. The final PCR cycle was followed by additional extension reaction at 72°C for 5 min. The PCR products were resolved on 4% NuSeive GTG agarose gel (Takara Bio Inc.) and then detected by fluorescent staining with ethidium bromide. After isolation of the PCR products on agarose gel, DNA sequencing of the PCR products was done on an ABI 3130 (Applied Biosystems, Foster City, CA) using a Big Dye Terminator v3.1 Cycle Sequencing kit (Applied Biosystems).

**RESULTS AND DISCUSSION**

**Identification of intronic splicing regulatory motifs by using genome alignments**

To systematically identify and predict sequence motifs that point to the regulation of alternative splicing, we focused on alternatively skipped exons as they are straightforward to measure and are the most frequent type of alternative splicing (36). We compared 21754 human mRNAs from the well-curated RefSeq database, and identified 1736 skipped exons in 1473 genes (see Supplementary Table S2 for the list of skipped exons).

It has been shown that a certain fraction of alternative splicing is not conserved between human and mouse (16). This means that some alternatively spliced exons that we identified by the comparison of human RefSeq mRNAs might not be alternatively spliced in the other species. However, if the existence of conserved sequences around the exons that are alternatively spliced in human, then this implicates that the conserved sequences is also involved in alternative splicing in other species. Recent studies have also shown that the inter-species comparison by using genome alignments makes it possible to identify many intronic splicing regulators (18–20). Based on these approaches, we retrieved flanking introns for each alternatively skipped exon and constructed alignments of 12 mammalian genomic sequences corresponding to the regions. After filtering out highly conserved 5’ donor and 3’ acceptor sites, polypyrimidine tracts and branch-point sites (Figure 1), we recorded all pentamers that were strictly conserved in at least 10 mammalian genomic sequences. We chose pentamers based on the fact that many splicing factors recognize specific sequences of 4–6 nt in length (37,38). Another reason to use pentamers is that the number of all possible patterns is higher (4^5 = 1024) than that for tetramers (4^4 = 256), which might help to reduce false counts of patterns. The statistical power comes at the cost of possibly missing some short and degenerated patterns, such as the Nova binding motif (YCYA) (39) or the CU-rich element (40). Longer patterns can be detected though by combining overlapping patterns (see below). To assess whether these pentamers are specifically conserved in the flanking introns of skipped exons, we compared the frequencies of these pentamers with those in the introns flanking constitutive exons as a control and calculated a statistical significance by Fisher’s exact test. Then we sorted the pentamers according to the Fisher’s P-values, and listed 18 pentamers with \( P < 1.3 \times 10^{-4} \) (Table 1; see Supplementary Table S3 for the complete list of motifs and the corresponding genes). The cutoff was selected as the highest P-value for the pentamer with experimental support (see below). Some of these pentamers are sub-strings of longer motifs; therefore, we clustered these according to their overlaps in the genomic sequence and arrived at a total of 11 distinct motifs.

Although we masked highly conserved regions longer than 50-residues (see ‘Materials and Methods’ section), there are only 64 intronic regions for the skipped exons out of the total of 1736 skipped exons. As a result, even without the masking, we obtained very similar statistics for the pentamers.

The most significant motif is GCATG (Table 1). This pentamer is conserved 157 times in the flanking introns of skipped exons, which corresponds 103 distinct exons from 98 genes. This pentamer, together with the second most significant one, forms a well-characterized hexameric motif, TGCATG, which has been identified as binding site for the Fox-1 protein (37). In some genes this motif is highly conserved among vertebrates (26), and it is also highly frequent in the neighboring introns of skipped exons in nematodes (38). The third and fourth of the pentamers (Table 1) form a hexameric motif, ACTAAC. This motif, which is similar to the branch-point consensus sequence, has been recently identified as specifically enriched in muscle-regulated alternative splicing (27). Among the other significantly overrepresented motifs, some are similar, but not identical, to some previously predicted motifs (Table 1). For example, the GTGTGGTG G motif is very similar to the purine-rich element that has been shown to be involved in the regulation of alternative splicing in thyroid hormone receptor mRNA (41). This motif is also similar to the G-rich motif, (A/U)GGG, which has been identified as an intronic splicing enhancer (42). Additional novel regulatory motifs are likely to be enriched in our list, but are below our stringent P-value cutoff (see Supplementary Table S4 for all the pentamers sorted by the P-values).
Table 1. Conserved pentamers with statistical significance

