A Collection of Pre-mRNA Splicing Mutants in Arabidopsis thaliana

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ABSTRACT To investigate factors influencing pre-mRNA splicing in plants, we conducted a forward genetic screen using an alternatively-spliced GFP reporter gene in Arabidopsis thaliana. This effort generated a collection of sixteen mutants impaired in various splicing-related proteins, many of which had not been recovered in any prior genetic screen or implicated in splicing in plants. The factors are predicted to act at different steps of the spliceosomal cycle, snRNP biogenesis pathway, transcription, and mRNA transport. We have described eleven of the mutants in recent publications. Here we present the final five mutants, which are defective, respectively, in RNA-BINDING PROTEIN 45D (rbp45d), DIGEORGE SYNDROME CRITICAL REGION 14 (dgcr14), CYCLIN-DEPENDENT KINASE G2 (cdkg2), INTERACTS WITH SPT6 (iws1) and CAP BINDING PROTEIN 80 (cbp80). We provide RNA-sequencing data and analyses of differential gene expression and alternative splicing patterns for the cbp80 mutant and for several previously published mutants, including smfa and new alleles of cwc16a, for which such information was not yet available. Sequencing of small RNAs from the cbp80 mutant highlighted the necessity of wild-type CBP80 for processing of microRNA (miRNA) precursors into mature miRNAs. Redundancy tests of paralogs encoding several of the splicing factors revealed their functional non-equivalence in the GFP reporter gene system. We discuss the cumulative findings and their implications for the regulation of pre-mRNA splicing efficiency and alternative splicing in plants. The mutant collection provides a unique resource for further studies on a coherent set of splicing factors and their roles in gene expression, alternative splicing and plant development.

KEYWORDS Arabidopsis thaliana CBP80 miRNAs mutant screen pre-mRNA splicing

Splicing of pre-mRNAs by the excision of introns and ligation of flanking exons is a prerequisite for the expression of most eukaryotic genes. Splicing entails two transesterification reactions carried out by the spliceosome, a large and dynamic ribonucleoprotein (RNP) machine located in the nucleus. At least six structurally and functionally distinct spliceosomal complexes containing core spliceosomal proteins, transiently-associated factors and different combinations of five different small nuclear (sn) RNAs - U1, U2, U4, U5 and U6 - act sequentially to execute the two catalytic steps of the splicing process (Matera and Wang 2014; Yan et al., 2017). The spliceosome is able to carry out constitutive splicing, in which the same splice sites are always used for a given intron, and alternative splicing, in which splice site usage for a given intron is variable. Alternative splicing increases transcriptome and proteome diversity (Nilsen and Graveley 2010; Syed et al., 2012; Reddy et al., 2013) and is important for development and stress adaptation in plants (Staiger and Brown 2013; Filichkin et al., 2015; Szakonyi and Duque 2018).

Most information on spliceosome composition and the splicing mechanism has been derived from genetic, biochemical and structural studies in yeasts and metazoan cells (Papasaikas and Vacárcel 2016). Structural and mechanistic insights into the splicing process in these organisms have relied heavily on the development of manipulable biochemical systems that perform the splicing reactions in vitro. Similar work has lagged in plants, largely owing to the lack of an efficient in vitro splicing system (although see recent progress in this area; Albaqami and Reddy 2018). Many of the approximately 430 predicted splicing-related proteins encoded in the Arabidopsis thaliana (Arabidopsis) genome have been identified through sequence similarity searches with yeast, Drosophila and human genes (Koncz et al., 2012). Genetic approaches have been useful for
revealing the physiological roles for some of these splicing factors. For example, forward genetic screens designed to investigate requirements for distinct processes, such as hormone responses (Hugouvieux et al., 2001; Zhan et al., 2015) and flowering time (Marquardt et al., 2014), have identified different splicing-related proteins. Reverse genetics has been used to study the consequences of specific splicing factor deficiencies at particular developmental stages (Ali et al., 2007; Szakonyi and Duque 2018) or under different environmental conditions (for example, Zhan et al., 2015; Laloum et al., 2018; Calixto et al., 2018. Cavallari et al. 2018; Huertas et al., 2019). However, to our knowledge, an unbiased forward genetic screen dedicated to identifying factors that influence alternative splicing of a well-defined alternatively spliced gene has not previously been conducted in any plant system.

We developed a novel GFP reporter system in Arabidopsis to carry out such a forward genetic screen. In this system, an intron-containing GFP reporter gene present in a wild-type ‘Target’ (WT T) line exhibits variable levels of GFP expression depending on the splicing pattern of its pre-mRNA. Of three major GFP splice variants present in wild-type plants, only one, which arises from splicing a U2-type intron with comparatively weak, non-canonical AU-AC splice sites, is present in wild-type plants, only one, which arises from splicing a U2-type intron with strong, canonical GU-AG splice sites, and an unspliced pre-mRNA – cannot be translated because they comprise premature termination codons. Our working hypothesis is that mutations in genes encoding splicing factors will alter the relative proportions of the three splice variants, leading to an increase or decrease in the level of translatable GFP mRNA. Theoretically, these changes should result, respectively, in either a Hyper-GFP (HGF) or GFP-weak (GFW) phenotype compared to the WT T line, which displays an intermediate level of GFP fluorescence (Kanno et al., 2016; 2017a, b; 2018a,b).

Using the GFP splicing reporter system, we conducted a classical forward genetic screen to identify mutants exhibiting altered splicing and GFP expression. By screening for changes in GFP fluorescence in seedlings derived from chemically mutagenized seed, we retrieved nine hgf mutants and seven gfw mutants. Eleven of the mutants, which indeed turned out to be defective in various splicing-related factors, have been published previously (Sasaki et al., 2015; Kanno et al., 2016; 2017a,b; 2018a,b). Here we report the identity of the final five mutants together with previously unpublished information on global gene expression and alternative splicing profiles in one new mutant and in several mutants for which this information has not been previously published. Our study provides foundational knowledge for further in depth investigations of the splicing factors retrieved in the screen, and allows an integrated analysis of a set of sixteen splicing-related proteins unified by their involvement in processing the same splicing reporter.

**MATERIALS AND METHODS**

**Plant materials**

All wild-type and mutant plants used in this study are in the ecotype Col-0 background and were cultivated under long-day conditions (22-23ºC, 16 hr light, 8 hr dark).

The T-DNA insertion mutants used in this study were as follows: SAIL_569_G05 (rpb45a; ATSG549000), SAIL_505_E03 (rpb45b; AT1G11650), SALK_063484C (rpb45c; AT4G27000), SALK_009736C, SAIL_527_G04, SALK_132471, SALK_053475 and SALK_152624 (cwc16b; AT1G17130), SAIL_608_B05 (smfb; AT2G14285). All the seeds were obtained from the Nottingham Arabidopsis Stock Center (NASC).

**Forward genetic screen**

The forward genetic screen based on an alternatively-spliced, intron-containing GFP reporter gene in Arabidopsis [T line; referred to herein as ‘wild-type’ (WT)] (Figure 1A) has been described in detail in prior publications (Sasaki et al., 2015; Kanno et al., 2016; 2017a, b; 2018a,b). Briefly, approximately 40,000 Arabidopsis seeds of the WT T line homozygous for the alternatively-spliced GFP reporter gene were treated with ethyl methane sulfonate (EMS) and sown on soil (M1 generation). From approximately 30,000 M1 plants that grew to maturity and produced self-fertilized seeds, 52 batches of M2 seeds (the first generation when a recessive mutation can be homozygous) were harvested. Surface-sterilized M2 seeds were germinated on solid Murashige and Skoog (MS) medium in plastic Petri dishes and examined under a fluorescence stereo microscope at seven days post-germination for GFP fluorescence. M2 seedlings showing enhanced or reduced GFP fluorescence relative to the WT T line were placed into hgf and gfw categories, respectively, and selected for further analysis (Figure 1B). Causal mutations were identified in the mutants by next generation mapping (NGM; James et al. 2013) using pooled DNA isolated from at least 50 BC1F2 progeny displaying the desired GFP phenotype. BC1F2 plants were produced by backcrossing the M2 plants with the WT T line followed by self-fertilization of the resulting BC1 progeny. Screening approximately 210,000 M2 seedlings (representing around seven M2 seedlings for each M1 plant) yielded nine hgf mutations and seven gfw mutations (Table 1). Mutations were confirmed by identification of multiple alleles and/or complementation analyses. All mutations reported here are recessive.

