Immune anticipation of mating in *Drosophila: Turandot M* promotes immunity against sexually transmitted fungal infections

Weihao Zhong, Colin D. McClure, Cara R. Evans, David T. Mlynski, Elina Immonen, Michael G. Ritchie and Nicholas K. Priest

*Proc. R. Soc. B* 2013 **280**, 20132018, published 30 October 2013

**Supplementary data**

"Data Supplement"


**References**

This article cites 60 articles, 18 of which can be accessed free

http://rspb.royalsocietypublishing.org/content/280/1773/20132018.full.html#ref-list-1

This article is free to access

**Subject collections**

Articles on similar topics can be found in the following collections

- ecology (1573 articles)
- genetics (110 articles)
- health and disease and epidemiology (233 articles)

**Email alerting service**

Receive free email alerts when new articles cite this article - sign up in the box at the top right-hand corner of the article or click here
Immune anticipation of mating in *Drosophila*: Turandot M promotes immunity against sexually transmitted fungal infections

Weihao Zhong¹, Colin D. McClure¹, Cara R. Evans¹, David T. Mlynski¹, Elina Immonen², Michael G. Ritchie³ and Nicholas K. Priest¹

1Department of Biology and Biochemistry, University of Bath, Bath BA2 7SW, UK
2Department of Ecology and Genetics, Animal Ecology, Evolutionary Biology Centre, Uppsala University, Norbyvägen 18 D, Uppsala 75236, Sweden
3School of Biology, Biomedical Sciences Research Complex, University of St Andrews, St Andrews, Fife KY16 9ST, UK

Although it is well known that mating increases the risk of infection, we do not know how females mitigate the fitness costs of sexually transmitted infections (STIs). It has recently been shown that female fruitflies, *Drosophila melanogaster*, specifically upregulate two members of the *Turandot* family of immune and stress response genes, *Turandot M* and *Turandot C* (*TotM* and *TotC*), when they hear male courtship song. Here, we use the Gal4/UAS RNAi gene knockdown system to test whether the expression of these genes provides fitness benefits for females infected with the entomopathogenic fungus, *Metarhizium robertsii* under sexual transmission. As a control, we also examined the immunity conferred by *Dorsal-related immunity factor* (*Dif*), a central component of the *Toll* signalling pathway thought to provide immunity against fungal infections. We show that *TotM*, but not *TotC* or *Dif*, provides survival benefits to females following STIs, but not after direct topical infections. We also show that though the expression of *TotM* provides fecundity benefits for healthy females, it comes at a cost to their survival, which helps to explain why *TotM* is not constitutively expressed. Together, these results show that the anticipatory expression of *TotM* promotes specific immunity against fungal STIs and suggest that immune anticipation is more common than currently appreciated.

1. Introduction

Mating is fraught with danger. In addition to the fitness costs associated with finding sexual partners, copulation and offspring production, mating increases the risk of acquiring sexually transmitted infections (STIs) [1–3]. In insects, STIs are often both highly prevalent and pathogenic [3,4]. It is generally thought that they exert a selective pressure strong enough to influence the evolution of mating systems, life histories, sexual conflict and sexual behaviour [3,5]. Yet, we have a poor understanding of how they have shaped the immune system [6].

Females could mitigate the risks of acquiring STIs through immune anticipation of mating, the activation of immune responses before sexual congress and potential exposure to pathogens [7]. Pre-emptive immune activation is predicted to be more advantageous than a purely reactive response because it shortens the time delay of the immune response, and thereby maximizes its efficiency ([7]; MT Siva-Jothy, E Harney, W Zhong 2013, unpublished data). We know that females upregulate a number of immunity-related genes in response to mating [8–12]. But even the act of courtship might stimulate immune activation. If immune genes expressed during courtship represent immune anticipation of
mating, then we would expect such responses to enhance immunity against STIs and to exhibit costs in some aspects of life history, because otherwise they would be constitutively expressed [13,14].

