



Invasion success of a widespread invasive predator may be explained by a high predatory efficacy but may be influenced by pathogen infection

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Abstract Invasive alien species (IAS) can drive community change through ecological interactions. Parasites and pathogens can play an important role in community function including mitigating or enhancing IAS impacts. Despite this, the degree to which pathogen pressure influences IAS impacts remains poorly understood. We quantified the predatory behaviour of the highly invasive alien harlequin ladybird (*Harmonia axyridis*) and two UK native species, the 7-spot (*Coccinella septempunctata*) and 2-spot (*Adalia bipunctata*) ladybirds, using comparative functional response experiments. We investigated the impacts of pathogen infection on the predatory ability of the ladybirds by exposing individuals to *Beauveria bassiana*, a widespread entomopathogen. Invasive *H. axyridis* was a more efficient predator than

both the native *A. bipunctata* and *C. septempunctata*, often having higher attack and/or lower prey handling time coefficients, whereas native *A. bipunctata* were the least efficient predators. These differences were found in both adult and larval life-stages. *Beauveria bassiana* infection significantly altered the predatory efficiency of adult and larval ladybird predators. The effects of pathogenic infection differed between species and life-stage but in many cases infection resulted in a reduced predatory ability. We suggest that the interactions between IAS and pathogens are integral to determining invasion success and impact.

Keywords *Beauveria bassiana* · Functional response · *Harmonia axyridis* · Invasive alien species · Parasite · Predator

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Introduction

The rate of species invasions has increased in recent decades with the expansion of global trade and movement, and further rate increases appear likely (Levine and D'Antonio 2003; Hulme 2009; Seebens et al. 2017; Pfieglers et al. 2018). As a result, understanding the impacts of species invasion events has rarely been more important. The impacts that IAS can impose on native systems vary with respect to their trophic position and key functional behaviours, such

as predation, which can also facilitate invader success (Salo et al. 2007; Bellard et al. 2016). Although characteristics of the invader can influence its effects, they can also differ according to characteristics of the recipient community. Parasites and pathogens play key roles within communities and can provide resistance to species invasions and modify the impacts of invading species, in addition to colonising novel areas as IAS themselves (Hatcher et al. 2014; Vilcinskas 2015; Roy et al. 2016b). Key functional roles are undertaken by parasites and pathogens through lethal and sub-lethal trait effects (Dunn and Hatcher 2015). Lethal effects of parasites can affect host population densities and result in population declines whereas the sub-lethal effects of infection can result in more complex impacts (Hatcher et al. 2014; Dunn and Hatcher 2015). For example, Roy et al. (2008b) provided evidence that harlequin ladybirds (*Harmonia axyridis*; Coleoptera, Coccinellidae) infected with *Beauveria bassiana* (Ascomycota, Cordycipitaceae) showed reduced egg production. Sub-lethal effects of parasites can also affect species with which hosts interact; for example, Dick et al. (2010) showed that *Gammarus pulex* (Crustacea, Amphipoda) infected with *Echinorhynchus truttae* acanthocephalan worms consume prey at an increased rate compared to uninfected conspecifics whereas Iltis et al. (2018) show similar results for infected *Dikerogammarus villosus* (Crustacea, Amphipoda). Despite their widespread presence within communities, parasites and pathogens are often absent from studies investigating the impacts of IAS (Hatcher et al. 2012), potentially resulting in oversimplified study systems that are unlikely to be representative of those in the field. However, it should be noted that efforts are ongoing to account for such interactions within some study systems (for example, with respect to *H. axyridis* see Haelewaters et al. 2017). Accounting for these effects not only provides insight as to species behaviours at suboptimal health, but also the role of parasites and pathogens during species invasions. Ultimately, understanding how species interact with parasites and pathogens is essential to accurately predict the ecological impacts of current and future invasive alien species.

Predation can be a key way in which IAS can influence native communities (e.g., Doherty et al. 2016). Predatory behaviour can be quantified through the use of predatory functional responses which

describe the relationship between a species' resource use and the availability of that resource (Holling 1959). Functional response studies aim to replicate one part of a complex interaction between predators and their prey in a simplistic form, specifically, how a predator's rate of prey consumption changes with respect to prey density. Functional response experiments have been shown to correctly predict the impact of multiple invasive alien species, for example the 'bloody red' shrimp (*Hemimysis anomala*) (Dick et al. 2013). Functional response experiments provide a *per-capita* measure of predatory ability and subsequently predatory pressure imposed on the prey species which can be compared between species and/or treatments and used to estimate population level impacts (Dick et al. 2017b; Laverly et al. 2017). Predatory functional responses have historically been used in population and community ecology as well as in pest management via biological control (for example O'Neill 1990; Sabelis 1992; Cuthbert et al. 2018). More recently, predatory functional responses have been applied within invasion ecology to understand and predict the impacts of invasive species (Alexander et al. 2014; Dick et al. 2017a; Uiterwaal and Delong 2018). Functional response experiments enable predation behaviour to be defined as one of three response types (I, II and III) and predatory coefficients to be calculated [handling time (h) and attack rate (a)] (Holling 1959) across a range of prey densities. The functional response type can inform the likely ecological impact of the predator on the prey population. For example, a predator displaying a type II relationship could be expected to have a destabilising impact on the prey community as prey would be reduced to low densities or result in localised extinction (Hassell 1978). Conversely, a type III relationship suggests that the predator is likely show prey switching behaviour when the primary prey species reaches low densities, suggesting a more stable relationship with the prey community (Murdoch and Oaten 1975). Ladybird assemblages within the UK provide a model system for studying the impacts of an IAS within a community context (Roy et al. 2016a).

