RESEARCH PAPER

Direct evidence for crossover and chromatid interference in meiosis of two plant hybrids (*Lolium multiflorum* × *Festuca pratensis* and *Allium cepa* × *A. roylei*)

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Abstract

Crossing over, in addition to its strictly genetic role, also performs a critical mechanical function, by bonding homologues in meiosis. Hence, it is responsible for an orderly reduction of the chromosome number. As such, it is strictly controlled in frequency and distribution. The well-known crossover control is positive crossover interference which reduces the probability of a crossover in the vicinity of an already formed crossover. A poorly studied aspect of the control is chromatid interference. Such analyses are possible in very few organisms as they require observation of all four products of a single meiosis. Here, we provide direct evidence of chromatid interference. Using *in situ* probing in two interspecific plant hybrids (*Lolium multiflorum* × *Festuca pratensis* and *Allium cepa* × *A. roylei*) during anaphase I, we demonstrate that the involvement of four chromatids in double crossovers is significantly more frequent than expected (64% versus 25%). We also provide a physical measure of the crossover interference distance, covering ~30–40% of the relative chromosome arm length, and show that the centromere acts as a barrier for crossover interference. The two arms of a chromosome appear to act as independent units in the process of crossing over. Chromatid interference has to be seriously addressed in genetic mapping approaches and further studies.

Keywords: Centromere, chromatid interference, crossover interference, homoeologous chromosome, hybrid, meiosis, recombination.

Introduction

Meiotic division is an integral part of sexual reproduction; it maintains stable chromosome numbers over generations. Chiasmata, the cytological expression of crossing over, form mechanical connections between homologous chromosomes during the first meiotic division and hence are essential elements of an orderly reduction of the chromosome number. As
such, crossovers are under strict genetic control, for both their number and distribution. This control is exercised via positive crossover interference which, in essence, has a suppressive effect on the formation of another crossover in the vicinity of an already established crossover. On the other hand, a mechanism of crossover assurance guarantees that each pair of homologues gets at least one, the so-called obligatory crossover (Jones, 1984; John, 1990; Hillers, 2004; Jones and Franklin, 2006). A single crossover between two homologues ensures their normal behaviour in meiosis. The number of potential sites of crossing over, represented by double strand breaks (DSBs), is always far greater than the number of eventual crossovers, and is presumably regulated genetically and epigenetically at the whole-genome level (Wang and Copenhaver, 2018; Modliszewski et al., 2018). Because of the tight control, the number of crossovers per bivalent varies between one and three for a majority of species, and only loosely correlates with chromosome length, offering more evidence for the regulation of crossover numbers (Mather, 1938; Fernandes et al., 2018; Otto and Payseur, 2019).

Crossover interference was observed and described already in the very first genetic mapping experiments in Drosophila melanogaster (Sturtevant, 1913; Muller, 1916) and is defined as a situation where the formation of one crossover reduces the probability of another crossover in its vicinity. Its existence can be readily verified in any genetic mapping exercise because the frequency of double crossovers between any sufficiently close loci on a chromosome is always lower than the product of frequencies of independent crossovers in these segments. The frequency of double crossovers is lower the closer the monitored intervals are to each other. As a consequence, crossover events are not randomly distributed along chromosomes (Portin, 2012; Zickler and Kleckner, 2016). Generally, the extent of crossover interference decreases with distance, understood in genetic and not physical terms. In polyploids such as wheat, genetic lengths of homoeologous chromosomes (their genetic maps) are essentially the same despite large differences in their length/DNA content, demonstrating that double crossovers, when they occur, are formed much more closely physically in shorter chromosomes than in longer chromosomes. Instances of negative crossover interference have been observed frequently in lower organisms such as fungi, but at times also in certain chromosome variants in Drosophila, barley, and maize (Auger and Sheridan, 2001; Esch and Weber, 2002). Whether negative interference exists in higher organisms with normal chromosome structure is debatable (Sill and Bengtsson, 1989). Widespread strong negative interference was invoked in genetic mapping in tetraploid wheat (Peng et al., 2000) but, with genetic maps roughly twice as long as can reasonably be expected based on chiasma frequencies and mapping in very similar populations (Lukaszewski and Curtis, 1993), marker scoring errors appear as an equally plausible explanation. Similarly, the issue of the independence of the two arms of a chromosome for crossover formation has not yet been resolved satisfactorily, and crossover interference across the centromere remains a point of dispute, with some evidence pointing in opposite directions (Laurie and Hultén, 1985; Zhao et al., 1995a; Berchowitz and Copenhaver, 2010).

The mechanism of crossover interference remains unclear, but probably involves some crossover-discouraging signal or substance that spreads along the chromosome arm. The mechanical stress model of crossover interference is based on the presumption that the stress drives formation of a crossover, but in doing so the local area is then relieved and there is not enough stress to drive formation of a second crossover nearby (Börner et al., 2004; Kleckner et al., 2004). The polymerization model assumes that once the crossover structure is attached along the synaptonemal complex, it has the same chance per unit time to initiate a bidirectional polymerization event. The structures responsible for it are presumably the late recombination nodules which can be observed in pachytene (King and Mortimer, 1990). Crossover interference clearly involves the synaptonemal complex as it is not transmitted via synaptonemal complex discontinuities, such as synaptic partner exchange points in translocation heterozygotes, or across separated centromeres in double ditelocentric lines (reviewed in Dawe, 1998). Interestingly, there appear to be two pathways leading to crossovers, one interference sensitive and the second interference insensitive (Osman et al., 2011; Mercier et al., 2005). In plants, the former is more abundant and accounts for ~80–85% of total crossovers (Berchowitz et al., 2007).

A bivalent in the first meiotic division is composed of two pairs of sister chromatids. Additional crossovers in a chromosome (or a chromosome arm) bring up the issue of chromatid choice for each event. It is generally accepted that for any given crossover, the chromatid choice is random, and hence a crossover involving two non-sister chromatids does not affect the choice of non-sister chromatids for another crossover in the same chromosome (Zhao et al., 1995a). This is, however, an assumption with very limited experimental support, particularly in higher organisms. Direct tests of chromatid interference require analyses of all four products of any given meiosis. This condition is met only in a very few instances, all in lower organisms, such as Aspergillus spp. where chromatid interference does not appear to operate (Whitehouse, 1958). In higher organisms, re-analysis of experimental data from several organisms (Zhao et al., 1995b) and mathematical modelling of genetic mapping data suggest that chromatid interference may exist (Teuscher et al., 2000), but no direct observation has been possible thus far.

