Downregulation of Barley Regulator of Telomere Elongation Helicase 1 Alters the Distribution of Meiotic Crossovers

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Programmed meiotic DNA double-strand breaks (DSBs), necessary for proper chromosomal segregation and viable gamete formation, are repaired by homologous recombination (HR) as crossovers (COs) or non-crossovers (NCOs). The mechanisms regulating the number and distribution of COs are still poorly understood. The regulator of telomere elongation helicase 1 (RTEL1) DNA helicase was previously shown to enforce the number of meiotic COs in Caenorhabditis elegans but its function in plants has been studied only in the vegetative phase. Here, we characterised barley RTEL1 gene structure and expression using RNA-seq data previously obtained from vegetative and reproductive organs and tissues. Using RNAi, we downregulated RTEL1 expression specifically in reproductive tissues and analysed its impact on recombination using a barley 50k iSelect SNP Array. Unlike in C. elegans, in a population segregating for RTEL1 downregulated by RNAi, high resolution genome-wide genetic analysis revealed a significant increase of COs at distal chromosomal regions of barley without a change in their total number. Our data reveal the important role of RTEL1 helicase in plant meiosis and control of recombination.

Keywords: barley, crossover, RTEL1, meiosis, recombination, RNAi

INTRODUCTION

Meiosis is a specialised cell division that generates haploid gametes through genome duplication followed by two rounds of chromosome segregation. At early prophase I, the homologs pair and synapse concomitantly with the introduction of a large number of DNA double-strand breaks (DSBs) by the topoisomerase type II-like protein Spo11 and its cofactors (Edlinger and Schlögelhofer, 2011; Panizza et al., 2011; Yadav and Bouuaert, 2021). The number of these programmed DSBs has been shown to be determined by the level of Spo11 activity (Xue et al., 2018) and restrained by the homolog engagement initiation during synaptonemal complex (SC) formation (Mu et al., 2020). Genome-wide mapping of meiotic DSB sites has been determined using chromatin immunoprecipitation followed by sequencing (ChIP-seq) and single-stranded DNA sequencing (SSDS;
Khil et al., 2012) in several species including mouse (Lange et al., 2016), Arabidopsis (Choi et al., 2018), and maize (He et al., 2017). These studies revealed that the meiotic DSB landscape is tightly controlled by multi-factorial local DNA and chromatin epigenetic features.

The repair of meiotic DSBs by homologous recombination (HR) is initiated by exonucleolytic resection, a process recently shown to be controlled by ATM kinase in mouse (Yamada et al., 2020). Progressively, meiotic DSBs are repaired by a dynamic interplay of multiple recombinases resulting in different joint molecules (Wild et al., 2019). Resolvases act on these DNA structures to yield crossover (CO) and non-crossover (NCO) recombinants (San-Segundo and Clemente-Blanco, 2020). Only a small subset of the initial large number of meiotic DSBs are stepwise matured to form COs and their quasi-constant final number per meiocyte is species dependent and under tight homeostatic control (Cole et al., 2012). Although still poorly understood, the mechanisms of CO designation involve multiple factors ranging from the local genomic sequence to the chromatin structure (reviewed by Gray and Cohen, 2016; Dluzewska et al., 2018). Most eukaryotes including plants generate two types of COs: class I and class II that are sensitive or not to interference control, respectively (Wang et al., 2015). The majority of COs (class I) are produced through the so-called ZMM pathway and spaced out along the chromosomes, whilst non-interfering COs are on the Mus81-dependent pathway (Lynn et al., 2007; Shinohara et al., 2008; Osman et al., 2011; Zhang et al., 2019). The mechanism of interference is still unknown, although some advance has been made by the discovery that topoisomerase II (Top2) plays a key role in this CO distribution pathway in yeast (Zhang et al., 2014; Wang et al., 2015). However, in Arabidopsis, this topoisomerase was found to play a role in the resolution of meiotic chromosomal interlocks (Martinez-Garcia et al., 2018) whilst the CO interference is imposed by the SC (Capilla-Pérez et al., 2021; France et al., 2021).