The harlequin ladybird (*H. axyridis*) is a highly invasive coccinellid predator that has invaded throughout the world aided by multiple releases as a biological control agent (Roy et al. 2016a; Camacho-cervantes and Ortega-iturriaga 2017). *Harmonia axyridis* will predate the immature stages of the

monarch butterfly (*Danaus plexippus*) (Koch et al. 2003). In addition to impacting prey species, *H. axyridis* has also led to declines of native ladybird populations through, at least in part, intra-guild predation (Ware and Majerus 2008; Katsanis et al. 2013). *Harmonia axyridis* now dominates many Coccinellidae assemblages throughout its invaded range (e.g. Brown and Roy 2017) which has resulted in reduced species diversity (Harmon et al. 2007; Koch and Galvan 2008; Bahlai et al. 2014; Grez et al. 2016). Following the arrival of *H. axyridis* in the UK in 2004, the 2-spot ladybird (*Adalia bipunctata*) showed a decline of 44% while 7-spot ladybird (*Coccinella septempunctata*) populations showed no significant change (Roy et al. 2012). Both of these species are historically common in the UK. The predatory ability of *H. axyridis* is believed to have been instrumental in the population declines of native Coccinellidae whilst giving the invasive species a competitive advantage, therefore facilitating its continued spread (for example Majerus et al. 2006; Xue et al. 2009; Abbott et al. 2014; Wu et al. 2018). *Harmonia axyridis* has been described as a voracious aphid predator (Majerus et al. 2006), however, the impacts on prey populations within the invaded range are less studied (Roy and Brown 2015; Roy et al. 2016a).

Beauveria bassiana is a widespread entomopathogenic fungus, found in multiple habitats including soils (Ormond et al. 2010), hedgerows (Meyling and Eilenberg 2006), and trees (Ormond et al. 2010; Howe et al. 2016), and is known to interact with UK Coccinellidae. Ormond et al. (2006) show that *B. bassiana* is a major cause of overwintering mortality for native *C. septempunctata*. While there have been no published field records of *B. bassiana* infection in *A. bipunctata*, Roy et al. (2008b) suggest this could be due to differences in habitat use, which result in *A. bipunctata* encountering *B. bassiana* less frequently. Roy et al. (2008b) also provide evidence, through a laboratory study, that *A. bipunctata* are similarly susceptible to *B. bassiana* infection as *C. septempunctata*, with both native species being more susceptible to pathogenic infection than *H. axyridis*. Howe et al. (2016) also report instances of infection in invasive alien *H. axyridis* field populations. Infection at lower doses can also be long lasting and result in sub-lethal trait mediated effects; for example, Roy et al. (2008b) showed that *B. bassiana* infection

reduces egg production of *H. axyridis*. Despite the recorded instances of infection, *H. axyridis* is known to be less susceptible to *B. bassiana* than many native ladybird species (Cottrell and Shapiro-Ilan 2003; Roy et al. 2008b; Verheggen et al. 2017; Haelewaters et al. 2018) and this is likely due to *H. axyridis*' advanced chemical defences (Koyama and Majerus 2007; Roy et al. 2008a; Shapiro-Ilan and Cottrell 2011; Verheggen et al. 2017; Gegner et al. 2018). Despite *H. axyridis* commonly interacting with *B. bassiana*, our understanding as to how *B. bassiana* infection changes *H. axyridis* behaviour in relation to those natives which have coevolved with this pathogen remains incomplete. For example, should *B. bassiana* infection reduce the ability *H. axyridis* to forage efficiently this could result in reduced pressure on aphids through predation and on native ladybird communities through competition and intra-guild predation.

In this study we aimed to compare the predatory behaviour of the invasive alien *H. axyridis*, and two UK native ladybird species; *A. bipunctata* and *C. septempunctata*, during their larval and adult life stages, so as to better understand the ecological impact of the *H. axyridis* invasion and any potential insights as to *H. axyridis*' invasion success. So as to correctly define predatory functional responses, a process that is often challenging, we used three statistical techniques; linear regression, LOESS curve fitting, and AICc scores. We compared adult and larval *H. axyridis* with adult and larval *A. bipunctata* and adult *C. septempunctata*. We also investigated how pathogen infection impacts the predatory ability of the three species across their larval and adult life stages by exposing individuals to a sub-lethal dose of *B. bassiana*. While recorded instances of *B. bassiana* infection in *H. axyridis* and *A. bipunctata* are rarely observed in the field, such infections are known to occur and could feasibly change in the future, potentially as a result of spillover from the invasive *H. axyridis*. Additionally, any changes in the susceptibility of either *H. axyridis* or *A. bipunctata* to *B. bassiana* would likely increase the exposure of the other co-occurring ladybird species to *B. bassiana*. We hypothesised that: the invasive alien *H. axyridis* would demonstrate more efficient predatory behaviour than the native species. Efficient predatory behaviour was defined as having a higher overall functional response relationship, increased attack rate or reduced handling time. We

further investigated how sub-lethal *B. bassiana* pathogenic infection would impact the predatory efficiency of the three ladybird species as this could shed light on the ecological impacts of *H. axyridis* and how these could change when exposed to a widespread pathogen.

Materials and methods

Insect cultures

We collected first and second larval instars of *H. axyridis* and adult *C. septempunctata* in Oxfordshire (51°60'N; – 1°11'W) in summer (June–August) 2016 through visual and sweep net sampling of vegetation. Due to their scarcity, we purchased *A. bipunctata* first and second instar larvae from an industrial supplier (Green Gardener, UK) and, as they were also scarce in this particular field season, we only collected adult *C. septempunctata*. Therefore, no larval trials were conducted with *C. septempunctata* larvae. All individuals were maintained at constant conditions (20 °C, 16:8 L:D cycle) for at least 7 days prior to experimentation. We reared *H. axyridis* and *A. bipunctata* larvae in control conditions until their use in either larval or adult experiments. We fed individuals a mixed diet of sycamore aphids (*Drepanosiphum platanoidis*; frozen, mixed age classes), *Ephestia kuehniella* (Lepidoptera, Pyralidae) eggs (Entofood, Koppert, the Netherlands) and an artificial diet (detailed by Roy et al. 2013). We purchased English grain aphids (*Sitobion avenae*) from a commercial supplier (Ervibank, Koppert, the Netherlands) and reared them in the same conditions on the wheat plants on which they were received. We sexed adult ladybirds using established physical characteristics (McCornack et al. 1980; Roy et al. 2011). We used females in experimental trials as they are known to consume prey at higher rates than males (Xue et al. 2009; Gupta et al. 2012; Honek et al. 2017). Due to the inability to sex ladybird larvae, the larval treatments were of mixed sex. All larval treatments used fourth instar ladybird larvae.

