A unified model of Hymenopteran preadaptations that trigger the evolutionary transition to eusociality

Andrés E. Quiñones1,† & Ido Pen1

Explaining the origin of eusociality, with strict division of labour between workers and reproductives, remains one of evolutionary biology’s greatest challenges. Specific combinations of genetic, behavioural and demographic traits in Hymenoptera are thought to explain their relatively high frequency of eusociality, but quantitative models integrating such preadaptations are lacking. Here we use mathematical models to show that the joint evolution of helping behaviour and maternal sex ratio adjustment can synergistically trigger both a behavioural change from solitary to eusocial breeding, and a demographic change from a life cycle with two reproductive broods to a life cycle in which an unmated cohort of female workers precedes a final generation of dispersing reproductives. Specific suits of preadaptations are particularly favourable to the evolution of eusociality: lifetime monogamy, bivoltinism with male generation overlap, hibernation of mated females and haplodiploidy with maternal sex ratio adjustment. The joint effects of these preadaptations may explain the abundance of eusociality in the Hymenoptera and its virtual absence in other haplodiploid lineages.
Eusociality characterized by physically differentiated castes is the most advanced form of social life found in the animal kingdom. Its hallmark is reproductive division of labour, where some individuals (workers) refrain from their own reproduction but instead increase the reproductive output of their parents. The first evolutionary hypothesis that attempted to explain the distribution of eusociality across animal taxa, the ‘haplodiploidy hypothesis’, was proposed by Hamilton using his then new inclusive fitness theory.\(^3\) Haplodiploidy is the sex determining mechanism whereby fertilized eggs become females, and unfertilized eggs males. As a consequence, full sisters are genetically more closely related to one another ($r = 3/4$) than to their own daughters ($r = 1/2$), thus favouring workers to channel more effort into raising the former rather than the latter. When Hamilton proposed his hypothesis, almost all the taxa known to have evolved eusociality—with the notable exception of ants, bees and wasps. Thus, it seemed at least plausible that haplodiploidy was an important preadaptation: a trait that arose before, and independently of evolving helpers at the nest, but that appeared to increase the chances for facultative helping, to ultimately produce a sterile worker caste. However, since then a number of additional diploid eusocial clades have been discovered, including ambrosia beetles,\(^4\) sponge-dwelling shrimp,\(^5\) and bathyergid mole rats,\(^6\) suggesting that the apparent importance of haplodiploidy is less obvious.\(^7\) The workers in these newly discovered diploid eusocial clades, despite showing reproductive altruism, have maintained their capacity to become reproductively active, at least under certain circumstances. In contrast, the advanced eusocial clades—all of them haplodiploid aculeate Hymenoptera except the termites—have obligate reproductive altruism; that is, the workers never mate and have lost their reproductive totipotency. Thus, haplodiploidy might still have an important facilitating role in the evolution of advanced stages of eusociality.

Trivers and Hare\(^8\) pointed out that Hamilton’s original model overlooked the fact that in haplodiploid species females are more closely related to their sons ($r = 1/2$) than their brothers ($r = 1/4$), and that this exactly cancels the genetic benefits of helping raise siblings instead of helping to raise offspring. Consequently, only when the production of the sexes is split among different nests (split sex ratios), will haplodiploidy favour worker behaviour, and it will do so only in those nests with female-biased sex ratios.\(^9\)–\(^10\) Thus, maternal ability to bias offspring sex ratios came to be seen as an additional preadaptation for the evolution of eusociality in Hymenoptera, in addition to any environmental conditions that would favour split sex ratios.\(^11\) Seger showed that temporally split sex ratios are promoted by the bivoltine life cycle found in many species of solitary insects closely related species with helpers at the nest.\(^11\) Bivoltinism, the production of two non-overlapping broods in one reproductive season, opens the possibility to split the production of the sexes between the two broods. Seger\(^11\) hypothesized that a female-biased second brood later in the season would promote the evolution of helping behaviour in the first brood, thus adding bivoltinism as yet another preadaptation for the early evolution of eusociality. A female-biased summer brood implies that if a spring-hatched female stays to help her mother, she would help raise siblings that are on average more closely related to her than her own offspring would be, if she would assist in raising an even sex ratio. However, Seger’s argument pertains only to the initial invasion of the helper trait, and does not account for possible changes in sex ratios driven by the presence of helping behaviour, once it has evolved.\(^12\) Sex allocation theory predicts evolutionary feedbacks between helping behaviour and sex ratios when only one of the sexes helps, as is the case in Hymenoptera,\(^13\)–\(^15\), thus making it necessary to jointly account for the co-evolution of helping behaviour and facultative sex ratio adjustment.

Besides haplodiploidy, maternal manipulation of offspring sex ratios and bivoltinism, several additional factors have been proposed to bias the odds in favour of eusociality, such as specific life cycle structures, ecological conditions\(^16\)–\(^17\) and last but not least a monogamous mating system.\(^18\)–\(^20\) However, it is unknown how these factors jointly affect the evolution of reproductive altruism, and in particular whether they act synergistically in promoting it. Here we develop a unified model, grounded in the life history of primitively social insects described by Seger\(^11\). The model integrates many of the proposed preadaptations, and allows the co-evolution of helping behaviour and sex ratios. We show that, indeed, specific combinations of traits, life history characteristics and ecological conditions strongly increase the likelihood that reproductive altruism evolves. Furthermore, we show that sex ratio evolution causes the production of a first brood of unmated workers before a brood of reproductives, leading to a univoltine life cycle reminiscent of annual colonies of bumblebees and vespine (yellowjacket) wasps that produce workers in the spring and early summer and a final brood of reproductive at the end of the season.

**Results**

**Partial bivoltinism.** We constructed models for populations with two partially overlapping generations per year, that is, partially bivoltine populations that are common in Hymenoptera\(^11\) (details in the Methods—Demography). The first generation of the year, or spring generation, gives rise to a summer generation consisting of offspring and survivors of the spring generation. The summer generation then gives rise to an autumn generation, some of which members overwinter to form a new spring generation, thus completing the life cycle (Fig. 1, top). In the models, females have three potentially evolvable traits: the probability $0 \leq h \leq 1$ of females hatching in the spring to forgo reproduction and stay at their natal nest to help their mother (for example, individuals with $h = 0$ never stay to help; of individuals with $h = 0.5$, half stay to help while the other half leave), and the offspring sex ratios: (proportion of sons) $0 \leq z_1 \leq 1$ and $0 \leq z_2 \leq 1$ produced in spring and summer broods, respectively. Each helper adds an additional $B$ offspring for each offspring produced by her mother. $B > 1$ implies that helpers are more efficient at raising siblings than at raising their own offspring, while $B < 1$ means that they are more efficient at raising their own offspring than they are at raising siblings.

**Unisexual or bisexual overwintering.** We followed Seger\(^11\) in considering two types of partially bivoltine life cycle: the ‘female overwinter’ (FH) type where only mated females from the autumn generation overwinter, and the ‘larval diapause’ (LD) type where both sexes overwinter as diapause larvae\(^10\),\(^11\) (Supplementary Figs 1 and 2). For both types of cycle, overwintering females reproduce first in the spring, and may reproduce (if they survive) a second time during summer, while females from the spring brood can only reproduce in the summer. In the FH model, males hatched in the spring can mate with females hatched in the spring, and if they survive (with probability $S_m$) they can mate with females hatched in the summer as well. In contrast, males hatched in the summer can only mate with females hatched in the summer. As a result, in the FH scenario, males hatched in the spring have an inherently higher expected reproductive success than males hatched in the summer. This causes natural selection to favour male-biased sex ratios in the spring and female-biased sex ratios in the summer.\(^11\) In the LD model the situation is reversed in that males from the summer brood overwinter and get the chance to
mate both in the following spring, with overwintering females, and, if they survive (with probability $S_m$), once again in the summer with females from spring broods; in contrast, males from the spring brood mate only in the summer with females from the spring brood. Thus, the LD life cycle favours male-biased sex ratios in the summer and female-biased sex ratios in the spring.$^{11}$

**Eusociality threshold.** We modelled the life cycles described above using the matrix population model approach. We first derived a transition matrix to track the dynamics of rare mutant phenotypes invading a resident population monomorphic for the three traits. Using the reproductive value approach$^{21,22}$, we derived the selection gradient for each of the traits. Our main interest was in studying the conditions under which altruistic helping ($h > 0$) evolves, and how this is affected by the presence or absence of specific preadaptations. To this end, we analysed different versions of our models, by varying: (1) the life cycle structure that depends on whether only females overwinter (FH), or both males and females overwinter (LD, see previous section for details); (2) the type of genetics (diploidy or haplodiploidy); (3) the mating system, with several alternative options (obligate monandry = lifetime monogamy with life-time storage of sperm from a single ejaculate, polyandry with females storing sperm of a variable number of males ($m_z$; see Methods—Polyandry) for life, serial monogamy with females needing to re-mate and store new sperm for producing a second brood; and lastly, (4) the presence or absence of maternal sex ratio control, that is, the ability to produce different offspring sex ratios for the first and the second broods. We analysed the effects of the preadaptations by deriving the minimal level of helper benefits ($B_{min}$) necessary for selection to favour the evolution of helping; we refer to this quantity as the eusociality threshold. This measure is equivalent to the ‘efficiency ratio’ of Charnov$^{23}$ and Grafen$^{9}$, and the ‘potential for altruism’ of Gardner$^{24,25}$, the relative efficiency of a helper in raising sibs as opposed to own offspring at which she is indifferent between those two options. Finally, we computed evolutionary dynamics of these traits using an adaptive dynamics approach$^{26,27}$ based on our inclusive fitness expressions$^{28}$ and we complement them with matching individual-based population genetic simulation models (see Methods—Individual-based simulations for details).