**Complementation**

For complementation of the rpb45d-1, dgerc14-1, cdkg2-3, iws1-2 and chb80-1 mutants, the respective coding sequences (https://www.arabidopsis.org/) were modified to add SalI sites and XbaI at the 5’ and 3’ ends, respectively, and to replace internal SalI or XbaI sites by silent mutations. The modified cDNAs were synthesized by Genscript (www.genscript.com) and cloned into pUC57 as SalI-XbaI fragments.

The binary vector BV-MpPAT, Sall (Matzke et al., 2010, PAT, phosphorothinic acetyltransferase, which confers resistance to DL-phosphinothricin) was altered to contain the 35S promoter from cauliflower mosaic virus (35Spro) (Pietrzyk et al., 1986) and 3C transcriptional terminator region (from the pea rbcS3C gene (Beney et al., 1990) between the PAT selection marker and the left T-DNA border region of the binary vector. Between the 35Spro and 3C terminator region, a chloramphenicol resistance marker was inserted.
Figure 1  Alternatively-spliced GFP reporter gene system. (A) The T-DNA construct introduced into Arabidopsis comprises a GFP reporter gene under the transcriptional control of a minimal promoter (TATA) and upstream viral (EPRV) enhancer. In the WT T line, however, the expected transcription initiation site (TSS) (light gray arrow) is not used. Rather transcription of GFP pre-mRNA initiates at an unanticipated upstream TSS (black bar and arrow). Alternative splicing yields three GFP splice variants: an unspliced transcript, a transcript resulting from splicing from a canonical GU-AG intron, and a transcript arising from splicing from a U2-type intron with non-canonical AU-AC splice sites (black asterisks). Hence only the AU-AC transcript can be translated into GFP protein. The coding sequence of GFP protein (green bars) uniquely contains a 27 amino acid extension (short green bars with black outline) compared to standard GFP (Fu et al., 2015; Kanno et al., 2016). Arrowheads denote a tandem repeat cluster upstream of the cryptic promoter. The black AUG designates the major translation initiation codon. The 3' AC splice site is only three nucleotides downstream of the 3' AG splice site (Kanno et al. 2008; 2016, 2017a, b). (B) Schematic of forward screen and GFP fluorescence phenotypes of newly germinated seedlings. The wild-type Target (T) line has an intermediate level of GFP fluorescence visible primarily in the stem (hypocotyl) and shoot and root apices. Mutants generated by EMS treatment of the T line exhibit either reduced (Weak) or enhanced (Hyper) GFP fluorescence relative to the T line. Cotyledons (first leaves appearing from a germinating seedling) appear red owing to auto-fluorescence of chlorophyll at the excitation wavelength for GFP. Figure adapted from Figure 1 in Kanno et al. (2018a) with permission from the Genetics Society of America.

between the SalI site after the 35Spro and the XbaI site positioned before the 3C terminator. By cutting with SalI and XbaI, the chloramphenicol resistance marker could be replaced by the modified CDS (under transcriptional control of the 35Spro) of genes to be complemented.

To permit immunoprecipitation of IWS1 in future experiments, the IWS1 cDNA (lacking a stop codon) was fused to three copies of a C-terminal FLAG tag. For this, an adaptor consisting of a SpeI fragment encoding three copies of a FLAG peptide sequence with a stop codon followed by XbaI NPTII XbaI SpeI fragment (NPTII, neomycin phosphotransferase conferring resistance to kanamycin) was synthesized by Genscript. This adaptor was inserted in the correct orientation into the XbaI site of the modified binary vector that contains an IWS1 cDNA lacking a stop codon, by selecting for kanamycin resistance in bacteria. The NPTII selection marker was deleted afterward with XbaI, thus reconstituting the 35Spro-IWS1-FLAG CDS.

Binary vectors containing the modified cDNAs were introduced into Agrobacterium tumefaciens strain ASE (Wu et al., 2018) via tri-parental mating (Matzke and Matzke, 1986) and Arabidopsis plants were transformed using the floral dip procedure (Clough and Bent 1998).

**RNA-sequencing and whole genome resequencing**

Total RNA was isolated from approximately 80 mg of two week-old seedlings (BC1F3 generation) of each mutant line tested in this study (EMS-generated mutants: cbp80, smfa-1, cwc16a-1, cwc16a-2, cwc16a-3; SALK T-DNA insertion line: smfb) and wild type T plants using a Plant Total RNA Miniprep kit (GMbiolab, Taiwan) according to the manufacturer’s protocol for the Lysis Solution B, which contains SDS/anti-oxidant. Briefly, the seedlings were ground into a fine powder in liquid nitrogen, then lysed in the Lysis Solution B for 10 min at 60°C, followed by column purification steps. The purified total RNA was extracted using nuclease-free water after the on-column DNase treatment. Construction of libraries and RNA-seq were carried out (biological triplicates for each sample) as described previously (Sasaki et al. 2015; Kanno et al. 2016).

Whole genome re-sequencing of the EMS-generated mutants cbp80, smfa-1, cwc16a-1, cwc16a-2, cwc16a-3 was performed to identify any remaining EMS-induced second-site mutations that change splice sites. Alternative splicing events containing mutations were excluded from further analysis. To prepare the sample for whole genome re-sequencing, genomic DNA was isolated by DCNasy plant mini kit (Qiagen, Taiwan) followed by concentrating by...
### Table 1 Mutants identified in a forward genetic screen based on an alternatively-spliced GFP reporter gene

| Hyper-GFP (hgf) mutant | Name         | AGI number | Predicted function in splicing                                                                 | No. of alleles | Effect of mutation on development | Reference |
|------------------------|--------------|------------|-------------------------------------------------------------------------------------------------|----------------|------------------------------------|-----------|
| hgf1                   | coilin       | At1g13030  | marker protein for Cajal bodies, which facilitate snRNP maturation                              | 12             | negligible                         | Kanno et al., 2016 |
| hgf2                   | CWC16a       | At1g25682  | step I factor                                                                                    | 3              | negligible                         | Kanno et al., 2017a |
| hgf3                   | SMU1         | At1g73720  | recruited prior to B° complex formation;                                                          | 1              | negligible                         | Kanno et al., 2017a |
| hgf4                   | SMFA         | At4g30220  | small nuclear ribonucleoprotein                                                                 | 1              | negligible                         | Kanno et al., 2017a |
| hgf5                   | PRP39A       | At1g04080  | U1 snRNP component                                                                               | 5              | negligible                         | Kanno et al., 2017b |
| hgf6                   | RBP45D       | At5g19350  | U1 snRNP component                                                                               | 2              | negligible                         | this study |
| hgf7                   | DG CR14-related | At3g07790 | spliceosomal C complex                                                                           | 2              | negligible                         | this study |
| hgf8                   | CDKG2        | At1g67580  | splicing-related protein kinase                                                                  | 1              | early flowering                   | this study |
| hgf9                   | IWS1         | At1g32130  | transcription elongation                                                                           | 2              | negligible                         | this study |
| gfw1                   | AtRTF2       | At5g58020  | contributes to ubiquitin-based regulation of the spliceosome?                                      | 2              | embryo lethal                      | Sasaki et al., 2015; Kanno et al., 2017a |
| gfw2                   | PRP8A        | At1g08070  | US snRNP component; acts at catalytic core of spliceosome                                         | 3              | embryo lethal                      | Sasaki et al., 2015; Kanno et al., 2017a |
| gfw3                   | RBM25        | At1g60200  | U1 snRNP component                                                                               | 2              | low seed set                       | Kanno et al., 2017b |
| gfw4                   | PRP18A       | At1g03140  | step II factor                                                                                    | 1              | short roots, small siliques       | Kanno et al., 2018a |
| gfw5                   | PRP4KA       | A3g25840   | recruited prior to B° complex formation; needed for catalytic activation of spliceosome           | 5              | broad rosettes, late flowering, tall stature, low seed set | Kanno et al., 2018b |
| gfw6                   | SAC3A        | At2g39340  | mRNA export factor                                                                                | 5              | negligible                         | Kanno et al., 2018b |
| gfw7                   | CBP80        | At2g13540  | multiple                                                                                        | 1              | Serrated leaves, early flowering  | this study |
| T line (WT)            | n.a.         | n.a.       | Wild-type line expressing GFP reporter gene; used for EMS mutagenesis                             | n.a.           | normal                            |           |