Beauveria bassiana infection

We cultured *B. bassiana* from a commercially available product (Botanigard WP, strain GHA) on

Sabouraud dextrose agar (SDA) in Petri dishes in darkness at 25 °C. We prepared single spore isolations from these cultures, and subsequently sub-cultured under the same conditions before being stored at – 20 °C in 10% glycerol (v/v sterile milli-Q water) as a cryoprotectant. Thawed sub-cultures were macerated, spread onto fresh SDA plates and cultured for approximately 14 days until sporulation. We prepared spore suspensions in 0.03% Tween 20 (v/v in sterile water) surfactant to reduce spores clumping together and the concentration of the resulting suspension was estimated using a Neubauer improved haemocytometer. We produced a 10⁶ spores ml⁻¹ dilution from the stock suspension approximately 16 h prior to the experiment, stored on-ice and homogenised before use in experiments. This dose was aimed to provide an ecologically relevant dose that could feasibly impact predatory behaviour (Roy et al. 2008b).

We inoculated ladybird predators with one of two treatment solutions; a control treatment of 0.03% Tween 20 or a 10⁶ spores ml⁻¹ *B. bassiana* spore suspension for infection treatments. Roy et al. (2008b) report the LD₅₀ (median lethal dose) of native *C. septempunctata* and *A. bipunctata* were similar at 10⁶ and 10^{6.2} spores ml⁻¹ respectively whereas invasive alien *H. axyridis* had an LD₅₀ of 10^{9.6} spores ml⁻¹. Individuals were inoculated by inversion (five times) in 1 ml of inoculum and were placed on filter paper (Whatman No. 1) in a Büchner funnel to remove excess inoculum. All equipment was cleaned with 95% ethanol between treatments. Following exposure to *B. bassiana*, treatment groups were housed separately to prevent contamination and starved for 8 h to standardise gut contents before the start of the experiment.

Experimental methods

Experimental arenas consisted of a Petri dish (90 mm) and contained blades of winter wheat (*Triticum aestivum*; ten strips, 40 mm in length) embedded in 2% water agar, approximately four mm in depth, so as to increase habitat heterogeneity. Filter papers (Whatman No. 1) were positioned in the lids to moderate moisture levels. Wheat was grown from seed (Syngenta) for 14 days before use. Grain aphids (*Sitobion avenae*) were provided as a prey resource at known densities of live second and third instar individuals.

Fourth instar larval treatments were provided with prey densities of; 1, 2, 4, 8, 16, 32, 64 and 128 individuals. Adult treatments received prey densities of 1, 2, 4, 8, 16, 32, 64, 128 and 256 individuals. Adults received an additional prey density treatment as they are known to consume more prey than larvae. We aimed to replicate each treatment combination five times, due to ladybird mortality the total number of treatment replicates varied between four and six (Supplementary Table 1). Each prey density was replicated five times with a control treatment, consisting of the same experimental areas and prey densities as the adult and larval treatments with no predator present.

Predators were weighed after their starvation period before being added to the experimental arenas. The experiment ran for 24 h at constant conditions during which time predation of aphid prey could occur. After 24 h the ladybirds were removed from the arenas and remaining prey were counted. No cases of partial consumption were observed. Individuals were starved for a further 12 h before resuming a mixed diet and were monitored for mortality over the next 14 days. Adult cadavers, collected within the 14 day post-experiment observation period, were surface sterilised using a 1% bleach solution to reduce contamination, before being plated out on 2% water agar and incubated in darkness at 25 °C. Incubated cadavers were visually checked for signs of fungal sporulation for a period of 14 days.

Statistical analysis

All statistical analyses were undertaken in R version 3.3.2 and RStudio version 1.0.136 (R Core Team 2016; RStudio 2016). We compared ladybird masses with respect to species and treatment using ANOVA and TukeyHSD post-hoc statistical tests for both life stages. The number of prey surviving in predator treatments was compared to the control treatments using linear regression with, in response to signs of overdispersion, a quasipoisson error structure. We compared the number of prey consumed in the predator treatments between species and treatments,

for both larvae and adult predators, using generalised linear models with quasipoisson error structures.

Functional responses

Functional response curve fitting was undertaken in R using the *bbmle* and *emdbook* statistical packages (Bolker and R Development Core Team 2014; Bolker 2016). Defining predatory functional response relationships can be difficult. In an attempt to overcome this, we used three methods to characterise predatory behaviour; linear regression of the proportion of prey eaten against prey density, second order Akaike information criterion (AICc) values and the qualitative inspection of non-parametric locally weighted scatter plot smoothing (LOESS) curves (Juliano 2001).

Functional response relationships were fitted using Holling’s original type I equation (Eq. 1), Rogers’ type II equation (Eq. 2), and Hassell’s type III equation (Eq. 3). Hassell’s type III and Rogers’ type II equations are similar however, while Rogers’ type II includes an attack rate parameter (a), Hassell’s type III assumes the attack rate varies with prey density via a hyperbolic relationship. Rogers’ type II and Hassell’s type III equations both account for prey depletion (Rogers 1972; Hassell 1978) and rely on the Lambert W function (Bolker 2016).

$$N_e = aTN_0 \tag{1}$$

$$N_e = N_0 \left(1 - e^{(a(N_e h - T))} \right) \tag{2}$$

$$N_e = N_0 \left(1 - e^{\frac{(d + bN_0)(hN_e - T)}{1 + cN_0}} \right) \tag{3}$$

In all equations N_e denotes prey consumed, N_0 is the number of prey provided, T is the time during which behaviours occurred, a and h are attack rate and handling time coefficients of the predators. In Hassell’s type III equation (Eq. 3) b , c and d are used to calculate the hyperbolic a . The attack rate constant (a) is defined as the rate of prey consumption and informs the gradient of the functional response curve whereas the handling time coefficient (h) is the rate of saturation and provides insight as to the time predators spend handling prey between attacks. Together these parameters define the predator’s overall functional response.

