When and how do sex-linked regions become sex chromosomes?

Deborah Charlesworth

Institute of Evolutionary Biology, University of Edinburgh, Edinburgh EH9 3JT, United Kingdom

E-mail: Deborah.Charlesworth@ed.ac.uk

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The attention given to heteromorphism and genetic degeneration of “classical sex chromosomes” (Y chromosomes in XY systems, and the W in ZW systems that were studied first and are best described) has perhaps created the impression that the absence of recombination between sex chromosomes is inevitable. I here argue that continued recombination is often to be expected, that absence of recombination is surprising and demands further study, and that the involvement of selection in reduced recombination is not yet well understood. Despite a long history of investigations of sex chromosome pairs, there is a need for more quantitative approaches to studying sex-linked regions. I describe a scheme to help understand the relationships between different properties of sex-linked regions. Specifically, I focus on their sizes (differentiating between small regions and extensive fully sex-linked ones), the times when they evolved, and their differentiation, and review studies using DNA sequencing in nonmodel organisms that are providing information about the processes causing these properties.

KEY WORDS: Genetic degeneration, hemizygosity, partially sex-linked region, pseudoautosomal region (PAR), sexual antagonism.

Despite great progress in understanding the evolution of nonrecombining sex-linked genome regions, surprisingly many interesting questions remain, including how often sex-determining regions have evolved suppressed recombination, or why they did so when this has occurred. The gaps in knowledge can create an apparently confusing picture. I here argue that much of the diversity reflects well-understood biological processes acting in a diversity of organisms that evolved separate sexes independently, at different times in the past, with sex-linked regions on different chromosomes, rather than “many exceptions to the rules” (Furman et al. 2020). Some striking similarities between sex-linked regions in different organisms stem from their lack of recombination (and subsequent genetic degeneration of Y and W chromosomes). Such sex-linked regions have, however, been the focus of so much attention that these features have sometimes been viewed as ubiquitous (Ponnikas et al. 2018) and regarded as inevitable. I focus on understanding that can be gained from younger systems and sex-determining regions that are in the process of evolving.

Figure 1 shows changes if a sex-linked region appears in a genome region, and Figure 2 summarizes different situations that generate systems with different ages and properties, from newly evolved sex-determining genes to XY or ZW sex chromosomes. This simplified framework focuses attention on the following well-established concepts:

1. An important general “rule” is that absence of recombination is crucial for evolution of distinct sex chromosomes (or extensive sex-linked regions). Even low crossover rates prevent differentiation (Pamilo et al. 1987; Blaser et al. 2014).
2. Absence of recombination can arise in different ways (Fig. 2). Newly evolved (young) sex-determining regions will often be small, although they can arise within physically large nonrecombining regions. Sex-linked regions could potentially remain as small as the initial sex-determining gene, or could subsequently evolve into large recombinationally suppressed regions. As explained below, the role of selection in reducing recombination is not yet fully understood.
3. If a sex-determining region does not recombine, Y-X differentiation will become detectable, over time, contrasting with adjacent recombining, or “pseudo-autosomal,” regions
D. CHARLESWORTH

Figure 1. Schematic diagram to illustrate the stages of evolution of a completely sex-linked region after recombination stops, in cases where it does stop. The left-hand end is a pseudoautosomal region (PAR) that continues to recombine. The absence of numbers on the x-axis, and the y-axis for divergence, is deliberate, as the diagrams are intended to show general concepts. Specific examples, and measures of divergence and degeneration, are described in the text. Importantly, the timescales for different processes differ (as indicated by the lines connecting the short segment of time corresponding to neutral divergence and the degeneration process); specifically, as drawn, loss of selective constraints is shown occurring in an evolutionary time corresponding to small, rather than large, neutral divergence, and complete degeneration might follow shortly after this stage, or might take much longer.

Many advances have occurred since molecular markers, and especially genome sequencing, became available. Many studies hope to discover sex-determining genes, and therefore start with locating the species’ fully sex-linked genome region. These studies also provide valuable information about differentiation. We can now discover whether a species with genetic sex-determination has a nonrecombining region, and, if so, how large it is, and when recombination between sex chromosomes became suppressed. Figure 1 shows the situation in very general terms, based on studies of animal and plant sex chromosomes over many years, and has no quantitative x-axis scale for either the time or the level of genetic degeneration.

