

# Predation by beetles (Carabidae, Staphylinidae) on eggs and juveniles of the Iberian slug *Arion lusitanicus* in the laboratory

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## Abstract

*Arion lusitanicus* has become a major pest species in western Norway in the last few years. This species originates from southern Europe but has been spread by humans over large parts of central and northern Europe during recent decades. Slugs have traditionally been controlled by the use of molluscicides; but, as these may have serious ecological side effects, biological control of slugs is highly desirable. Potential biological control agents include nematodes, gastropods and arthropods. In laboratory experiments, we tested whether five common predator beetles would feed on eggs and juveniles of *A. lusitanicus*. The species *Carabus nemoralis*, *Nebria brevicollis*, *Pterostichus melanarius* and *Pterostichus niger* (Carabidae) as well as *Staphylinus erythropterus* (Staphylinidae) were tested, of which only *P. melanarius* has been tested on *A. lusitanicus* previously. *Nebria brevicollis* did not feed on slug eggs or newly hatched slugs, but the remaining four species all killed and ate a large proportion of the eggs and hatchlings offered. Both *P. melanarius* and *P. niger* also destroyed *A. lusitanicus* eggs and hatchlings under conditions emulating those in the field. Prey size choice experiments were conducted by feeding *C. nemoralis*, *P. niger* and *S. erythropterus* on different sizes of *A. lusitanicus*. *Carabus nemoralis* was also given a choice between two slug species, *A. lusitanicus* and *Deroceras reticulatum*. A significant preference for slugs smaller than one gram was evident for *C. nemoralis*, while the other beetles struggled much more to overcome the mucus of juvenile slugs. No significant preference was found between *A. lusitanicus* and *D. reticulatum* as prey for *C. nemoralis*. We also discuss the feasibility of biological control of *A. lusitanicus* using beetle predators.

**Keywords:** *Arion*, *Deroceras*, biological control, Carabidae, invasive species, predation, Staphylinidae

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## Introduction

Many animal species have been spread by international trade in the last few centuries. An increasing exchange of soil and plant material over large distances has led to the introduction of alien species into areas where they have become pests (Keller *et al.*, 1999). In particular, alien slugs (Gastropoda) cause increasing problems in horticulture and agriculture (South, 1992). Slugs become pest species mainly in areas where precipitation in spring is high (Keller *et al.*, 1999; Hommay, 2002), rarely causing any problems in areas with dry spring weather. Gastropods may affect the species composition of natural plant communities as well as seriously reducing the yields and devaluing a wide range of crops by feeding damage (Port & Port, 1986; South, 1992; Barker, 2002). Another reason for concern is that introduced gastropods may endanger native species (Essl & Rabitsch, 2002).

The invasive form of the slug *Arion lusitanicus* Mabilbe 1868 (also regarded as *A. vulgaris* Moquin-Tandon 1855; Anderson, 2005) (Pulmonata: Arionidae) originates from the Iberian Peninsula and southern France but has been introduced into central and northern Europe, where it has become a major pest in the last 30 years (von Proschwitz, 1996; Dolmen & Winge, 1997; Grimm *et al.*, 2000). The slug was first recorded in Norway in 1988 (von Proschwitz & Winge, 1994) and has subsequently spread along much of the coast, reaching high densities locally (Dirks, I., Tomasgård, T.E.H., do Amaral, M.J.A., Solhøy, T., Skartveit, J. & Mong, C., unpublished data) and causing considerable damage to garden plants and strawberry cultivation.

Slug control in agriculture and horticulture is mainly through the application of pesticides (South, 1992; Barker, 2002; Iglesias *et al.*, 2002), but the effect of molluscicides is often variable and short-lived, and they may affect non-target organisms negatively, including natural enemies of slugs and other pests (Langan *et al.*, 2004). Integrated management of slugs, therefore, is highly desirable, including maximising regulation by natural enemies.

Promising candidates for natural biological control of the Iberian slug include carabid beetles (Symondson, 2004). However, adult specimens of the Iberian slug secrete a thick covering of sticky mucus when irritated, which may render them immune to attack from many predators. Further, the Iberian slug has been shown to produce a defensive compound (Schroeder *et al.*, 1999), which may deter some predators. However, it is probable that eggs and newly hatched slugs are more susceptible to predation (Paill, 2000) and, thus, to biological control.