Comparing predatory behaviours

Predatory statistics of attack rates (a) and handling times (h) were calculated and compared using non-linear least squares regression (as described by Juliano 2001). The number of prey consumed was regressed against the initial density, density² and density³. Type I and II responses would be indicated by a significant and negative first order term (density) and a type III response would be indicated by a significant and positive first order (density) and quadratic term (density²) or a significant third order term (density³) (Juliano 2001). Confidence intervals were calculated for each functional response relationship through bootstrapping ($n = 999$). Separate models were fitted for fourth instar larvae and adult predator treatments.

Results

Prey survival in control treatments, containing no predators or fungal pathogen, was 86.9%, which was significantly higher than predator treatments ($H.axyridis = 48.8\%$, $C.septempunctata = 50\%$ and $A.bipunctata = 70.8\%$) (Table 1). Prey mortality was therefore attributed to predatory behaviour of the focal predators. *Beauveria bassiana* infection was confirmed in 63% of adult and 48.5% of larvae infection treatment individuals that died following experiments. 5.9% of uninfected treatment adults and no larvae showed infection.

The masses of adult ladybird species did not differ between the infection treatment groups (ANOVA; $F_{2,257} = 0.836$, $P = 0.434$) and for each of the three ladybird species, ladybird masses were not significantly different between treatment groups (ANOVA;

$F_{1,259} = 1.117$, $P = 0.291$), as a result we did not account for mass in subsequent models. Adults of each species differed significantly in mass (ANOVA; $F_{2,260} = 500.9$, $P < 0.001$) and TukeyHSD results indicated this is driven by *A. bipunctata* (mean \pm SD, 9.97mg \pm 2.4, $n = 89$) being significantly smaller than *H. axyridis* (36.05mg \pm 7.38, $n = 88$) and *C. septempunctata* (37.22mg \pm 8.34, $n = 86$). We found no evidence that larvae masses varied significantly with *B. bassiana* infection treatments whether we accounted for species differences or not (ANOVA; $F_{1,152} = 2.912$, $P = 0.09$ and $F_{1,153} = 2.461$, $P = 0.119$). As with adult predators, larval *A. bipunctata* (5.87mg \pm 2.41, $n = 80$) were significantly smaller than *H. axyridis* (mean \pm SD, 19.51mg \pm 8.91, $n = 76$) (ANOVA; $F_{1,154} = 174.2$, $P < 0.001$).

In adult treatments, the three ladybird species consumed prey at significantly different rates (GLM; $F_{(2,259)} = 11.952$, $P < 0.001$), with invasive *H. axyridis* consuming the most and *A. bipunctata* consuming the least, and more prey were consumed with the increasing prey density treatments (GLM; $F_{(2,259)} = 268.848$, $P < 0.001$). Pathogen exposure did not significantly impact the number of prey consumed by adult ladybirds with each of the interaction terms containing the pathogen treatment (density * species * pathogen, species * pathogen, density * pathogen and density * species) and the main effect all being removed at $P > 0.05$. Although ultimately removed from the final model, a marginally significant species * pathogen interaction term (GLM; $F_{(2,256)} = 2.965$, $P = 0.054$) suggested that the pathogen exposure might have changed the prey consumption of the ladybird species differently. In larval treatments, the number of prey consumed with

Table 1 Results of a logistic regression of the total prey consumed in each prey density treatment for each species

	Coefficient (\pm SE)	t	P
Intercept	- 3.266 (\pm 2.021)	- 1.616	0.107
Density	0.689 (\pm 0.030)	23.029	< 0.001***
<i>Adalia bipunctata</i>	5.029 (\pm 2.023)	2.487	0.014*
<i>Coccinella septempunctata</i>	5.708 (\pm 2.022)	2.823	0.005**
<i>Harmonia axyridis</i>	5.907 (\pm 2.021)	2.922	0.004**

Results indicate each species consumed significantly more prey than control treatments in which predators were absent. Prey density was also found to be significant, with more prey consumed at higher prey densities. The analysis was carried out using a quasi-poisson error structure with prey density values scaled and centred. Asterisks denote significance of P values; * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$

increasing density treatment changed significantly when ladybird predators were subject to pathogen exposure (GLM; $F_{(1,151)} = 1075.8$, $P = 0.010$). Similar to the adult treatments, larvae of the two ladybird species consumed significantly different numbers of prey (GLM; $F_{(1,153)} = 65.962$, $P < 0.001$), with *H. axyridis* larvae consuming more than *A. bipunctata* larvae. As with the adult ladybird analyses, all other terms were removed from the final model at significance values (P) of more than 0.05.

Functional responses

All species treatments showed type II functional responses (Fig. 1).

Logistic regression of the proportion of prey consumed against prey density indicated that 7 of the

10 treatments revealed a significant and negative first order term (density) (Supplementary Tables 2a and 2b), indicating a type II functional response. Two of these analyses showed a significant second order term (density²), however, these were positive and did not indicate a type III response. No density terms were significant in three treatments; uninfected *A. bipunctata* and infected *C. septempunctata* adults and infected *H. axyridis* larvae. This could suggest either a type I relationship or that the functional response relationship was undetectable. Further investigation of these treatments using AICc values of the fitted functional response equations (Eqs. 1–3) suggested a type II response for uninfected *A. bipunctata* adults (Supplementary Table 3). AICc values for infected *C. septempunctata* adults suggested a type III relationship and an indistinguishable type II/III relationship