To categorize species with genetic sex-determination into the types in Figure 2, Figure 1 distinguishes between “sex-linked regions,” which span a wide range of ages, sizes, and levels of differentiation, and differentiated “sex chromosomes” (potentially with cytologically detectable heteromorphism) to categorize species with genetic sex-determination into the types in Figure 2. As has long been suspected, some organisms are in the early stages of sex chromosome evolution. For example, the plant *Mercurialis annua* shows minimal sequence divergence between Y- and X-linked sequences, and little sign of genetic degeneration (Veltsos et al. 2019). In some animals, new sex-determining regions evolved recently through “turnover events” (reviewed in Vicoso 2019). In contrast, other well-studied animals, including mammals, *Drosophila*, and many birds, have very old-established sex chromosomes where most of the XY or ZW pair is nonrecombining, and the Y or W is highly degenerated, having lost most of the genes carried on the X or Z counterpart. In plants, separate sexes evolved more recently (see below), but the sex chromosomes of several distantly related dioecious plants share these properties, including *Silene latifolia* (Papadopulos et al. 2015), *Rumex* species (Grabowska-Joachimiak et al. 2015; Crowson et al. 2017), *Cannabis sativa* (Prentout et al. 2020), and *Coccinia grandis* (Fruchard et al. 2020).

Detecting Completely Sex-Linked Regions, and Estimating Their Sizes, When They are Present

Cytogenetic studies and genome sequencing

Cytogenetic studies were important in the discovery of many of the old-established, highly degenerated, sex chromosomes examples just mentioned (Swanson 1957; Westergaard 1958; Zrzava et al. 2018). Loss of genes’ functions allowed deletions of large parts of the Y or W chromosomes, creating detectable YX heteromorphism, or even XO systems, as in many grasshoppers and nematodes, where the sex-determining genes have been lost (and maleness or femaleness no longer involves an active

(par Fig. 1). Importantly, increased sequence divergence, accumulation of repetitive sequences (expanding the region’s physical size), and genetic degeneration (potentially deleting genes, shrinking the Y-linked region) occur on different timescales. Sex chromosome heteromorphism mainly reflects the net effects of accumulation of repeats and genetic degeneration (other changes that can contribute will be mentioned below).
| Situations producing new sex determining regions | Direct effect on recombination | Creation of large non-recombining regions |
|---------------------------------------------|-----------------------------|------------------------------------------|
| Male- or female-determining factor on the Y or W chromosome; sex is then determined by an X-autosome balance system, as in *Drosophila*; see Swanson 1957). Heteromorphism can also occur by accumulation of repetitive sequences. In some birds, accumulation appears to have restored homomorphism after ZW heteromorphism had evolved, and the W chromosome had lost most genes (Rutkowska et al. 2012; Furo et al. 2017). Several different approaches have been developed for detecting such regions in genome sequences, and discovering their sizes. Detailed information is now available about large completely sex-linked regions in species where cytogenetic studies had already established their existence: mammals (Skaletsky et al. 2003; Cortez et al. 2014) birds (Pigozzi 2011; Zhou et al. 2014; Schmid et al. 2015), reptiles (Schield et al. 2019), fish, including sticklebacks (Varadhharajan et al. 2019; Peichel et al. 2020), Dipteran and Lepidopteran insects (Vicoso and Bachtrog 2015; Fraisse et al. 2017), and plants, including bryophytes (Allen 1917, 1932; Okada et al. 2001; Ishizaki et al. 2002; Marks et al. 2019). Physically smaller completely sex-linked regions that also carry multiple genes have been found in a moss (McDaniel et al. 2007; Carey et al. 2020), brown algae (Ahmed et al. 2014), the flowering plant papaya (Wang et al. 2012), and some cichlids (Gammerdinger and Kocher 2018). With just a single individual of each sex, fully sex-linked regions that have undergone genetic degeneration and lost genes can be ascertained because the Y- or W-linked region is “hemizygous” and shows sex-specific haploid depth of coverage compared with the other sex, or with genome regions that are not completely sex linked. This approach can be used even in non-model species such as Paleognathous birds (Zhou et al. 2014), snakes (Schield et al. 2019) and Schistosomes (Vicoso and Bachtrog 2011). However, it will only detect Y- and W-linked regions that have been nonrecombining for long enough to have lost many genes, or for sequences to have diverged sufficiently that they do not map to a reference genome of the other sex (“old” systems in Fig. 1). This approach may therefore fail to detect nonrecombining regions in species with younger sex

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**Figure 2.** Scheme to show the different ways in which a small region carrying the sex determining gene or genes can arise, and the different situations where these regions could expand in size to form extensive fully sex-linked regions that, given enough time, would be predicted to evolve the complete set of sex chromosome properties (divergence of Y- or W-linked sequences, accumulation of repetitive sequences, chromosome rearrangements, and genetic degeneration). In the two situations involving polymorphisms in two genes that create selection for closer linkage (right-hand columns), an inversion could suppress recombination across a physically large region, or “evolutionary stratum,” that includes many genes without sex-related functions.
chromosomes, or whose sex-linked regions include only a few genes.

