The interaction of antlions and ants is postulated to be a predator-prey interaction in which the involved parties coevolve. Here, we investigated two issues of potential significance in terms of antlions and ants imposing selective pressures on one another. First, we determined whether trap-building antlions and sand-dwelling ants closely co-occurred in an area inhabited by both. In the field, we found that ants were the main potential prey items in artificial traps placed inside aggregation zones of antlions and that *Formica cinerea* workers comprised the majority of these ants. Second, we checked whether rescue behavior, a type of prosocial behavior displayed by *F. cinerea* workers and performed towards nestmates captured by antlions, reduced the hunting success of the latter. In the laboratory, we found that rescue attempts were very rarely successful. Overall, caution must be used when considering the coevolution of antlions and ants. Clearly, even though these two organisms can closely co-occur, the rescue behavior of ants seems to be unrelated to the predatory threat from antlions.

**Keywords** Antlions · Ants · *Euroleon* · *Formica* · *Myrmeleon* · Predator-prey interaction

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**Introduction**

Predator-prey interactions are considered to be among the main reasons for coevolution, i.e., a phenomenon in which organisms exert selective pressures on one another (Janzen 1980). Indeed, from the perspective of the prey community, it is important to develop and improve the abilities to survive and increase fitness (Moore and Biewener 2015). Avoiding death by predation can be accomplished in various ways, sometimes quite spectacularly. For example, a moth prey of the genus *Bertholdia* broadened its defensive repertoire over the course of evolution by developing ultrasonic clicks that jam sonars used by their bat predators in the genus *Eptesicus* (Corcoran et al. 2009).

Some antlions (Neuroptera: Myrmeleontidae) are “sit and wait” predators (Perry and Pianka 1997) with larvae that use traps while feeding under natural conditions. Trap building has evolved rarely in the animal kingdom (Ruxton and Hansell 2009). The uniqueness of this foraging strategy has long attracted attention to antlions, which possess several traits that make them seemingly efficient predators (Miler et al. 2017a). These traits include strong mandibles used for injecting paralytics (Matsuda et al. 1995; Nishiwaki et al. 2007), high vibration sensitivity that helps them detect approaching prey (Devetak et al. 2007; Fertin and Casas 2007) and learning abilities that allow them to improve foraging strategies (Kuszewska et al. 2016; Miler et al. 2017b). However, the diet of antlions is not well known, mostly mentioned anecdotally in studies on other questions (Heinrich and Heinrich 1984; Matsura 1986, 1987;
Cylindrical plastic vessels (3 cm in diameter, 11 cm in length) were used as artificial traps, covered on the inside with Fluon (Sigma-Aldrich, Germany). These vessels were put in the sand, 5 within the *M. bore* antlion zone and 5 within the *E. nostras* antlion zone, collected once per week and replaced by fresh ones from May to September. The vessels were buried at 30 cm intervals and were not baited. Collected vessels were closed with a cap and dried, and then their contents were examined. If arthropods were present, they were identified to the family level. If identification was not possible, the arthropods were listed together as “unidentifiable”. The data were pooled into four-week periods. For each period, the percentage of Formicidae among all identified potential prey was calculated separately for each species of antlion. Additionally, the percentage of *F. cinerea* individuals among Formicidae was calculated for each species of antlion.

Tests of rescue behavior were performed in the laboratory on animals collected in the field at the same location. We collected 50 *M. bore* and 50 *E. nostras* antlion larvae. Approximately 500 *F. cinerea* foragers from each of two colonies were also collected. The taxonomic identity of animals was confirmed in the laboratory, following appropriate keys (Czechowski et al. 2012; Badano and Pantaleoni 2014). Ants were kept in two separate plastic boxes (25 × 17 × 10 cm each), with walls covered with Fluon (Sigma-Aldrich, Germany), and food and water were provided ad libitum. All larvae were placed in paper test boxes (a separate box for each larva; 13 × 13 × 4.5 cm) filled with dry, sieved, fine-grained sand collected from the original habitat. Each larva was placed in the middle of its box, and then a plastic ring (7 cm in diameter) was placed on top of the sand to force the larva to build its trap within. This ring was removed after approximately 20 h, and tests started shortly thereafter. For the tests, a plastic ring (10 cm in diameter) covered in Fluon (Sigma-Aldrich, Germany) was placed in the box with an antlion equipped with a functional trap. A group of six ants was introduced into the ring, on the flattened surface around the antlion trap. We noted for 3 min whether any of these ants were captured, and if so, then we prolonged the observational period for 3 min since capture. During that time, we noted whether any signs of rescue behavior occurred in any of the remaining ants (0/1 response), measured the latency to the first episode of rescue (if one occurred), and whether rescue was successful. This type of testing can be considered ecologically relevant based

