Animal, fungi, and plant genome sequences harbour different non-canonical splice sites

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Abstract

Most protein encoding genes in eukaryotes contain introns which are inter-5 woven with exons. After transcription, introns need to be removed in order 6 to generate the final mRNA which can be translated into an amino acid 7 sequence by the ribosome. Precise excision of introns by the spliceosome 8 requires conserved dinucleotides which mark the splice sites. However, 9 there are variations of the highly conserved combination of GT at the 5' 10 end and AG at the 3' end of an intron in the genome. GC-AG and AT-AC 11 are two major non-canonical splice site combinations which are known for 12 many years. During the last few years, various minor non-canonical splice 13 site combinations were detected with all possible dinucleotide permuta-14 tions. Here we expand systematic investigations of non-canonical splice 15 site combinations in plant genomes to all eukarvotes by analysing fungal 16 and animal genome sequences. Comparisons of splice site combinations 17 between these three kingdoms revealed several differences such as a sub-18 stantially increased CT-AC frequency in fungal genomes. In addition, high 19 numbers of GA-AG splice site combinations were observed in two animal 20 species. In depth investigation of splice site usage based on RNA-Seq 21 read mappings indicates a generally higher flexibility of the 3' splice site 22 compared to the 5' splice site. 23

²⁴ Introduction

Splicing, the removal of introns after transcription, is an essential step dur-25 ing the generation of mature mRNAs in eukaryotes. This process allows 26 variation which provides the basis for quick adaptation to changing con-27 ditions [1,2]. Alternative splicing, e.g. skipping exons, results in an enor-28 mous diversity of synthesized proteins and therefore substantially expands 29 the diversity of products encoded in eukaryotic genomes [3-6]. The full 30 range of functions as well as the evolutionary relevance of introns are still 31 under discussion [7]. However, introns are energetically expensive for the 32 cell to maintain as the transcription of introns costs time and energy and 33 the removal of introns has to be exactly regulated [8]. Dinucleotides at 34 both intron/exon borders mark the splice sites and are therefore highly 35 conserved [9]. GT at the 5' end and AG at the 3' end of an intron form the 36 canonical splice site combination on DNA level. More complexity arises 37 through non-canonical splice site combinations, which deviate from the 38 highly conserved canonical one. Besides the major non-canonical splice 39 site combinations GC-AG and AT-AC, several minor non-canonical splice 40 site combinations have been detected before [9, 10]. 41

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Furthermore, the position of introns in homologous genes across organ-43 isms, which diverged 500-1500 million years ago, are not conserved [11]. 44 In addition, many intron sequences mutate at a higher rate due to hav-45 ing much less of an impact on an organism's reproductive fitness com-46 pared to a mutation located within an exon [12]. These factors, along with 47 the existence of several non-cannonical splice sites, make the complete 48 prediction of introns, even in non-complex organisms like yeast, almost 49 impossible [13, 14]. Moreover, most introns which can be predicted com-50 putationally still lack experimental support [15]. 51

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Splice sites are recognised during the splicing process by a complex of
 snRNAs and proteins, the spliceosome [16]. U2-spliceosome and U12 spliceosome are two subtypes of this complex which comprise slightly dif-

ferent proteins with equivalent functions [17–19]. Although the terminal dinucleotides are important for the splicing process, these splice sites are not sufficient to determine which spliceosome is processing the enclosed intron [20]. This demonstrates the complexity of the splicing process which involves additional signals present in the DNA. Even though multiple mechanisms could explain the splicing process, the exact mechanism of noncanonical splicing is still not completely resolved [5].

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⁶⁴ Branching reaction and exon ligation are the two major steps of splic-⁶⁵ ing [21,22]. In the branching reaction, the 2'-hydroxyl group of the branch-⁶⁶ point adenosine initiates an attack on the 5'-phosphate of the donor splice ⁶⁷ site [23,24]. This process leads to the formation of a lariat structure. Next, ⁶⁸ the exons are ligated and the intron is released through activity of the 3'-⁶⁹ hydroxyl group of the 5'exon at the acceptor splice site [21].