**GENETIC MAPPING**

Genetic mapping within families can detect variants that co-segregate with the sex of the progeny. Combined with genome sequencing, very large numbers of variants in DNA (or RNA transcripts) can be mapped to positions in chromosomes’ physical maps in a species’ reference genome assembly. With dense markers, even small regions may be detectable. Genetic studies can also reveal that sex-determining genes are on different chromosomes in related species (suggesting creation of new sex-determining regions by turnover events or repeated evolution of new sex-determining genes, as mentioned in Fig. 2). Turnovers have been detected in many animals (Vicoso 2019), and are starting to be discovered in plants (Tennessen et al. 2018; Xue et al. 2020). Finally, genetic maps are increasingly being used to separate the two haplotypes of the sex-linked region, by genotyping progeny individuals and a parent of the heterozygous sex (Zhou et al. 2020). Such “phasing” of variants in a fully sex-linked region is necessary for describing Y-X differences and estimating Y-linked regions’ ages (see below). However, the haplotypes in families refer only to variants in the phased individual, which may not be consistently sex specific.

Genetic mapping has established that the plant *Mercurialis annua* has a genetic sex-determination system, with co-segregating Y-linked markers across a physically large region, at least in families (Veltsos et al. 2019). Family analysis will, however, overestimate the physical size of fully sex-linked regions, because, in a small family, some partially sex-linked markers will co-segregate with fully sex-linked ones and be classified incorrectly. The true sizes are therefore often not accurately known, but only upper size estimates.

**GENOME-WIDE ASSOCIATION APPROACHES (GWAS)**

**USING SEX AS THE PHENOTYPE**

To ascertain sex-specific variants, samples of multiple unrelated individuals from natural populations are needed, to ensure many past generations during which recombination can occur. Such genome-wide analyses often employ $F_{ST}$ estimates between the sexes for sequences in nondegenerated fully sex-linked regions (Natri et al. 2013; Schultheiß et al. 2015; Gammerdinger and Kocher 2018), similarly to analyses aimed at discovering genes affecting other phenotypes. The sex-determining region can potentially be narrowed down to a physically small part of a chromosome, as in many fish species (Kamiya et al. 2012; Myoshio et al. 2015; Gammerdinger and Kocher 2018; Conte et al. 2019). Figure 3 shows the wide range of recent size estimates in flowering plants.

When the Y- and X-linked regions can be distinguished, this can reveal heteromorphism in species whose chromosomes are too small for cytological detection. This approach can also detect “micro-heteromorphism,” when the two haplotypes of a physically small sex-linked region differ by rearrangements, such as inversions, as in papaya (Wang et al. 2012), and/or show deletion/duplication differences (Fig. 5 below shows one example).

Studies of the physical sizes of fully sex-linked regions also provide information about the numbers of “ancestral” (X- or Z-linked) genes. Genetic degeneration can then be quantified as the proportion of genes that remaining on the Y or W chromosome, and the proportion that still appear to be functional. These important data are currently available only in humans (Sayres and Makova 2013) and *Drosophila* (e.g., Zhou and Bachtrog 2015), and a few plants (Papadopulos et al. 2015; Wu and Moore 2015; Fruchard et al. 2020; Prentout et al. 2020).

**The Ages of Sex-Linked Regions, Including Nonrecombining Regions, When Present**

As already explained, the different ways in which sex-determining genes can appear, and different evolutionary times
since they originated (Fig. 2), can, in principle, explain much of the otherwise puzzling differences in differentiation between Y- and X-linked regions. It is therefore important to estimate ages of such regions, when they are present, and to test whether some sex-determining regions have simply not had enough time for suppressed recombination to evolve.

The appearance of a sex-determining locus on a chromosome defines its oldest fully sex-linked region. A single sex-determining gene is necessarily sex-linked, and, if the alleles controlling the two sexes differ by a single mutation, this mutation is completely sex-linked. Other sequence variants in the gene might, however, recombine (albeit with a low recombination rate, as they are physically close). Such variants will show incomplete associations with the sexes, as in fugu; in this fish, recombination must occur often enough to separate variants in the 17.5-kb region identified (Kamiya et al. 2012).