One way to address this possibility is to identify candidate immune genes associated with courtship and perform infection and fitness assays in which the expression levels of the genes are manipulated. Recently, Turandot C and Turandot M (TotC and TotM), members of the Turandot family of immune and stress response genes, were shown to be upregulated in the heads of female Drosophila melanogaster stimulated by male courtship songs independent of any physical encounter with males [15]. Of the two, TotM is probably the better candidate for anticipatory immunity against STIs, as it is poorly induced by non-immune-related stress [16] but strongly induced by both fungal infections [16–18] and mating [9,19,20]. In addition, induction of TotM by natural fungal infection exhibits similar fold-change in expression to well-known antifungal antimicrobial peptides (AMPs), including Drosomycin and Metchnikowin [18]. Surprisingly, there is little evidence that courtship stimulates the upregulation of the canonical Toll and Imd pathway immune genes, such as Dorsal-related immunity factor (Dif), an NF-kB-like factor that regulates Toll-dependent immune responses thought to provide immunity specifically against Gram-positive bacteria and fungi [15,21,22].

Previous efforts in establishing D. melanogaster as a model laboratory system for studying insect STIs have focused on bacterial pathogens [23,24]. However, entomopathogenic fungi might be more appropriate. First, entomopathogenic fungi are widespread across diverse environments causing a large proportion of all known insect STIs, and indeed the majority of all insect diseases [3,25]. Second, because fungal spores cause infection through direct contact with the cuticle [26,27], they are amenable for comparisons between sexual and non-sexual horizontal transmission. Finally, studying the sexual transmission potential of entomopathogenic fungi in the laboratory have important implications for their application in the field as agents of biocontrol [28–30].

Here, we examine the hypothesis that TotM provides protection against sexually transmitted Metarhizium robertsi, a generalist soil-borne entomopathogenic fungus, which exhibits both sexual and non-sexual transmission in dipterans and has been used extensively in biocontrol [25,31,32]. Specifically, we test the predictions that (i) Metarhizium can be sexually transmitted in D. melanogaster; that (ii) expression of TotM helps to mitigate the cost of infections under sexual transmission, but not direct modes of transmission; and that (iii) the expression of TotM has fitness costs in the absence of sexually transmitted Metarhizium. To address these questions, we use the Ga4/UAS RNAi-targeted gene knockdown approach [33], in conjunction with large-scale demographic analysis, to estimate the immunity and fitness conferred by TotM, TotC and Dif under both STIs and high-dose direct topical infections (DTIs) of M. robertsi.

2. Material and methods

(a) Fly strains and fungal culture maintenance

A wild-type Dahomey strain of D. melanogaster (provided by Dr Stuart Wigby, University of Oxford) was kept in large population cages (1 m³) with overlapping generations for 2 years prior to the start of the experiments. RNAi strains were obtained from Vienna Drosophila RNAi Center (UAS-TotM-IR, transformant ID 106727; UAS-TotC-IR, transformant ID 106379; UAS-Dif-IR, transformant ID 30579). We used the non-tissue-specific Act5C promoter to drive ubiquitous expression of Ga4 and UAS constructs (Act5C-Ga4/Cyo, Bloomington Stock Center stock number 44114). We crossed Act5C-Ga4/Cyo females with males carrying one of the UAS constructs to generate the active knockdown genotypes (Act5C-Ga4/UAS-TotM-IR; Act5C-Ga4/UAS-TotC-IR; Act5C-Ga4/UAS-Dif-IR). As a control for the presence of the UAS transgene, we crossed w1118 wild-type females (the genetic background for all RNAi lines, obtained from Bloomington Stock Center) with males carrying one of the UAS constructs (UAS-TotM-IR/+; UAS-TotC-IR/+; UAS-Dif-IR/+). As a control for the presence of the Ga4 driver, we crossed Act5C-Ga4/Cyo females with w1118 males (Act5C-Ga4/+). The effectiveness of RNAi knockdowns of TotM and TotC was confirmed by semi-quantitative PCR [34]. All experimental animals were maintained at 25 °C with 12 L:12 D cycle in standard Drosophila vials at low densities (approx. 50 flies/vial) for at least two generations prior to the start of experiments. We used an oatmeal–molasses–agar media with added live baker’s yeast and an antifungal agent (Nipagin), which inhibited the growth of naturally occurring saprophytic fungi. All experimental flies used were collected as virgins over a period of 24 h.