Crossover and chromatid interference can be analysed in several different ways. Generally, the former can be studied directly, and copious raw data for such analyses are generated in each genetic mapping effort, especially when mapped loci can be directly placed on the DNA sequence assemblies. Chromatid interference, on the other hand, in a great majority...
of organisms can be studied only indirectly, such as by mathematical modelling (Teuscher et al., 2000) mentioned above. However, recent technological advances offer a chance to study both phenomena directly by cytological observation in meiosis. One such approach is to visualize sites of crossovers in meiotic cells with fluorescently labelled probes, for example for the MLH1 mismatch repair protein (Barlow and Hultén, 1998; Phillips et al., 2013; Anderson, 2014). This approach, however, may lack the resolution of individual chromatids. Another way is the analysis of the anaphase I chromosomes in hybrids where the DNA sequence divergence between the parental genomes permits chromosome painting, hence direct visualization of the crossover points. Thus, it is possible not only to score the overall level and distribution of crossovers in individual chromosome arms, chromosomes, and entire genomes, but also to study their distribution among chromatids. Another advantage of this approach is that the results are not biased by potential gametophytic selection, which may distort the ratios of recombinant and parental chromosomes transmitted to the progeny. Its weakness is in the selection of parents: these need to be sufficiently different at the DNA sequence level to enable unambiguous discrimination of homologues by chromosome painting, and yet sufficiently close genetically to provide for regular chromosome pairing and disjunction. Such hybrids do exist among plant species. Here we focus on the F1 hybrids between representatives of Lolium and Festuca, known for a high (87–97%) metaphase I chromosome pairing and crossing over (Jauhar, 1975; Kopecký et al., 2008). Scoring crossovers in anaphase I, where individual chromatids are clearly visible, offers direct evidence not only for crossover interference but also for chromatid interference and possible effects across the centromere. Cytogenetic stocks from our past experiments (Kopecký et al., 2008) made it possible to analyse the physical attributes of crossing over in individual chromosomes and their arms, while high-density chromosome genetic maps (Kopecký et al., 2010) made it possible to relate these observations to the DNA sequence across the entire genome. Similar observations were made on wide hybrids in the genus Allium (A. cepa×A. roylei) displaying frequent pairing and recombination between homoeologous chromosomes (Khrustaleva and Kik, 2000).

Materials and methods

Plant material and chromosome preparations

The frequency of possible types of crossover configurations during meiosis was evaluated in diploid F1 hybrids of Festuca pratensis×Lolium multiflorum. Chromosome substitution lines of F. pratensis/L. multiflorum hybrids were used to estimate the frequency and distribution of double crossovers for all seven F. pratensis chromosomes individually. The tetraploid monosomic substitution lines (2n=4x=28; 27L+1F) were developed in previous studies (Kopecký et al., 2008, 2010). Plants with the same substituted F. pratensis chromosome were intercrossed and their progenies were germinated and individually analysed. Similarly, the F1 hybrids of A. cepa×A. roylei (Scholten et al., 2016) were intercrossed and their progenies were individually screened by genomic in situ hybridization.

Individual anthers confirmed to be in meiotic anaphase I stage and root tips were fixed in Carnoy’s solution (absolute ethanol/glacial acetic acid, 3:1 v/v) at 37 °C for 7 d and microscope preparations were made according to Masoudi-Nejad et al. (2002).

Genomic in situ hybridization

Genomic in situ hybridization analyses were done on mitotic and meiotic chromosome spreads according to Kopecký et al. (2008). Total genomic DNA (gDNA) of L. multiflorum and A. cepa was used as blocking DNA and total gDNAs of F. pratensis and A. roylei were labelled with digoxigenin (DIG) using the DIG-Nick Translation Kit (Roche Applied Science, USA) according to the manufacturer’s instructions, and used as probes for grass and onion hybrids, respectively. The probe/blocking DNA ratio was ~1:150. Signal detection was made with anti-DIG–fluorescein isothiocyanate (FITC) conjugate (Roche Applied Science). Chromosomes were counterstained with DAPI in Vectashield (Vector Laboratories, Oberkochen, Germany). Chromosome analysis was done under an Olympus AX70 microscope equipped with epifluorescence and a SensiCam B/W camera. Images were captured with Micro Image and processed with Adobe Photoshop v.6 software.

Scoring and statistical treatment

All statistical analyses were done in R 3.4.3 (R Core Team, 2019). For meiotic analyses of diploid F1 hybrids of L. multiflorum×F. pratensis, we evaluated exclusively the anaphase I stage. Each cell has seven pairs of chromosomes. Each arm was evaluated separately. Each pair of homologous arms displayed either no crossover or one of the possible crossover exchanges: single crossover or double crossover with two, three, or four chromatids involved. Theoretically, with no chromatid interference, the proportion of the three crossover types should be 1:2:1. We statistically treated the observed frequencies for this ratio using the multinomial test from the ‘stats’ library (R Core Team). Proportions of different numbers of crossovers on one arm between chromosomes with one and two crossovers on the other arm were compared using the two-sided Fisher’s exact test. For calculation, the function Fisher.test with P-values computed by Monte Carlo simulations based on 2000 replicates from the ‘stats’ library was used.

To determine the positions of the crossover events among the progenies of onion and grass hybrids, we measured the lengths of the introgressed segments and the lengths of both arms of recombinant chromosomes using ScionImage software. Calculations of the distances (in megabases) between two crossovers on one arm were done based on the length of individual F. pratensis chromosomes (Kopecký et al., 2010). The difference in distribution of different crossover types was evaluated by comparing their distributions along chromosome arms divided into 10 segments (bins) of 10% of their length. Two empirical distributions were compared using the function ks.boot in the ‘Matching’ library in R. The function uses a bootstrap version of the Kolmogorov–Smirnov test, providing accurate coverage even when the distributions being compared are not entirely continuous and ties occur in the dataset (Sekhon, 2011).