Many fully sex-linked regions include multiple genes. This could be because the sex-determining locus evolved in a non-recombining genome region carrying other genes (Fig. 2). In species where the region was ancestrally nonrecombining, the time when the sex-determining locus appeared defines the age. However, recombination may have become suppressed subsequently (see the next section).

AGE ESTIMATES USING PHYLOGENETIC DATA
The ages of a sex-linked region in a set of related dioecious species can be estimated from a phylogeny based on divergence of sequences from those of their closest nondioecious relatives. However, this relies on assuming that the dioecious species share the same oldest fully sex-linked region. When turnovers have occurred, replacing one sex-determining region with a new one, this approach could overestimate ages. This creates problems for estimating the ages of single-gene sex-determining systems, where approaches using a nonrecombining genome region (see the next section) are unavailable. To exclude turnover events, evidence is needed that all the species share the same sex-determining gene, in the same genomic location. In fish taxa, small sex-determining regions are often found in different physical locations, or different chromosomes (e.g., Kamiya et al. 2012; Myosho et al. 2015; Iida et al. 2018), and in Diptera (Post 1985; Mahajan and Bachtrog 2017; Meisel et al. 2020). Among flowering plants, it is currently unclear which species have old-established yet small sex-determining regions, and consequently it is not known how often small regions fail to evolve into large recombinationally suppressed regions.

Y-X DIVERGENCE ESTIMATES
Once Y-linkage is established, by whatever process, Y sequences will start diverging from their X counterparts, providing direct information about the time since Y-X recombination stopped (Fig. 1). If sex-linked genes have been ascertained in a dioecious species, one can use Y-X sequence divergence to test whether the Y became isolated before the split from the most closely related nondioecious species (Lawson-Handley et al. 2004; Dixon et al. 2018). Even within a single dioecious species, Y-X and W-Z divergence estimates reflect the times when recombination stopped. However, some points should be noted. First, times should ideally be estimated in terms of synonymous site divergence ($K_s$) values, as such sites are often weakly selected (although their divergence is not completely neutral, because selection opposes substitutions in some synonymous sites, e.g., Chamary et al. 2006; Parmley et al. 2006; Walsh et al. 2020). $K_s$ is nevertheless roughly proportional to the natural evolutionary time unit, the number of generations (at least until many sites may have undergone multiple substitutions, an effect termed “saturation,” illustrated in Fig. 1). Second, the common practice of translating synonymous site divergence values into estimated numbers of years should be avoided, as it precludes comparisons between species with different generation times (which are often poorly known).

There are also difficulties. In functional genes, selective constraints acting on many nonsynonymous sites make divergence initially slower than for synonymous sites. Although this obviates the need to correct for saturation, this advantage is outweighed because these estimates are confounded with degeneration, as nonsynonymous substitutions increase as genes lose functions (Fig. 1). Estimating divergence using all sites, without distinguishing between synonymous and nonsynonymous sites, is therefore problematic. It is preferable to separate genome sequences into coding regions, and to estimate synonymous site divergence.

Furthermore, highly degenerated sex-linked regions often contain few genes, making Y-X or W-Z divergence estimates unreliable or impossible, for example, in Drosophila species (except when fusions with autosomes in species without recombination in males have created neo-Y regions carrying many newly Y-linked genes; see Bachtrog et al. 2008). Old sex-linked regions with low gene density and high repeat density may even remain undetected unless the complete genome can be assembled and sex differences in coverage assessed (see above). A further problem arises when an XY male genome sequence is assembled using a female reference assembly (XX, avoiding assembly problems with diverged Y-linked sequences). If the Y-linked region includes genes not present on the X chromosome, such reference-based assembly will either leave these sequences unmapped or incorrectly map them to the most similar autosomal sequences. Reference-quality genome sequences of both sexes are therefore needed (Wei and Bachtrog 2019; Xue et al. 2020).
**EVOLUTIONARY STRATA**