Metarhizium robertsi (isolate 2575, previously known as Metarhizium anisopliae strain ME1) was obtained from the Agricultural Research Service Collection of Entomopathogenic Fungal Cultures (ARSEF, United States Department of Agriculture). We inoculated quarter-strength sabouraud dextrose agar (SDA) with Metarhizium anisopliae (isolate 2575, previously known as Metarhizium anisopliae strain ME1) was obtained from the Agricultural Research Service Collection of Entomopathogenic Fungal Cultures (ARSEF, United States Department of Agriculture). We inoculated quarter-strength sabouraud dextrose agar (SDA) with M. robertsi conidia (asexual fungal spores) and incubated the plates at 28 °C for four weeks before storing at 4 °C for up to three months. Conidia were collected by scraping the surface of the sporulating culture with an inoculating loop.

(b) Sexual transmission of fungal pathogen

We assessed the transmission potential of M. robertsi by exposing healthy Dahomey females to males that had been topically inoculated with the fungus. At adult age day 4, groups of 10 virgin males were topically inoculated with 6 mg of conidia without CO2 anaesthesia by shaking in a 250 ml conical flask for 20 s. Inoculated flies were held in temporary holding vials for 24 h, ensuring that they had opportunities to groom themselves, which has previously been shown to be effective at removing fine dust particles [35]. At adult age day 5, each infected male fly was introduced into a new vial containing 10 uninjected virgin females of the same age and removed after 24 h. The logic of giving males time to groom and subsequently using a fresh vial was to allow male to adopt a more natural behaviour [32] and to minimize the probability of females contracting infection from conidia that had been dislodged during grooming. We then transferred and held treated females in individual vials for a further 24 h to allow egg-laying. The presence of larvae 4 days after oviposition indicated that the female had mated with an infected male. We assessed the infection status of females by the presence of Metarhizium-like fungal growth on cadavers. Flies were briefly immersed in 70% ethanol before being gently crushed and placed in Petri dishes on moistened filter paper at the end of the egg-laying period. After an
incubation period of 5 days at 28 °C, we examined all cadavers for signs of Metarhizium-like fungal growth (either hyphae or conidia) with a low-power dissection microscope. Because high levels of horizontal transmission of conidia between infected and naive flies owing to non-sexual contact could confound our interpretation, we also assessed the potential for non-sexual horizontal transmission of M. robertsi using the same procedures described above by exposing naive males and females to infected flies of the same sex.

(c) Survival assays under direct topical infection and sexually transmitted infection

We assessed the effects of gene knockdowns on survival under high-dose DTIs and sexual transmission (STI) using adult flies for all genotypes. For DTI, at adult age day 7, we infected groups of approximately 300 mixed-sex flies of each genotype with 20 mg of conidia, or kept as uninfected control, following the protocol described previously. Inoculated flies were held in temporary holding vials for 30 min before being transferred to demography cages (10 × 15 cm). For STI, we first inoculated 6-day-old w1118 males in groups of 20 with 12 mg of conidia, and then transferred 20 infected or control males with 20 uninfected females to demography cages at adult age day 7. As infected males in STI treatment suffered much greater mortalities than control males, we restored the original complement of 20 infected males by adding freshly infected w1118 males at day 12 and 24 postinoculation. For both DTI and STI, we removed and recorded dead flies daily until day 9 postinoculation and every 2 days thereafter. We also tracked the changes in pathogen loads in the first 24 h following DTI by sampling inoculated Dahomey wild-type flies at three time points postinoculation (0, 2.5 and 24 h; n = 9). Sampled flies were individually homogenized in 200 μl of 0.04% Tween80, diluted by a factor of 103 and spread onto standard SDA plates. Pathogen loads were assessed by counting the numbers of colony forming units (CFUs) following incubation at 28 °C for 24 h.