The mutants retrieved in a forward genetic screen based on an alternatively-spliced GFP reporter gene in Arabidopsis (Figure 1) include a predicted core spliceosomal protein (SMFa); putative components of the U1 snRNP (PRP39a, RBM25, RBPs54d) and U5 snRNP (PRP8); putative step I and step II factors transiently associated with the spliceosome (CWC16a and PRP18a, respectively); a predicted complex C protein (DGCR14); putative splicing regulatory proteins (RTF2, SMU1, PRP4ka, CDKG2); one structural protein presumed to be important for snRNP maturation (coilin), putative mRNA export factors (SAC3a, CBP80) and a predicted transcription elongation factor (IWS1). Developmental phenotypes are primarily observed in six (of seven) identified gfh mutations, two of which are embryo-lethal.

Further screening of the M2 population after publication of the first alleles of coilin, PRP39a and CWC16a has identified three new alleles of coilin (RRH; first intron, 3’ splice site; second intron, 5’ splice site), one new prp39a allele (R226) and two new cwc16a alleles (W18; fifth intron, 3’ splice site). These unpublished alleles are counted in the number of alleles shown here.

**Abbreviation:** SRA, Sequence Read Archive (NCBI); ABRC, Arabidopsis Biological Resource Center; T (or ST) refers to the WT T line harboring the alternatively-spliced GFP reporter gene. If the sequencing data from T line has a separate SRA number, it is noted in the table; n.d., not done; n.a., not available.

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**Analysis of RNA-seq data for differential gene expression and differential alternative splicing**

**Differential expression analysis:** To determine differential expression of the cbp80, coilin, cwc16a, and smfa and smfb mutants compared to their respective WT T controls, we estimated the transcript per million (TPM) expression with Salmon (version 0.13.1; Patro et al., 2017) for the Reference Transcript Dataset for *Arabidopsis thaliana* 2 (ArTND2)-Quantification of Alternatively Spliced Isoforms (QUASI) (ArTND2-QUASI) annotation (Zhang et al. 2017). Transcript read counts were grouped per gene using tximport (Soneson et al. 2015) and differentially expressed genes were determined using edgeR with the exactTest (version 3.18.1; Robinson et al. 2010). Genes were considered differentially expressed for a false discovery rate (FDR) < 0.05.

**Read alignment:** Reads were mapped to the index based on the TAIR10 genome release (Lamesch, et al. 2012) and the ArTND2 Transcriptome with STAR (version 2.6.0c; Dobin et al. 2013) using a 2-pass mapping. The following parameters were used: -outSAMprimaryFlag AllBestScore,-outFilterMismatchNmax 2/0 (first/second pass),-outPossMatchNmax 2000, -outFilterIntronMotifs RemoveNoncanonical,-alignIntronMin 0,-alignIntronMax 0,-outSAMstrandField none,
–outSAMtype BAM SortedByCoordinate. During the second pass, the splice junction files of the relevant control and test samples were passed to the mapping via the--sjdbFileChrStartEnd flag.

**Alternative splicing analysis:** Alternative splicing events were obtained and quantified using Whippet (version 0.11; Sterne-Weiler et al. 2018). Two separate splice graph indices were generated; one for the detection and quantification of exon skipping (ES), alternative acceptor (AA) and alternative donor (AD) events, and another for the retained introns (RI) and exitron (EI) events. Exitrons are alternatively spliced internal regions of protein-coding exons (Marquez et al., 2015). Both indices were based on the AtRTD2 transcriptome annotation (Zhang et al., 2017), supplemented with the relevant STAR RNA-seq alignments and were generated with the--bam-both-novel and--bam-min-reads 3 flags. The RI/EI index was further supplemented with ‘pre-mRNA’ coordinates of the genes and the exitron splice junctions detected using an in-house script. The ‘pre-mRNA’ coordinates range from the start to the end of the gene and allow us to quantify the retention levels of all annotated introns in a gene. The Whippet delta step was run with default parameters, except for the--min-samples 3 flag. The alternative acceptor and donor events were filtered, assuring that at least both (alternative) junctions were detected in the Whippet data. The RI events were required to be covered by at least one read for either all control and/or test samples. All events with a probability $\geq 0.9$ and an absolute delta percent-spliced-in (PSI) $\geq 0.1$ were considered significant differential alternative splicing events.

**SNP/indel calling:** SNPs and indels were identified using the Genome Analysis Toolkit (GATK) pipeline (Van der Auwerla et al. 2013). Picard (version 2.10.9, http://broadinstitute.github.io/picard) was used to generate the sequence data for the TAIR10 genome release. Reads were aligned to the TAIR10 genome using BWA-MEM (0.7.16a-r1181; Li 2013), with the added -M flag. The resulting SAM file was converted to BAM format, sorted, and duplicates were marked using Picard tools. The GATK (version 3.8-0-ge9d806836) HaplotypeCaller was used to obtain the raw variants and the Select-Vara function was used to extract the SNPs and indels. SNPs were filtered using the following filter expression: "$QD < 2.0 || FS > 60.0 || MQ < 40.0 || MQRankSum < -12.5 || ReadPosRankSum < -8.0."$ The filter expression for indels was as follows: "$QD < 2.0 || FS > 200.0 || ReadPosRankSum < -20.0."$

SNPs and indels were intersected with the ArT2D2 annotated transcripts and the significantly regulated alternative splicing nodes from Whippet using in-house scripts.

**RT-PCR to detect GFP splice variants:** Total RNA was isolated as described above but without the on-column DNase treatment. Twenty-five microliters of the RNA solution were then treated with two units of RQ1 RNase-Free DNase (Promega, USA) in a total reaction volume of 50 μl according to the manufacturer’s instructions. The DNase treated RNA was purified by NucleoSpin RNA Clean-up kit (Macherey-Nagel, Germany) and eluted with 60 μl of nuclease-free water. Following the manufacturer’s protocol, cDNA was made by Transcriptor First Strand cDNA Synthesis Kit (Roche, USA) from 1 μg of the purified RNA and an oligo d(T) primer as a template and a primer, respectively. RT-PCR was carried out under the following conditions: [94° for 2 min followed by 28 cycles of 94° for 10 s, 58° for 20 s, and 72° for 90 sec, and finally 72° for 7 min] or [94° for 2 min followed by 24 cycles of 94° for 10 s, 58° for 20 s, and 72° for 30 s, and finally 72° for 7 min] for detecting GFP transcripts or Actin transcripts, respectively. Primers are shown in Table S1.

