Does polyploidy inhibit sex chromosome evolution in angiosperms?

Li He¹ and Elvira Hörandl²*

¹Eastern China Conservation Centre for Wild Endangered Plant Resources, Shanghai Chenshan Botanical Garden, Shanghai, China, ²Department of Systematics, Biodiversity and Evolution of Plants, University of Göttingen, Göttingen, Germany

Dioecy is rare in flowering plants (5–6% of species), but is often controlled genetically by sex-linked regions (SLRs). It has so far been unclear whether, polyploidy affects sex chromosome evolution, as it does in animals, though polyploidy is quite common in angiosperms, including in dioecious species. Plants could be different, as, unlike many animal systems, degenerated sex chromosomes, are uncommon in plants. Here we consider sex determination in plants and plant-specific factors, and propose that constraints created at the origin of polyploids limit successful polyploidization of species with SLRs. We consider the most likely case of a polyploid of a dioecious diploid with an established SLR, and discuss the outcome in autopolyploids and allopolyploids. The most stable system possibly has an SLR on just one chromosome, with a strongly dominant genetic factor in the heterogametic sex (e.g., xxxY male in a tetraploid). If recombination occurs with its homolog, this will prevent Y chromosome degeneration. Polyploidy may also allow for reversibility of multiplied Z or X chromosomes into autosomes. Otherwise, low dosage of Y-linked SLRs compared to their multiple homologous x copies may cause loss of reliable sex-determination at higher ploidy levels. We discuss some questions that can be studied using genome sequencing, chromosome level-assemblies, gene expression studies and analysis of loci under selection.

Introduction

Dioecy, the sexual system with separate female and male individuals, is phylogenetically widespread in angiosperms, but occurs only in 5–6% of species (Ming et al., 2011; Renner, 2014). Functional hermaphroditism with hermaphroditic flowers is the most common system, and the transition to dioecy can involve gynodioecy, monoecy, or other sexual systems (Darwin, 1877; Richards, 1997). Dioecy in many plants is under genetic control, with a sex-linked locus or region (SLR). We term
such chromosomes "SLR bearing," but here restrict the definition of sex chromosomes (following: Charlesworth, 2015) to mean chromosomes carrying completely sex-linked regions that include many genes and do not recombine. This distinction recognizes a key aspect of such “classical” sex chromosomes, their lack of recombination, which can arise in several different ways (Charlesworth, 2022), and which allows accumulation of deleterious mutations, resulting in degenerated sex chromosomes like those in mammals and Drosophila (Muller, 1918; Bachtrog, 2008). One hypothesis involves selection favoring the suppression of recombination between the sex-determining locus and linked sexually antagonistic polymorphic genes (Fisher, 1931; Rice, 1987). Suppression of recombination, however, might also result from neutral sequence divergence (Jeffries et al., 2021). Lenormand and Roze (2022) recently proposed a model for stepwise evolution of sex chromosomes via “lucky inversions” that prevent recombination. Genetic sex-determination has evolved independently multiple times in plants, but many have remained in the homomorphic state, suggesting that recombination has often not become suppressed, and that their SLRs carry just a sex-determining gene (or a few physically close and closely clinked sex-determining genes), explaining why degeneration have not occurred in these species. Eventually, ancestral sex chromosomes can be replaced by a new set of sex-determining chromosomes, and revert to autosomes (Vicoso, 2019; Figure 1A).

Many angiosperms are polyploid, having more than two chromosome sets (Van De Peer et al., 2017). Polyploidy is regarded an important evolutionary mechanism for angiosperms (e.g., Comai, 2005; Soltis and Soltis, 2009, 2016; Abbott et al., 2013; Hörandl, 2022). All angiosperms share an ancient genome duplication (Jiao et al., 2011), and 25–35% of species are more recently evolved polyploids of this already polyploid state (Landis et al., 2018). Unlike most animal polyploids, most polyploid plants are viable and can reproduce sexually (e.g., Levin, 2002; Comai, 2005). Asexual reproduction (apomixis) occurs in less than 1% of species and is not necessarily correlated to polyploidy (Mogie, 1992; Hojsgaard et al., 2014; Hojsgaard and Hörandl, 2019). In animals, the reason for the rarity of polyploidization is thought to be because it disturbs the “balance” system of sex determination by the X/autosome ratio (Muller, 1925). Such systems are found in species with extremely degenerated sex chromosomes in the heterogametic sex like the Drosophila Y (Orr, 1990). In this case, the heterogametic sex in a tetraploid (XXYY) will only produce heterozygous (XY) gametes if X-X and Y-Y pairs form preferentially at meiosis; since YY gametes are not viable, the heterogametic sex will be lost, resulting in rapid extinction (Orr, 1990; Figure 1B). Although sex chromosomes of angiosperms have been studied for a century (reviewed by Westergaard, 1958), less attention has been given to polyploidy in dioecious plants.