Ground beetles (Coleoptera, Carabidae) are common predators that feed on a number of invertebrate groups (Sunderland, 2002; Symondson, 2002; Toft & Bilde, 2002). Within carabids, there are a few mollusc specialists (notably *Cychrus caraboides* (L.)) and also species (e.g. *Carabus* spp.) which are predominantly mollusc feeders (Toft & Bilde, 2002). In addition, many generalist species also include molluscs as part of their diet (Thiele, 1977; Sunderland, 2002). Field studies have demonstrated that ground beetles, notably *Pterostichus melanarius* (Illiger, 1798), can effectively reduce gastropod populations and damage to crops (Asteraki, 1993; Bohan *et al.*, 2000; Symondson *et al.*, 2002; McKemey *et al.*, 2003; Oberholzer *et al.*, 2003; Symondson, 2004). The present study investigated whether some of the common beetle species would eat eggs, hatchlings and

juveniles of *A. lusitanicus* under laboratory conditions. It is necessary to know which species are able to kill and consume slugs of the different size classes found in the field before doing larger experiments (Kaiser *et al.*, 1993), as well as developing molecular methods for detection of predation in the field (Harper *et al.*, 2005; Thomas *et al.*, 2009).

Four widespread predatory carabid species were tested and compared: *Carabus nemoralis*, *Nebria brevicollis*, *Pterostichus niger* and *P. melanarius*, as well as the ubiquitous staphylinid *Staphylinus erythropterus*. The beetles were selected on the basis of being relatively large and common predators in gardens and arable fields. The *Carabus* species are generally thought to feed on soft-bodied prey, such as earthworms and gastropods (Hengeveld, 1980a,b; Luff, 2007), preventing the latter from producing large amounts of mucus by killing the slugs efficiently (Pakarinen, 1994). *Carabus violaceus* has been found to prey mainly on *A. lusitanicus* when juvenile slugs were available (Paill, 2000), but this species mostly occurs in forests in Scandinavia rather than open fields (Lindroth, 1985). By contrast, *C. nemoralis* is common in anthropogenically altered habitats (e.g. parks, gardens) and has been found to prey on molluscs (Tod, 1973), including the slug *Arion ater* (Poulin & O'Neil, 1969). *Nebria brevicollis*, on the other hand, is regarded as mainly a springtail predator (Thiele, 1977), but it has also been shown to prey on molluscs (Tod, 1973), including juvenile *Deroceras reticulatum* (Ayre, 2001). *Pterostichus melanarius* has, in particular, been found to be an effective slug predator both in laboratory experiments (McKemey *et al.*, 2001; Oberholzer & Frank, 2003) and under field conditions (Symondson, 1993, 2004; Symondson *et al.*, 1996, 2002; McKemey *et al.*, 2003) and has been found to feed on *A. lusitanicus* in the field (Paill, 2004). In addition, *P. niger* has been found to prey on gastropods (Tod, 1973; Pakarinen, 1994). The food preferences of the large staphylinid beetle *S. erythropterus* are less known, but it is regarded as a generalist predator (V. Gusarov, Natural History Museum in Oslo, personal communication).

In the present work, we wanted to test the hypothesis that all the selected beetles, *C. nemoralis*, *P. niger*, *P. melanarius* and *S. erythropterus*, consume eggs and newly hatched slugs. Further, we aimed to test the hypothesis that *C. nemoralis* prefer *D. reticulatum* over *A. lusitanicus*, based on the much more sticky mucus of arionid slugs compared to *D. reticulatum* (Pakarinen, 1994). In addition, we wanted to test the hypothesis that beetles prefer smaller slugs over larger ones also due to less mucus production. The terminology for biological control we use follows Eilenberg *et al.* (2001).

## Materials and methods

### Study sites

All slugs and beetles studied were collected in the vicinity of Bergen, western Norway. Beetles and slugs were sampled separately for the different experiments in 2003, 2004, 2006 and 2007. Thus, the beetles used had different ages and feeding histories, but we attempted to mitigate any problem that this could cause by using beetles and slugs from the same sampling site and time period, and made sufficient replicates to control for these factors when testing their ability to consume slugs. The area has an oceanic climate with high precipitation throughout the year, mild

Table 1. Number of predator-prey combinations in the Petri-dish experiments with eggs and newly hatched slugs. Each Petri dish contained one beetle and the number of prey offered to each beetle are presented in the columns 'eggs offered' and 'hatchlings offered'.

	Individuals/ replicates	Treatment	Eggs offered	Hatchlings offered
<i>Carabus nemoralis</i>	9	Eggs only	10	0
	1	Eggs only	15	0
	1	Hatchlings only	0	10
	6	Eggs & hatchlings	5	5
	2	Eggs & hatchlings	10	5
	2	Eggs & hatchlings	10	3
<i>Pterostichus melanarius</i>	3	Eggs only	10	0
	3	Eggs only	15	0
	4	Hatchlings only	0	10
	2	Eggs & hatchlings	10	10
<i>Pterostichus niger</i>	2	Eggs only	10	0
	5	Eggs only	15	0
	5	Hatchlings only	0	10
	1	Eggs & hatchlings	5	5
	1	Eggs & hatchlings	10	5
	4	Eggs & hatchlings	10	10
<i>Nebria brevicollis</i>	5	Eggs only	5	0
	14	Eggs only	10	0
	3	Eggs only	15	0
	2	Eggs & hatchlings	5	5
	2	Eggs & hatchlings	10	3
	1	Eggs & hatchlings	10	5
<i>Staphylinus erythropterus</i>	2	Eggs only	10	0
	1	Eggs only	15	0
	2	Hatchlings only	0	10
	1	Eggs & hatchlings	5	3
	1	Eggs & hatchlings	5	5
	3	Eggs & hatchlings	10	5
	1	Eggs & hatchlings	10	10
	1	Eggs & hatchlings	15	3