**Detecting GFP protein by Western blotting**

Approximately 100 mg of two-week-old seedlings grown on solid MS medium in plastic Petri dishes were frozen in liquid nitrogen, disrupted into a fine powder by TissueLyser II (Qiagen, USA) and resuspended in 100 μl of extraction buffer A (50 mM HEPES-KOH pH 7.9, 400 mM KCl, 2.5 mM MgCl2, 1 mM EDTA, 1 mM DTT, 0.1% Triton X-100) supplemented with EDTA-free protease inhibitor cocktail (Roche, USA). The suspension was vortexed three times for 15 sec and centrifuged for 10 min at maximum speed at 4°. The supernatants were mixed with equal volumes of extraction buffer A without KCl (sample A). The pellet was resuspended in 200 μl of extraction buffer B (50 mM HEPES-KOH pH 7.9, 200 mM KCl, 2.5 mM MgCl2, 1 mM EDTA, 1 mM DTT, 0.1% Triton X-100) supplemented with EDTA-free protease inhibitor cocktail (Roche, USA) and sonicated three times for eight seconds, 10% duty cycle and 20% power (Bandelin Sonopuls HD 2070 with MS 73 probe), followed by centrifugation for 10 min at maximum speed at 4°. The supernatants were mixed with sample A. Protein concentrations in the samples were measured using the Bradford assay. Five hundred nanograms of protein were separated by ExpressPlus PAGE Gel, 4–12% (Genescript, Taiwan), transferred to Amersham Hybond P 0.2 μm PVDF Membrane (GE Healthcare, USA), followed by Western blotting according to standard procedures. Rabbit anti-tubulin (AS10 680; Agrisera, Sweden), and mouse anti-GFP (CPA9022; Cohesion bioscience, Taiwan) antibodies were used at 1:1,000 dilutions. Secondary antibody, goat anti-rabbit IgG-conjugated with horseradish peroxidase (Agrisera, Sweden) and goat anti-mouse IgG-conjugated with horseradish peroxidase (Biorad, USA), were used, respectively, at a 1:10,000 dilution. The blots were developed using Amersham ECL Select Western Blotting Detection Reagent kit (GE Healthcare, USA).

**PCR to identify gene knock-outs of T-DNA insertion mutants**

To isolate genomic DNA from the T-DNA insertion mutant lines, several two-week-old seedlings were frozen in liquid nitrogen, disrupted into fine powder by TissueLyser II (Qiagen, USA) and resuspended in 250 μl of extraction buffer (100 mM Tris-HCl pH 8.0, 500 mM NaCl, 50 mM EDTA, 10 mM β-mercaptoethanol). After adding 35 μl of 10% SDS, the samples were incubated for 10 min at 65°. After ammonium acetate precipitation and isopropanol precipitation followed by washing and drying steps, the DNA pellet was dissolved in an appropriate volume of TE buffer (10 mM Tris-HCl pH 8.0, 1 mM EDTA). Genotyping was carried out by PCR using specific primers listed in Table S1. PCR conditions were as follows [94° for 2 min followed by 35 cycles of 94° for 10 s, 58° for 20 s, and 72° for 1 min, and finally 72° for 7 min]. When genotyping by using CAPS or dCAPS methods, the PCR products were digested by a suitable restriction enzyme (Table S1).

**Small RNA analysis in the cbp80-1 mutant**

Total RNA was isolated from two-week-old seedlings of the WT T line and cbp80-1 mutants using a PureLink Plant RNA Reagent (Thermo Fisher, USA) and MaxTract high-density gel tubes (Qiagen, USA) following the manufacturer’s instructions. The quality and quantity of isolated RNA were checked with Agilent Bioanalyzer prior to usage. RNA concentrations were assessed by NanoDrop (ND-1000 spectrophotometer). Ten micrograms of total RNA of each mutant line
were used for library preparation and sRNA sequencing (Illumina HiSeq 2500 system) by an in house Genomic Technology Core Facility.

After the quality and adapter trimming, the clean sRNA tags were processed according to a previously published procedure (Wu et al. 2017). In brief, the sRNA tags of 18–26 nt were mapped to the Arabidopsis genome (Release TAIR10) (Lamesch, et al. 2012) with Bowtie (Langmead et al. 2009). Tags that exhibited more than 20 genomic hits or were mapped to chloroplast genome, mitochondria genome, rRNAs, tRNAs, snRNAs or snoRNAs were discarded. To carry out cross-library comparison, the read numbers are used with normalization in transcripts per fifty million (TP50M). The normalized read counts of tag sequences from individual sequencing libraries were calculated by dividing the raw value by the total abundance of adjusted total raw counts of each library, and then multiplied by 50 million.

Northern blotting of sRNA was performed as described previously (Lee et al., 2015) with a few modifications. Briefly, ten micrograms of total RNA were separated on 10% denaturing polyacrylamide TBE-Urea gels (Thermo Fischer, USA) and transferred to Hybond-N+ membranes (GE Healthcare, USA) using Electro Blot Mini System (Major Science, Taiwan). The membrane was then UV cross-linked with 120-mJ energy and baked for 1 h at 80°. DNA oligonucleotides complementary to miRNA (miRBase version 21) (Kozomara et al., 2019) were used as probes to determine the expression of miRNA (sequences of probes in legend of Table S6). The probes were γ-32P end-labeled using T4 polynucleotide kinase (New England Biolabs, USA). Probe hybridizations were performed using Ultrahyb-Oligo buffer (Thermo Fischer, USA) at 37° overnight. After washing with buffer containing 2× SSC and 0.1% SDS, the membranes were exposed on Phosphor-Imager screens, and scanned using the Typhoon Scanner (GE Healthcare, USA). These membranes were also exposed to X-ray film for 1-7 days.

Data availability
Figure S1 contains data on the rbp45d mutants; Figure S2: contains data on the dgrc14 mutants Figure S3 contains data on the cdkg2 mutant; Figure S4 contains data on the iws1 mutant; Figure S5 contains data on the cbp80 mutant; Figure S6 contains a Northern blot analysis of selected miRNAs in the cbp80-1 mutant; Figure S7 shows data on a redundancy test of SMFA and SMFb; Figure S8 contains data on the cwc16a mutants; CWC16; Table S1 contains primer sequences; Table S2 contains comparative phenotypic data on the cbp80-1 mutant, cbp80 complemented plants and wild-type; Table S3 contains an analysis of differential alternative splicing events in the cbp80 mutant; Table S4 contains an analysis of differentially expressed genes and miRNAs in the cbp80 mutant; Table S5 contains an analysis of differential alternative splicing events in the cbp80 mutant; Table S6 contains an analysis of differentially expressed genes in the cwc16a mutants; Table S7 contains an analysis of differential alternative splicing events in the smfa and smfb single mutants; Table S10 contains an analysis of differential alternative splicing events in the smfa and smfb single mutants; Table S11 contains an analysis of differential alternative splicing events in the smfa smfb double mutant; Table S12 contains an analysis of differentially expressed genes in the smfa smfb double mutant.

