

Sex ratio adjustment and kin discrimination in malaria parasites

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Malaria parasites and related Apicomplexans are the causative agents of some of the most serious infectious diseases of humans, companion animals, livestock and wildlife. These parasites must undergo sexual reproduction to transmit from vertebrate hosts to vectors, and their sex ratios are consistently female-biased. Sex allocation theory, a cornerstone of evolutionary biology, is remarkably successful at explaining female-biased sex ratios in multicellular taxa, but has proved controversial when applied to malaria parasites. Here we show that, as predicted by theory, sex ratio is an important fitness-determining trait and *Plasmodium chabaudi* parasites adjust their sex allocation in response to the presence of unrelated conspecifics. This suggests that *P. chabaudi* parasites use kin discrimination to evaluate the genetic diversity of their infections, and they adjust their behaviour in response to environmental cues. Malaria parasites provide a novel way to test evolutionary theory, and support the generality and power of a darwinian approach.

The allocation of resources to male versus female offspring is one of the most well understood topics in evolutionary biology, and provides some of the best evidence for darwinian adaptation in the natural world^{1–5}. In many cases, simple theory can successfully predict when, why and by how much organisms should adjust their offspring sex ratio in response to changes in their environment. However, despite over a century of research effort directed at malaria (*Plasmodium*) and related Apicomplexan (including *Haemoproteus* and *Leucocytozoon*) blood parasites, evolutionary biologists have not been able to explain their sex ratios^{6–10}. These parasites replicate asexually, and a proportion of these asexually produced parasites develop into sexual stages, termed gametocytes. When taken up in an insect vector's blood meal, gametocytes differentiate into gametes and mate. In *Plasmodium* species, male gametocytes can differentiate into a maximum of eight gametes and each female differentiates into a single gamete. Sex allocation in *Plasmodium* is consistently female-biased when measured across populations, but sex ratios vary extensively when measured throughout individual infections^{11–15}. Sex is crucial for transmission to vectors, and so there is a drive to identify drugs and develop vaccines that block this process. Yet remarkably little is understood as to why such female-biased and variable sex allocation strategies have evolved.

Hamilton's theory of 'local mate competition' (LMC) predicts that female-biased sex allocation is favoured when genetically related males compete for mates; because males can each fertilize more than one female, an equal sex ratio would result in a wasteful surfeit of male gametes⁴. For malaria parasites, LMC predicts that the unbeatable sex allocation strategy (investment into males relative to females) depends on the inbreeding rate, and is given by the equation $z^* = (1 - f)/2$, where z^* is the proportional allocation into males versus females and f is Wright's coefficient of inbreeding^{16–19}. When inbreeding does not occur, owing to a large number of genotypes represented in mating groups (no inbreeding; $f = 0$), the unbeatable strategy is to produce an equal number of males and females²⁰ ($z^* = 1/2$). Conversely, when only one or a small number of genotypes are present (inbreeding; $f > 0$), female-biased sex allocation is favoured⁴ ($z^* < 1/2$).

LMC is one of the most successful theories in evolutionary biology and its explanatory power has been demonstrated in a variety of taxa²¹ including plants, snakes, insects, mites, worms and fish, but the application of LMC theory to Apicomplexan parasites has proved controversial and inconclusive. There is considerable variation in the inbreeding rate (as allowed by the number of multiple infections) experienced by parasites within and across species of Apicomplexa^{22–24}. LMC is supported by correlations between observed sex ratio and inbreeding rate across several Apicomplexan taxa, with female-biased sex ratios occurring in populations with higher rates of inbreeding^{19,25,26}. However, contradictory data are as numerous: (1) sex ratios in populations of *Haemoproteus* bird parasites do not correlate with the inferred genetic diversity of their infections and are consistently less female biased than expected⁹; (2) LMC theory cannot explain the considerable variation in sex ratios observed during experimental infections^{8,19,26–28}; (3) the inbreeding rate is not the only factor shaping sex allocation, because *Plasmodium* parasites facultatively alter their sex ratio in response to changes in host anaemia^{11,13}; and (4) successful transmission to vectors appears unrelated to mating group sex ratio^{29,30}.