(d) Fecundity assay under sexually transmitted infection

We assessed the effects of gene knockdowns on survival and fecundity of females exposed to fungus-infected males using flies from the same cohort collected for survival assays. In the fecundity assay, we first infected 2-day-old w1118 wild-type adult males (the genetic background of our RNAi strains). At 24 h-postinoculation, infected or uninfected control males were transferred to individual vials containing a single uninfected virgin female for each genotype. The mating pairs were assigned positions in randomized blocks and transferred to new vials after 24 h, and thereafter every 2 days until day 9 (n = 55/treatment/genotype). Used food vials were frozen 18 days after collection and the numbers of eclosed pupae were counted giving a combined measure of fecundity and larval viability. We assessed the proportion of females that became infected through mating with infected males by sampling all surviving females at the end of day 9 postinoculation (96.8%, 701/724) and checking for signs of Metarhizium-like fungal growth after incubation at 28 °C for up to two months.

(e) Statistical analysis

All statistical analyses were performed with R version 2.15 [36]. We assessed the contribution of mating to the transmission of STIs by comparing the proportions of flies that displayed Metarhizium-like fungal growth for mated females, and those that were kept with infected males but remained virgin using χ2-tests with continuity correction. We used student’s t-test on CFUs to directly compare pathogen loads immediately after inoculation and after 24 h.

Cox proportional hazard regressions were used to analyse all survival data. The full model (including all genotypes) contained age at death and censoring information as the response variables—genotype, infection treatment and their interaction—were included as predictor variables. A separate Cox regression was performed for each gene of interest that only included the relevant knockdown and control genotypes (e.g. for TotM, the data included these genotypes: Ac5C-Gal4/UAS-TotM-IR, Act5C-Gal4/+ and +/UAS-TotM-IR). For each gene of interest, we first extracted the hazard ratios (the fold-increase in risk of death in infected animals relative to uninfected controls) for the knockdown genotype and its combined control genotype (by pooling raw survival data of the relevant control genotypes) from the Cox models. Because the mortality rate in the DTI treatment is substantially higher than that in the STI treatment, it is difficult to directly compare the effect of immune gene knockdowns in the two treatments. To overcome this problem, we calculated normalized hazard ratios by dividing the hazard ratios of each knockdown by its associated combined control genotype. Unlike simple metrics of lifespan, this measure describes the effect of each gene knockdown on immunity after accounting for its genetic background, which allows us to directly compare the immune properties conferred by genes under STIs and DTIs, despite great differences in effect size. We assessed the survival cost of gene expression in the absence of infections by comparing the hazard ratios of each gene knockdown relative to its combined control genotype under uninfected control conditions.

We used mixed effects models to assess the effects of genotype and infection on fecundity across time. The full model included the number of eclosed pupae produced at each time point as the response variable; genotype, treatment, time and all associated two-way interactions as fixed effects (three-way interaction was non-significant when fitted, and thus dropped from the full model), and individual females as random effect (intercepts). We also included the age at death of male partners as a covariate in the full model to account for the possibility that females might have lower fecundity under STI simply owing to a lack of remating opportunities as infected males die at earlier ages than uninfected controls. Female fecundity in the first 24 h was excluded from the model as the fecundity was much lower than that at other time points and previous experiments suggested minimal in vivo fungal growth in this period (VL Hunt, W Zhong, CD McClure, DT Mlynski, EML Duxbury, AK Charnley, NK Priest 2013, unpublished data). We assessed the fecundity cost of gene expression in the absence of infections by comparing the mean total pupae productions of the gene knockdown (day 0–9 posttreatment) and the combined control genotype using one-way analysis of variance.