**Co-evolutionary dynamics.** An analysis of the co-evolutionary dynamics of helping and sex ratios shows that the FH life cycle, with overlapping generations of males, initially favours the evolution of male-biased broods in the spring and female-biased broods in the summer (Fig. 2), as predicted by Seger$^{11}$. Thus, sex ratio manipulation under the FH life cycle strongly promotes a transition from solitary breeding to reproductive altruism, while the LD life cycle tends to inhibit it (Methods—Selection on helping behaviour). Moreover, once helping behaviour is present and daughters increase the fecundity of their mother, natural selection favours mothers that allocate more resources into the production of more helpers. This is achieved by shifting the sex ratio of the spring brood to produce more females. Eventually, the spring brood becomes 100% female helpers which, due the complete lack of males in their cohort, all remain unmated and thus represent the start of an obligate worker caste. Moreover, the same lack of spring males causes the summer sex ratio to evolve back to fifty–fifty (Fig. 2). This amounts to a major life history transition from a partially bivoltine (Fig. 1, top) to a univoltine life cycle cycle, with a specialized breeder and life-time unmated workers (Fig. 2, bottom), triggered by an evolutionary feedback between social behaviour and sex ratios$^{12}$. This evolutionary feedback is robust to assumptions in the mutational structure of the three evolving traits (Supplementary Fig. 3).

**Preadaptations and synergies in the evolution of helpers.** Our models, unlike Hamilton’s original haplodiploidy hypothesis$^3$, account for the joint effects of genetics (ploidy) and life history/ecological traits (overwintering, mating system, sex ratios, sex-specific survival). However, the relative importance of the different traits differs in magnitude and consistency. Lifetime monogamy$^{16}$, for instance, due to its positive effect on within-brood genetic relatedness, unambiguously reduces the eusociality threshold to one half of the threshold under serial monogamy and up to one half of the threshold for varying degrees of polyandry (Fig. 3). In contrast, haplodiploidy$^{15}$ and sex ratio manipulation can both favour and harm the evolution of helping behaviour$^{25}$, depending on the specific life cycle (FH or LD, Fig. 3). However, the simultaneous presence of lifetime monogamy, haplodiploidy, and sex ratio manipulation in a FH life cycle can reduce the threshold to only 2/3 of their productivity as solitary breeders (Fig. 3, Methods—Selection on helping behaviour). Specifically,
for haplodiploids with an FH life cycle, where females on average have an effective number of mates $m$ and use the sperm of the same males for both broods, the condition for altruistic helping behaviour to be favoured by selection is

$$B > \frac{2 - O_m}{1 + 1/m_e - 4O_m}$$

(1)

Here $0 \leq O_m \leq 1$ is a measure of generation overlap between males born in the first brood and males born in the second brood (see Methods—Class-specific individual reproductive values). If no males from the first brood mate with females from the second brood ($O_m = 0$), then (1) reduces to $B > 1$ at best (monogamy: $m_e = 1$) and $B > 2$ at worst (extreme polyandry: $m_e \to \infty$). Conversely, if males from the first brood monopolize females from the second brood ($O_m \to 1$), as a result of strongly split sex ratios between males in the first brood and females in the second brood, then (1) reduces to $B > 2/3$ at best ($m_e = 1$) and again $B > 2$ at worst ($m_e \to \infty$). The right-hand side of (1) strictly increases with the effective number of mates $m_e$; therefore, any increase in polyandry (higher $m_e$), all else being equal, makes it harder for altruistic helping behaviour to evolve (Supplementary Fig. 4).

For diploids, the equivalent condition is

$$B > \frac{2}{1 + 1/m_e}$$

(2)

Note that the right-hand side of (2) is always larger than the right-hand side of (1) and never smaller than one and that every deviation from monogamy ($m_e > 1$) will make helping behaviour less likely to evolve. For the LD life cycle, the corresponding conditions (1) and (2) are always less favourable for the evolution of reproductive altruism (see Methods—Selection on helping behaviour).

Combining the results for haplodiploids and diploids, in order for the threshold value of $B$ to dip below unity, the necessary conditions are haplodiploidy, female hibernation, partial survival of spring males until the summer cohort hatches ($O_m > 0$) and limited promiscuity ($m_e < 1/(1 - 4O_m)$); together these conditions are
sufficient. Thus, in taxa possessing combinations of specific preadaptations (lifetime monogamy, haplodiploidy, sex ratio manipulation and unisexual overwintering), the likelihood of the evolution of altruistic helping is significantly increased. Note that this does not imply the evolution of a caste of unmated workers. Coevolution of sex ratios and helping behaviour leads eliminates the production of males in the spring brood (Fig. 2, $z_1 = 0$), and with this, the end of male generation overlap ($O_m = 0$; see Methods—Demography). Thus, the final step of the evolutionary transition to a specialized unmated worker caste requires the benefits of helpers to be larger than 1.

**Discussion**

The two types of life history considered here represent distinctive characteristics found in different taxonomic groups of Hymenoptera. The female hibernation life cycle is a common feature of Halictine bees, bumblebees Vespine and Polistes wasps, where despite large diversity in life history and social behaviour, life cycles usually start with a solitary mated female or (in Polistes) occasionally groups of mated females. Overwintering characterizes the life cycle of many temperate species due to seasonality, but tropical species often also have periods of inactivity driven by the cycles of rainy and dry weather. Halictine bees are the group of animals in which helper reproductive altruism has evolved more times than in any other group. In primitively eusocial Halictines, despite workers keeping the potential to reproduce, the life cycle resembles the one described by our model: colonies first produce a female-biased helper brood followed by a reproductive brood at the end of the season. In contrast, the initial stage in our model, although present, is rare among solitary Halictines. Most solitary Halictine species produce only a single brood per reproductive season. Some species, such as *Halictus sexcinctus* and *H. rubicundus*, can both nest solitarily and with helpers depending on their geographical location; solitary populations are univoltine, but if they live in locations where the length of the season allows more than one brood, they recruit daughter helpers as described by our model. The rarity of the bivoltine life cycle and the corresponding facultative sex ratios, both in solitary and social Halictines, has been used as an argument against the broader applicability of Seger’s model to understand the evolution of helper and worker caste. Our model resolves that issue by showing that strict bivoltinism and a female-biased summer brood are no longer expected once helpers become unmated workers, due to the feedback between sex ratio evolution and the helping behaviour. Turning the argument upside down, we expect the bivoltine life cycle with female hibernation to be rare in groups with helping behaviour and female-biased summer broods. In agreement with our model predictions, comparative analysis has shown that a female-biased sex ratio in the first brood correlates positively with the degree of sociality across insect lineages with helpers at the nest.

The larval diapause condition is found in sphecid wasps, for which—all although less well studied—there is only a single report of helpers at the nest. However, there is some evidence of the sex ratio biases predicted by the Seger model. Larval diapause is also common among Megachilidae bees, the almost exclusively solitary sister lineage of the corbiculate bees (honey bees, stingless bees, bumblebees, orchid bees). Lifetime monogamy is a preadaptation that extends beyond the Hymenopteran scenario portrayed by our model. The favourable effects that lifetime monogamy has on the evolution of reproductive altruism and cooperative breeding have both considerable theoretical and empirical support. Strict parental monogamy has now been well documented as the ancestral state across lineages of the social Hymenoptera and termites with permanent worker castes, but the mechanisms are very different. In the Hymenoptera lifetime monogamy is facilitated by the presence of lifetime sperm storage, which could be considered a morphological preadaptation. In higher termites, lifetime monogamy is facilitated by the royal chamber inside the nests. In our model, any departure from lifetime monogamy reduces the prospects for the emergence of reproductive altruism, and the presence of monogamy enhances the favourable effects of haplodiploidy and sex ratio manipulation on the likelihood of retaining helpers at the nest. Our analysis also shows that strict monogamy is not necessary for the threshold value of the benefits per helper to be smaller than unity; for sufficiently low levels of polyandry, the combined presence of haplodiploidy, female hibernation, sex ratio manipulation and male generation overlap may cause the threshold $B < 1$ facilitating the evolution of reproductive altruism (Supplementary Fig. 4). However, in the final stages of the evolutionary transition, due to the lack of males, strict monogamy is necessary for the threshold to be as low as $B = 1$.

Some recent models have questioned the general positive effect of monogamy for the evolution of eusociality. However, a more general analysis has shown that putative exceptions regarding the effect of monogamy are restricted to cases where altruism is determined uniquely by a single allele of strong effects. Moreover, these models focus on the rate of initial increase of alleles for helping behaviour—where multiple mating by males carrying mutant alleles allows for faster spreading of mutant alleles across colonies—but not on equilibrium levels of altruism after multiple successive invasions of mutant alleles of smaller effects. Our models, in contrast, assume that gradual substitution of many alleles of small effect is driving the evolution of social behaviour, consistent with empirical data on life-history traits being generally polygenic and heritable in a quantitative genetics sense. Thus, we believe that the evidence, both theoretical and empirical, supports the role of monogamy as being crucial for the evolution of a worker caste.

