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Bread Wheat TaSPO11-1 exhibits evolutionary conserved function in meiotic recombination across distant plant species.

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The manipulation of meiotic recombination in crops is essential to rapidly develop new plant varieties, producing more in a sustainable manner. One option is to control the formation and repair of the meiosis-specific DNA double strand breaks (DSBs) that initiate recombination between the homologous chromosomes and ultimately lead to crossovers. These DSBs are introduced by the evolutionarily conserved topoisomerase-like protein SPO11 and associated proteins. Here, we characterized the homoeologous copies of the SPO11-1 protein in hexaploid bread wheat (*Triticum aestivum* L.). The genome contains three *SPO11-1* gene copies that exhibit 93-95% identity at the nucleotide level and clearly, the A and D copies originated from the diploid ancestors *Triticum urartu* and *Aegilops tauschii*, respectively. Further, phylogenetic analysis of 105 plant genomes revealed a clear partitioning between monocots and dicots, with the seven main motifs being almost fully conserved even between clades. The functional similarity of the proteins among monocots was confirmed through complementation analysis of the rice *spo11-1* mutant by the wheat *TaSPO11-1-5D* coding sequence. Also, remarkably, albeit the wheat and Arabidopsis SPO11-1 proteins share only 55% identity and the partner proteins also differ, the *TaSPO11-1-5D* cDNA significantly restored the fertility of the Arabidopsis *spo11-1* mutant indicating a robust functional conservation of the SPO11-1 protein activity, across distant plants. These successful heterologous complementation assays, using the Arabidopsis and rice hosts, are good surrogates to validate the functionality of candidate genes and cDNA, as well as variant constructs, when the transformation and mutant production in wheat is much longer and tedious.

We analysed the three homoeologous copies of *SPO11-1* genes in the bread wheat genome. They are very similar but not identical, revealing their parental origin from diploid wheats. These SPO11-1 proteins contain the seven essential and evolutionary conserved motifs, now found in 105 plant species. The coding sequence of the wheat D-copy significantly complemented the rice and...
Arabidopsis spo11-1 mutants indicating a strong functional conservation between highly divergent species.

**Keywords**

SPO11-1; Meiotic recombination; Protein evolution; Wheat; Grasses.
Introduction

The improvement of genetic diversity in cultivated species is of utmost importance especially towards the development of a sustainable agriculture using for example lower amounts of water, fertilizers, pesticides or fungicides. One way to reach this challenge is to explore the tremendous gene resources that already exist in nature. This can be achieved by crossing the elite lines with related and/or exotic varieties bearing advantageous alleles of agronomical interest. Such genetic admixture occurs naturally through meiotic recombination and gamete formation. The mechanisms of meiotic recombination leading to crossover products (COs) have long been studied in model organisms, including plants (reviewed by (Mercier et al. 2015)). Importantly, crossover recombination not only generates new combinations of parental alleles, but also ensures the faithful distribution of the homologous chromosomes during the reductive meiotic division (MI), key to the success of sexual reproduction and fertility (Hunter 2015, Mercier, et al. 2015).

Meiotic recombination is initiated by the formation of programmed DNA double-strand breaks (DSBs) catalysed by the topoisomerase-like protein SPO11 (Bergerat et al. 1997, Keeney et al. 1997); for reviews, see (de Massy 2013, Lam and Keeney 2014, Robert et al. 2016b)). Resembling the archaeal topoisomerase VI complex, SPO11 (ortholog of the archaea A subunit) is likely working in a protein complex with one or several other proteins (Lam and Keeney 2014), in particular with the MTOPVIB protein (orthologous to the archaea B subunit) to form an active catalytic complex (Figure 1) (Robert et al. 2016a, Vrielynck et al. 2016).

SPO11 is evolutionarily conserved among sexually reproducing organisms and is encoded by a single gene in most organisms (Malik et al. 2007). However, there are several SPO11 genes in plant genomes. The Arabidopsis genome carries three related genes named SPO11-1, SPO11-2 and SPO11-3 (Hartung and Puchta 2000, Grelon et al. 2001, Hartung and Puchta 2001, Sprink and Hartung 2014). Both SPO11-1 and SPO11-2 are essential for meiotic recombination (Grelon, et al. 2001, Hartung and Puchta 2001, Stacey et al. 2006, Hartung et al. 2007) while SPO11-3 is involved in somatic endoreduplication (Hartung et al. 2002, Sugimoto-Shirasu et al. 2002, Yin et al. 2002). The situation in rice (Oryza sativa) is even more complex with five SPO11-related genes (Jain et al. 2006, Yu et al. 2010,
An et al. 2011). Of these rice genes, to date only OsSPO11-1 and OsSPO11-2 have been clearly
demonstrated to be required for meiotic recombination (Yu, et al. 2010, Fayos et al. 2019). The
interactions between SPO11-1, SPO11-2 and MTOPVIB proteins have recently been described in
Arabidopsis, rice and mice (Figure 1) (Fu et al. 2016, Robert, et al. 2016a, Vrielynck, et al. 2016, Xue
et al. 2016).

Another layer of complexity in meiosis in plants is polyploidy, a common feature of plant kingdom,
which concerns about 30% of existing flowering plants (Wood et al. 2009), including most of the
world’s important crops. Polyploid species can be divided into autopolyploid species derived from
whole genome duplication, such as potato (Solanum tuberosum, 4x), strawberry (Fragaria ananassa,
8x), and allopolyploid species originating from crosses between closely related species such as oilseed
rape (Brassica napus, 2n = 4x = 38, AACC), cotton (Gossypium arboreum, 2n = 4x = 52, AADD) or
tobacco (Nicotiana tabacum, 2n = 4x = 48). Allopolyploid species thus contain different sets of related
but not completely homologous chromosomes called homoeologues. It is now clearly established that
all flowering plants have experienced at least one and usually several rounds of polyploidy (also called
whole genome duplications; WGD) during the course of their evolution (Soltis et al. 2009, Van de Peer et al. 2009). Thus, other angiosperms, including Arabidopsis, rice or Brassica crops, are ancient
polyploids (paleo-polyploids) that returned to a functionally diploid state by a massive elimination of
some but not all duplicated genes post-WGD (a process called fractionation; (Doyle et al. 2008)).