3. Results

(a) Sexual transmission of fungal pathogen

We found that M. robertsi can be sexually transmitted in the fruitfly, with approximately one in five (55/263) naive females displaying Metarhizium-like fungal growth on their cadavers.
EMERGING FROM INFECTED FLY CADAVER AND (HIZIUM MELANOGLASTER
We found that (b) Effects of sexually transmitted infection and direct
after being placed with a topically infected male for 24 h
(figure 1a,b). Further analysis showed that fungal transmission
was driven primarily by mating, as the proportion of cadavers
with fungal growth was higher in gravid females than that
infertile females ($\chi^2 = 8.96, p = 0.0028; $figure 1c). The dose
received by females was likely to be low as the pathogen
load of the topically infected males was only approximately
5000 CFU, which had declined by grooming from the initial
load of approximately 20 000 CFUs ($t = 7.69, p = 0.006; $electronic supplementary material, $figure S1). Finally, we also
found that Metarhizium could be transmitted among same-
sex flies ($7/277$ for male-to-male transmission and $7/266$ for
female-to-female transmission; $electronic supplementary material, $figure S2). Nevertheless, naive flies were much more
likely to be infected through sexual transmission than through
non-sexual transmission, 20.9 versus 2.6%, respectively.

(b) Effects of sexually transmitted infection and direct
topical infection on survival across RNAI strains
We found that TotM promotes immunity against Metarhizium
when it is sexually transmitted (STI), but not when it is applied
as a DTI. The effect of STIs on the hazard ratio, which esti-
mates the risk of death in infected treatments relative to
control treatments, was highly dependent on the host geno-
type (overall: genotype $\times$ treatment, $\chi^2 = 26.4, p = 0.0002; $figure 2a). Specifically, TotM knockdown flies (Act5C-Gal4/
UAS-TotM-IR) were susceptible to STIs, but there was
no evidence of susceptibility in either of $+/Act5C-Gal4$ or
$+/UAS-TotM-IR$ control genotypes (genotype $\times$ treatment; $\chi^2 = 15.8, p = 0.00037; $figure 2a). By contrast, there was no
difference in susceptibility to STIs among Dif knockdown
flies (Act5C-Gal4/UAS-Dif-IR) and its associated control geno-
types $+/Act5C-Gal4$ and $+/UAS-Dif-IR$ (genotype $\times$
treatment, $\chi^2 = 0.05, p = 0.98$). Surprisingly, TotC knockdown
flies (Act5C-Gal4/UAS-TotC-IR) had slightly higher survival
postexposure than both of control $+/Act5C-Gal4$ and
$+/UAS-TotC-IR$ genotype flies (genotype $\times$ treatment,
$\chi^2 = 9.1, p = 0.011; $figure 2a).

We found different patterns under DTI. While DTIs gener-
ally caused very rapid mortalities such that 95% of flies died
within 9 days, some genotypes were much more susceptible
(overall: genotype $\times$ treatment, $\chi^2 = 751, p < 0.0001; $figure 2b). As expected [22], Dif knockdown (Act5C-Gal4/UAS-Dif-
IR) females were significantly more susceptible to DTIs than
either of its control genotypes (genotype $\times$ treatment,
$\chi^2 = 545, p < 0.0001; $figure 2b). However, neither TotM nor
TotC knockdown was more susceptible to DTIs than their
respective control genotypes (figure 2b). Interestingly,
although the hazard ratio of the Dif knockdown line under
DTI was more than 16 times higher than that of TotM knock-
down under STI (46.2 $\pm$ 6.2 versus 2.8 $\pm$ 0.6), their hazard
ratios were comparable after they were normalized to account
for the susceptibility of their control genotypes (2.4 $\pm$ 0.4
versus 2.7 $\pm$ 0.7; $figure 2c).