Seeds of all mutant and wild-type Arabidopsis thaliana lines listed in Table 1 are available at the Arabidopsis Biological Resource center (ABRC, Ohio, USA) and all DNA and RNA sequence data for selected mutants and the wild-type T line are available at NCBI under the respective accession numbers listed as follows. hgf1/coilin/At1g13030/ ABRC stock number: CS69632, CS69639; NCBI accession numbers: R40/hgf1-1 and P439L/hgf1-8; SRR071829 T line: SAMN12817582, P439L/hgf1-8;SAMN12817583, R40/hgf1-1: SAMN12817584, T line, P439L/hgf1-8, and R40*:hgf1-1: SRRP089352 and SRP089656. hgf2/CWC16a/At125682/ ABRC stock number: CS69846, CS72366, CS72367; NCBI accession numbers: this study, cwc16a-1: SRRP09352, T line: SAMN12817589, cwc16a-1: SAMN12817590, cwc16a-2: SAMN12817591, cwc16a-3: SAMN12817592. hgf3/SMU1/At1g3720/ ABRC stock number: N623852; NCBI accession numbers: smu1-1: SRRP09352. hgf4/SMFA/At4g302220/ ABRC stock number: CS69848; NCBI accession numbers: T line: SAMN12817585, smfa: SAMN12817586, smfb: SAMN12817587, smfab: SAMN12817588. hgf5/PRP39A/ At1g04080/ ABRC stock number: CS69936, CS69937, CS69960; NCBI accession numbers: prp39a-3 and prp39a-4: SRP108084 T line: SRP093528. hgf6/RBP45D/At5g19350/ ABRC stock number: CS72358, CS72359; NCBI accession numbers not determined. hgf7/DG CR1- related: At3g07790/ ABRC stock number: CS72360, CS72361; NCBI accession numbers not determined. hgf9/1W51/At1g32130/ ABRC stock number: CS72363, CS72364; NCBI accession numbers not determined. gfw1/AtRTF2/At5g58020/ ABRC stock number: CS69596, N340515; NCBI accession numbers: T line: SR1652313, atrtf2-1: SR1652314, atrtf2-2 heterozygous: SR1652316, atrtf2-2 homozygous: SR1652317. gfw2/PRP8A/At1g08070/ ABRC stock number: CS69597; NCBI accession numbers: prp8-7: SR1652315. gfw3/RBM25/At1g60200/ ABRC stock number: CS69940, CS69941; NCBI accession numbers not determined. gfw4/PRP18A/At1g03140/ ABRC stock number: CS69984; NCBI accession numbers: prp18a-1: SR119240, T line: SRRP093582 and SRP119240. gfw5/PRP4KA/ At3g25840/ ABRC stock number: CS71818; NCBI accession numbers: prp4ka-4: SRP117313, gfw6/SAC3A/At2g39340/ ABRC stock number: CS71820; NCBI accession numbers: sac3a-6: SRP117313, gfw7/ CBP80/At2g13540/ ABRC stock number: CS72565; NCBI accession numbers: T line: SAMN12817580, cbp80: SAMN12817581, T line and cbp80: SRP089656 and SRP089665. The T line (WT)/ ABRC stock number: CS69640 is the control for all the RNA-seq experiments. Supplemental material available at figshare: https://doi.org/10.25387/g3.11369361.

RESULTS
The sixteen factors identified in the forward genetic screen are listed in Table 1. All correspond to putative splicing-related proteins predicted to act at a number of steps of the spliceosomal cycle and sNRP biogenesis pathway (Figure 2 and Figure 3, respectively). More than one allele was retrieved for ten of the mutants, suggesting that the screen is close to saturation. Eleven of the mutants have been reported previously. The published hgf mutants include coilin/hgf1 (Kanno et al., 2016); cwc16a/hgf2, smu1/hgf3, smfa/hgf4 (Kanno et al., 2017a); and prp39a/hgf5 (Kanno et al., 2017b). The gfw category includes rtf2/gfw1 and prp8a/gfw2 (Kanno et al., 2017a); rtm25/gfw3 (Kanno et al., 2017b); prp18a/gfw4 (Kanno et al., 2018b); and prp4ka/ gfw5 and sac3a/gw6 (Kanno et al., 2018b). Here we present the final five mutants: four in the hgf category (rbp45d, dgrc14, cdkg2, and iws1) and one in the gfw group (cbp80).
New hgf mutants

*rbp45d/hgf6 – At1g19350*: NGM analysis of two mutants placed in the hgf6 group revealed mutations in the gene encoding RBP45D (RNA-Binding Protein 45D). Arabidopsis RBP45D, which is 425 amino acids in length, is a nuclear RNA binding protein that contains three RRM (RNA Recognition Motif) domains and preferentially binds to nuclear poly(A) RNA *in vitro* (Lorković et al., 2000). The two *rbp45d* mutations that we recovered result, respectively, in a premature termination codon (W306; *rbp45d-1*) and a frame shift caused by a mutation in the splice site acceptor of the 6th (out of seven) intron (*rbp45d-2*). (Figure S1A). A wild-type copy of the RBP45D gene complemented the hyper-GFP phenotype when introduced into an *rbp45d* mutant (Figure S1B). The two *rbp45d* mutants did not show any obvious growth or developmental abnormalities.

The RBP45D orthologs in budding yeast and humans - Nam8p and TIA-1, respectively - are U1 snRNP components that stimulate splicing at weak splice sites (Gottschalk et al., 1998; Förch et al., 2000, 2002). Although RBP45D has been shown previously to lack this capability, an Arabidopsis protein similar to RBP45D, termed UBP1, appears to enhance splicing at otherwise inefficiently spliced introns (Lorković et al., 2000). In the alternatively-spliced GFP reporter system, somewhat increased levels of both the translatable GFP AU-AC transcript, which may contribute to the hyper-GFP phenotype of the *rbp45d* mutant, and the non-translatable GU-AG transcript were observed (Figure S1C). These results might indicate that wild-type RBP45D can, in some cases, suppress splicing at both weak and strong splice sites. Another possibility is that mutations in RBP45D contribute to a hyper-GFP phenotype through a mechanism other than splicing regulation. Further work is needed to investigate these possibilities.

RBP45D has three paralogs in Arabidopsis: RBP45A (At5g54900), RBP45B (At1g1650) and RBP45C (At1g27000). All four RBP45 genes are ubiquitously expressed, although RBP45D is expressed at a lower level than its three paralogs (http://bar.utoronto.ca/). We tested whether mutations in *RBP45A, RBP45B* and *RBP45C* would affect expression of the GFP reporter gene by crossing the respective T-DNA insertion mutants with the WT T line. The F2 seedlings descending from these crosses were genotyped for homozygosity of the desired T-DNA insertion mutation. Unlike the *rbp45d* mutants, the *rbp45a, rbp45b* or *rbp45c* mutants, which represent T-DNA knockout insertions (Figure S1D), did not display a hyper-GFP phenotype or increased accumulation of GFP protein relative to the WT T line (Figure S1B and S1E, respectively). The *rbp45* paralogs are thus functionally non-equivalent in splicing in the GFP splicing reporter system.

dgcr14/hgf7 – At1g307790: NGM analysis of two mutants placed in the hgf7 group revealed mutations in the gene encoding DGCR14-like (DiGeorge Syndrome Critical Region, also termed ESS2 (Expression Studies)2) in humans (Lindsay et al., 1996) and EsS-2 (ES2-Similar) in *Caenorhabditis elegans* (Noma et al., 2014). DiGeorge Syndrome is a human pleiotropic developmental disorder that is caused by a chromosome 22 deletion, the shortest of which contains the DGCR14 gene. DGCR14-related proteins, which belong to the ESS2 super-family of proteins, are evolutionarily conserved from fission yeast to humans. ESS2 proteins are typically around 500 amino acids in length and contain two predicted coiled-coil domains but no other recognizable functional domains.

There are not yet any published reports on DGCR14-like proteins in plants. In Arabidopsis, DGCR14-like is a single copy, intronless gene that encodes a protein 509 amino acids in length. We recovered two alleles in the screen: *dgcr14-1* (Q80*) and *dgcr14-2* (W365*) (Figure S2A). The corresponding mutants show a hyper-GFP phenotype and increased accumulation of GFP protein, which can be complemented with a wild-type copy of the DGCR14-like gene (Figure S2B and S2C). No obvious growth, morphological or reproductive defects were observed in the *dgcr14* mutants. DGCR-like is ubiquitously expressed in Arabidopsis (http://bar.utoronto.ca/) and predicted to encode a nuclear-localized protein (http://suba.live/).

