The model role of a non-model organism

The genus *Silene* (Caryophyllaceae), with a tradition of genetical and ecological studies dating back to Mendel and Darwin, has remarkably many interesting features. First, the species in the genus vary widely in their breeding systems and ecology. Second, several members of this mainly holarctic genus can be easily bred, and have short life cycles, and are thus convenient for experimental and field studies. Genomic resources are now becoming increasingly available in *Silene*, making genetic, quantitative genetic and molecular studies possible. A strength of *Silene* as a model system, compared with many classical model organisms, is that researchers can rely on a large number of ecological studies encompassing biotic interactions with sexually transmitted fungi, pollinators and herbivores. It is the wealth of ecological and other earlier knowledge that makes the genus *Silene* important for studying many biological questions, including the suppression of recombination during sex chromosome evolution, sexually antagonistic selection in an organism that is not an animal, epigenetic processes in flower development, speciation and reproductive isolation, multi-trophic interactions, disease ecology and biological invasions.

Thanks to the classical genetic and ecological work on *Silene*, new progress can now be made in studying some of these important unresolved questions in biology with the aid of modern molecular tools. That new model systems with accessible and well-studied ecology open fruitful avenues for investigation is illustrated also by *Mimulus* (Wu et al., 2008) and the ongoing efforts to develop genomic resources in an increasing number of different systems. In the following, we outline the unique features of the *Silene* system, and describe active research areas and future directions, highlighting new advances and work in progress.

Evolution of sexual systems

Plants show remarkable diversity in their sexual and mating systems, ranging from hermaphroditism to dioecy, from self-incompatible hermaphroditism to hermaphroditism in which individuals possess a capacity to self-fertilize, but which often show complex mechanisms to promote outcrossing (Darwin, 1877; Barrett, 2002). Understanding the origin and maintenance of this diversity has been a central goal of plant evolutionary biology, and much can be learned from studies that compare closely related species or populations that vary in their sexual system. The genus *Silene* was recognized since the early days of evolutionary biology as important for studying sexual and mating systems. The genus includes members that are self-fertile hermaphrodites, some of which are cleistogamous, self-incompatible hermaphrodites, but also sexually polymorphic species
expressing dioecy (separate sexes) and unusual systems such as gynodioecy (polymorphism for females and hermaphrodites) (Darwin, 1876, 1877; Barrett, 2002). Indeed Silene (then Lychnis) was used by Darwin to illustrate dioecy as a state derived from ancestral hermaphroditism (Darwin, 1876, pp 410–411) and Silene has continued to be central in studies of sexual and mating system evolution (Baker, 1963) and more recently even evolution of sex determination systems because some of the dioecious species of this genus have even evolved heteromorphic sex chromosomes (see below).

DNA sequence data allow reliable phylogenetic reconstructions in many taxa of biological interest, including Silene (Oxelman et al., 1997). These reveal that dioecy evolved independently at least twice in the genus (Desfeux et al., 1996). In the clade that includes Silene latifolia, sex chromosome heteromorphism has also evolved (see below, (Armstrong and Filatov, 2008); Figure 1), whereas in another dioecious clade, Silene colpophylla has no heteromorphic sex chromosomes, and genes that are sex-linked in S. latifolia are not linked to the S. colpophylla sex-determining locus (Mrackova et al., 2008). Dioecy probably evolved through gynodioecy (Maurice et al., 1993; Schultz, 1994), therefore the common occurrence of gynodioecy in the genus Silene has long been of interest. Nucleo-cytoplasmic inheritance of male sterility (interaction between cytoplasmic male sterility (CMS) factors and nuclear restorers of fertility) was established by classical genetic work (see below), and theoretical work has shown how intra-genomic conflict between maternally and biparentally inherited genes can lead to the maintenance of such gender polymorphism. In some Silene species, male-sterile individuals are rare, and some females produce perfect hermaphrodite flowers (Desfeux et al., 1996). In other species, females are very common (Shykoff, 1988; Delph and Mutikainen, 2003).