(c) Effect of sexually transmitted infection on fecundity
across RNAI strains
Sexually transmitted Metarhizium infections resulted in repro-
ductive costs for female flies. Exposure to topically infected
male partners initially had little impact on female reproduc-
tion, but over time, female fecundity in the infected treatment
declined relative to uninfected controls (treatment $\times$ time,
$F_{1,2030} = 30.5, p < 0.0001; $electronic supplementary material,
$figure S3). This pattern was consistent in all lines as there
was no evidence that TotM or indeed any gene knockdown
strain suffered greater fecundity reduction than their control
genotypes (treatment $\times$ genotype, $F_{6,705} = 1.49, p = 0.19$).
The reduction in female fecundity under STIs could not be
explained by a lack of remating opportunities owing to
increased mortalities of infected male partners, because male
longevity did not significantly contribute to female fecundity
over the course of the experiment ($F_{1,705} = 3.5, p = 0.062$).
In addition, while the cadavers of females that had been exposed
to infected males were more likely to exhibit Metarhizium-
lke fungal growth than those exposed to control males
($\chi^2 = 5.69, p = 0.017$), there was no evidence that the RNAi
knockdown genotypes influenced the probability of fungal
growth ($\chi^2 = 0.001, p = 0.97; $electronic supplementary material, $figure S4).

(d) Effect of immune gene expression on survival and
fecundity in uninfected flies
We found that the expression of TotM and Dif, but not TotC,
results in survival costs for uninfected females. Both TotM
and Dif knockdown flies (Act5C-Gal4/UAS-TotM-IR and
Act5C-Gal4/UAS-Dif-IR), but not TotC knockdown flies
(Act5C-Gal4/UAS-TotC-IR), showed enhanced survival rela-
tive to their control genotypes (TotM: $\chi^2 = 8.58, p = 0.0034;$
By contrast, we found evidence for reproductive benefits of TotM and TotC expression, but reproductive costs of Dif expression. Both TotM and TotC knockdown females had lower total reproduction than their respective controls, whereas Dif knockdown females were more fecund than its control.

Figure 2. TotM is required for enhanced survival under STI, but not under DTI. (a) Cox proportional hazard ratios of STI relative to uninfected controls, (b) Cox proportional hazard ratios of DTI relative to uninfected controls and (c) the susceptibility of TotM and Dif under both STI and DTI after normalization for differences in the influence of mode of infection on hazard of the control genotypes. Dotted lines indicate hazard ratio of 1, which indicate both infected and uninfected controls had the same risk of death. Asterisks (*) indicate the level of statistical significance of hazard ratios (*p < 0.05; **p < 0.01; ***p < 0.001).

Dif: χ² = 26.6, p < 0.0001; TotC: χ² = 0.88, p = 0.35; figure 3a).
genotypes (TotM: $F_{1,135} = 44.8, p < 0.0001$; TotC: $F_{1,127} = 7.6, p = 0.0068$; Dif: $F_{1,128} = 6.3, p = 0.014$; figure 3b).

4. Discussion

Mechanisms of insect immunity are known to be pathogen-specific [37,38]. However, the extent to which insects use ecological cues to inform which responses to mount is not known. Our study shows that a gene that is upregulated in anticipation of mating provides protection against sexually transmitted *Metarhizium* infections. This finding is important because it illuminates the molecular mechanisms as well as the life-history costs and benefits which underpin immunity against STIs. In combination with previous results [15], our results imply that fruitflies demonstrate immune anticipation of mating and that immune anticipation could be a general mechanism for achieving immune specificity.

(a) A Turandot gene that enhances immunity against sexually transmitted infections

Hundreds of *Drosophila* genes, including TotM, have been identified on the basis of elevated expression following immune challenges, but the functional consequences of these genes are rarely established [16–18]. This is a problem because gene expression does not necessarily translate into immunity against live pathogens [39–41]. We show that TotM confers protection against fungal STIs and its effects are similar in magnitude to that conferred by Dif to fungal DTIs.