Results

Frequency of crossovers in anaphase I configurations in Lolium×Festuca hybrids

Using in situ probing with labelled total gDNA, we monitored crossover events involving individual chromatids in each chromosome of 96 meiocytes of L. multiflorum×F. pratensis (2n=2x=14; seven L. multiflorum chromosomes+seven...
in the progeny as single events and, thus, are placed in the category of single crossovers.

Out of 629 and 328 recombined chromosomes of grasses and onions, respectively, the majority demonstrated single crossovers (63.8% and 65.8%), followed by two crossovers, one on each arm (23.2% and 21.7%) and double crossovers, sometimes with single or double crossover on the other arm (13.0% and 13.1%), respectively. The proportions of chromosomes in each of the three categories varied for individual Festuca chromosomes (Table 1).

Using progenies of single chromosome substitution lines, it was possible not only to estimate the frequencies of double crossovers for each of the Festuca chromosomes, but also to estimate the mean distances in megabases between adjacent crossovers in each chromosome (Fig. 3; Table 2). The frequencies of double crossovers differed markedly between the short and long arms. In the shortest arm of the F. pratensis karyotype, that of chromosome 5, we did not observe any products of double crossover (involving two or three chromatids). With the obligatory chiasma and the positive crossover interference, this is a typical distribution as affected by the chromosome arm length. It was clearly evident in wheat during the genetic mapping of the physical attributes of chromosomes (Lukaszewski and Curtis, 1993), and was confirmed by ultra-high-resolution analysis (Jordan et al., 2018).

On average, two adjacent crossover events on one arm of a single chromatid were separated by ~97 Mb. However, this distance showed considerable variation from one chromosome arm to another. The shortest distance between two adjacent crossovers in an arm was ~26 Mb. There was no correlation between chromosome/chromosome arm length (CL), the frequency of double crossover (FdCO), and the average distance (AD) between two crossovers on one arm (double crossovers involving two or three chromatids, but not four) as revealed by Spearman correlation coefficient (for arms, n=14: CL versus FdCO r=0.42, P=0.149; CL versus AD r=0.19, P=0.535; FdCO versus AD r=−0.15, P=0.635; for chromosomes, n=7, CL versus FdCO r=−0.43, P=0.333; CL versus AD r=−0.29, P=0.535; FdCO versus AD r=−0.20, P=0.670).

The distribution of single and double crossovers differed in both grass and onion hybrids (Fig. 4). The distribution of single crossovers in both hybrids showed similar patterns, with the highest frequency in the intervals from 20% to 40% of the arm length from the telomeres, and with a significant decline in frequencies nearing the centromeres (proximal 30% of the arm). Relatively low frequencies were also evident in the most distal 10% of the arm (next to telomeres). However, this may be an artefact of the method used, as short terminal segments from the non-probe parent may be undetectable by in situ probing with labelled DNA (Lukaszewski et al., 2005). On the other hand, in grasses, double crossovers had the highest frequency in the intervals of 10–20% (the first crossover) and...
Fig. 1. Crossovers in grasses and onions visualized by genomic in situ hybridization. Various types of crossovers can be seen directly during anaphase I in F₁ hybrids of *Lolium multiflorum* × *Festuca pratensis* (A–D), in the mitotic cell of the progeny of *F. pratensis*/*L. multiflorum* single chromosome substitution lines (E), and in the mitotic cell of the progeny of *F.* *Allium cepa* × *A. roylei* hybrids (F). Insets provided in the upper right figure (B) are enlargements of the pairing partners in the metaphase I plate (A) with the description of crossover type. Similarly, white lines indicate pairing partners in the duplicated figure (D) of the metaphase plate (C). Total gDNA of *F. pratensis* and *A. roylei* was labelled with digoxigenin (green/yellow colour) and sheared DNA of *L. multiflorum* and *A. cepa* was used as blocking DNA (red pseudocolour).
40–50% (the second crossover) of the arm measured from telomeres. This indicates that crossover interference in grasses acts up to a distance of ~30% of the average arm length. A similar crossover distribution was observed in onions, even though no clear frequency peaks were evident. Once we calculated the frequencies of the theoretical and actual appearance of double crossovers for all combinations of intervals (an interval equalled 10% of the physical arm length), the positive crossover interference was observed between intervals 0–10%, 10–20%, 10–20% and 20–30%, 10–20% and 30–40%, 20–30% and 30–40%, 20–30% and 40–50%, 20–30% and 50–60%, 30–40% and 40–50%, and between 30–40% and 50–60% (data not shown). On average, two crossovers in one arm were separated by 36% and 41% of the arm length (ranging from 13% to 78% and from 9% to 83%) in onions and grasses, respectively (Fig. 4). This supports the indication that in both hybrids, crossover interference acts at a physical distance of 30–40% of an average chromosome arm. This does rule out the possibility that the crossover interference is governed by mechanisms modulated by the absolute physical distance between two crossovers.

In general, the average frequency of chromosomes with double crossovers in grasses was 13% (Table 1). The highest frequency of such events was ~20% for chromosome 2, which also had the lowest average distance between two adjacent crossovers (77 Mb) (Table 2), and the lowest was 8% for chromosome 4, which is the longest chromosome in the genome (Kopecký et al., 2010) and seems to have strong crossover interference (average length