winters and relatively cool summers. Average annual precipitation in Bergen is 2250 mm, and monthly mean temperatures are 1.3°C in January (coldest month) and 14.6°C in July (warmest month) (data from Norwegian Meteorological Institute, 2002).

#### Feeding trials

The beetles, *C. nemoralis*, *N. brevicollis*, *P. niger*, *P. melanarius* and *S. erythropterus*, were kept at 4°C in plastic boxes with mosses, which gave cover as well as helping to prevent desiccation. The beetles were fed earthworms (Lumbricidae) once a week. Prior to each experiment, the beetles were starved. In 2003, they were starved for nine days at 4°C following McKemey *et al.* (2003); while, in 2004 and in the size choice experiment in 2006, they were starved for two days at room temperature (19°C) following Oberholzer & Frank (2003). In the prey choice experiment in 2007, different starvation periods were used to test for the effect of starving. Slug eggs were kept in plastic boxes with moss and grass to avoid desiccation and stored at 4°C until needed. The feeding trials were carried out in Petri dishes, 9 cm diameter by 6 cm tall. The bottom of each dish was lined with wet filter paper to maintain high humidity. In each experiment, one beetle specimen was added to half of the Petri dishes while the remained contained no beetles and served as controls.

#### Predation on eggs and newly hatched slugs in Petri dishes

Eggs were placed on a thin layer of soil, covered by moss, placed at room temperature and inspected daily for hatching. Hatchlings were fed with carrot and stored at 4°C until the start of the experiments when they were 1–4 days old. The mean total biomass of eggs and hatchlings provided for each beetle was 31 ± 4 mg and 19 ± 2 fresh weight, respectively. The walls of the Petri dishes were smeared with FLUON® (polytetrafluoroethylene) to prevent the slugs from escaping out of reach of the beetles (Symondson, 1993). The number of eggs and hatchlings offered to the beetles varied between experiments (table 1), which had to be taken into account in the numerical analyses. The number of beetles (replicates) used in the different experiments also varied (table 1), but the total for each species was thought to be sufficient to analyse the capability for handling eggs and newly hatched slugs. Furthermore, a predation index (PI) was defined, as

$$PI = \text{number of prey attacked} / \sqrt{\text{number of prey offered}}$$

This gives higher scores to beetles which consumed a given percentage of the eggs or hatchlings offered, if higher numbers were offered. Thus, a higher index score is given for a beetle consuming ten out of 20 slug eggs compared to a beetle consuming five out of ten, even though proportionally both fed the same. In this way, we can score predation by a beetle more accurately. To provide a scale for the predation

Table 2. Calculation of biomass (mg) of consumed eggs and newly hatched slugs within the different consumption indices (CI): 0, no eggs or slugs; 1, 1–25% consumed; 2, 26–50% consumed; 3, 51–75% consumed; and 4, 76–100% consumed.

CI	0	1	2	3	4
Egg	0	7.75	15.5	23.25	31.0
Hatchlings	0	4.75	9.5	14.25	19.0

indices, to make them comparable to the other indices which are fractions, they were standardised by dividing all indices by the largest PI, so that the final index would range between 0–1.

After 24 h in darkness at 19°C, the beetles were removed and the state of damage to eggs and hatchlings assessed. The beetles were used in 1–3 experiments due to the limited availability of beetles. The same starving period and storing conditions were used between experiments when the same beetle was used on multiple occasions. In order to estimate the biomass of slug eggs or hatchlings consumed, each egg or hatchling was given a consumption index (CI) ranging between 0–4 (Oberholzer & Frank, 2003) as follows: 0, not consumed or destroyed; 1, up to 25% of content consumed; 2, 26–50% of contents consumed; 3, 51–75% of content consumed; and 4, 76–100% of content consumed. Biomass of eggs and hatchlings consumed was calculated by multiplying the number of eggs or hatchlings in each consumption category by the mass for each category and added together (table 2). This sum was divided by the total biomass offered to give the fraction consumed. The number of replicates differed due to the limited availability of beetles, eggs and newly hatched slugs (table 1).