Given the prevalence of WGD in plants, it is critical to integrate and extend our knowledge of
important biological processes such as meiotic recombination into the field of polyploidy. From this
point of view, (Lloyd et al. 2014) showed that meiotic genes return to a single copy more rapidly than
genome-wide average in Angiosperms. Analysis of the presence and expression of meiotic genes in
two recent polyploid species (oilseed rape and bread wheat; ~10,000 years ago) suggested that their
loss is passive and is a long-term process.

Among allopolyploid species, hexaploid bread wheat (Triticum aestivum L.; 2n = 6x = 42) is derived
from two successive interspecific crosses (Blake et al. 1999, Huang et al. 2002). These involved the
three diploid species T. monococcum ssp urartu (AA), a yet unknown species related to the Sitopsis
section (BB; the closest one being Aegilops speltoides; S genome) and Aegilops tauschii (DD). The
recent release of the anchored and annotated wheat whole-genome sequence (IWGSC 2018) thus revealed that more than 60% of the 39,474 genes are present in triplicates and more than 90% have at least two homoeologous copies.

The large size (17 Gb) and complexity of the hexaploid wheat genome makes it challenging for functional analyses of gene, in particular for gene involved in complex biological complex such as meiosis. Until now, only a dozen genes (TaRAD51, TaRAD51C, TaRAD51D, TaDMC1, TaMRE11, TaRAD50, TaASY1, TaZYP1, TaPHS1, TaPH1, TaREC8, TaRECQ-7), among more than 100 known as involved in meiotic recombination in plants, have been cloned and significantly analysed in wheat. These analyses have mainly been limited to comparisons of sequences of homoeologous copies, expression analyses and immunolocalization (Boden et al. 2007, de Bustos et al. 2007, Boden et al. 2009, Devisetty et al. 2010, Perez et al. 2011, Khoo et al. 2012a, Khoo et al. 2012b, Ma et al. 2018, Gardiner et al. 2019). TaPh1, which controls homoeologous recombination in wheat, remains the best-defined locus involved in meiosis in wheat (Griffiths et al. 2006, Moore 2014, Martin et al. 2017).

To date, there is no study analysing SPO11-1 from a polyploid species. SPO11-1 is a key gene at the onset of the recombination process. It is therefore essential to understand how it behaves in a polyploid context such as that of bread wheat. In this work, we wanted to know to what extent wheat SPO11-1 homoeologous copies (TaSPO11-1-5A, TaSPO11-1-5B and TaSPO11-1-5D) are conserved between each other and with those of other plants. We thus sought to identify the hexaploid wheat SPO11-1 gene and to assess its functionality during meiosis. We isolated the homoeologous (A, B and D) copies of TaSPO11-1 and performed a phylogenetic analysis with SPO11-1 from different species. The three copies were compared in detail with those of Arabidopsis and Rice to estimate their conservation. We extend these comparisons in demonstrating the functionality and evolutionary conservation of wheat TaSPO11-1 through heterologous complementation of the corresponding Arabidopsis and Rice mutant lines, i.e. across distant plant species.
Results

*SPO11-1 is conserved between the three homoeologous genomes of the hexaploid wheat.*

To identify the orthologous copies of SPO11-1 within the bread wheat genome, we used the newly released wheat genome sequence (IWGSC 2018) and the Arabidopsis SPO11-1 protein sequence as a query (At3g13170). Three copies were identified, one on each of the three homoeologous genomes from group 5: TraesCS5A02G391400, TraesCS5B02G396300 and TraesCS5D02G401100 further named TaSPO11-1.5A, TaSPO11-1.5B and TaSPO11-1.5D, respectively. The length of the loci was similar with approximately 3.9 kb for the genomic sequences and 1.2 kb for the coding sequences (CDS) (Table 1). The gene structures, previously described for land plants (Sprink and Hartung 2014) with 15 exons and 14 introns, are identical and exhibit 93-95% identity including the introns (98% for the coding sequences). The three coding sequences which share 99% identity between each other encode proteins of 387, 386 and 387 amino acids, respectively (Table 1; Figure S1). The single Amino acid Polymorphism (SAP) mapped at position 45 and 74 for the 5A copy, 189 and 291 for the 5B copy and 64 and 66 for the 5D copy. In addition, the 5B copy carries an in-frame deletion of 3 nucleotides, removing an Alanine amino acid at position 72 (Figure S1).

The *SPO11-1 genes are conserved between wheat and wheat ancestors.*

We then sought to assess the sequences of the bread wheat *SPO11-1* copies with their ancestors. The ancestor of the B genome remains unknown but the wheat *SPO11-1*-5A and 5D sequences could be compared with their respective ancestors *Triticum urartu* and *Aegilops tauschii*, respectively. The *Triticum urartu* (TRIUR3_12346) and the *Aegilops tauschii* (LOC109743941) *SPO11-1* gene and the corresponding CDS sequences are reported in Table 1. The *Triticum urartu* and *Aegilops tauschii* *SPO11-1* genes share the conserved structure of 15 exons and 14 introns. Then, we re-annotated these sequences manually using the newly released wheat sequence as a basis for splicing sites and obtained two newly annotated CDS sequences of 1173 bp for *Triticum urartu* and 1164 bp for *Aegilops tauschii* encoding two proteins of 390 and 387 aa, respectively (Table 1).
Close comparison of the wheat TaSPO11-1-5A and -5D CDS and protein sequences with their ancestors showed that wheat TaSPO11-1-5D and *Aegilops tauschii*’s are 100% identical while TaSPO11-1-5A and *Triticum urartu*’s SPO11-1 share 99% identity (Figure S2). Furthermore, the SAP Cys/Ser at position 64 and Ser/Asp at position 66 between *Triticum urartu* and *Aegilops tauschii*’s SPO11-1 are conserved between the wheat A and D genomes (see above; Figures S1 and S2). These results suggest that the wheat SPO11-1 sequence has not changed since the polyploidization event. The extreme conservation of the homoeologous *SPO11*-1 genes and proteins within the Triticeae tribe suggests that they are all functional and presumably under a strong selection pressure in regard to their essential role in meiosis. In accordance with this, expression data showed that all three homoeologous genes are equivalently expressed during meiosis (Martin *et al.* 2018).