Table 1. Numbers or chromosomes with and without homoeologous crossovers and frequencies (in parentheses) of different crossover types among the progeny of single chromosome substitution lines of F. pratensis into L. multiflorum (individually for each chromosome)

| Fp/Lm chromosome | No crossover | Single crossover (%) | Two crossovers, one on each arm (%) | Double crossover (%) |
|------------------|--------------|----------------------|------------------------------------|---------------------|
| 1                | 5            | 63 (63)              | 21 (21)                            | 16 (16)             |
| 2                | 7            | 59 (57)              | 24 (23)                            | 21 (20)             |
| 3                | 4            | 58 (62)              | 25 (27)                            | 10 (11)             |
| 4                | 6            | 56 (70)              | 18 (23)                            | 6 (8)               |
| 5                | 3            | 62 (78)              | 10 (13)                            | 7 (9)               |
| 6                | 2            | 26 (40)              | 27 (42)                            | 12 (18)             |
| 7                | 6            | 67 (77)              | 10 (11)                            | 10 (11)             |
| Average          | (64)         | (23)                 | (13)                               |                     |
between two events spanning a double crossover was 112 Mb) (Table 2). On the other hand, chromosome 1 is the shortest in the *F. pratensis/L. multiflorum* genomes (Kopecký et al., 2010) but it showed the highest average distance between adjacent crossovers. This suggests that the interference distance in the system studied here may be chromosome specific and not genome wide.
There was a significant difference in the distribution of single crossovers along the chromosome arms and of the first crossover when two crossovers were present on an arm, both in grasses (Kolmogorov–Smirnov test, $D=0.542$, bootstrap $P<0.001$) and in onions ($D=0.455$, bootstrap $P<0.001$) (Fig. 4). When two crossovers were present, the first crossovers were much more distal, with a peak at 10–20% of the arm length, and dropped rapidly towards the centromere. In grasses, the first crossovers of double crossover arms were limited to the distal half of the arms. A similar distribution
Table 3. Average positions of crossovers based on their abundance (in % of the arm length, measured from the telomere)

| Type of crossover                                      | Average position of crossover (% of arm from telomere) |
|--------------------------------------------------------|--------------------------------------------------------|
|                                                        | Onions | Grasses |
| Single crossover                                       | 41.32  | 43.5    |
| Two crossovers (one on each arm)                       | 43.85  | 40.6    |
| Single crossover with double crossover on the other arm | 46.49  | 38.6    |

Table 4. The relationship between the number of crossovers in one chromosome arm and the number of crossovers in the other arm

| No. of crossovers on one arm | Grasses | Onions |
|-----------------------------|---------|--------|
|                            | Number of crossovers on the other arm |
|                            | 0       | 1      | 2      | 3      |
| 0                           | 308     | 133    | 0      |        |
| 1                           | 308     | 66     | 26     | 0      |
| 2                           | 133     | 26     | 6      | 0      |
| 3                           | 0       | 0      | 0      | 0      |

was observed in onions, with a gradual decrease from the telomere toward the centromere. It is evident that second crossovers on an arm are possible only when the first crossover is sufficiently distal.

Crossover interference does not act across the centromere

To test if crossover interference acts across the centromere, or if the two arms of a chromosome are independent for crossover formation, we performed three comparisons between the expected values and the experimental data. If the interference acts across the centromere, the following should apply. (i) The distribution of single crossovers and of two crossovers (each on one of the arms) should differ, with the latter located more distally towards the telomeres. Indeed, we observed that the distribution of single crossovers per chromosome and two crossovers per chromosome with one at each arm differed significantly in grasses (Kolmogorov–Smirnov test, D=0.101, bootstrap P=0.002), but not in onions (D=0.079, bootstrap P=0.334). In grasses, however, single crossovers were distributed primarily in the interstitial parts of the chromosome arms, while two crossovers were more spread over the arms and were localized relatively frequently outside the interstitial regions. In onions, single crossovers and two crossovers were spread over the arms, except for a rapid decline in the centromeric and pericentromeric regions. (ii) When a double crossover is formed on one arm, the distribution of crossovers on the other arm should be more distal. Although there were not enough such cases for statistical tests, the average positions of crossovers were similar between a single crossover, two crossovers (one on each arm), and a single crossover with a double crossover on the other arm (Table 3). (iii) With two crossovers in one arm, there should be fewer crossovers in the other arm as compared with a single crossover in the first arm. No difference was observed in the distribution of crossovers in the second arm when comparing one and two crossovers in the first arm in both grasses (Fisher’s exact test, P=0.415) and onions (Fisher’s exact test, P=0.754) (Table 4). These three tests indicate that in grasses and in onions, crossover interference does not act across the centromere; two arms of a single chromosome appear to be independent for establishment of the crossover.

Discussion

This study was performed on wide hybrids and so observations and results may not be fully representative of strictly homologous chromosome pairing and recombination. Wide hybrids made direct observations possible and offered a unique chance of studying the immediate effects of crossing over as early as anaphase I, but at the same time they might have skewed the results in some unpredictable direction. However, these hybrids show surprisingly high regularity of meiotic chromosome pairing and segregation (Jauhar, 1975; Kopecky et al., 2008), suggesting that the results may be of wider significance. On the other hand, there is a chance that the distribution of crossovers, as judged by the exchange points in recombined chromosomes, may be wider than what could be expected based on the chiasma distribution in the parental species (Karp and Jones, 1983). It also must be pointed out that data on the physical distribution of crossovers along chromosomes and chromosome arms do carry an inherent error. There are differences in genome sizes of the parental species of both hybrids, and these translate into differences in chromosome lengths. Whether these differences are evenly distributed along all chromosomes is very far from clear. For that reason, most of the analyses are based on relative length values.

Chromatid interference

The ability to distinguish parental chromatin by chromosome painting allowed us to analyse the frequency and distribution
of crossovers directly during the meiotic division, immediately after bivalent separation, and ensured that all products of every individual meiosis could be scored (Fig. 1). This avoided any possible bias associated with uneven chromosome transmission and gametic or zygotic selection. These direct observations were supplemented with earlier observations of the structure of chromosomes among the progeny.

Ever since Barbara McClintock’s demonstration of the relationship between chiasmata and crossing over (Creighton and McClintock, 1931), chiasmata alone indicated that crossovers were not distributed evenly along chromosomes. This has been demonstrated in many organisms and in several different ways (reviewed in Otto and Payseur, 2019). The pattern of crossovers is affected by genetic and presumably epigenetic systems, with a general preference for distal location, and perhaps by the chromatin structure, which makes some parts of the chromosomes inaccessible to crossing over, regardless of the position on the telomere–centromere axis (Lukaszewski, 2008; Lukaszewski et al., 2012; Higgins et al., 2014). Within this general pattern, if more crossovers are formed in an arm, their frequencies and distribution tend to form a sinusoid pattern, reflecting the presence and strength of the positive crossover interference (Muller, 1916; Mather, 1938).