#### Predation on eggs and newly hatched slugs in mini-plots

These experiments were carried out in two Styrofoam boxes measuring 46 × 75 × 20 cm (base area 0.345 m<sup>2</sup>). The boxes were filled with ~10 cm of substrate, consisting of vegetation with attached soil. The substrate was collected randomly from the same site where beetles and slugs were collected in 2003. The vegetation consisted mainly of grasses (Poaceae) and the moss *Rhytidadelphus squarrosus* and was cut down to 8–10 cm. No effort was made to extract alternative prey from the vegetation in order to make the experimental design as natural as possible. Each box was divided by aluminium sheets into five experimental arenas, each 0.069 m<sup>2</sup> (15 × 46 cm). The dividing walls, as well as the walls of the Styrofoam boxes, were smeared with a 2 cm wide strip of FLUON® to prevent the slugs from escaping (Symondson, 1993). Twenty eggs of *A. lusitanicus* were transferred to each experimental arena. The eggs were divided into two groups of ten eggs each and covered with vegetation to mimic, as closely as possible, the way that eggs can be found in clusters in the field. In addition, 20 newly hatched slugs were distributed throughout each experimental arena. The boxes were covered by transparent plastic sheets to maintain humidity. Since the beetles used in this experiment are nocturnal, the experiment was carried out in a darkened room. The beetles were kept in the arenas for 72 h at 12–14°C. They were then removed, the remaining eggs and hatchlings were collected and the numbers lost

were scored. A total of six *P. melanarius* and six *P. niger* were tested.

#### Prey size choice by carabids feeding on *A. lusitanicus*

Each beetle (26 specimens of *C. nemoralis*, 9 *P. niger* and 11 *S. erythropterus*) were offered three relatively different sized *A. lusitanicus*: one small (0.1–0.3 g), one medium (0.3–0.9 g) and one large-sized slug (0.6–2.4 g). The Petri dishes were covered by lids to avoid slugs or beetles escaping. The experiment progressed for 2 h under low light at 19°C, and choice of slugs was detected during the experiment. The first slug that was killed and partly or completely consumed was recorded for further analyses.

#### Prey choice experiment with *C. nemoralis* feeding on *A. lusitanicus* and *D. reticulatum*

Each *C. nemoralis* was offered two slugs of the same size, one *A. lusitanicus* and one *D. reticulatum*. The weight of the slugs ranged from 0.1 to 0.6 g fresh weight. The experiment progressed under the same conditions as the size choice experiment and the results were also obtained in the same way, except that different starving periods (0, 3, 6, 9, 13, 14 and 17 days) were used to look for any effects of starving on prey choice.

#### Statistical analyses

All statistical analyses were performed utilizing the free software R (version 2.8.0) (R Development Core Team, 2008). There was no reason to anticipate a parametric distribution of the predation index (PI) since the data consisted of counts, and the data did not follow a normal distribution according to the Kolmogorov-Smirnov test. Thus, the non-parametric Kruskal-Wallis test was employed to indicate significant differences in PI between the pooled data for the carabids as well as differences between the experiments conducted with different starving regimes in 2003 and 2004. When several groups are compared in statistical tests, multiple comparison tests will avoid the increased probability of a type I error that occurs if more than two groups are incorporated in the same test. To identify the species of carabids that most efficiently preyed on or killed eggs or newly hatched slugs, we employed the non-parametric multiple comparisons for unequal sample sizes test described in Zar (1999). This procedure is based on the Nemenyi test but arranges the mean ranks, rather than the rank sums, in order of magnitude. Our comparisons of predation indices (PI) between different species in the feeding trials, therefore, were based on pooled data.

A generalised linear mixed-effect model (GLMM) was applied for the size-choice experiment by using the function *glmmPQL*, which is available in the MASS package for R. The observed first choice that led to consuming a slug by a given beetle was used as the response variable, and the three different size categories (small, medium, large) were used as the explanatory variable. A mixed model, using the beetles as a random factor, was included since the choice of slug was dependent on the other two slugs available for each beetle.

Table 3. Mean fraction of eggs and newly hatched slugs attacked and consumed during the feeding trials for eggs and hatchlings offered alone, predation on eggs when offered with hatchlings, predation on hatchlings when offered with eggs and hatchlings together. PI, predation index.