**The SPO11-1 proteins are highly conserved and shares key protein domains throughout plant kingdom.**

To get deeper insight into the evolution of SPO11-1 within plants, we first compared the *Arabidopsis thaliana* and wheat SPO11-1 amino acid sequences (Figure S3). We found significant identity (54-55%) similar to the BLAST-P analysis (e-value = 4e-156) (Table 1; Figure S3). This relies on the presence of seven highly conserved motifs present on the archaebacterial subunit A of topoisomerase VI (Bergerat, *et al.* 1997, Keeney, *et al.* 1997, Diaz *et al.* 2002, Malik, *et al.* 2007), *Arabidopsis thaliana* (Hartung and Puchta 2000, Hartung, *et al.* 2007, Shingu *et al.* 2010), wheat (Figure 2 and Figure S3), and many other organisms (Malik, *et al.* 2007). We note that the first motif that contains the catalytic Tyrosine residue for DSB formation (Tyr103 in Arabidopsis) (Bergerat, *et al.* 1997) is located at position 129 in the SPO11-1 A and D homoeologous wheat copies and at position 128 in the B copy. The second motif contains the invariant Arginine 130 (R156 in the A and D copies and R155 in the B copy) that is essential for the function of SPO11-1 *in vivo* (Diaz, *et al.* 2002, Shingu, *et al.* 2010). The fourth motif contains conserved residues implicated in TopoVI DNA binding activity (Glycine 215 and Arginine 222, 223 and 226 in Arabidopsis) (Shingu, *et al.* 2010). Finally, the seventh motif (10 amino acids) is fully conserved between Arabidopsis and wheat (Figure 2 and
Figure S3). This Toprim domain contains two conserved residues invariant in all species examined in this study (Lysine 332 and Glutamic acid 334 in Arabidopsis) that affect DSB formation in yeast when mutated (Diaz, et al. 2002). Interestingly, a 3D predictive structural modelling of the TaSPO11-1-5D protein shows that all seven motifs are linked to each other (Figure 2). This also shows that the essential DNA binding (5Y-CAP) and cleavage (Toprim) domains are clearly physically distinguishable from one another (Figure 3). Altogether, these analyses indicate that all SPO11 key residues and domains are conserved in the wheat homoeologous SPO11-1 proteins.

Then to extend our analysis to other plant species, we retrieved 155 SPO11-1 protein sequences from 153 plant species including monocots and dicots. A curation step based on the accuracy of the ATG and STOP codon position, the splicing and the integrity of the sequences allowed to retain and aligned 107 robust sequences from 105 species (Supplementary material S4). The amino acid sequence similarity ranged from 51% (Cucumis melo) to 100% (Aegilops tauschii). Compared to the wheat SPO11-1-5D copy, the mean similarity between these plant SPO11-1 proteins reach 65%. They all contain the seven most conserved motifs that landmark the SPO11 orthologs. We thus built a consensus sequence for each motif based on the alignment of the sequences (Figure 4) and calculated the average identities of each motif from the 107 sequences (Figure 4B). The motifs show very strong conservation with identities of 90.3, 89.1, 89.6, 93.3, 95.1, 99.1 and 96.5% for motifs 1, 2, 3, 4, 5, 6 and 7, respectively (Figure 4B). In particular, the active residues described above are fully conserved, except the last arginine in motif 4 (Arg226 in Arabidopsis) that is variant in 11/105 (10.5%) of the sequences. Among these, the six variants with a Glycine instead of Arginine, specifically belong to the Rosids clade and specifically to the Rosales order. Finally, the alignment of the 107 complete sequences led us to generate a phylogenetic tree which revealed a perfect separation into two groups corresponding to monocots (blue, Figure 5) and dicots (red, Figure 5). Interestingly, Amborella trichopoda, sister of the angiosperms (flowering plants), is at the frontier of the groups, suggesting it shares properties of both groups. Altogether, our extensive computational analysis of the SPO11-1 protein sequences in plants highlighted extensive conservation of the 7 key protein sequence motifs as well as limited evolution since the separation of monocots and dicots.
Heterologous expression of the Wheat TaSPO11-1-5D coding region restores the fertility of the Rice spo11-1 mutant.

The strong conservation of SPO11-1 protein sequences across plant species does not, in itself, prove that they are functionally interchangeable between species, especially given that they form parts of a multi-protein complex. To address this issue, we sought to determine whether or not the expression of the bread wheat SPO11-1 protein would rescue the meiotic phenotypes and sterility of the spo11-1 mutant of another monocotyledonous plant, namely rice. To perform this heterologous complementation assay, the wheat TaSPO11-1-5D coding sequence placed under the maize Ubiquitin promoter was introduced in our Oryza sativa ssp japonica var. Kitaake Osspo11-1-1 mutant line generated through CRISPR/Cas9 mutagenesis (see Methods; Fayos et al., unpublished). The Osspo11-1-1 mutation is a frameshift resulting from a single nucleotide (A) insertion in the ATG sequence (ATAG). The homozygous Osspo11-1-1 mutant lines fail to form chiasma during meiosis and are sterile (Yu, et al. 2010, Fayos, et al. 2019). Functional complementation can thus be easily visualized as an increase in seed production. Three spo11-1-1 homozygous plants (out of 33 primary transformants) carrying a single copy of the UBI::TaSPO11-1-5D transgene were obtained by transformation and remarkably, were fertile (Figure 6A). In the following T1 generation, the restoration of the rice spo11-1-1 fertility strictly co-segregated with the presence of the UBI::TaSPO11-1-5D transgene. Indeed, comparison of the number of filled spikelets in wild-type plants and in the progeny of two transformants grown in similar conditions shows that Rice spo11-1-1 mutant plants expressing the UBI::TaSPO11-1-5D transgene exhibit a fertility comparable to that of wild-type plants (Figure 6B). Altogether, these results demonstrate that the TaSPO11-1-5D coding sequence is functional and the wheat protein can functionally replace the rice SPO11-1 in its essential meiotic function.

