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Analysis of pollen-specific alternative splicing in *Arabidopsis thaliana* via semi-quantitative PCR

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ABSTRACT

Alternative splicing enables a single gene to produce multiple mRNA isoforms by varying splice site selection. In animals, alternative splicing of mRNA isoforms between cell types is widespread and supports cellular differentiation. In plants, at least 20% of multi-exon genes are alternatively spliced, but the extent and significance of tissue-specific splicing is less well understood, partly because it is difficult to isolate cells of a single type. Pollen is a useful model system to study tissue-specific splicing in higher plants because pollen grains contain only two cell types and can be collected in large amounts without damaging cells. Previously, we identified pollen-specific splicing patterns by comparing RNA-Seq data from *Arabidopsis* pollen and leaves. Here, we used semi-quantitative PCR to validate pollen-specific splicing patterns among genes where RNA-Seq data analysis indicated splicing was most different between pollen and leaves. PCR testing confirmed eight of nine alternative splicing patterns, and results from the ninth were inconclusive. In four genes, alternative transcriptional start sites coincided with alternative splicing. This study highlights the value of the low-cost PCR assay as a method of validating RNA-Seq results.

INTRODUCTION

Alternative splicing is a form of post- and co-transcriptional regulation in which pre-mRNA transcripts from the same gene are spliced in different ways. Alternative splicing can affect any part of a transcript, and it has been well known since the early 2000s that alternative splicing in mammalian genomes frequently remodels conserved or functional motifs in the encoded proteins (Loraine et al., 2002; Cline et al., 2004). These differences can have important functional consequences; in animals as diverse as fruit flies and mammals, differential expression of alternative splice variants supports and enforces cellular differentiation and development. Textbook examples of regulatory processes that depend on differential expression of splice

variants include sex determination in insects, neuronal differentiation in mammals, and aspects of hormone signaling (reviewed in (Li et al., 2007; Zheng and Black, 2013)).

In higher plants, far less is known about tissue or cell-type specific alternative splicing. This is partly due to a lack of data, as well as the technical difficulty in collecting large amounts of the same cell type from plants. Until recently, ESTs were the main source of information about splice variants, and several studies used ESTs to investigate the diversity and prevalence of alternative splicing in plants (Ewing et al., 1999; Campbell et al., 2006; Schindler et al., 2008; Wang et al., 2008; English et al., 2010). However, in contrast to human and mouse genomes, for which tens of millions of ESTs are available, the best-studied plants (rice, maize, and Arabidopsis) have fewer than four million ESTs combined, which meant that far less has been learned about alternative splicing in plants compared to animals. Now, thanks to high throughput cDNA sequencing (RNA-Seq) and improving bioinformatics approaches, limited data is becoming less of a problem. However, another challenge arises from the unique biology of plants in comparison to animals; plant cells inhabit a complex matrix of cell wall materials that make it difficult to prepare homogenous cell populations. The integrity and composition of the cell wall plays a key role in regulation of plant cellular processes, especially defense responses, which involve changes in splicing patterns (Howard et al., 2013). Thus, there is a possibility that cell isolation methods that involve degradation of the cell wall may introduce artifacts leading to over- or under-estimation of tissue- or cell-type specific alternative splicing.

Fortunately, there are some tissue or cell types in plants where the sample collection problem can be overcome. Pollen grains are one example; in most plants, pollen is well suited to collecting cellular materials in amounts required for preparing cDNA libraries for RNA-Seq (Johnson-Brousseau and McCormick, 2004). During pollen development, the pollen grain becomes dehydrated and enters a dormant state, and so mature pollen can be collected in large amounts with minimal risk of introducing artifacts, in part because collection procedures are typically no harsher than natural mechanisms by which pollen is spread by insects or wind. In Arabidopsis, similar to other flowering plants, pollen grains contain three cells: two sperm cells and a vegetative cell, also called the pollen tube cell, which encloses the sperm cells and gives rise to the pollen tube (reviewed in (Dresselhaus and Franklin-Tong, 2013). When a pollen grain lands on a compatible stigma, the grain rehydrates and germinates, extruding a pollen tube that grows through the style toward the ovary, carrying the sperm cells. Upon entering the ovary, the pollen tube bursts, releasing the sperm, which merge with the central cell and the egg, giving rise to endosperm and embryo. Experiments with *in vitro* pollen germination and inhibitors of transcription have shown that Arabidopsis pollen grains contain sufficient mRNA to support the early stages of germination and tube growth, but synthesis of new RNA is required to complete fertilization (Honys and Twell, 2003; Wang et al., 2008) with gene expression changing during tube growth (Honys and Twell, 2003, 2004; Pina et al., 2005; Wang et al., 2008; Qin et al., 2009; Boavida et al., 2011). Thus pollen offers an opportunity to

69 identify and study tissue-specific splicing in plants and how splicing changes during a well-studied
70 developmental process.

71 Previously we used high throughput sequencing of cDNA (RNA-Seq) to investigate gene expression and
72 splicing patterns in Arabidopsis pollen (Loraine et al., 2013). By comparing gene models and RNA-Seq read
73 alignments, we identified around thirty genes that were annotated as alternatively spliced, were expressed in
74 both pollen and leaves, and where the relative abundance of splice site use differed between pollen and leaves.
75 Thus, these genes represented candidates for tissue-specific splicing in Arabidopsis. Here, we report results from
76 new experiments that tested alternative splicing of a subset of these genes using semi-quantitative PCR. Our
77 results confirmed tissue-specific splicing patterns, and also indicated that for many genes where alternative
78 splicing affected the 5' region, alternative transcriptional start sites (TSSs) may also play an important role in
79 generating transcript diversity.

81 MATERIALS AND METHODS

83 Bioinformatic analysis of splicing patterns

84 Our previously published report described using RNA-Seq alignments to identify differential alternative
85 splicing in pollen and leaf RNA-Seq data sets (Loraine et al., 2013). Briefly, alternative splicing (AS) events
86 annotated as part of the TAIR10 gene models from *Arabidopsis thaliana* were identified by comparing regions
87 of overlap between exons and introns in different gene models belonging to the same gene, as described in
88 (English et al., 2010). For each alternative splicing event, the number of spliced reads supporting mutually
89 exclusive choices was identified for each of two leaf libraries and one pollen library. For simplicity, only
90 annotated alternative donor and acceptor sites were considered, and alternative splice forms were compared as
91 pairs. For each AS event, the percentage of gapped reads in each library that supported each form was calculated
92 and compared between pollen and leaf samples. Alternative splicing events were identified in which percentage
93 of reads supporting a given splicing choice varied more than 20% between the two sample types. The AS events
94 were then ordered by this percent difference; the listing of these events is shown in Supplemental Data File S1.
95 The ten most differentially spliced genes were identified for testing by PCR; the list included At-U2AF65A,
96 which was tested previously and is not discussed again here (Loraine et al., 2013). Note also that previously we
97 referred to the leaf samples as “seedlings,” but here refer to them as “leaf” samples. This is because the older
98 terminology was confusing to some readers. For many people the term “seedling” means: whole seedlings that
99 are two weeks old or younger. The samples used in here and in the previous study were from three-week old,
100 soil-grown plants and included leaves and not roots.

102 PCR primer design for analysis of splicing patterns

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PCR amplifications included an initial “hot start” 4 minute denaturation incubation at 94°C followed by 30 cycles of denaturation (94°C), annealing, and elongation (72°C) incubations. Annealing temperatures were selected based on T_m predicted by the Primer 3 software, and when primers failed to amplify a band, gradient PCR was performed to identify the optimal annealing temperature. Amplifications were typically performed in volumes of 20 μ L per template/primer combination with 2 μ L each of 5 μ M forward and reverse primer stock solutions, .08 μ L of taq polymerase (Bioline, catalog number 21086), and 0.5 μ L aliquots of pollen or leaf cDNA reactions that had been diluted 1:20. Cycle numbers were estimated to be in logarithmic phase based on prior analysis of qPCR analysis of genes of varying starting RPKMs (Loraine et al., 2013).

Analysis of gel images for semi-quantitative PCR

The relative abundance of splice variants amplified by PCR was estimated using methods presented in (Palusa et al., 2007; Venables et al., 2012). Equal amounts of PCR reaction mixtures were fractionated by agarose gel electrophoresis containing ethidium bromide. Gels were visualized under short wave UV light in a UVP Biospectrum AC Imaging System GelDoc station and photographed using a digital camera. Each gel was typically photographed at different run times and using different exposure times to obtain high quality images for analysis.

Gel images were analyzed using the VisionWorksLS version 7.0.1 software program. Rectangular regions surrounding ethidium-stained bands in each lane were identified, and for each band, the software was used to estimate molecular weight along with a background-corrected band intensity value, called “I-vol” in the software. Visualization of lane profiles was also done to ensure that band intensity measurements were within the range of detection of the instrument. Gels also included lanes containing molecular mass standards (Hyperladder 1 kb, Bioline catalog number BIO-33053). Band intensities of the molecular size/mass markers were plotted against their masses and only gels where this relationship was close to linear ($R^2 \geq 0.90$) were used. Band intensities and size estimates were then imported into the R statistical programming environment for analysis.