A correlation analysis of polyploidy and sexual systems on c. 1000 angiosperm species found that polyploidy is associated with sexual dimorphism (gynodioecy, androdioecy, and dioecy), but not specifically with dioecy, and that patterns are highly clade-specific (Glick et al., 2016). For many dioecious tropical trees and shrubs, however, chromosome numbers and ploidy levels are unknown, and more work is needed.

Knowledge about sex chromosome evolution in polyploid plants is also scanty. Modern review articles about SLRs and genetic sex determination in plants focus on diploid systems (Feng et al., 2020; Charlesworth, 2021; Renner and Müller, 2021). The recently launched database,1 included 166 angiosperm species with sex chromosomes or SLR-regions, of which 110 (66.3%) are classified as diploids, vs. only 20 (12%) as polyploids (Baránková et al., 2020), suggesting that polyploidy would not be highly correlated with sex chromosome evolution. However, the data are not yet clear enough, as 36 of the dioecious species (21.7%) have unknown ploidy levels (Baránková et al., 2020). Moreover, species-rich plant orders and families that experienced genome duplications before their diversification (Soltis et al., 2009), are hardly represented (only six cases in Brassicales, four in Poaceae, zero in Fabaceae, and Solanaceae), yet dioecy is present in all of them.

On the other hand, it is already known that several entirely dioecious lineages include polyploid series (Westergaard, 1958). Polyploidy may trigger shifts to dioecy (via different pathways reviewed by Ashman et al., 2013; Henry et al., 2018) speculated that whole-genome duplication could favor shifts to dioecy via evolution of sex-determining loci in duplicated genes. However, few such shifts are documented, except in Fragaria (Tennesen et al., 2016) where dioecy evolved in highly diploidized octoploids that arose from diploid, hermaphroditic progenitors (Tennesen et al., 2014). Most dioecious polyploids probably arose from diploid species with established SLR systems (e.g., Salix polyclona, He et al., 2022). Almost all angiosperms in which sex chromosome evolution is currently studied, including kiwi, papaya, persimmon, asparagus, strawberry, are cultivated plants (Henry et al., 2018), and little is known about sex chromosome evolution in natural polyploid systems. However, some examples (Table 1) include the tetraploids Rumex acetosella (Cunado et al., 2007) and hexaploid Mercurialis annua (Gerchen et al., 2022) all with male heterogamy. We briefly summarize sex chromosome evolution in diploids, and focus here on models of polyploidization from progenitors with established diploid non-degenerated systems (XY), and discuss how the presence of SLR-bearing chromosomes may affect new polyploids, and the cytological consequences, and whether polyploidy favors breakdown of dioecy, or causes reversals of X (or Z) chromosomes back to autosomes.

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1 https://www.sexchrom.csic.es/
A. Evolution of a sex chromosome in diploid plants

1) No degeneration small SLR
2) reduced recombination mutation accumulation TE invasion reversible degeneration
3) irreversible gene loss X/Y divergence

B. X/A ratio (A = autosomal ploidy); e.g., tetraploid (4A):

| Adults | Viable gametes |
|--------|----------------|
| 4X : 4A = 1.0 | female |
| 3X : 4A = 0.75 | intersexual |
| 2X : 4A = 0.5 | male |
| 1X : 4A = 0.25 | male |

C. Autopolyploid origin; x and y codominant

D. Autopolyploid origin; Y dominant over x; supernumerary x revert to autosomes (A)

E. Allopolyploid origin with both unreduced gamete formation; Y dominant; preferential pairing of homologs

FIGURE 1 (Continued)
Sex chromosome evolution in diploid plants

In sessile organisms, separation of the sexes in different individuals reduces the chance of successful matings (Darwin, 1876; Hörandl and Hadacek, 2020). One factor favoring the evolution of dioecy in plants, and maintaining it over long evolutionary times, is prevention of self-fertilization and consequent inbreeding depression (Darwin, 1877; Renner, 2014). This is particularly important in wind-pollinated, long-lived plants, including tropical trees.