Several CO antagonists have been identified in different forward genetic screens in Arabidopsis. Generally, genetic manipulation of these meiotic genes reduces fertility due to disturbance of the SC and/or meiotic recombination with changes in CO frequency and distribution. It has been widely proposed that fertile plants with increased recombination would be useful for crop improvement. In Arabidopsis, Crismani et al. (2012) identified suppressor mutations that restored fertility to a class of semi-sterile meiotic mutants in the ZMM pathway. They identified independent fertility restorers, including mutations in a conserved Fanconi anemia complementation group M (FANCM) helicase that had increased recombination (Crismani et al., 2012; Knoll et al., 2012) and FANCM-associated proteins MHF1 and MHF2 (Girard et al., 2014). The FANCM ortholog was found to direct NCO in competition with the pathway of class II COs (Lorenz et al., 2012). Several other fertility restorers in different meiotic genes and their interactors including RECQ4 (Séguèla-Arnaud et al., 2015), Top3 α (Séguèla-Arnaud et al., 2015, 2017), and FIGL1 (Girard et al., 2015; Fernandes et al., 2018a) have now been identified and shown to increase the overall recombination frequency in Arabidopsis (Séguèla-Arnaud et al., 2015; Fernandes et al., 2018a,b) and have been tested for their breeding utility in crops like Brassica (Blary et al., 2018), rice (Zhang et al., 2017; Fayos et al., 2019), and tomato (de Maagd et al., 2020).

In addition to FANCM and RecQ4, several other DNA helicases including the regulator of telomere elongation helicase 1 (RTEL1) are essential in ensuring genome stability during both vegetative and reproductive stages in plants (Dorn and Puchta, 2019). RTEL1 is a multifunctional DNA helicase essential for telomere maintenance, DNA replication, repair, and recombination; mutations in RTEL1 have been associated with several severe human diseases (Vannier et al., 2014). Its role as a suppressor of homologous recombination analogous to yeast SRS2 helicase was first reported in C. elegans and human cells (Barber et al., 2008; Villeneuve, 2008). The same group went on to demonstrate its function as an enforcer of meiotic CO interference and homeostasis in C. elegans (Youds et al., 2010). Genetic analysis of C. elegans rtle1 mutants showed they lacked CO interference resulting in more COs in both chromosome arm and regions even though the number of DSBs was not affected. This work suggested that RTEL1 regulates meiotic recombination by dismantling transient strand invasions in D-loop joint intermediates and channelling DSBs repair by the synthesis-dependent strand annealing (SDSA) pathway promoting NCOs.

Based on the findings of Youds et al. (2010), RTEL1 is therefore a potential target for improving crop breeding by altering patterns of recombination. However, its role in meiotic recombination in plants has not been reported. The two studies of RTEL1 in plants were limited to the vegetative stage in Arabidopsis (Recker et al., 2014; Hu et al., 2015). These studies showed that RTEL1 helicase plays an essential role in safeguarding genome stability by repairing replication defects and telomere maintenance. Using a highly sensitive β-glucoronidase (GUS) based HR reporter system, it was demonstrated that RTEL1 helicase suppresses HR in somatic cells. Here, we studied the RTEL1 gene structure and expression in barley (Hordeum vulgare) and downregulated its expression specifically in reproductive tissues of barley cultivar Golden Promise using an RNAi construct under the control of meiotic DMC1 promoter. Selected RNAi lines were crossed to a different cultivar (cv. Bowman) and the derived F2 population analysed using a 50k iSelect SNP Array to determine at high resolution the impact of RTEL1 downregulation on the landscape of meiotic recombination.

**MATERIALS AND METHODS**

**Plant Material**

Barley (Hordeum vulgare) cultivars (cv.) Bowman (Bw) and Golden Promise (GP) and RTEL1RNAi transgenics were grown in compost in a standard heated greenhouse as described by Barakate et al. (2014).
Multiple Alignment and Phylogenetic Analysis

For RTEL1 phylogeny, orthologous sequences were identified from human (*Homo sapiens*), mouse (*Mus musculus*), African clawed frog (*Xenopus laevis*), Zebra fish (*Danio rerio*), *C. elegans*, fruit fly (*Drosophila melanogaster*), *Physcomitrella patens*, Amborella trichopoda, tobacco (*Nicotiana tabacum*), tomato (*Solanum lycopersicum*), soy bean (*Glycine max*), maize (*Zea mays*), and pineapple (*Ananas comosus*). The resulting T1 transgene was generated in the glasshouse and crossed to Bowman cultivar. The obtained T2:F1 seeds of the line *HvRTEL1RNAi* were germinated in the presence of hygromycin, and resistant seedlings were grown in the glasshouse and the crosses were confirmed using Kompetitive Allele-Specific PCR (KASP™) assay (Rasheed et al. 2016) and F1 polymorphic markers (Supplementary Table 1). The resulting T2:F2 seeds of *Bw×HvRTEL1RNAi* were collected and used in KASP assay with F1 polymorphic markers (Supplementary Table 1) and 50K iSelect SNP Array (Bayer et al., 2017).