Heterologous expression of the Wheat TaSPO11-1-5D coding region restores the fertility of the Arabidopsis spo11-1 mutant.

The success of the wheat-rice interspecies complementation prompted us to investigate whether or not
the wheat TaSPO11-1-5D complements the spo11-1 meiotic defects in Arabidopsis, a more distant
species belonging to a different clade. In Arabidopsis, the Atspo11-1 mutant fails to form meiotic
DSBs and exhibits a severe reduction in fertility (Grelon, et al. 2001). Thus, we placed the TaSPO11-
1-5D coding sequence under the control of the Arabidopsis RAD51 promoter and introduced this
construct into the Arabidopsis SPO11-1/spo11-1-2 heterozygous plants (Figure 7A). The RAD51
promoter is well expressed in Arabidopsis meiocytes (Chen et al. 2010, Yang et al. 2011, Walker et al.
2018) and known to drive successful complementation of other Arabidopsis meiotic mutants (Da Ines
et al. 2013). PCR genotyping of the SPO11-1 locus of 41 TaSPO11-1 primary transformants showed
that 15 were homozygous for the spo11-1-2 allele (spo11-1-2/spo11-1-2), 15 heterozygous (SPO11-
1/spo11-1-2) and 11 were wild-type (SPO11-1/SPO11-1). Remarkably, 14/15 spo11-1-2 homozygous
plants carrying the TaSPO11-1-5D transgene exhibited 15% to 70% fertility, instead of the 5%
residual fertility observed in the absence of the transgene (Figure 7C and Table S1). Then, we
monitored fertility in the progeny (T2 generation) of four randomly selected T1 plants. Consistently,
the restoration of fertility (30 to 80%) strictly co-segregated with the presence of the transgene (Figure
7D and Table S2). Thus, the heterologous expression of the Wheat TaSPO11-1-5D protein is able to
restore fertility in the Arabidopsis spo11-1 mutant.

**Restoration of the wild-type meiotic progression in the Arabidopsis spo11-1 mutant
expressing the TaSPO11-1-5D transgene.**

In wild-type plants, the meiotic chromosomes condense, recombine and synapse during prophase I
(Figure 8A-E). Full synapsis of the homologs is seen at late prophase I (Figure 8B). The chromosomes
then further condense and five bivalents (homologous chromosome pairs attached together by
chiasmata and sister chromatid cohesion) are observed at metaphase I (Figure 8C). Homologous
chromosomes then segregate to opposite poles to give two sets of five chromosomes at metaphase II
(Figure 8D). Meiosis II then proceeds and gives rise to four balanced haploid nuclei (Figure 8E). In
contrast, the spo11-1 mutants lack DSBs formation, hence recombination, pairing and synapsis of the
homologs (Figure 8F-G), manifested by the presence of 10 univalents instead of bivalents at
metaphase I (Figure 8H). Chromosome mis-segregation eventually produces unbalanced metaphase II (Figure 8I) and polyads (Figure 8J). In sharp contrast, the cytogenetic analysis of pollen mother cells from the spo11-1 plants expressing the TaSPO11-1-5D transgene revealed the presence of normal meiotic figures (Figure 8K-O). In particular, 5 bivalents were readily observed at metaphase I (Figure 8M). Subsequent proper homologous chromosome segregation at anaphase I (Figure 8N) followed by separation of sister chromatids during the second equational division resulted in 4 balanced meiotic products (Figure 8O). We note however that in accordance with the partial restoration of the fertility in the complemented plants, most metaphases I exhibited a mixture of bivalents and univalents (Figure 9A). Thus, to examine whether the expression of two copies of the transgene will quantitatively improve the faithful progression of meiosis, we characterized two independent transgenic lines homozygous for the TaSPO11-1-5D transgene. Their fertility (~ 40%) remained in the average range of the single copy transgene lines (line 3 and 36, see Figure 7). Cytologically, in the two complemented lines, we observed a mean of 1.7 (n=57) and 2.6 (n=74) bivalents per cell instead of 0.05 bivalents/cell (n = 19) in the spo11-1 mutants and 5 (n=36) in the wild-type meiocytes (Figure 9B and C). We also noted that a majority of bivalents in the complemented lines exhibited a rod-shaped structure with a single chiasma, although ring-shaped bivalents with at least two chiasmata were also observed (10 to 30% of bivalents for line 3 and 36, respectively). Accordingly, in both complemented lines, the number of chiasmata per cell significantly increased reaching a mean of 2 (± 1.7, n = 57) and 3.6 (± 2.6, n = 74), respectively. This is a strong increase compared to the spo11-1 mutant (0.05 ± 0.2, n = 19), yet lower than wild-type plants (9.3 ± 0.8, n = 36).

Altogether, these results demonstrate that the wheat SPO11-1 can functionally replace the absence of the Arabidopsis SPO11-1 ortholog, substantially restoring meiotic recombination and normal meiotic progression. Beyond the protein sequence homology, these results demonstrated its evolutionary conserved function.
Meiotic DSBs are formed in Arabidopsis spo11-1 mutants expressing Wheat TaSPO11-1-5D transgene.

The presence of bivalents and chiasmata indicate that meiotic recombination occurs and hence that meiotic DSBs are formed in plants expressing TaSPO11-1-5D transgene. Given that fertility is not fully restored, this however suggest that less DSBs might be produced in the complemented lines or, alternatively, that DSBs are repaired without forming COs. We thus sought to analyse the ability of TaSPO11-1-5D to form DSBs in meiotic cells. We performed immunolocalization of the strand-exchange protein RAD51 as a marker for DSB formation in both wild-type and spo11-1 + TaSPO11-1 complemented plants (line 36). As expected, numerous RAD51 foci were observed in early prophase I cells of wild-type plants (mean of 92 foci per cell, n = 39; Figure 10). RAD51 foci were also observed in pollen mother cell nuclei of spo11-1 mutant plants expressing TaSPO11-1-5D. However, a strong two-fold reduction in RAD51 foci formation was detected in these plants (mean of 45 foci per cell, n = 47; Figure 10). This strongly suggests that DSB levels are reduced in the complemented lines and this may explain the limited complementation by TaSPO11-1.