Of the two components of genetic interference, crossover interference can be easily scored in any genetic mapping experiment. The second component, chromatid interference, is far more difficult to observe. It informs if the chromatids involved in the first crossover affect the choice of chromatids for a second crossover on the same chromosome. Chromatid interference can only be reliably and directly scored when all four products of a single meiosis are available, and this happens infrequently. Consequently, the current knowledge of chromatid interference stems from statistical models, and the general assumption is that chromatid interference does not exist. Geneticists generally assume its absence while studying recombination and creating genetic maps (Strickland, 1958; Broman and Weber, 2000; Broman et al., 2002).

Early studies on Neurospora crassa, Saccharomyces cerevisiae, and Aspergillus nidulans indicated that the choice of chromatids for second crossovers may not be random (Lindegren and Lindegren, 1942; Hawthorne and Mortimer, 1960). Similarly, the model of Teuscher et al. (2000), based on data from a study on mouse and two studies on Drosophila, suggested that chromatid interference may play an important role in meiosis. Similarly, weak chromatid interference was deduced in maize, human oocytes, and Caenorhabditis elegans (Hou et al., 2013; Li et al., 2015; Li et al., 2018). Our study offers clear and unambiguous evidence for a strong positive chromatid interference: the second crossover in an arm was formed more frequently between chromatids not involved in the first crossover (Fig. 2). In other words, all four chromatids of two homoeologous chromosomes were involved in double crossovers much more frequently than expected (64% versus 25%) and the most abundant class expected for random chromatid selection, that with three chromatids involved, was seriously under-represented, accounting for only 21% of cases versus 50% of those theoretically expected. Thus, our results provide strong support for the model of Teuscher (2000) and deliver interesting results for further studies on crossover formation and recombination in plants. They also point out a need to consider chromatid interference in genetic mapping. We envisage that newly developed visual assays for yeasts and the model plant Arabidopsis (Berchowitz and Copenhaver, 2008), which enable scoring all four products of meiosis, could increase our understanding of chromatid interference in the near future.

The molecular mechanism underlying chromatid interference is unknown. However, a study on C. elegans suggests that an orthologue of the breast cancer susceptibility gene 1 (BRCA-1) may be involved in this pathway. Numbers of double crossovers with two chromatids involved were reduced and numbers of single crossovers were elevated in the brc-1; zim-1 mutant compared with the zim-1 single mutant. This indicates that BRCA-1 may counteract chromatid interference under meiotic dysfunction, such that more crossovers with the same two chromatids involved occur (Li et al., 2018). Some studies also pointed to the role of Tel1, a protein kinase that responds to DNA damage, in both crossover and chromatid interference (Zhang et al., 2011; Garcia et al., 2015; Cooper et al., 2016). According to Fowler et al. (2018), Tel1 is activated when the formation of DSBs occurs in a clustering hotspot, promoting the phosphorylation and inactivation of some components of the DSB-forming complex, which prevents the formation of a second DSB in the vicinity. In the absence of Tel1, multiple breaks are formed, increasing the frequency of crossovers. The clustering of DSB hotspots can involve each homologous chromosome separately (two chromatids) or both together (four chromatids), and in species where clustering involves the homologues pair the interference is stronger (Smith and Nambiar, 2020). As such, cluster gathering of four chromatids may explain the chromatid interference found in our study.

Crossover interference

Apart from a few exceptions, such as A. nidulans and S. pombe discussed above, crossover interference is a widespread phenomenon in eukaryotes (Strickland, 1958; Snow, 1979), but its strength and spatial efficiency differ among species. In roundworm C. elegans, the absence of double crossovers may be taken as an indication of complete crossover interference (Meneely et al., 2002). Similarly, crossover interference appears to be extremely strong in mouse (Broman et al., 2002). In other species, it may be strong in short intervals: it is almost complete in D. melanogaster and very strong in Neurospora crassa (Foss et al., 1993).

Similar to genetic studies based on screening of progeny to assess the crossover interference, our approach did not allow
incorporation of a double crossover where four and three chromatids were involved. Thus, it provides detailed characterization of crossover interference arising exclusively from double crossovers involving two chromatids. In grasses and onions, we never observed double crossovers at distances shorter than 9% and 12% of the arm length, respectively. Using genome-wide numbers, this translates to ~37 Mb and 241 Mb (Labani and Elkington, 1987; Doležel et al., 1992; Kopecký et al., 2010). As the distance from the initial crossover increases, the interference strength weakens and chances for a second crossover increase. Crossover interference is effective up to 15–20 Mb in Drosophila, corresponding to about one half of a chromosome arm, and up to 0.9 Mb in Neurospora, corresponding to ~30% of an average arm length (Foss et al., 1993). Similarly, Lawrie et al. (1995) observed crossover interference extending over ~68% and 77% of a chromosome length in mouse autosomes 5 and 15, corresponding to about one-third of the arms. Complete crossover interference over distances equalling 25–30% of a chromosome arm has been confirmed in Chorthippus borreus (Jones, 1987). In budding yeast, Malkova et al. (2004) detected interference over distances of about a quarter of the total length of a chromosome arm.