RNAti Construct

A 750bp fragment corresponding to the 3’ end of *HvRTEL1* coding sequence was PCR-amplified using two *HvRTEL1* specific primers with Gateway sites (attB1-HvRTEL1 and attB2-HvRTEL1) and the full-length *HvRTEL1* cDNA plasmid as the DNA template. The obtained PCR fragment was cloned into the entry vector pDONR207 Gateway BP Clonase (Life Technologies). Using Gateway LR Clonase (Life Technologies), *HvRTEL1* fragment was then transferred into the destination vector pBRACT207m-HvDMC1p, a derivative of pBRACT207, 1 where the maize ubiquitin 1 promoter was replaced by the barley DMC1 promoter. The final *HvRTEL1RNAi* construct therefore has the RNAi cassette under the transcriptional control of the barley DMC1 promoter. As a control, a partial fragment of 715 bp of β-glucuronidase (GUS) gene was cloned into the original pBRACT207 destination vector to make the construct GUSRNAi. A list of oligonucleotides used for plasmid construct preparation can be found in Supplementary Table 4.

Preparation and Screening of Transgenic Plants

The strategy for regenerating and screening the transgenic plants is summarised in Supplementary Method 1. *HvRTEL1RNAi* construct was transferred into *Agrobacterium tumefaciens* AGL1 strain containing the plasmid helper pSOUP. Barley cv. Golden Promise transformation was performed by the Functional Genomics Facility at The James Hutton Institute, Dundee, United Kingdom using immature embryos under hygromycin selection as described previously (Barakate et al., 2014; Colas et al., 2019). Hygromycin resistant T0 *HvRTEL1RNAi* transgenic lines were grown in the glasshouse to maturity. Twenty T1 seeds per line were germinated on plates containing 0.5% phytgal in the presence of 100μg/ml of hygromycin (Jacobsen et al., 2006) to detect the presence of the transgene. Eight hygromycin resistant transgenic seedlings were transferred into 15cm pots of soil and grown in the glasshouse and their vegetative growth and fertility were monitored. In the T1 generation, fertility was scored for all transgenic lines and semi-sterile lines were selected for further analysis. Reduced fertility was used as a proxy for monitoring the impact of *RTEL1* down-regulation on meiosis as we demonstrated previously for several other meiotic genes in barley (Barakate et al., 2014; Colas et al., 2019).

Crosses and Recombination Assays

T1 seeds of the selected lines were germinated in the presence of hygromycin and resistant seedlings with the *HvRTEL1RNAi* transgene were grown in the glasshouse and crossed to Bowman cultivar. The obtained T2:F1 seeds of the line *HvRTEL1RNAi* were germinated in the presence of hygromycin, and resistant seedlings were grown in the glasshouse and the crosses were confirmed using Kompetitive Allele-Specific PCR (KASP™) assay (Rasheed et al. 2016) and F1 polymorphic markers (Supplementary Table 1). The resulting T2:F2 seeds of *Bw×HvRTEL1RNAi* were collected and used in KASP assay with F1 polymorphic markers (Supplementary Table 1) and 50K iSelect SNP Array (Bayer et al., 2017).

For KASP recombination assay, genomic DNA was extracted from glasshouse grown 2-week-old leaf tissue from 96 T2:F1 individuals containing the *HvRTEL1RNAi* transgene and 96 wild type (*Bw×GP*) F1 individuals using Qiagen QIAamp™ 96 DNA QIAcube HT kits on a QIAcube HT automated platform (Qiagen, Hilden, Germany). DNA quality was assessed using a Nanodrop 2000 (Thermo Scientific, Massachusetts, United States) with a requirement of >1.8 for 260/280 and 260/230 absorbance ratios. DNA was quantified using Picogreen (Thermo Scientific, Massachusetts, United States). Two independent ca. 10-cM intervals on 1H (centromeric) and 3H (distal) were studied using KASP assays (LCG, Biosearch, Hoddesdon, United Kingdom).

For 50K iSelect SNP Array based recombination assays, a total of 300 ng of lyophilised DNA was sent to Geneseek (Neogen Corporation, Auchincruive, United Kingdom) for Illumina HTS processing and HiScan chip imaging (Illumina, San Diego, United States). SNP R and Theta values were extracted using GenomeStudio Genotyping Module v2.0.2 (Illumina, San Diego, California) and allele scores were created using paRsnp (an in-house software package for clustering, visualising and comparing Illumina SNP genotyping data).

Recombination Data Analysis

Data cleaning

Genotypic data were cleaned with a custom R script which filtered out monomorphic and low-quality markers. Similarly, individuals with excessive missing or low-quality data were also removed from the analysis.

50K Recombination Analysis

The physical order of the markers and their respective physical positions on the barley genome were obtained from the current
physical assembly (MorexV2, Monat et al., 2019). Crossovers were counted as the change of allele between two consecutive markers, from either a parental allele to heterozygous or the other way around. Double recombination events were counted with a score of 2 when the allele call was switching from one parental homozygous genotype to the other. To ensure the number of crossovers was not inflated by the isolated allele switches produced by genotyping errors, the changes of alleles were only counted as valid if they were maintained in the following and previous three markers. The effect of missing data in the three-marker window on CO calling was corrected manually.