For each band, its abundance relative to other bands in the same lane was calculated and used to test for differences between pollen and leaf samples. The relative abundance of per band was calculated as its percent contribution the total amount of DNA in each lane. The percent contribution per band was calculated as $100 \times b/T$, where b was the band’s size-normalized intensity and T was the sum of all size-normalized band intensities in the lane, calculated as $\sum_{i=1}^N b_i$. Size-based normalization of band intensity was done to account for how ethidium staining (and band brightness) depended on both amplicon abundance and amplicon size, since larger amplicons contained more sites (base pairs) for ethidium to bind (Green and Sambrook, 2012). For each gel containing replicate pollen and leaf samples, a t test was used to determine whether there was a statistically significant difference in a band’s percentage between pollen and leaf samples. As the prior RNA-Seq data

analysis indicated the whether pollen or leaf percentages were greater and since there was no reason to expect unequal variance, one-sided t-tests assuming equal variance were used. R code used to calculate percentages and perform statistical testing are available as Markdown (.Rmd) in the folder named “GelAnalysis” in the project repository at <http://www.bitbucket.org/lorainelab/pollenas>. For each gene tested, a Markdown file and its corresponding HTML report file is available; readers can re-run the code or open the HTML file in a Web browser to view the code, the output of the code, and accompanying figures.

Sanger sequencing of PCR products

For higher-abundance amplicons that did not match annotated gene models or that suggested use of alternative transcription start sites, PCR products were extracted from the gels and sequenced by conventional Sanger sequencing to confirm band identity. Sequence reads obtained from Sanger sequencing were aligned on the Arabidopsis genome using blat (Kent, 2002) and visualized in Integrated Genome Browser for comparison with gene models. Blat was run with default settings except for the maxIntron parameter, which was set to 2000, since 99% of Arabidopsis introns are 2000 bases or smaller. Sequencing results are described in detail in the folder named “AmpliconSequencing” in the project repository at <http://www.bitbucket.org/lorainelab/pollenas>.

Data Access

The RNA-Seq data are available from the Sequence Read Archive under SRA accessions SRR847501, SRR847502 (pollen library); SRR847503, SRR847504 (seedling library 1); and SRR847505, SRR847506 (seedling library 2). Read and junction feature alignments depicted in figures are available from the IGBQuickLoad data source named “pollen” under the Arabidopsis June 2009 (TAIR9/10) genome assembly release. A tutorial introducing Integrated Genome Browser and describing how to perform a visual splicing analysis using the pollen RNA-Seq data set is available at <http://tinyurl.com/pk25nbm>.

RESULTS

To further investigate pollen-specific splicing patterns, we first revisited the analysis described in our previous study of gene expression in pollen using RNA-Seq data (Loraine et al., 2013). We identified 30 genes (Data S1) that were expressed in both pollen and leaves where the prevalence of at least one splicing variant differed 20% or more between the two samples types. From this list, we selected the most highly differentially spliced genes for further testing via semi-quantitative PCR, an established method for assaying alternative splicing (Ali et al., 2007; Venables et al., 2012). In semi-quantitative PCR, primers that flank the differentially spliced region are used to amplify splice variants from cDNA prepared from different samples, ideally multiple biological replicates. PCR products are run out on a gel, visualized by staining with ethidium bromide, and then the relative brightness of each amplicon (band) is quantified using image analysis software (Green and

Sambrook, 2012). The percent contribution of each band to the total signal in each lane is compared across lanes; if the percent abundance of a band increases, for example, from 10% to 90%, this indicates a change in splicing.

An advantage of semi-quantitative PCR for assessing splicing changes is that because the values being compared are percentages, expressed relative to the total amount of product per lane, the amount of cDNA used as template in each reaction can vary between lanes; no across-lane normalization is required. Another advantage is that PCR and gel electrophoresis are inexpensive and do not require specialized equipment. Using this method, we previously confirmed that one of the ten most differentially spliced genes (U2AF65A) was differentially spliced between pollen versus leaves. Here, we used semi-quantitative PCR supplemented with Sanger sequencing of amplicons to test differential splicing of the nine other most differently spliced genes identified by RNA-Seq.

Splice variants from Arabidopsis SR genes At-SR30 (AT1G09140) and At-RS41 (AT5G52040) are differentially spliced in pollen and leaves

At-SR30 and At-RS41 encode two of eighteen serine/arginine-rich (SR) RNA-binding proteins in Arabidopsis (Barta et al., 2010) thought to play a role in regulation of splice site choice. According to the RNA-Seq data, At-SR30 (Fig. 1A) and At-RS41 (Fig. 2A) were differentially spliced in pollen and leaf. Figures 1A and 2A show junction features aligned onto each gene, where features summarize the number of spliced reads in each sample that supported the corresponding intron in the gene models. In both genes, the relative abundance of spliced reads supporting alternative splicing events was clearly different in leaf versus pollen. We also observed a small number of unspliced reads that aligned to the differentially spliced intron, indicating a low level of intron retention (not shown), a common form of alternative splicing in plants (English et al., 2010).

To validate the RNA-Seq results, we designed PCR primers flanking the differentially spliced region and performed semi-quantitative PCR using newly prepared pollen and leaf cDNA samples. For both genes, semi-quantitative PCR confirmed the differential splicing patterns found by RNA-Seq. Figure 1B-C shows that two splice variants of At-SR30 were expressed in both pollen and leaf samples, but the relative abundance of the two variants was different. We also observed a low abundance, higher molecular weight amplicon in pollen and leaf samples consistent with intron retention. For At-RS41, we observed similar results; PCR amplified four splice variants, the relative abundance of the splice variants was different between pollen and leaves, and low abundance amplicons consistent with intron retention were apparent (Fig. 2B-C).

In both genes, differential splicing affected the protein-coding region of the transcripts, but in different ways. In At-SR30, the leaf-preferred isoform introduced a premature stop codon that deleted 23 amino acids from the carboxyl end of the protein, including five SR repeats, which are possible phosphorylation targets and mediate interactions with other proteins (reviewed in (Shepard and Hertel, 2009). For At-RS41, the pollen-

preferred isoform included an alternative exon that introduced a new translation start site leading to loss of 35 N-terminal amino acids encoding part of an RNA recognition motif. Thus, the pollen-preferred form of At-SR30 encoded the full-length protein, while the pollen-preferred form of At-RS41 encoded a truncated form lacking part of the amino terminal region of the protein.

Alternative splicing in pollen remodels starch synthase gene AT5G65685

AT5G65685, encoding a starch synthase-like protein, is highly alternatively spliced in the 5' region (Fig. S1A). Visualization of RNA-Seq alignments suggested there were many possible splice variants affecting the amino terminus of the protein and that the splicing patterns in pollen and leaves were different. According to the RNA-Seq data, the first intron (relative to isoform AT5G65685.1) contained two alternative donor sites that were differentially spliced in pollen compared to leaves, with leaf samples using the upstream site and pollen samples using the downstream site. The RNA-Seq data also suggested there were several unannotated variants containing retained introns. Semi-quantitative PCR corroborated all of these observations (Fig. S1B-C). PCR primers designed around the alternatively spliced region amplified multiple bands corresponding to annotated variants and to many novel, lower abundance variants corresponding to retained intron isoforms.

Alternative splicing of AT4G21720, gene of unknown function

AT4G21720 encodes a conserved protein of unknown function with two annotated splice variants., with alternative acceptor sites affecting the 3' end of the second intron (Fig. S2A). Visualization of RNA-Seq read alignments in Integrated Genome Browser indicated that the upstream acceptor site, which introduced a frameshift that eliminated 44 amino acids, was preferred in pollen, and the downstream acceptor site, which allowed translation of the full-length protein, was preferred in leaves.

As before, primers flanking the alternatively spliced region were designed and used to amplify cDNAs prepared from pollen and leaves (Fig. S2B-C). The PCR assay results supported differential splicing of AT4G21720 in pollen versus leaves. PCR products consistent with both acceptor sites were observed and the relative abundance of the products was similar to what was observed in the RNA-Seq data (Fig. S2B-C and Data S1). An additional higher molecular weight band was also observed in both samples, and its size suggested it was derived from occasional retention of the alternatively splice intron. However, its abundance was extremely low.

5' Alternative splicing coincided with alternative TSSs in four of nine genes tested

AT3G17120, another gene of unknown function, is annotated as producing two splice variants due to exon skipping in the 5' UTR of the gene, and the RNA-Seq data analysis indicated that the relative abundance of splice variants was different in pollen and leaves (Data S1). However, visual inspection of RNA-Seq alignments

alongside the gene models suggested the gene contained an alternative transcription start site (TSS) located within the first annotated intron (Fig. 3A). We tested this in PCR experiments using variant specific primers F1 and F2 and a common primer R. In leaves, F1 with R amplified a product, but F2 and R did not (Fig. 3B). In pollen, F2 and R amplified a product, but F1 with R did not. Combining all three primers in the same reaction amplified leaf and pollen-specific products, and sequencing confirmed the identity of products shown in Figure 3B. This confirmed that pollen and leaves expressed distinct isoforms and strongly suggested that the alternative transcripts were from alternative transcriptional start sites. Because the putative alternative TSS preferred in pollen was downstream of the donor site in the first intron, that donor site was not present in the pollen-specific transcript, which meant that splicing of the first intron in pollen required a different, alternative donor site.