Dioecy and sex-linked regions evolved multiple times in angiosperms (Renner, 2014; Feng et al., 2020). Evolution from a functionally hermaphroditic ancestor requires two mutations, and both currently known plant sex-determining systems involve a gene whose action promotes one sex function and suppresses the other (review e.g., in Renner and Müller, 2021). Evolution via a turnover event in an ancestrally dioecious species will also usually involve an active maleness or femaleness factor, and both currently known plant sex-determining systems, as in many animals (including Drosophila, nematodes, and, with female heterogamy, Lepidoptera) (Bachtrog et al., 2022), showing that polyploids cannot maintain stable X/Y ratios nor restore the heterogametic sex. This form of sex-determination appears to be rare in angiosperms. (C) An autopolyploid plant originated via unreduced female gametes termed a “triploid bridge,” because a triploid generation is initially formed by unreduced gametes of one parent, and can produce haploid, diploid, and triploid gametes; the * denotes low fertility of this generation. The diagram shows the expected progeny types assuming codominant expression of a Y-linked male-determining factor, and no preferential pairing of homologs (possible xyy genotypes not shown). (D) Autopolyploid with a dominant Y-linked factor; xxy genotypes represent fertile males. After some time two of the x may revert to autosomes A, whereas recombination between the xY pair prevents degeneration. (E) Allopolyploid origins via unreduced gametes of both parents (shown in green and gray), and with a dominant Y-linked maleness factor. Homologs of the same parent will preferentially pair at meiosis (see also Supplementary Figure 1). Initially only xxy male allopolyploids can be formed and produce heterozygous male gametes (see text). Backcrossing with unreduced gametes of female parents can produce various tetraploid hybrid genotypes, including both xxx females and xxy males, resulting in stable populations. Here also two of the x may revert to autosomes (A), and only one possible recombining xY pair (derived from the same progenitor) remains. Further hybridization with a third species can result in a hexaploid, which eventually results in reversal to a non-dioecious system.
Mank, 2022). Hence, haploid selection is probably much stronger in plants than in animals (Otto et al., 2015), and could prevent sex chromosome degeneration. Indeed, YY (or WW) genotypes are viable in diploid dioecious plants at first phase (Ming et al., 2011), which is consistent with purifying selection eliminating deleterious mutations in haploid Y (or W) chromosomes in gametophytes (though other alternatives, including recent origins, or small sizes of sex-linked regions, are probably also important).

Degeneration will not happen if recombination occurs. Dioecy in plants restricts sexual selection mostly to post-pollination processes (Hörandl and Hadacek, 2020). Hence, selection might favor sexually antagonistic mutations in sex-linked genes less often than in animals. Therefore factors other than sexual antagonism may be required for dioecious plants to evolve extensive fully sex-linked regions and undergo degeneration (e.g., Lenormand and Roze, 2022).

Sex chromosomes in polyploid plants

Cytological mechanisms

Polyploids are divided into two major types, autopolyploids that originate within one species, and those originating after hybridization between species (allopolyploids). The chromosomes of new autopolyploids may form multivalents or rings, as they are not diverged and do not pair preferentially at meiosis (e.g., Stebbins, 1947; Comai, 2005). Allopolyploids have two diverged chromosome sets (homoeologs); chromosomes derived from the same parent species show preferential pairing, resulting in more or less regular bivalent formation, whereas homoeolog pairing is less frequent (Mason and Wendel, 2020; Supplementary Figure 1). Therefore, allopolyploids maintain fixed heterozygosity for the different alleles derived from the two parental species and carried by the homoeologous chromosomes (Comai, 2005; Glover et al., 2016). These mechanisms have consequences for SLR-bearing chromosomes.