The recombination fraction ($r$) was calculated by dividing the number of crossovers between consecutive markers by the number of individuals that had data for those markers. Finally, the genetic distance between two markers was calculated by using the Kosambi formula as follows:

$$Kosambi \text{ genetic distance} \left( cm \right) = \frac{\ln \left[ 1 + 2 \ast \left( \frac{r}{2} \right) \right]}{4} \ast 100$$

Recombination differences were compared using the Wilcoxon's signed rank test (as published in Devaux et al., 1995) using the R ggpubr package.

RESULTS
Barley RTE1 (HvRTEL1) Gene Structure and Expression

A BLAST search was performed with the Arabidopsis RTE1 amino acid sequence (AT1G79950.1) to find the rice RTE1 mRNA sequence (Os01g0592900). This monocot mRNA sequence was then used in a BLAST search of our inhouse transcriptome database of meiotic anthers of barley cv. Optic and one partial cDNA sequence (TrinityTranscript16036) of 2,393 bp was found. Barley RTE1-specific primers were used in rapid amplification of cDNA ends (RACE) and reverse transcription PCR (RT-PCR) to clone the full-length cDNA of 3,370 bp (NCBI accession number: MW689197). This cDNA is comprised of 2,964 bp of open reading frame (ORF) flanked by 88 bp and 318 bp corresponding to the 5'-UTR and the 3'-UTR, respectively. Like other barley meiotic genes that we have characterised (Higgins et al., 2012; Colas et al., 2016, 2019; Barakate et al., 2021), a high GC content of up to 80% was detected at the 5' region in a GC scan of the transcript BAnTr.GP.3HG008068.1 using a 50 bp window (data not shown).

A barley genomic contig of 34,464 Kbp that contained the entire RTE1 gene was found (RTEL1.mrna1_gpv1_chr3H_317419392_317444340_1000) in the barley Golden Promise genomic database (Schreiber et al., 2020). The structure of barley RTE1 (HvRTEL1) was determined by alignment of the full-length cDNA and gene sequences (Figure 1A). HvRTEL1 contains 23 exons and 24 introns including two in the 3'-UTR. Interestingly, the HvRTEL1 gene contains two large introns in the coding sequence (Intron 16: 6,690 Kbp and Intron 19: 10,744 Kbp) and one at the 3'-UTR (Intron 23: 4,140 Kbp).

Based on transcriptomic data from 16 different tissues and developmental stages of barley (International Barley Genome Sequencing Consortium, 2012), RTE1 is most highly expressed in young inflorescences compared to other vegetative tissues (Figure 1B). Expression was also detected in the transcriptomic data of isolated barley anthers and meiocytes (Barakate et al., 2021). Although RTE1 is expressed during all meiotic stages (and in germinating embryos), the highest transcript level was detected in anthers at the pre-meiosis stage (Figure 1C). In addition, a total of eight predicted RTE1 transcript isoforms were found in this Golden Promise barley anther and meiocyte transcriptomic data. Except for the isoforms 1 and 4 that are full-length (2,964 nt coding sequence), all six transcripts were alternatively spliced. Isoform 3 resulted from a partial exclusion of exon 10, an alternative splicing (AS) event that was detected in all analysed samples of anther and meiocyte developmental stages and germinating embryos. A similar partial exclusion of exon 12 was found in the transcript 6. In isoform 2, only the first 14 introns were spliced out whilst isoform 8 retained the first 5 introns. Isoform 5 retained 111 nt of intron 17 and isoform 7 was missing 10 nt of exon 21. Consequently, an ORF was maintained of only four transcripts (1, 3, 4, and 6). With the exception of transcript 4 (BAnTr.GP.3HG008068.4), the remaining seven isoforms showed negligible expression in all samples (Supplementary Figure 1). HvRTEL1 encodes a protein of 987 amino acid residues with a predicted MW of 110.5 kDa and an overall pI of 8.78. HvRTEL1 contains ResIII (15–67 aa), DEAD_2 (99–261 aa), Helicase_C_2 (515–710 aa), and Harmonin_N_Like (891–987 aa) pfam domains as previously reported for its Arabidopsis and mammalian orthologues (Recker et al., 2014; Figure 1D). A phylogenetic tree shows clear separation of the monocotyledon and dicotyledon RTE1 sequences with the barley protein closest to those of wheat and goat grass (Supplementary Figure 2). The Plant RTE1 protein sequences used to build the phylogenetic tree are relatively well conserved. Their analysis with GLAM2 online software (Frith et al., 2008) revealed eight highly conserved motifs with the best at 480–534 aa of the protein sequence (Figure 1D). None of them overlaps with the AS event of exon 10. Analysis of natural variation in barley (Russell et al., 2016) detected three synonymous and one nonsynonymous (D750E) variant, a conserved amino acid in monocotyledons. According to cultivar Morex genome assembly version 2, barley chromosomes are divided into three zones: zone 1 (distal) enriched in rapidly evolving genes, zone 2 (interstitial) and zone 3 (proximal) enriched in housekeeping genes (Mascher et al., 2017). The RTE1 gene is on chromosome 3 (343,594,707–343,619,631 bp) close to the centromeric end of interstitial Zone 2 where recombination is low.
Suppression of Barley RTEL1 During Meiosis Affects Plant Fertility