Failure to understand sex allocation in malaria parasites poses problems for both medical science and evolutionary biology. If an evolutionary framework cannot explain a relatively simple trait like parasite sex allocation, there is no reason to believe that it can be usefully applied to more complex traits such as virulence. Also, organisms such as parasites and microbes present a novel and independent test for the explanatory power and generality of an evolutionary theory that has been largely developed to explain the biology of metazoan taxa such as insects, birds and mammals³¹. Explicitly testing whether sex allocation in malaria parasites is shaped by LMC has not previously been possible, because the required experimental methods and techniques have only just become available.

First, we test the basic assumptions of LMC as applied to malaria parasites. We test whether females are the more limiting sex for mating group productivity and whether the relationship between sex ratio and fitness varies in the manner predicted by theory^{1,4}. Even though LMC has been so successfully applied to other taxa, this

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fundamental assumption has yet to be properly tested in any species. Such data are lacking owing to the difficulties in manipulating this trait independently of confounding variables, and the very success of LMC theory has suggested that such a test is unnecessary. We also test whether there is genetic variation for patterns of within-infection sex allocation in malaria parasites, and examine whether these patterns follow the predictions of theory^{32,33}. Second, we test the key predictions of LMC by manipulating the number of genotypes present in infections (inbreeding rate) and investigate whether focal genotypes adjust their sex allocation strategy in response^{13,26,34,35}. Facultative sex-ratio adjustment in response to genetic diversity would confirm that sex allocation strategies in malaria parasites are as sophisticated as those observed in multicellular taxa, that they can be explained by evolutionary theory, and that malaria parasites can discriminate kin from non-kin.

Fitness consequences of sex allocation

We used two genetically transformed lines of the rodent malaria *Plasmodium berghei* to examine the fitness consequences of sex ratio variation^{36,37}. One parasite line (*pb48/45-ko*) cannot express genes essential for male-gamete fertility, so it is limited to female function; the other (*pb47-ko*) cannot express genes essential for female-gamete fertility, so it has only male function. Otherwise, both lines are genetically identical, exhibit normal infection dynamics and their gametocyte and gamete production is the same as their wild-type ancestor. By mixing parasites from these lines in different proportions and using *in vitro* fertilization culture methods, we directly manipulated mating-group sex ratio (proportion of male gametocytes) and measured the resulting reproductive success as the number of zygotes (ookinetes) produced³⁸. As predicted by theory, mating success is maximized at intermediate sex ratios, indicating that sex allocation in malaria parasites is likely to be under stabilizing selection (Fig. 1: linear term, $F_{1,90} = 40.97$, $P < 0.0001$; and quadratic term, $F_{1,90} = 13.81$, $P = 0.004$). These data also support LMC theory, as mating group reproductive success is maximized at female-biased sex ratios (maximum 33% males: 95% confidence index (CI), 20–39%; see Supplementary Information). We estimate that each male produces an average of 2.03 viable gametes (95% CI, 1.56–4.00), revealing that, on average, male gametocytes fail to realize their potential fecundity of eight gametes and will become limiting to mating group productivity at extremely female-biased sex ratios. Owing to biological constraints, difficulty controlling for other relevant trade-offs, and genotype-by-environment interactions, mapping fitness across the range of sex allocation strategies has not

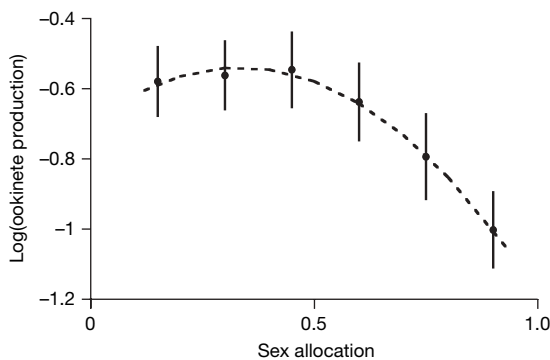


Figure 1 | The fitness consequences of sex ratio variation. The relationship between sex allocation (given as proportion male) and fitness (given as log(ookinete production)) varies in the manner predicted by theory, revealing that this life-history trait of malaria parasites is important and under selection. Shown is log-transformed mean ookinetes ($\times 10^6 \text{ ml}^{-1}$) produced in 19 cross-factored sets of *P. berghei* cultures spanning 0–100% males ($R^2 = 0.77$); dashed line is the fitted relationship. As expected, no ookinetes were produced in our control groups of 0% and 100% males, and we excluded these data from our analysis. Error bars, \pm s.e.m.

previously been possible for malaria parasites or any other taxa. Therefore, these data are the first to unequivocally support a fundamental assumption of sex allocation as applied to any species.