Autopolyploids originate mainly via the “triploid bridge” process (Figures 1C,D; Ramsey and Schemske, 1998). The major constraint preventing establishment of such autopolyploids is male or female bias and reduced fertility in the triploid generation and possible instability of meiosis, resulting in aneuploid progeny and low fertility (e.g., Comai, 2005). But its unreduced (triploid) gametes can fuse during fertilization with haploid gametes from a parent of the other sex, potentially establishing a cytologically stable autotetraploid. Various tetraploid sex chromosome configurations are possible, including xxyy (small captions mean codominant or recessive versus dominant Y) genotypes, which could produce xx, xy, and yy gametophytes and gametes (Figure 1C). If Y is dominant, xxxx females and xxxY males may emerge, constituting a stable population (Figure 1D), though sex-determination in autotetraploid dioecious plants with homomorphic sex chromosomes have been little studied (Spoelhof et al., 2017). In the heterogametic sex, the SLR-bearing chromosome can pair and (if the Y-linked region is not rearranged or otherwise prevented from recombining with the x) it may continue to recombine with any of the other three x chromosomes during meiosis, preventing degeneration (Figures 1C,D).

Allopolyploidization rarely involves a triploid bridge, but mostly arises after fertilization between unreduced gametes derived from two parental species or hybrids (Ramsey and Schemske, 1998). A first factor limiting production of such populations is that both parents must have the same XY or WZ system, which can differ within a genus (e.g., Salix and Silene) (Balounova et al., 2019; Martin et al., 2019; Gulyaev et al., 2022). The second, newly formed polyploid will be in the minority and, as single individuals of dioecious plants cannot self-fertilize, they will backcross to the parents, which creates low fitness progeny (Levin, 1975; Ramsey and Schemske, 1998). Third, reproduction of the newly formed polyploid requires fertilization by a plant of the other sex (see Figures 1C,E); thus, they must live long enough to overlap with offspring generations. They will also have to form unreduced gametes, because only fusion of unreduced male and female gametes from the two dioecious parent species will produce tetraploids with the heterogametic sex (e.g., xx1xY2 male, 1 from species 1 and 2 from species 2). To produce males, the Y-linked male-determining gene must be dominant, because male xx1xY2 genotypes cannot be formed via unreduced gametes of allopolyploids because the female parent can only produce xx eggs and the male parent mostly xY pollen (Figure 1E). Somatic doubling is rare in nature (Ramsey and Schemske, 1998), and hence doubling of a x1y2 male diploid interspecific hybrid together with doubling a xx female is unlikely to happen frequently enough to establish populations. The xx1xY2 allotetraploid will produce male flowers and x1Y2 and x1x2 gametes. Its SLR-bearing chromosome Y will preferentially pair with the x from the same parent, resulting in four male heterozygous gamete classes. The formation of the next hybrid generation requires crossing with an unreduced female gamete derived from the other parent; only if this occurs, a stable cohort of allotetraploids with xxxY males and xxxx females can form. Most plants can produce fertile heterogametic males with only one copy of the Y chromosomes (XXXY) and females (XXXX) (e.g., Melandrium album = Silene latifolia) (Warmke and Blakeslee, 1939; Westergaard, 1958; Henry et al., 2018). This system can potentially be maintained for higher polyploids (Westergaard, 1958). Similar considerations apply to female heterogamy.

As in autopolyploids (Figure 1E), less differentiated Y chromosomes might recombine with the conspecific X and prevent further degeneration in polyploids (Charlesworth et al., 2021). On the other hand, the presence of duplicated sex chromosomes might increase recombination rates between the SLRs of autopolyploids (Pecinka et al., 2011; Glover et al., 2016). Furthermore, polyploidization can also increase the
recombination rate of X or Z chromosomes, maintaining their gene content (Wilson and Makova, 2009). However, natural selection likely reduces crossover rates of tetraploid homologs. Gates (1926) and Wright et al. (2015) suggested that the multiplied sex chromosomes (Z or X) in polyploids can gradually transform into autosomes, and this appears to have occurred in polyploid Rumex acetosella, which each retained have only one pair of sex chromosomes (XY for males) (Singh, 1968; Cunado et al., 2007). The dioecious octoploid Fragaria (e.g., Fragaria chiloensis) species that arose from hermaphroditic diploid progenitors also each have only one SLR on one of the homologues in each species studied (Tennessen et al., 2016; Cauret et al., 2022), while the others may have become autosomes. The “new” autosomes could still be just like the ancestral X or Z chromosome, or their genes might be lost due to diploidization.