Alternative splicing of AT3G06330, encoding a putative RING-type ubiquitin ligase, also appeared to coincide with alternative TSSs. AT3G06330 is annotated as having an alternative donor site in the 5' end of the gene that changes the amino terminal region of the protein. According to the RNA-Seq data, the downstream donor site was preferred in pollen, while the upstream donor site was preferred in leaves (Fig. S3A). As before, we designed variant-specific forward primers and used them together with a common reverse primer in PCR reactions with pollen and leaf cDNAs. Reactions that included all three primers amplified an approximately 455 bp product in both leaves and pollen and an approximately 395 bp product in pollen only. Comparison with gene models suggested that the lower-molecular weight pollen product might correspond to a transcript with splicing pattern similar to AT3G06330.2 but which initiated transcription at different location, as if the annotated start site were used, we would have observed an additional larger band corresponding to primers F1 and R. These results suggested the models shown in Figure S3B, and Sanger sequencing of the PCR products confirmed them. These results suggest two alternative TSSs were used in pollen but only the upstream site was used in leaves (Fig. S3B-C).

A similar result was obtained for RH11, an alternatively spliced gene encoding a DEAD/H-box RNA helicase. Inspection of the RNA-Seq alignments suggested there was at least one alternative TSS, and this alternative TSS enforced the use of an alternative donor site in the first intron (Fig. S4A). Variant-specific forward primers were used with a common reverse primer to amplify the differentially spliced regions. The variant-specific primers amplified product in both leaves and pollen, indicating that both samples used the alternative TSSs, but pollen and leaf samples each favored a different variant (Fig. S4B-C).

Lastly, we also found evidence for alternative TSSs in AT1G67480, which encodes a galactose oxidase/kelch repeat superfamily protein. The RNA-Seq data indicated that the annotated gene models were correct, but also suggested that other isoforms were expressed (Fig. S5A). The pollen data contained reads indicating a novel splice form, a retained-intron form, and two possible alternative TSSs. Semi-quantitative PCR amplification supported these alternative forms (data not shown). Primers designed to amplify the annotated, differentially spliced region produced amplicons with sizes that were consistent with predictions from

the RNA-Seq data (Fig. S5B). Multiple variants were co-expressed in pollen and leaves, but their relative abundance was different between the sample types (Fig. S5C).

Semi-quantitative PCR analysis of AT2G19480, a NAP1 histone chaperone, failed to show pollen specific splicing

AT2G19480 (NAP1;4) encodes a NAP1 histone chaperone (Liu et al., 2009; Gao et al., 2012) with three splice variants that differ in the three prime region. According the RNA-Seq data, an exon-skipped form was preferred in pollen and the non-skipped, retained form was preferred in leaves (Fig. S6A). PCR primers were designed to amplify all forms (F and R2 in Fig. S6A) or just the exon-included form (F and R1 in Fig. S6A). However, the PCR amplifications contradicted the RNA-Seq results in that we found evidence for the exon-skipped form only in each sample (data not shown). Thus, the PCR assessment of alternative splicing confirmed existence and expression of the exon-skipped form in both pollen and leaves but was unable to identify the exon-retained form.

On-line data repository and RNA-Seq visualization resource

To enable readers to reproduce analyses, we assembled a git-based data and source code repository at <http://www.bitbucket.org/lorainelab/pollenas>. Readers familiar with the git version control program can clone the repository onto their local desktop, thus enabling them to view and retrieve older versions of the repository or use it as a basis for their own work. Tutorials on using git are available on the bitbucket.org Web site and many others. However, it is not necessary to use the git program to obtain a copy. Readers can also download the repository as a single “zip” by clicking the “download” icon on the project home page. Files can also be downloaded individually by clicking the “source” icon and then browsing the repository files and folders.

Analyses described above are organized into folders named for their contents. Typically, each folder contains a “results” sub-folder containing computer-generated results files and a “data” sub-folder containing primary data files obtained from the VisionWorks software or sequencing facility. Most analyses reported here were done using the R programming language and are documented in R “Markdown” (.Rmd) report-generation files that enable results and analysis code to be shown together in the same report file. Each “Markdown” file has a corresponding HTML report output file that was generated using the knitr R library, a tool for literate programming in R. As such, readers can either re-run the code themselves or can open the report file in a Web browser, after downloading the repository. Note that re-running analyses will also require installing some libraries that are not part of the base R distribution; this can be done using the “install packages” command within R.

We also provided RNA-Seq alignment files for visualization in Integrated Genome Browser as part of a pollen-specific IGBQuickLoad site at <http://igbquickload.org/pollen>. To view the data in IGB, readers first should obtain a copy of the software from the IGB Web site at <http://www.bioviz.org/igb>. The software can be

downloaded for free from the site; installers are available for Apple, Windows, and Linux computers. Once IGB is running, users can click the Arabidopsis image on the IGB home screen; this opens the Arabidopsis TAIR10 genome and loads the Arabidopsis reference gene annotations. To load RNA-Seq data shown in the figures, users can open the Pollen data source under the IGB Data Access tab. The data sets described here are named Pollen, Leaf 1, and Leaf 2. Alignments of sequences obtained from Sanger sequencing of amplicons can also be loaded. To view these data, readers can download the alignments file (named Amplicons.pslx) from the data and source code repository and open them in IGB as with any other file.

To illustrate the potential usefulness of this resource, we examined RNA-Seq alignments overlapping U2AF65B, one of two genes encoding U2AF65, the large subunit of the U2 snRNP accessory factor dimer, which helps to define 3' splice sites by recruiting the U2 snRNP to the pre-mRNA. Genes that encode U2AF65 in Arabidopsis include U2AF65A (AT4G36690) and U2AF65B (AT1G60900). Previously, we showed that U2AF65A is differentially spliced in pollen as compared to leaf, but we did not report on alternative splicing of U2AF65B because there is only one model associated with this gene, and our splicing analysis pipeline detected differential splicing only in genes with multiple models. We therefore visually inspected RNA-Seq read alignments overlapping U2AF65B. Visualization of the RNA-Seq read alignments suggested that U2AF65B encodes at least one other splice variant thanks to alternative splicing of the fourth intron (Figure 4). However, because the gene was not abundantly expressed in pollen, it was impossible to determine from RNA-Seq data alone whether the gene is differently spliced in pollen versus leaf.

DISCUSSION

To better characterize alternative splicing patterns in pollen, we performed semi-quantitative PCR testing of several genes that prior bioinformatic analysis of RNA-Seq data had indicated were differentially spliced between pollen and Arabidopsis leaves. For each gene, read alignments and gene models were visualized using the Integrated Genome Browser (Nicol et al., 2009). In each case, visual inspection led to deeper understanding of splicing patterns and their possible effects on protein sequence and biological function. PCR testing of independently prepared pollen and leaf samples confirmed pollen-specific splicing patterns for eight of nine genes tested. Two genes with validated pollen-specific splicing patterns encoded SR proteins (splicing regulators that influence splice site choice) and alternative splicing of the pollen-enriched forms produced isoforms with altered conserved motifs important for function. In four genes, PCR analysis showed that alternative splicing choices coincided with possible alternative transcriptional start sites, which enforced selection of alternative donor sites at the 5' ends of the affected genes. Table 1 summarizes the results.

Of the nine genes tested, only one gene (AT2G19480, nucleosome assembly protein) was not confirmed as differentially spliced between pollen and leaves. However, this negative result may have been due to limitations of the PCR-based assay. The differentially spliced variants differed by only 18 bases, which made

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REFERENCES

- Ali GS, Palusa SG, Golovkin M, Prasad J, Manley JL, Reddy AS** (2007) Regulation of plant developmental processes by a novel splicing factor. *PLoS One* **2**: e471
- Barta A, Kalyna M, Reddy AS** (2010) Implementing a rational and consistent nomenclature for serine/arginine-rich protein splicing factors (SR proteins) in plants. *Plant Cell* **22**: 2926-2929
- Boavida LC, Borges F, Becker JD, Feijo JA** (2011) Whole genome analysis of gene expression reveals coordinated activation of signaling and metabolic pathways during pollen-pistil interactions in *Arabidopsis*. *Plant Physiol* **155**: 2066-2080
- Campbell MA, Haas BJ, Hamilton JP, Mount SM, Buell CR** (2006) Comprehensive analysis of alternative splicing in rice and comparative analyses with *Arabidopsis*. *BMC Genomics* **7**: 327
- Cline MS, Shigeta R, Wheeler RL, Siani-Rose MA, Kulp D, Loraine AE** (2004) The effects of alternative splicing on transmembrane proteins in the mouse genome. *Pac Symp Biocomput*: 17-28
- Dresselhaus T, Franklin-Tong N** (2013) Male-female crosstalk during pollen germination, tube growth and guidance, and double fertilization. *Molecular plant* **6**: 1018-1036
- English AC, Patel KS, Loraine AE** (2010) Prevalence of alternative splicing choices in *Arabidopsis thaliana*. *BMC Plant Biol* **10**: 102
- Ewing RM, Ben Kahla A, Poirot O, Lopez F, Audic S, Claverie JM** (1999) Large-scale statistical analyses of rice ESTs reveal correlated patterns of gene expression. *Genome Res* **9**: 950-959
- Gao J, Zhu Y, Zhou W, Molinier J, Dong A, Shen WH** (2012) NAP1 family histone chaperones are required for somatic homologous recombination in *Arabidopsis*. *Plant Cell* **24**: 1437-1447
- Green MR, Sambrook J** (2012) *Molecular Cloning: A Laboratory Manual*, 4th Edition, Vol 1. Cold Spring Harbor Laboratory Press, Cold Spring Harbor
- Gulledge AA, Roberts AD, Vora H, Patel K, Loraine AE** (2012) Mining *Arabidopsis thaliana* RNA-seq data with Integrated Genome Browser reveals stress-induced alternative splicing of the putative splicing regulator SR45a. *Am J Bot* **99**: 219-231
- Honys D, Twell D** (2003) Comparative analysis of the *Arabidopsis* pollen transcriptome. *Plant Physiol* **132**: 640-652
- Honys D, Twell D** (2004) Transcriptome analysis of haploid male gametophyte development in *Arabidopsis*. *Genome Biol* **5**: R85
- Howard BE, Hu Q, Babaoglu AC, Chandra M, Borghi M, Tan X, He L, Winter-Sederoff H, Gassmann W, Veronese P, Heber S** (2013) High-throughput RNA sequencing of pseudomonas-infected *Arabidopsis* reveals hidden transcriptome complexity and novel splice variants. *PLoS One* **8**: e74183
- Johnson-Brousseau SA, McCormick S** (2004) A compendium of methods useful for characterizing *Arabidopsis* pollen mutants and gametophytically-expressed genes. *Plant J* **39**: 761-775
- Kent WJ** (2002) BLAT--the BLAST-like alignment tool. *Genome Res* **12**: 656-664
- Li Q, Lee JA, Black DL** (2007) Neuronal regulation of alternative pre-mRNA splicing. *Nat Rev Neurosci* **8**: 819-831