Apart from the specific predictions about sex ratios in different broods, our model highlights the importance of the production of more than one brood per year. The production of two broods opens the possibility to have opposite sex ratio biases in each one of them, a form of temporal split sex ratios. Bivoltinism has also been proposed as crucial for the origin of a developmental preadaptation that favours evolution of the worker phenotype. The origin of helper and worker phenotypes requires a developmental system that expresses reproductive altruism at the right moment of the life cycle. Bivoltinism requires tuning the expression of two metabolically different phenotypes: one that stores reserves and enters into hibernation or aestivation diapause (summer brood), and one that reproduces immediately (spring brood). The developmental system that tunes the expression of these two distinctive bivoltine phenotypes could then be co-opted to express the worker and queen phenotypes. Thus, multiple sources of evidence point towards bivoltinism as a predecessor a facilitating factor for retaining helpers at the nest. Multivariate statistical analysis shows that indeed the number of broods, governed by breeding season length and developmental time, is positively correlated with the level of sociality among facultative and obligately eusocial species. Moreover, environmental factors related to season length appear to be important determinants for the expression of helping behaviour in species that can either have helpers or be solitary, depending on geographical location. Hamilton’s idea that haplodiploidy favours the evolution of reproductive altruism has been challenged on both empirical and theoretical grounds.
theoretical grounds. The empirical grounds are that reproductive altruism occurs also in several diploid species. However, up until now there has been no formal phylogenetically corrected statistical analysis to assess whether eusociality is more often found in haplodiploid lineages than diploid ones; but, a quick look still seems to suggest there is. The theoretical grounds are that when both sexes are properly accounted for in the inclusive fitness calculations, haplodiploidy requires female-biased sex ratios to favour eusociality. However, recent evidence suggests that the conditions (polyandry, see above) which favour such biased sex ratios are unlikely to have been present in the early evolution of eusociality. Here we have shown that, depending on the type of bivoltine life cycle, haplodiploidy can both inhibit and promote the evolution of reproductive altruism. When the bivoltine life cycle starts with mated females emerging from hibernation, haplodiploidy strongly favours the evolution of helpers at the nest. Furthermore, haplodiploidy is a sex determination system that allows flexibility in maternal control of offspring sex ratios. Given that we identified maternal sex ratio manipulation to be another preadaptation, haplodiploidy has a two-fold effect in the evolution of reproductive altruism in the Hymenoptera. This does not mean that our model predicts in general that helpers at the nest evolved more often in species with haplodiploid genetics; but, rather, that reproductive altruism will be found more often in species with haplodiploid genetics that present also the other preadaptations. Furthermore, a closer look at species with diploid genetics might lead to the discovery of preadaptations specific to that case.

The evolutionary scenario encapsulated by our model starts with a population living in a seasonal environment that allows the production of two broods, associated with two mating episodes in one reproductive season. Our model predicts that evolution drives such populations towards a single synchronous mating episode in the life cycle, a condition predominant in all taxa with obligatorily unmated workers, even when living in tropical environments and with secondarily evolved obligate polyandry in that single episode. Once a univoltine life cycle with a first brood of workers has evolved, it is possible to envision evolutionary, demographic and ecological changes that extend the specialization of the first (worker) brood, leading eventually to physically differentiated castes. For starters, under the predicted fully female first brood, females from the first brood cannot mate. Thus, reproductive competition inside the nests does not compromise the social endeavour, paving the way for further adaptation of females to their worker role. The evolution of fully committed workers, which have lost reproductive totipotency, increases the benefits they provide to the colony. Such more elaborate commitments requires drastic behavioural and developmental changes, during the large number of generations required to decisively modify developmental pathways towards queen and worker phenotypes. For seconds, environmental changes that increase the length of the reproductive season may allow the production of a larger worker brood. These changes could occur simply by range expansion toward lower latitudes or altitudes. A larger worker brood would then possibly enable colonies to gather enough resources to transition towards a perennial life cycle. Evidence for such a transition can be found in bumblebees, where temperate species invariably have the univoltine eusocial life cycle predicted by our model, while some tropical relatives have a perennial life cycle, while maintaining the female hibernation condition and a single-synchronous mating episode. However, as long as unmatedness of the first brood females has not gone to fixation, reductions in the length of the breeding season may reconstitute solitary life, which appears to have happened more often in Halictines than in any other lineages with helpers at the nest. Yet other evolutionary scenarios might be driven by conflicts inside the colony; for example, queen-worker conflict over control of the sex ratio of the second brood, or conflict over worker production of male eggs, might lead to alternative social organizations and life histories. Taking into account those potential scenarios will probably give a better prediction of the distribution of eusociality in the Hymenoptera.

The evolutionary transition to eusociality, the most advanced form of social life, encompasses radical and complex changes in many facets of a species’ biology. Rather than one unique causal factor, we showed how specific combinations of them can drive the transition. Monogamy, haplodiploidy with maternal sex ratio manipulation, bivoltinism with male generation overlap, and hibernation of mated females combine to provide the most favourable conditions for the evolution of reproductive altruism. The Hymenoptera seem to have serendipitously ended up with such a set of traits, and because of them, have achieved their supreme ecological position.

Methods

General modelling approach. We consider populations with two partially overlapping generations per year, that is, partially bivoltine populations with a spring generation and a summer generation. Each generation has a specific class structure, that is, a frequency distribution of females and males, reproductive and helpers. The vector \( N_i(t) \) keeps track of the class distribution for the spring generation at year \( t \), while \( N_i(t+1) \) tracks the summer generation in year \( t+1 \). Transitions between generations are modelled with demographic matrices \( D \), that track survivors and the offspring produced by females, and \( O \), that track the next generation.

At demographic equilibrium (DE: when \( N_i(t+1) = N_i(t) \)), all transitions in a single year can be described by a block matrix of the form:

\[
D = \begin{pmatrix} D_1 & D_2 \\
D_3 & O_1 & O_2 \\
D_4 & O_3 & O_4 \\
\end{pmatrix}
\]

The \( O \) are matrices of appropriate dimensions filled with zeroes. The dominant eigenvalue of \( D \) must be \( \lambda = 1 \). To ensure this we scale winter survival by a factor \( z \) (see ‘Demography’ below). The corresponding dominant right eigenvector \( u \) contains for both generations the stable class distributions in DE, which will be needed for the inclusive fitness calculations outlined below.

Class-specific individual reproductive values, the long-term genetic contributions of individuals to future generations, are derived from a gene flow matrix. This matrix keeps track of female and male reproduction and survival, where each contributor to a newly born or survivor in the next time step gets credit according to the proportion of genes derived from the contributor. For example, in haplodiploidy a female gets 100% credit for her male offspring, while females and males both get 50% credit for each of their female offspring. In diploids both parents get 50% credit for both sexes. The gene flow matrices have the same block structure as the demographic matrices:

\[
A = \begin{pmatrix} O_1 & O_2 \\
A_1 & A_2 \\
\end{pmatrix}
\]

The dominant left eigenvector \( v \) of \( A \), that is, the solution of \( v^T A = v^T \), where \( T \) denotes transposition, contains the class specific individual reproductive values.

To model the evolution of a trait \( x \), where \( x \) is a sex ratio or helping tendency, we analysed the inclusive fitness of a focal mutant individual with trait value \( x^* \). The focal individual belongs to class \( k \), and contributes \( w_{jk}(x,x^*) \) individuals to class \( j \) with average relatedness \( r_{jk} \) to the focal individual, then the focal individual’s inclusive fitness is given by

\[
W(x,x^*) = \sum_{jk} w_{jk}(x,x^*) r_{jk} v_j^*
\]

where \( v_j^* \) is the reproductive value of class-\( j \) individuals in the resident population, \( w_{jk} \) is the contribution from class-\( j \) to class-\( k \) and \( r_{jk} \) is the average relatedness. The selection differentials with respect to \( x \) are then given by

\[
\frac{\partial W}{\partial x} \bigg|_{x=x^*} = \sum_j w_{jk}(x,x^*) r_{jk} \frac{\partial v_j^*}{\partial x} \bigg|_{x=x^*}
\]

If a trait is expressed by individuals in more than one class, then the selection differentials are obtained by summing over the appropriate classes, weighing each class according to its normalized class frequency \( n_{jk} \).

Evolutionary equilibria are calculated by setting the selection differentials to zero and solving for \( x^* \). Evolutionary dynamics are modelled using a standard adaptive dynamics approach, where the rate of change of \( x^* \) over evolutionary
time is proportional to the selection differential:
\[
\frac{dc}{dt} \propto \frac{\partial W}{\partial x} \bigg|_{x=x^*}
\] (8)

We used individual-based simulations to check the results of the analytical inclusive fitness analyses; details of the simulations are in section Individual based simulations.

Demography. Here we construct the demographic models that keep track of the class frequencies in the partially bivoltine populations. A summary of our notation is in Table 1. We consider two kinds of life history: the Female Hibernation (FH; Supplementary Fig. 1) life history and the Larval Diapause (LD; Supplementary Fig. 2) life history. For the moment, we assume female lifetime monogamy, i.e. each female mates once in her life with a single male; later (section ‘Polyandry hampers the evolution of helping’) we drop this assumption.