Genetic variation for sex allocation

We then used wild-type clonal genotypes of the rodent malaria *P. chabaudi* to test, for the first time, whether there is within-species genetic variation for patterns of sex allocation in malaria parasites. We sequenced six *P. chabaudi* genotypes and determined that our recently developed quantitative reverse-transcription PCR assays³⁹ for measuring sex ratios could be applied to all of them (see Supplementary Information). We initiated replicate controlled infections of each of these genotypes and measured infection parameters daily throughout the acute phase. We found that our six genotypes followed four significantly different sex allocation patterns throughout their infections (Fig. 2: $\chi^2_{55} = 159.55$, $P < 0.0001$; see Supplementary Table 1). Such genetic variation is required for selection to act, but could itself be adaptive and reflect differences in the number of gametes produced by males of different genotypes, a parameter which may also vary throughout infections^{30,32}. If this is the case, then we can estimate the fecundity of male gametocytes from sex ratio (see Supplementary Information). In addition to controlling for variation across days, our analyses controlled for infection parameters (including virulence, anaemia, asexual parasite and gametocyte density) that could influence sex ratio and confound differences between the genotypes. Only parasite density remained in the minimal model, which correlated positively with sex ratio ($\chi^2_1 = 5.25$; $P = 0.022$; slope = $(27.35 \pm 13.41) \times 10^6 \text{ ml}^{-1}$).

Sex ratio variation during infections

In addition to the presence of co-infecting genotypes, malaria parasites are predicted to allocate sex in response to factors that compromise their fertilization success in blood meals^{11,26,32,33}. Specifically, a lower female-bias is predicted for a given inbreeding rate when the fertility of males is reduced by factors such as low gametocyte density, host anaemia and immunity. There are two broad reasons why these factors are expected to reduce fertilization success. First, when gametocyte density is low and/or hosts are anaemic, there is a stochastic risk of too few male gametocytes being taken up in blood meals to fertilize the females. Second, if the appearance of host factors reduces the ability of male gametocytes to produce viable gametes, there will

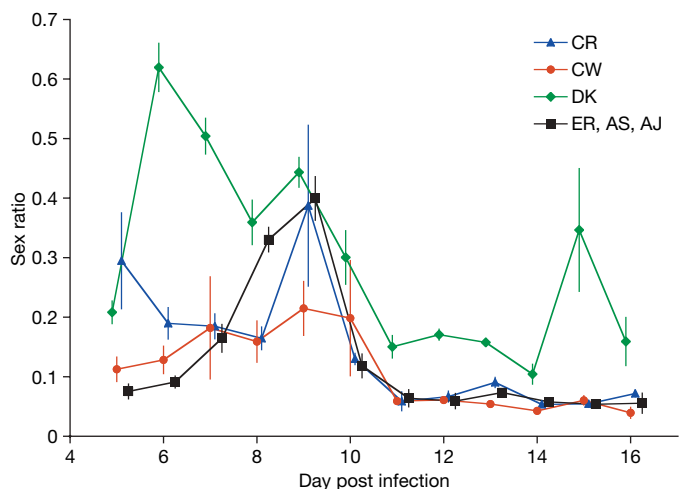


Figure 2 | Genetic variation in patterns of sex allocation. *P. chabaudi* genotypes exhibit significant genetic variation in the sex ratios produced throughout their infections. Genotypes DK, CW and CR all followed significantly different sex allocation patterns but AS, AJ and ER could be grouped together. Here, and in Figs 4–6, sex ratio is given as proportion male. The means from 30 independent infections are presented and the x-axis is jittered for clarity. Error bars, \pm s.e.m.

not be enough male gametes in the blood meal to fertilize the females. When few gametocytes are able to interact in a blood meal and/or male gamete production is low, parasites are expected to increase their investment in male gametocytes to ensure their females are fertilized. Data from our single genotype infections support these predictions, and show that throughout infections sex ratios correlate negatively with red blood cell density ($\chi^2_3 = 15.86$; $P = 0.0012$) and gametocyte density ($\chi^2_3 = 22.11$; $P < 0.0001$), but positively with parasite density (which is related to the strength of host immune responses; $\chi^2_3 = 35.35$; $P < 0.0001$), and that there is genetic variation for these patterns (Fig. 3 and Supplementary Table 2).