- 453 **Liu Z, Zhu Y, Gao J, Yu F, Dong A, Shen WH** (2009) Molecular and reverse genetic characterization of
- 454 NUCLEOSOME ASSEMBLY PROTEIN1 (NAP1) genes unravels their function in transcription and
- 455 nucleotide excision repair in *Arabidopsis thaliana*. *Plant J* **59**: 27-38
- 456 **Loraine AE, Helt GA, Cline MS, Siani-Rose MA** (2002) Protein-based analysis of alternative splicing in the
- 457 human genome. *Proc IEEE Comput Soc Bioinform Conf* **1**: 118-124
- 458 **Loraine AE, McCormick S, Estrada A, Patel K, Qin P** (2013) RNA-seq of *Arabidopsis* pollen uncovers novel
- 459 transcription and alternative splicing. *Plant Physiol* **162**: 1092-1109
- 460 **Nicol JW, Helt GA, Blanchard SG, Jr., Raja A, Loraine AE** (2009) The Integrated Genome Browser: free
- 461 software for distribution and exploration of genome-scale datasets. *Bioinformatics* **25**: 2730-2731
- 462 **Palusa SG, Ali GS, Reddy AS** (2007) Alternative splicing of pre-mRNAs of *Arabidopsis* serine/arginine-rich
- 463 proteins: regulation by hormones and stresses. *Plant J* **49**: 1091-1107
- 464 **Pina C, Pinto F, Feijo JA, Becker JD** (2005) Gene family analysis of the *Arabidopsis* pollen transcriptome
- 465 reveals biological implications for cell growth, division control, and gene expression regulation. *Plant*
- 466 *Physiol* **138**: 744-756
- 467 **Qin Y, Leydon AR, Manziello A, Pandey R, Mount D, Denic S, Vasic B, Johnson MA, Palanivelu R** (2009)
- 468 Penetration of the stigma and style elicits a novel transcriptome in pollen tubes, pointing to genes critical
- 469 for growth in a pistil. *PLoS Genet* **5**: e1000621
- 470 **Schindler S, Szafranski K, Hiller M, Ali GS, Palusa SG, Backofen R, Platzer M, Reddy AS** (2008)
- 471 Alternative splicing at NAGNAG acceptors in *Arabidopsis thaliana* SR and SR-related protein-coding
- 472 genes. *BMC Genomics* **9**: 159
- 473 **Shepard PJ, Hertel KJ** (2009) The SR protein family. *Genome biology* **10**: 242
- 474 **Simpson CG, Fuller J, Maronova M, Kalyna M, Davidson D, McNicol J, Barta A, Brown JW** (2008)
- 475 Monitoring changes in alternative precursor messenger RNA splicing in multiple gene transcripts. *The*
- 476 *Plant journal : for cell and molecular biology* **53**: 1035-1048
- 477 **Takahashi H, Lassmann T, Murata M, Carninci P** (2012) 5' end-centered expression profiling using cap-
- 478 analysis gene expression and next-generation sequencing. *Nat Protoc* **7**: 542-561
- 479 **Venables JP, Vignal E, Baghdiguian S, Fort P, Tazi J** (2012) Tissue-specific alternative splicing of Tak1 is
- 480 conserved in deuterostomes. *Mol Biol Evol* **29**: 261-269
- 481 **Wang BB, O'Toole M, Brendel V, Young ND** (2008) Cross-species EST alignments reveal novel and
- 482 conserved alternative splicing events in legumes. *BMC Plant Biol* **8**: 17
- 483 **Wang Y, Zhang WZ, Song LF, Zou JJ, Su Z, Wu WH** (2008) Transcriptome analyses show changes in gene
- 484 expression to accompany pollen germination and tube growth in *Arabidopsis*. *Plant Physiol* **148**: 1201-
- 485 1211
- 486 **Zheng S, Black DL** (2013) Alternative pre-mRNA splicing in neurons: growing up and extending its reach.
- 487 *Trends Genet* **29**: 442-448
- 488
- 489

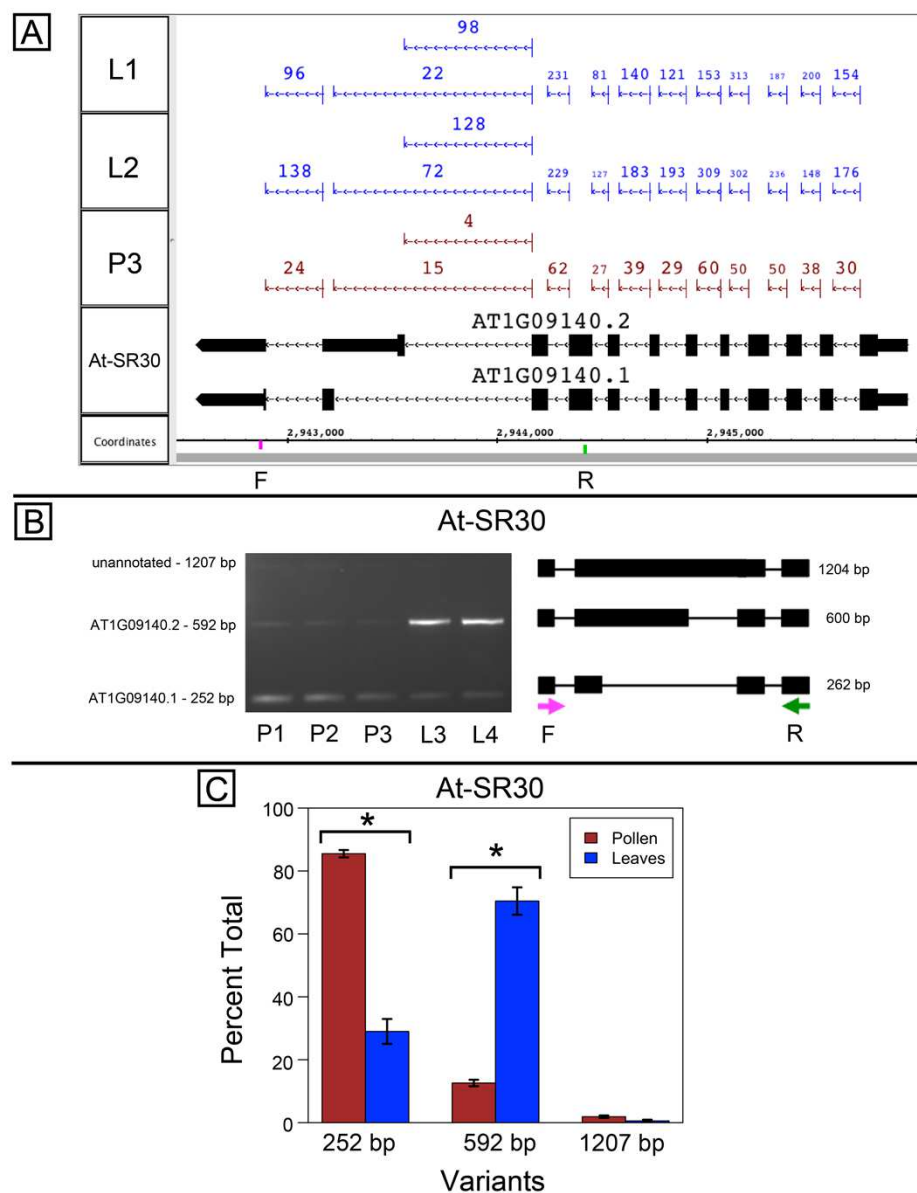


Figure 1. Pollen-specific splicing in At-SR30. (A) Junction features from RNA-Seq reads for leaf (L1, L2) and pollen (P3) samples alongside annotated gene models. Numbers indicate how many spliced reads supported the indicated junction in pollen and leaf RNA-Seq libraries. Arrows indicate the direction of transcription and taller blocks show translated regions. Primers used in semi-quantitative PCR are shown. Primer sequences were CCAGTGGCCAGTTTTTCATTT (F) and GTGTGAGTCGAAGCCCAGAT (R). (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs and corresponding model of alternative splice variants. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.