Demography for female hibernation. We keep track of seven classes of individuals (Supplementary Fig. 1). The spring generation consists of mated females, the ‘foundresses’, which have survived hibernation (class 1) and the sperm they carry (class 2). Each mated female produces \(F_1\) offspring, a fraction \(z_1\) of which are sons (class 7) and \(1 - z_1\) daughters (classes 3 and 4). A fraction \(h\) of daughters remain at the natal nest and become helpers (class 4), while \(1 - h\) become independent breeders (class 3). Mated females themselves survive until summer to breed again with probability \(S_1\) (class 5), along with the sperm they still carry (class 6).

These transitions between spring and summer generations are encapsulated by the matrix
\[
D_1 = \begin{bmatrix}
1 & 2 \\
3 & 4 \\
5 & 6 \\
7 & 0
\end{bmatrix}
\] (9)

Numbers above the columns indicate contributing classes, numbers along rows indicates ‘receiving’ classes. Note that column 2 contains zeroes only since we only count offspring of females in order to prevent double counting.

In the summer generation, each class 3 female mates with a single class 7 male and produces \(F_2\) offspring, with sex ratio \(z_2\). Each class 5 female, which may have helpers at her nest, produces \(F_3\) offspring with sex ratio \(z_2\). We sometimes refer to the offspring produced by the summer generation as the ‘autumn offspring’. These autumn offspring mate in the autumn, as do a fraction \(S_0\) of class 7 males of the summer generation. The mated females go into hibernation and have density-dependent survival such that the number of class 1 females in the next spring is the same as in the previous spring. These transitions are governed by the matrix
\[
D_2 = \frac{1}{2} \begin{bmatrix}
3 & 4 & 5 & 6 & 7 \\
2z_2F_1 & 0 & x_2F_1 & 0 & 0 \\
2z_2F_3 & 0 & x_2F_3 & 0 & 0
\end{bmatrix}
\] (10)

Both demographic matrices can be combined into a single matrix that keeps track of the class frequencies in both generations at DE:
\[
D = \begin{bmatrix}
O_1 & D_1 \\
D_2 & O_2
\end{bmatrix}
\] (11)

Here \(O_1\) and \(O_2\) are respectively \(2 \times 2\) and \(5 \times 5\) matrices filled with zeroes. The DE relative class frequencies are solutions of \(Du = u\) and they are
\[
\begin{align*}
    u_1 & = 1 = x_2F_1u_3 + x_2F_3u_5 \\
    u_2 & = u_1 = 1 \\
    u_3 & = \tilde{h}_2F_1 \\
    u_4 & = S_1\tilde{h}_2F_1 \\
    u_5 & = S_1 \\
    u_6 & = S_1 \\
    u_7 & = z_1F_1
\end{align*}
\] (12)

Note that we have set the relative frequency \(u_1\) of class 1 to unity and expressed the other class frequencies as multiples of the class 1 frequency. This normalization determines the density-dependent female winter survival, as follows:
\[
\alpha = \frac{1}{z_2(u_2F_3 + u_3F_3)} \equiv \frac{1}{z_2F_A}.
\] (13)

\(F_A\) can be regarded as the total number of autumn offspring per spring female, and \(\tilde{z}_2F_A\) therefore the corresponding number of females among autumn offspring. The scaling ensures that on average every spring female is exactly replaced by another spring female one year later.

Given the DE class frequencies, the average number of mates per female during the autumn mating season, respectively during the summer mating season, are...
given by:

\[ Q_1 = \frac{z_2 F_A}{z_2 F_A + S_m z_1 F_1} \]  

\[ Q_2 = \frac{u_1}{u_1 + u_6} = \frac{h z_1 F_1}{z_1 F_1 + S_m z_2 F_1} \]  

In the autumn, males produced by the summer generation \( (z_2 F_A) \) and the surviving males of the summer generation \( (S_m z_1 F_1) \) compete together for \( z_2 F_A \) females, hence on average \( Q_1 \) mates per male. In the summer, the class 7 males compete for the non-helping class 3 females, hence \( Q_2 \) mates per male.

**Demography for larval diapause.** For the LD life history, the spring generation consists of unmated females and males that survived winter diapause. This first generation mates randomly and produces the summer generation, consisting of surviving adults and new offspring (Supplementary Fig. 2). It is conceptually convenient to split summer males into surviving spring males and the sons of spring females, hence we now have 8 classes in total. Following the same approach as for the FH life history, the DE class frequencies are tracked by the block matrix

\[ D = \begin{bmatrix} 0 & 0 & z_2 F_1 & z_2 F_1 & 0 & 0 & 0 & 0 \\ 0 & 0 & z_2 F_1 & z_2 F_1 & 0 & 0 & 0 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ S_f h z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ 0 & S_m & 0 & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \]  

The stable class distribution follows again from \( Du = u \):

\[\begin{align*}
  u_1 & = 1 = z_2 F_A \\
  u_2 & = z_1 / z_2 \\
  u_3 & = h z_1 F_1 \\
  u_4 & = S_f h z_1 F_1 \\
  u_5 & = S_f \\
  u_6 & = S_f \\
  u_7 & = z_1 F_1 \\
  u_8 & = S_m z_2 / z_2 
\end{align*}\]

The average number of mates for spring and summer males are, respectively,

\[ Q_1 = \frac{u_1 - u_2}{u_1 + u_8} = \frac{h z_1 F_1}{z_1 F_1 + S_m z_2 / z_2} \]  

\[ Q_2 = \frac{u_1}{u_1 + u_8} = \frac{h z_1 F_1}{z_1 F_1 + S_m z_2 / z_2} \]  

**Effect of helping.** For both FH and LD life histories, the expected number of helpers for a class 5 summer female is given by

\[ H = h z_1 F_1. \]

We assume that each helper increases the number of offspring produced by her mother by an amount \( b F_A \), hence

\[ F_A = F_A (1 + b H) = F_A (1 + b h z_1 F_1). \]

Note that a class 5 female's output increases linearly with the number of daughters\( z_1 F_1 \) produced during the spring.

At the moment a helper 'decides' to stay at her natal nest, her expected contribution \( B \) for each of her mother's future offspring is conditional on her mother's survival:

\[ B = S_f b. \]

**Class-specific individual reproductive values.** Individuals of different classes typically differ in their relative contributions to the future gene pool, and individual reproductive values (RVs) quantify these differences\(^4\). In matrix population models, the individual RVs are the elements of the dominant left eigenvector of a gene flow matrix, and they are used as weights in inclusive fitness calculations\(^28,29\).

In this section, we first derive the class-specific RVs and then use them to derive the RVs of daughters and sons for the spring and summer generations.

**Reproductive values of haplodiploids with female hibernation.** The gene-flow matrix is easily derived from Supplementary Fig. 1 by inspecting the outgoing edges from each node and determining what proportion of genes in a 'receiving' node can be attributed to the 'donating' node. Surviving individuals obviously contribute 100% of their genes to their surviving selves, while under haplodiploidy a male offspring derives 100% of his genes from his mother and for a female offspring both her mother and her father get credit for 50% of her gene content. For the haplodiploid FH life history, this gives rise to the following gene flow matrix:

\[ A = \begin{bmatrix} 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & S_m Q_1 \\ 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 \\ \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & 0 \\ \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \]  

Note that the columns that correspond to contributions by males contain the \( Q_1 \) defined in equation (17a,b), which are the mean numbers of mates per male. The RVs are solutions of \( v^T = v^T A \):

\[ v_1 = \frac{1}{2} z_2 F_1 v_1 + S_f v_5 + z_1 F_1 v_7 \]  

\[ v_2 = \frac{1}{2} z_2 F_1 v_1 + S_f v_6 \]  

\[ v_3 = \frac{1}{2} z_2 F_1 v_1 + z_1 F_1 v_7 \]  

\[ v_4 = 0 \]  

\[ v_5 = \frac{1}{2} z_2 F_1 v_1 + z_1 F_1 Q_1 v_2 \]  

\[ v_6 = \frac{1}{2} z_2 F_1 v_1 \]  

\[ v_7 = \frac{1}{2} z_2 F_1 Q_1 v_1 + S_m Q_1 v_2 \]

The RVs of daughters and sons born in the spring and the summer, respectively, can then be calculated as

\[ v_{m1} = v_1 = V (2 - O_m) \]  

\[ v_{m2} = v_2 = \frac{z_1}{z_2} (1 - O_m) \]

Here

\[ O_m = \frac{S_m z_1 F_1}{z_1 G + S_m z_1 F_1} \]  

\[ V = \frac{h z_1 F_1}{z_1 G + S_m z_1 F_1} \]  

\[ G = S_f F_A + \frac{h z_1 F_1}{z_1 G}. \]

Note that the RVs (23a–d) are normalized such that the RV of a daughter born in summer, \( v_{m2} \), is set to unity and the other RVs are multiples of \( v_{m2} \). A crucial quantity is \( 0 \leq O_m \leq 1 \), which can be regarded as measure of generation overlap between spring and summer males. If \( S_m = 0 \) and/or \( Z_1 = 0 \), male generations do not overlap, and the relative within-generation RV of males to females reduce to the familiar \( \frac{Z_1}{Z_2} / Z_2 \) for haplodiploids\(^30\). On the other hand, if there is male generation overlap (\( O_m > 0 \)), then in the spring the RV of males increases with respect to that of females, while in the summer the RV of males decreases with respect to that of females. This explains selection for male-biased spring sex ratios and female-biased summer sex ratios for the FH life cycle, at least as long as helping is rare (see section 'Sex ratios selection').