Y-X divergence estimates have revealed that, even in species with long-established sex-linked regions, parts of these chromosomes sometimes also subsequently stopped recombining, forming so called “evolutionary strata,” first detected in humans (Laht and Page 1999; Skaletsky et al. 2003). This observation is important, because it shows that recombination has undergone an evolutionary change, rather than being the ancestral state for the region. The oldest human strata (Y\(\times\)X synonymous site divergence, \(K_s\), much higher than 20\%) are shared with other mammals (Sandstedt and Tucker 2004; Cortez et al. 2014), and include large numbers of ancestrally X-linked genes (369, 84, and 128; see Sayres and Makova 2013). Smaller nonrecombining strata adjacent to the PAR boundary are younger, with \(K_s\) around 20\%, and some are specific to only some mammal lineages. Strata with multiple genes are also found in birds (Zhou et al. 2014), snakes (Schield et al. 2019), and plants (Bergero et al. 2007; Wang et al. 2012; Zhang et al. 2020). However, Y-X and W-Z divergence estimates remain scarce.

**OTHER CHANGES AFTER RECOMBINATION BETWEEN THE SEX CHROMOSOME PAIR BECOMES SUPPRESSED**

Regions of suppressed recombination carrying new sex-determining genes also start accumulating Y-linked mutations causing genetic degeneration, repetitive sequences (reviewed by Bachtrog 2008), and adaptive changes, including the evolution of dosage compensation in response to degeneration (e.g., Mank et al. 2011; Disthe 2016; Gu and Walters 2017). Gene movements into the sex-linked region may also occur, or from the sex-linked region to an autosome (Bellott et al. 2010). These changes also accumulate over evolutionary times (Fig. 1), but probably do not greatly affect the sizes of nonrecombining regions (although more detailed future studies are needed). Their rates depend on many factors other than evolutionary times, and do not estimate such times. These changes can, however, be used to infer changes in recombination in nonmodel species where sex linkage cannot be tested genetically, or in old, highly degenerated systems, where divergence time estimates are not possible (see Fig. 1). For example, similarly to the phylogenetic approach described above, analysis of sex differences in coverage revealed degeneration in Schistosome lineages, initially in an old fully sex-linked region, and later independently in adjacent, but different, regions in two derived lineages (Picard et al. 2018). Clearly, recombination became suppressed in several distinct events.

Similarly, accumulation of repetitive sequences compared with a suitable “outgroup” suggests that a species has evolved a new nonrecombining region. Such changes may be very fast once recombination stops (Charlesworth et al. 1994). Prominent accumulation of repetitive sequences is detected in recently evolved *Drosophila* neo-Y chromosomes (Bachtrog 2003), grasshopper neo-Y-linked regions (Palacios-Gimenez et al. 2020), and before loss of genes in plants including papaya (Wang et al. 2012), and in lizards (Matsubara et al. 2014).

**Genetic Degeneration**

Although the extent of genetic degeneration increases with the time a region has been evolving under full sex linkage, theoretical modeling has identified other important factors (reviewed by Bachtrog 2008). Degeneration rates may therefore differ greatly between different organisms. Together with the scarcity of quantitative degeneration data and divergence time estimates, this contributes to the seemingly confusing picture mentioned above. Many studies describe depth of coverage ratios in the two sexes, which merely detects regions with degenerated sequences. Few indicate the proportion that are hemizygous in males, and the number of XY gene pairs whose Y copy is a pseudogene, and species with partially degenerated sex-linked regions or strata have been little studied.

Testable predictions are nevertheless available. First, most models predict that degeneration will be faster in sex-linked regions with many genes (although a recent model predicts degenerate regions with few genes; Lenormand et al. 2020). Estimates of numbers of “ancestral” genes should allow tests of these ideas. Single-gene systems, and small chromosomes that acquire a sex-determining gene, such as microchromosomes of lizards (Matsubara et al. 2014), might be expected to degenerate slowly, and data from such nonmodel species should become available.

Second, degeneration has a nonlinear time-course. Genes are predicted to initially lose functions rapidly by major effect mutations, followed by slower changes, and eventually deletion of sets of genes (Fig. 1). Data from sex-linked regions at all degeneration stages are therefore needed. Plants, which include many species with small or young sex-linked regions, may be less suitable than animals, because selection in the haploid phase, including the pollen of flowering plants, may oppose degeneration (Bergero and Charlesworth 2011; Chibalina and Filatov 2011; Hough et al. 2014). However, considerable degeneration has been documented in several plants (see above), so data from plants are still needed.