Genetic degeneration

Tetraploid plants will regularly form diploid gametes, in which the effects of recessive deleterious mutations in one chromosome copy will be lessened by the function provided by the unmutated chromosome; this may mask mutations from selection (e.g., Otto and Gerstein, 2008), especially in high ploidy levels (e.g., hexaploid xxxxxY). Whether genes in a Y-linked region will degenerate more slowly in a polyploid than in a diploid plant, will probably depend on several factors, including the mutations’ dominance coefficients, as well as any differences in recombination. Whether the masking effect could lead to an accumulation of deleterious recessive mutations in the pseudoautosomal region of sex chromosome in the heterozygous state needs to be investigated.

Consequences of polyploidy for the maintenance of dioecy

Polyploids have been observed to be associated with loss of dioecy of diploid progenitors, with changes occurring to monoecy, e.g., in hexaploid Diospyros kaki (Akagi et al., 2016), to hermaphroditism, e.g., in tetraploid Empetrum (Anderberg, 1994), or to monoecy and androdioecy in tetra- to dodecaploid Mercurialis (Pannell et al., 2004; Gerchen et al., 2022). In willows (Salix), shifts to diverse sexual systems occur in polyploids (Mirska, 2014; Mirski et al., 2017). These transitions to non-dioecious systems might be favored by selection for reproductive assurance by uniparental reproduction, especially in colonization scenarios (Ashman et al., 2013). However, also gene dosage effects and differential homeolog expression could cause lability if only one active Y interacts with multiple copies of x (unless all but one have turned into autosomes). Interestingly, introgression of a new Y chromosome from distantly related species happened at the origin of hexaploid, androecious Mercurialis annua (Gerchen et al., 2022). Such reversals to non-dioecious sexual systems may be indicators that polyploidization can prevent the maintenance of sex-determining systems for long evolutionary times, and this may be a further factor contributing to the rarity of degenerated sex chromosomes in plants. However, the shifts from dioecy to non-dioecious systems in several cultivated plants, including grape vines (Badouin et al., 2020) and papaya (Ming et al., 2007), do not involve ploidy changes, and could be due to human selection for major mutations (Henry et al., 2018). Loss of active sex-determiners, such as the maleness factors discussed above, could involve Y chromosome deletions, as observed in Silene latifolia (Kazama et al., 2016).

Discussion and outlook

Further work on polyploid dioecious plants with SLRs is clearly needed. Studies on neopolyploids (or synthetic polyploids) would help test whether dioecy in polyploids is rare because of constraints preventing their origination. Second, it should be tested whether auto- and allopolyploids differ in their heterogametic chromosome constitution, as predicted. Third, recombination between sex chromosome homologs, with non-recombining regions restricted in size, probably often prevents degeneration of plant sex chromosomes, alongside selection in the haploid pollen. The sizes of regions should be investigated for both XY and ZW systems, and the times when they evolved. Finally, the role of genetic or epigenetic effects in lability of sexual systems needs further study, specifically whether lability tends to be greater in higher polyploids with only one copy of the heterogametic chromosomes (XY or ZW).

Cytogenetic studies can identify heteromorphic sex chromosomes, and these will often be degenerated W or Y chromosomes, and also detectable from sex differences in genomic coverage, but in plants these appear to be the exception. Homomorphic sex chromosomes are more difficult to detect (Palmer et al., 2019; Baránková et al., 2020), sometimes not even with genome sequencing. Genome-wide association studies (GWAS) can detect sex-linked regions, but will often find large partially sex-linked regions as well as completely sex-linked ones. Genetic differentiation between the sexes (FST) can provide a finer scale for use with less differentiated SLR-bearing chromosomes (Palmer et al., 2019; He et al., 2021) and even for polyploids (Kühl et al., 2021). The highly accurate long HiFi (high-fidelity) reads produced by PacBio, combined with Hi-C, can generate haplotype-resolved genome assemblies of polyploids (Zhang et al., 2019; Cheng et al., 2021). Genome coverage, and of Ka/Ks ratios in divergence from related species to detect weakened efficacy of purifying selection, can detect and quantify degeneration of SLR protein-coding genes (as in examples outlined above). Analyses of gene expression via RNA seq, and epigenetic mechanisms controlling expression.
(including cytosin-methylation and small RNAs activity) are important for understanding the expression of different sex phenotypes.

Author contributions

Both authors listed have made a substantial, direct, and intellectual contribution to the work, and approved it for publication.