**Reproductive values of diploids with female hibernation.** The gene-flow matrix differs from the haplodiploid case in that both parents now get credit for 50% of male offspring:

\[ A = \begin{bmatrix} 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & S_m Q_1 \\ 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 \\ \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & 0 \\ \frac{1}{2} z_2 F_1 & \frac{1}{2} z_2 F_1 & 0 & 0 & \frac{1}{2} z_2 F_1 & 0 & \frac{1}{2} z_2 F_1 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ S_f & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_1 & 0 & 0 & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \]  

(27)
The class-specific RVs are then given by:

\[ v_1 = \frac{1}{2} z_2 F_1 v_3 + S_0 v_5 + \frac{1}{2} z_1 F_1 v_7 \]  
(28a)
\[ v_2 = v_1 \]  
(28b)
\[ v_3 = \frac{1}{2} z_2 F_1 v_3 + \frac{1}{2} z_2 F_3 v_7 \]  
(28c)
\[ v_4 = 0 \]  
(28d)
\[ v_5 = (F_2/F_1) v_3 \]  
(28e)
\[ v_6 = \frac{1}{2} z_2 F_1 v_3 \]  
(28f)
\[ v_7 = \frac{1}{2} z_2 F_1 v_3 \]  
(28g)

The spring and summer RVs of daughters and sons are now

\[ v_{11} = \tilde{h} v_1 = V (2 + O_m) \]  
(29a)
\[ v_{1m} = v_1 = V \frac{z_1}{z_4} \left( 2 + O_m \right) + S_0 \frac{z_2}{z_4} (1 - O_m) \]  
(29b)
\[ v_{2} = 2 v_1 = 1 \]  
(29c)
\[ v_{2} = 2 v_1 = 1 \]  
(29d)
Here

\[ O_m = \frac{S_m z_2}{z_4 v F_0 z_2} \]  
(30)

is again a measure of the degree of generation overlap in males. It is slightly smaller than the corresponding quantity for haplodiploids; thus, in diplods the divergence in RVs between females and males will be slightly smaller as well. If there is no male generation overlap, that is, \( O_m = 0 \), then the relative RVs of males to females reduces to the familiar \( z_i / z_i \) for diplods\(^{36}\).

### Reproductive values of haplodiploids with larval diapause

The gene-flow matrix can be derived from Supplementary Fig. 2, as follows:

\[ A = \begin{bmatrix} 0 & 0 & \frac{1}{2} z_2 z_3 F_3 & 0 & \frac{1}{2} z_2 z_3 F_5 & \frac{1}{2} z_2 z_3 F_7 \\ 0 & 0 & \frac{1}{2} z_2 z_3 F_3 & 0 & \frac{1}{2} z_2 z_3 F_5 & \frac{1}{2} z_2 z_3 F_7 \\ \frac{1}{2} z_2 z_3 F_3 & \frac{1}{2} z_2 z_3 F_5 & 0 & 0 & 0 & 0 \\ \frac{1}{2} z_2 z_3 F_3 & \frac{1}{2} z_2 z_3 F_5 & 0 & 0 & 0 & 0 \\ z_1 F_3 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_3 & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \]  
(31)

The corresponding RVs are given by

\[ v_1 = \frac{1}{2} z_2 z_3 F_3 v_3 + S_0 v_5 + z_1 F_1 v_7 \]  
(32a)
\[ v_2 = \frac{1}{2} z_2 z_3 F_3 v_3 + S_0 Q_1 v_6 + z_1 v_5 \]  
(32b)
\[ v_3 = \frac{1}{2} z_2 z_3 F_3 v_3 + \frac{1}{2} z_2 z_3 F_5 v_7 \]  
(32c)
\[ v_4 = 0 \]  
(32d)
\[ v_5 = (F_2/F_1) v_3 \]  
(32e)
\[ v_6 = \frac{1}{2} z_2 z_3 F_1 v_3 \]  
(32f)
\[ v_7 = \frac{1}{2} z_2 z_3 F_1 v_3 \]  
(32g)
\[ v_8 = v_7. \]  
(32h)

The RVs of daughters and sons in spring and summer are now given by

\[ v_{11} = \tilde{h} v_1 = V (2 + O_m) \]  
(33a)
\[ v_{1m} = v_1 = V \frac{z_1}{z_4} \left( 2 + O_m \right) + S_0 \frac{z_2}{z_4} (1 - O_m) \]  
(33b)
\[ v_{2} = 2 v_1 = 1 \]  
(33c)
\[ v_{2} = 2 v_1 = 1 \]  
(33d)

Here

\[ O_m = \frac{S_m z_2}{z_4 v F_0 z_2} \]  
(34a)
\[ O_m = \frac{S_m z_2}{z_4 v F_0 z_2} \]  
(34b)

are measures of generation overlap in females and males, respectively. If \( O_m = 0 \), there is no overlap between male generations, and the relative RVs of males to females again reduces to the familiar \( z_i / z_i \) for haplodiploids. In contrast to the case for the FH life cycle, for the LD life cycle male generation overlap reduces the RV of spring males relative to that of spring females, while the RV of summer males is increased with respect to that of summer females. This explains why selection favours female-biased spring sex ratios and male-biased summer sex ratios (see ‘Sex ratios selection’).

#### Reproductive values of diplods with larval diapause

The gene-flow matrix is now given by

\[ A = \begin{bmatrix} 0 & 0 & \frac{1}{2} z_2 z_3 F_3 & 0 & \frac{1}{2} z_2 z_3 F_5 & \frac{1}{2} z_2 z_3 F_7 \\ 0 & 0 & \frac{1}{2} z_2 z_3 F_3 & 0 & \frac{1}{2} z_2 z_3 F_5 & \frac{1}{2} z_2 z_3 F_7 \\ \frac{1}{2} z_2 z_3 F_3 & \frac{1}{2} z_2 z_3 F_5 & 0 & 0 & 0 & 0 \\ \frac{1}{2} z_2 z_3 F_3 & \frac{1}{2} z_2 z_3 F_5 & 0 & 0 & 0 & 0 \\ z_1 F_3 & 0 & 0 & 0 & 0 & 0 \\ z_1 F_3 & 0 & 0 & 0 & 0 & 0 \end{bmatrix} \]  
(35)

The RVs are then given by the equations

\[ v_1 = \frac{1}{2} z_2 z_3 F_1 v_3 + S_0 v_5 + \frac{1}{2} z_1 F_1 v_7 \]  
(36a)
\[ v_2 = \frac{1}{2} z_2 z_3 F_1 v_3 + S_0 Q_1 v_6 + \frac{1}{2} z_1 F_1 v_7 + S_m v_8 \]  
(36b)
\[ v_3 = \frac{1}{2} z_2 z_3 F_1 v_3 + \frac{1}{2} z_2 z_3 F_5 v_7 \]  
(36c)
\[ v_4 = 0 \]  
(36d)
\[ v_5 = (F_2/F_1) v_3 \]  
(36e)
\[ v_6 = v_5 \]  
(36f)
\[ v_7 = Q_1 v_6 \]  
(36g)
\[ v_8 = v_7. \]  
(36h)

The RVs of daughters and sons in spring and summer are now given by

\[ v_{11} = \tilde{h} v_1 = V (1 + T) \]  
(37a)
\[ v_{1m} = v_1 = V \frac{z_1}{z_4} (1 - O_m) (1 + T) \]  
(37b)
\[ v_{2} = 2 v_1 = 1 \]  
(37c)
\[ v_{2} = 2 v_1 = 1 \]  
(37d)

Here male generation overlap \( O_m \) is given by equation (34b) and

\[ T = \frac{F_a + \frac{1}{2} z_2 z_3 F_1 F_3 O_m}{F_a - \frac{1}{2} z_2 z_3 F_1 O_m} \]  
(38)

Note that \( T \geq 1 \) with equality if there is no male generation overlap. As before, if there is no male generation overlap, that is, \( O_m = 0 \), then the relative RVs of males to females reduces to the familiar \( z_i / z_i \) for diplods.

### Sex ratios selection

In this section, we derive inclusive fitness expressions and corresponding selection differentials for females controlling the spring sex ratio \( z_1 \) and summer sex ratio \( z_2 \).