Wheat SPO11-1-5D functionally interacts with Arabidopsis SPO11-2 and MTOPVIB to induce meiotic recombination.

Current knowledge suggests that SPO11-1 does not exhibit DNA cleavage activity alone but acts in a protein complex, physically interacting with the SPO11-2 and MTOPVIB proteins in Arabidopsis and functionally related orthologs in other organisms, in order to form an active topoisomerase VI-like complex that catalyses meiotic DSB formation (Figure 1, Robert et al., 2016; Vrielynck et al., 2016). So, to determine whether TaSPO11-5D also needs the presence of the Arabidopsis SPO11-2 and MTOPVIB proteins to induce meiotic recombination, we crossed our spo11-1_TaSPO11-1-5D transgenic plants with Arabidopsis spo11-2 or mtovIB mutant lines and analysed the fertility of the double mutants. Clearly, as shown in Figure 11, the presence of the TaSPO11-1-5D transgene did not rescue the sterility of the spo11-1 spo11-2 and spo11-1 mtovIB double mutant plants. This excludes
the possibility that TaSPO11-1 induce DSBs independently of SPO11-2 (non-plant organisms have a single SPO11 protein) and confirms the need for the MTOPVIB to form DSBs.

**Discussion**

TaSPO11-1 homoeologous copies are highly similar between each other and to those from angiosperms.

To identify SPO11-1 genes from wheat (*T. aestivum*), we exploited the first assembled and annotated pseudomolecule sequence of the wheat genome (IWGSC 2018). Using *in silico* assignment of the Arabidopsis SPO11-1 protein (At3g13170), we readily identified the three homoeologous copies, mapping on chromosomes 5A, 5B and 5D and indicating good conservation of the protein sequence between the two species. The three homoeologous wheat copies are highly similar with ~95% identity at the nucleotide level for the genomic sequences. This is consistent with data from expression of wheat genes (Ramirez-Gonzalez *et al.* 2018) showing an homoeologous SNP diversity ranging from 95.0% to 97.2% within triads (genes present in only three homoeologous copies).

We also observed that the D copy (TaSPO11-1-5D) is identical to the copy from *Ae. tauschii*, the donor of the D genome, while the A copy (TaSPO11-1-5A) is only very slightly different from that of *T. monococcum ssp urartu*, the donor of the A genome. Divergence between the A and the B genome lineages occurred ~7 million years ago (MYA; (Marcussen *et al.* 2014)) while the D genome diverged from the A and B genomes, 1 to 2 million years after. The two successive polyploidization events giving rise to *T. aestivum* occurred at least 0.58 to 0.82 MYA for the first one and 0.23 to 0.43 MYA for the second one. In addition, it is suggested that only a few accessions of *Ae. tauschii* contributed to the D genome of bread wheat (Giles and Brown 2006).

SPO11-1 proteins are characterized by the presence of several conserved domains (Bergerat, *et al.* 1997, Keeney, *et al.* 1997). Accordingly, these seven domains are highly conserved in all the plant sequences that we examined. In particular, the essential residues for DNA cleavage (Tyr103 in Arabidopsis) or binding (Gly215, Arg222, Arg223 and Arg226 in Arabidopsis) are highly conserved although their position changed slightly according to the total size of the protein, which indeed varies
among species for yet unknown reasons. A previous study using 42 SPO11-1 sequences from land plants, but not wheat, indicated that SPO11-1 is highly conserved in plants (Sprink and Hartung 2014). Here, we analysed SPO11-1 sequences from more than 100 plants, including wheat, and show that SPO11-1 exhibits high sequence identities. In particular, more than 90% identity was observed in the most broadly evolutionary conserved functional domains. Overall, our extensive phylogenetic analyses based on sequence comparison of plant SPO11-1 indicates that this protein evolved slowly and exhibits an evolutionary pattern consistent with known relationships between plant species.

Heterologous complementation analyses reveal functional conservation of the SPO11 complex features.

Beyond the computational analyses of the SPO11-1 genes, we asked to what extent the function of SPO11-1 is also evolutionary conserved throughout plants, testing the complementation of the rice and Arabidopsis spo11-1 mutants with the SPO11-1-5D cDNA from bread wheat. Wheat and rice are monocots while Arabidopsis thaliana is a more distantly related dicot. Remarkably, our data show that expression of the wheat gene was able to complement both rice and Arabidopsis mutants, with full complementation in the former.

Furthermore, our analyses in Arabidopsis show that the complementation by TaSPO11-1 restores DSB formation (RAD51 foci) and recombination (chiasmata) and still requires the presence of the wild-type Arabidopsis SPO11-2 and MTOPVIB genes. Wheat SPO11-1 and Arabidopsis SPO11-2 and MTOPVIB are able to interact and form a functional inter-species complex, albeit resulting in only a partial restoration (10 to 70% with most lines showing 20-40% restoration). Immunolocalization of the RAD51 recombinase indicates that less DSBs are formed in the complemented plants (50% of wild-type level in the tested line) and this likely explains the partial complementation.