The examples listed above indicate that the effectiveness of crossover interference is surprisingly similar among the species tested, and it extends up to about one-third of a chromosome arm. Even though we worked with wide hybrids, the results follow the same pattern: in both cases, crossover interference acts efficiently up to ~30–40% of the chromosome arms (Fig. 3). The same range was observed in the B genome of wheat (Łukaszewski and Curtis, 1993). Interestingly, it appears that it is the relative distance along the chromosome arm and not the physical or genetic distance that matters. The two hybrids used in our study differ significantly in their genome sizes and lengths of chromosomes in megabases: an average chromosome in onions is ~2 Gb in length, while it is ~0.4 Gb in grasses (Doležel et al., 1992; Kopecký et al., 2010), corresponding to a physical crossover interference distance of ~350 Mb and 100 Mb, respectively (Table 2). Mean interchiasma distances in mouse and S. cerevisiae chromosomes gave the impression of being correlated with chromosome lengths so that the mean interchiasma distance was greater in longer chromosomes (Lawrie et al., 1995). The possible mechanism explaining this phenomenon involving a conformational chain reaction leading to the allosteric blocks of recombination in neighbouring regions in a time-dependent manner has been described by Kaback et al. (1999). This indicates that the processes underlying crossover interference may be evolutionarily conserved. We should note that despite the general mechanism of crossover interference acting at the whole-genome level, there are exceptions where crossover interference has variable intensities and distributions in different regions of a chromosome, such as observed in budding yeast and in Populus euphratica (Malkova et al., 2004; Wang et al., 2019).

As in other cases studied earlier (see John, 1990), we observed that second crossovers on an arm are predominantly formed only when the first crossovers are sufficiently distal (Fig. 4). This may be related to the progress of synopsis, from the telomere toward the centromere, with the distal crossover (presumably formed first) leaving enough room on the arm for crossover interference strength to drop before the end of the recombination process, hence increasing the opportunity for an additional crossover to form. The role of the centromere as a barrier for crossover interference remains unclarified (reviewed in Nambiar et al., 2019). The question is: are the two arms of a chromosome independent in crossover formation or can a crossover on one arm interfere with crossovers on the other arm? The current evidence is unclear. No positive interference was observed across the centromere in N. crassa (Strickland, 1961; Perkins, 1962). On the other hand, in the grasshopper, Colombo and Jones (1997) provided evidence that interference did act across the centromere, and the level of interference in the region spanning centromeres appeared to be no different from that seen in any other chromosome segment. Similarly, the chiasmata numbers on two arms of a single chromosome do not appear to be independent in humans (Broman and Weber, 2000) or in species such as Culex and Pnauen (Callan and Montalenti, 1947; Harte, 1956). In this study, we did not observe any significant effect of crossovers in one arm on the crossover frequency and distribution in the other arm, implying that in our material crossover interference may not extend across the centromere (Tables 3, 4). However, crossover distribution might have also played a role here: with predominantly distal crossovers, the distance across the centromere in most cases would be greater than the interference distance.

In conclusion, our study clearly illustrates that chromatid interference does operate in higher Eukaryotes, along with crossover interference. Both play an important role in meiotic division, the latter acting over a distance corresponding to 30–40% of the physical length of a chromosome arm. In materials studied here, the centromere appears to act as an effective barrier for crossover interference and thus the two arms of a chromosome seem to act as independent units for crossovers. We are aware that our study is based on observations of crossovers between homoeologous chromosomes in wide hybrids and these may not completely correspond to interactions between perfectly homologous chromosomes. However, meiotic chromosome pairing in these hybrids is essentially normal (for illustrations of meiosis in the Lolium×Festuca hybrid see https://olomouc.uib.cas.cz/getattachment/Research-groups/Kopecky-group/Meiosis-With-Labeled-Parental-Genomes.pdf.aspx?lang=en-US). Moreover, all crossovers must meet certain DNA criteria, both for the substrate length and for the level of homology (Shen and Huang, 1986, 1989; Datta et al., 1997) and there is no reason to suspect that the crossovers observed here
violate them. Given the recent progress of technology, we envisage that it will be possible to monitor both homologous and homoeologous crossovers in the near future using tetrad analysis based on the *quartet1* (*qrt1*) mutation, haplotype-specific chromosome painting as shown in maize, or the DeepTetrad technology currently available only for *Arabidopsis thaliana* (Copenhaver et al., 2000; Martins et al., 2019; Lim et al., 2020).

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**Author contributions**

Conceptualization: AJL, DK; data curation: MTMF, MG, AJL, DK; formal analysis: MG, MD, DK; funding acquisition: JD, DK; investigation: MTMF, KP, MK, AJL, DK; methodology: AJL, DK; project administration: DK; resources: OS, VHT, AJL, DK; supervision: VHT, JD, AJL, DK; validation: AJL, DK; visualization: MTMF, MG, KP, AJL, DK; writing—original draft preparation: MTMF, AJL, DK; writing—review and editing: OS, VHT, JD, AJL, DK.

**Data availability**

All data supporting the findings of this study are available within the paper.

**References**

Anderson LK, Lohmiller LD, Tang X, Hammond DB, Javernick L, Shearer L, Basu-Roy S, Martin OC, Falque M. 2014. Combined fluorescent and electron microscopic imaging unveils the specific properties of two classes of meiotic crossovers. Proceedings of the National Academy of Sciences, USA 111, 13415–13420.

Auger DL, Sheridan WF. 2001. Negative crossover interference in maize translocation heterozygotes. Genetics 159, 1717–1726.

Barlow AL, Hultén MA. 1998. Crossing over analysis at pachytene in man. European Journal of Human Genetics 6, 350–358.

Berchowitz LE, Copenhaver GP. 2008. Fluorescent *Arabidopsis* tetrads: a visual assay for quickly developing large crossover and crossover interference data sets. Nature Protocols 3, 41–50.

Berchowitz LE, Copenhaver GP. 2010. Genetic interference: don’t stand so close to me. Current Genomics 11, 91–102.

Berchowitz LE, Francis KE, Bey AL, Copenhaver GP. 2007. The role of *AIMUS81* in interference-insensitive crossovers in *A. thaliana*. PLoS Genetics 3, e132.

Börner GV, Kleckner N, Hunter N. 2004. Crossover/noncrossover differentiation, synaptonemal complex formation, and regulatory surveillance at the leptotene/zygotene transition of meiosis. Cell 117, 29–45.

Broman KW, Rowe LB, Churchill GA, Paigen K. 2002. Crossover interference in the mouse. Genetics 160, 1123–1131.

Broman KW, Weber JL. 2000. Characterization of human crossover interference. American Journal of Human Genetics 66, 1911–1926.

Callan HG, Montalenti G. 1947. Chiasma interference in mosquitofish. Journal of Genetics 48, 119–134.