The inclusive fitness expressions are a focal female’s expected lifetime number of daughters and sons, each weighted by a corresponding coefficient of relatedness and RV, given the focal female’s sex ratios \( z_1 \) and \( z_2 \) in a resident population with sex ratios \( z_1^* \) and \( z_2^* \):

\[ W_1 (z_1, z_1^*) = F_1 (z_1 F_1 v_1 + z_1 O_m v_3^*) + S_1 F_1 (1 + h z_1 z_2^*) (z_1 F_1 v_3 + z_2 F_1 O_m v_3^*) \]  
(39a)
\[ W_2 (z_2, z_2^*) = \frac{h z_1}{z_2} \left( \frac{1}{z_1} F_1 v_1 + \frac{1}{z_2} F_1 v_3 \right) (z_2 F_2 v_3 + z_2 F_2 O_m v_3^*) \]  
(39b)
Variables equipped with an asterisk are evaluated at their resident values $z_i^*$ and $z_j^*$. In outbred populations, the relatedness of a daughter to her mother is $r_{dm} = z_j/z_i$ for both haplodiploids and diplodips, while a son is related to his mother by $r_{ms} = 1$ for haplodiploids and $r_{ms} = 1/2$ for diplodips. The first term in the inclusive fitness (37a) of a spring female correspond to her $F_1$ offspring produced in the spring, while the second term corresponds to her expected number of offspring $F_2 = F_1(1 + bh; F_1)$ produced in the summer, provided she survives with probability $S_t$. Note that her total inclusive fitness increases with the number of female helpers, given by equation (18), which in turn increases with the proportion of daughters $z_1 = 1 - z_i$ produced in the spring.

The corresponding selection differentials are given by:

$$\frac{\partial W_1}{\partial z_1} |_{z_1 = z_i^*} = F_1(1 - r_{dau} v_{11} + r_{son} v_{m1} - Bh(z_r dza v_{12} + z_r r_m v_{m1})), \quad (40a)$$

$$\frac{\partial W_2}{\partial z_2} |_{z_2 = z_j^*} = 1 + f_s F_1 + f_m F_2 - (1 - r_{dau} v_{12} + r_{son} v_{m2}). \quad (40b)$$

Note that in the absence of help ($h = 0$), the sex ratios $z_i$ are in equilibrium if and only if daughters and sons yield the same ‘life-for-life’ relatedness to their mother: $r_{dau} v_{11} = r_{son} v_{m1}$. Otherwise, all else being equal, the presence of female helpers shifts selection in the spring towards more daughters.

**Sex ratio selection in haplodiploids with female hibernation.** Plugging in the reproductive values (23a–d) and the relatedness coefficients, the selection differential (40a) for $z_i$ can be written as

$$\frac{\partial W_1}{\partial z_i} |_{z_1 = z_i^*} = F_1 \left[ \left( -1 + \frac{z_i^*}{z_i} + \frac{4}{z_i} \right) + \frac{S_m}{z_i} \left( 1 - z_i^* \right) - \frac{4}{z_i} \right] \quad (41)$$

The parameter $O_m$ is given by equation (24) and measures overlap of male generations. It is easy to see that in the absence of help ($h = 0$), and with non-overlapping generations of males ($S_m = O_m = 0$), all terms between brackets but the first vanish and the equilibrium sex ratio is $z_i^* = z_i^* = 1/2$. Without help but with overlapping male generations ($S_m > O_m > 0$), the equilibrium spring sex ratio is male-biased, that is, $z_i^* < z_i^*$. With help ($h > 0$), the last term on the right of equation (41) may come to dominate and a female-biased sex ratio may be favoured.

The selection differential (40b) simplifies to

$$\frac{\partial W_2}{\partial z_2} |_{z_2 = z_j^*} = 1 + f_s F_1 + f_m F_2 \left[ -1 + \frac{z_j^*}{z_j} \left( 1 - O_m \right) \right] \quad (42)$$

Without overlapping generations of males ($S_m = O_m = 0$) the equilibrium sex ratio is again unbiased: $z_i^* = z_j^* = 1/2$. If male generations do overlap ($S_m, O_m > 0$) daughters are overproduced in equilibrium: $z_i^* > z_i^*$, that is, a female-biased summer sex ratio.

**Sex ratio selection in diplodips with female hibernation.** The selection differentials (40a and b) simplify to

$$\frac{\partial W_1}{\partial z_1} |_{z_1 = z_i^*} = \frac{z_i^*}{z_i} F_1 \left[ \left( 2 + O_m \right) + \frac{S_m}{z_i} \left( 1 - z_i^* \right) - \frac{4}{z_i} \right] \quad (43a)$$

$$\frac{\partial W_2}{\partial z_2} |_{z_2 = z_j^*} = \frac{z_j^*}{z_j} F_2 \left[ -1 + \frac{z_j^*}{z_j} \left( 1 - O_m \right) \right] \quad (43b)$$

Now the overlapping-generation parameter $O_m$ is given by equation (30). Qualitatively, the same results hold as for haplodiploids: in the absence of help ($h = 0$) and with non-overlapping generations of males ($S_m = O_m = 0$), the equilibrium sex ratios are unbiased ($z_i^* = z_j^* = 1/2$). If male generations overlap ($S_m, O_m > 0$), the equilibrium spring sex ratio is male-biased and the summer sex ratio is female-biased. However, again, a sufficiently large benefit from help favours female-biased sex ratios in the spring.

**Sex ratio selection in haplodiploids with larval diapause.** The selection differentials (40a and b) now simplify to

$$\frac{\partial W_1}{\partial z_1} |_{z_1 = z_i^*} = \frac{z_i^*}{z_i} F_1 \left[ \left( -1 + \frac{z_i^*}{z_i} - \frac{1}{z_i} \right) + \frac{S_m}{z_i} \left( 1 - z_i^* \right) - \frac{4}{z_i} \right] \quad (44a)$$

$$\frac{\partial W_2}{\partial z_2} |_{z_2 = z_j^*} = \frac{z_j^*}{z_j} F_2 \left[ -1 + \frac{z_j^*}{z_j} \left( 1 - O_m \right) \right] \quad (44b)$$

The overlapping generation parameters $O_m$ and $O_n$ are given by equations (34a) and (34b), respectively. In the absence of help ($h = 0$), and with non-overlapping generations of males ($S_m = O_m = 0$), clearly the equilibrium sex ratios are again unbiased: $z_i^* = z_j^* = 1/2$. Without help but with overlapping male generations ($S_m, O_n > 0$), the equilibrium spring sex ratio is female-biased, that is $z_i^* > z_j^*$, while the equilibrium summer sex ratio is male-biased, $z_j^* < z_j^*$, in contrast to the results for the FH life cycle. With help ($h > 0$), a female-biased spring sex ratio is favoured regardless of generation overlap.

**Sex ratio selection in diplodips with larval diapause.** The selection differentials (40a and b) now simplify to

$$\frac{\partial W_1}{\partial z_1} |_{z_1 = z_i^*} = \frac{z_i^*}{z_i} F_1 \left[ \left( -1 + \frac{z_i^*}{z_i} - \frac{1}{z_i} \right) + \frac{S_m}{z_i} \left( 1 - z_i^* \right) - \frac{4}{z_i} \right] \quad (45a)$$

$$\frac{\partial W_2}{\partial z_2} |_{z_2 = z_j^*} = \frac{z_j^*}{z_j} F_2 \left[ -1 + \frac{z_j^*}{z_j} \left( 1 - O_m \right) \right] \quad (45b)$$

The overlapping generation parameter $O_n$ is given by equation (34b) and by equation (38). Qualitatively, the same results hold as for haplodiploids with the LD life cycle: in the absence of help ($h = 0$), and with non-overlapping generations of males ($S_m = O_n = 0, T = 1$), the equilibrium sex ratios are $z_i^* = z_j^* = 1/2$. In the absence of help but with overlapping male generations ($S_m, O_n > 0$), the equilibrium spring sex ratio is female-biased ($z_i^* > z_j^*$), while the equilibrium summer sex ratio is male-biased ($z_j^* < z_j^*$). With help ($h > 0$), a female-biased spring sex ratio is always favoured.

**Selection on helping behaviour.** The inclusive fitness of a focal daughter with helping tendency $h$ in a resident population with helping tendency $h'$ is given by

$$W_h(h, h') = F_1 \left[ (z_r dza v_{12} + z_r r_m v_{m2}) + hB(z_r r_m v_{m2}) \right]. \quad (46)$$

The first term between brackets is the inclusive fitness through daughters and sons obtained by not helping (with probability $h$), while the second term is the inclusive fitness through additional sisters and brothers obtained by helping (with probability $h'$). The appropriate coefficients of relatedness and RVs depend on the specific scenario regarding genetics and life history and will be derived below. The corresponding selection differential is then:

$$\frac{\partial W_h}{\partial h} |_{h = h'} = F_1 \left[ - (z_r dza v_{12} + z_r r_m v_{m2}) + hB(z_r r_m v_{m2}) \right]. \quad (47)$$

Therefore helping will be selected for whenever

$$B > \frac{4 - 2O_m}{4 - O_m} \equiv B_{\min} \quad (49)$$

Since

$$0 \leq O_m = \text{San}z_i F_1 z_j G + \text{San}z_j F_1 \leq 1, \quad (50)$$

it follows that

$$\frac{1}{2} \leq B_{\min} \leq 1 \quad (51)$$

A greater generation overlap in males favours a lower benefit threshold for helping behaviour to evolve, which in turn is favoured by a male-biased spring sex ratio and female-biased summer sex ratio—precisely the sex ratios favoured by selection in the FH life cycle.

**Selection on helping behaviour in diplodips with female hibernation.** Using the RVs (27) and the relatedness coefficients $r_{san} = 1/2$, the selection differential (45) now reduces to

$$\frac{\partial W_h}{\partial h} |_{h = h'} = \frac{z_i^* F_1}{2} \left[ (2 - O_n) + \frac{hB}{4 - O_n} \right] \quad (52)$$

Obviously helping will be selected for whenever

$$B > 1 \quad (53)$$

Unlike in haplodiploids, for diplodips with the FH life cycle generation overlap in males does not favour a lower helping threshold and helping is always more difficult to evolve than in haplodiploids.

