Infection avoidance behaviour: female fruit flies adjust foraging effort in response to internal and external cues of viral infection

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Infection avoidance behaviours are the first line of defence against pathogenic encounters. Behavioural plasticity in response to internal or external cues can therefore generate potentially significant heterogeneity in infection. We tested whether *Drosophila melanogaster* exhibits infection avoidance behaviour during foraging, and whether this behaviour is modified by prior exposure to *Drosophila C Virus* (DCV) and by the risk of DCV encounter. We examined two measures of infection avoidance: (1) the motivation to feed in the presence of an infection risk and (2) the preference to feed on a clean food source over a potentially infectious source. While we found no clear evidence for preference of clean food sources over potentially infectious ones, female flies were less motivated to feed when presented with a risk of encountering DCV, but only if they had been previously exposed to the virus. We discuss the relevance of behavioural plasticity during foraging for host fitness and pathogen spread.
Introduction

Hosts vary considerably in their ability to acquire and transmit infection 1–3, and much of this variation is caused by differences in the contact rate between susceptible individuals and sources of infection 4,5. For example, viruses of Drosophila fruit flies are not only widely distributed, they also show very broad host range 6. Given the high viral prevalence of pathogens in natural environments, mounting a timely and efficient immune response to all possible pathogenic challenges would be physiologically costly and ultimately ineffective. Hosts capable of reducing the probability of contacting parasites, infected conspecifics or infectious environments can therefore not only prevent the deleterious effects of infection, but also circumvent the undesirable energetic costs of immune responses, including immunopathology 4,7. Avoiding infection is therefore the first line of non-immunological defence against infection 8, and is known to occur across a broad range of host taxa, including humans 7,9.

Like most traits, infection avoidance behaviours are likely to vary according to the context of infection, and pathogens are major drivers this context 4,7,9–11. Pathogens may alter host responses in two ways. By altering the immunophysiology of the host during infection, pathogens can alter host behaviour 12,13. Pathogens also modify the host external environment by increasing the likelihood of exposure to novel infections, and these external cues of infection risk are also known to influence host behavioural responses 4,7. Understanding variation in infection avoidance behaviours therefore provides an important functional link between the neurological, behavioural and immunological processes that together govern the spread of disease 12.
Insects are ideal systems to investigate the interplay between infection and behaviour. The fruit fly *Drosophila* is especially amenable to these studies, as it is one of the best developed model systems for host-pathogen interactions and behavioural ecology and genetics. One of the most studied pathogenic interactions in *Drosophila* is the host response to systemic and enteric infection with *Drosophila C Virus* (DCV). DCV is a horizontally transmitted +ssRNA virus that naturally infects the fly gut, causing intestinal obstruction, severe metabolic dysfunction and eventually death. As a consequence of its pathology, female flies infected with DCV are also known to exhibit behavioural modifications, such as reduced locomotion and increased sleep. The *Drosophila*-DCV interaction therefore offers a powerful system to investigate the ecological consequences that may arise from the physiological and behavioural effects of enteric viral infections.

In the present study we used a combination of controlled experimental infections and foraging choice assays, to test whether adult *D. melanogaster* are able to avoid potentially infectious environments when foraging for food, and if avoidance behaviour is modified in response to virus exposure history and to different risks of acquiring DCV infection. We find evidence for avoidance behaviours in the form of reduced motivation to feed according to the risk of infection. However, these effects were only present in female flies previously exposed to DCV, indicating potentially important sexual dimorphism in infection avoidance.

**Materials and methods**
Fly and virus stocks

All flies used were from a long-term laboratory stock of Wolbachia-free Drosophila melanogaster Oregon R line, maintained on Lewis medium in standard conditions: 25°C, with a 12:12h light:dark cycle. Fly stocks were routinely kept on a 14-day cycle with non-overlapping generations under low larval densities. The DCV culture used in this experiment was grown in Schneider Drosophila Line 2 (DL2) as described in 24. Ten-fold serial dilutions of this culture (diluted in Ringers buffer solution) were aliquoted and frozen at -80°C for long-term storage before use.