In theory, nucleo-cytoplasmic gynodioecy might sometimes lead to the evolution of separate sexes (Maurice et al., 1993; Schultz, 1994). Empirical tests may, in the future, be possible in Silene. It is not yet clear whether, or how, quantitative variation in sex functions relates to the evolution of fully separate sexes, but Silene species are excellent for quantitative genetic studies, having a fairly rapid life cycle and relatively small adult size. Current topics of quantitative genetic studies in Silene are mainly focussing on the evolution of sexual dimorphism and sexually antagonistic selection (see below). Also in this context, it is particularly the dioecious Silene that provide a test case for comparing shared/parallel evolutionary processes in plant and animal systems.

**Figure 1** (a) Male mitosis prepared from germinating Silene latifolia seeds. (b) Meiotic metaphase I in pollen mother cells of Silene latifolia. Although the autosomes are still pairing, a precocious separation of the X and Y chromosomes begins. Chromosomes are counterstained with 4',6-diamidino-2-phenylindole (DAPI); the sex chromosomes (X and Y) are indicated. Scale bar: 10 µm (B. Vyskot).

**Sex chromosome evolution**

The existence of species with sex chromosomes within the broad variation of sexual systems found within the genus allows comparison with other well-studied sex chromosomes (for example, those of mammals) for investigating the generality of processes acting during sex chromosome evolution. Sequence and genetic mapping data have shown that the sex chromosomes in S. latifolia and S. dioica evolved recently from a pair of autosomes (Filatov, 2005a). Divergence between the sequences of homologous genes on the X and Y chromosomes indicates an age of <10 million years (Filatov, 2005a), yet marked chromosome heteromorphism has already evolved, with remarkable similarity to the human XY pair. In mammals, the X chromosome carries over one thousand genes and is mostly euchromatic, whereas the Y chromosome has few functional genes and is mostly heterochromatic. The sex chromosomes pair during male meiosis, but undergo genetic recombination only in small pseudoautosomal regions. Gene loss from the Y chromosome (‘genetic degeneration’) accompanied suppression of recombination in different regions of the mammalian and bird XY chromosome pairs, creating ‘evolutionary strata’. The oldest stratum includes highly diverged genes that ceased recombining long ago, whereas genes in the youngest stratum are much less diverged (Lahn and Page, 1999). Over the last decade, several S. latifolia sex-linked genes have been identified (for example, Matsunaga et al., 2003; Filatov, 2005b), and they reveal that Silene sex chromosomes also have evolutionary strata with different divergence levels, indicating that recombination between these chromosomes also stopped in a step-wise manner over time (Nicolas et al., 2005; Bergero et al., 2007).

The recently evolved Y chromosome of S. latifolia is more suitable than the ancient, highly degenerated, human Y chromosome for studying the evolutionary processes causing the loss of functional genes from the Y chromosome. S. latifolia Y genes show signs of degeneration,
including fast protein evolution, mutations damaging sites known to be important for protein activity (Filatov, 2005c), low expression level and insertions of transposable elements in introns (Marais et al., 2008) and severely reduced polymorphism, an indirect effect of the processes that cause degeneration (Filatov et al., 2000). Cytogenetic analysis of *S. latifolia* chromosomes shows accumulation of repeat sequences on the Y chromosome (Kubat et al., 2007).

The numbers of genes on the *Silene* sex chromosomes are still unknown, but probably greatly exceed the 10 currently known sex-linked genes. The *S. latifolia* genome is very large (its DNA content is similar to that of the human genome), consistent with a high repetitive content throughout the genome (Cermak et al., 2008). Sequencing and assembling this genome is currently infeasible, but partial sequencing is possible using BAC libraries, and comparative analysis of genomic DNA sequences from the X and Y chromosomes can potentially provide estimates of gene loss from the Y chromosome. Expressed sequence tag databases are also being developed, and will provide genetic markers for mapping the sex chromosomes, for fluorescent in situ hybridization (Armstrong and Filatov, 2008) and for comparing expression differences between X- and Y-linked alleles (Marais et al., 2008). Expression studies may also help to identify sex-determining genes in *S. latifolia*, about which we still know very little, despite having some knowledge of the pathways involved in the floral developmental programs in *Silene* (Zluvova et al., 2007) and the Y chromosomal regions involved in male-determination (Lardon et al., 1999; Lebel-Hardenack et al., 2002; Bergero et al., 2008).