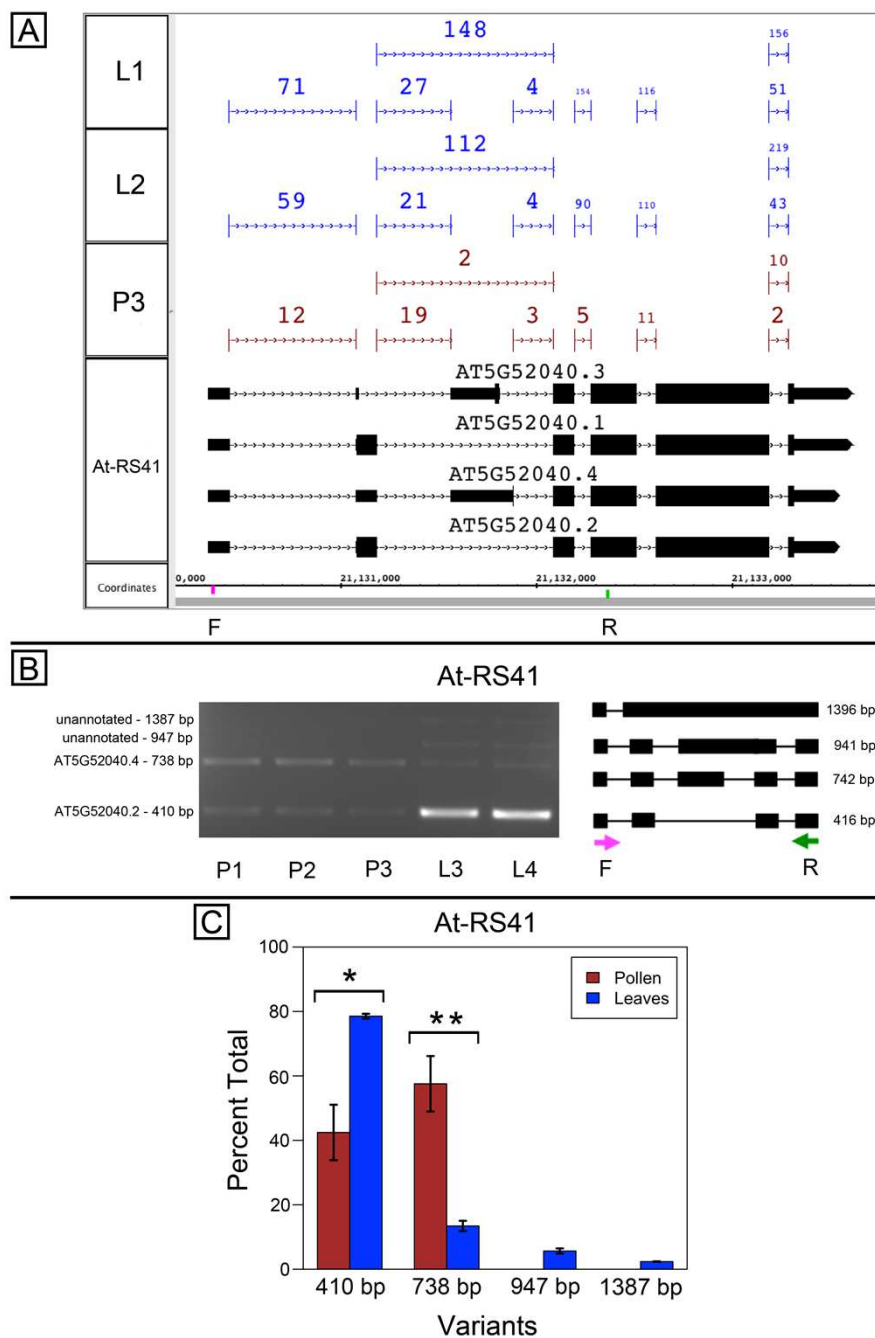


Figure 2. Pollen-specific splicing in At-RS41. (A) Junction features from RNA-Seq reads for leaf (L1, L2) and pollen (P3) samples alongside annotated gene models. Numbers indicate how many spliced reads supported the indicated junction in pollen and leaf RNA-Seq libraries. Primers used in semi-quantitative PCR are shown. (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs and corresponding model of alternative splice variants. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.

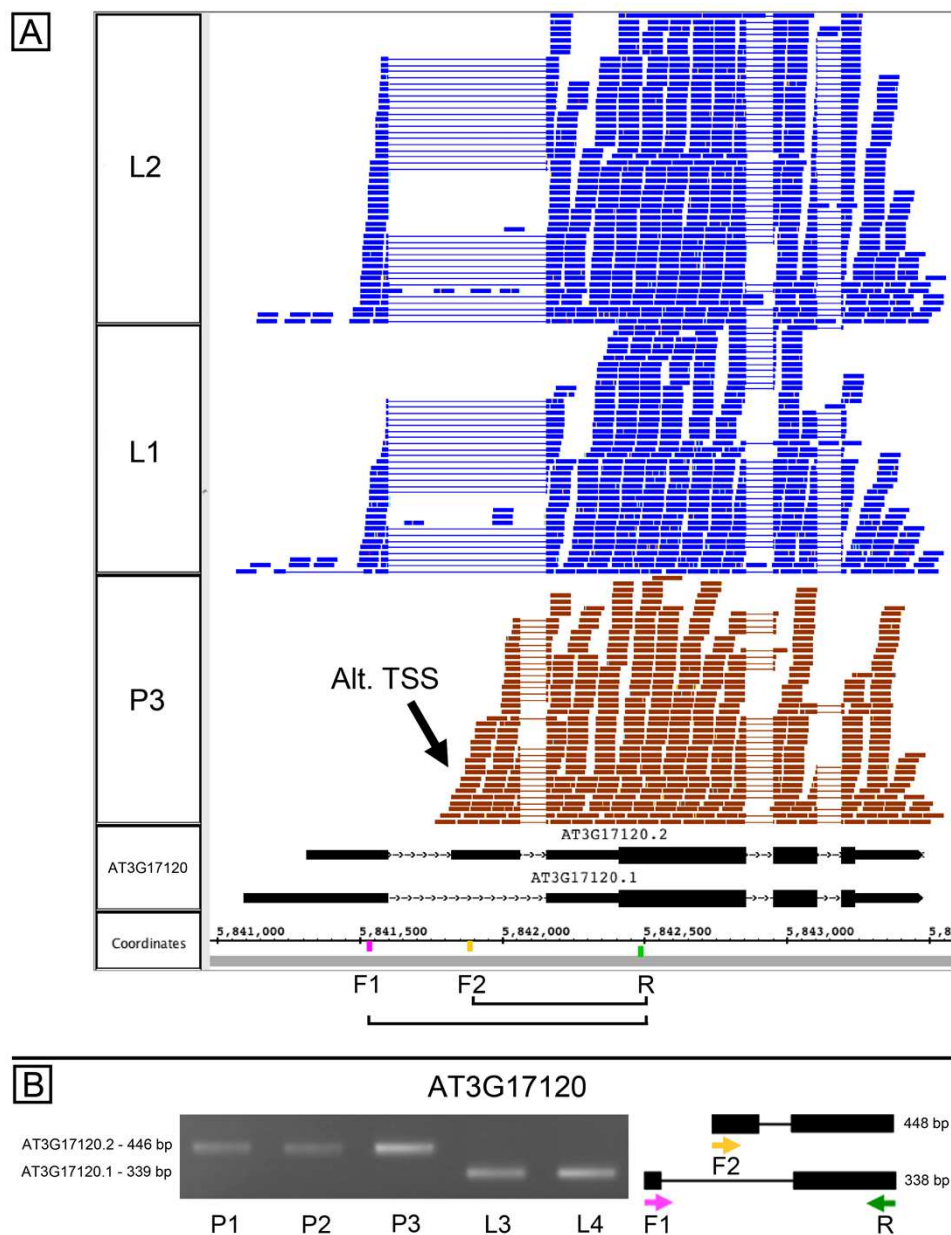


Figure 3. Alternative splicing and putative alternative transcriptional start site in AT3G17120, a gene of unknown function. (A) Visualization of RNA-Seq read alignments from leaf (L2, L1) and pollen (P3). Locations of PCR primers are indicated on the coordinates track. Primer sequences from left (5') to right (3') were AGAATCCGCCATTTTCTCCT (F1), CCTTGCTTCAAGCCCGAGAT (F2), and TTCCCATTTATCTCCCCAGATT (R). (B) Gel electrophoresis of PCR products from amplifying pollen (P1, P2, and P3) and leaf (L1, L2) cDNAs in reactions that included primers F1, F2, and R. Corresponding model of alternative splice variants to right of gel. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.