To avoid potential negative impact of constitutive downregulation of RTEL1 on vegetative growth of barley transgenics, such as the stunting phenotype previously reported for barley ZYP1 (Barakate et al., 2014) and MSH7 (Lloyd et al., 2007) genes, the maize ubiquitin 1 promoter in the transformation vector pBRCT207 was replaced with the barley DMC1 promoter to drive the expression of RTEL1 RNAi specifically in spikes and meiocytes (Colas et al., 2019). Stunted vegetative growth phenotypes were absent for transgenics containing a GUS RNAi construct, where a fragment of E. coli β-glucuronidase (GUS) was used as a control. A total of 23 T0 independent HvRTEL1RNAi lines were recovered in barley cv. Golden Promise and transgene integration in T0 plants was confirmed by PCR using their genomic DNA and a pair of oligonucleotides specific to the selectable hygromycin marker gene. Six T0 lines were completely sterile and the number of seeds of the remaining 17 T0 lines were scored and compared to the control GUS RNAi. Whilst HvRTEL1RNAi lines showed different degrees of semi-sterility, 23 regenerated GUS RNAi lines were fully fertile producing an average of 22.4 ± 5.5 seeds per tiller (Supplementary Figure 3A). This result is in agreement with the previous study showing semi-sterility of RTEL1 T-DNA knockout in Arabidopsis (Röhrig et al., 2016). Twenty T1 seeds from each of HvRTEL1RNAi and GUS RNAi lines were germinated in the presence of hygromycin and eight resistant T1 seedlings
per line were grown to maturity and screened for reduced fertility as an indicator of a possible meiotic defect. In total, seven 
HvRTEL1RNAi lines that consistently showed reduced fertility in 
all of their corresponding T1 plants compared to the control 
were selected for further analysis (Supplementary Figure 3B).
The seeds of the HvRTEL1RNAi#8 line were very thin and did 
not germinate. The HvRTEL1RNAi#9 line was also excluded since 
50% of its plants produced very few seeds (0–28). Four lines 
(HvRTEL1RNAi#1, HvRTEL1RNAi#2, HvRTEL1RNAi#11, and 
HvRTEL1RNAi#15) were selected and fertilised using pollen from 
cultivar Bowman and their F1 seeds collected. The reciprocal 
cross was not performed as it was shown to be less successful in 
Arabidopsis rtel1 mutant (Röhrig et al., 2016). Untransformed 
barley Golden Promise was also crossed to Bowman to provide a 
(GP×Bw) control. True F1 progenies were confirmed using 
Competitive Allele-Specific PCR (KASP™) assays (Rasheed et al., 
2016) for specific SNPs on barley chromosomes 2H and 7H 
(Supplementary Table 1). Since HvRTEL1RNAi#11 was less fertile 
than the other three lines, its corresponding F1 seeds were selected 
in the presence of hygromycin to maintain the transgene and 
their F2 seeds collected. A total of 18 F1 lines were generated and 
F2 seeds of a single F1 individual HvRTEL1RNAi#11×Bw1 
(RTEL1RNAi×Bw1) were chosen to assess the effect of RTEL1 
down-regulation on meiotic recombination.

**Downregulation of RTEL1 Increases Recombination at Sub-Telomeric Regions**

A total of 96 F2 seeds were germinated for both RTEL1RNAi×Bw1 
and GP×Bw1 and their genomic DNA extracted. A preliminary 
assessment of recombination rate was performed using KASP 
assays for SNPs markers that bounded the 1H pericentromeric 
and 3H telomeric regions (Supplementary Table 1). Compared to 
GP×Bw1 control, RTEL1RNAi×Bw1 line showed a 
reduction (from 6.90 to 3.30%) and increase (from 7.70 to 14.50%) 
of recombination in 1H pericentromeric and 3H telomeric regions, 
respectively (Supplementary Table 2). This pattern of recombination 
was probed further using the barley 50k iSelect SNP Array (Bayer 
et al., 2017). From the initial 43,799 markers, 11,125 were used for 
analysis after filtering out all monomorphic and low-quality 
markers. Twelve individuals out of 192 samples were removed 
from the analysis for reasons of quality control as their data 
revealed a high number of missing data and heterozygous calls. 
This left 90 individuals per population for the genetic analysis.