The young age of the *Silene* sex chromosomes may provide insights into another very interesting set of questions concerning how and why X–Y recombination is suppressed. As in mammals, recombination between the *Silene* X and Y chromosomes is restricted to small ‘pseudoautosomal’ regions (Figure 2). A plausible hypothesis for the evolution of the non-recombining regions is that, as sex chromosomes evolve, sexually antagonistic genes (enhancing male fertility, but with negative effects in females) accumulate, particularly on Y chromosomes (see also below, ‘sexual dimorphism and sexual antagonism’). Polymorphism for sexually antagonistic alleles, maintained by such conflicts, creates selection against recombination (which would move such alleles onto the X). A young sex chromosome system in which genetic mapping is possible offers opportunities for testing for such genetic variants, which have not yet been identified in any organism; studies in *Silene* have a promise for finding such genes. Mammalian Y chromosomes are rearranged, and do not permit inferences about whether non-recombining regions evolved through inversions suppressing recombination, but data in *Silene* are suggesting that inversions alone are not responsible (Zluvova et al., 2005; Bergero et al., 2008).

**Reproductive isolation and species boundaries**

Studies on animals have established that sex chromosomes play a major role in the evolution of reproductive isolation between species, but little is known about plants, and *Silene* is ideal for investigating this. In animals, ‘two rules of speciation’ have been stated in relation to postzygotic reproductive isolation (Coyne and Orr, 1989). ‘Haldane’s rule’ summarizes the observation that, in crosses between species with chromosomal sex determination, when one sex is absent, rare or sterile, it is usually the heterogametic sex. ‘The large X-effect’ is the frequently large role of the X chromosome in hybrid sterility (Coyne and Orr, 1989).

As sex chromosomes evolved recently in *Silene*, we may be able to discover genomic regions associated with reproductive isolation in related species pairs with sex chromosomes. This will allow direct tests of whether genes and coevolved gene complexes involved in reproductive isolation have accumulated on the sex chromosomes.

Recombination is suppressed across most of the *Silene* Y chromosome and the X chromosome recombines only in females. Recombination thus cannot disrupt co-adapted gene complexes on the Y chromosome and is less likely to do so on the X chromosome compared with the autosomes. Gene introgression after hybridization between cross-compatible species is thus predicted to be lower for the Y chromosome than the X chromosome (and should be highest for the autosomes). Indeed, isolation between *S. latifolia* and *S. dioica* (assessed by FST for DNA sequence variants in two genes), is significantly stronger on the non-recombining Y chromosome, compared with the recombining X chromosome, even taking into account the lower within-population variability of Y than X sequences (Ironside and Filatov, 2005).

Sex chromosomes are not the only genetic factors in *Silene* that may link sex determination and reproductive isolation. Gynodioecy, the coexistence of hermaphrodite and female individuals, is probably ancestral in the genus (Desfeux et al., 1996) and is typically controlled by interactions between mitochondrial CMS factors and nuclear fertility restorers. CMS factors can also influence compatibility between species through the loss of coordination between nuclear and organellar genes inherited from different parents (Chase, 2007). The species-rich genus *Silene* provides an excellent system

**Figure 2** Evolutionary strata in *Silene latifolia*. The synonymous divergence between X and Y copies (y axis, % d,X–Y) and the position of the X copy of the X chromosome genetic map (x axis) are shown for the 10 known sex-linked gene pairs (circles). This plot shows that at least three events of recombination suppression between X and Y have occurred in *S. latifolia*. The dates of these events calculated assuming molecular clock (1.8 × 10⁻⁸ syn. substitutions/syn. site/year) are shown. Adapted from Nicolas et al. (2005) and Bergero et al. (2007).
for assessing the contribution of nuclear–cytoplasmic interactions to reproductive isolation between species.

Epigenetic processes

The existence of sex chromosomes also raises questions about dosage compensation of X-linked genes for maintaining correct gene expression levels in XY-male plants and suppression of pistil development in flowers of male plants. The role of epigenetic processes in the control of gene and sex expression is thus also being studied in the dioecious *S. latifolia*.