At all stages of degeneration of a nonrecombining region, the rates also depend on the specific properties of the genes present (e.g., Kramer et al. 2016; Rifkin et al. 2020; Bellott and Page 2021). A striking example is the neo-Y of *D. busckii*, which is more degenerated (with 58\% nonfunctional genes) than the larger and older one in *D. miranda* (only 34\% nonfunctional genes), probably because the latter evolved from a “dot” chromosome, whose genes show low selective constraints (Zhou and Bachtrog 2015).
The data currently available suggest that most animal strata with Y-X or W-Z $K_s$ values above 20% show essentially complete degeneration of most ancestral genes. With $K_s$ below this value, 50% or more of the ancestral genes present on the X are generally also present as likely functional copies on the Y, consistent with theoretical predictions (Bachtrog 2008). However, the $K_s$ level and evolutionary time needed for strata to reach the stage of major gene loss, and for de novo evolution of dosage compensation, remain unclear. Deletions within fully sex-linked regions, contributing to heteromorphism, probably occur only in the late stages of degeneration, as large deletions are generally highly deleterious (Bull 1983; Manna et al. 2012; Bazrgar et al. 2013), unless the genes are all under weak selection, or the region has already degenerated and become a “gene desert” (Nóbrega et al. 2004).

### Why are Sex-Determining Regions Often Nonrecombining?

I next outline different situations that can account for lack of recombination in sex-determining regions. First, I describe two situations that involve selection for reduced recombination in systems with polymorphisms for a sex-determining gene and a second gene. Collectively they can be termed “the sexually antagonistic polymorphism hypothesis.”

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**Figure 4.** Situations where selection favors recombination suppression. (A) When the ancestral population is cosexual, females can arise by mutations in a male-function gene ($M \rightarrow m$), and males by a femaleness-suppressing mutation ($F \rightarrow SuF$) in the same genome region; the Y haplotype will then carry $M$ and $SuF$. This is a “supergene” model for sex determination. (B) In a situation after a turnover event that creates a new maleness ($M$) factor in an ancestral population of males and females, the $M$ factor defines a Y haplotype, and a sexually antagonistic maleness-enhancing factor, $A$, establishes a polymorphism in the same genome region. (C) In either case, recombinants between the two haplotypes will have disfavored combinations of alleles at the two loci, and a nonrecombining region (possibly extensive) could evolve.

**APPEARANCE OF NEW SEX-DETERMINING REGIONS: TWO-GENE SYSTEMS**

A major hypothesis to explain why XY and ZW chromosome pairs do not recombine involves selection for closer linkage between two polymorphic loci. One such situation (Fig. 4A) arises when separate sexes evolve de novo in an ancestrally cosexual species (for instance dioecious flowering plants that evolved from ancestors with hermaphrodite flowers, or in monoecious species, with each individual producing both male and female flowers). The evolution of females and males requires two mutations. In one model (Charlesworth and Charlesworth 1978), the two mutations are in separate genes, and each mutation causes sterility of one sex, and therefore acts antagonistically in the other sex. As reviewed by Westergaard (1958), dioecy in plants often involves a recessive loss-of-function male-sterility mutation that produces the females, and a femaleness-suppressing mutation that produces males (Fig. 2A); hypothetically both mutations could occur in a single gene, although no example has yet been reported. Sex-determination then resembles supergenes controlling other polymorphisms that probably involve separate polymorphic mutations affecting different traits (Schwander et al. 2014; Charlesworth 2016). Unless the second mutation acts specifically only in one genotype of the first gene, loose linkage prevents establishment of a polymorphism. However, if a closely linked mutation does establish a polymorphism, tighter linkage may evolve.
SEXUALLY ANTAGONISTIC POLYMORPHISM NEAR A SEX-DETERMINING GENE

Figure 2B illustrates the other plausible biological situation involving a two-gene polymorphism: when a sexually antagonistic (SA) polymorphism establishes in a gene closely linked to a sex-determining gene (Kirkpatrick and Guerrero 2014), or a turnover event involves a sex-determining gene appearing near a SA polymorphism (Bull 1983; van Doorn and Kirkpatrick 2007). As mentioned above, turnovers are well documented in animals and plants. The first of these situations can also arise after de novo evolution of separate sexes has generated a single sex-determining gene. Figure 5 illustrates a naturally evolved case in the persimmon, Diospyros lotus; in this plant, an active (genetically dominant) maleness factor has evolved; females are the “default” sex, developing only when the maleness gene is absent (Akagi et al. 2014; Akagi and Charlesworth 2019).