Partial complementation in Arabidopsis with Arabidopsis clones has been frequently observed, as for Arabidopsis SPO11-1 (Xue et al. 2018). Many factors could influence the efficacy of complementation: the use of a recipient T-DNA mutant plant, T-DNA integration, the choice of the promoter or the use of CDS or genomic sequences. Although this cannot be excluded, we do not think that reduced DSBs formation in our transgenic plants result from lower expression of TaSPO11-1.
gene. Using the same strategy as for TaSPO11-1, we could show that expression of TaSPO11-2 cDNA driven by the RAD51 promoter is able to fully restore fertility of the Arabidopsis spo11-2 mutant (Benyahya et al., unpublished). This indicates that RAD51 promoter allows sufficient transcription of TaSPO11 genes. Conversely, translation may be affected. This is particularly true since codon usage bias is well known to be different in monocots and dicots (Plotkin and Kudla 2011, Camiolo et al. 2015). However, without specific TaSPO11-1 antibodies this hypothesis cannot be tested. In the present case of expression in an heterologous species, an additional key factor for the incomplete complementation and the reduced DSB formation is the amino acid sequence divergence of the transgenic and endogenous proteins that needs to interact in a multi-protein complex. Sufficient restoration of the mutant phenotypes has been obtained to conclude on the formation of functional interactions but the incomplete complementation uncovered subtle deficiencies of interest. For instance, TaSPO11-1 may be less prone to interact with other DSB-associated proteins and to form an active complex. In this context, it will be interesting to individually assay the other members of the SPO11 complex and attempt to co-express in Arabidopsis mutants the wheat TaSPO11-1, TaSPO11-2 and TaMTOPVIB once the likely homoeologous genes and coding regions have been well identified. It will also be interesting to more extensively analyse amino acid sequence divergence and its effect on interaction with other essential DSB-associated endogenous proteins. Eventually, working with hypomorphic mutants of the meiotic SPO11/MTOPVIB complex with reduced DSBs could be very valuable to better understand the relationship between DSB formation and CO regulation (CO assurance, homeostasis and interference).

**Conclusion**

In this study, we isolated the three wheat homoeologous copies for SPO11-1: TaSPO11-1-5A, TaSPO11-1-5B, TaSPO11-15D. The three copies are highly similar between each other and with those from diploid ancestors, *T. urartu* and *Ae. tauschii*, suggesting that all three are functional. SPO11-1 protein is very well conserved across angiosperms with conserved domains. Remarkably, due to the high level of similarity, TaSPO11-1-5D protein was able to restore the fertility of rice and Arabidopsis
spo11-1 mutants. This also showed that the wheat proteins could be used (and hence further studied) in other more tractable model plants. This is of particular interest in polyploid species in which the redundancy of function brought by the homoeologous genes and variant alleles adds an additional level of genetic complexity in the wild type context, and where the construction of appropriate mutants remains technically difficult and time-consuming due to the polyploidy of the genome (Ramirez-Gonzalez, et al. 2018).
Experimental Procedures

Plant material and growth conditions

The rice mutant was obtained through CRISPR/Cas9 genome editing as described in Fayos et al., (unpublished). Rice plants were cultivated in controlled conditions with a temperature of 28°C during the day and 24°C during the night, with 60% hygrometry. The natural light is completed by artificial sodium light (700μmol/m²/s). The Arabidopsis thaliana spo11-1-2, spo11-2-3 and mtovIB-2 mutants used in this work have been described previously (Grelon, et al. 2001, Hartung, et al. 2007, Vrielynck, et al. 2016). Arabidopsis plants were grown under the following standard conditions: seeds were stratified in water at 4°C for 2 days and grown on soil or in vitro on 0.8% agar plates, 1% sucrose and half-strength Murashige and Skoog salts (M0255; Duchefa Biochemie). Plants were cultivated in a greenhouse or a growth chamber with a 16/8 hour light/dark cycle, at 23°C and 60% relative humidity.

Recovery and synthesis of TaSPO11-1

Wheat genome D SPO11-1 DNA sequence (TraesCS5A02G391400) was first retrieved through a BLAST analysis research on the newly annotated wheat genome sequence (IWGSC 2018) using the Arabidopsis thaliana SPO11-1 protein sequence (At3g13170) as an input with basic BLAST parameters. The annotated CDS sequence was determined using Triannot pipeline (Leroy et al. 2012). CDS sequence was synthesized with a short additional sequence at the 5’ end of the gene (coding for the peptide PEFMAMEAPGIR) and flanked with GATEWAY attB sites. Synthesized product was inserted into pDONR/Zeo and further verified by sequencing.

3D modelling and rendering

3D structural model of TaSPO11-1-5D protein was generated by homology modelling on PHYRE2 online pipeline ((Kelley et al. 2015), http://www.sbg.bio.ic.ac.uk/~phyre2/html/page.cgi?id=index) with intensive modelling mode and TaSPO11-1-5D full sequence as amino acid sequence. Rendering was made with PyMOL 2.3.3 software.
Recovery of SPO11-1 protein sequences from multiple plant species, alignment, logo and phylogeny

We retrieved all plant SPO11-1 sequences using Wheat and Arabidopsis SPO11-1 sequences as query on NCBI BLAST website (https://blast.ncbi.nlm.nih.gov/Blast.cgi) with nr database. The Barley SPO11-1 sequence was retrieved using IPK barley BLAST server website (https://webblast.ipk-gatersleben.de/barley_ibsc/viroblast.php) and the barley AA (HC and LC) Morex v2.0 database. Multiple Sequence Alignment was done using ClustalW with basic parameters. Motifs logo were built using WebLogo website (https://weblogo.berkeley.edu/) (Crooks et al. 2004). Phylogeny was done with the following settings: ClustalW alignment and PHYLIP Neighbor Joining for the construction of the tree.

Cloning of TaSPO11-1 and plant transformation

For rice complementation, a LR GATEWAY recombination cassette was inserted in a pZmUBI-tNos vector at MCS location using BamH1 restriction sites to form a pZmUBI-LR-tNos vector. The complete TaSPO11-1 CDS fragment was inserted in this vector under the control of the UBI promoter using GATEWAY cloning sites. Rice seed embryo-derived callus from line segregating the OsSPO11-1-1 mutation were then transformed accordingly to the method described in (Sallaud et al. 2003). Primary transformants expressing the TaSPO11-1 transgene were selected on hygromycin selection medium. Mutation in the ATG of OsSPO11-1 (Loc_Os03g54091) was ascertain by PCR (primers: SPO11-R1 ccaaaatcttgtgggtgct and SPO11-F2 cggaggagcagtagttctgg) and sequencing. Presence and integrity of the transgene was also verified by PCR (primers: pUBI-F cttgatacttggatgatggc and tNOS-R cgcaagaccggcaacaggttc) and sequencing.