Genetic diversity and sex allocation

Having verified that the key assumptions of sex allocation theory are appropriate to the biology of malaria parasites, we tested whether they can facultatively respond to LMC. We compared sex ratios produced by our six single-genotype infections to sex ratios in mixed-infections consisting of all six genotypes. Our above analyses support the prediction that parasite sex ratios are influenced by other parameters^{11,26,32,33}, so we predicted that any response to the presence of co-infecting genotypes is most likely to be detected before these

factors exert their confounding influences. For all of our treatment groups, day 8 post infection (PI) was the modal day of peak parasite density, and parasite density was subsequently reduced by immune factors, anaemia and competition between genotypes. Therefore, we restricted our analyses to the period between infections becoming patent and reaching their peaks (days 5–7 PI). As predicted by LMC, sex ratios in six-genotype infections were less female-biased than in single-genotype infections, and the magnitude of this effect varied through infections (Fig. 4: $\chi^2_2 = 19.93$, $P < 0.0001$; see Supplementary Table 3). As with the inbreeding rate and sex ratio, we found a significant negative correlation between gametocyte density and sex ratio ($\chi^2_1 = 11.94$; $P < 0.0001$; slope = $(-28.20 \pm 8.18) \times 10^{-3} \text{ ml}^{-1}$). Using the model of ref. 25, and assuming that male gametocytes produce up to 8 gametes on average, the predicted evolutionary stable sex ratios for single- and six-genotype infections are 0.11 (proportion male) and 0.42 respectively. At the start of infections, our data support these predictions: on days 5 and 6 PI, sex ratios were respectively 0.14 ± 0.02 and 0.20 ± 0.04 in the single-genotype infections, and 0.46 ± 0.09 and 0.42 ± 0.07 in the six-genotype infections.

Sex allocation in focal genotypes

The best way to test whether parasites respond to LMC is to test whether focal genotypes alter their sex allocation in response to the presence of co-infecting genotypes. We were able to examine this for the three *P. chabaudi* genotypes AJ, AS and ER in single- and double-genotype infections. Our quantitative PCR assays could distinguish genotype AJ from genotypes AS and ER but not between AS and ER (see Supplementary Information). Therefore, we tested whether AS and ER independently altered their sex allocation when co-infecting with AJ, and whether AJ altered its sex allocation when co-infecting with either AS or ER. As we could confirm that parasites from both genotypes were present throughout the mixed infections, we were able to analyse sex ratios when parasites are in the growth phase (parasite density increasing) and post-peak phase (parasite density declining) of their infections. We split our analyses into the pre-peak period (days 5–7 PI) and the post-peak phase (days 8–12 PI). Because sex ratios produced by AJ when co-infecting with AS were not significantly different to those produced when co-infecting with ER (growth phase; $\chi^2_1 = 1.66$, $P = 0.198$; post-peak phase; $\chi^2_1 = 0.25$, $P = 0.617$), we grouped these AJ data. In the growth phase of their infections, both AJ and ER increased their investment in males when co-infecting with another genotype, as predicted by LMC, but AS did not (Fig. 5; $\chi^2_2 = 9.98$, $P = 0.007$; see Supplementary Table 4a). Sex ratio adjustment was greater in AJ than ER and all other infection parameters fitted in the maximal model were non-significant. In contrast, during the post-peak phase only AJ increased investment in males when infecting with a second genotype; AS decreased