**50K Polymorphism Coverage**

There was an average of 1,589 SNP loci per chromosome 
(Supplementary Table 3) with chromosome 1H having considerably 
fewer (772) compared to the rest. Following the genomic zone 
distribution described by Mascher et al. (2017), most loci were in 
the interstitial Zone 2 (61.2%), which is the largest physical zone, 
with the distal Zone 1 containing 29.1% of the SNP loci, whilst 
very few were found in the centromeric Zone 3 (3.5%). The overall 
density of markers per chromosome showed that the distal areas 
(Zone 1), despite having a smaller number of markers, had the 
densest coverage (9.1 SNP/Mbp) compared to the interstitial areas 
(Zone 2) with 2.5 SNP/Mbp, whilst zone 3 had in total 0.4 SNP/Mbp.

The positions of the first and last loci on each chromosome were verified to determine the potential missing coverage of the genetic map and very distal CO (Supplementary Table 3). The largest uncovered regions were the short arms of chromosomes 5H, 6H, and 7H with 0.33, 0.26, and 0.24% of the physical chromosome not covered, respectively, and the long arms of chromosomes 3H and 4H with the terminal 1.1 and 0.35% missing, respectively.

**Recombination Frequency**

The overall correlation of genetic and physical maps is similar in 
both GP×Bw1 and RTEL1RNAi×Bw1 populations. However, the 
control F1 cross showed higher total recombination frequency 
compared to the RTEL1RNAi×Bw1 line with a total genetic map 
of 1034.3 cM compared to 987.6 cM. This would correspond to 
20.6 and 19.7 COs in the control and RTEL1RNAi×Bw1 populations, 
respectively (assuming 50 cM/CO). However, this difference of 
46.7 cM is not significant according to Wilcoxon’s signed rank test. 
At the chromosome level, the control population showed higher 
recombination in all chromosomes except for the chromosomes 
2H and 5H where recombination was higher in RTEL1RNAi×Bw1 line.

**Recombination Distribution**

Analysis of the recombination distribution was initially carried 
out by utilising the divisions of the chromosomes into the three 
genomic zones described by Mascher et al. (2017). The three 
genomic zones were defined based on the median of 20-mr 
frequencies, Zone 1 being characterised by distal regions high in 
gene content and low-copy elements; Zone 2 by large interstitial 
regions with intermediate gene-density and high 20-mer elements; 
and Zone 3 by proximal regions, with unique conserved sequences 
and low recombination (Mascher et al., 2017). At a genome-wide 
level, across all chromosomes, RTEL1RNAi×Bw1 population showed 
higher recombination (570.3 cM) than the control (524.7 cM) in 
the distal Zone 1 although this was not significantly different. 
However, for Zone 2 the RTEL1RNAi×Bw1 population exhibited 
significantly lower (p<0.05) recombination than the control with 
totals of 416.7 and 509.6 cM, respectively. Zone 3 exhibited practically 
no recombination in either population, with the RTEL1RNAi×Bw1 
only exhibiting 0.6 cM for this region. The relative change of 
crossover distribution of the RTEL1RNAi×Bw1 population from 
Zone 2 to Zone 1 was tested with a γ-squared test by comparing 
the proportions of crossovers between the two populations for 
every zone with a contingency table (Figure 2A). Zones 1 and 
2 showed significant differences (p<0.001) in the relative change 
of distribution of crossovers between populations, as RTEL1RNAi×Bw1 
population showed a relatively higher proportion of crossovers 
in Zone 1 compared to Zone 2 (57.7/42.2%), whilst the control 
had a similar proportion of crossovers in both zones (50.6/49.4% 
respectively). At the chromosomal scale, this pattern in Zone 2 
(less recombination in the RTEL1RNAi×Bw1 line) was maintained for all chromosomes, whilst the higher recombination 
in Zone 1 of RTEL1RNAi×Bw1 was detected in all chromosomes 
except 4H and 6H (Figure 2B, Supplementary Figure 4).