The mechanisms through which TotM enhances immunity are currently unknown. All protein products encoded by the *Turandot* gene family are thought to be actively produced in the *Drosophila* fat bodies and secreted into the haemolymph, where they are hypothesized to act as protein chaperones or as signalling molecules [16,42]. Though direct tests are needed, it seems unlikely that TotM possesses direct antimicrobial activities similar to known antifungal AMPs, such as *Drosomycin* and *Metchkinokwin*. Instead, overexpression of another *Turandot* gene, TotA, does not provide increased protection against Gram-negative bacterial infections [42,43]. Instead, TotM might help the fly to tolerate persistent fungal infections by mitigating the negative effects of the infection without actively suppressing pathogen growth [44–46]. Consistent with a role in enhancing tolerance, not resistance, we found that fungi were as likely to emerge from the control genotype flies as they were from TotM knockdown flies.

(b) Mode of transmission and immunity

Fruitflies have a remarkable ability to mount immune responses which are specific to the pathogens they encounter [37,38]. Our work shows that the efficacies of their immune responses are also specific to the mode of infection transmission. STIs differ from other modes of transmission in that they tend to cause chronic low-level infections, which do not result in rapid septicaemia and increased host mortality—consequences typically associated with acute immune challenges [2]. The lower initial inoculums in our STI treatment is evidenced by the proportion of flies that exhibit fungal growth on female cadavers (5–25% for STIs and 80–95% for DTIs; VL Hunt, W Zhong, CD McClure, DT Mlynski, EML Duxbury, AK Charnley, NK Priest 2013); and the increased grooming activities we observed in the DTI treatment, which efficiently reduced pathogen load (this study; [35]). Consistent with the differences in pathogen dose between the two infection treatments, we found that sexually transmitted *Metarhizium* infections cause weak, though significant, fitness costs for females and that the expression of TotM, but not Dif, ameliorates the survival costs associated with STIs. By contrast, we found that direct topical *Metarhizium* infections cause substantial fitness costs for females and that the expression of Dif, but not TotM, helps ameliorate those survival costs. Taken together, these findings show that fruitflies have a specific mechanism for immunity against low-dose STIs and against high-dose DTIs, even for the same pathogen.

It is important to acknowledge that though we have established a role for TotM in immunity against low-dose STIs, we do not know whether TotM confers immunity against STIs per se or to low-dose infections more generally. We cannot dismiss
the possibility that high fungal doses overwhelmed the fine-tuned protective effects provided by TotM or that low fungal doses masked the susceptibility of the Dif knockdown. Similarly, the choice of diet could confound our results, as the fecundity benefits of TotM and TotC expression might have resulted from the ad libitum access to dietary yeast in this study [47]. Another potential problem is that genetic constructs, such as the Act5C driver and UAS element, may have pleiotropic effects on the life history of the fly, which could confound direct comparisons with the knockout genotype. However, these problems are unlikely to influence our interpretations. The response to topical fungal infection in our Dif knockdowns was similar to that of the classic Dif knockout mutant [22]. Because our experiments were conducted under the same dietary conditions and because our analysis included normalizations to control genotypes, we can confidently attribute the survival reduction in TotM knockdown to the effect of gene expression, rather than to potential confounding factors such as diet, genetic pleiotropy or the general frailty of immune gene knockdown lines [48]. Regardless of how they confer immunity, our findings provide clear evidence that TotM and Dif are specific for different modes of fungal transmission and that their expressions have different life-history consequences for the host.

It is important to stress that we are not arguing that M. robertsi is predominantly transmitted sexually or claiming that it is transmitted internally during copulation. Given the proclivity of Metarhizium for topical transmission, we would expect there to be some non-sexual transmission, even in our STI treatments. Drosophila tends to aggregate on food sources, which could have increased contacts and fungal transmission in this study [49]. However, non-sexual transmission is unlikely to be substantial enough to change the interpretation of the data. First, males had been given 24 h for grooming and were subsequently placed in fresh vials, which reduced the risk of females indirectly picking up dislodged spores. Second, we found that females who mated with infected males were more likely to be infected than those that did not. And, finally, in independent experiments, infection success was substantially lower in same-sex transmission trials than in trials involving sexual transmission (21 versus 3%). Thus, although we documented that the fungus can be transmitted non-sexually, sexual transmission is primarily responsible for the observed infections in our STI treatments.