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References

Abbott, R., Albach, D., Ansell, S., Arntzen, J. W., Baird, S. J. E., Bierne, N., et al. (2013). Hybridization and speciation. J. Evol. Biol. 26, 229–246. doi: 10.1111/j. 1420-9101.2012.02598.x

Akagi, T., Henry, I., Kawai, T., Comai, L., and Tao, R. (2016). Epigenetic regulation of the sex determination gene MeGI in polyploid persimmon. Plant Cell 28, 2905–2915. doi: 10.1105/tpc.16.00532

Anderberg, A. (1994). Phylogeny of the empetraceae, with special emphasis on character evolution in the genus empetrum. Syst. Biol. 19, 35–46. doi: 10.2307/ 2419710

Ashman, T. L., Kwok, A., and Husband, B. C. (2013). Revisiting the dioecy-polyplody association: Alternate pathways and research opportunities. Cytog. Gen. Rev. 140, 241–255. doi: 10.1159/000355306

Bachtrog, D. (2008). The temporal dynamics of processes underlying Y chromosome degeneration. Genetics 179, 1513–1525. doi: 10.1534/genetics.107. 084012

Bachtrog, D., Mank, J. E., Peichl, C. L., Kirkpatrick, M., Otto, S. P., Ashman, T. L., et al. (2014). Sex determination: Why so many ways of doing it? PLoS Biol. 12:e1001899. doi: 10.1371/journal.pbio.1001899

Badouin, H., Veh, A., Gindraud, F., Flutre, T., Dumas, V., Vautrin, S., et al. (2020). The wild grape genome sequence provides insights into the transition from dioecy to hermaphroditism during grape domestication. Gen. Biol. 21:223. doi: 10.1186/s13059-020-02131-y

Balounová, V., Gogela, R., Cegan, R., Cangren, P., Zluvová, J., Safar, J., et al. (2019). Evolution of sex determination and heterogamy changes in section otites of the genus silene. Sci. Rep. 9:1045. doi: 10.1038/s41598-018-37412-x

Baránková, S., Pascual Díaz, J., Sultana, N., Alonso Lifante, M., Balant, M., Balounova, V., Cegan, R., Cangren, P., Zluvová, J., Safar, J., et al. (2020). Sex-chrom, a database on plant sex chromosomes. New Phytol. 227, 1594–1604. doi: 10.1111/nph.16635

Bull, J. J. (1983). Evolution of Sex Determining Mechanisms. Menlo Park, CA: The Benjamin/Cummings Publishing Company, Inc.

Cauret, C. M. S., Mortimer, S. M. E., Roberti, M. C., Ashman, T.-L., and Liston, A. (2022). Chromosome-scale assembly with a phased sex-determining region resolves features of early Z and W chromosome differentiation in a wild octoploid strawberry. G3 (Bethesda, Md) 12:jkac139. doi: 10.1093/g3journal/jkac139

Charlesworth, D. (2015). Plant contributions to our understanding of sex chromosome evolution. New Phytol. 208, 52–65. doi: 10.1111/nph.13497

Charlesworth, D. (2021). When and how do sex-linked regions become sex chromosomes? Evolution 75, 569–581. doi: 10.1111/ero.14196

Charlesworth, D. (2022). Some thoughts about the words we use for thinking about sex chromosome evolution. Philos. Trans. R. Soc. London Ser. B Biol. Sci. 377:20210314. doi: 10.1098/rstb.2021.0314

Charlesworth, D., Bergero, R., Graham, C., Gardner, J., Keegan, K., and Dyer, K. A. (2021). How did the guppy Y chromosome evolve? PLoS Genet. 17:e1009704. doi: 10.1371/journal.pgen.1009704

Cheng, H., Concepcion, G., Feng, X., Zhang, H., and Li, H. (2021). Haplotype-resolved de novo assembly using phased assembly graphs with hifiasm. Nat. Methods 18, 170–175. doi: 10.1038/s41592-020-01056-5

Comai, L. (2005). The advantages and disadvantages of being polyploid. Nat. Rev. Genet. 6, 836–846. doi: 10.1038/nrg1711

Cronk, Q. (2022). Some sexual consequences of being a plant. Philos. Trans. R. Soc. London Ser. B Biol. Sci. 377, 20210213–20210213. doi: 10.1098/rstb.2021.0213

Cunado, N., Cuñado, N., Navajas Pérez, R., de la Herrán, R., Ruiz Rejón, C., Ruiz Rejón, M., et al. (2007). The evolution of sex chromosomes in the genus rumex (polygonaceae): Identification of a new species with heteromorphic sex chromosomes. Chrom. Res. 15, 825–833. doi: 10.1007/s10577-007-166-6

Darwin, C. (1876). Cross and Self-Fertilization of Plants. London: John Murray.

