Parasite and predator risk assessment: nuanced use of olfactory cues

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Foraging herbivores face twin threats of predation and parasite infection, but the risk of predation has received much more attention. We evaluated, experimentally, the role of olfactory cues in predator and parasite risk assessment on the foraging behaviour of a population of marked, free-ranging, red-necked wallabies (Macropus rufogriseus). The wallabies adjusted their behaviour according to these olfactory cues. They foraged less, were more vigilant and spent less time at feeders placed in the vicinity of faeces from dogs that had consumed wallaby or kangaroo meat compared with that of dogs feeding on sheep, rabbit or possum meat. Wallabies also showed a species-specific faecal aversion by consuming less food from feeders contaminated with wallaby faeces compared with sympatric kangaroo faeces, whose gastrointestinal parasite fauna differs from that of the wallabies. Combining both parasite and predation cues in a single field experiment revealed that these risks had an additive effect, rather than the wallabies compromising their response to one risk at the expense of the other.

1. Introduction

It is widely understood that foraging animals trade-off the risk of predation against the risk of insufficient resources for maintenance and reproduction, typically by increasing their investment in predator vigilance and/or avoiding feeding in locations where they are vulnerable to attack [1,2]. Less well appreciated is whether animals also incorporate the risk of parasite infection into such foraging trade-offs [1], even though parasitism presents significant challenges to host fitness through negative impacts on growth, reproductive success or survival [3,4]. While several studies have investigated the combined effects of parasites and predators on fitness [5,6] and general behaviours, such as grooming [7,8], the combined effect on foraging behaviour is largely unknown. Yet foraging animals may often face these risks in combination, especially in times of peak parasite activity.

Accurate risk assessment of both predation and parasite infection requires a nuanced recognition of appropriate cues. The faeces of carnivorous vertebrates contain odour cues that are frequently used by herbivores to assess the risk of predation [9–13]. While the precise nature of these cues is probably complex, there is evidence that predator diet affects these volatile compounds, allowing prey to perceive differences between predator species, and even between individuals of the same species that have fed on different prey [14–16]. Similarly, many grazing herbivores avoid areas adjacent to faeces to reduce the risks of parasite transmission [1,17–20]. Herbivores infected with gastrointestinal parasites produce faeces containing parasite eggs that hatch into infective larvae on the sward, where the larvae may be ingested by other herbivores [4,21]. The parasite larvae are probably undetectable by the herbivore, so faeces are used as a cue for infection risk [20]. While a generalized aversion to all faeces is advantageous for herbivores foraging in the same areas as conspecifics, it may result in less efficient foraging if heterospecifics share foraging grounds but not parasites [22,23].

Here, we experimentally test the responses of free-ranging, red-necked wallabies (Macropus rufogriseus) to cues associated with the risks of predators, parasites and both in combination. Macropodid marsupials (kangaroos and wallabies) have been exposed to predation from past and present predators [24] and are typically wary of predator faeces [25,26]. Macropodids also have
diverse and highly species-specific gastrointestinal parasite faunas [27] and often share foraging grounds (e.g. [28]). We asked the following questions: (i) do red-necked wallabies vary their response to predator olfactory cues according to variation in the predator’s diet; (ii) is faecal aversion in red-necked wallabies host-species-specific; and most importantly: (iii) how do the combined risks of predators and parasites impact on the foraging decisions of red-necked wallabies?

2. Material and methods

(a) Study site

We conducted this study at the Victoria Valley Airbase (Grampians National Park, Australia; 37°11’S, 142°20’E), which comprises two 80 × 800 m runways of mown native grasses and heath. Red-necked wallabies and grey kangaroos (Macropus giganteus and Macropus fuliginosus) are common, and they forage and defecate throughout the runways. Dingoes (Canis lupus dingo) preyed on wallabies in the past [24], but have been extirpated from the Grampians region; the main extant predator is the red fox (Vulpes vulpes). In the Grampians region, red-necked wallabies have up to 22 conspecifics (Bennett’s wallaby, Trichosurus vulpecula), 11 congenerics (a mix of kangaroos, Macropus rufus, M. giganteus and M. fuliginosus), 14 larger non-native eutherians (sheep, Ovis aries); 11 smaller, native marsupials (common brush-tail possum, Trichosurus vulpecula); and 11 smaller, non-native eutherians (European rabbit, Oryctolagus cuniculus). The meat was obtained from Yarra Valley Game Meats (Healesville, Victoria, Australia). We fed four adult dogs, of varying breed, sex and size, with meat of one of the five prey types for 48 h to eliminate traces of previous diets, and then collected faeces produced while on that diet. We collected the faeces immediately on deposition and immediately froze them in airtight containers. Containers were later pooled into one set of experimental faeces for each diet.