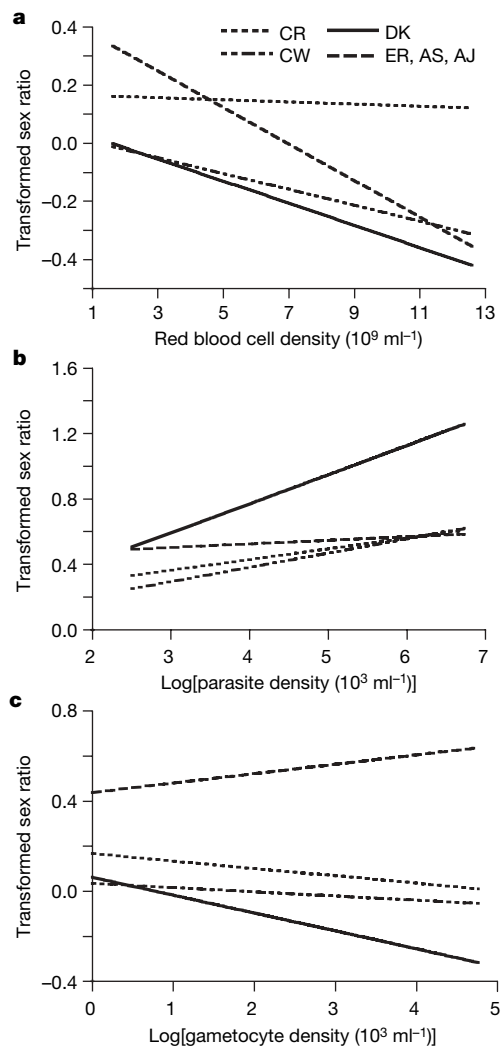


Figure 3 | Explaining sex ratio variation throughout infections. Sex ratios (arcsin square-root transformed) of *P. chabaudi* correlate with the density of: **a**, red blood cells; **b**, parasites; and **c**, gametocytes. Lines are fitted from the estimates predicted by the minimal model using infection parameters observed 48 h before sex ratios (see Supplementary Information). Genotypes are grouped according to the four different sex ratio patterns followed throughout 30 independent infections (Fig. 2).

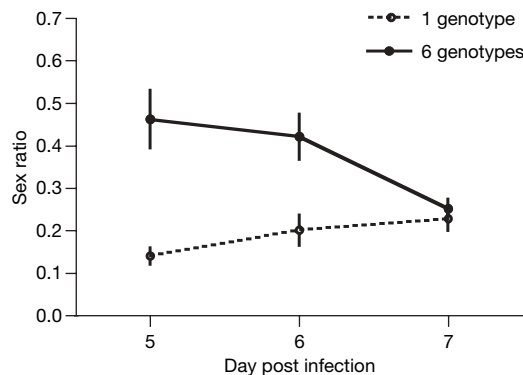


Figure 4 | Sex ratio varies with the genetic diversity of *P. chabaudi* infections. Infections with six genotypes produced significantly less female biased sex ratios than those with one genotype, but only at the start of infections. Means are presented from 40 independent infections. Error bars, \pm s.e.m.

investment in males and sex ratios were not significantly altered in ER (Fig. 6; $\chi^2_2 = 27.33$, $P < 0.0001$; see Supplementary Table 5a). All other infection parameters fitted in the maximal model were non-significant apart from red blood cell density, which correlated positively with sex ratio ($\chi^2_1 = 6.89$, $P = 0.009$; slope = $(32.91 \pm 13.06) \times 10^6 \text{ ml}^{-1}$).

Facultative sex allocation in response to infection genetic diversity of genotypes ER and AJ (when in two different co-infection scenarios) provides unequivocal support for LMC. Given this result, we extended our analysis to test whether sex allocation of focal genotypes correlates with their relative representation in infections. Theory predicts that genotypes making a small relative contribution of gametocytes to the mating group should invest in more males than when making a relatively large contribution⁴⁰. Using our co-infection data, we show that there is a negative correlation between sex ratio and the proportion of gametocytes contributed by focal genotypes during the pre-peak period ($\chi^2_1 = 16.08$, $P < 0.0001$; slope = -0.52 ± 0.12 ; see Supplementary Table 4b), but not during the post-peak period ($\chi^2_1 = 2.68$, $P = 0.102$; Supplementary Table 5b). Using proportional representation within the mating group as a proxy of inbreeding rate, LMC theory predicts a sex ratio of $z^* = (1 - f)/2$, and hence a slope of $dz^*/df = -0.5$; thus, the pre-peak data are quantitatively consistent with LMC theory. These data support the possibility that genotypes

may be able to infer their own relative frequency in infections. It is not clear why genotype AS did not produce less female-biased sex ratios in response to the presence of a co-infecting genotype. However, this genotype is substantially less virulent than genotypes AJ and ER. The least virulent genotype in our panel is DK⁴¹, and this genotype produced the least female-biased sex ratios in single infections. Virulence is a life-history trait unique to infectious organisms, and could influence reproductive strategies in malaria parasites. Understanding how malaria parasites evaluate the genetic diversity of their infections may also explain why genotype AS did not behave as predicted by LMC.