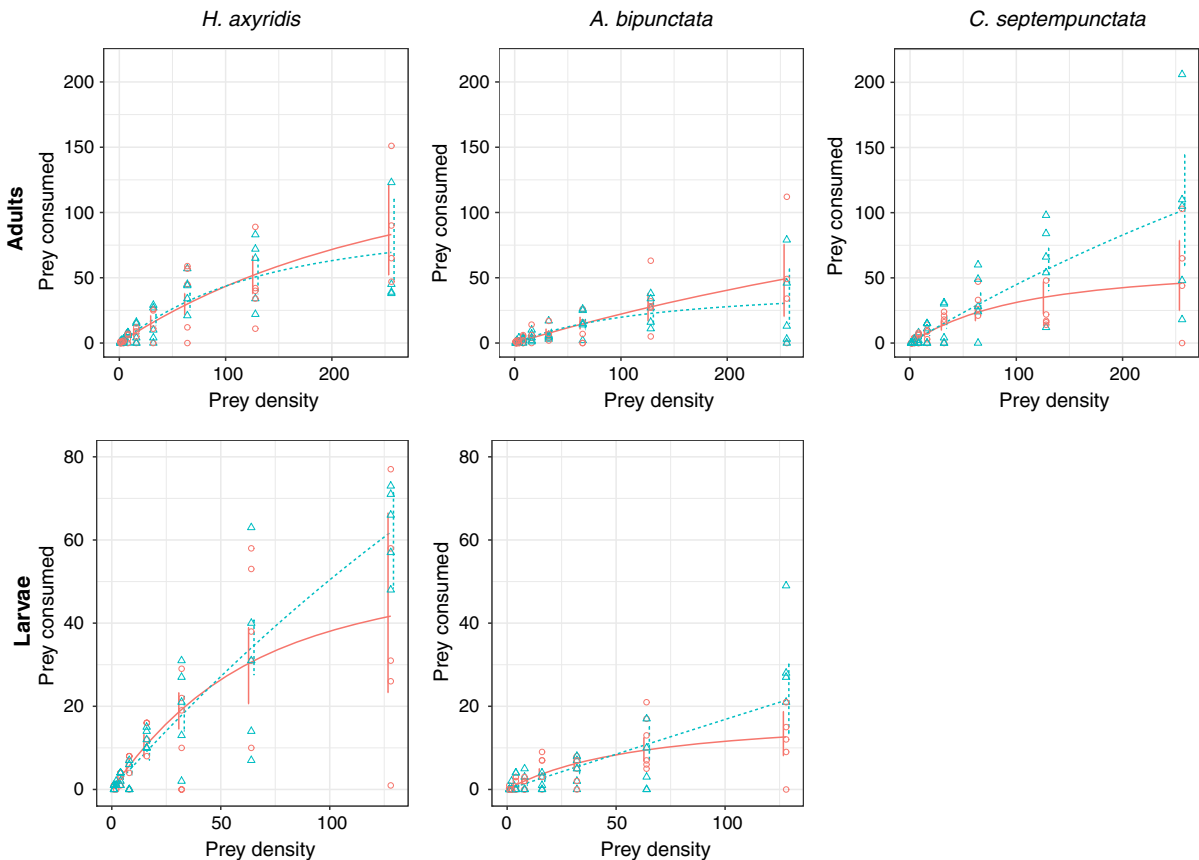


Fig. 1 Predatory functional response curves for three ladybird predators; invasive alien *H. axyridis* (left) and native *A. bipunctata* (middle) and *C. septempunctata* (right) across their adult (top) and larval (bottom) life-stages. Functional response curves (lines) are displayed with replicate data (points; Table 1)

and bootstrapped 95% confidence intervals ($n = 999$) (vertical lines). Uninfected predators (red solid lines and circles) were inoculated with a control dose of Tween 20 and *B. bassiana* infected predators (blue dashed lines and triangles) were inoculated with a 10^6 suspended spore solution

for infected *H. axyridis* larvae. Visual inspection of fitted LOESS curves provided qualitatively similar results (Supplementary Figure 1). As the majority of methods and treatments showed type II responses, this was accepted for all species-treatment combinations.

Comparing predatory behaviours

Visual inspection of functional response curves suggested between-species differences in predatory behaviour, as well as different responses to infection. The functional response curves suggested that *H. axyridis* consumed prey at a higher rate than native *A. bipunctata* and *C. septempunctata* (Fig. 1) and this was associated with increased attack rate and handling time coefficients which suggest a greater forage ability (Table 2). A similar result was also seen in larval treatments with invasive *H. axyridis* consuming more prey than native *A. bipunctata* (Fig. 1).

Predators responded to *B. bassiana* infection differently, varying with species and life-stage (Table 3). Infected *H. axyridis* and *A. bipunctata* adults showed lower functional response curves than uninfected conspecifics. In contrast, larval treatments showed the opposite response with infected individuals consuming more prey than uninfected individuals. Adult *C. septempunctata* showed an opposing response to infection than other adult treatments, instead infected individuals ate more than uninfected individuals. Pathogenic infection also increased the variation in predation, with infected individuals eating at both higher and lower rates than uninfected treatments. In all pairwise comparisons between infected and uninfected treatments, functional response curves differed the most in the higher prey density treatments (Fig. 1).

Predatory behaviour appeared to differ between treatments but as the confidence intervals for the fitted functional response relationships overlapped we explored these relationships further through comparison of predatory statistics [attack rates (*a*) and handling times (*h*)]. It is important to note that the predatory coefficients (*a*, *h*, and maximum feeding rates) are intrinsically linked and combined result in the overall predatory behaviour exhibited by the species. For example, an increase in a predators handling time will result in a decreased maximum feeding rate. Pairwise comparisons of attack rate (*a*) and handling time (*h*) showed significant

differences between species treatment combinations (Table 2). Species differed significantly with respect to their predatory behaviour with 36 or 42 pairwise comparisons between handling time and attack rate coefficients being significantly different ($P < 0.001$) and in each comparison at least one of the predatory statistics (*a* and *h*) was significantly different (Table 2).

Within species treatments, *B. bassiana* infection resulted in increased attack rates (*a*) in adult *H. axyridis* (uninfected = 0.762, infected = 1.005, $z = -2.696$, $P = 0.007$) and *A. bipunctata* (uninfected = 0.281, infected = 0.392, $z = 2.189$, $P = 0.029$) (Table 3). Adult *C. septempunctata* showed no significant change in attack rate when subjected to pathogen pressure ($P = 0.323$). Conversely, infected ladybird larvae showed lower attack rates in comparison to their uninfected conspecifics (Table 3). However, when adult ladybirds were subjected to pathogen infection *C. septempunctata* showed a shortening of handling times whereas *A. bipunctata* and *H. axyridis* both showed increases (Table 3). Larval treatments of both species showed shorter handling times when exposed to the pathogen (Table 3).