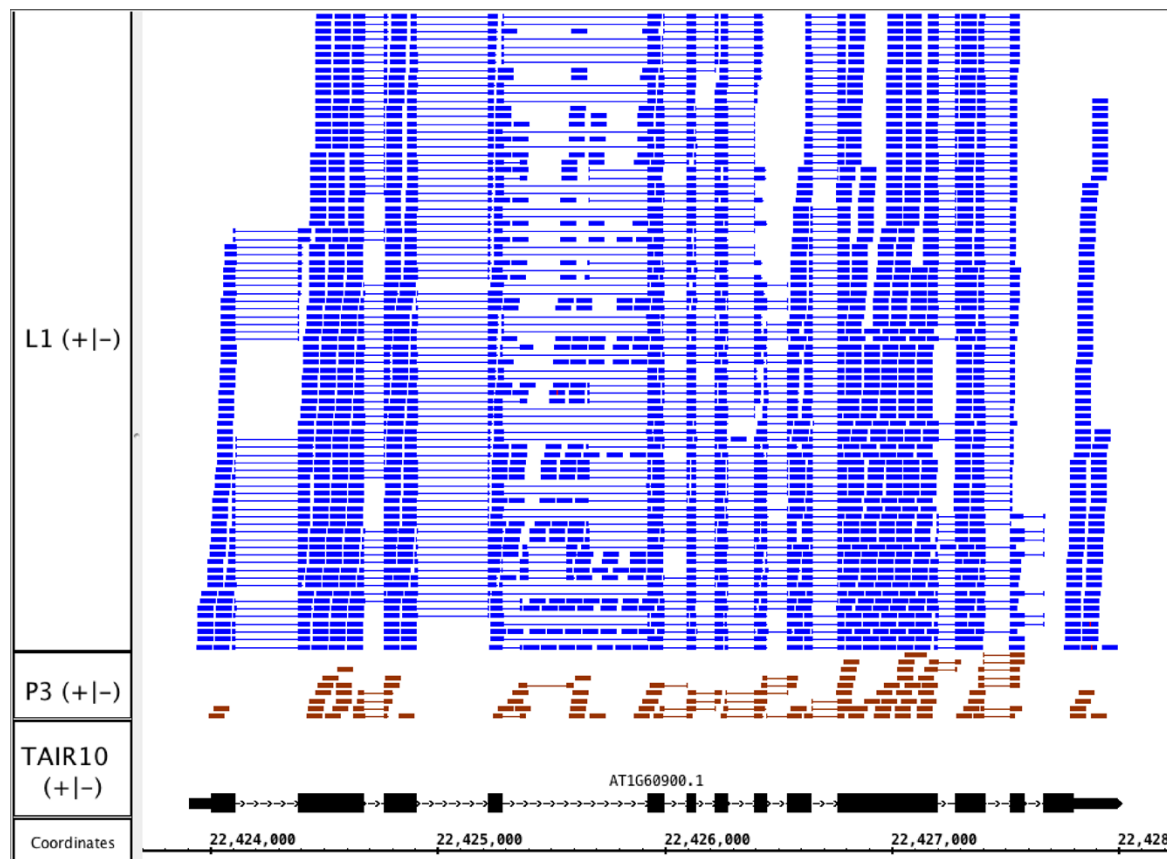
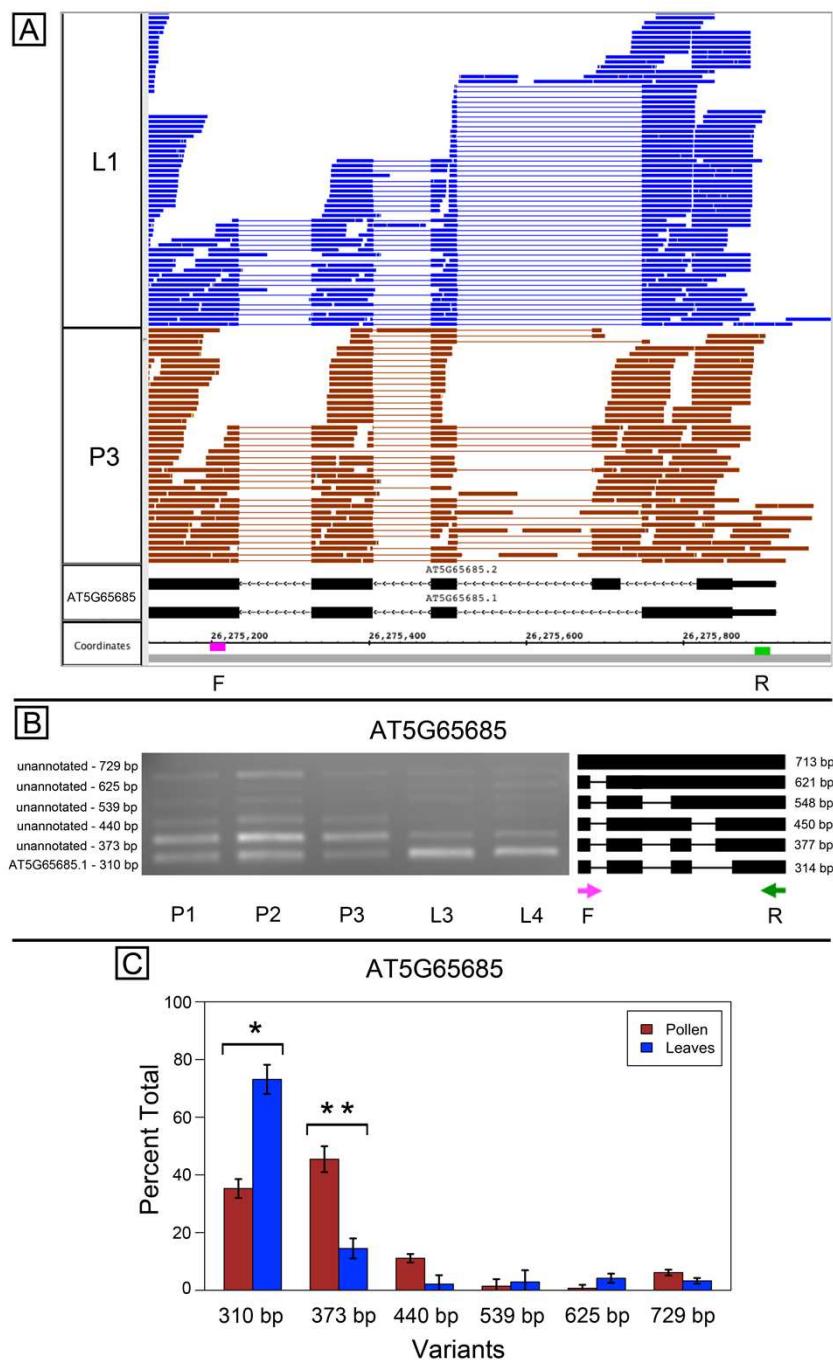


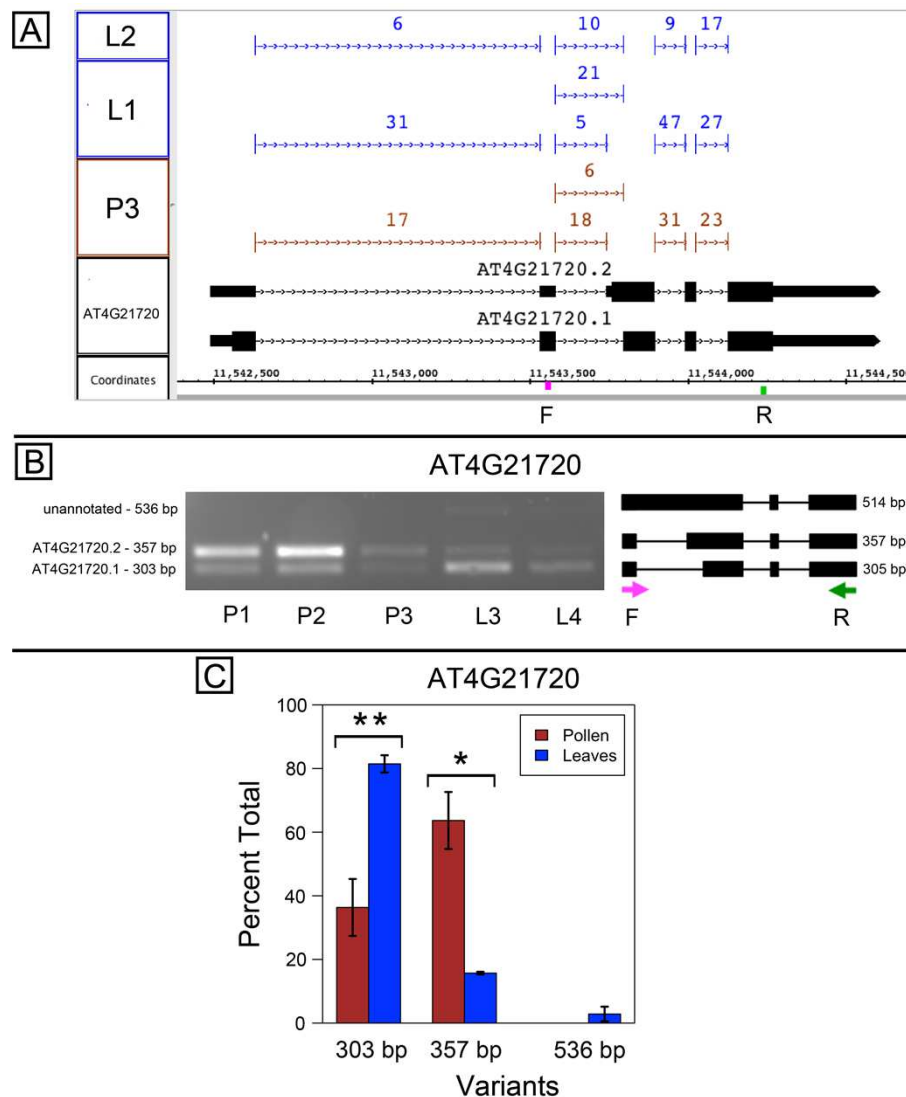
Figure 4. RNA-Seq read alignments indicating that the fourth intron in Arabidopsis U2AF65B is alternatively spliced. Reads from pollen sample P3 and leaf sample L1 are shown.