Virus exposure

Flies used in the foraging choice assays were obtained by preparing 10 vials of Lewis medium and yeast containing ten mated females. Flies were allowed to lay eggs for 48 hours resulting in age-matched progeny reared in similar larval densities. To test the effect of previous exposure to virus on avoidance behaviour during foraging, We exposed these progeny to DCV via the oral route of infection two to three days after eclosion. Oral DCV infection causes small but significant reduction in fly survival19 and in we have found that orally infected flies experience changes affects fly mortality, fecundity, fecal shedding (Vale, unpublished data), activity and sleep24. Briefly, single-sex groups of 20 flies were placed in vials containing agar previously sprayed with DCV (“exposed” to 50 µl of 10⁸ viral copies/ml) or the equivalent volume of Ringers buffer solution as a control (“not exposed”). This procedure produced 10 replicate vials of either healthy or virus-exposed male or female flies. The viral dose used here was lower than previously reported methods19, so we first tested this dose was sufficient to result in viable DCV infections by
measuring changes in virus titres and fly survival in separate experiments (Fig. 1). Fly survival was monitored on 5 replicate groups of 10 OreR female flies per vial for 11 days following oral exposure. To measure changes in DCV titre, twenty-five, 2-3 day-old female flies were individually housed in vials previously sprayed with DCV as described above for 3 days. Five flies were collected 1, 3, 6, 9 or 13 days after exposure and total RNA was extracted from flies homogenised in Tri Reagent (Ambion), reverse-transcribed with M-MLV reverse transcriptase (Promega) and random hexamer primers, and then diluted 1:10 with nuclease free water. qRT-PCR was performed on an Applied Biosystems StepOnePlus system using Fast SYBR Green Master Mix (Applied Biosystems). We measured the relative fold change in DCV RNA relative to rp49, an internal Drosophila control gene, calculated as $2^{-\Delta \Delta Ct}$ as described in 25.

Foraging choice assays

Following 3 days of virus exposure, we set up independent foraging choice assays in cages - cylindrical transparent plastic containers (12 cm in diameter) containing two equally spaced plastic vials of standard Lewis fly medium supplemented with dry yeast. For each combination of "DCV exposed" and "not exposed" male or female flies, we set up two sets of cages to simulate different risks of infection: a "no risk" environment, with two clean vials (sprayed with sterile Ringers solution), and a "high-risk" environment where one of the vials was sprayed with DCV, as described above. Six replicate 20-fly groups were allocated to the "high-risk" chambers and four replicates to the "no risk" chambers, resulting in a total of 40 independent foraging choice cages. Flies were added to the chamber from a neutrally placed hole in the lid, and the number of flies that settled on each vial was recorded every 30
minutes for five hours. Care was taken to randomise the position of the cages so that the orientation of the light did not influence the choice of the flies in any systematic way.

Statistical Analysis

To measure infection avoidance, we took two approaches. First, we hypothesised that the motivation to feed would be lower in environments where the risk of infection is higher. We therefore compared the motivation to feed between the “no risk” and “high-risk” cages, measured as the proportion of flies inside each replicate cage that chose to feed on any of the provided food sources. We also asked whether flies that chose to feed showed any evidence of avoiding potentially infectious food sources. For this analysis we focused on the “high risk” cages and recorded the proportion of flies choosing the clean food source over the infectious food source in each replicate cage. In both analyses of ‘motivation to feed’ and ‘infection avoidance’, data on the proportion of flies choosing each food source within each replicate cage were analysed with a generalised linear model assuming binomial error and logit link function, and included fly ‘sex’, ‘previous exposure’ and ‘infection risk’ as fixed effects. ‘Replicate cage’ was included as a random effect, nested within treatments. We also analysed the average motivation to feed and infection avoidance across all time points, in a model that included “time” as a random effect. Treatment specific contrasts were used to test the significance of pairwise comparisons. Analyses were carried out using JMP 12.

Results and Discussion
The ability to detect and discriminate between clean and potentially infectious environments is vital to avoid the adverse consequences of infection. In this study we tested if infection avoidance behaviour in *Drosophila melanogaster* is modified by its previous exposure to a viral pathogen and by the risk of infection with that same pathogen when encountered during foraging. Viral exposure prior to the behavioural assay was achieved by placing flies in a DCV contaminated environment for 3 days, allowing flies to acquire DCV infection orally. DCV acquired through the oral route using this protocol continued to replicate within the fly, increasing by 10-100 fold by day 13 following oral exposure (F4,19 = 8.78, p=0.0003; Fig. 1A) and resulted in up to 20% mortality within this period (Fig. 1B).

In the foraging choice assay, only a fraction of flies chose either of the food sources provided, and this motivation to feed increased over time for flies in all treatment groups (χ²₁ = 11.00, p=0.001; Fig. 2A). The rate at which motivation increased differed between sexes ('Time × Sex' interaction, χ²₁ = 12.47, p=0.0004), and on average female flies showed greater motivation to feed than males (χ²₁ = 5.01, p=0.025), with 67% of female and 36% of male flies making a choice to feed on any of the provided substrates during the observation period. Once flies had made the choice to feed on one of the provided food sources, the choice between a clean and a potentially infectious food source was not affected by previous exposure to DCV ('previous exposure', χ²₁ = 0.513, p=0.47) in either male or female flies ('sex', χ²₁ = 0.595, p=0.44).
Across the entire observation period, the motivation to feed differed between sexes, and depended both on their previous exposure and on their current risk of infection ('Sex' × 'risk of infection' × 'Previous exposure' interaction, $\chi^2_{1} = 21.82$, $p<0.0001$). The proportion of males choosing any food substrate did not vary with previous exposure to DCV in either high-risk ($\chi^2_{1} = 2.21$, $p=0.137$) or no-risk environments ($\chi^2_{1} = 0.09$, $p=0.764$; Fig. 1). In female flies however, previous exposure and current infection risk affected the motivation to feed on the provided food sources. When there was no risk of infection (Fig. 2B, light grey bars) the motivation to feed was greater in females that were previously exposed to DCV than in otherwise healthy, non-exposed females ($\chi^2_{1} = 104.11$, $p<0.001$). Among females that were previously exposed to infection, we found that the presence of a risk of acquiring infection resulted in lower foraging effort - with just over 50% of flies making the choice to feed - compared to females in cages where there was no risk of acquiring infection, where over 80% of flies made the choice to feed (Fig. 2B; $\chi^2_{1} = 168.48$, $p<0.001$).