Discussion

We have tested and confirmed the assumptions and predictions of sex allocation theory in malaria parasites. The female bias typical of single-genotype infections declines towards an equal sex ratio in multiple infections. We also show that: (1) sex ratio is important and transmission studies must consider this trait alongside gametocyte density; (2) mating group productivity decreases at extremely female-biased sex ratios because male gametes become limiting; (3) within-infection sex ratio patterns are explained by variation in anaemia and parasite densities (patterns for which there is significant genetic variation); and (4) female-biased sex allocation in response to LMC decreases as infections progress and parasite densities decline due to competition, anaemia and the host's immune response. These data support 'fertility insurance theory', which predicts that malaria parasites should adjust their reproductive strategies to maximize transmission opportunities throughout their infections^{11,26,32,33}. Furthermore, our data do not support the hypothesis that malaria

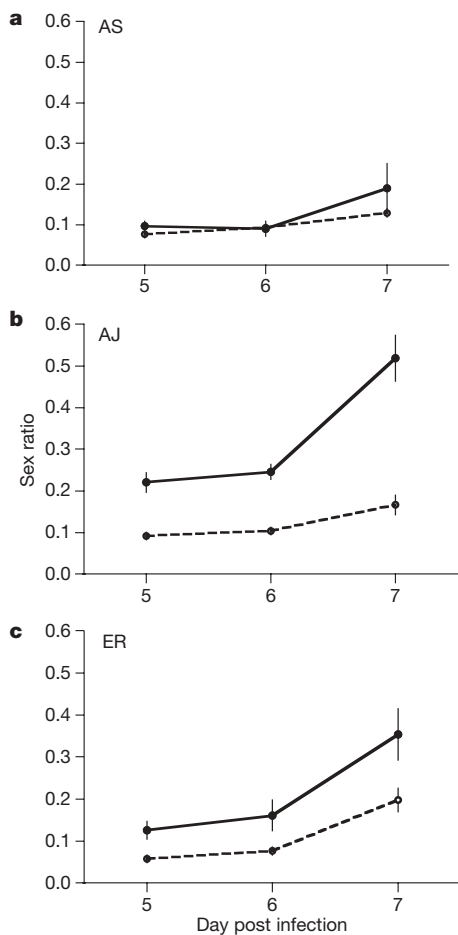


Figure 5 | Sex ratios of focal genotypes during the growth phase of infections. Mean sex ratios throughout the growth phase of infections for *P. chabaudi* focal genotypes when alone (dashed line) and co-infecting with a second genotype (solid line). **a**, AS; **b**, AJ; **c**, ER. Sex ratios of genotypes AS and ER could be distinguished from sex ratios of AJ but AS and ER could not be distinguished from each other (see Supplementary Information). Sex ratios produced by AJ when co-infecting with AS or ER were not significantly different so these infections are grouped. We followed 5 independent infections for each genotype combination. Error bars, \pm s.e.m.

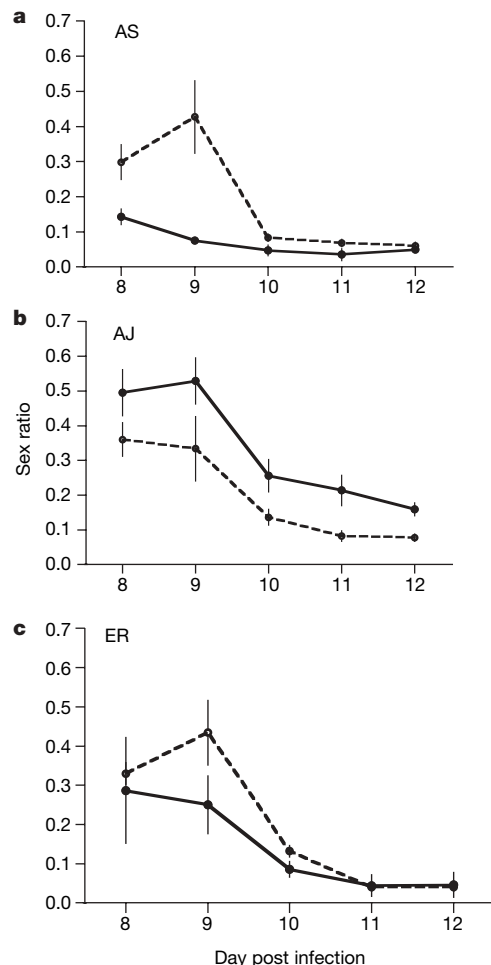


Figure 6 | Sex ratios of focal genotypes during the post-peak phase of infections. As Fig. 5 but for the post-peak phase.

parasites preferentially mate with clone-mates, regardless of the genetic make-up of their mating groups⁴². We have shown that an evolutionary approach can be successfully applied to sex allocation in malaria parasites, suggesting that researchers in pure and applied fields can be optimistic about understanding more complex parasite life-history traits, such as virulence, that are of such economic, medical and veterinary importance⁴³.