Species	Treatment	Repeats	Attacked	Consumed	P.I.
<i>C. nemoralis</i>	Eggs only	10	0.80	0.61	0.58
	Hatch. only	1	0.40	0.28	0.40
	Eggs with hatch.	10	0.81		
	Hatch. with eggs	10	0.90		
	Eggs & hatch.	10	0.82	0.74	0.59
<i>N. brevicollis</i>	Eggs only	22	0.05	0.05	0.03
	Hatch. only	0	–		
	Eggs with hatch.	5	0.03		
	Hatch. with eggs	5	0.00		
	Eggs & hatch.	5	0.14	0.02	0.03
<i>P. melanarius</i>	Eggs only	6	0.79	0.63	0.62
	Hatch. only	4	0.93	0.46	0.65
	Eggs with hatch.	2	0.75		
	Hatch. with eggs	2	0.75		
	Eggs & hatch.	2	0.75	0.61	0.75
<i>P. niger</i>	Eggs only	7	0.95	0.80	0.78
	Hatch. only	5	0.92	0.86	0.65
	Eggs with hatch.	6	1.00		
	Hatch. with eggs	6	1.00		
	Eggs & hatch.	6	1.00	0.86	0.93
<i>S. erythropterus</i>	Eggs only	3	0.91	0.54	0.65
	Hatch. only	2	0.45	0.31	0.32
	Eggs with hatch.	7	0.82		
	Hatch. with eggs	7	0.94		
	Eggs & hatch.	7	0.72	0.68	0.75

## Results

### Predation on eggs and newly hatched slugs

A Kruskal-Wallis test found, despite the difference in starvation periods, no difference in predation index between the experiments of 2003 and 2004 ( $\chi = 2.6461$ ,  $df = 1$ ,  $P > 0.05$ ); and, consequently, they were analysed together. No loss of, or damage to, eggs or newly hatched slugs was observed in the control dishes. The maximum biomasses of eggs and newly hatched slugs consumed were 31 mg and 19 mg, respectively (table 2). The results of the feeding trials are summarised in table 3. A Kruskal-Wallis test revealed significant differences in predation indices between the species tested ( $\chi = 61.008$ ,  $df = 4$ ,  $P < 0.001$ ). Pairwise comparisons revealed that *N. brevicollis* killed and consumed significantly less prey than the other species; thus, this species was excluded from the non-parametric multiple comparison models. The multiple comparison models revealed a significantly higher predation index ( $P < 0.05$ ) for *P. niger* compared to *P. melanarius*, *S. erythropterus* and *C. nemoralis*. The results were qualitatively similar when comparing the proportions attacked and consumed (data not shown), expect that the differences between *C. nemoralis* and *P. niger* were not significant. Comparisons between *P. melanarius*, *S. erythropterus* and *C. nemoralis* were not significantly different in terms of attacks and consumptions.

### Predation on eggs and hatchlings under semi-natural conditions

No significant difference was found between *P. melanarius* and *P. niger* (table 4; Kruskal-Wallis test,  $\chi = 0.1961$ ,

Table 4. Predation on eggs and hatchlings under semi-natural conditions. Results are given as proportion of available prey consumed.

Species	<i>Pterostichus niger</i>	<i>Pterostichus melanarius</i>
Eggs	0.58	0.57
Hatchlings	0.10	0.12

$df = 1$ ,  $P > 0.05$ ). Overall, the numbers preyed upon were lower than in the Petri-dish experiments. On average for all beetles, 15% of hatchlings and 55% of eggs were attacked during the 72 h experiment.

### Prey-size-choice experiments using *A. lusitanicus*

*Carabus nemoralis* showed size preferences when feeding on *A. lusitanicus*, preferring slugs smaller than 1 g (fig. 1). Predation on slugs termed 'large' was significantly lower compared with the ones termed 'small' (GLMM,  $P$ -value = 0.0019) and the ones termed 'medium' ( $P$ -value = 0.0365). However, predation on these two latter groups was not significantly different from each other ( $P$ -value = 0.1113). *P. niger* seems to have less appetite for juvenile slugs than the larger beetle *C. nemoralis*. Only half of the former beetles consumed slugs while more than 80% of the latter beetles consumed slugs, of which 40% consumed more than one slug during the 2 h long experiment and 20% ate all three slugs. *Staphylinus erythropterus* did not kill any of the juvenile slugs weighing 0.1 g or more, suggesting that the slugs were too large for this beetle to handle.

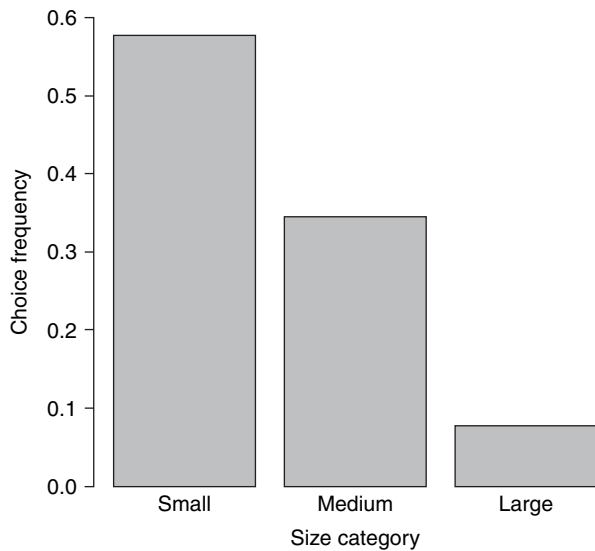


Fig. 1. Beetles choice in size categories of *Arion lusitanicus*. Results are given as proportions of beetles choosing the different sizes of slugs. Small, 0.1–0.3 g; medium, 0.3–0.9 g; and large, 0.6–2.4 g.