For Arabidopsis complementation, the complete SPO11-1 CDS fragment was cloned into the GATEWAY destination vector pMDC32 (Curtis and Grossniklaus 2003) in which the 35S promoter was replaced with the Arabidopsis RAD51 promoter (1031bp upstream of the RAD51 ATG; (Da Ines, et al. 2013) with a HindIII/AscI digest. The plasmid was then inserted in an Agrobacterium tumefaciens C58C1 strain which was subsequently used to transform Atspo11-1-2 heterozygous
mutant plants by the floral dip method (Clough and Bent 1998). T1 seeds from the Agrobacterium-
transformed plants were sown on soil and T1 transformants were selected for Hygromycin resistance
on 0.5X MS/ 1% sucrose/ 0.8% agar plates containing 15µg/ml Hygromycin B Gold (InvivoGen).
Presence of the transgene and genotypes of transformants were verified by PCR.

**Arabidopsis male meiotic chromosome spreads**

Chromosome spreads were prepared according to (Ross et al. 1996). Whole inflorescences were fixed
in ice-cold ethanol/glacial acetic acid (3:1) for 3 x 30 min and stored at -20°C until further use.
Immature flower buds were rinsed twice at room temperature in distilled water for 5 min. This was
followed by two washes in citrate buffer for 5 min. Buds of appropriate size were selected under a
binocular microscope and incubated for 75 to 90 minutes on a slide in 100µL of enzyme mixture
(0.3% w/v cellulase (Sigma), 0.3% w/v pectolyase (Sigma) and 0.3% cytohelicase (Sigma)) in a moist
chamber at 37°C. Each bud was then softened for 1 minute in 15µL of acetic acid (60%) on a
microscope slide at 45°C, fixed with ice-cold ethanol/glacial acetic acid (3:1) and air-dried.
Eventually, slides were mounted in Vectashield mounting medium with DAPI (1.5 µg.mL⁻¹; Vector
Laboratories Inc.).

For chiasma counting, number of chiasmata at metaphase I was estimated based on bivalent
configuration: rod-shaped bivalents were considered to contain a single chiasma and ring-shaped
bivalents, two (one on each arm) (Sanchez Moran et al. 2001).

**Immunolocalization of proteins in pollen mother cells (PMCs)**

Spreads of PMCs for immunolocalization of RAD51 were performed as described previously
(Armstrong et al. 2002). Primary antibodies used for immunostaining were: anti-ASY1 raised in
guinea Pig (1:500) (Higgins et al. 2004) and anti-RAD51 raised in rat (1:500) (Kurzbauer et al. 2012).

**Microscopy**
All observations were made with a motorized Zeiss AxioImager.Z1 epifluorescence microscope (Zeiss) using a PL Apochromat 100X/1.40 oil objective. Photographs were taken with an AxioCam MRm camera (Zeiss) driven by ZEN Pro software (Zeiss). Captured images were further processed and adjusted for brightness and contrast on ZEN Pro and ImageJ/FIJI software.

**Statistical analysis**

All graphs and statistical analyses were performed using software GraphPad PRISM 6. To determine whether differences between two groups were statistically significant, groups were compared using ordinary one-way ANOVA and Holm-Sidak test to account for multiple comparisons. A *P*-value of 0.05 or less was considered to be statistically significant.
Acknowledgements

We thank Mathilde Grelon for providing spo11-1, spo11-2 and mtopVIB mutants, and Chris Franklin and Peter Schlögelhofer for ASY1 and RAD51 antibodies, respectively. The authors thank members of the GeCO group and members of the recombination group for helpful discussions. This work was supported by CNRS, INRA, INSERM, Université Clermont Auvergne, and the French government IDEX-ISITE initiative 16-IDEX-0001 (CAP20-25). RM was funded by a PhD grant from the Association Nationale Recherche Technologie (ANRT CIFRE Grant No. 2014/1020).

Supplementary Materials Legends

Figure S1. Sequence alignment of Bread wheat SPO11-1 proteins. Alignment was generated using ClustalW. Numbers indicate amino acid positions. Red letters indicate single amino acid polymorphism and blue letters denote amino acid deleted in the B genome. Identical residues are highlighted in black and asterisks, colons and dots under the sequences, indicate identical, conserved and semi-conserved residues, respectively.

Figure S2. Sequence alignment of SPO11-1 proteins from bread wheat and ancestors. Alignment of SPO11-1 from genome A, D, and their ancestors Triticum urartu and Aegilops tauschii was generated using ClustalW. Numbers indicate amino acid positions. Amino acid highlighted in cyan and yellow designate single amino acid polymorphism.

Figure S3. Sequence alignment of bread wheat and Arabidopsis proteins. Alignment was generated using ClustalW. Numbers indicate amino acid positions. Red squares with roman numerals indicate the conserved motifs. Identical residues are highlighted in black and asterisks, colons and dots under the sequences, indicate identical, conserved and semi-conserved residues, respectively.
Figure S4. Sequence alignment of 107 plant SPO11-1 proteins. Alignment was generated using ClustalW 2.0. Numbers indicate amino acid positions. Asterisks, colons and dots under the sequences, indicate identical, conserved and semi-conserved residues, respectively.

Table S1. Seed number per silique in wild-type, Atspo11-1 and Atspo11-1_TaSPO11-1 primary transformants. 10 to 12 fruits were counted per plant. (n.d. : not determined). P-value were calculated using ordinary one-way ANOVA and Holm-Sidak test to account for multiple comparisons.

Table S2. Seed number per silique in wild-type and in the progeny of Atspo11-1-2_TaSPO11-1 transformants. Seeds were counted in 4 plants per genotype and 8 fruits per plant. (n.d. : not determined). P-value were calculated using ordinary one-way ANOVA and Holm-Sidak test to account for multiple comparisons.
References


Table 1: Wheat, its ancestors and *Arabidopsis thaliana* *SPO11-1* gene characteristics. gDNA: genomic DNA; CDS: coding sequence; bp: base pair; aa: amino acid.

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Figures Legends

Figure 1: Schematic representation of the putative SPO11 meiotic DSB catalytic complex.
This protein complex is suggested to be an heterotetramer composed of one heterodimer (SPO11-1 and SPO11-2; green and blue, respectively) and one homodimer (MTOPVIB; orange) (Robert, et al. 2016b).