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Previous in-depth analyses of non-canonical splice sites in fungi and an-71 imals were often focused on a single or a small number of species [9, 72 25, 26]. Several studies focused on canonical GT-AG splice sites but ne-73 glected non-canonical splice sites [27, 28]. Our understanding of splice 74 site combinations is more developed in plants compared to other king-75 doms [10, 29-33]. Previous works reported 98 % GT-AG splice site com-76 binations in fungi [25], 98.7 % in plants [10] and 98.71 % in animals [9]. 77 Consequently, the proportion of non-canonical splice sites is around or be-78 low 2 % [9,10,25]. To the best of our knowledge, it is not known if the value 79 reported for mammals is representative for all animals. The combined pro-80 portion of minor non-canonical splice sites is even lower e.g. 0.09 % in 81 plants, but still exceeding the frequency of the major non-canonical AT-82 AC splice sites [10]. Despite this apparently low frequency, non-canonical 83 splice site combinations have a substantial impact on gene products, es-84 pecially on exon-rich genes [10]. About 40 % of genes with 40 exons are 85 affected (AdditionalFile 11). 86

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88 Consideration of non-canonical splice sites is important for gene predic-

tion approaches, because these sites cannot be identified ab initio [29]. 89 Moreover, as many human pathogenic mutations occur at the donor splice 90 site [34], it is of great interest to understand the occurence and usage of 91 non-canonical splice sites. Therefore, several non-canonical splice sites 92 containing AG at the acceptor site were investigated in human fibrob-93 lasts [34]. Alongside this, fungi are interesting due to pathogenic proper-94 ties and importance in the food industry [35]. Since splicing leads to high 95 protein diversity [3-6], the analysis of splicing in fungi is important with re-96 spect to biotechnological applications e.g. development of new products. 97

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In this study, a collection of annotated genome sequences from 130 fungi 99 and 489 animal species was screened for canonical and non-canonical 100 splice site combinations. RNA-Seq data sets were harnessed to identify 101 biologically relevant and actually used splice sites. Non-canonical splice 102 site combinations, which appeared at substantially higher frequency in a 103 certain kingdom or species, were analysed in detail. As knowledge about 104 splice sites in plants was available from previous investigations [10, 29], 105 a comparison between splice sites in fungi, animals and plants was per-106 formed. 107

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Results and Discussion

110 Analysis of non-canonical splice sites

In total, 64,756,412 and 2,302,340 splice site combinations in animals and fungi, respectively, were investigated based on annotated genome sequences (AdditionalFile 1 and 2). The average frequency of the canonical splice site combination GT-AG is 98.3 % in animals and 98.7 % in fungi, respectively. These values exceed the 97.9 % previously reported for plants [10], thus indicating a generally higher frequency of non-canonical splice site combinations in plants. As previously speculated [10], a gen-

erally more complex splicing system in plants could be an adaptation to 118 changing environments. Since most plants are not able to change their 119 geographic location, the tolerance for unfavourable conditions should be 120 stronger than in animals. The lower proportion of non-canonical splice 121 sites in fungi compared to animals seems to contradict this hypothesis. 122 However, the genome size and complexity needs to be taken into account 123 here. The average animal genome is significantly larger than the average 124 fungal genome (Mann-Whitney U-Test; p=5.64e-68) (AdditionalFile 3). 125

Average percentages of the most important splice site combinations were 126 summarized per kingdom and over all analysed genomes (Table 1). The 127 number of canonical and non-canonical splice site combinations per species 128 was also summarized (AdditionalFile 4 and 5). A higher percentage of 129 non-canonical splice sites was observed in animals in comparison to fungi. 130 Several species strongly exceeded the average values for major and minor 131 non-canonical splice sites. The fungal species Meyerozyma guilliermondi 132 shows approximately 6.67 % major and 13.33 % minor non-canonical 133 splice sites. Eurytemora affinis and Oikopleura dioica reveal approximately 134 10 % minor non-canonical splice sites. In summary, the observed frequen-135 cies of canonical and major non-canonical splice site combinations are 136 similar to the pattern previously reported for plants [10], but some essen-137 tial differences and exceptions were found in animals and fungi. 138

Table 1: Splice site combination frequencies in animals, fungi, and plants. Only the most frequent combinations are displayed here and all minor non-canonical splice site combinations are summarized as one group ("others"). A full list of all splice site combinations is available (AdditionalFile 6 and 7).