When the heterogametic sex is achiasmate (or genome regions that undergo crossovers are strongly localized in one sex), a new sex-determining gene can define a new fully sex-linked region. Such male-specific crossover patterns are observed in distantly related fish, including the guppy (Bergero et al. 2019) and the stickleback (Sardell and Kirkpatrick 2019), and in frogs (Rodrigues et al. 2018). As reviewed recently (Charlesworth 2019), they may be commoner than is currently realized, as genetic maps are rarely estimated separately for the two sexes, and physical maps, to locate the genome regions where crossovers occur, are available from only a few species.

CAVEATS CONCERNING THE SA POLYMORPHISM HYPOTHESIS

Although both situations in Figure 4 generate selection for closer linkage with the sex-determining gene, a response to this selection will not happen unless heritable variation exists for recombination rates in the region. Second, the selection and dominance coefficients under which SA polymorphisms can be maintained are restrictive (Fry 2010). In theoretical studies with given selection and dominance coefficients, maintenance is most likely in regions very closely linked to sex-determining loci (Jordan and Charlesworth 2012; Kirkpatrick and Guerrero 2014). Therefore, even though mutations with SA effects can probably occur in many genes across the genomes of dioecious animals and plants (Connallon and Clark 2014), the numbers of genes in which SA polymorphisms could potentially be established are probably small (unless a sex-determining locus evolves within a large ancestrally nonrecombining region). The waiting time until a suitable SA polymorphism becomes established will therefore often be long. The same applies to the model in Figure 4A (Charlesworth and Charlesworth 1978). However, if linkage is already close, closer linkage will evolve slowly (although an inversion preventing crossovers could nevertheless establish an extensive new Y-linked region; see Fig. 2C).

On the other hand, SA conflicts can be resolved by evolving sex differences in gene expression, and need not lead to suppressed recombination. Vicoso et al. (2013) suggested that this might account for the persistence of large partially sex-linked regions in the Paleognathous birds (whereas recombination has become suppressed in the homologous regions of the ZW pair of Neognathous birds). Sex-biased expression has not been confirmed for emu partially sex-linked genes (Xu et al. 2019).
Nevertheless, the idea is plausible, as conditions for expression level changes are less stringent than for changes in linkage, because there is no requirement for a SA polymorphism to be maintained. Indeed, after fixation of an allele that benefits one sex, but reduces fitness in the other, sex-specific control of its expression is favored. Overall, therefore, chromosome regions carrying sex-determining genes are not always expected to evolve suppressed recombination.

Surprisingly few studies have examined the evolution of sex limitation of expression, although expression differences are detected at many loci. Genes encoding testis or egg proteins, or anther-specific proteins in plants, inevitably have sex-specific expression, but many genes expressed in other tissues might change from being expressed in both sexes, to having sex-biased expression if conflicts have evolved. Changes producing over- or underrepresentation of genes with sex-biased expression (termed “masculinization” and “demasculinization” of sex chromosomes) have been documented, and may reflect the different balance of selection on mutations with sexually antagonistic effects (as reviewed, e.g., by Vicoso and Charlesworth 2006; Bachtrog et al. 2010; Mank and Wright 2012).

ALTERNATIVES TO THE SA POLYMORPHISM HYPOTHESIS
As Figure 2 shows, the SA polymorphism hypothesis is only one possible explanation for suppressed recombination in sex-determining regions. I have already mentioned that a large fully sex-linked region could evolve without any selection favoring close linkage if a sex-determining gene arises within a region that already had a low recombination rate. Large “recombination deserts” are known in many plant genomes (Charlesworth 2019), and the sex-determining locus is within such a region in a species of the plant Rumex (Rifkin et al. 2020). Such a situation would also facilitate the establishment of SA polymorphisms because many genes within such a region will be closely linked to the sex-determining locus.

SMALL NONRECOMBINING REGIONS
At least two other possibilities can create small nonrecombining regions, again without selection for loss of recombination. An insertion causing “micro-heteromorphism” (such as the duplication creating the nonpaired maleness factor in Fig. 5) might directly prevent pairing in the regions flanking the insertion, causing a few neighboring genes to also become fully Y- or X-linked (pink). Single-gene systems in poplar species appear to involve a male-determining gene that, like that in the persimmon, arose as a duplication that silenced a female-promoting gene (Müller et al. 2020; Xue et al. 2020). Small genome regions carrying multiple male-specific genes have been detected in two other plants, Asparagus officinalis (Harkess et al. 2020) and the date palm (Torres et al. 2018). A duplication also created a new sex-determining gene in the fish, medaka, Oryzias latipes (Kondo et al. 2006). These male-specific regions have no homologous counterpart with which to pair (formally they are hemizygous fully Y-linked regions, but distinct from the hemizygosity caused by genetic degeneration, where genes were lost from a Y-linked region).