If the *S. latifolia* Y chromosome is genetically degenerated, gene dosage may be compensated, either by silencing of one X chromosome copy in females (as in mammals) or by increasing the activity of the single X chromosome copy in males (as in *Drosophila*). Cytogenetic experiments indicate reproducible changes in chromatin structure of the *S. latifolia* sex chromosomes (Vyskot *et al*., 1993), including DNA replication timing differences and methylation differences between the two X chromosomes in female somatic cells, reminiscent of the mammalian dosage compensation mechanism (lyonisation), which also involves DNA methylation (Figure 3). With the availability of sex-linked *S. latifolia* genes, it becomes possible to test whether dosage compensation has also evolved in dioecious plants.

Epigenetic processes represent mechanisms by which gene expression can be reversibly suppressed. In large plant nuclear genomes, such as *S. latifolia* (2C ~ 5.8 pg DNA), up to 30% of all cytosine residues are methylated, both in CpG doublets and in CpNpG triplets. A simple treatment of plant meristems with a cytosine analog, 5-azacytidine, can dramatically reduce the nuclear genome 5-methylcytosine content. It is surprising that the only phenotypic change observed with this treatment in male *S. latifolia* was sex reversal, manifested as frequent bisexuality of flowers in karyotypically male (XY) individuals (Figure 4) (Janousek *et al*., 1996). The inflorescences of these plants are mosaics of staminate and perfect flowers, consistent with epigenetic control (Vyskot, 1999). The effect is heritable, and the ‘epimutation’ involved showed Y-linked inheritance in reciprocal crosses with wild-type plants. 5-azacytidine blocks DNA methylation, and generally activates genes. In this case, however, it inactivates a Y-linked female suppressor function, perhaps through DNA overall hypomethylation-induced silencing. Frequent sex reversal from male to hermaphroditism was also observed when apical meristems were treated with trichostatin A, an inhibitor of histone deacetylase (Negrutiu *et al*., 2001), supporting the possibility of epigenetic control of sexual development in the dioecious *S. latifolia*.

Sexual dimorphism and sexual antagonism

Yet another consequence of having separate sexes in several *Silene* species is sexual dimorphism (phenotypic differences between males and females), which is widespread in gonochoristic animals and dioecious plants (Geber *et al*., 1999). Its occurrence suggests that selection acts on traits in a sex-specific manner. However,
homologous genes that are expressed in both sexes lead to overall positive genetic correlations between the sexes, and this can prevent the evolution of sexual dimorphism and lead to sexually antagonistic gene expression (Bedhomme and Chippindale, 2007).

Dioecious *Silene* species are excellent for studying this phenomenon. As dioecy has arisen more than once within *Silene*, there are opportunities for comparative studies to test hypotheses about its evolution. Although dioecy is not rare among flowering plants, many dioecious species are woody, long-lived trees (Renner and Ricklefs, 1995), which makes genetic studies difficult, even in crop species. In contrast, several dioecious *Silene* species are short-lived perennials that often flower in their first year, allowing experiments to be done within a short time frame. Moreover, it gives the opportunity for comparing many life-history traits between the sexes, including longevity.

Good measures of sexual dimorphism for a variety of traits, together with the underlying genetic variation and covariation within and between the sexes, can be obtained by established quantitative-genetic crossing designs or artificial-selection experiments. Such experiments have shown remarkable sexual dimorphism and genetic correlations between the sexes for multiple fitness-related traits in *S. latifolia* (Meagher, 1999; Delph, 2007). Flower size, flower number and leaf traits have been extensively studied; these traits vary in their means and degree of sexual dimorphism among populations, and genetic correlations within and between the sexes are high (Steven et al., 2007). Sex-specific quantitative trait loci have been found for these traits, especially in males; the finding that the major ones are on the sex chromosomes (Scotti and Delph, 2006) suggests that the studies on the evolution of sexual dimorphism in this species may uncover examples of sexual antagonism.

Correlations between traits within the sexes can also constrain the evolution of sexual dimorphism. In *Silene* (and plants more generally), ecologists can often use proxies for the fitness of individuals in experimental and/or natural settings, such as flower and fruit production, and relate this to variation in traits of interest, such as sexually dimorphic traits. Phenotypic-selection experiments in populations that differ in their degree of sexual dimorphism have revealed the extent to which correlations between traits within the sexes constrain the evolution of sexual dimorphism. Another way to study how trait variation affects fitness is to plant experimental arrays of individuals with extreme phenotypes and with known genetic markers, and test the different phenotypes’ performance over time. Given the important interspecific interactions between various dioecious *Silene* species and their pollinators/seeds predators and pathogens, it will be interesting to see how the various players contribute to the existence of sexual antagonism.