	GT-AG	GC-AG	AT-AC	others
animals	98.334 %	0.983 %	0.106 %	0.577 %
fungi	98.715 %	1.009 %	0.019 %	0.257 %
plants	97.886 %	1.488 %	0.092 %	0.534 %
all	98.265 %	1.074 %	0.101 %	0.560 %

Different properties of the genomes of all investigated species were anal-139 ysed to identify potential explanations for the splice site differences (Ad-140 ditionalFile 8 and 9). In fungi, the average number of introns per gene 141 is 1.49 and the average GC content is 47.1 % (\pm 7.39). In animals, each 142 gene contains on average 6.95 introns and the average GC content is 39.4 143 % (\pm 3.87). This difference in the GC content could be associated with the 144 much lower frequency of AT-AC splice site combinations and the higher fre-145 quency of CT-AC splice site combinations in fungi (Figure 1). CT-AC has a 146 higher GC content than the AT rich AT-AC splice site combination. A gen-147 erally higher GC content could result in the higher GC content within splice 148 site combinations due to the overall mutations rates in these species. 149

A comparison of the genome-wide GC content to the GC content of all 150 splice sites revealed a weak correlation in the analysed fungi (r \approx 0.236, 151 $p \approx 0.008$). Species with a high genomic GC content tend to show a high 152 GC content in the splice site combinations in the respective species. A 153 similar correlation ($r \approx 0.4$, p<0.001) was found in plant and animal species 154 as well (AdditionalFile 10). Additionally, the GC content in fungal genomes 155 is substantially exceeding the average GC content of plant and animal 156 genomes. 157

The most frequent non-canonical splice site combinations show differences between animals, fungi, and plants (Figure 1). In fungal species, the splice site CT-AC is more frequent than the splice site combination AT-AC. Regarding the splice site combination GA-AG in animals, two outliers are clearly visible: *Eurytemora affinis* and *Oikopleura dioica* show more GA-AG splice site combinations than GC-AG splice site combinations.

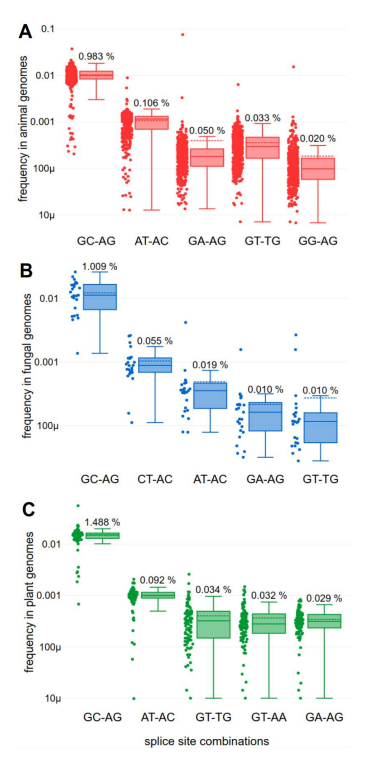


Figure 1: Frequencies of non-canonical splice site combinations in animals, fungi, and plants. The frequency of non-canonical splice site combinations across the 489 animal (red), 130 fungal (blue) and 121 plant (green) genomes is shown. Normalization of the absolute number of each splice site combination was performed per species based on the total number of splice sites. The frequency of the respective splice site combination of each species is shown on the left hand side and the percentage of the respective splice site combination on top of each box plot.

Despite overall similarity in the pattern of non-canonical splice site combi-164 nations between kingdoms, specific minor non-canonical splice sites were 165 identified at much higher frequency in some fungal and animal species. 166 First, RNA-Seq data was harnessed to validate these unexpected splice 167 site combinations. Next, the frequencies of selected splice site combina-168 tions across all species of the respective kingdom were calculated. The 169 correlation between the size of the incorporated RNA-Seq data sets and 170 the number of supported splice sites was examined as well (AdditionalFile 171 11). In animals, there is a correlation ($r \approx 0.417$, $p \approx 0.022$) between num-172 ber of supported splice sites and total number of sequenced nucleotides 173 in RNA-Seg data. For fungi, no correlation between number of splice sites 174 and size of the RNA-Seg data sets could be observed. It is important 175 to note that the the number of available RNA-Seq data sets from fungi 176 was substantially lower. Further, analysis of introns with canonical and 177 non-canonical splice site combinations, respectively, revealed that a higher 178 number of introns is associated with a higher proportion of non-canonical 179 splice sites (AdditionalFile 12). 180