**Selection on helping behaviour in haplodiploids with larval diapause.** Using the RVs (33c and d) and the relatedness coefficients $r_{san} = 1/2$ and $r_{son} = 1/2$, the
Table 2 | Parameter values used in the individual based simulations.

| Parameter | Value |
|-----------|-------|
| \( N \)   | 5,000 |
| \( \mu \) | 0.01  |
| \( \sigma \)| 0.01  |
| \( t \)  | 25,000|

Description:
- Number of nests or founding females in the spring
- Mutation rate
- s.d. of normal distribution from which mutation values are drawn
- Number of years the simulation was run

Selection differential (47) now becomes

\[
\frac{\partial W_h}{\partial h_{\text{bro}}} = \frac{1}{2} z_h F_1 \left[ - (2 + (1 - O_h)O_m) + \frac{1}{2} B (4 + (1 - O_h)O_m) \right]
\]

where the generation-overlap parameters \( O_h \) and \( O_m \) are given by equation (34a and b). Now helping will be selected for whenever

\[
B > \frac{4 + 2(1 - O_h)O_m}{4 + (1 - O_h)O_m} = B_{\text{min}}
\]

Whenever there is some generation overlap, \( B_{\text{min}} > 1 \) and in general

\[
1 \leq B_{\text{min}} \leq \frac{4}{3}
\]

In general, therefore, helping in haplodiploids with the LD life cycle is harder to evolve than in haplodiploids with the FH life cycle.

Selection on helping behaviour in diplods with larval diapause. Using the RVs (37c,d) and the relatedness coefficients \( r_{\text{mam}} = r_{\text{mam}} = \frac{1}{2} \), the selection differential (47) now becomes

\[
\frac{\partial W_h}{\partial h_{\text{bro}}} = \frac{1}{2} z_h F_1 (1 + T) (-1 + B)
\]

Just like in diplods with the FH life cycle, helping is selected for whenever

\[
B > 1
\]

Therefore, in contrast to the FH life cycle, for the LD life cycle haplodiploidy makes helping more difficult to evolve.

Coevolution of sex ratios and helping behaviour. Coevolution of sex ratios and helping behaviour was modelled using a standard adaptive dynamics approach. For each trait \( x \in \{ z_1, z_2, h_t \} \), the dynamics over evolutionary time \( t \) is given by

\[
\frac{dx}{dt} = K \frac{\partial W_h}{\partial x}
\]

The scaling constant \( K = 0.115 \) was chosen to make the adaptive dynamics results commensurate with the results from individual-based simulations (see section 'Individual-based simulations'). The selection differentials are given above.

Numerical integration of differential equation (59) was carried out with R 3.1.0 (ref. 67), using the package deSolve.68

Polyandry hampers the evolution of helping. We look at two types of polyandry:
(1) serial monogamy, where surviving spring females mate for a second time with a different male; sperm from the first mating is not stored. Thus, females from the first and second brood are half-sisters, and the relatedness coefficient must be replaced accordingly \((r_{\text{mam}} = \frac{1}{4})\). (2) Polyandry, where autumn females mate with more than one male. Their sperm is stored and used by surviving females to produce a second brood. The number of males that females mate with determines the coefficient of relatedness between females. We show calculations only for the FH life cycle.

Serial monogamy. Since a female does not store sperm, class 6 disappears; otherwise the demography remains the same. The reproductive values are now given by

\[
\begin{align*}
v_1 &= \frac{1}{2} z_1 F_1 v_3 + z_1 z_2 F_1 v_3 \\
v_2 &= \frac{1}{2} z_1 z_2 F_1 v_3 \\
v_3 &= \frac{1}{2} z_2 z_2 F_1 v_3 + z_2 z_2 z_1 Q_2 v_2 \\
v_4 &= 0 \\
v_5 &= (F_3 / F_3) v_3 \\
v_6 &= \frac{1}{2} z_2 z_2 F_3 v_1 + \frac{1}{2} z_2 z_2 Q_1 v_1 + z_2 z_2 Q_2 v_2
\end{align*}
\]

Here \( Q_1 = u_1 / u_2 = S_1 / (z_1 F_1) \). To analyse the evolution of helping we only need the RVs of summer daughters and summer sons, where we normalize the former to unity:

\[
\begin{align*}
v_{21} &= 1 \\
v_{32} &= \frac{1}{z_2} (1 - O_m - O_t) \\
O_h &= \frac{z_1 z_2 S_1 F_1}{z_2 G + S_2 z_1 F_1}
\end{align*}
\]

The selection differential is the same as (47) except that \( r_{\text{mam}} \) is replaced by \( r_{\text{mam}} \) and it simplifies to

\[
\frac{\partial W_h}{\partial h_{\text{bro}}} = \frac{1}{2} z_h F_1 (1 - O_m - O_t) (-1 + \frac{1}{2} B)
\]

Clearly, positive selection for helping requires

\[
B > 1
\]

Effect of polyandry on selection in haplodiploids. The coefficient of relatedness between female offspring of the same mother, for haplodiploids in a fully outbred population is given by

\[
r_{\text{mam}} = \frac{1}{4} + \frac{1}{2} p = \frac{1}{4} + \frac{1}{2} \frac{m \text{r}}{m + m_{\text{r}}},
\]

where \( p = \sum_i p_i^2 \) is the probability that two females with the same mother share the same father, which equals the sum of squared paternity shares \( p_i \) of all males that have mated with the same female. The inverse of \( p \) in turn defines the effective number of mates \( m \text{r} \) per female, which is bounded above by the average number of mates per female.29,69 Replacing the coefficient of relatedness (65) into the selection gradient (47) we get

\[
\frac{\partial W_h}{\partial h_{\text{bro}}} = \frac{1}{2} z_h F_1 \left[ - (2 - O_m) + \frac{1}{2} B (2(1 + 1/m_\text{r}) - O_m) \right],
\]

and the condition for the evolution of helping is

\[
B > \frac{4 - 2O_m}{2(1 + 1/m_\text{r}) - O_m}.
\]

If we assume that females only mate with a single male \((m_\text{r} = 1)\), we obtain the result for monogamy (49). If we assume \( m_\text{r} > 1 \), then \( B_{\text{min}} > 1 \), but if \( 1 < m_{\text{r}} < 2 \), then it is possible that \( B_{\text{min}} < 1 \) as long as \( O_m > 0 \) (Supplementary Fig. 4). In the limit of infinitely many males, such that all females with the same mother are half-sisters, we obtain the same result as in serial monogamy that helping requires helpers to be more twice as efficient \((B_{\text{min}} > 2)\) at raising sibling as at raising offspring. Condition (67) shows that haplodiploidy can benefit the evolution of helping under the female hibernation scenario even if females are not strictly monogamous, but this effect diminishes as females are more promiscuous (Supplementary Fig. 4).

Effect of polyandry on selection in diplods. The coefficient of relatedness under diplod between two sisters is given by

\[
r_{\text{mam}} = \frac{1}{4} (1 + 1/m_\text{r}),
\]

where, like before, \( m_\text{r} \) is the effective number of males females mate with. The helping selection differential then reduces to

\[
\frac{\partial W_h}{\partial h_{\text{bro}}} = \frac{1}{2} z_h F_1 (2 - O_m)(-1 + \frac{1}{2} B (1 + 1/m_\text{r})),
\]
and the condition for the evolution of helping is

\[ B > \frac{2}{1 + 1/m} \]  

(70)

From condition (70) it is clear that if females mate with only one male \( m = 1 \) we recover the previous result (53) that helpers need to be more efficient at raising their sibling than their offspring \( (B > 1) \). If females mate with infinitely many males, the evolution of helping requires helpers to be more than twice as efficient at raising sibling as offspring \( (B > 2) \). All in all, higher levels of female promiscuity increases the necessary benefits for the evolution of helping behaviour.

**Individual-based simulations.** Evolutionary dynamics were also analysed using individual based simulations. We simulated a population that has the life cycle exemplified by Supplementary Figs 1 and 2. Each individual is characterized by three loci. The loci represent proportion of males in the spring \( (s_L) \) and summer \( (s_S) \), and the probability that a female born in the spring stays to become a helper \( (h) \). We assume that the phenotypic effect of an allele in each locus is a continuous quantity that ranges between 0 and 1. Sex ratios in the colony are in full control \( (12) \), and the condition for the evolution of helping is a hierarchy of models. Proc. Natl Acad. Sci. 88, 10993 (1991).

12. Fawcett, T. W., Kuiper, B., Weissing, F. J. & Pen, I. Sex-ratio control erodes sexual selection, revealing evolutionary feedback from adaptive plasticity. Proc. Natl Acad. Sci. 108, 15925–15930 (2011).

13. Pen, I. & Weissing, F. J. Sex-ratio optimization with helpers at the nest. Proc. R. Soc. B Biol. Sci. 267, 539–543 (2000).

14. Stubblefield, J. W. & Orzack, S. H. Resource transfers and evolution: helpful offspring and sex allocation. Theor. Popul. Biol. 83, 64–81 (2013).

15. Ross, L., Gardner, A., Hardy, N. & West, S. A. Ecology, not the genetics of sex determination, determines who helps in eusocial populations. Curr. Biol. 23, 2385–2387 (2013).