(c) The cost of immune expression

Though many studies have documented the costs of immunity [14,47,50], the molecular and physiological basis of such costs are often poorly understood [13]. We found that under uninfected control conditions Dif is generally deleterious in the absence of infections. The expression of Dif entails both significant survival and fecundity costs, which is also supported by a previous study of Dif knockout mutant [48]. The costs of Dif expression are likely to arise from its control of AMP induction through the Toll pathway [22], though Dif might also function in other non-immunity-related processes [21]. These strong fitness costs could help to explain why Dif only appears to be modestly induced by direct topical fungal infections [17] and why it was not up-regulated in females in response to male courtship songs (at least in their heads) [15].

By contrast, our findings for the Turandot genes are only partially consistent with the predicted costs of immune gene expression. We found that TotM has an antagonistic pleiotropic influence on the life history of the fly: though it is costly for survival, expression of TotM also substantially enhances female fecundity. In addition, while there was no evidence that TotC conferred immunity against Metarhizium, it did not contribute to survival cost and even enhanced female fecundity. However, unlike Dif, there is evidence that TotC and TotM play additional roles in reproduction. In particular, TotC and TotM are upregulated in response to exposure to male accessory gland proteins [8–12]. Perhaps TotM could mediate the trade-off between late-age survival and early-age reproduction, a key component of fitness in populations with fluctuating growth rates [51]. Thus, though we cannot easily tease apart the cost of expression from the additional roles played by TotM, the fact that its expression induces survival costs indicates that TotM has a long-term detrimental effect, which is an important facet of the explanation for why it is not constitutively expressed. Interestingly, TotM and TotC appear to evolve more rapidly than Dif [52], suggesting that they have experienced divergent or relaxed selection, perhaps as a consequence of their lower cost of expression [14,47,53].

(d) Mating and immune anticipation in insects

Mating is frequently associated with heightened risk of contracting both ‘pure’ STIs and other opportunistic infections [3,54–56]. Such threats could be countered by upregulating immunity-related genes postmating [8–12]. However, because of the full deployment of immune responses can often take a considerable amount of time [57,58], selection is expected to favour immune anticipation of mating [7]. Though there have been few well-documented cases, immune anticipation is likely to be far more common than currently appreciated. Our study supports the hypothesis that female fruitflies can mitigate the risk of contracting sexually transmitted fungal infections during mating by pre-emptively upregulating TotM [15]. More generally, there are many other biological scenarios associated with elevated disease risk for which we would expect immune anticipation to be advantageous, such as feeding (as has been documented in bed bugs; MT Siva-Jothy, E Harney, W Zhong 2013, unpublished data) and crowding of conspecifics [59–61]. A particularly tantalizing possibility is that the control of many immune genes including TotM [62] by circadian clock genes might reflect ‘anticipation’ of predictable fluctuations of disease risk over the course of 24 h. Thus, the courtship-induced, pre-emptive upregulation of TotM might be representative of a general pattern of immune anticipation in insects, underlining the intimate link between brain, behaviour and immunity [63,64].

Acknowledgements. We thank S. Reynolds, D. Henk and two anonymous reviewers for comments and V. Hunt for the photos of infected fruitflies. For assistance with running the project, we thank Fiona Chapman, Chloe Clark, Sarah Duxbury, Freyja Prentice, Matthew Turner and Owain Williams.

Data accessibility. All supporting data are accessible in the electronic supplementary material.

Funding statement. This work was funded by BBSRC studentships to W.Z., C.D.M. and D.T.M. and BBSRC grant BB/I000836/1 to N.K.P.