As infections consist of billions of parasites, the ability to distinguish clone-mates from unrelated conspecifics requires kin discrimination^{44,45}. This discovery raises important questions. Does within-infection relatedness influence competitive interactions between genotypes and shape patterns of virulence^{46–48}? Low relatedness between co-infecting malaria parasites is expected to favour the most virulent competitors, but related parasites could also cooperate to facilitate growth or immune evasion and result in more virulent infections. Apoptosis has recently been described in protozoan parasites⁴⁹, and kin discrimination supports the possibility that this is actually an altruistic behaviour. Whether malaria parasites are using indirect (environmental) or direct (genetic) cues to discriminate kin must now be investigated. Given that *P. chabaudi* parasites are able to detect the presence of conspecifics in a non-natural host environment (see Supplementary Information), they may be using direct cues rather than changes to their environment. Genetic kin discrimination is rare, but can evolve in situations where marker diversity will be maintained by extrinsic processes^{45,50}. Host–parasite interactions may provide a strong enough source of balancing selection; does pressure to evade immune recognition also enable malaria parasites to employ their sophisticated social behaviour?

METHODS SUMMARY

We used recently developed genetic modification and molecular techniques to carry out these experiments. The fitness experiment used *P. berghei* lines from the same genetic background in which genes essential to either male or female function have been knocked out. Methods for *in vitro* culture of mating groups are based on previously published protocols³⁸. We measured reproductive success by counting ookinetes (zygotes at 18 h of age). Morphological identification is straightforward, as unfertilized female gametes and gametocytes are spherical and ookinetes are elongated banana-shaped cells. We used *P. chabaudi* for our sex ratio experiments for two reasons. First, natural infections vary in genetic diversity and second, a bank of distinct and characterized clonal genotypes are available for this species. We used the genotypes AS, AJ, ER, CR, CW and DK (WHO Registry of Standard Malaria Parasites, The University of Edinburgh) and our experimental design was based on previously published competition treatments. We used previously developed quantitative reverse-transcription PCR assays for a gametocyte-specific and a male-specific gene to calculate sex ratios (proportion of gametocytes that are male). We used 6–8 week old male MF1 mice (in-house supplier, University of Edinburgh). All mice received an intraperitoneal (i.p.) inoculation of parasitized red blood cells in a 0.1 µl dose as previously described¹¹. Mice were housed at 21 °C with a 12 h light cycle, and maintained on a diet of SDS41B food pellets (Harlan Scientific) and 0.05% PABA supplemented drinking water to enhance parasite growth.

Full Methods and any associated references are available in the online version of the paper at www.nature.com/nature.

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Author Contributions S.E.R. conceived and designed the experiments, carried out the fitness consequences experiment, analysed sex ratio data and prepared the manuscript. D.R.D. developed the PCR assays, carried out the sex ratio experiments and data collection. A.G. analysed the fitness data and contributed to discussions and manuscript preparation.

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METHODS

The fitness consequences of sex ratio variation. To produce *P. berghei* parasites for experimental cultures, we i.p. inoculated doses of 1×10^7 parasites into mice that had been pre-treated with phenylhydrazine (60 mg kg^{-1}), to increase gametocyte production, 3 days before receiving their parasites. Parasites were harvested on day 3 PI to set up experimental cultures and also initiate the next set of infections. Each *pb48/45*-ko infection was paired with a *pb47*-ko infection and within each pair, parasites from the two lines were mixed in the appropriate ratios to produce each of our eight sex allocation treatments. Each pair of infections contributed parasites to one replicate of each of the following sex ratios (% male gametocytes): 0, 15, 30, 45, 60, 70, 90 and 100. We paired infections based on their similarity in gametocyte densities and sex ratio to keep haematocrit at 2% for all cultures. We initiated 24 pairs of infections, over 6 blocks, and chose the 19 closest matched pairs to set up our experimental cultures. By cross-factoring each pair of infections with all sex ratio treatments we avoided confounding sex ratio with infection specific factors that could influence fertilization success (for example, gametocyte age, asexual parasite density, anaemia and immune factors in serum) and could control for pair identity ($F_{18,90} = 15.39$; $P < 0.0001$). To calculate the volume of blood required from every *pb47*-ko infection (to contribute males), we calculated the density (per ml of mouse blood) of exflagellating males using a haemocytometer. To calculate the volume of blood required from every *pb48/45*-ko infection (to contribute females) we used blood smear and red blood cell density data (proportion of red blood cells containing female gametocytes \times red cells per ml). Cultures contained an average of $(0.5 \pm 0.04) \times 10^6$ gametocytes ml^{-1} and this variation was controlled for in our analysis ($F_{1,90} = 60.93$; $P < 0.0001$). We cultured 100 μl blood in RPMI with 10% calf serum, pH 8, for 20 h at 20 °C. To count ookinetes, we vortex mixed each culture before counting the number observed in a haemocytometer.