Discussion

Consistent with our hypothesis, we have shown that a widespread invasive alien predator (*Harmonia axyridis*) consumes more prey than native *Adalia bipunctata* and *Coccinella septempunctata*. The higher consumption rate of *H. axyridis* was linked with better forage ability including higher attack rate and shorter handling time coefficients. Typical efficient predatory behaviour would consist of high rates of attack on prey and short periods of time spent handling and consuming prey. We suggest this *per-capita* difference in predatory consumption and forage ability between native and IAS could shed light on the documented ecological impacts of *H. axyridis*. Specifically, these attributes could give *H. axyridis* an ecological advantage over native competitors (e.g. other Coccinellidae) and prey (e.g. aphid) species. Previous literature has suggested that the invasive *H. axyridis* is an efficient predator of aphid pests (Xue et al. 2009; Abbott et al. 2014; Wu et al. 2018) and this is likely to have facilitated the species' spread following multiple

Table 2 Maximum likelihood comparisons of functional response parameters [attack rate (*a*) and handling time (*h*)] between species and treatments

Life stage	Base species-treatment		Contrast species-treatment		Metric	Estimate (± SE)	z	P		
Adult	Infected	<i>A. bipunctata</i>	Infected	<i>C. septempunctata</i>	<i>a</i>	- 0.273 (±0.06)	- 4.520	< 0.001***		
					<i>h</i>	0.020 (± 0.003)	6.925	< 0.001***		
				<i>H. axyridis</i>	<i>a</i>	- 0.614 (± 0.083)	- 7.404	< 0.001***		
					<i>h</i>	0.012 (± 0.003)	4.259	< 0.001***		
					Uninfected	<i>A. bipunctata</i>	<i>a</i>	0.110 (± 0.050)	2.189	0.029*
							<i>h</i>	0.017 (± 0.004)	4.834	< 0.001***
		<i>C. septempunctata</i>	<i>a</i>	<i>a</i>	- 0.354 (± 0.083)	- 4.284	< 0.001***			
				<i>h</i>	0.006 (± 0.003)	1.943	0.052.			
			<i>H. axyridis</i>	<i>a</i>	- 0.371 (± 0.070)	- 5.280	< 0.001***			
				<i>h</i>	0.016 (± 0.003)	5.628	< 0.001***			
				<i>C. septempunctata</i>	Infected	<i>H. axyridis</i>	<i>a</i>	- 0.342 (± 0.083)	- 4.128	< 0.001***
							<i>h</i>	- 0.008 (± 0.001)	- 7.138	< 0.001***
	Uninfected	<i>A. bipunctata</i>	<i>a</i>		0.383 (± 0.050)	7.642	< 0.001***			
			<i>h</i>		- 0.003 (± 0.002)	- 1.036	0.300			
		<i>C. septempunctata</i>	<i>a</i>		- 0.082 (± 0.083)	- 0.989	0.323			
			<i>h</i>		- 0.014 (± 0.002)	- 8.105	< 0.001***			
	<i>H. axyridis</i>	<i>a</i>	- 0.097 (± 0.070)	- 1.392	0.164					
		<i>h</i>	- 0.004 (± 0.001)	- 3.168	0.002**					
		Uninfected	<i>A. bipunctata</i>	Infected	<i>H. axyridis</i>	<i>a</i>	- 0.724 (± 0.076)	- 9.553	< 0.001***	
						<i>h</i>	- 0.005 (± 0.002)	- 2.099	0.036*	
			Uninfected	<i>C. septempunctata</i>	<i>a</i>	- 0.464 (±0.076)	- 6.147	< 0.001***		
					<i>h</i>	- 0.011 (± 0.003)	- 4.075	< 0.001***		
	<i>H. axyridis</i>			<i>a</i>	- 0.481 (± 0.061)	- 7.802	< 0.001***			
				<i>h</i>	- 0.001 (± 0.002)	- 0.409	0.683			
<i>C. septempunctata</i>	Infected	<i>H. axyridis</i>	<i>a</i>	- 0.259 (± 0.100)	- 2.586	0.010**				
			<i>h</i>	0.006 (± 0.002)	3.544	< 0.001***				
	Uninfected	<i>H. axyridis</i>	<i>a</i>	- 0.017 (± 0.090)	- 0.183	0.855				
			<i>h</i>	0.010 (± 0.002)	5.797	< 0.001***				
		<i>H. axyridis</i>	Infected	<i>H. axyridis</i>	<i>a</i>	- 0.243 (± 0.090)	- 2.696	0.007**		
					<i>h</i>	- 0.004 (± 0.001)	- 3.475	< 0.001***		
Larvae	Infected	<i>H. axyridis</i>	Infected	<i>A. bipunctata</i>	<i>a</i>	0.781 (± 0.079)	9.929	< 0.001***		
					<i>h</i>	0.052 (± 0.013)	3.900	< 0.001***		
				Uninfected	<i>A. bipunctata</i>	<i>a</i>	0.546 (± 0.095)	5.747	< 0.001***	
						<i>h</i>	- 0.050 (± 0.010)	- 4.829	< 0.001***	
					<i>H. axyridis</i>	<i>a</i>	- 0.486 (± 0.151)	- 3.219	0.001**	
						<i>h</i>	- 0.013 (± 0.002)	- 5.833	< 0.001***	
	Uninfected	<i>H. axyridis</i>	Infected	<i>A. bipunctata</i>	<i>a</i>	0.816 (± 0.078)	10.545	< 0.001***		
					<i>h</i>	0.049 (± 0.012)	4.276	< 0.001***		
		Uninfected	<i>A. bipunctata</i>	<i>a</i>	1.033 (± 0.143)	7.237	< 0.001***			
				<i>h</i>	- 0.037 (± 0.011)	- 3.574	< 0.001***			
			<i>A. bipunctata</i>	Infected	<i>a</i>	0.224 (± 0.060)	3.770	< 0.001***		
					<i>h</i>	0.103 (± 0.017)	6.079	< 0.001***		

Functional response parameters were calculated through the fitting of the Rogers' 'random predator' type II functional response equation (Eq. 2). Maximum likelihood comparisons are made using methods described by Juliano (2001). Asterisks denote significance of *P* values; **P* < 0.05; ***P* < 0.01; ****P* < 0.001

Table 3 Comparison of predicted attack rate (*a*) and handling time (*h*) coefficients between infected and uninfected predator treatments