Finally, the observation of slight changes in the location of the boundary between the completely and partially sex-linked (pseudoautosomal or PAR) regions suggests that small new sex-linked regions may arise in another manner. Two sets of genes (totaling only 17 X-linked genes; Sayres and Makova 2013) have become completely sex-linked genes in humans, but are PAR genes in other mammals, and the PAR boundaries in different mammals also differ slightly (Van Laere et al. 2008; Skinner et al. 2013), including among mouse species (White et al. 2012; Morgan et al. 2019). In plants, similar variation has been found between strains in the plant Carica papaya (Lappin et al. 2015), and in Silene latifolia, the boundary is not sharply defined, also suggesting a recent change (Krasovec et al. 2020). The reasons for such shifts are unknown, but heterochromatic regions enriched for repetitive sequences that evolved after a recombination-suppressed region evolved might create micro-heteromorphism or actively inhibit crossing over in large fully sex-linked regions (Phillips and Ihssen 1985; Charlesworth et al. 1994). I am not aware of any explicit model or empirical evidence supporting such an idea, but detailed studies of PAR boundary changes may help understand such changes.

Conclusions: Why Do Some Sex Chromosome Pairs Remain Homomorphic?
I have argued that sex-determining regions need not necessarily evolve into large, multigene nonrecombining regions, although young sex-linked regions with small Y-X divergence values can sometimes be extensive. Large suppressed recombination regions that still retain many Y-or W-linked genes, particularly those with more than a single stratum of divergence time, support the long-accepted view that the appearance of sex-determining genes has repeatedly triggered such changes in a remarkable diversity of different types of organisms, leading to differentiation. However, very recently evolved systems will initially be homomorphic (unless recombination is suppressed and there has been enough time for repetitive sequence accumulation), and some such chromosome pairs may continue to undergo crossovers.

The evolution of heteromorphism is not expected to be clock-like, or even a monotonic change. Systems old enough to have undergone genetic degeneration may sometimes remain homomorphic, as in the recently discovered case of skinks (Kostmann et al. 2021). The frequency of such cases is not yet
clear, nor why heteromorphism is lacking. Some such situations may reflect technical difficulties in detecting differences in physically small chromosomes, and others could be taxa that (for unknown reasons) rarely undergo chromosome rearrangements, or where major repetitive sequence accumulation has not happened, or secondary loss of heteromorphism (see above).

In contrast, genetic degeneration is an inevitable decline, starting when recombination stopped (although different degeneration levels cannot yet be explained in precise quantitative terms). Although neither the proportion of species where extensive reduced recombination regions evolved in response to the presence of a sex-determining locus nor the role of selection in such changes (the SA polymorphism hypothesis) is yet fully understood, it is already clear that the times when sex-determining regions stopped recombining explain much of the variation in degeneration levels.

Chromosome rearrangements such as inversions may often be involved in recombination suppression. However, their presence does not demonstrate that suppressed recombination has evolved, as this need not involve rearrangements, and, rearrangements readily spread after recombination is suppressed. Moreover, a Y- or W-linked region’s lack of recombination in might reflect rearrangements that spread by genetic drift in a small population (reviewed by Ironside 2010; Ponnikas et al. 2018). This cannot explain an overrepresentation on sex chromosomes unless some further mechanism creates a higher rearrangement input rate on these chromosomes (such as the remarkable apparent difference in the Dipteran blackflies; see Adler et al. 2016). Unfortunately, comparative tests of whether chromosomes carrying sex-determining loci have a special tendency to subsequently evolve nonrecombining regions are hampered by a reporting bias: sex chromosome differences are readily detectable cytologically, without laborious surveys of all chromosomes in multiple individuals of a species. Even work to test whether sex chromosome rearrangement polymorphisms are commoner than autosomal ones is only just beginning (Anderson et al. 2020). Genome sequencing can potentially give unbiased information for such tests. However, such data are not yet easily obtainable in nonmodel species, as multiple individuals of both sexes are still required to discover sex-specific variants. Cytogenetic information can help choose suitable species for genome sequencing targeted at testing ideas such as the neutrality hypothesis, and classifying sex-determining regions into the types suggested here. In turn, results from the different types should lead to a better understanding of evolutionary changes in sex-determining loci and regions, and their timings.

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