Table 1. Summary of alternative splicing assay results. All genes except AT2G19480 were confirmed through PCR to have differential splicing patterns in pollen versus leaf sample types. Table indicates whether AS affected predicted protein, and if putative alternative transcriptional start sites (Alt. TSS) were identified. P – pollen, L – leaves, N – no, Y - yes.

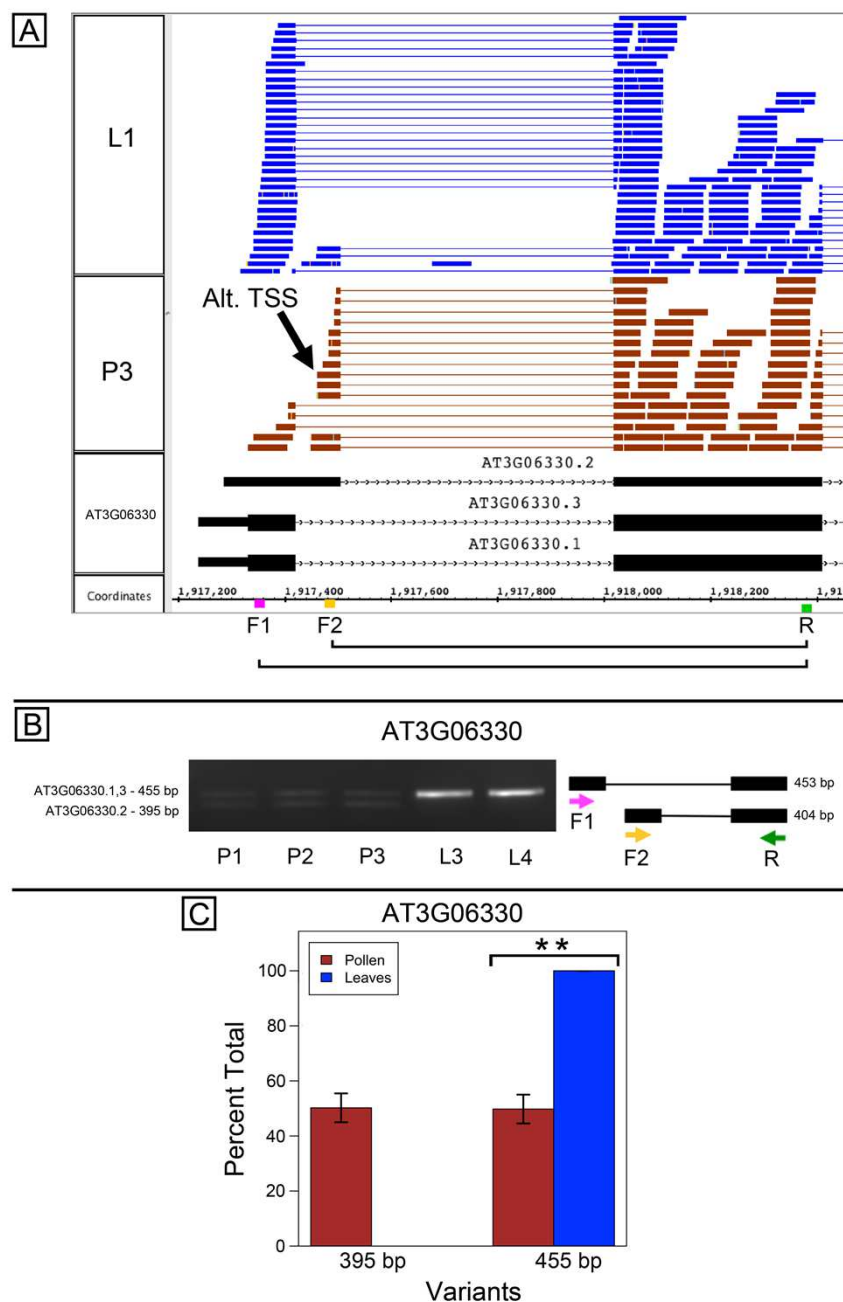
Locus	Annotation	AS affects protein	Putative Alt. TSS
AT1G09140	At-SR30, involved in splicing	Y	N
AT5G52040	At-RS41, involved in splicing	Y	N
AT5G65685	Starch synthase	Y	N
AT4G21720	Unknown	Y	N
AT3G17120	Unknown	N	Y
AT3G06330	Zn-finger, RING-CH-type	N	Y
AT3G58510	RH11, RNA helicase	N	Y
AT1G67480	Galactose oxidase/kelch repeat	N	Y
AT2G19480	Nucleosome assembly protein	Y	N



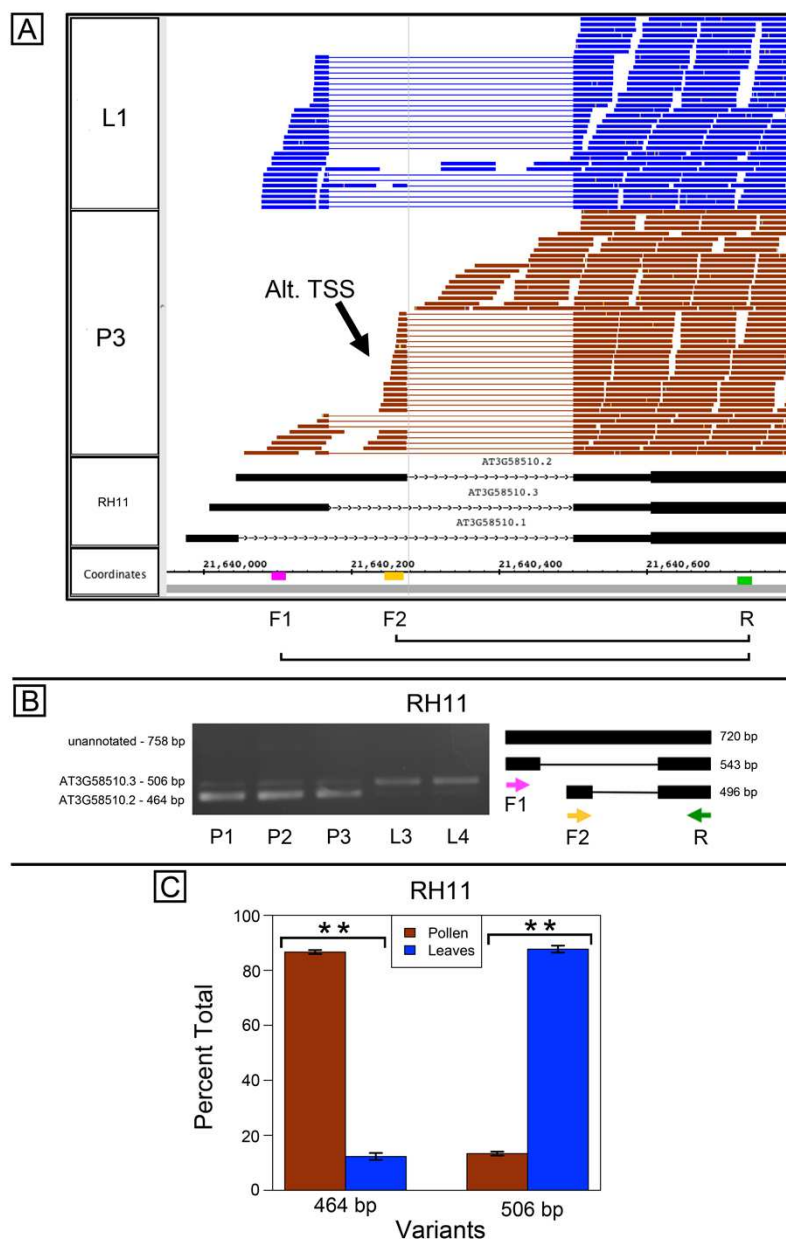
Supplemental Figure 1. Alternative splicing in AT5G65685, a putative starch synthase. (A) RNA-Seq read alignments for leaf (L1) and pollen (P3) samples alongside annotated gene models with PCR primer locations of primers indicated on the coordinates track. Primer sequences were ACTGCAGCTAGGCGTTGTTT (F) and GTCGTTCCCGGTAAATTTTG (R) (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs and corresponding model of alternative splice variants. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.



Supplemental Figure 2. Alternative splicing in AT4G21720, a gene of unknown function. (A) Junction features from RNA-Seq reads for leaf (L2, L1) and pollen (P3) samples alongside annotated gene models. Number of spliced reads supporting each junction indicated above junction. Locations of primers indicated on the coordinates track. Primer sequences were AAGGGATGTGATGCCGATAG (F) and TCCTCAGTAGGAGGCTGCAT (R) (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs and corresponding model of alternative splice variants. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.