<table>
<thead>
<tr>
<th>Rank</th>
<th>Conserved pentamer</th>
<th>Clustered motif</th>
<th>P-value$^a$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>GCATG</td>
<td>TGCAT</td>
<td>1.0 x 10^-29</td>
</tr>
<tr>
<td>2</td>
<td>TGCTG</td>
<td>CTGCTG</td>
<td>1.4 x 10^-20</td>
</tr>
<tr>
<td>3</td>
<td>ACTAA</td>
<td>ACTAAC</td>
<td>7.5 x 10^-12</td>
</tr>
<tr>
<td>4</td>
<td>CTAAAC</td>
<td>ACTAAC</td>
<td>9.8 x 10^-10</td>
</tr>
<tr>
<td>5</td>
<td>TGCTG</td>
<td>CTGCTG</td>
<td>8.7 x 10^-08</td>
</tr>
<tr>
<td>6</td>
<td>GCTGC</td>
<td>CTGCTG</td>
<td>3.3 x 10^-07</td>
</tr>
<tr>
<td>7</td>
<td>TGCTT</td>
<td>CTTGCCTT</td>
<td>7.2 x 10^-06</td>
</tr>
<tr>
<td>8</td>
<td>CTGGC</td>
<td>CTTGCCTT</td>
<td>8.2 x 10^-06</td>
</tr>
<tr>
<td>9</td>
<td>GTGGG</td>
<td>GTGGTGGG</td>
<td>1.1 x 10^-05</td>
</tr>
<tr>
<td>10</td>
<td>TTTCT</td>
<td>TTTCT</td>
<td>1.2 x 10^-05</td>
</tr>
<tr>
<td>11</td>
<td>AAGAT</td>
<td>AAGAT</td>
<td>3.6 x 10^-05</td>
</tr>
<tr>
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<td>TGGAA</td>
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<td>CTGCTG</td>
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<td>AAAGG</td>
<td>6.3 x 10^-05</td>
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<td>TCTTG</td>
<td>TCTTG</td>
<td>1.2 x 10^-04</td>
</tr>
<tr>
<td>18</td>
<td>GTGTTG</td>
<td>GTGTTGGG</td>
<td>1.3 x 10^-04</td>
</tr>
</tbody>
</table>

Pentamers are sorted by P-values. The pentamers that cover only a part of known motifs are excluded from the list (see Supplementary Table S4 for a complete list of the pentamers sorted by P-value).

$^a$P-values are calculated by Fisher’s exact test.

The length of flanking introns that are considered by different studies varies a lot, e.g. including 100 bases (21) or up to 300 bases (9) from the splice sites. To see the distribution of conserved pentamers around the flanking introns of alternatively skipped exons, we counted the number of 18 pentamers listed in Table 1 (Figure 2). For both upstream of the 3' splice sites and downstream of the 5' splice sites, the conserved pentamers are more frequent within 300 bases from the splice sites, but we still see some conserved motifs even up to 1000 bases. Although there must be bona fide motifs in the regions >1 kb apart from the splice sites, the inclusion of those compromises the selectivity of the analysis.

Comparison with the other data sets

We compared the 18 significant pentamers with recently predicted intronic splicing regulators (6,18–20,43) (Table 2). Although all the statistically significant pentamers are also listed in the other data sets at least as a sub-string of the longer patterns, there is no single data set that covers all the 18 pentamers. Four pentamers, GTGGG, AAAGG, GTGTTG and GGTGGG, which corresponds to two clustered motifs, GTGGTGGG and AAA GG, exist only as a sub-string of the longer patterns in the other data sets. In some data sets, even well-known patterns are missing. For example, the most significant pentamer in our list is missing in three out of four sets of motifs identified by Churbanov et al. (20). Another example is ACTAAAC, which is missing not only in the three sets of the motifs identified by Churbanov et al. (20), but also missing in the list of intron-identity elements identified by Zhang et al. (43). Most of the pentamers we identified, especially the ones with lower P-values, are also represented in the donor/acceptor intrinsic flanking region by Voelker et al. (18) and the conserved words located in the upstream/downstream of internal exons analyzed by Yeo et al. (19).