High diversity of non-canonical splice sites in animals

Kupfer et al. suggested that splicing may differ between fungi and ver-182 tebrates [25]. Our results indicate substantial differences in the diver-183 sity of splice site combinations other than GT-AG and GC-AG in fungi 184 $(H' \approx 0.0277)$ and animals $(H' \approx 0.0637)$ (Kruskal-Wallis: p ≈ 0.00000). Be-185 sides the overall high proportion of minor non-canonical splice sites (Table 186 1), differences between species are high (Figure 1). The slightly higher in-187 terguartile range of splice site combination frequencies in animal species 188 and especially in plant species (Figure 1A and C), together with the rel-189 atively high frequency of "other" splice sites in animals and plants (Table 190 1) suggest more variation of splice sites in the kingdoms of animals and 191 plants compared to the investigated fungal species. Thus, the high di-192 versity of splice sites could be associated with the higher complexity of 193 animal and plant genomes. In addition, the difference in prevalence be-194

tween the major non-canonical splice site combination GC-AG and minor
 non-canonical splice site combinations is smaller in animals compared to
 fungi and plants (Figure 1).

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GA-AG is a frequent non-canonical splice site combination in some an-199 imal species. Two species, namely Eurytemora affinis and Oikopleura 200 dioica, showed a much higher abundance of GA-AG splice site combi-201 nations compared to the other investigated species (Figure 1A). RNA-Seq 202 reads support 5,795 (28.68 %) of all GA-AG splice site combinations of 203 these species. In both species, the number of the GA-AG splice site com-204 bination exceeds the number of the major non-canonical splice site com-205 bination GC-AG. 206

For *Eurytemora affinis*, the high frequency of the GA-AG splice site combinations was described previously for 36 introns [36]. We quantified the proportion of GA-AG splice site combinations to 3.2 % (5,345) of all 166,392 supported splice site combinations in this species. The donor splice site GA is flanked by highly conserved upstream AG and a downstream A (Figure 2).

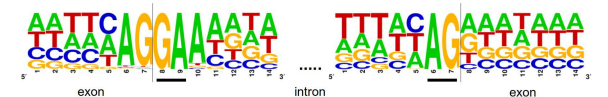


Figure 2: Flanking positions of GA-AG splice site combinations in *Eurytemora affinis* and *Oikopleura dioica*. All 5,795 supported splice site combinations of these two species were investigated. Seven exonic and seven intronic positions are displayed at the donor and acceptor splice sites. Underlined bases represent the terminal dinucleotides of the intron i.e. the donor and acceptor splice site.

Efficient splicing of the splice site combination GA-AG was detected in human fibroblast growth factor receptor genes [37]. Further, it was suggested
that this splicing event is, among other sequence properties, dependent on
a canonical splice site six nucleotides upstream [37], which does not exist

in the species investigated here (Figure 2). An analysis of all five potential 217 U1 snRNAs in this species did reveal one single nucleotide polymorphism 218 in the binding site of the 5' splice site from C to T in one of these U1 219 snRNAs. This could result in the binding of AG/GGAAGT or AGG/GAAGT 220 instead of AG/GTAAGT. Although this would imply an elegant way for the 221 splicing of GA-AG splice sites, the same variation was also detected in 222 putative human U1 snRNAs. Therefore, another mechanism seems to be 223 responsible for splicing of introns containing the GA-AG splice site combi-224 nation. 225

²²⁶ CT-AC is a frequent splice site combination in fungi

Although the general frequency pattern of fungal splice site combinations 227 is similar to plants and animals, several fungal species displayed a high 228 frequency of minor non-canonical CT-AC splice site combinations. This 229 co-occurres with a lower frequency of AT-AC splice site combinations. 230 Non-canonical splice sites in fungi were, so far, only described in stud-231 ies which focussed on a single or a few species. An analysis in the 232 oomycota species Phytophthora sojae, which is a fungus-like microorgan-233 ism [38, 39], revealed 3.4 % non-canonical splice site combinations GC-234 AG and CT-AC [40]. Our findings indicate, that the minor non-canonical 235 splice site combination CT-AC occurs with a significantly (Mann-Whitney 236 U-Test; $p \approx 0.00035$) higher frequency than the major non-canonical splice 237 site combination AT-AC. In contrast, the frequency of AT-AC in animals 238 and plants exceeds the CT-AC frequency significantly (p<0.001) (Figure 239 3A). For the splice site combination CT-AC a sequence logo, which shows 240 the conservation of this splice site in four selected species, was designed 241 (Figure 3B). In summary, we conclude that CT-AC is a major non-canonical 242 splice site combination in fungi, while AT-AC is not. 243