**Organelle DNA**

Organelle DNA is known to play a role in sex determination in *Silene* and the properties of the chloroplast and mitochondrial genomes of several *Silene* species have recently received considerable attention. Interest in the mitochondrial genome traces largely to the cytonuclear sex determination in gynodioecious *Silene*. Cytonuclear gynodioecious systems are a prime example of genetic conflict. When the inheritance of CMS elements is maternal (as is usually assumed for mitochondrial genes), evolutionary conflicts occur between the fitness interests of the mitochondrial genome (more and/or better seeds produced by females enhances mitochondrial fitness) and those of the nuclear genome (higher transmission of genes through pollen by hermaphrodites enhances nuclear fitness). Recent studies of *Silene acaulis* and *Silene vulgaris* have shown that inheritance of the mitochondrial genome is not strictly maternal (Städdler and Delph, 2002; McCauley et al., 2005). The evidence is largely indirect and is on the basis of observations suggesting intra- and intergeneric recombination within the mitochondrial genome that is detectable when the recombining sequences differ, thus implying mitochondrial heteroplasmy through bi-parental inheritance, which has recently been documented in *S. vulgaris* (Welch et al., 2006). However, the considerable (though imperfect) linkage disequilibrium between mitochondrial genes (McCauley and Ellis, 2008) and a nearly perfect statistical association between mitochondrial and chloroplast genes (Storchova and Olson, 2004) suggests that such ‘paternal leakage’ is rare.

Another interesting property of *Silene* mitochondrial genomes is the high level of DNA sequence diversity in the coding regions of some species (Städdler and Delph, 2002; McCauley et al., 2007), and the high interspecific divergence of mitochondrial gene sequences of others.
These results contrast with the observation that coding regions of most plant mitochondrial genes are highly conserved. Differences in standing levels of variation and apparent differences in mutation rates between mitochondrial genes may be caused by rare mitochondrial recombination (Barr et al., 2007), but the high level of intra-specific variation sometimes observed is probably a consequence of long-term balancing selection inherent to persistence of CMS genotypes in gynodioecious systems (Städler and Delph, 2002). In *S. vulgaris*, however, there is suggestive evidence from chloroplast variation for frequent turnover of lineages owing to recurrent selective sweeps in the organellar genomes (Ingvarsson and Taylor, 2002). Theory shows that each of these contrasting outcomes is possible.

*Silene* chloroplast genes (cpDNA) have been used extensively as markers of seed movement at the metapopulation and phylogeographical levels. Given the potential importance of colonization events for determining the local genetic structure of plant metapopulations with local patchy distribution and frequent turnover, genetic markers that reflect seed movement (such as variable regions of cpDNA, assuming maternal inheritance) are particularly useful and can be compared with nuclear genetic markers for making inferences on the rates of gene flow through pollen and seeds (McCauley, 1994).

### Biological invasions

Beyond the study of mating systems, sex determination, their mechanisms and their consequences, the genus *Silene* has been instrumental as a biological model for other biological and ecological phenomena, such as biological invasions and trophic interactions. Biological invasions are increasingly being viewed as a major contributor to the loss of biodiversity around the globe, and *Silene* species are emerging as an outstanding system for research into this phenomenon. *S. latifolia* and *S. vulgaris* are native to Eurasia, but both are now widespread in parts of North America. The genetic diversity in the species’ native and introduced ranges must be determined by both the mode of invasion (a single versus multiple introductions from one or multiple origins), together with the genetic structure in the native range.

Analysis of cpDNA sequences indeed suggests that the differing phylogeographical histories of the two species in Europe affected their respective genetic structures in North America: in *S. latifolia*, few, discrete invasion events from strongly genetically structured populations in the native range resulted in genetic bottlenecks followed by admixture, whereas invasion in *S. vulgaris* involved more representative sampling of diversity, with little detectable admixture (Taylor and Keller, 2007).