Analysis of genome alignments with various groups of species

To measure the changes in the power of motif detection with several groups of mammalian species, we analyzed the genome alignments for each group of species. Here, we analyzed the following five groups of species: (i) human and mouse; (ii) Euarchontoglires; (iii) Boreouetheria; (iv) placental mammals; and (v) mammals (see Table 3 for the species in each group). Since we do not know bona fide cis-elements in the genome-wide scale, we focused only on the pentamer, GCATG, which is the part of the well-known cis-element for alternative splicing, and counted the number of the conserved pentamers for each groups of species (Table 3). If genome alignments of only human and mouse are analyzed, we obtain more than 300 motifs around skipped exon. But, at the same time, higher number of motifs are found in the introns around constitutive exons, indicating that analyzing only two species, human and mouse, would give a higher ratio of false positive predictions. As a result, we have a higher P-value by Fisher’s exact test comparing to the sets with more species.

Although the divergence within the group is similar between the set consists of human and mouse (the first
row in Table 3) and Euarchontoglires (the second row in Table 3), number of the GCATG pentamers and the associated P-value are very different. This indicates that adding more internal species can drastically improve the power of the cis-element detection. By comparing Euarchontoglires and Boreouetheria, number of the conserved GCATG pentamers around the skipped exons drops from 177 to 158. The difference in the number of motifs might represent the existence of species-specific motifs, i.e. the motifs exist only in Euarchontoglires but do not exist neither in dog nor in cow genomes. Or another explanation for this is that the motifs might not be aligned properly because of the inclusion of distant species.

Motifs are often co-occur in the same flanking introns

When analyzing the genomic positions of these motifs, we found that a significant number of motifs co-occur in the same flanking introns of skipped exons. One such example is found in the downstream of the skipped exon of the nuclear distribution element-like protein (NDEL1; Figure 3A). The two motifs, ACTAAC and TGCAAT, are strictly conserved in the 12 mammalian species, and there are no highly conserved segments around this region except for these two motifs. It has been shown that the exon is selectively included in the transcript in melanocyte but not in melanoma (44). This cell line-specific differential expression of the exon suggests a tightly regulated mechanism for the alternative splicing of the exon, and the co-occurring motifs are the most likely candidates to mediate this process. Another example is found in the downstream of the skipped exon of myosin binding protein C (MYBPC1), where TGCAAT and TGCTT, are separated by 19–21 residues (Figure 3B). The two examples illustrate that binding of the splicing regulator to the individually well-characterized TGCAAT motif appears to be fine-tuned by other factors binding to other neighboring sites. This led to the hypothesis that different motif pair combinations might provide the signature for context-dependent splicing.

By applying a cumulative hypergeometric distribution function [Equation (1)], we obtained statistically significant co-occurring motifs. To determine the cutoff of co-occurrence P-value, we compared the distributions of the P-values calculated from all possible pairs among the 18 highly significant motifs and the P-values calculated from randomly selected pairs of pentamers (Figure 4). Based on these distributions, we selected the two cutoffs, \( P < \frac{1.0}{C2_{10}} \) and \( P < \frac{1.0}{C2_{10}} \), for statistically significant co-occurrence of motifs. Although we restricted our analysis only to the highly significant motifs listed in Table 1, various combinations of co-occurring motifs are significant, forming a complex network (Figure 5), likely to be the basis for context-dependent regulation. Very recently, Barash et al. (9) presented three networks, which consist of motifs and other features, to predict tissue-dependent alternative splicing. Some of the edges in our network are also present in their networks. For example, the edge showing the frequent association of TGCAAT and ACTAAC in our