Life-stage	Species	Metric	Uninfected (\pm SE)	Infected (\pm SE)	<i>P</i>
Adult	<i>A. bipunctata</i>	<i>a</i>	0.281 (\pm 0.027)	0.392 (\pm 0.043)	0.029*
		<i>h</i>	0.005 (\pm 0.002)	0.022 (\pm 0.003)	< 0.001***
	<i>C. septempunctata</i>	<i>a</i>	0.746 (\pm 0.071)	0.664 (\pm 0.042)	0.323
		<i>h</i>	0.016 (\pm 0.002)	0.002 (\pm 0.001)	< 0.001***
	<i>H. axyridis</i>	<i>a</i>	0.762 (\pm 0.056)	1.005 (\pm 0.071)	0.007**
		<i>h</i>	0.006 (\pm 0.001)	0.010 (\pm 0.001)	< 0.001***
Larvae	<i>A. bipunctata</i>	<i>a</i>	0.333 (\pm 0.057)	0.192 (\pm 0.023)	0.032*
		<i>h</i>	0.054 (\pm 0.010)	0.003 (\pm 0.009)	< 0.001***
	<i>H. axyridis</i>	<i>a</i>	1.366 (\pm 0.131)	0.879 (\pm 0.076)	0.001**
		<i>h</i>	0.017 (\pm 0.002)	0.004 (\pm 0.002)	< 0.001***

Coefficients were calculated by fitting Rogers' 'random predator' type II functional response equation (Eq. 2) and compared using maximum likelihood. Asterisks denote significance of *P* values; **P* < 0.05; ***P* < 0.01; ****P* < 0.001

releases as a biological control agent. We show that *H. axyridis* is indeed an effective predator, in keeping with our initial hypothesis and previous literature.

Predatory behaviours differ between species

We have also provided evidence that the two larger ladybird species (*H. axyridis* and *C. septempunctata*) were more similar, with respect to their predation rates, than the smaller *A. bipunctata*. These findings are consistent with both our hypotheses and the wider literature. For example, Xue et al. (2009) show that *H. axyridis* and *C. septempunctata* both show type II functional responses when provided with soybean aphids (*Aphis glycines*) during larval and adult life stages. Similarly, Jalali et al. (2010) provide evidence that adult *A. bipunctata* also show a type II functional response curves when provided a diet of pea aphids (*Acyrtosiphon pisum*). As far as we are aware, we are among the first to compare the predatory abilities of *H. axyridis*, *C. septempunctata*, and *A. bipunctata* and to consider both adult and larval life-stages. Furthering our understanding as to the relative predatory abilities of these three species is important to understand the potential impacts of *H. axyridis* on native ladybird species, through competition and intra-guild predation, and native prey sources (e.g. aphids), through predation. We suggest that the predatory behaviour of *H. axyridis* could, at least in part, be due to their size. Invasive *H. axyridis* and native *C. septempunctata* are both large ladybirds and were found to be generally more similar in their predatory behaviours than the

smaller native *A. bipunctata*, which consumed prey at lower rates than both larger species. Features that facilitate more efficient predatory behaviour are known to scale with predator size. For example, size commonly correlates with greater predator speed, which can increase predator attack rates, and less time spent consuming and digesting prey, which will reduce a predators handling time (Woodward and Warren 2007; Gergs and Ratte 2009). A similar relationship was also noted in larval treatments, with *H. axyridis* and *A. bipunctata* predatory behaviours being significantly different. Metabolic theory, as discussed by Brown et al. (2004), suggests that the energetic demands of an organism is correlated with the organism's mass. While this is in keeping with our findings, no further investigation of the relationship between consumption rate and predator mass was undertaken as while *H. axyridis* and *C. septempunctata* do overlap with respect to their masses, neither overlap with the masses of *A. bipunctata* and this would result in complete separation in statistical analyses.

Pathogenic infection changes predatory ability

For the first time, to our knowledge, we also show the impact of a widespread pathogen on the predatory ability of ladybirds and, specifically, how this impacts the relative predatory abilities of native and invasive ladybirds. *Beauveria bassiana* infection resulted in significant changes in predator forage ability. Invasive alien *H. axyridis* and native *A. bipunctata* adults

showed an increase in attack rate and handling time coefficients when exposed to the pathogen. While an increase in the attack rate coefficient would suggest an increase in prey consumption, the increase in the handling time coefficient would suggest the opposite with the predator spending more time handling and consuming prey individuals. Visual inspection of the functional response relationship shows these coefficients result in a lower overall functional response curve. Contrary to our expectations, native *C. septempunctata* adults showed reduced prey handling times when exposed to the pathogen and no significant change in attack rates. Both larval treatments (*H. axyridis* and *A. bipunctata*) demonstrated significantly reduced attack rate and handling time coefficients when subject to the pathogen treatment. It could have been expected that the ladybird species would have responded differently to *B. bassiana* due to their different evolutionary histories with the pathogen. For example, native ladybird species are likely to have an evolutionary history with *B. bassiana* and are therefore more likely to have behavioural or chemical defences while non-native species are more likely to be naïve to the novel pathogen (Hatcher and Dunn 2011). Evidence for this hypothesis is the fact that *B. bassiana* is a significant cause of *C. septempunctata* overwintering mortality and, while unable to demonstrate such avoidance behaviours within this experiment, *C. septempunctata* are known to avoid *B. bassiana* infected cadavers in the field (Ormond et al. 2011). It is also likely that *B. bassiana* infection could persist within *C. septempunctata* populations and therefore impact behaviours, such a predation, as we have described here. However, *H. axyridis* is known to have a highly efficient immune system, consisting of antimicrobial peptides, which results in *H. axyridis* being less susceptible to many natural enemies including bacteria, fungi, yeasts, and parasitoid flies and wasps (Roy et al. 2008b; Berkvens et al. 2010; Gross et al. 2010; Vilcinskas et al. 2013; Verheggen et al. 2017; Ceryngier et al. 2018; Gegner et al. 2018). Gross et al. (2010) report that the antimicrobial activity of *H. axyridis* hemolymph was approximately one thousand times greater than *C. septempunctata* hemolymph. It is likely that this defence against natural enemies will have facilitated the spread of *H. axyridis*, in addition to other features of the species, such as high predatory ability.