A major question in biology is why invasive species that are relatively benign competitors in their native ecosystem become dominant after introduction. One explanation is that natural selection in the absence of enemies shifts the optimal investment of resources from defence to enhanced growth and reproduction. *S. latifolia* was introduced to North America approximately 200 years ago, and there are large differences in the species’ biology between the two continents: (1) *S. latifolia* has escaped some specialist enemies during colonization, and damage levels are higher in Europe (Wolfe, 2002); (2) North American plants are ‘weedier’ and more reproductively prolific (Blair and Wolfe, 2004), and they invest less in defence and have lower resistance to fungal infection and fruit predation (Wolfe et al., 2004). The development of genomic tools for *Silene* will soon allow dissection of the genetic changes associated with successful invasions.

### Biotic interactions: *Microbotryum* and the evolutionary ecology of disease

Another important field of ecology in which *Silene* is an important model is disease evolution. Many members of the genus *Silene* are attacked by anther-smut disease caused by the fungus *Microbotryum* (Figure 5a). This fungus invades the plant and produces spores in the anthers, which replace the normal pollen. The disease sterilizes its host and is transmitted when insects carry spores from infected to healthy flowers during pollination (Antonovics, 2005). In females of dioecious host species, anther development is induced by the pathogen, and this remarkable symptom caught the attention of early researchers including Darwin (1863) to be followed by a series of studies using the pathogen to understand the host’s sexual development (for example, Strasburger, 1871).
1900; Shull, 1910). Thus, utility of this disease for studying Silene continues to be shown and is enhanced by modern molecular approaches to gene expression during infection (Uchida et al., 2003; Kazama et al., 2005). Moreover, the smut diseases provide some of the strongest arguments for the value of ‘model systems’ in their own right. As early as 1755, Mathieu Tillet showed experimentally that grain smut of wheat was transmitted by spores produced in the infected kernels. Had the full implications of this work been realized by the broader scientific world, thousands of lives might have been saved by a much earlier acceptance of the germ theory of disease.

The Silene–Microbotryum system has become a model for studying the evolution of sexually transmitted diseases (Nunn et al., 2000), as well as for studying the forces responsible for host shifts (Antonovics et al., 2002), and the importance of genetic variation in the regional epidemiology of the disease (Antonovics et al., 1997). Ecologists have often ignored the impact of sterilizing and sexually transmitted diseases on the size and distribution of individuals in natural populations, and interesting parallels can be drawn between anther-smut disease and the impact of AIDS on human populations (Figure 6). Anthersmut disease has many advantages for the study of the epidemiology and evolutionary genetics of disease. It has almost no direct agricultural or economic impact, and can therefore readily be studied in many species in natural populations, and experimental populations are easily established in the field (Antonovics et al., 2002). It has a yeast-like stage with two mating types, is easily grown in the lab and has a small genome (ca. 25 Mb) with chromosomes readily separable by pulse-field gel electrophoresis. Multilocus pathogen and host phylogenies are providing the basis for comparative studies of disease characteristics, including the ecological and evolutionary determinants of host shifts, disease emergence and the evolution of virulence.

Microbotryum also has heteromorphic chromosomes that carry the mating type locus and are sometimes called sex chromosomes. These are the first heteromorphic chromosomes to be discovered in fungi. As in Silene, dimorphism of these chromosomes has repeatedly evolved independently in the genus, they seem to have a region of suppressed recombination, in which transposable elements have accumulated (Hood et al., 2004). However, fungal mating compatibility acts in the haploid stage, so that diploid individuals are always heterozygous and there is no sexual asymmetry in recombination opportunity or exposure of recessives, such as exist for the sex chromosomes of diploids. Similarities in the evolutionary forces acting on diploid sex chromosomes and haploid mating-type chromosomes are currently under study (Hood et al., 2004). As is often the case in biology, parasites have their own parasites. In the case of Microbotryum, these include many transposable elements in its genome. Comparative study of the coevolution of transposable elements with fungal genomic defences (repeat induced point mutations; Hood et al., 2005) is another exciting future research area.

Several other aspects of Microbotryum ecology and genetics also require joint study with its Silene hosts to benefit understanding of both. Microbotryum has recently been shown to consist of different species, highly specialized onto one or two host plants (Le Gac et al., 2007a). Partial premating isolation among host plants also affects Microbotryum because the pathogen transmission depends on the host plants’ habitats and pollinators (van Putten et al., 2007). There has however been little, if any, cospeciation between the plants and the fungi (Refregier et al., 2008). Premating barriers between both close plant and fungal species are weak, making this system an ideal one to evaluate the importance of host and pathogen hybridization in disease evolution (Le Gac et al., 2007b; Karrenberg and Favre, 2008; Minder and Widmer, 2008; Sloan et al., 2008a).