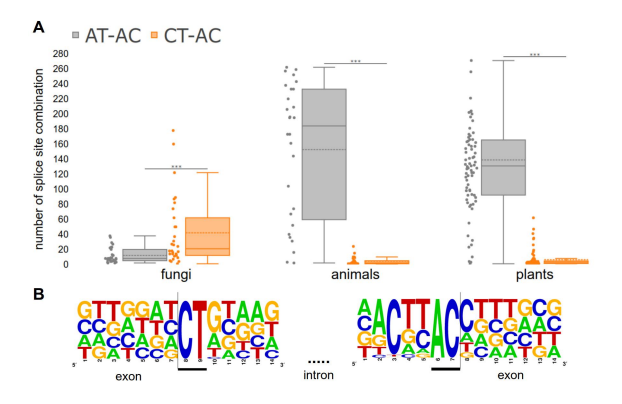


Figure 3: CT-AC frequency exceeds AT-AC frequency in fungi. A) Number of the minor non-canonical splice site combination CT-AC in comparison to the major non-canonical splice site combination AT-AC in each kingdom (p<0.001). B) Sequence logo for the splice site combination CT-AC in four selected fungal species (*Alternaria alternata, Aspergillus brasiliensis, Fomitopsis pinicola* and *Zymoseptoria tritici*). In total, 67 supported splice sites with this combination were used to generate the sequence logo.

- ²⁴⁴ The highest frequencies of the splice site combination CT-AC, supported
- ²⁴⁵ by RNA-Seq reads, were observed in Alternaria alternata, Aspergillus brasilien-
- sis, Fomitopsis pinicola and Zymoseptoria tritici (approx. 0.08 0.09 %).
- ²⁴⁷ As AT-AC was described as major non-canonical splice site, these findings
- ²⁴⁸ indicate a different splice site pattern in fungi compared to animals and ²⁴⁹ plants (Figure 3).

Intron size analysis

In total, 8,060,924, 737,783 and 2,785,484 transcripts across animals, 251 fungi and plants, respectively, were selected to check whether the intron 252 lengths are multiples of three. Introns with this property could be kept in 253 the final transcript without causing a shift in the reading frame. There is 254 no significant difference between introns with different splice site combina-255 tions (Table 2). The ratio of introns with a length divisible by 3 is very close 256 to 33.3 % which would be expected based on an equal distribution. The 257 only exception are minor non-canonical splice site combinations in fungi 258 which are slightly less likely to occur in introns with a length divisible by 3. 259

	splice site combination	frequency of introns divisible by 3	total number of introns divisible by 3
animals	GT-AG	0.333862150381	n=63677347
	AT-AC	0.325106284189	n=68919
	GC-AG	0.330352389911	n=636823
	others	0.327633755094	n=496411
fungi	GT-AG	0.33932356858	n=2273756
	AT-AC	0.331775700935	n=428
	GC-AG	0.333577333793	n=23224
	others	0.3125	n=6240
plants	GT-AG	0.332967299596	n=14227286
	AT-AC	0.326150175229	n=13411
	GC-AG	0.329271562364	n=216326
	others	0.323971037399	n=93638

Table 2: Proportion of introns with length divisible by 3. The results of intron length analysis for selected splice site combinations for animals, fungi and plants are shown.

Conservation of non-canonical splice site combinations across species

In total, A. thaliana transcripts containing 1,073 GC-AG, 64 AT-AC and 19 262 minor non-canonical splice sites were aligned to transcripts of all plant 263 species. Homologous intron positions were checked for non-canonical 264 splice sites. GC-AG splice site combinations were conserved in 9,830 265 sequences, matched with other non-canonical splice site combinations in 266 121 cases, and aligned to GT-AG in 13,045 sequences. Given that the 267 dominance of GT-AG splice sites was around 98 %, the number observed 268 here indicates a strong conservation of GC-AG splice site combinations. 269 AT-AC splice site combinations were conserved in 967 other sequences, 270 matched with other non-canonical splice site combinations in 93 cases, 271 and aligned to GT-AG in 157 sequences. These numbers indicate a con-272 servation of AT-AC splice site combinations, which exceeds the conserva-273 tion of GC-AG splice site combinations substantially. Minor non-canonical 274 splice sites were conserved in 48 other sequences, matched with other 275 non-canonical splice site combinations in 64 cases, and were aligned to 276 a canonical GT-AG splice site in 213 cases. This pattern suggests that 277 most non-canonical splice site combinations are either (A) mutations of 278 the canonical ones or (B) mutated towards GT-AG splice site combina-279 tions. 280

The power of this analysis is currently limited by the quality of the alignment. Although splice site combinations should be aligned properly in most cases, small differences in the number could be caused by ambiguous situations. It is likely that both hypothesis stated above are partly valid. To assign each splice site combination to A or B, a manual inspection of the observed phylogenetic pattern would be required.