### Table 2. Comparison of the 18 pentamers with the patterns in other data sets

<table>
<thead>
<tr>
<th>Voelker et al. (18)</th>
<th>Yeo et al. (19)</th>
<th>Churbanov et al. (20)</th>
<th>Wang et al. (6)</th>
<th>Zhang et al. (43)</th>
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</thead>
<tbody>
<tr>
<td><strong>Donor</strong>&lt;sup&gt;a&lt;/sup&gt;</td>
<td><strong>Acceptorb</strong></td>
<td>3'-ISE&lt;sup&gt;c&lt;/sup&gt;</td>
<td>3'-ISS&lt;sup&gt;c&lt;/sup&gt;</td>
<td>5'-ISE&lt;sup&gt;d&lt;/sup&gt;</td>
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<tr>
<td>(819)</td>
<td>(1007)</td>
<td>(814)</td>
<td>(1032)</td>
<td>(478)</td>
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<td>GCATG</td>
<td>M</td>
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<td>–</td>
<td>s</td>
<td>–</td>
<td>s</td>
</tr>
<tr>
<td>TCTTG</td>
<td>s</td>
<td>M</td>
<td>M</td>
<td>s</td>
</tr>
<tr>
<td>GGGTG</td>
<td>–</td>
<td>s</td>
<td>–</td>
<td>s</td>
</tr>
<tr>
<td>GCTGG</td>
<td>s</td>
<td>–</td>
<td>s</td>
<td>s</td>
</tr>
</tbody>
</table>

The total number of predicted motifs are indicated in the parenthesis under each data set.

<sup>a</sup>Significantly conserved n-mers found in the donor/acceptor intronic flanking region (18).

<sup>b</sup>Conserved words that are located upstream/downstream of internal exons (19).

<sup>c</sup>3'-/5'-splice site-related intronic splicing silencers/enhancers (20).

<sup>d</sup>3'-/5'-splice site intronic motifs (6).

<sup>e</sup>Intron-identity elements (43).

M, exact match; s, the pentamer is included as a sub-string of the longer pattern; –, no match found.
Table 3. GCATG motif in the genome alignments of various groups of species

<table>
<thead>
<tr>
<th>Species</th>
<th>Conserved GCATG in the introns</th>
<th>P-value*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Around skipped exons</td>
<td>Around constitutive exons</td>
</tr>
<tr>
<td>Human + mouse</td>
<td>303</td>
<td>136</td>
</tr>
<tr>
<td>Euarchontoglires</td>
<td>177</td>
<td>29</td>
</tr>
<tr>
<td>Boreoeutheria</td>
<td>158</td>
<td>17</td>
</tr>
<tr>
<td>Placental mammals</td>
<td>140</td>
<td>21</td>
</tr>
<tr>
<td>Mammals</td>
<td>157</td>
<td>26</td>
</tr>
</tbody>
</table>

In the mammals set, we count the motif if it is strictly conserved at least in 10 species.

*P-values are calculated by Fisher’s exact test.

**Human, chimpanzee, macaque, mouse, rat, rabbit.**

***Human, chimpanzee, macaque, mouse, rat, rabbit, dog, cow.**

****Human, chimpanzee, macaque, mouse, rat, rabbit, dog, cow, armadillo, elephant, tenrec.****

******Human, chimpanzee, macaque, mouse, rat, rabbit, dog, cow, armadillo, elephant, tenrec, opossum.******

Prediction of alternatively skipped exons by motif co-occurrence

We then tested whether the co-occurrence of the identified motifs in flanking introns of an exon can be applied to predict novel alternatively skipped exons. For this, we extended our analysis to the flanking introns of all annotated exons in the human Ensembl gene set, network is also present in the network for muscle-specific inclusion of alternative exons (9). Another pair of motifs in our network, TTTCT and TGCCATG, can be found in the network for central nervous system-specific inclusion of alternative exons (9). We have these edges (i.e. pairs of motifs) in the same network because we do not take tissue-specificity into account. We consequently analyzed functional GO terms and tissue-specific isoforms in EST and cDNA databases, but did not see any significant association of the co-occurring motifs and functional classes or tissue-specificity (data not shown).