The differences in recombination distribution were also 
analysed by dissecting the genome into physical intervals
of 2% of each chromosome, allowing a Wilcoxon’s signed rank test to be carried out. The distribution changes were visualised by merging all the chromosomes together dividing the genome in 50 bins (Figure 3). This analysis allowed a higher resolution of recombination distribution, showing the increase of the distal recombination in \textit{RTEL1}RNAi × Bw.1 compared to the control was located in the most distal 2% in both the short (increase of 34.8 cM, \( p < 0.05 \)) and the long arms (16.6 cM, \( p < 0.05 \)) of the chromosomes. Similarly, the recombination levels in the interstitial regions were significantly lower (17.8 cM, \( p < 0.05 \)) in the 84–86% interval in the long arms. This increase in recombination in the distal regions is not evenly shared between chromosomes (Figure 2A), with the short arms of chromosomes 2H and 6H and the long of 5H contributing the most, with increases of 10, 7.9, and 10.6 cM, respectively.

**DISCUSSION**

Meiotic recombination, which is necessary for proper chromosomal segregation, is tightly regulated by the intricate integration of programmed DSB induction, HR repair machinery and chromosomal remodelling resulting in the right proportions
of CO and NCO recombinants (Altendorfer et al., 2020). The frequency and distribution of COs are regulated by the still poorly understood phenomena of CO homeostasis and interference, respectively (Wang et al., 2015). Understanding the molecular mechanisms that limit the number and close spacing of COs is of interest not least for the potential applications in crop breeding (Taagen et al., 2020). Several DNA helicases including RTEL1 have been found to dismantle most of the D-loop joint molecules of recombination channeling their repair through the NCO pathway (Dorn and Puchta, 2019). Given its importance in genome stability and its relevance to human health (Vannier et al., 2014), RTEL1 helicase has been well studied in animals including its role in meiotic CO homeostasis and interference imposition in C. elegans (Youlds et al., 2010). Surprisingly, studies of this important helicase in plants have been limited to the vegetative stage in Arabidopsis (Recker et al., 2014; Hu et al., 2015; Röhrig et al., 2016; Olivier et al., 2018; Dorn et al., 2019; Aklilu et al., 2020). In this study, we characterised the barley RTEL1 gene structure and expression and downregulated its expression in reproductive tissues using RNAi to determine its function in meiotic CO frequency and distribution.

The barley RTEL1 helicase is encoded by a single gene that contains 24 introns including two large ones in the coding sequence and one in the 3′-UTR. We have found such large introns in other genes involved in meiosis such as FIGL1, HEI10, KU70, Met1A, and MSH2 but their significance remains to be determined. It is possible that they could relate to gene expression regulation as reported in Drosophila where large introns can promote alternative splicing (Kandul and Noor, 2009) although only 16 such large introns (>5 kbp) were found in Arabidopsis (Chang et al., 2017). Although we detected eight RTEL1 transcripts in barley anther and meiocyte transcriptome (Barakate et al., 2021), only one isoform was expressed. Interestingly, we detected several AS events of partial exon skipping (exon 10, exon 12, and exon 21) and intron retention in all samples. Such AS events were also detected for Arabidopsis RTEL1 gene (Martin et al., 2021) but remain to be confirmed by isoform specific PCR amplification and sequencing. These AS events would be of potential interest to detect and quantify at tissue level resolution to determine whether they are involved in the regulation of RTEL1 expression in plants. Indeed, AS and in particular intron retention was shown to control the timely usage of transcripts during meiosis in mice (Naro et al., 2017) and Brassica (Golicz et al., 2021). RTEL1 expression was found to be highest in spikes compared to other barley tissues and developmental stages and highest in isolated anthers at the pre-meiosis stage, concurrent with its known function in DNA replication. Its expression in isolated meiocytes indicates a potential function in meiotic recombination as previously shown in C. elegans (Youlds et al., 2010).

We generated barley transgenic plants, where RTEL1 was downregulated specifically in reproductive tissues using a previously described barley DMC1 promoter thus avoiding potential deleterious effects on vegetative growth (Colas et al., 2019). Although half of the T0 lines were completely sterile, other lines maintained semi-sterility in the T1 generation and enabled the development of a mapping population for recombination analysis. The genetic map generated for the control population GP × Bw (1034.3 cM) approximates other barley genetic maps calculated in previous studies including for GP × Morex population (Bayer et al., 2017). Although the overall number of COs and total genetic map was not significantly altered by RTEL1 downregulation, the density of marker coverage used allowed the detection of subtler changes of recombination distribution. At a genomic zone delimitation-scale (Mascher et al., 2017), RTEL1 downregulation showed significantly less recombination in the interstitial Zone 2 and non-significant but higher recombination in the distal Zone 1 compared to the control. Interestingly, when dissecting the genome into physical intervals of 2%, the increase of distal recombination under the RTEL1 downregulation was significant and concentrated mainly in the first and last 2% physical intervals of the genome. This increase of recombination was attributed mainly to chromosome 2H, 6H short arm, and 5H long arm, but given the extremely distal nature of it, it would be assumable that recombination in these regions is likely to be underestimated and highly affected by the lack of polymorphism in the very distal ends of chromosome 3H, 4H, 5H, and 6H (Supplementary Table 3). It is potentially of relevance that RTEL1 inhibits HR during replication and at telomere ends (Dorn and Puchta, 2019), which could align with this very distal increase in recombination detected when downregulating the gene.