²⁸⁷ Usage of non-canonical splice sites

²⁸⁸ Non-canonical splice site combinations were described to have regula-²⁸⁹ tory roles by slowing down the splicing process [41]. Previous reports

also indicated that non-canonical splice site combinations might appear in 290 pseudogenes [9, 10]. To analyse a possible correlation of non-canonical 291 splice sites with low transcriptional activity, we compared the transcript 292 abundance of genes with non-canonical splice site combinations to genes 293 with only canonical GT-AG splice site combinations (Figure 4A). Genes 294 with at least one non-canonical splice site combination are generally less 295 likely to be lowly expressed than genes with only canonical splice sites. 296 While this trend holds true for all analysed non-canonical splice site com-297 bination groups, GC-AG and AT-AC containing genes display especially 298 low proportions of genes with low FPKMs. We speculate that a stronger 299 transcriptional activity of genes with non-canonical splice sites compen-300 sates for lower turnover rates in the splicing process. The regulation of the 301 genes might be shifted from the transcriptional to the post-transcriptional 302 level. This trend is similar for animals and plants (AdditionalFile 13). In 303 fungi, genes with minor non-canonical splice sites display relatively high 304 proportions of genes with low FPKMs. 305

Moreover, a higher number of non-canonical splice sites per gene is associated with a lower expression. This leads to the suggestion, that noncanonical splice sites occur more often within pseudogenes.

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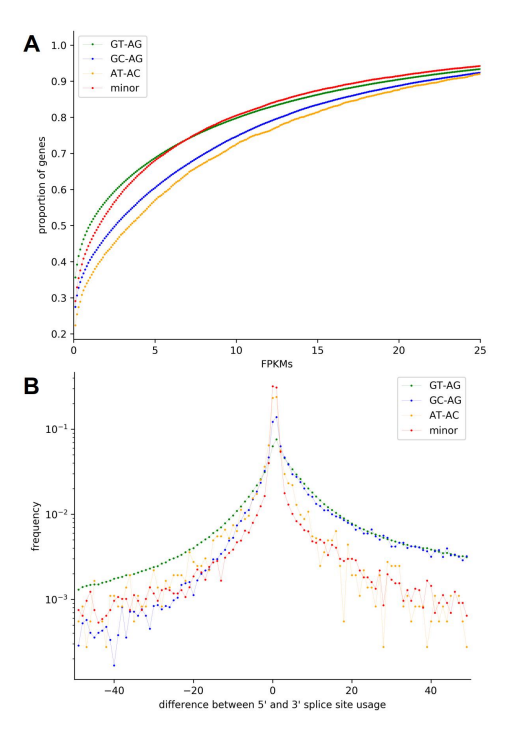


Figure 4: Usage of non-canonical splice sites in plant species. A) Comparison of the transcript abundance (FPKMs) of genes with non-canonical splice site combinations to genes with only canonical GT-AG splice site combinations. GC-AG and AT-AC containing genes display especially low proportions of genes with low FPKMs. This leads to a higher transcript abundance of genes with low FPKMs. B) Comparison of the usage of 5' and 3' splice sites. On the x-axis, the difference between the 5' splice site usage and the usage of the 3' splice site is shown. A fast drop of values when going to the negative side of the x-axis indicates that the 3' splice site is probably more flexible than the 5' splice site.

Introns are mostly defined by phylogenetically conserved splice sites, but 310 nevertheless some variation of these splice sites is possible [9, 10, 25, 26, 311 40]. To understand the amount of flexibility in respect to different terminal 312 dinucleotides, we compared the usage of donor and acceptor splice sites 313 over 4,141,196 introns in plants, 3,915,559 introns in animals and 340,619 314 introns in fungi (Figure 4B). The plot shows that the 3' splice site seems 315 to be more fexible than the 5' splice site which was observed in all three 316 kingdoms. Our observations align well with previous findings of a higher 317 flexibility at the 3' splice site compared to the 5' splice site. A mutated 5' 318 splice site represses the removal of the upstream intron [10, 42, 43]. Fur-319 ther, for plants and animals, the difference between the usage of the 5' 320 splice site and the 3' splice site is notably higher for introns with the splice 321 site combination GC-AG. 322