Figure 3. Examples of co-occurring motifs. (A) ACTAAC and TGCATG in the flanking intron of the skipped exon of NDEL1 and (B) TGCCATG and TGCCATG in the flanking intron of the skipped exon of MYBPC1. The part of the gene structure around the skipped exon is shown above the alignment with the exon numbers. The format of the alignment: the first column, the name of the species; second column, chromosome or scaffold identifier; the third column, direction of the gene on the chromosome, the fourth column, position on the chromosome. Conserved residues are indicated by asterisks under the alignments. Conserved motifs are indicated by colored boxes.
exons with co-occurring motifs in the flanking introns ($P = 3.7 \times 10^{-9}$ by $\chi^2$ test). We further looked for the supporting evidence for the exon skipping in our predictions in recently published RNA-Seq data from six tissue samples (5), and, in addition to the 29 annotated skipping events, RNA-Seq data confirmed 8 additional instances (Supplementary Table S5). To see if more tissue data can improve the detection rate of exon skipping, we applied a re-sampling method to the existing data. The results show that if we have RNA-Seq data for more tissue samples, even more predictions of alternatively skipped exons will be supported although the data are still too sparse to develop a tissue code for exon skipping (Figure 6). For the predictions with no EST and mRNA support, we also randomly selected 10 predicted alternatively skipped exons together with 10 internal exons unlikely to undergo exon skipping and carried out RT–PCR to actively search for the evidence of exon skipping. In 12 tissue libraries, 3 of the 10 exons predicted to be skipped were confirmed to be skipped in certain tissues, whereas no exon skipping was detected for the 10 unpredicted exons as far as we examined (Supplementary Table S6). For example, the skipped form of the third exon of ENST00000256858, as well as the included form, is detected in fetal brain and two regions of adult brain (hippocampus and amygdala) (Figure 7), while it is not skipped in all the other tissues we tested (see Supplementary Figure S1 for all the results of the PCR experiments). This not only confirms the predictive power of motif combinations, but also the low resolution of current transcriptome data with respect to cell type and organism age. For the predictions that the skipped forms have not been confirmed by PCR experiments, we cannot exclude the possibility that the co-occurring motifs might work in a competitive manner. In the future, the addition of other features, such as exon/intron length and RNA secondary structure, should improve the prediction accuracy (9).
In total, 39 exons out of 118 predictions are confirmed to be alternatively spliced (Supplementary Table S5). To see if these 39 cases have some additional features comparing to the rest of the 79 cases, first we calculated the distances of the motifs pairs. The average distances for the confirmed 39 cases and the rest of the 79 cases are 518.1 and 321.3 bases, respectively. Applying Mann–Whitney U-test (45) to these two distributions of the motif distances, we obtained $z' = 1.97$, which corresponds to $P > 0.01$. Then we calculated secondary structure forming potential around the motifs. For each motif, we extracted the sequence from 50 bases upstream to 50 bases downstream of the motif, and applied the RNAfold program (46) to calculate possible folding energy. The average folding energy values for the confirmed 39 cases and the rest of the 79 cases are $-19.6$ and $-17.8$ kcal/mol, respectively. The two energy distributions are compared by Mann–Whitney U-test, which gives $z' = 2.56$ ($P > 0.01$). From these analyses, we do not see any statistically significant differences between the experimentally confirmed set and the set yet to be confirmed at least for these two features.

CONCLUSIONS

It has been shown that some distinct splicing factors work together to regulate alternative splicing in some instances (47–49). We show that such co-occurrence of the cis-elements is not a special case for a certain pair of motifs but it might be a common principle as almost all motifs identified here form a network that might be the basis for context-specific intronic splicing regulation. The co-occurring elements might act in a cooperative or competing manner to provide complexity in the regulation of alternative splicing, for example, to achieve fine-grained tissue- or developmental stage-specific control. Recently an experimental system has been devised to analyze coordinated regulation of tissue-specific alternative splicing (50), which will allow a more detailed analysis of the regulation provided by the co-occurring elements identified in this study. This kind of combinatorial regulation is analogous to that of transcription, in which combinatorial usage of distinct transcription factors provide a large variety of possibilities for gene expression patterns (51), and this principle might, thus, widespread in genomic regulation. Co-occurrence of cis-regulatory elements also suggests that, although one element might be sufficient to explain the regulatory mechanism of alternative splicing under a given condition, other elements might be necessary to fully explain the complex regulation of alternative splicing.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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