### Methods

The potential prey of larvae was assessed in the field near Klucze (southern Poland, Błędowska Desert). Błędowska Desert is inhabited by at least two species of antlions, *M. bore* and *E. nostras*, with former being abundant and occurring in open space exposed to direct sunlight, with only lower instars sometimes seeking protection from wind and rain under hanging branches, and the latter always occurring in isolated patches highly protected from wind, rain and direct sunlight. *F. cinerea* sand-dwelling ants are common throughout the area.
on previous field observations (Czechowski et al. 2002). We used pulling at the limbs/antennae/mandibles of the captured individual, digging around it, removal of the sand covering it, and direct attack on the antlion mandibles as signs of rescue behavior (Miler 2016). For the antlions that captured prey, we compared the number of tests with rescue between the two antlion species using Fisher’s exact test (FET) in STATISTICA 13 software (Tibco, USA). Then, the median latency to the first episode of rescue in all tests in which rescue occurred was calculated. Data were not further analyzed due to a lack of successful rescues (see below).

Results

Arthropods captured in artificial traps were identified to the family level (Table 1). During the entire period of the survey, Formicidae comprised a gross majority of the potential prey for M. bore (~ 98%). In the case of E. nostras, Formicidae comprised the majority in July and August (~ 73–99%) but not during the remaining months (~ 29–50%). Importantly, F. cinerea workers comprised approximately 99 and 97% of the total captured Formicidae in M. bore and E. nostras aggregation zones, respectively.

Among the 50 M. bore and 50 E. nostras antlion larvae collected, 47 of the former and 33 of the latter constructed functional traps under laboratory conditions. Only these individuals participated in rescue behavior tests. Ants were captured by either antlion species in the case of 48 tests in total. Rescue attempts occurred in 28 of these 48 tests (58%), and there was no difference in the proportion of tests with rescue attempts between antlion species (16 of 28 in M. bore and 12 of 20 in E. nostras; FET yielded $p = 1.000$). The first episodes of rescue behavior occurred after a median latency ± SD of 42 ± 31 s. No rescue attempt was successful. In 3 tests, the captured individual was released dead, and another ant was captured in its place.

Discussion

We show here that ants, especially F. cinerea workers, were generally abundant in the artificial traps placed in the antlion-inhabited areas throughout the season. Interestingly, the largest numbers of these ants coincided with the highest point of their colony development (Czechowski et al. 2002) as well as the highest ambient temperatures, favoring the physiological capacity of trap building in antlions (Antol et al. 2018). Even if our data overestimated what antlions can truly catch as prey and only a subset of the potential prey presented herein were within their reach (Humeau et al. 2015), it is highly likely that ants are nevertheless the main prey of antlions, as is often assumed.

Earlier observations of Tetramorium ants facing unknown species of antlions indicated that these ants rarely rescue their nestmates successfully (Taylor et al. 2013). The present results also suggest that rescue behavior is without ecological relevance in the context of antlion larva capture. This conclusion is supported by a very narrow time frame available for rescue in this context. Antlions inject paralytics into their captured prey; these paralytics cause paralysis in ants at a maximum of 2 min after injection (Griffiths 1980; Matsuda et al. 1995; Nishiwaki et al. 2007). Taking into account the fact that the captured ant has to “call for help” to elicit rescue in its nestmates (Czechowski et al. 2002; Frank et al. 2017; Miler and Kuszewska 2017), these 2 min leave little room for a successful rescue action, which is expressed after a delay (here, median latency ± SD: 42 ± 31 s). It seems plausible that rescue behavioral patterns in F. cinerea, and possibly other sand-dwelling ants, are more of an artefact in the context of antlion larva capture (stemming, e.g., from ants’ inability to differentiate between entrapment contexts). It is worth noting that F. polycycta ants, forest groundcover species that never meet antlions in nature but probably face other entrapment situations, show rescue behaviors in an antlion larva capture context (Miler et al. 2017c), supporting the idea that ants do not differentiate between entrapment contexts. Nest cave-ins are likely much more important than antlion encounters for the evolution of rescue behavioral patterns in F. cinerea species, as well as in other sand-dwelling ants. Such a conclusion, however, would need additional support, preferably from studies conducted under field conditions. Such studies could additionally aim to estimate the predation rate of ants by antlions since, if low, this rate could provide another evidence that rescue attempts are artefacts in the context of antlion larva capture.