#### Prey choice experiments using *A. lusitanicus* and *D. reticulatum*

A Kruskal-Wallis test suggested no significant difference in prey choice by *C. nemoralis* ( $\chi = 3.4893$ ,  $df = 2$ ,  $P$ -value = 0.1747) when choosing between *A. lusitanicus* and *D. reticulatum*. A total of 30 beetles chose *A. lusitanicus*, while 24 beetles fed on *D. reticulatum* and 12 did not feed. No prey preference between the two slugs was persistent for all the different starvation periods.

#### Behaviour of beetles and slugs

*Carabus nemoralis* showed an extraordinary variety of behaviours in the Petri-dish experiments. Some individuals attacked the first encountered slug or egg when put into the experimental arena, while others did not attack any prey the first 20–60 minutes. They attacked the slugs at the posterior end and often killed their prey during the first attack. The other carabid species, as well as the staphylinid, were in most cases stressed during the first minutes of the experiments. These species did not direct their attacks to any specific part of the slug and normally attacked repeatedly the larger juveniles at intervals, with frequent cleaning of their mandibles to remove mucus. *Pterostichus niger* attacked juveniles larger than 0.1 g, but often gave up after several failed attempts.

## Discussion

#### Capacity to eat *A. lusitanicus* eggs and hatchlings

We have demonstrated that the common beetles, *C. nemoralis*, *P. melanarius*, *P. niger* and *S. erythropterus*, will all eat eggs and newly hatched *A. lusitanicus* when offered them in the laboratory. *Nebria brevicollis* did not eat *A. lusitanicus* eggs and hatchlings under our experimental conditions. While *N. brevicollis* did destroy some eggs, a closer

examination revealed that the eggs were just bitten into and hardly anything consumed. The hungry beetles seem to have tasted the eggs but then rejected them. As *N. brevicollis* is regarded to feed mainly on springtails (Thiele, 1977), it is possible that eggs and juveniles of the Iberian slug are outside the size range of suitable prey and/or are unpalatable to this species. Tod (1973) found a correlation between size of beetle and slug predation in the field, where *N. brevicollis* crops rarely contained slugs, while the larger species, like *C. nemoralis*, fed on slugs frequently. Ayre (2001) found that *N. brevicollis* preyed upon *D. reticulatum* juveniles, but these were considerably smaller (ca. 4 mm) than those of *A. lusitanicus* used in our study (8–10 mm).

In the Petri-dish experiments, the fraction of prey consumed was lower than the fraction attacked for all beetle species; however, the differences were small and the two measures highly correlated. This suggests that eggs and juveniles of *A. lusitanicus* were edible to all these beetle species, except *N. brevicollis*. However, slugs like *A. lusitanicus* may be less palatable than other prey (e.g. earthworms), and more palatable prey may be preferred under field conditions where a range of prey are on offer.

The mini-plot experiment, under semi-natural conditions, addressed one of the shortcomings of the Petri-dish experiments by introducing search time; the eggs and newly hatched slugs were presented in conditions emulating those in the field. The setting was not entirely realistic since the area over which the beetles could roam was limited. The mini-plot experiments showed that a substantially lower fraction of eggs and hatchlings were preyed upon under such conditions than in the highly unnatural conditions of the Petri dishes. However, over 50% of eggs and 15% of hatchlings were preyed upon during 72 h, demonstrating that the beetles were able to find and eat slug eggs and newly hatched slugs under these conditions. In the laboratory, hatching of *A. lusitanicus* eggs took 20 days in room temperature (19–21°C) and 150 days at low temperatures (4–6°C), with large variations between egg clutches held at the same temperature (Tomasgård, 2005). The eggs are thus exposed in the field to potential predation for periods 7–50 times longer than the duration of the mini-plot experiments, giving ample time for predation.

#### Predation on juvenile *A. lusitanicus* and *D. reticulatum*

*Carabus nemoralis* consumed juveniles of up to 1.3 g, although a significant preference for slugs less than 1 g was found. *Pterostichus niger*, on the other hand, appeared to have difficulties handling the mucus of juvenile slugs and seems to be restricted to eggs and newly hatched *A. lusitanicus*. This is partly in accordance with a previous study by Kaiser *et al.* (1993), who found a preference for juvenile slugs of *A. lusitanicus* less than 0.1 g in *Carabus cancellatus*, while *P. melanarius* preferred eggs and *C. granulatus* showed no particular preference. Furthermore, Paill (2000) found that *C. violaceus* preferred smaller *A. lusitanicus* in the field. However, a preference for smaller slugs may be counterbalanced in the field since larger juveniles might be easier to find, which has been shown by McKemey *et al.* (2003) for *P. melanarius* feeding on *D. reticulatum*.