The results suggest that the change of distribution is more likely to be driven predominantly by a shift of recombination, skewing the distribution from Zone 2 towards Zone 1 regions under the RTEL1 downregulation, rather than an absolute increase and reduction of recombination. Alterations in the timing of synapsis could play a role in this shift as this has been previously related to crossover distribution changes in barley (Higgins et al., 2012), though further studies using immunocytology would be necessary to correlate the spatiotemporal binding of RTEL1 with DSB repair and CO distribution and chromosome pairing. It will be interesting to confirm whether the newly discovered end-adjacent regions (EARs) in yeast (Subramanian et al., 2019) are also present in plants and their potential contribution to CO increase in the RTEL1 RNAi population.

The overall pattern of recombination in the barley RTEL1 RNAi line was different from that in a C. elegans knockout mutant where the CO number and distribution were strongly altered (Youlds et al., 2010). This potentially is due to differences in the mutant as well as the recombination landscape and control in the two species.

Earlier studies of RTEL1 in plants demonstrated its important role in genome stability during vegetative growth of Arabidopsis (Recker et al., 2014; Hu et al., 2015). RTEL1 was found to play essential roles during replication-associated DNA damage, stabilising repetitive elements and rDNA repeats and antagonising homologous recombination (Röhrig et al., 2016; Dorn et al., 2019; Dorn and Puchta, 2019).
This function seems to be conserved in plants as shown in *Physcomitrella patens* (Kamisugi et al., 2016; Goffová et al., 2019). Using a terminal restriction fragment assay (TRF) in *Arabidopsis*, Hu et al. (2015) concluded that RTEL1 loss does not result in telomere shortening whilst Recker et al. (2014) showed that the telomeres might even be longer in the mutants. This apparent discrepancy could be addressed using recent high-resolution methods for telomere length measurement (Lai et al., 2018; Kahl et al., 2020; Luo et al., 2020). More recently, RTEL1 was found to compensate telomere shortening in somatic cells in the absence of telomerase or replication protein A in *Arabidopsis* mutants (Olivier et al., 2018; Aklilu et al., 2020). Instead of knock-out mutants and RNAi downregulation, RTEL1 domain specific mutants could be generated using CRISPR-based gene editing or TILLING populations (Schreiber et al., 2019) helping to fully characterise the functions of RTEL1 in both somatic and meiotic phases.

It is possible that there is some redundancy in recombination control as RTEL1 can rescue yeast *srs2* (suppressor of RAD Six-screen mutant 2) mutants in the presence of methyl methanesulfonate (Frizzell et al., 2014). SRS2 is another helicase that was shown recently to be involved in meiosis by preventing late prophase DNA damage in yeast (Sasanuma et al., 2019) and so it would be interesting to study its potential RTEL1 complementation in plants. *In vitro* biochemical characterisation of the *Arabidopsis* SRS2 homolog demonstrated its 3′- to 5′-helicase activity and its involvement in processing different recombination intermediates (Blanck et al., 2009). barley SRS2 (BAnTr.GP.2HG007838, HORVU2Hr1G051300) is expressed in different tissues including anthers and meiocytes (Barakate et al., 2021; Milne et al., 2021) but its role in meiosis remains to be studied.

In conclusion, this study revealed the role of RTEL1 helicase in plant meiosis and control of recombination for the first time. The downregulation of RTEL1 expression specifically in reproductive tissues impacted on recombination distribution but not overall rates. This effect is unlike the phenotype observed in *C. elegans* but may relate to the choice of knock-down mutants in this study as much as to the radically different genome architectures of the two species.

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**DATA AVAILABILITY STATEMENT**

The SNPs data presented in the study are deposited in the European Variation Archive repository, accession number PRJEB47529 (Project) and ERZ3471485 (Analyses).

**AUTHOR CONTRIBUTIONS**

AB, CH, and RW contributed to experimental design. AB and SV contributed to RNAi construct preparation, screening and crosses of transgenic plants, and material collection. DD and JS produced the transgenic plants. MM, MA, and LR performed and analysed KASP and 50K genotyping data. MS performed expression data analysis. JO contributed to phylogenetic tree preparation. AB, MA, and RW contributed to manuscript preparation. All authors contributed to the article and approved the submitted version.

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**SUPPLEMENTARY MATERIAL**

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2021.745070/full#supplementary-material