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Although bona fide non-canonical splice site combinations are present in 324 many plant transcripts [10], additional isoforms of the genes might exist. 325 To evaluate the relevance of such alternative isoforms, we assessed the 326 contribution of isoforms to the overall abundance of transcripts of a gene. 327 Therefore, the usage of splice sites flanking an intron was compared to 328 the average usage of splice sites. This reveals how often a certain intron 329 is removed by splicing. Introns with low usage values might only be in-330 volved in minor transcript isoforms. While most introns display no or very 331 small differences, GT-AG introns deviate from this trend. This indicates 332 that non-canonical splice site combinations are frequently part of the dom-333 inant isoform. Again, these findings were similar for all of the investigated 334 kingdoms. 335

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337 Conclusion

Our investigation of non-canonical splice sites in animals, fungi and plants revealed kingdom specific differences. Animal species with a high proportion of GA-AG splice site combinations were examined. Further, properties

of introns and splice sites were analysed. One aspect of this analysis is, 341 that the 3' splice site seems to be more flexible than the 5' splice site, 342 which was observed in all three kingdoms. In fungi, the splice site com-343 bination CT-AC is more frequent than the splice site combination AT-AC. 344 This makes CT-AC a major non-canonical splice site combination in fungal 345 species, while AT-AC should be considered a minor non-canonical splice 346 site in fungi. Overall, our findings demonstrate the importance of con-347 sidering non-canonical splice sites despite their low relative frequency in 348 comparison to the canonical splice site combination GT-AG. RNA-Seg data 349 confirmed the existence and usage of numerous non-canonical splice site 350 combinations. By neglecting non-canonical splice sites, bona fide genes 351 might be excluded or at least structurally altered. 352

353 Methods

Analysis and validation of splice site combinations

Genome sequences (FASTA) and corresponding annotations (GFF3) of 355 130 fungal species and 489 animal species were retrieved from the 356 NCBI. Representative transcript and peptide sequences were extracted 357 as described before [10]. General statistics were calculated using a 358 Python script [10]. The completeness of all data sets was assessed with 359 BUSCO v3 [44] using the reference data sets 'fungi odb9' and 'meta-360 zoa odb9', respectively [45] (AdditionalFile 14 and 15). To validate the 361 detected splice site combinations, paired-end RNA-Seq data sets were 362 retrieved from the Sequence Read Archive [46] (AdditionalFile 16 and 363 17). The following validation approach [10] utilized STAR v2.5.1b [47] 364 for the read mapping and Python scripts for downstream processing 365 (https://doi.org/10.5281/zenodo.2586989). An overview of the RNA-Seq 366 read coverage depth of splice sites in animals [48] and fungi [49] is avail-367 able. RNA-Seg read mappings with STAR and HiSat2 were compared 368 based on a gold standard generated by exonerate, because a previ-369

ous report [50] indicated a superiority of STAR. All transcripts with non-370 canonical splice sites in A. thaliana and Oryza sativa were considered. 371 When investigating the alignment of RNA-Seq reads over non-canonical 372 splice sites, we observed a high accuracy for both mappers without a 373 clear difference between them. Previously described scripts [10] were 374 adjusted for this analysis and updated versions are available on github 375 (https://doi.org/10.5281/zenodo.2586989). The distribution of genome 376 sizes was analysed using the Python package dabest [51]. Sequence 377 logos for the analysed splice sites were designed at http://weblogo. 378 berkeley.edu/logo.cgi [52]. 379

Calculation of the splice site diversity

A custom Python script was applied to calculate the Shannon diversity index (H') [53] of all splice site combinations in fungi, animals and plants (https://doi.org/10.5281/zenodo.2586989). To determine the significance of the obtained results, a Kruskal-Wallis test [54] was calculated using the Python package scipy [55]. Further, the interquartile range of all distributions was examined.

Investigation of a common non-canonical splice site in fungi

A Mann-Whitney U Test implemented in the Python package scipy was performed to analyse differences in the number of minor non-canonical splice site combinations. The observed distributions were visualized in a boxplot (https://doi.org/10.5281/zenodo.2586989) constructed with the Python package plotly [56].

³⁹⁴ Detection of potential U1 snRNAs

A potential U1 snRNA of *Pan troglodytes* (obtained from the NCBI) was subjected to BLASTn [57] against the genome sequences of selected

species. Hits with a score above 100, with at least 80 % similarity and
 with the conserved sequence at the 5' end of the snRNA [58] were in vestigated, as these sequences are potential U1 snRNAs. The obtained
 sequences were compared and small nucleotide variants were detected.