A number of studies have shown that ground beetles will prey upon gastropods irrespective of the length of the starving period prior to the experiment (Ayre, 2001; Mair & Port, 2001; McKemey *et al.*, 2001, 2003; Oberholzer & Frank,

2003; Oberholzer *et al.*, 2003), so eating slugs seems not to depend upon duration of starvation. Further, starving seems to have no effect on choice of slug species by *C. nemoralis*, and no preference existed for *D. reticulatum* vs. *A. lusitanicus*. The same lack of preference of slug species has been found for *Carabus problematicus* and *Abax parallelepipedus* when given *Arion subfuscus*, *A. intermedius*, *A. circumscriptus*, *A. ater* and *Malacolimax tenellus* (Bless, 1977).

#### *Predation in the field on A. lusitanicus*

We have demonstrated that several beetle species will eat eggs and juveniles of *A. lusitanicus* under experimental laboratory conditions, but more research is needed to determine to what extent beetles actually eat slugs in the field. Many small-sized slugs are found up in the vegetation, where they may be out of reach of some beetles, particularly large species like *C. nemoralis*. Except for *S. erythropterus*, the beetles we studied are all primarily nocturnal, and slugs might thus escape predation by climbing up into the vegetation at night. It is also questionable how palatable the eggs and juveniles of *A. lusitanicus* are to beetles, in particular, when alternative prey is available. In a rather questionable experiment, Schroeder *et al.* (1999) isolated the defensive diterpene miriamin from *A. lusitanicus* eggs and showed that the substance deterred the coccinellid *Harmonia axyridis* from feeding on moth eggs coated with this extract. As these coccinellids do not eat mollusc eggs (and no comparison was made with extracts from the eggs of other mollusc species), this tells us nothing about the potential deterrence of slug predators, such as *Carabus* species. Oberholzer & Frank (2003) found that *P. melanarius* fed on *A. lusitanicus* eggs with no harm to the beetles. Similarly, *P. melanarius* fed exclusively on *A. lusitanicus* eggs over several weeks and showed no mortality (W.O.C. Symondson, unpublished data). We did not observe anything indicating that the beetles suffered any harm from feeding on slug eggs and juveniles, suggesting that these carabids and staphylinids are insensitive to diterpene miriamin.

Direct observation of predation in the field is difficult since both the beetles and the slugs are nocturnal and spend much time in dense vegetation. Molecular markers to identify *A. lusitanicus* DNA in beetle guts will be useful in revealing food preferences together with DNA-based markers and methods already developed by Harper *et al.* (2005), Dodd (2004) and Hatteland, B.A., Noble, L.R., Schander, C., Skage, T. & Solhøy, T. (in prep.). Appropriate markers for *A. lusitanicus* DNA have been optimised for foregut analyses (Hatteland, B.A., King, R.A., Symondson, W.O.C. & Solhøy, T., in prep.); thus, it will be possible for a range of beetle species to be screened for evidence of feeding on *A. lusitanicus* as well as other slugs. DNA-based markers will not be able to distinguish between predation on slugs and their eggs, however, although this has been shown to be possible using monoclonal antibodies (Symondson *et al.*, 1995; Mendis *et al.*, 1996).

The activity peak of a potentially useful predator should coincide with the egg and juvenile phases of *A. lusitanicus*; the egg phase takes place from September to November and the juvenile phase is mainly from October to June in Norway (Dirks *et al.*, in prep.). The potential of the predator species tested in this study only partly fulfil this criterion. *Carabus nemoralis* is mainly active as adults during spring and early summer with a smaller activity peak in early autumn

(Lindroth, 1985; Turin *et al.*, 2003), when the abundance of juvenile slugs is highest. This is unlike other *Carabus* species, as well as *P. melanarius*, *P. niger*, *N. brevicollis* and *S. erythropterus*, that all have an activity peak in the latter half of the summer (July–August) when the juvenile slugs are less abundant (Hatteland *et al.*, in prep.). *Pterostichus melanarius* has been found to feed mainly on *A. lusitanicus* in spring and autumn when juvenile slugs of <200 mg are abundant (Paill, 2004). However, the larvae of the two *Pterostichus* species are active in autumn (Kaiser *et al.*, 1993; Thomas, 2002) and *N. brevicollis* larvae are active in spring (Traugott, 1998). *Pterostichus melanarius* larvae have been shown to feed on both *D. reticulatum* and *Arion intermedius* under semi-field conditions (Thomas *et al.*, 2009), and orientation towards slugs in the soil is possible by olfaction (Thomas *et al.*, 2008).

