Insufficient evidence of infection-induced phototactic behaviour in *Spodoptera exigua*: a comment on van Houte *et al.* (2014)

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Some of the most spectacular interactions between hosts and parasites occur when parasites manipulate their hosts’ behaviour. Acanthocephalan worms that infect gammarid shrimps induce host behaviours which elevate predation vulnerability when they need to transmit to their final vertebrate host [1]. *Ophiocordyceps* fungi similarly increase transmission by forcing ants to clamp their jaws around leaves in elevated positions before killing them [2]. However, the mechanisms underlying such manipulations remain relatively obscure.

A recent paper by van Houte *et al.* [3] claims to demonstrate that: 1) infection with the baculovirus *Spodoptera exigua* multiple nucleopolyhedrovirus (SeMNPV) causes *S. exigua* larvae to die in an elevated position; and 2) this is achieved by the virus triggering a positive phototactic response in its larval host. Their study is grounded in knowledge that baculoviruses manipulate climbing behaviour in some lepidopteran species [4]. Here we argue van Houte *et al.*’s study is flawed: the experimental design cannot test the authors’ hypotheses and the data presented are open to other interpretations that do not support the authors’ conclusions.

Death in elevated positions?

First we consider van Houte *et al.*’s evidence that the virus SeMNPV induces death at an elevated height. The authors placed infected larvae in jars and recorded larval height over
several days until all pupated or died of infection. Baculovirus infection caused substantial mortality after 3-4 days. van Houte et al. show that larval height increased during an early climbing phase and that mean height of infected larvae remained high until the end of the experiment (their figure 1a). However, many of the larvae included in this data set had died of infection. The authors repeatedly recorded the height of dead larvae, despite the fact these larvae were clearly incapable of moving. We re-plotted their data excluding those that had previously died (our figure 1a) and show the association between infection and climbing is anything but clear cut. We also note that ongoing larval death means that the true sample size declined from 31 to 2 during van Houte et al.’s experiment. The observation that infected larvae “die at elevated positions” could be adequately explained by two simple facts: (i) larvae naturally climb; and (ii) viruses kill them, but not instantaneously.

Is phototaxis in infected larvae caused by viral infection?

Next we question the evidence the authors use to justify their conclusion that SeMNPV causes a change in host behaviour by inducing phototaxis. van Houte et al. placed SeMNPV infected larvae in three different light regimes: ‘continuous dark’, ‘lit from above’, and ‘lit from below’. The height of each larva at death was later recorded. The authors conclude that because larval height at death differs strikingly between these lighting treatments, SeMNPV infection induces phototaxis. This conclusion is undermined by the absence of suitable control experiments on uninfected larvae. These controls are necessary to demonstrate that the response of infected larvae to light regime change is caused by viral infection and does not also occur in uninfected individuals. Whilst the authors did study uninfected larvae (which were mock-infected), they did not subject them to the more informative ‘lit from below’ treatment (for reasons that were not mentioned). We therefore only have measures of phototactic behaviour for both infection classes for two of the lighting treatments (‘lit from above’ and ‘continuous dark’). Unfortunately, even these treatments are not comparable because the authors report completely different behavioural metrics in the two infection classes (uninfected: ‘height twice daily until pupation’; infected: ‘height at death’). These metrics cannot be directly compared without information about the time at which larvae died in the infected treatment.
The authors state that climbing in uninfected larvae is “not light-dependent”. This assertion is crucial to their argument that the virus induces phototaxis. However, van Houte et al. restrict their comparison of uninfected larvae in the light and dark treatment to two specific times at which the larvae moulted. At other times climbing behaviour in uninfected larvae differs markedly between the lighting regimes (our figure 1b). Indeed, the peak of climbing occurs 69 hours earlier in the ‘dark’ than in the ‘lit from above’ treatment. By these metrics the climbing of uninfected larvae is influenced by light. Therefore, it seems premature for the authors to conclude that viral infection drives the observed phototaxis in the infected treatment.

Perspective

We would like to be more constructive than simply to point out problems in van Houte et al.’s paper. Their data are compatible with a different hypothesis that does not require viral manipulation. SeMNPV may simply induce larval death during a peak in natural climbing behaviour, meaning that larvae die in elevated positions. This could potentially represent an interesting example of optimally timed host-killing by a pathogen [5] (but further experiments are necessary to properly test this). Whilst virally-induced host climbing has been demonstrated in another system [6], ‘tree top disease phenomena’ have been reported in numerous host species. In each of these cases it is important to determine whether this phenomenon results from viral manipulation of climbing behaviour, or from optimally timed larval killing, or both. To show that host behavioural changes actually result from parasite manipulation, future studies should endeavour to rule out plausible alternative explanations, including changes resulting from morbidity associated with infection, or adaptive host responses to parasitism [7]. Demonstration that the behaviour of infected hosts changes at a specific time which favours the parasite’s own fitness can provide definitive evidence of behavioural manipulation [8].

We do not doubt that behavioural manipulation of lepidopteran larvae by baculoviruses occurs in some host-virus systems, potentially including this one. Unfortunately, van Houte et al.‘s experiments lack sufficient comparable controls and cannot rule out possible alternative
explanations. It is our opinion that the conclusions of van Houte et al.'s paper are not supported by the data they present.

References


**Figure Legend**

**Figure 1.** Mean height of baculovirus-infected and healthy larvae. (a) Height of larvae following exposure to the baculovirus. Closed circles: original data. Open circles: data where larvae were excluded after the first point at which they are found to be dead (and therefore lose the ability to move). The dotted line represents % survival (left hand axis). (b) Height of uninfected larvae in light (12 L: 10 D) and dark (0 L: 24 D) treatments (closed and open circles, respectively) (data unchanged from van Houte *et al.*). Error bars show SE.