From biochemical studies in human cells, DGCR14 appears to be most abundant in the splicesomal C complex (Hegele et al., 2012) (Figure 2). This affiliation remains to be confirmed in plants. The mechanism by which DGCR14 acts in splicing is not yet known. In *C. elegans*, ESS-2 has been found to foster accurate mRNA splicing when a splice site contains non-canonical sequences (Noma et al., 2014). The increased level of the translatable AU-AC GFP transcript in *dgcr14* mutants suggests that the wild-type DGCR14 protein can suppress splicing at weak splice sites (Figure S2D), but additional experiments are required to confirm this idea.

cdkg2/hgf8 – At1g67580: NGM analysis of the hgf8 mutant identified a mutation in the gene encoding CDKG2 (Cyclin-Dependent Kinase G2). CDKs constitute an evolutionarily conserved group of serine/threonine kinases that have diverse roles in eukaryotes including cell cycle regulation, transcriptional modulation, pre-mRNA splicing, and translation (Doonan and Kitsios 2009). In humans CDKG1, which together with CDKG10 is most closely related to plant CDKGs, is particularly noted for its role in splicing (Doonan and Kitsios 2009; Cavallari et al., 2018). Arabidopsis CDKG2 has not yet been implicated in mRNA processing; however, its close homolog, CDKG1 (At5g63370), was reported to associate with the splicesome and regulate splicing of a gene involved in pollen wall formation (Huang et al., 2013). CDKG1 was also found to modulate temperature-sensitive alternative splicing of a factor involved in regulation of flowering time in a pathway that involves CDKG2 in the thermosensing mechanism (Cavallari et al., 2018).

CDKG2 is a ubiquitously expressed, nuclear protein that is 752 amino acids in length (http://bar.utoronto.ca; http://suba.live/). Two T-DNA insertion alleles, *cdkg2-1* and *cdkg2-2* (SALK_012428 and SALK_090262) have been published previously (Ma et al., 2015). The allele we isolated in the screen, *cdkg2-3*, results in a D530N substitution that alters a highly conserved amino acid in the highly conserved kinase domain (Figure S3A). The position of the *cdkg2-3* mutation suggests the kinase activity of CDKG2 is important for its function in splicing in the GFP reporter gene system but this remains to be tested in the future. The level of the translatable AU-AC GFP transcript increases substantially in the *cdkg2-3* mutant, which presumably accounts for the hyper-GFP phenotype of the mutant (Figure S3B). By contrast only trace amounts of the non-translatable GU-AG and unspliced transcripts can be detected by RT-PCR (Figure S3C). The *cdkg2-3* mutant has a normal appearance but it is somewhat early flowering (approximately four days before WT plants under our long-day growth conditions), as has been shown previously with the *cdkg2-1* and *cdkg2-2* T-DNA insertion alleles mentioned above (Ma et al., 2015).

*iwsl/hgf9 – At1g32130*: The *hgf9-1* mutation was identified by NGM as an allele of the evolutionarily conserved transcription factor IWS1 (Interacts With Spt6). In yeast, Iws1 has been shown to act in a complex with RNA polymerase II and Spt6 to facilitate pre-mRNA splicing, efficient mRNA export, and transcription elongation.
accompanied by histone H3K36 methylation, which is a mark of transcriptionally active chromatin (Yoh et al., 2008).

The IWS1 protein in Arabidopsis, which is 502 amino acids in length, has a negligibly-expressed paralog, IWS2 (At4g19000) (Li et al., 2010). Two alleles of iws1, iws1-2 and iws1-3, were recovered in the screen. The iws1-2 mutation alters the acceptor site of the 9th intron, which is the penultimate intron, and the mutation in iws1-3 results in a P446L amino acid substitution (Figure S4A). The hyper-GFP phenotype of iws1-2 can be complemented by a wild-type copy of the IWS1 gene (Figure S4B and S4C). The iws1-2 mutant plants are viable and do not display any obvious defects in growth, development or reproduction under normal growth conditions. Mutations in IWS1 were also isolated in independent forward screens in Arabidopsis for factors required for brassinosteroid-induced gene expression (seb1; suppressor of bes1-d) (Li et al., 2010) and for High Nitrogen Insensitive (HNI) plants (Widiez et al., 2011). IWS1 thus appears to participate in multiple physiological processes in plants, which probably reflects its predicted participation in transcription elongation, pre-mRNA splicing, epigenetic modulation, and mRNA export (Yoh et al., 2008, Li et al., 2010; Widiez et al., 2011).

It is unclear how the iws1 mutations we identified confer a hyper-GFP phenotype. Epigenetic modifications, including various histone marks, are known to affect splicing efficiency (Luco et al., 2011; Naftelberg et al., 2015; Godoy Herz and Kornbluht 2019). Conceivably, IWS1-dependent histone modifications in the GFP transcribed region could be altered in the iws1 mutants, thus potentially affecting splicing of GFP pre-mRNA. However, the splicing pattern of GFP pre-mRNA changes only slightly in the iws1-2 and iws1-3 mutants (Figure S4D). An alternative role for IWS1 in modulating transport of GFP mRNA from the nucleus to the cytoplasm is conceivable but remains to be further examined in the context of the hyper-GFP phenotype of the iws1 mutants.

New gfw mutant

cbp80/gfw7 – A12g13540: NMG analysis of the gfw7 mutant identified a mutation in the gene encoding CBP80 (Cap Binding Protein 80). CBP80, together with CBP20, forms the heterodimeric cap binding complex (CBC), which binds the 7-methylguanosine cap at the 5’-end of eukaryotic mRNAs. The CBC participates in multiple processes in the cell, including transcription, splicing, transcript export, and translation (Figure 2 and Figure 3) (Kuhn et al., 2008; Gonatopoulos-Pournazis and Cowling 2014). The CBC was initially isolated biochemically from HeLa cells, where it was shown to be important for splicing (Izaurralde et al., 1994). In Arabidopsis, the ortholog of CBP80 was first identified genetically in a forward screen.
for abscisic acid (ABA) hypersensitive (abh1) mutants (Hugouvieux et al., 2001).

The Arabidopsis CBP80 protein is 848 amino acids in length. The mutation identified in this screen, cbp80-1 (= abh1-9), creates a premature termination codon (W630*) in the middle of a conserved MIF4G (Middle domain of eIF4G), type II domain (Figure S5A). It is not clear which step(s) in GFP pre-mRNA processing and transport is affected in the cbp80-1 mutant. A role in splicing is suggested by an increase in the untranslatable, unspliced GFP transcript and decrease in the translatable GFP transcript resulting from splicing at the AU-AC splice sites in the cbp80-1 mutant (Figure S5B). However, one or more other CBP80-dependent steps important for GFP
Table 2 Numbers of DEGs and DAS events in different splicing-related factors recovered in the forward genetic screen

| Direction of change | cbp80  | coilin * | cwc16a * | smfa  | sfmb  |
|---------------------|--------|----------|----------|-------|-------|
| Up                  | 2105   | 334      | 676      | 3121  | 1093  |
| Down                | 567    | 435      | 265      | 1937  | 1299  |
| Total Number of DEGs| 2672   | 769      | 941      | 5058  | 2392  |
| Down                | 2105   | 334      | 676      | 3121  | 1093  |
| Total Number of DAS genes | 1507  | 255      | 240      | 1465  | 287   |
| Total Number of DAS events | 2064 (100) | 287 (100) | 318 (100) | 2184 (100) | 353 (100) |
| Event               | cbp80  | coilin * | cwc16a * | smfa  | sfmb  |
| AA                  | 168 (8.1) | 129 (44.9) | 45 (14.2) | 292 (13.4) | 69 (19.5) |
| AD                  | 199 (9.6) | 17 (5.9) | 27 (8.5) | 244 (11.2) | 37 (10.5) |
| CE                  | 158 (7.7) | 20 (7.0) | 20 (6.3) | 307 (14.1) | 29 (8.2) |
| EI                  | 83 (4.0) | 13 (4.5) | 26 (8.2) | 158 (7.2) | 37 (10.5) |
| RI                  | 1456 (70.5) | 108 (37.6) | 200 (62.9) | 1183 (54.2) | 181 (51.3) |
| Total Number of DAS events | 2064 (100) | 287 (100) | 318 (100) | 2184 (100) | 353 (100) |
| Total Number of DAS genes | 1507  | 255      | 240      | 1465  | 287   |

Abbreviations: AA, alternative acceptor/alternative 3′ splice site; AD, alternative donor/alternative 5′ splice site; CE, cassette exon; EI, extron; RI, retained intron.