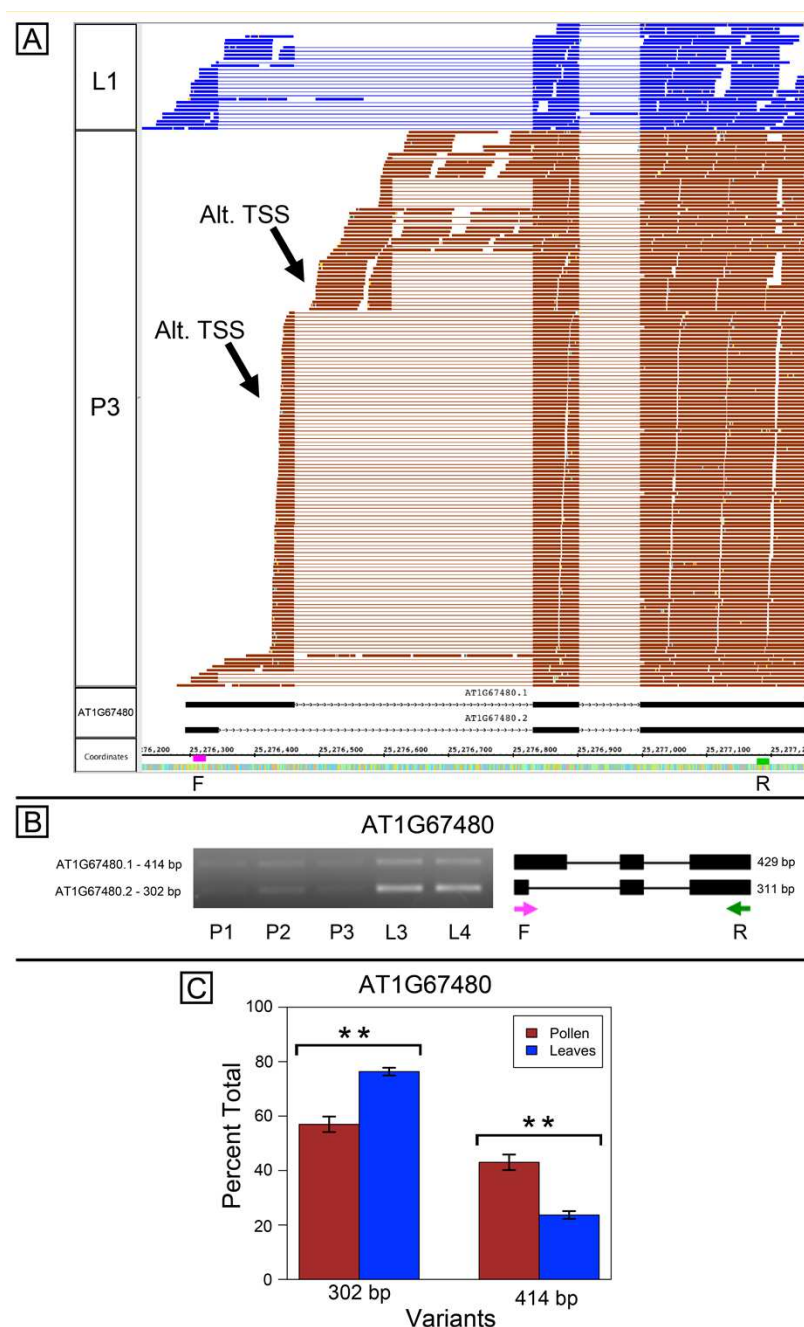


Supplemental Figure 3. RNA-Seq and PCR amplification of AT3G06330 transcript variants. (A) Zoomed-in view of the 5' region of RNA-Seq read alignments from leaf (L1) and pollen (P3) alongside annotated gene models. Location of alternative transcription start site (Alt. TSS) indicated by arrow. PCR primers shown on the coordinates axis. Primer sequences were AGATTCGGCAGCTCAAGAAC (F1), TTTTCCGGGGTTGTGATAGA (F2), and ACCCCAGCTGAAAGTTGTGT (R). (B) Gel electrophoresis of PCR amplification from pollen (P1, P2, P3) and leaf (L3, L4) cDNAs from reactions that included all three primers F1, F2, and R. Corresponding model of alternative splice variants to right of gel. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.

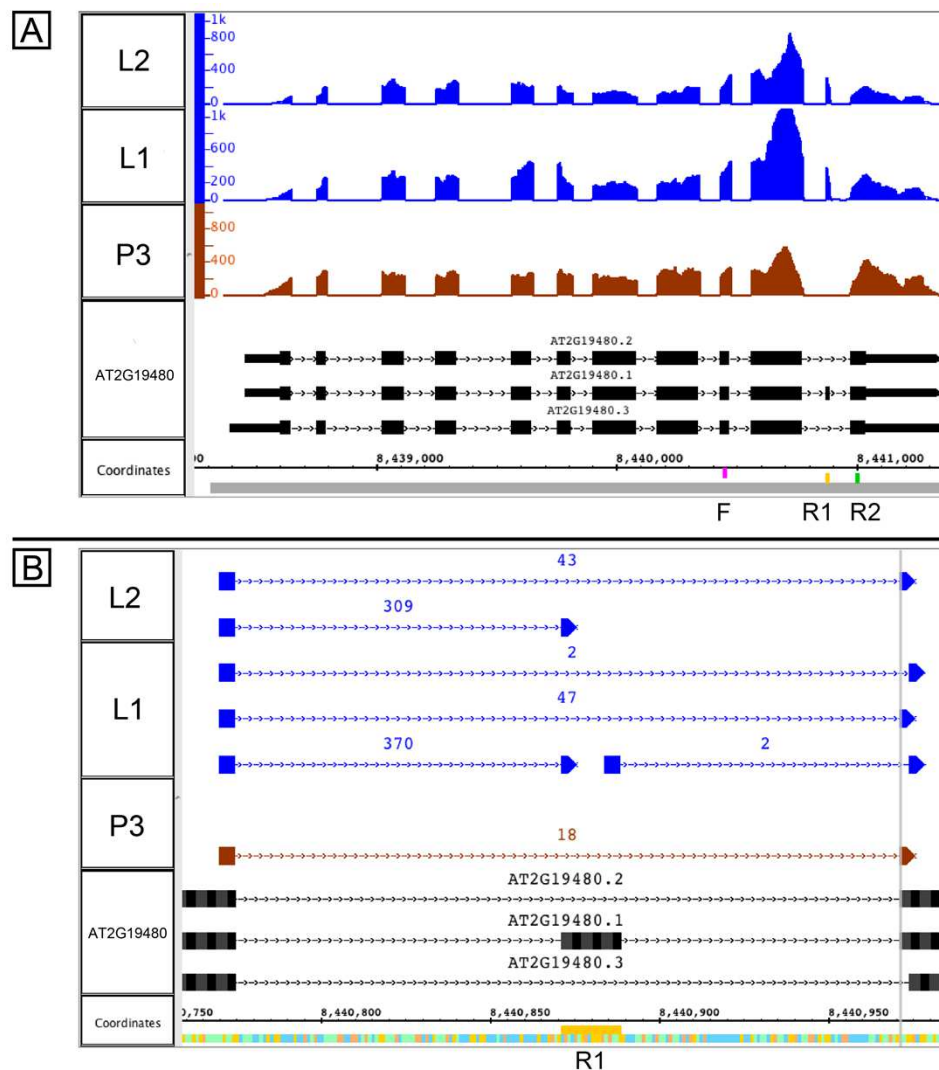


Supplemental Figure 4. Alternative splicing and alternative transcriptional start sites in RH11

(AT3G58510) (A) RNA-Seq read alignments covering the differentially spliced 5' UTR region from leaf (L1) and pollen (P3) alongside annotated gene models. Location of alternative transcription start site (Alt. TSS) indicated by arrow. Locations of primers indicated on the coordinates track. Primer sequences were TCCGTTGGGTGAACGACTAC (F1), AGGGTTCCTTAATTGGTTTATTCGT (F2), and ACCGTTCTAACTCAGCATC (R). (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs from reactions that included all three primers F1, F2, and R. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Unannotated 758 bp fragment was below threshold for quantification. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.



Supplemental Figure 5. Alternative splicing in AT1G67480, a putative galactose oxidase. (A) RNA-Seq read alignments from leaf (L1) and pollen (P3) alongside annotated gene models. Location of two alternative transcription start sites (Alt. TSS) indicated by arrows. Locations of primers indicated on the coordinates track. Primer sequences were ACAGAACCCAAAAACCGACA (F) and ACTTTCGTGAAACCCGTCAC (R) (B) Gel electrophoresis of PCR amplification of pollen (P1, P2, P3) and leaf (L3, L4) cDNAs and corresponding model of alternative splice variants. Estimated fragment size from gel and theoretical fragment size based on splice model found to left and right, respectively, in base pairs (bp). (C) Percent total of each observed splice variant in pollen and leaf samples quantified from gel electrophoresis in B. Values are averages of replicate samples. Error bars indicate two standard deviations. Asterisk indicates p-value less than 0.05, double asterisk less than 0.01.



Supplemental Figure 6. Exon skipping in AT2G19480 (NAP1;4). (A) Coverage graphs for leaf (L2, L1) and pollen (P3) RNA-Seq data showing exon skipping or exon inclusion in the final intron. Values on the y axis are the number of reads per base pair positions indicated in the coordinates track. Locations of PCR primers are indicated on the coordinates track. Primer sequences were CAAGGACAAATGGAGCATGA (F), CTTGTGCCAGCTGATGA (R1), and TCTCACCTGCTTGACCTTCC (R2). (B) A close-up view of the junction tracks for the alternatively spliced exon in the 3' region. Number of spliced reads supporting each junction indicated above junction.