Naturally, the present results do not mean that sand-dwelling ants are helpless in terms of antlion larva capture. Ants can minimize the risk of predation by avoiding antlion aggregation zones to some extent (Gotelli 1996; Morrison 2004; Hollis et al. 2017), which
### Table 1: The potential prey of *M. bore* and *E. nostras* throughout the season. Data show arthropods captured in artificial pits placed inside antlion zones of both antlion species. Unidentifiable specimens were degraded and mostly comprised of Diptera.

| Dates of collection (2018) | Captured arthropods in *M. bore* (in count order) | Captured arthropods in *E. nostras* (in count order) |
|---------------------------|---------------------------------------------------|---------------------------------------------------|
|                           | family | count | family | count |
| **05.05–26.05**           | Formicidae | 326   | Formicidae | 6   |
|                           | Carabidae | 3     | Curculionidae | 4   |
|                           | Curculionidae | 3   | Carabidae | 3   |
|                           | Acrididae | 1     | Staphylinidae | 3   |
|                           | Cleridae | 1     | Myrmeleontidae | 2   |
|                           | Elateridae | 1    | Nitidulidae | 2   |
|                           | Unidentifiable | 1    | Scarabaeidae | 1   |
| **02.06–23.06**           | Formicidae | 192   | Formicidae | 9   |
|                           | Carabidae | 2     | Myrmeleontidae | 8   |
|                           | Staphylinidae | 1  | Curculionidae | 6   |
|                           | Unidentifiable | 2    | Aphididae | 2   |
| **30.06–21.07**           | Formicidae | 357   | Formicidae | 604  |
|                           | Carabidae | 3     | Curculionidae | 5   |
|                           | Staphylinidae | 2  | Myrmeleontidae | 2   |
|                           | Acrididae | 1     | Cerambycidae | 1   |
|                           | Stratiomyidae | 1 | Chrysomelidae | 1   |
|                           | Unidentifiable | 14   | Unidentifiable | 3   |
| **28.07–18.08**           | Formicidae | 472   | Formicidae | 36   |
|                           | Byrrhidae | 3     | Myrmeleontidae | 2   |
|                           | Carabidae | 3     | Curculionidae | 3   |
|                           | Sphecidae | 2     | Carabidae | 2   |
|                           | Chrysomelidae | 1 | Chrysomelidae | 2   |
|                           | Curculionidae | 1 | Aradidae | 1   |
|                           | Unidentifiable | 4    | Cerambycidae | 1   |
|                           |           |       | Chrysomelidae | 1   |
|                           |           |       | Vespidae | 1   |
|                           |           |       | Unidentifiable | 1   |
| **25.08–15.09**           | Formicidae | 89    | Formicidae | 6   |
|                           | Carabidae | 2     | Myrmeleontidae | 4   |
|                           | Geotrupidae | 1    | Carabidae | 1   |
|                           | Unidentifiable | 3    | Throscidae | 1   |
|                           |           |       | Unidentifiable | 1   |
is their first “line of defense” against ant lions. Additionally, earlier results obtained in rescue behavior investigations utilizing ant lions (see, e.g., Miler 2016) are not invalidated by the present findings since rescue is a general behavioral category and major factors contributing to its occurrence (e.g., life expectancy) may very well be highly context independent.

In summary, it is tempting to look for evidence of coevolution between ant lions and ants (Hollis et al. 2015; Hollis 2017). As we show here with *F. cinerea* workers, sand-dwelling ants may closely co-occur with trap-building ant lions. However, it seems that the rescue actions of these ants are unlikely displayed as a countermeasure to strong selection pressure from ant lions. The question of antlion and ant mutual dependence remains open, however, even in light of the present results. An important further step in investigating this issue would be to conduct a broader-scale study to analyze the geographical distribution of both taxa with a focus on their degree of overlap.

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