16. Gadagkar, R. Demographic predisposition to the evolution of eusociality: a hierarchy of models. Proc. Natl Acad. Sci. 88, 10993 (1991).

17. Hunt, J. H. & Amdam, G. V. Bivoltinism as an antecedent to eusociality in the paper wasp genus Polistes. Science 308, 264–267 (2005).

18. Boomsma, J. J. Lifetime mating patterns and the evolution of eusociality. Philos. Trans. R. Soc. B Biol. Sci. 364, 3191–3207 (2009).

19. Herrera, E. A. Ecological and genetic distribution of eusociality: the case for kin selection. Behav. Ecol. 24, 331–333 (2013).

20. Fromhage, L. & Kokko, H. Monogamy and haplodiploidy act in synergy to promote the evolution of eusociality. Nat. Commun. 2, 597 (2011).

21. Taylor, P. D. The selection differential in quantitative genetics and ESS models. Evolution 50, 2106–2110 (1996).

22. Pen, I. & Weissing, F. J. Towards a unified theory of cooperative breeding: the role of ecology and life history re-examined. Proc. R. Soc. B Biol. Sci. 267, 2411–2418 (2000).

23. Charnov, E. L. Evolution of eusocial behavior: offspring choice or parental optimization? J. Theor. Biol. 66, 451–456 (1977).

24. Gardner, A. Sex-biased dispersal of adults mediates the evolution of altruism among juveniles. J. Theor. Biol. 262, 339–345 (2010).

25. Gardner, A., Alpedrinha, J. & West, S. A. Haplodiploidy and the evolution of eusociality: split sex ratios. Am. Nat. 179, 240–256 (2012).

26. Dieckmann, U. & Law, R. The dynamical theory of coevolution: a derivation from stochastic ecological processes. J. Math. Biol. 34, 579–612 (1996).

27. Pen, I. & Taylor, P. D. Modelling information exchange in worker–queen conflict over sex allocation. Proc. R. Soc. B Biol. Sci. 272, 2403–2408 (2005).

28. Taylor, P. D., Wild, G. & Gardner, A. Direct fitness or inclusive fitness: how shall we model kin selection? J. Evol. Biol. 20, 301–309 (2007).

29. Boomsma, J. J. & Ratniek, F. L. W. Paternity in eusocial insects. Philos. Trans. R. Soc. B Biol. Sci. 351, 947–975 (1996).

30. Schwarz, M. P., Richards, M. H. & Danforth, B. N. Changing paradigms in insect social evolution: insights from Halictine and Allophine bees. Annu. Rev. Entomol. 52, 127–150 (2007).

31. Prys-Jones, O. E. & Corbet, S. A. Bumblebees (Pelagic Publishing, 2011).

32. Sakagami, S. F. Specific differences in the bionomic characters of bumblebees: a comparative review. J. Fac. Sci. Hokkaido Univ. Ser. VI Zool. 20, 390–447 (1976).

33. Hunt, J. H. Trait mapping and salience in the evolution of eusocial vespid wasps. Evolution 53, 225–237 (1999).

34. Smith, A. R., Wcislo, W. T. & O’Donnell, S. Assured fitness returns favor sociality in a mass-provisioning sweat bee, Megalopa genalis (Hymenoptera: Halictidae). Behav. Ecol. Sociobiol. 51, 14–21 (2003).

35. Danforth, B. N. Evolution of sociality in a primitive eusocial lineage of bees. Proc. Natl Acad. Sci. 99, 286–290 (2001).

36. Richards, M. H. Nesting biology and social organization of Halictus sexcinctus (Fabricius) in southern Greece. Can. J. Zool. 79, 2211 (2001).

37. Hogendoorn, K. & Leys, R. Life-cycle of Halictus rubicundus Christ (Hymenoptera: Halictidae) in the Netherlands: comparison of two populations. J. Kans. Entomol. Soc. 72, 160–166 (1999).

38. Tepedino, V. J. & Parker, F. D. Alternation of sex ratio in a partially bivoltine wasp, Trypoxylon (Trypoxylul) politum (Hymenoptera: Sphexidae). Behav. Ecol. Sociobiol. 53, 23–30 (2003).

39. Alpedrinha, J., West, S. A. & Gardner, A. Haplodiploidy and the evolution of eusociality: worker reproduction. Proc. Natl Acad. Sci. 108, 421–438 (2013).
46. Alpedrinha, J., Gardner, Andy & West, S. A. Haplodiploidy and the evolution of eusociality: worker revolution. Am. Nat. 184, 303–317 (2014).

47. Hughes, W. O. H., Oldroyd, B. P., Beekman, M. & Ratnieks, F. L. W. Ancestral monogamy shows kin selection is key to the evolution of eusociality. Science 320, 1213–1216 (2008).

48. Boomsma, J. J. Beyond promiscuity: mate-choice commitments in social breeding. Philos. Trans. R. Soc. B: Biol. Sci. 368, 20120050 (2013).

49. Nonacs, P. Monogamy and high relatedness do not preferentially favor the evolution of cooperation. BMC Evol. Biol. 11, 58 (2011).

50. Olejarz, J. W., Allen, B., Veller, C. & Nowak, M. A. The evolution of non-reproductive workers in insect colonies with haplodiploid genetics. eLife 4, e08918 (2015).

51. Davies, N. G. & Gardner, A. Monogamy promotes worker sterility in insect societies. Preprint at bioRxiv http://biorxiv.org/content/early/2016/06/16/059154 (2016).

52. Grafen, A. in Behavioural Ecology: An Evolutionary Approach (eds Krebs, J. R. & Davies, N. B.) (Sinauer, 1984).

53. Hunt, J. H. A conceptual model for the origin of worker behaviour and adaptation of eusociality. J. Evol. Biol. 25, 1–19 (2012).

54. West-Eberhard, M. J. in Proceedings of Tenth International Congress of Animal Societies: Theories and Facts (eds Brown, J. L., Kikkawa, J. & Ito, Y.) (Int. Union Study of Social Insects, 1987).

55. West-Eberhard, M. J. in Chemistry and biology of social insects (eds Eder, J. & Rembold, H.) (J. Peperny, 1987).

56. Shintani, Y. Quantitative short-day photoperiodic response in larval development and its adaptive significance in an adult-overwintering cerambycid beetle, Phytoeca rufiventris. J. Insect Physiol. 57, 1053–1059 (2011).

57. Kocher, S. D. et al. Transitions in social complexity along elevational gradients reveal a combined impact of season length and development time on social evolution. Proc. R. Soc. B Biol. Sci. 281, 20140627 (2014).

58. Field, J., Paxton, R. J., Soro, A. & Bridge, C. Cryptic plasticity underlies a major evolutionary transition. Curr. Biol. 20, 2028–2031 (2010).

59. Fucini, S., Di Bona, V., Mola, F., Piccaluga, C. & Lorenzi, M. C. Social wasps without workers: geographic variation of caste expression in the paper wasp Polistes biglumis. Insectes Soc. 56, 347–358 (2009).

60. Crozier, R. H. Advanced eusociality, kin selection and male haploidy. Asct. J. Entomol. 47, 2–8 (2008).

61. Nonacs, P. Resolving the evolution of sterile worker castes: a window on the advantages and disadvantages of monogamy. Biol. Lett. 10, 20140089 (2014).

62. Caswell, H. Matrix Population Models (Sinauer Associates, 2000).

63. Brommer, J., Kokko, Hanna & Pietiäinen, H. Reproductive effort and reproductive values in periodic environments. Am. Nat. 155, 454–472 (2000).

64. Fisher, R. A. The Genetical Theory of Natural Selection (Clarendon, 1930).

65. Taylor, P. D. Allele frequency change in a class-structured population. Am. Nat. 135, 95–106 (1990).

66. Taylor, P. D. Inclusive fitness models with two sexes. Theor. Pop. Biol. 34, 145–168 (1988).

67. R Development Core Team. A Language and Environment for Statistical Computing (R foundation for statistical computing Vienna, 2014) http://www.R-project.org.

68. Soetaert, K., Petzoldt, T. & Setzer, R. W. Solving differential equations in R: package deSolve. J. Stat. Soft. 33, 1–25 (2010).

69. Schärer, L. & Pen, I. Sex allocation and investment into pre- and post-copulatory traits in simultaneous hermaphrodites: the role of polyandry and local sperm competition. Philos. Trans. R. Soc. B 268, 20120052 (2013).

Acknowledgements
We thank Franzo Weissing, Bram Kuijper for fruitful discussions and Sandra Rehan for commenting on an earlier version of the manuscript.

Author contributions
A.E.Q. and I.P. initiated, planned and coordinated the project; A.E.Q. and I.P. constructed the model and analysed its outcome. A.E.Q. and I.P. wrote the paper.

Additional information
Supplementary Information accompanies this paper at http://www.nature.com/naturecommunications

Competing interests: The authors declare no competing financial interests.

Reprints and permission information is available online at http://npg.nature.com/reprintsandpermissions/

How to cite this article: Quiniones, A. E. & Pen, I. A unified model of Hymenopteran preadaptations that trigger the evolutionary transition to eusociality. Nat. Commun. 8, 15920 doi: 10.1038/ncomms15920 (2017).

Publisher’s note: Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.