Colombo PC, Jones GH. 1997. Chiasma interference is blind to centromeres. Heredity 79, 214–227.

Cooper TJ, Garcia V, Neale MJ. 2016. Meiotic DSB patterning: a multifaceted process. Cell Cycle 15, 13–21.

Copenhaver GP, Keith KC, Preuss D. 2000. Tetrad analysis in higher plants. A budding technology. Plant Physiology 124, 7–16.

Creighton HB, McClintock B. 1931. A correlation of cytological and genetic crossing-over in *Zea mays*. Proceedings of the National Academy of Sciences, USA 17, 492–497.

Datta A, Hendrix M, Lipsitch M, Jinks-Robertson S. 1997. Dual roles for DNA sequence identity and the mismatch repair system in the regulation of mitotic crossing-over in yeast. Proceedings of the National Academy of Sciences, USA 94, 9757–9762.

Daw RK. 1998. Meiotic chromosome organization and segregation in plants. Annual Review of Plant Physiology and Plant Molecular Biology 49, 371–395.

Dolezel J, Sgròbati S, Lucretti S. 1992. Comparison of three DNA fluorochromes for flow cytometric estimation of nuclear DNA content in plants. Physiological Plantarum 85, 625–631.

Escar E, Weber W. 2002. Investigation of crossover interference in barley (*Hordeum vulgare* L.) using the coefficient of coincidence. Theoretical and Applied Genetics 104, 786–796.

Fernandes JB, Séguela-Arnaud M, Larchevêque C, Lloyd AH, Mercier R. 2018. Unleashing meiotic crossovers in hybrid plants: Proceedings of the National Academy of Sciences, USA 115, 2431–2436.

Foss E, Lande R, Stahl FW, Steinberg CM. 1993. Chiasma interference as a function of genetic distance. Genetics 133, 681–691.

Fowler KR, Hyppa RW, Cromie GA, Smith GR. 2018. Physical basis for long-distance communication along meiotic chromosomes. Proceedings of the National Academy of Sciences, USA 115, E9333–E9342.

Garcia V, Gray S, Allison RM, Cooper TJ, Neale MJ. 2015. Tel1(AtM)-mediated interference suppresses clustered meiotic double-strand-break formation. Nature 520, 114–118.

Harte C. 1956. Die Variabilität der Chiasmenbildung bei *Paeonia tenuifolia*. Chromosoma 8, 152–182.

Hawthorne DC, Mortimer RK. 1960. Chromosome mapping in *Saccharomyces*: centromere-linked genes. Genetics 45, 1085–1110.

Higgins JD, Osman K, Jones GH, Franklin FC. 2014. Factors underlying restricted crossover localization in barley meiosis. Annual Review of Genetics 48, 29–47.

Hillers KJ. 2004. Crossover interference. Current Biology 14, R1036–R1037.

Hou Y, Fan W, Yan L, et al. 2013. Genome analyses of single human oocytes. Cell 155, 1492–1506.

Jauhar PP. 1975. Chromosome relationships between *Festuca* and *Paeonia tenuifolia*. Cell 126, 293–320.

Jones G. 1987. Chiasmata. In: Moens PB, ed. Meiosis. Orlando: Academic Press, 213–244.

Jones GH. 1984. The control of chiasma distribution. Symposia of the Society for Experimental Biology 38, 293–320.

Jones GH, Franklin FC. 2006. Meiotic crossing-over: obligation and interference. Cell 126, 246–248.
Jordan KW, Wang S, He F, et al. 2018. The genetic architecture of genome-wide recombination rate variation in allotetraploid wheat revealed by nested association mapping. The Plant Journal 95, 1039–1054.

Kaback DB, Barber D, Mahon J, Lamb J, You J. 1999. Chromosome size-dependent control of meiotic reciprocal recombination in Saccharomyces cerevisiae: the role of crossover interference. Genetics 152, 1475–1486.

Karp A, Jones RN. 1983. Cytogenetics of Lolium perenne. Part 2. Chiasma distribution in inbred lines. Theoretical and Applied Genetics 64, 137–145.

Khristovaeva LI, Kik C. 2000. Intergression of Allium fistulosum into A. cepa mediated by A. roylei. Theoretical and Applied Genetics 100, 17–26.

King JS, Mortimer RK. 1990. A polymerization model of chiasma interference and corresponding computer simulation. Genetics 126, 1127–1138.

Kleckner N, Zickler D, Jones GH, Dekker J, Padmore R, Henle J, Hutchinson J. 2004. A mechanical basis for chromosome function. Proceedings of the National Academy of Sciences, USA 101, 12592–12597.

Kopecký D, Havránková M, Loureiro J, Castro S, Lukaszewski AJ, Bartos J, Kopecká J, Doelez J. 2010. Physical distribution of homoeologous recombination in individual chromosomes of Festuca pratensis in Lolium multiflorum. Cytogenetic and Genome Research 129, 162–172.

Kopecký D, Lukaszewski AJ, Doelez J. 2008. Meiotic behaviour of individual chromosomes of Festuca pratensis in tetraploid Lolium multiflorum. Chromosome Research 16, 987–998.

Labani R, Elkington R. 1987. Nuclear DNA variation in the genus Allium L. (Liliaceae). Heredity 59, 119–128.

Laurie DA, Hultén MA. 1985. Further studies on bivalent chiasma frequency in human males with normal karyotypes. Annals of Human Genetics 49, 193–201.

Lawrie NM, Tease C, Hultén MA. 1995. Chiasma frequency, distribution and interference maps of mouse autosomes. Chromosome 104, 308–314.

Li Q, Saito TT, Martinez-Garcia M, Deshong AJ, Nadarajan S, Lawrence KS, Checchi PM, Colaiacovo MP, Engebret J. 2018. The tumor suppressor BRC1A1-BARD1 complex localizes to the synaptonemal complex and regulates recombination under meiotic dysfunction in Caenorhabditis elegans. PLoS Genetics 14, e1007701.

Li X, Li Y, Yan J. 2015. Dissecting meiotic recombination based on tetrad analysis by single-microspore sequencing in maize. Nature Communications 6, 6648.