**Biotic interactions: pollination and seed predation**

Pollinators play a role in prezygotic reproductive isolation and genetic structure (Richards, 2000; Teixeira and Bernasconi, 2007), but are also important seed predators in Silene (Jolivet and Bernasconi, 2006; Kephart, 2006). Most species are pollinated by moths or bumblebees and have extensively differentiated floral morphologies and scent profiles (Jurgens et al., 2002; Kazama et al., 2005).

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**Figure 6** Relationship between population growth and incidence of anther smut in Silene latifolia and HIV in humans. (a) Silene annual growth rates measured as ln(N_{t+1}/N_{t}) for 1989–1993 in Giles County, Virginia; data grouped by population size (classes 8–15, 16–31, and 32–63), dot sizes are proportional to the number of populations (range 3–239). (b) Data for humans are predicted annual per capita growth rates for 2000–2025 in sub-Saharan African countries, compiled from United Nations, World Population Prospects. The 2002 Revision. Volume III: Analytical Report.
Jürgens, 2006). Plant–pollinator interactions are generally seen as mutualistic, but antagonism also occurs. Several *Silene* species are pollinated by nocturnal moths (the lepidopteran fruit predators *Hadena*, Noctuidae and *Perizoma*, Geometridae) that lay eggs in the flowers, using the developing seeds as a resource for their progeny (*nursery pollination*; Figure 5b). But unlike the *Yucca/Yucca* moth and fig/fig wasp associations (obligate mutualisms), *Silene* flowers can be pollinated by other insects both in the native (Gimenez-Benavides et al., 2007) and in the introduced ranges (Barthelmess et al., 2006). This offers ideal opportunities for testing how ecological variation in co-pollinator efficiency influences the outcome of the interaction with the seed predator. *Silene–Hadena* interactions may represent a form of nursery pollination with low or incipient specialization (Kephart, 2006). Recent evidence indicates that fruit abortion (Jolivet and Bernasconi, 2006) and rapid post-pollination changes in scent emission (Mühlemann et al., 2006) may regulate seed damage at this early stage of plant–mutualist interaction.

Interactions between herbivores and the pathogens they carry may range from mutualistic to antagonistic. In *Silene*, nursery pollinators, including *Hadena*, play a major role in the spread of spores of the anther-smut fungus, affecting pollination and seed set in complex multi-trophic interactions (Carlsson-Graner et al., 1998). Fungal attack reduces the benefits of the interaction to the moth. It is interesting that this is mediated not only by reduced resource quality of infected plants, but also by reduced ‘refuge space’ for moth larvae (Biere et al., 2002), a factor known to be important in stabilizing trophic interactions in natural communities. Moths respond by avoiding oviposition on infected plants (Biere and Honders, 2006). This system also illustrates that the often complex interactions between organisms at different trophic levels are also affected by human activities. These often lead to fragmentation of *S. latifolia* habitats in the native range. Reduced population size and increased isolation between plant patches are predicted to differentially affect organisms at different trophic levels. In fragmented native *S. latifolia* populations, isolation and host population size indeed hardly affect the frequency of *Hadena* in flowers, but reduce the incidence of some of its parasitoid species (Elzinga et al., 2005).

**Conclusions**

The detailed knowledge that has been obtained about *Silene* and its interacting organisms over more than a century provides a rich source of interesting unresolved questions in plant ecology and evolution. With increasing availability of genetic tools, further progress in *Silene* should be possible on the range of fundamental questions outlined here. We have highlighted several areas in which new studies are needed; for example, the suppression of recombination during sex chromosome evolution and the role of sexually antagonistic selection in this process. The ecological knowledge available in species of this genus will prove to be an important advantage in comparison with many other systems that were developed as models because of molecular tractability alone, but without information about natural populations. In the age of PCR and genome-wide sequencing of DNA and mRNA, as also illustrated by *Mimulus* (Wu et al., 2008), molecular non-model species in which interesting biological questions are waiting to be solved attract renewed interest and allow new advances.

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**References**