Only the DEGs/DAS events changed in all alleles, with the same direction of change, are listed here. For the full overview of alternative splicing events per allele, see the respective supplementary tables.

expression could also be affected (Figures 2 and 3). Further work in the future is needed to investigate this question.

In addition to its GFP-weak phenotype (Figure S5C), the cbp80-1 mutant has a visible developmental phenotype featuring serrated rosette leaves, decreased rosette diameter, short stature and reduced seed set (Figure S5D, Table S2). These phenotypes, some of which have been noted previously in other cbp80 mutants (Kuhn et al., 2008; Montgomery and Carrington 2008), can be at least partially complemented by introducing a wild-type copy of the CBP80 gene into the cbp80-1 mutant, as can the GFP-weak phenotype (Figure S5C and S5D; Table S2).

To determine the effects of the cbp80-1 mutation on global gene expression and alternative splicing patterns, we carried out triplicate RNA-seq and analyzed the sequencing data for differentially expressed genes (DEGs) and differential alternative splicing (DAS) events in the cbp80-1 mutant. In agreement with earlier results obtained from another cbp80 mutant using an RT-PCR alternative splicing panel (Raczynska et al., 2010), we found that a number of transcripts are differentially spliced in the cbp80-1 mutant (Table 2; Table S3). A notable feature of the DEG list in the cbp80 mutant is the increased accumulation of 53 microRNA (miRNA) precursors (pri-miRNAs) (Table S4, sheet miRNA). The high representation of pri-miRNAs (Table S4, sheet miRNA) was paralleled by a decrease in many of the corresponding mature miRNAs (Figure S6). These results expand on prior findings of decreased AU-AC mRNA splicing pattern in the cbp80 mutant (P16) introduced into the wild-type T line by crossing and F2 seedlings descending from the resulting F1 plants were screened for homozygous smfb progeny in a homozygous TT background. Unlike smfa homozygous seedlings, homozygous smfb seedlings displayed neither a hyper-GFP phenotype nor increased amounts of GFP protein (Figure S7B and S7C). Moreover, in contrast to smfa, which accumulated elevated levels of the translatable AU-AC GFP transcript relative to WT plants, the GFP pre-mRNA splicing pattern in the smfb mutant was virtually unchanged from the WT T line (Figure S7D). We thus conclude that SMFA and SMFB are functionally non-equivalent in the GFP reporter gene system.

A mutation in the gene encoding putative step 2 factor CWC16a (Figure 2) was initially identified in the GFP splicing reporter screen as hgf2/cwc16a-1 (F50) (Table 1) and two new alleles, cwc16a-2 (W18) and cwc16a-3 (frame shift from amino acid 194), are reported here (Figure S8A). Attempts to test mutations in CWC16b, the paralog of CWC16a, were unsuccessful because of the Translatability AU-AC GFP transcript insertion lines of cwc16b available from seed stock centers, three did not contain the T-DNA at the expected location and two did not appear to be cwc16b knockout alleles (Figure S8B). However, the fact that we recovered three alleles of cwc16a and none of cwc16b in our screen suggests a special role for CWC16a in splicing GFP pre-mRNA splicing. Indeed, the levels of the translatable AU-AC GFP transcript and the amount of GFP protein increase substantially in the cwc16a mutants (Figure S8C and S8D, respectively), which is fully consistent with their strong hyper-GFP phenotypes (Figure S8D). The lack of a convincing cwc16b knockout mutation prevented testing the splicing pattern of GFP pre-mRNA in a cwc16b mutant and the viability of a double cwc16a cwc16b mutant.

Redundancy tests

We previously carried out tests for functional redundancy of paralogs of PRP39A, PRP18A and PRP4KA, which were all identified in the GFP splicing reporter screen (Table 1). Similar to results obtained for RBP45D paralogs (see above), none of the paralogs tested previously (PRP39B, PRP18B and PRP4KB) were functionally equivalent to the A forms of the respective genes (Kanno et al., 2017b, 2018a, b). In the present study, we tested SMFB for functional redundancy with its paralog, SMFA, which was identified in the screen as the hgf4-1 mutant (P16) (Table 1; Figure S7A and S7B). A knockout T-DNA insertion allele of smfb (Figure S7C) was introduced into the wild-type T line by crossing and F2 seedlings descending from the resulting F1 plants were screened for homozygous smfb progeny in a homozygous TT background. Unlike smfa homozygous seedlings, homozygous smfb seedlings displayed neither a hyper-GFP phenotype nor increased amounts of GFP protein (Figure S7B and S7C). Moreover, in contrast to smfa, which accumulated elevated levels of the translatable AU-AC GFP transcript relative to WT plants, the GFP pre-mRNA splicing pattern in the smfb mutant was virtually unchanged from the WT T line (Figure S7D). We thus conclude that SMFA and SMFB are functionally non-equivalent in the GFP reporter gene system.

Additional RNA-seq analyses from previously published mutants

We previously published RNA-seq data and analyses of DEGs and DAS events in two coilin mutants ([hgf1-1 (R40), hgf1-8 (P439L)]
and one allele of cwc16a (cwc16a-1). However, these earlier analyses lacked either an optimal number of biological replicates (coilin alleles; Kanno et al., 2016) or were performed on only a single allele (cwc16a-1; Kanno et al., 2017a). Here we expand on these prior results by reporting new RNA-seq data (in triplicate) and DEG and DAS analyses from the two coilin alleles mentioned above and from two newly identified alleles of cwc16a: cwc16a-2 and cwc16a-3 (Figure S8A-D). The more comprehensive results obtained by using multiple alleles of each mutant can be compared and lists of DEGs and DAS events shared by all alleles compiled. These comparisons hone the findings and provide a more accurate assessment of genes that robustly change in expression and alternative splicing profiles in a given mutant. The total number of shared DAS and DEG events among multiple alleles is shown in Table 2, with details, respectively, in Tables S5 and S6 (coilin/hgf1 alleles) and Tables S7 and S8 (cwc16a alleles). A notable feature of the coilin DEG data are the highly significant representation of genes involved in metabolism of secondary compounds and responses to various stresses and environmental stimuli (Table S6, GO_share_UP_P). A previous albeit more limited transcriptome analysis of coilin mutants similarly suggested a prominent role for coilin in stress responses (Kanno et al., 2016).

We also present here new triplicated RNA-seq data and DEG/ DAS analyses for smfa-1, smfa-1, and smfa-1 double mutants. This information was not reported in a previous publication on smfa-1 mutants (Kanno et al., 2017a). An analysis of the RNA-seq data obtained with smfa-1 and smfb-1 single mutants revealed about twice as many DEGs (FDR < 0.05) in smfa-1 compared to smfb-1 (5058 and 2392, respectively, with 827 shared between the two paralogs) (Table 2; Table S9). More than 100 DEGs in smfa-1, compared to only seven in smfb-1, correspond to known or predicted splicing factors in Arabidopsis (numbering around 430; Koncz et al., 2012) (Table S9_a_Koncz share and b_Koncz share). These results suggest that wild-type SMFA-1 is involved more frequently than SMFB-1 in modulating the expression of splicing-related factors. When considering DAS events, the difference between the two paralogs is even more pronounced (approximately sixfold). We identified 2184 cases of DAS for smfa-1 vs. 353 for smfb-1, with 98 instances of DAS shared between the two paralogs (Table 2; Table S10). The limited overlaps in DEGs and DAS events in the smfa-1 and smfb-1 mutants suggest that SMFA-1 is the predominant paralog acting in both gene expression and in splicing in Arabidopsis. These findings extend to the genome-wide level the functional non-equivalence that was observed with the two smf paralogs in the GFP splicing reporter system. Double smfa-1 smfb-1 mutants are viable, and information about DAS and DEG events in the double mutant, which are essentially a sum of the individual smfa-1 and smfb-1 results, is shown in Tables S11 and S12, respectively.