Lim EC, Kim J, Park J, et al. 2020. DeepTetrad: high-throughput image analysis of meiotic tetrads by deep learning in Arabidopsis thaliana. The Plant Journal 101, 473–483.

Lindegren CC, Lindegren G. 1942. Locally specific patterns of chromatid interference with applications to recombination data. Genetics 37, 441–457.

Lukaszewski AJ, Curtis CA. 1993. Physical distribution of recombination in B-genome chromosomes of tetraploid wheat. Theoretical and Applied Genetics 86, 121–127.

Lukaszewski AJ, Kopecky D, Linc G. 2012. Inversions of chromosome arms 4AL and 2BS in wheat invert the patterns of chiasma distribution. Chromosome Research 10, 266 | Ferreira et al.

Mather K. 1938. Crossing-over. Biological Reviews 13, 252–292.

Meneely PM, Farago AF, Kauffman TM. 2002. Crossover distribution and high interference for both the X chromosome and an autosome during oogenesis and spermatogenesis in Caenorhabditis elegans. Genetics 162, 1169–1177.

Mercier R, Jolivet S, Yezon D, et al. 2005. Two meiotic crossover classes cohabit in Arabidopsis. Current Biology 15, 692–701.

Modliszewski JL, Wang H, Albright AR, Lewis SM, Bennett AR, Huang J, Ma H, Wang Y, Copenhaver GP. 2018. Elevated temperature increases meiotic crossover frequency via the interfering (Type I) pathway in Arabidopsis thaliana. PLoS Genetics 14, e1007384.

Muller HJ. 1916. The mechanism of crossing-over. The American Naturalist 50, 195–221.

Nambari M, Chuan YC, Smith GR. 2019. Distributing meiotic crossovers for optimal fertility and evolution. DNA Repair 81, 102648.

Osman K, Higgins JD, Sanchez-Moran E, Armstrong SJ, Franklin FC. 2011. Pathways to meiotic recombination in Arabidopsis thaliana. New Phytologist 190, 523–544.

Otto SP, Payseur BA. 2019. Crossover interference: shedding light on the evolution of recombination. Annual Review of Genetics 53, 19–44.

Peng J, Korol AB, Fahima T, Röder MS, Ronin YI, Li YC, Nevo E. 2000. Molecular genetic maps in wild emmer wheat. Triticum dicoccoides: genome-wide coverage, massive negative interference, and putative quasi-linkage. Genome Research 10, 1590–1531.

Perkins DD. 1962. The frequency in Neurospora tetrads of multiple exchanges within short intervals. Genetic Research 3, 315–327.

Phillips D, Wnetrzak J, Nibau C, Barakate A, Ramsay L, Wright F, Higgins JD, Perry RM, Jenkins G. 2013. Quantitative high resolution mapping of HvMLH3 foci in barley pachytene nuclei reveals a strong distal bias and weak interference. Journal of Experimental Botany 64, 2139–2154.

Portin P. 2012. Further evidence for the theory that crossover interference in Drosophila melanogaster is dependent on genetic rather than physical distance between adjacent crossover points. Open Journal of Genetics 02, 155–162.

R Core Team. 2019. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing.

Säll T, Bengtsson BO. 1989. Apparent negative interference due to variation in recombination frequencies. Genetics 122, 935–942.

Scholten OE, van Kaauwen MP, Shahin A, Hendrickx PM, Keizer LC, Burger K, van Heusden AW, van der Linden CG, Vosman B. 2016. SNP-markers in Allium species to facilitate introgression breeding in onion. BMC Plant Biology 16, 187.

Sekhon JS. 2011. Multivariate and propensity score matching software with automated balance optimization: the matching package for R. Journal of Statistical Software 42, 1–52.

Shen P, Huang HV. 1986. Homologous recombination in Escherichia coli: dependence on substrate length and homology. Genetics 112, 441–457.

Shen P, Huang HV. 1989. Effect of base pair mismatches on recombination via the RecBCD pathway. Molecular & General Genetics 218, 358–360.

Snow R. 1979. Maximum likelihood estimation of linkage and interference from tetrad data. Genetics 92, 231–245.

Smith GR, Nambari M. 2020. New solutions to old problems: molecular mechanisms of meiotic crossover control. Trends in Genetics 36, 337–346.

Strickland WN. 1958. An analysis of interference in Aspergillus nidulans. Proceedings of the Royal Society B: Biological Sciences 149, 82–101.

Strickland WN. 1961. Tetrad analysis of short chromosome regions of Neurospora crassa. Genetics 46, 1125–1141.

Sturtevant AH. 1913. A third group of linked genes in Drosophila melanogaster. Science 37, 990–992.

Teuscher F, Brockmann GA, Rudolph PE, Swalve HH, Guiard V. 2000. Models for chromatic interference with applications to recombination data. Genetics 156, 1449–1460.
Wang P, Jiang L, Ye M, Zhu X, Wu R. 2019. The genomic landscape of crossover interference in the desert tree *Populus euphratica*. Frontiers in Genetics 10, 440.

Wang Y, Copenhaver GP. 2018. Meiotic recombination: mixing it up in plants. Annual Review of Plant Biology 69, 577–609.

Whitehouse HLK. 1958. Use of loosely linked genes to estimate chromatid interference by tetrad analysis. Nature 182, 1173–1174.

Zhang L, Kim KP, Kleckner NE, Storlazzi A. 2011. Meiotic double-strand breaks occur once per pair of (sister) chromatids and, via Mec1/ATR and Tel1/ATM, once per quartet of chromatids. Proceedings of the National Academy of Sciences, USA 108, 20036–20041.

Zhao H, McPeek MS, Speed TP. 1995b. Statistical analysis of chromatid interference. Genetics 139, 1057–1065.

Zhao H, Speed TP, McPeek MS. 1995a. Statistical analysis of crossover interference using the chi-square model. Genetics 139, 1045–1056.

Zickler D, Kleckner N. 2016. A few of our favorite things: pairing, the bouquet, crossover interference and evolution of meiosis. Seminars in Cell & Developmental Biology 54, 135–148.