We prepared feeders by placing 200 g of crushed maize and 100 g of an inedible matrix (eucalypt woodchips) in clear plastic freezer trays (36 × 25 cm) placed on the open runways. We placed four clear plastic lids, each with approximately 50 g of defrosted experimental predator faeces, 30 cm from the feeder in the four cardinal directions. Procedural controls had rocks on the plastic lids to imitate faeces. Each trial involved a randomly selected type of faeces or control; the observer was always blind to the type of faeces.

We observed wallabies using a Kowa spotting scope (20 × 60) from a distance of less than or equal to 40 m in a concealed position, first recording the distance between the feeder and the nearest cover, the weather conditions and the time of day. When a wallaby entered a 3 m radius around the feeder, we recorded its identity and time to subsequently arrive at the feeder. Once feeding, we noted the number of times it exhibited vigilance bouts and amount of food eaten, with parasite type (kangaroo or wallaby), prey type (kangaroo, wallaby or possum), distance to nearest cover, time of day (morning or evening) and their interactions as fixed effects, and animal ID as a random factor. The approach time variation in foraging behaviours (approach time, number of vigilance bouts and amount of food eaten), with treatment (faecal source), distance to nearest cover, time of day (morning or evening) and their interactions as fixed effects, and animal identification (ID) as a random factor to account for multiple tests of the same individual. For the herbivore faecal aversion experiment, we used a paired t-test to compare the amount of food consumed from feeders contaminated with the faeces of either kangaroos or wallabies. In the combined faecal aversion experiment, we used REML to explore the sources of variation in foraging behaviours (approach time, number of vigilance bouts and amount of food eaten), with parasite type (kangaroo or wallaby), prey type (kangaroo, wallaby or possum), distance to nearest cover, time of day and the interaction terms as main effects, and animal ID as a random factor. The approach time and number of vigilance bouts were log transformed to improve the distribution. In all analyses we initially included the full model, but we report (see the electronic supplementary material) reduced models in which we sequentially removed interaction terms where they were not significant (p > 0.1).

(c) Herbivore faecal aversion

We confirmed that the wallabies exhibited herbivore faecal aversion by presenting animals with paired feeders (as described below) containing 200 g of crushed maize contaminated with 100 g of mixed macropod faeces. Pilot studies (72 replicates from 14 individuals) indicated that wallabies always chose uncontaminated trays when presented with a choice, so we did not include uncontaminated trays in subsequent experiments.

We conducted faeces, immediately after observing deposition, from animals foraging around the airbase, separating them by species. Each trial comprised a pair of adjacent feeders: one contaminated with 100 g of faeces of red-necked wallabies (con-specific feeder) and the other with 100 g of mixed faeces of eastern and western grey kangaroos (heterospecific feeder). As before, we observed wallabies from a concealed position. We recorded the identity of wallabies that approached and fed at the feeders, together with their choice of feeder and any switching between feeders. Observations lasted until the animal left the feeder or 1 h had elapsed, when we collected and weighed the feeders to determine the amount of food consumed. We discarded any remaining food and faeces as before. We only included in our analysis trials where the wallaby sniffed both feeders, the animal contributed to fewer than five trials, and rain did not fall before we collected the feeder. In total, 19 individuals contributed to 46 trials.

(e) Statistical analysis

We analysed data using JMP v. 10 (SAS institute, USA). For the predator faecal aversion experiment, we used restricted maximum-likelihood (REML) models to determine the source of variation in foraging behaviours (approach time, number of vigilance bouts and amount of food eaten), with treatment (faecal source), distance to nearest cover, time of day (morning or evening) and their interactions as fixed effects, and animal identification (ID) as a random factor to account for multiple tests of the same individual. For the herbivore faecal aversion experiment, we used a paired t-test to compare the amount of food consumed from feeders contaminated with the faeces of either kangaroos or wallabies. In the combined faecal aversion experiment, we used REML to explore the sources of variation in foraging behaviours (approach time, number of vigilance bouts and amount of food eaten), with parasite type (kangaroo or wallaby), prey type (kangaroo, wallaby or possum), distance to nearest cover, time of day and the interaction terms as main effects, and animal ID as a random factor. The approach time and number of vigilance bouts were log transformed to improve the distribution. In all analyses we initially included the full model, but we report (see the electronic supplementary material) reduced models in which we sequentially removed interaction terms where they were not significant (p > 0.1).
3. Results

In the predator faecal aversion experiment, wallabies adjusted their foraging behaviour according to type of prey consumed. The wallabies took significantly longer ($p < 0.05$) to approach feeders surrounded with faeces of dogs fed on wallaby and kangaroo compared with the other prey types, were more vigilant (figure 1a), and consumed less food at these feeders (figure 1b and electronic supplementary material, table S3).