Darwin, C. (1877). The Different Forms of Flowers on Plants of the Same Species. London: John Murray. doi: 10.5962/bhl.title.11054

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Supplementary material

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Mank, J. E. (2022). Are plant and animal sex chromosomes really all that different? Philos. Trans. R. Soc. Lond. Ser. B. Biol. Sci. 377:20210218. doi: 10.1098/rstb.2021.0218

Martin, H., Carpenter, F., Gallina, S., Godé, C., Schmitt, E., Moyse, A., et al. (2019). Evolution of young sex chromosomes in two dioecious sister plant species with distinct sex determination systems. Gen. Biol. Evol. 11, 350–361. doi: 10.1093/gbe/evz001

Mason, A. S., and Wendell, J. F. (2020). Homologous exchanges, segmental allopolyploidy, and polyploid genome evolution. Front. Genet. 11:1014. doi: 10.3389/fgene.2020.01014

Ming, R., Bendahan, M., and Renner, S. S. (2011). Sex chromosomes in land plants. Ann. Rev. Plant Biol. 62, 485–514. doi: 10.1146/annurev-plant-043009-103914

Ming, R., Yu, Q., Moore, H. P., and Moore, P. (2007). Sex determination in papaya. Semin Cell Dev. Biol. 18, 401–408. doi: 10.1016/j.semcdb.2006.11.013

Mirsik, P. (2014). Exceptions from dioecy and sex liability in genus salix. Dendrobiology 71, 167–171. doi: 10.23657/dentbio.071.017

Mirsik, P., Brzozko, E., Jedrzejczyk, I., Kotowicka, J., Ostrowiecka, B., and Wroblewska, A. (2017). Genetic structure of dioecious and triecious salix myrsinifolia populations at the border of geographic range. Tree Genet. Genom. 13:15. doi: 10.1007/s11295-016-1096-6

Mogie, M. (1992). The Evolution of Asexual Reproduction in Plants. London: Chapman and Hall.

Mulder, H. J. (1918). Genetic variability, twin hybrids and constant hybrids, in a case of balanced lethal factors. Genetics 3, 422–499. doi: 10.1093/genetics/3.3.422

Mulder, H. J. (1925). Why polyploidy is rarer in animals than in plants. Am. Nat. 59, 346–353. doi: 10.1086/280047

Orr, H. A. (1990). Why polyploidy is rarer in animals than in plants revisited. Am. Nat. 136, 759–770. doi: 10.1086/285130

Otto, S. P., and Geisert, A. C. (2008). The evolution of haploidy and diploidy. Curr. Biol. 18, R1121–R1124. doi: 10.1016/j.cub.2008.09.039

Otto, S. P., Scott, M. F., and Immler, S. (2015). Evolution of haploid selection in predominantly diploid organisms. Proc. Natl. Acad. Sci. U.S.A. 112, 15952–15957. doi: 10.1073/pnas.1520041112

Palmer, D., Rogers, T., Dean, R., Wright, A. E., and Wright, A. (2019). How to identify sex chromosomes and their turnover. Mol. Ecol. 28, 4709–4724. doi: 10.1111/mec.15245

Pannell, J., Obbard, D., Buggs, R. J. A., and Buggs, R. J. A. (2004). Polyploidy and the sexual system: what can we learn from mercurialis annua? Mol. Ecol. 13, 441–447. doi: 10.1111/j.1365-294x.2004.02088.x

Ramsey, J., and Schemske, D. W. (1998). Pathways, mechanisms, and rates of polyploid formation in flowering plants. Ann. Rev. Ecol. Syst. 29, 467–501. doi: 10.1146/annurev.ecolsys.29.1.467

Renners, S., and Muller, N. A. (2021). Plant sex chromosomes defy evolutionary models of expanding recombination suppression and genetic degeneration. Nat. Plants 7, 392–402. doi: 10.1038/s41477-021-00884-3