Beauveria bassiana infection resulted in different outcomes for adult and larval life-stages. It is possible that *B. bassiana* infection, through its hyphal growth throughout the host, would impose physiological damage that would impede the predatory ability of individuals, resulting in reduced prey consumption, attack rates and an increase in handling times. However, predatory behaviour may either increase, as the host attempts to mitigate the costs of infection (for example Dick et al. 2010; Bunke et al. 2015), or decrease as the costs of infection rise from either the damage or increased metabolic demand associated with the infection process (for example, MacNeil et al. 2003; Haddaway et al. 2012; Toscano et al. 2014). We suggest that the physiological damage and increased metabolic demand resulted in a decreased ability to consume prey in adult treatments whereas the mechanism driving the observed changes in infected larvae is less clear. We suggest that desiccation or other fungal, viral or bacterial infections could be a contributing factor. Upon infection, *B. bassiana* conidia germinate and penetrate the hosts outer integument before commencing extensive hyphal growth throughout the host's internal cavity. Vey and Jacques (1977) and Poprawski et al. (1999) suggest the repeated penetration of the outer cuticle or soft body of the host can result in an increased risk of desiccation and subsequent infections which would result in additional costs to the host and subsequently affect the host's predatory behaviours. We suggest the larvae are responding to these increased costs, specifically desiccation, by increasing their consumption rates however, further investigation would be required to explicitly establish this relationship.

Implications for invaded communities

In light of our findings, we propose that the invasion of *H. axyridis* is likely to have imposed an increased level of novel predatory pressure on prey species (e.g. aphids) and indirect effects on competitors (e.g. other Coccinellidae). While *B. bassiana* is known to result in approximately 10–15% *C. septempunctata* overwintering mortality (Ormond et al. 2006), instances of infection in *H. axyridis* and *A. bipunctata* are considered rare. We suggest our findings are of interest with respect to current field interactions, for example the predatory differences between the three Coccinellidae, and the impact *B. bassiana* has on the predatory ability

of *C. septempunctata*, but also within a more theoretical framework whereby interactions between *H. axyridis*, *A. bipunctata*, and *B. bassiana* become more frequent than is currently documented. *Harmonia axyridis* is known to be highly abundant and commonly dominates invaded Coccinellidae assemblages (Brown and Roy 2017), it is likely therefore that the *per-capita* differences identified here will scale up and result in larger community impacts in field populations as *H. axyridis*' numerical response to prey density is taken into account (see Dick et al. 2017b). While not quantified here, the numerical response, a measure of how predator abundance changes with respect to changes in prey abundance, is likely to impact native prey in a similar way to the predators *per capita* functional response. Accounting for the demographics of wild populations is essential to further our understanding of any potential impacts posed by a predator and overcome limitations within functional response studies. For example, functional response studies do not commonly account for prey switching which could otherwise be expected in natural communities. However, it has been suggested that generalist predators, such as Coccinellidae, are likely to show less pronounced prey switching than predators foraging on a few prey (Van Leeuwen et al. 2013). Additionally, measures of predatory behaviour attained through functional response studies rely on those focal individuals being representative of the wider predator community. The Coccinellid predators used as part of this study were all of similar ages (fourth instar larvae or recently emerged adults) however, it could be expected that unhealthy or otherwise suboptimal individuals, including those becoming increasingly moribund, could show lower rates of predatory behaviour.

It is likely that *H. axyridis* will impact some species more than others, for example Roy et al. (2012) attribute the decline of native *A. bipunctata* (44% in Britain and 30% in Belgium) to the arrival and subsequent spread of *H. axyridis*. In contrast, *C. septempunctata* populations showed no significant change. Kenis et al. (2017) use a collection of risk measures (for example, the likelihood of encountering *H. axyridis*) to predict the native species most at risk from *H. axyridis*. Native *A. bipunctata* were identified as being at 'very high' risk while native *C. septempunctata* were identified as being at 'medium risk'. We have shown *H. axyridis* to be an efficient predator

which, when compared to additional characteristics of the species, could further our understanding as to how *H. axyridis* impacts native communities. For example, in addition to being a relatively large ladybird (Hodek et al. 2012) and consuming a wide variety of prey species (see Roy and Brown 2015), including engaging in specialised predatory behaviours (e.g. Intra-guild predation; e.g. Pell et al. 2008), we have provided further evidence as to the efficient predatory ability of *H. axyridis*. We also suggest the increased predatory behaviour exhibited by *H. axyridis* could have facilitated the species' initial spread and continued success throughout its invaded range.

Concluding remarks

We have provided evidence that pathogenic infection impacted the predatory behaviour of ladybirds in a species and life-stage specific way. Despite being known to mediate invasion success and impact through their lethal and sub-lethal effects (Dunn et al. 2012; Strauss et al. 2012), the effects of parasites and pathogens are rarely accounted for within invasion ecology regardless of literature showing that their impacts can vary. For example, invasive *Gammarus pulex* harbouring acanthocephalan infection show increased intake of prey (Dick et al. 2010). Conversely, infection has also been shown to result in reduced consumption rates (Wright et al. 2006; Toscano et al. 2014). While instances of infection by native parasites and pathogens may be low in *H. axyridis*, due to its efficient immune system, understanding how infection can modify the key functional behaviours of this widespread invasive alien species is key to furthering our understanding of the effects of infection on the success and impacts of invasion events (Brook et al. 2008; Strayer 2010). Here we demonstrate that pathogenic infection affects a key functional trait, predation, of *H. axyridis* and two native species (*A. bipunctata* and *C. septempunctata*).

We have shown that the invasive alien *H. axyridis* displays significantly more efficient predatory behaviour than two native predators in both adult and larval life-stages. Pathogenic infection significantly changed the foraging ability of ladybird predators in a species and life-stage specific way but resulted in no measurable change in overall prey consumption. We suggest the impacts of *H. axyridis* are at least partially

explained by the more efficient predatory behaviour detailed here.

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Compliance with ethical standards

Author contributions WF, AD, LB, HH, and HR designed the experiment. WF and HH collected the data. WF analysed the data and wrote the manuscript. All authors critiqued the manuscript for intellectual content.

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