Figure 2: Protein sequence alignment of *Triticum aestivum* and *Arabidopsis thaliana* SPO11-1.
Only the seven conserved motifs are shown. Conserved amino acids are highlighted in green and similar amino acids are highlighted in yellow. Catalytically active tyrosine and conserved glycine and arginine involved in DNA-binding activity are in red.

Figure 3: 3D modelling of TaSPO11-1-5D protein.
A 3D structural model of SPO11-1 obtained by sequence homology modelling with PHYRE2 online pipeline. This SPO11-1 model consists of amino acid residues 1 to 387. 336 residues over 387 (87%) were modelled with >90% accuracy. A and B are mirror views of the structural model rendered with PyMol software. The seven conserved motifs are shown in green, the Toprim domain in light pink and the 5Y-CAP domain in light yellow. Catalytic tyrosine is depicted in magenta and the glycine and arginine essential for DNA-binding activity are depicted in red and blue, respectively.

Figure 4: Sequence and conservation level of the seven conserved motifs in SPO11-1 proteins from 105 species.
(A) Consensus sequence of the seven conserved motifs extracted from 107 SPO11-1 sequences. (B) Conservation level of the seven motifs within 107 SPO11-1 sequences. White boxes: lower and upper quartile of conservation rate for each motif. Black lines: mean conservation rate for each motif. Dashed lines: maximum and minimum conservation rate values. Circles: outliers.

Figure 5: Phylogenetic tree of plant SPO11-1 proteins from 105 species.
Dicotyledons are shown in red, monocotyledons in blue and *Amborella trichopoda*, sister of the angiosperms, is shown in purple.

**Figure 6: Wheat TaSPO11-1-5D restores fertility of the Rice spo11-1 mutant.**

(A) Pictures of rice *spo11-1* mutant plants and panicles expressing or not the wheat TaSPO11-1-5D transgene. Rice *spo11-1* plants are sterile and develop panicles with empty spikelets (right panel). In contrast, *spo11-1* mutant plants expressing wheat TaSPO11-1-5D are fertile and develop panicles with filled spikelets (left panel). (B) Percentage of filled spikelet per panicle in wild-type plants and progeny of two *spo11-1* transformants expressing wheat TaSPO11-1-5D (T2 and T64). In the box and whiskers, each dot represents the percentage of filled spikelet per panicle in one plant (n = 6 for wild-type plants, n = 5 for *spo11-1*, and n = 5 for *spo11-1* TaSPO11-1-5D plants). Means are represented by a + and horizontal bars denote medians.

**Figure 7: Wheat TaSPO11-1-5D restores fertility of the Arabidopsis spo11-1 mutant**

(A) Schematic representation of the pRAD51:TaSPO11-1 construct. (B) Wild-type plants have long siliques full of seeds, while Atspo11-1 mutants are sterile and exhibit very short siliques. Expression of the TaSPO11-1 in Atspo11-1 mutants restores fertility. (C) number of seeds per silique in Wild-type, Atspo11-1, and 15 Atspo11-1 + pRAD51:TaSPO11-1 independent primary transformants. Each dot represents the number of seeds in one silique. (D) number of seeds per silique in Wild-type, and T2 progeny of 4 Atspo11-1 + pRAD51:TaSPO11-1 independent primary transformants (T3, T12, T13 and T36 as indicated under graph). Each dot represents the number of seeds in one silique. Blue dots show number of seeds per silique in wild-type, red dots Atspo11-1 mutants and black dots represent Atspo11-1 mutants expressing TaSPO11-1.

**Figure 8: Meiotic progression in wild-type, Atspo11-1 mutants and Atspo11-1 mutants complemented with wheat TaSPO11-1.**

DAPI staining of chromosomes during meiosis in Arabidopsis (A-E) wild-type, (F-J) *spo11-1* and (K-O) *spo11-1* + TaSPO11-1 plants. (A, F, K) Early prophase I, (B, G, L) Late prophase I, (C, H, M)
Metaphase I, (D, I, N) Metaphase II, and (E, J, O) Tetrad. In wild-type, cells show pairing and
synapsis of homologous chromosomes at late prophase I (B), five bivalents at metaphase I (C), two
groups of five chromosomes at Metaphase II (D) and balanced tetrads (E). spo11-1 mutants exhibit
defective synapsis (G), univalent in Metaphase I (H) and unbalance Metaphase II (I) and polyads (J).
In spo11-1 expressing wheat TaSPO11-1 (K-O), wild-type meiotic figures can be observed. (Scale
Bar: 10 µm).

Figure 9: Expression of wheat TaSPO11-1 in Atspo11-1 mutant promotes bivalent formation
(A) Representative images of Metaphase I are shown (Scale Bar: 10 µm).
(B) Mean number of bivalents (dark grey) and pairs of univalent (grey) per meiosis
(C) Bivalents per cell (in percentage). Atspo11-1 plants expressing TaSPO11-1 show a significant
increase of bivalent formation when compared to Atspo11-1 mutants. Number of cells analyzed is
indicated in parentheses.

Figure 10: Reduced numbers of RAD51 foci in Atspo11-1 mutants complemented with wheat
TaSPO11-1.
(A) Immunolocalization of RAD51 (green) and the chromosome axis protein ASY1 (red) on
leptotene/zygotene meiotic chromosome spreads. (Scale Bars: 5 µm). (B) Quantification of RAD51
foci per positive cell throughout prophase I in both wild-type and Atspo11-1 mutants expressing
TaSPO11-1 (T36). (p-value < 0.0001, Mann-Whitney test).

Figure 11: Complementation of Atspo11-1 by TaSPO11-1 requires presence of AtSPO11-2 and
AtMTOPVIB.
(A) Pictures of siliques from wild-type, and mutant plants expressing or not TaSPO11-1. Genotype of
the mutants are indicated above pictures. Fertile plants have long siliques while sterile plants have
short siliques. Fertility of Atspo11-1 + TaSPO11-1 is lost by deletion of either AtSPO11-2 or
AtMTOPVIB. (B): Mean number of seeds per silique. Each dot represents the number of seeds in one
silique.