Sequencing qRT-PCR target genes in *P. chabaudi* parasite lines. We have previously developed qRT-PCR to quantify total parasites, total gametocytes and male gametocytes, based on the detection of the *P. chabaudi* common gametocyte gene 1 (*CG1*) and male gametocyte gene 1 (*MGI*). To determine what combinations of genotypes in genetically diverse infections would enable us to follow a focal genotype we sequenced *CG1* and *MGI* genes in CR, ER, DK and CW *P. chabaudi* parasite lines. A 659 bp region of the *CG1* gene and a 924 bp region of the *MGI* gene were then amplified from DNA extracted from each clone and sequenced. Our AJ specific primer sets could discriminate and quantify AJ parasites, gametocytes and male gametocytes in infections with AS and ER. Our AS specific primer sets could discriminate and quantify either AS or ER parasites, gametocytes and male gametocytes in infections with AJ but could not distinguish AS from ER parasites because AS and ER share the same *CG1* and

MGI alleles. The other 3 genotypes each had an AS/ER allele and an AJ allele so could not be distinguished from any others, in any combination.

Experimental design. Male MF1 mice (in-house supplier, University of Edinburgh) were infected with the clonal *P. chabaudi* genotypes AS, AJ, ER, CR, CW or DK (WHO Registry of Standard Malaria Parasites, The University of Edinburgh). We initiated infections with each of our six *P. chabaudi* genotypes on their own, in combinations of two genotypes, three genotypes and all together. We infected five mice for each of 11 treatment groups as follows: (1) six groups of single-genotype infections, consisting of 1×10^6 AJ, AS, ER, CR, CW or DK parasites; (2) two groups of two-genotype infections, one group with 1×10^6 AJ + 1×10^6 AS parasites and a second group with 1×10^6 AJ + 1×10^6 ER parasites; (3) one group of three-genotype infections with 1×10^6 AJ + 1×10^6 AS + 1×10^6 ER parasites; and (4) two identical groups of six-genotype infections with 1×10^5 AJ + 1×10^5 AS + 1×10^5 ER + 1×10^5 CR + 1×10^5 CW + 1×10^5 DK parasites. All infections were sampled in the morning when the circulating parasites were in ring or early trophozoite stages from days 5 to 16 PI. Red blood cell densities were estimated using flow cytometry and reticulocyte densities were estimated from thin blood smears as previously described. **Analyses.** We used R version 2.5.0 (The R Foundation for Statistical Computing; <http://www.R-project.org>) for all analyses. In all analyses sex ratios were arcsine square root transformed. Owing to the number of host deaths in mixed-genotype infections we maximized data resolution by restricting analyses of these data to days 5–12 PI. Statistics are presented from linear mixed-effects models of sex ratio variation throughout infections to overcome problems associated with pseudoreplication in longitudinal analysis. We evaluated the significance of fixed effects by comparing models using log-likelihood ratio tests following stepwise deletion of the least significant term. Specifically, we compared the change in model deviance, following term deletion, to χ^2 distributions with degrees of freedom corresponding to the difference in number of terms in the models. We simplified maximal models using maximum likelihood techniques, until only significant terms remained in the model ($\alpha < 0.05$). We then re-ran minimal models using restricted maximum likelihood to estimate the effect sizes reported in the text. We do not present results for the main effects of terms contained in significant interactions.

Models included the identity of genotypes, the number of genotypes in infections and day PI as factors (as sex ratio variation over time is nonlinear), as well as relevant interactions between these terms. Where possible we grouped genotypes together when there was no significant difference in their sex ratio patterns across infections. Covariates known or suspected to influence sex ratio were fitted, including: (1) measures of virulence and anaemia (mass, red blood cells and reticulocytes); (2) relative and absolute contributions of focal genotypes' gametocytes and asexuals to their infections; and (3) absolute number of gametocytes and asexuals in infections.