In the herbivore faecal aversion experiment, wallabies typically approached the feeders, sniffed each, then began to feed from one of them, which we deemed their ‘choice’. Once the wallabies had made a choice, they switched feeders on only six (13%) occasions. The wallabies consumed five times more from heterospecific feeders (70.3 ± 7.1 g) than from conspecific feeders (13.8 ± 3.9 g; $t_{45} = 6.4$, $p < 0.0001$).

In the combined faecal aversion experiment, the time taken to approach the feeder was influenced by predator cues, but not parasite risk (figure 2a). The time spent vigilant was influenced by both predator and parasite cues (figure 2b), and an interaction term including predator cues, parasite cues and distance from cover (electronic supplementary material, table S3). The greater time spent vigilant in the wallaby faecal treatment may reflect a general unwillingness to feed, since there was no choice in this experiment. The amount of food consumed was similarly affected by cues indicating the risk of both predators and parasites (figure 2c), and the distance from cover (electronic supplementary material, table S3). The predator cue × parasite cue interaction term was not significant ($F_{2,52.7} = 0.022$, $p = 0.98$).

4. Discussion

Red-necked wallabies make impressive use of olfactory cues to gauge the risk of predation and parasite infection. The wallabies approached feeders surrounded by the faeces of dogs that had fed on macropodids more warily than if the faeces came from dogs that had fed on other herbivores. Further, herbivore faecal aversion in red-necked wallabies is host species-specific; wallabies ate five times more from feeders contaminated by heterospecific than conspecific faeces. Finally, the wallabies’ food consumption decreased additively when faced with the combined faecal cues for the risk of predation and parasite infection. While herbivore faeces may include social information (including identity, gender, dominance), this is unlikely to explain our results because the faeces included in each trial derive from numerous individuals, and therefore provide a non-specific cocktail of odours. We also think it
unlikely that red-necked wallabies use conspecific faeces as a cue to prior foraging by conspecifics, thereby improving foraging efficiency by faecal avoidance. First, herbivores select foliage by direct visual and olfactory assessment, which provides more precise cues of forage quality and quantity. Second, field experiments indicate that congers will not trade-off their aversion to faecal contamination for a gain in foraging efficiency [20].

The precise chemical nature of the olfactory cues present in carnivore faeces is not known in this system, but they are clearly influenced by diet [34–36]. Such effects are also common in other chemical sensory systems [37]. The results of our field experiments caution against inferring that prey respond to different predator species per se, because the predators may also have different diets. Red-necked wallabies most strongly avoided faeces produced from diets that contained either conspecifics or closely related macropodids, rather than faeces from diets of larger rather than smaller prey, or from faeces containing other marsupial rather than eutherian material. Perhaps, the responses of red-necked wallabies are learnt, with the response reflecting the frequency of encounter [38,39].

Our results suggest that herbivore faecal aversion as a means of minimizing gastrointestinal parasite infection levels [18–20] may be more nuanced than previously thought. Although wallabies fed more from uncontaminated than contaminated food, their aversion was stronger if the food was contaminated by the faeces of conspecifics rather than heterospecifics. This finding suggests strong selection for parasite avoidance on the one hand, and efficient foraging on the other. Our results contrast with findings that reindeer (Rangifer tarandus) do not avoid sheep faeces less than conspecific faeces [40], perhaps reflecting relaxed selection either because these two species share a sufficiently broad gastrointestinal parasite fauna, or there has been little history of the two species foraging in the same location.

Studies report both additive and interactive effects of parasites and predators on fitness [5,6]. The wallabies might be expected to avoid feeders associated with predator cues, regardless of whether they had greater risk of parasite infection, if the more immediate threat of predation outweighed that of parasitism. This was not the case. Rather, avoidance behaviour was most evident at feeders that were associated with the greatest combined danger of predators and parasites. This suggests that while the fitness consequences of predation and parasites are often different, wallabies do not compromise their response to the risks of either predators or parasites in the face of the other. More complete analyses of foraging behaviour should consider these risks simultaneously: foraging herbivores are likely to trade-off foraging efficiency against avoiding both predators and parasites.

Figure 2. The combined effects of predator risk (the presence of dog faeces derived from different diets) and parasite risk (presence of conspecific and heterospecific macropod faeces) on: (a) the approach time (s); (b) the number of vigilance bouts; and (c) the amount eaten (g) by red-necked wallabies. Values are means with standard error bars. Connecting lines above the bars indicate non-significant (p > 0.05) differences between predator risk treatments, and an asterisk indicates significant differences between parasite risk treatments within predator risk treatments. (Online version in colour.)

References


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