In addition to responding to external cues of infection (infection risk), internal physiological cues (in this case, previous exposure to DCV) may also modify avoidance behaviour. Behavioural modifications due to infection are widely reported among animals \(^9,27\), and can be classified into (i) parasitic manipulation that enhances parasite transmission \(^9\) (ii) sickness behaviours that benefit the host by conserving energetic resources during infection \(^13\), or (iii) side-effects of pathogenicity that do not benefit the host or the parasite \(^27\). Female flies infected orally with DCV are known to experience increased lethargy and sleep \(^24\), so these effects could also explain the reduced feeding
activity we detected in female flies that had been previously exposed to DCV.

Another potential explanation for reduced motivation to feed in previously exposed flies is infection-induced anorexia, a commonly described sickness behaviour. However, it is unlikely that a lower motivation to feed is simply a symptom of a "sick" fly, because it varied according to the risk of infection, and even reached 80% in exposed flies when foraging in a 'no risk' environment (Fig. 2). This suggests that flies are actively avoiding contact with the potentially contagious food source by lowering their foraging effort.

The higher motivation to feed of some female flies when the risk of infection was absent (Fig. 2) suggests flies were able to identify external cues of infection risk. Identifying infection cues is a general prerequisite of avoidance behaviours and occurs across a wide range of different taxa. For example, lobsters are known to detect and avoid virus-infected conspecifics; fruit flies and nematodes are capable of avoiding pathogenic bacteria; gypsy moth larvae are able to detect and avoid virus-contaminated foliage; sheep have been found to prefer to graze in parasite-poor patches; and it has been argued that the disgust response in humans has evolved because it decreases contact with potential infection. It is unclear how flies are able to identify food sources contaminated with a viral pathogen. In Drosophila and C. elegans avoidance of pathogenic bacteria is enabled by evolutionary conserved olfactory and chemosensory pathways, while avoidance of parasitic wasps appears to be mainly enabled by the visual sensory system. While avoiding virus infected conspecifics is probably driven by visual cues of infection, it remains unclear how virus-contaminated environments may trigger a lower motivation to feed in Drosophila.
The fact that only female flies demonstrated avoidance is an indication that any potentially adaptive effects of avoiding infection may be related to oviposition, which coincides with feeding. For flies previously exposed to DCV, avoiding infection would not confer substantial direct benefits given the physiological and behavioural costs of this infection, but would however reduce the exposure of future offspring to infection. While flies previously exposed to DCV do not appear to immune primed following an initial viral exposure, our results point to a sort of behavioural priming, where females previously exposed to infection avoid foraging in potentially infectious environments.

In summary, using a combination of experimental infections and behavioural assays, we find evidence for infection avoidance in Drosophila in the form of reduced motivation to feed, which was most pronounced when flies were faced with an increased risk of encountering an infectious food source. However, these effects were only present in female flies, indicating potentially important sexual dimorphism in infection avoidance.

Understanding how avoidance behaviours may vary is therefore important for our understanding of how disease will spread in natural populations, and more broadly how pathogens might evolve in response to variation in host infection avoidance strategies.

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Competing interests

The authors declare that they have no competing interests.

Author contributions

PFV conceived the study. PFV and MDJ designed the experiment. MDJ and PFV carried out the experimental work. PFV analysed the data, wrote the manuscript and provided all research consumables.
References


26. JMP. (SAS Institute Inc.).


Fig.1. Exposing flies to DCV by placing them in DCV-contaminated vials for three days resulted in flies acquiring replicating virus as shown by the increase in DCV titres over time (Fig. 1A). Grey points shown are individual replicate female flies, black bars are mean titres. This orally acquired DCV infection had a moderate effect on fly survival (full black line) compared to uninfected control flies (dotted line) (Fig. 1B).
Fig. 2. Single sex-groups of flies that had been previously exposed either to DCV or to a sterile inoculum were tested in a ‘no-risk’ environment (choice between two clean vials; light grey) or a ‘high-risk’ environment (choice between a clean vial and a DCV-contaminated vial; dark grey). The motivation to feed, measured as the proportion of flies in the cage that fed on any of the provided food sources, increased over time (Fig 2A). Fig 2B shows the average of motivation to feed taken across the whole observation period for each combination of fly sex, prior DCV exposure and current exposure risk (‘no-risk’ environment (light grey) or a ‘high-risk’ environment (dark grey). Data show means ± SE.