Renners, S. S. (2014). The relative and absolute frequencies of angiosperm sexual systems. Dioecy, monoecy, gynodioecy, and an updated online database. Am. J. Bot. 101, 1588–1596. doi: 10.3732/ajb.1400196

Rice, W. R. (1987). The accumulation of sexually antagonistic genes as a selective agent promoting the evolution of reduced recombination between primitive sex chromosomes. Evolution 41, 911–914. doi: 10.1111/j.1558-5646.1987.tb0864x

Richards, J. A. (1997). Plant Breeding Systems. London: Chapman and Hall. doi: 10.1007/978-1-4899-3043-9

Singh, R. B. (1968). A dioecious polyploid in rumex acetosella. J. Heredity 59, 168–170. doi: 10.1093/jhered/59.4.168

Solits, D. E., Albert, V. A., Leebens-Mack, J., Bell, C. D., Paterson, A. H., Zheng, C. F., et al. (2009). Polyploidy and angiosperm diversification. Am. J. Bot. 96, 336–348. doi: 10.3732/ajb.0800079

Solits, P. S., and Solits, D. E. (2009). The role of hybridization in plant speciation. Ann. Rev. Plant Biol. 60, 561–588. doi: 10.1146/annurev.arplant.043008.020395

Solits, P. S., and Solits, D. E. (2016). Ancient WGD events as drivers of key innovations in angiosperms. Curr. Opin. Plant Biol. 30, 159–165. doi: 10.1016/j.copb.2016.03.015
Spoelhof, J., Soltis, P., Soltis, D. E., and Soltis, D. (2017). Pure polyploidy: Closing the gaps in autoploidy research. J. Syst. Evol. 55, 340–352. doi: 10.1111/jse.12253

Stebbins, G. L. (1947). Types of polyploids: their classification and significance. Adv. Genet. 1, 403–429. doi: 10.1016/S0065-2660(08)60490-3

Tennessen, J., Govindarajulu, R., Ashman, T.-L., and Liston, A. (2014). Evolutionary origins and dynamics of octoploid strawberry subgenomes revealed by dense targeted capture linkage maps. Gen. Biol. Evol. 6, 3295–3313. doi: 10.1093/gbe/evu261

Tennessen, J., Govindarajulu, R., Liston, A., and Ashman, T.-L. (2016). Homomorphic ZW chromosomes in a wild strawberry show distinct recombination heterogeneity but a small sex-determining region. New Phytol. 211, 1412–1423. doi: 10.1111/nph.13983

Van De Peer, Y., Mizrachi, E., and Marchal, K. (2017). The evolutionary significance of polyploidy. Nat. Rev. Genet. 18, 411–424. doi: 10.1038/nrg.2017.26

Vicoso, B. (2019). Molecular and evolutionary dynamics of animal sex chromosomes. Nat. Rev. Genet. 3, 1632–1641. doi: 10.1038/s41559-019-1050-8

Warmke, H. E., and Blakeslee, A. F. (1939). Sex mechanism in polyploids of melandrium. Science 89, 391–392. doi: 10.1126/science.89.2313.391

Westergaard, M. (1958). The mechanism of sex determination in dioecious flowering plants. Adv. Genet. 9, 217–281. doi: 10.1016/S0065-2660(08)60306-7

Wilson, M., and Makova, K. (2009). Genomic analyses of sex chromosome evolution. Annu. Rev. Genom. Hum. Genet. 10, 333–354. doi: 10.1146/annurev-genom-082908-150105

Wright, K. M., Arnold, B., Xue, K., Surinova, M., O’Connell, J., and Bomblies, K. (2015). Selection on meiosis genes in diploid and tetraploid Arabidopsis arenosa. Mol. Biol. Evol. 32, 944–955. doi: 10.1093/molbev/msv398

Zhang, X., Zhang, S., Zhao, Q., Ming, R., Tang, H., and Tang, H. (2019). Assembly of allele-aware, chromosomal-scale autoploidy genomes based on Hi-C data. Nat. Plants 5, 833–845. doi: 10.1038/s41477-019-0487-8

Zuk, J. (1963). An investigation on polyploidy and sex determination within the genus rumex. Acta Soc. Bot. Polocy 32, 5–67. doi: 10.5586/asbp.196 3.001