401 Correlation between the GC content of the genome and 402 the GC content of the splice sites

The Pearson correlation coefficient between the GC content of the genome sequence of each species and the GC content of the respective splice site combination was calculated using the Python package scipy. Splice site combinations were weighted with the number of occurences while calculating the GC content. Finally, the correlation coefficient and the p-value were determined. For better visualization, a scatter plot was constructed with the Python package plotly [56].

⁴¹⁰ Phylogeny of non-canonical splice sites

All A. thaliana transcripts with non-canonical splice sites were subjected 411 to BLASTn searches against the transcript sequences of all other plant 412 species previously studied [10]. The best hit per species was selected for 413 an alignment against the respective genomic region with exonerate [59]. 414 Next, splice site combinations were extracted and aligned. This align-415 ment utilized MAFFT v7 [60] by representing different splice site com-416 binations as amino acids. Finally, splice site combinations aligned with 417 the non-canonical splice site combinations of A. thaliana were analysed 418 (https://doi.org/10.5281/zenodo.2586989). 419

420 Usage of non-canonical splice sites

Genes were classified based on the presence/absence of non-canonical splice combinations into four groups: GT-AG, GC-AG, AT-AC, and minor non-canonical splice site genes. When having different non-canonical

splice sites, genes were assigned into multiple groups. Next, the tran-424 scription of these genes was quantified based on RNA-Seq using feature-425 Counts [61] based on the RNA-Seg read mapping generated with STAR. 426 Binning of the genes was performed based on the fragments per kilobase 427 transcript length per million assigned reads (FPKMs). Despite various 428 shortcomings [62], we consider FPKMs to be acceptable for this analysis. 429 Outlier genes with extremely high values were excluded from this analysis 430 and the visualization. Next, a cumulative sum of the relative bin sizes was 431 calculated. The aim was to compare the transcriptional activity of genes 432 with different splice site combinations i.e. to test whether non-canonical 433 splice site combinations are enriched in lowly transcribed genes. 434

Usage of splice sites was calculated per intron as previously described 436 [10]. The difference between both ends of an intron was calculated. The 437 distribution of these differences per splice site type were analysed. In-438 trons were grouped by their splice site combination. The average of both 439 coverage values of the directly flanking exon positions was calculated as 440 estimate of the local expression around a splice site combination. Next, 441 the sequencing coverage of a transcript was estimated by multiplying 200 442 bp (assuming 2x100 nt reads) with the number of read counts per gene 443 and normalization to the transcript length. The difference between both 444 values was calculated for each intron to assess its presence in the major 445 isoform. 446

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Additional Files

AdditionalFile 1. List of genome sequences and annotations of the investigated animal
 species.

AdditionalFile 2. List of genome sequences and annotations of the investigated fungal species.

652 AdditionalFile 3. Distribution of genome sizes of all species.

AdditionalFile 4. Distribution of canonical and non-canonical splice sites per species in
 the animal kingdom.

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AdditionalFile 5. Distribution of canonical and non-canonical splice sites per species in
 the fungal kingdom.

660 AdditionalFile 6. List of all possible splice site combinations in animal species.

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662 AdditionalFile 7. List of all possible splice site combinations in fungal species.

664 AdditionalFile 8. Genome statistics concerning each analysed animal species.

666 **AdditionalFile 9.** Genome statistics concerning each analysed fungal species.

667 668 **AdditionalFile 10.** Correlation between the GC content of the genome and the GC con-669 tent of the splice sites per kingdom.

AdditionalFile 11. Correlation between the size of the used RNA-Seq data sets and the
 number of supported splice sites.

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AdditionalFile 12. Proportion of genes with non-canonical splice sites in dependence of the number of introns.

677 AdditionalFile 13. Usage of non-canonical splice sites in animals and fungi.

AdditionalFile 14. Non-canonical splice sites in BUSCOs and in all genes were assessed per species in the animal kingdom.

AdditionalFile 15. Non-canonical splice sites in BUSCOs and in all genes were assessed per species in the fungal kingdom.

AdditionalFile 16. List of Sequence Read Archive accession numbers of the investigated animal RNA-Seq data sets.

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AdditionalFile 17. List of Sequence Read Archive accession numbers of the investigated
 fungal RNA-Seq data sets.

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