**DISCUSSION**

In a forward genetic screen designed to identify factors involved in alternative splicing of a GFP reporter gene in Arabidopsis, we recovered sixteen splicing-related proteins that are predicted to have a variety of roles in the spliceosomal cycle and snRNP biogenesis pathway (Table 1). Based on the functions of their orthologs in other organisms, some of the factors we identified are likely to be components of a particular snRNP (U1: PRP39A, RBM25, RBP45D; U5: PRP8A). Others are predicted to be associated with a specific spliceosomal complex (complex C: DGC14), or to be required at a specific catalytic step of splicing (step 1: CWC16A; step 2: PRP18A). Several factors are potentially splicing regulators involved in catalyzing or targeting various post-translational modifications, including protein phosphorylation (PRP4KA, CDKG2) and ubiquitination (RTF2, SMU1). We also identified factors presumed to be important for snRNP biogenesis (SMFA, coilin); mRNA transport (SAC3A); and transcription elongation/histone methylation (IWS1). One factor, CBP80, is potentially involved in multiple steps including splicing in the nucleus, mRNA export, and snRNA maturation in the cytoplasm. The data we gathered have verified a role for a number of previously uncharacterized proteins in pre-mRNA splicing in plants and revealed novel morphological and developmental phenotypes conferred by specific mutations. Our work has also generated new information on genome-wide gene expression and alternative splicing profiles of endogenous genes in the respective mutants. The overall findings provide foundational knowledge that can underpin more in-depth investigations of the splicing-related proteins in the future.

In addition to its foundational aspects, our study is strengthened and expanded in scope by merging and comparing data from the complete collection of mutants. A combined analysis permits a broader understanding of alternative splicing that cannot be gleaned from the examination of single mutants alone. For example, an overview of the current data set already illustrates the extraordinary complexity of splicing regulation. Even though different mutations often have the same effect on GFP pre-mRNA splicing and GFP expression, there is little overlap in the population of endogenous genes exhibiting splicing defects in the respective mutants (Kanno et al., 2017a; Kanno et al., 2018b). Substantial overlap would be expected if the mutations were revealing distinct, common features of introns affected in both mutants. Therefore, it has not been possible to discern fixed rules that govern splicing from the extant data. The findings rather indicate that each intron represents a unique context for splicing to occur, such that the effects of specific mutations on the splicing outcome are largely unpredictable (Pleis et al., 2007). The application of various post-translational modifications to splicing factors, which is suggested by our identification of putative protein kinases and ubiquitination targeting factors in the screen, adds a further significant dimension to the regulation of alternative splicing that is only beginning to be understood (de la Fuente van Bentem et al., 2006).

Another interesting point arising from the cumulative findings is that obvious developmental phenotypes are observed primarily with GFP-weak mutants (Table 1). Although the biological significance of these findings is still uncertain, they may reflect different developmental roles for proteins required for the splicing reactions to take place at all vs. proteins that influence splice site selection but do not affect the occurrence of splicing per se. The latter category may be more important in plants for modulating splicing patterns in response to stress conditions or other transitory signals that induce an adaptive response.

Unlike their counterparts in yeasts and metazoans, many genes encoding splicing factors in Arabidopsis and other higher plants are duplicated but the extent of functional redundancy or functional divergence has not been clear in most cases (Kalyna and Barta 2004; Koncz et al., 2012). In our mutant collection, we found a number of cases in which paralogs have a non-redundant function in GFP pre-mRNA splicing (Kanno et al., 2017b, 2018a; b: this study). Paralogs of genetically-identified factors that we tested directly and found to be non-redundant in the GFP reporter gene system include PRP39B, PRP4KB, PRP18B, SMFB, and RBP45A, B and C. Although a direct examination of CWC16B was not possible owing to the lack of a suitable knockout T-DNA insertion allele, we presume that CWC16A is the main form involved in splicing GFP pre-mRNA because our screen identified three alleles of cwc16a and none of cwc16b. Similarly,
SAC3A and PRP8A were the only paralogs of the respective genes retrieved in the screen, suggesting a specific role for the A forms in the GFP splicing reporter system. (Kanno et al., 2017b, 2018a, b: this study). Although null mutations in these factors may be lethal, weak mutations that affect GFP splicing and expression could conceivably have been identified in the screen. For example, weak alleles of rtf2 and prp8 were recovered in the screen even though null mutations in these factors are embryo-lethal (Sasaki et al., 2015; Kanno et al., 2017a). Analysis of the RNA-seq data for both A and B forms of SMF revealed only modest overlaps in DEGs and DAS events between the paralogs, extending the functional non-equivalence of SMFA and SMFB that was observed with the GFP reporter gene to the genome-wide level. Extensive functional divergence of paralogous plant genes encoding splicing factors increases the complexity and capabilities of the splicing machinery and is likely to be another feature of splicing in plants that contributes to their developmental plasticity and ability to adjust to a constantly changing environment.

Another noteworthy aspect emerging from the combined analysis is that mutations were retrieved in splicing factors predicted to act throughout the splicesomal cycle and snRNP maturation pathways and not just at a single crucial step of GFP pre-mRNA splicing. The only possible bias concerns predicted constituents of the U1 snRNP, which is involved in recognizing the 5’ splice site early in the splicesomal cycle. Of these, we identified three putative U1 snRNA components in the screen: RBP45D, PRP39A and RBM25 (Figure 2). The otherwise widespread distribution of the identified mutations as well as the participation of only single members of paralogous gene pairs at different steps of the splicing process hint that our system is illuminating a specialized pathway comprising a set of dedicated components.

The features of the alternatively-spliced GFP reporter gene that permit the genetic identification of a coherent set of splicing factors that cooperate in splicing of GFP pre-mRNA are not known. Potentially, however, they could be a subject for future investigation using CRISPR-Cas9-mediated genome editing to alter nucleotides within the spliced regions of the GFP gene. To our knowledge, many of the splicing-related proteins we identified have not been picked up in any other forward screen focusing on any type of process in plants, which again points toward one or more distinctive but as yet unidentified features of our GFP splicing reporter that render it particularly sensitive to mutations in the genetically identified factors. Based on the strong preference of their cognate mutants for the weaker AU-AC splice sites in GFP pre-mRNA, some HGF factors, such as CWC16A, SMU1, DGCR14 and CDKG2, might be important for discriminating between strong and sub-optimal/weak splice sites in the GFP pre-mRNA. Whether this observation extends to endogenous genes remains to be determined by scrutinizing in more detail the genome-wide RNA-seq data to assess both canonical and non-canonical splice site usage.

In summary, we used a unique alternative splicing system with both strong (GU-AG) and weak (AU-AC) splice sites, and an easy readout (GFP expression) to identify a coherent set of splicing factors that act at different stages of GFP pre-mRNA splicing and expression in Arabidopsis. The GFP pre-mRNA system is versatile and, similarly to other plant splicing reporters (Simpson et al., 2012, 2014), can be used for other purposes, for example, to test mutations in other splicing factors or under different growth/environmental conditions. The resources generated in this study, including seeds of all mutants and high-throughput sequencing data, have been deposited in public repositories (Table 1) and hence are available to the international plant science community. We anticipate that additional global insights into the regulation of alternative splicing in plants will emerge once an entire data set containing detailed phenotypic and RNA-seq data from all mutants is available.

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