#### *The potential of beetle predators for biological control of A. lusitanicus*

Since we found that several carabid species ate eggs and juveniles of *A. lusitanicus*, any measures which promote abundance of these beetles (and possibly also staphylinids) are likely to be beneficial in terms of slug population reduction. The most realistic biological control of slugs by beetles is often not classical nor inundative biological control but conservation biological control (Symondson, 2004). Introducing alien beetles is too risky and culturing these beetles is very expensive due to cannibalism among larvae. The only species that has so far been found possible to mass culture economically is the woodland-edge species *Abax parallelepipedus* (Symondson, 1994), which is common in many parts of Europe but not present in western Norway (Lindroth, 1986). Crop management practices should rather take into account the beetle fauna by reducing use of insecticides and those molluscicides that are toxic to carabids, especially in spring when *C. nemoralis* is active. Provision of refugia may also increase the number of beetles (Altieri *et al.*, 1982). Alternatively, or additionally, hedges may be planted to connect surrounding woodlands to arable fields since *C. nemoralis* has been found to disperse along such habitat strips (Glück & Kreisel, 1986; Gruttke, 1994). Future studies should address the numbers of beetles needed to control slugs like *A. lusitanicus*, and whether these numbers are present in arable fields and surrounding habitats, or if they need to be increased. *Arion lusitanicus* has been found to be numerous and causes problems for strawberry cultivation even though beetles like *C. nemoralis* were present (Hatteland *et al.*, in prep.), which suggests that some sort of manipulation is necessary. On the other hand, we do not know how the exclusion of such predators will affect the slug populations. Clearly, manipulative studies under field conditions are needed to explore how the beetles are affecting pest species like *A. lusitanicus*. However, studies of predation in the field (as mentioned above) should first be carried out to determine to what extent these beetles are feeding on *A. lusitanicus*.

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## References

- Altieri, M.A., Hagen, K.S., Trujillo, J. & Caltagirone, L.E.** (1982) Biological control of *Limax maximus* and *Helix aspersa* by indigenous predators in a daisy field in central coastal California. *Acta Oecologia* **3**, 387–390.
- Anderson, R.** (2005) An annotated list of the non-marine Mollusca of Britain and Ireland. *Journal of Conchology* **38**, 607.
- Asteraki, E.J.** (1993) The potential of carabid beetles to control slugs in grass/clover swards. *Entomophaga* **38**, 193–198.
- Ayre, K.** (2001) Effect of predator size and temperature on the predation of *Deroceras reticulatum* (Müller) (Mollusca) by carabid beetles. *Journal of Applied Entomology* **125**, 389–395.
- Barker, G.M.** (2002) *Molluscs as Crop Pests*. Wallingford, UK, CABI Publishing.
- Bless, R.** (1977) Studies on the relationships of carabids as predators to gastropods as prey. *Anzeiger für Schadlingskunde, Pflanzenschutz, Umweltschutz* **50**, 55–57.
- Bohan, D.A., Bohan, A.C., Glen, D.M., Symondson, W.O.C., Wiltshire, C.W. & Hughes, L.** (2000) Spatial dynamics of predation by carabid beetles on slugs. *Journal of Animal Ecology* **69**, 367–379.
- Dodd, C.S.** (2004) Development and optimisation of PCR-based techniques in predator gut analysis. PhD thesis, Cardiff University, Cardiff, UK.
- Dolmen, D. & Winge, K.** (1997) Boasneglen (*Limax maximus*) og iberiasneglen (*Arion lusitanicus*) i Norge; utbredelse, spredning og skadevirkninger (In Norwegian). 'Limax maximus and Arion lusitanicus in Norway: distribution, expansion and injurious effects', Report 4, 4–24. Trondheim, Norway, Vitenskapsmuseet.
- Eilenberg, J., Hajek, A. & Lomer, C.** (2001) Suggestions for unifying the terminology in biological control. *BioControl* **46**, 387–400.
- Essl, F. & Rabitsch, W.** (2002) *Neobiota in Österreich*. Vienna, Austria, Umweltbundesamt.
- Glück, E. & Kreisel, A.** (1986) Die Hecke als Lebensraum, Refugium und Vernetzungsstruktur und ihre Bedeutung für die Dispersion von Waldcarabidenarten. *Laufener Seminarbeiträge* **10**, 64–83.
- Grimm, B., Paill, W. & Kaiser, H.** (2000) Daily activities of the pest slug *Arion lusitanicus*. *Journal of Molluscan Studies* **66**, 125–130.
- Gruttko, H.** (1994) Dispersal of carabid species along a linear sequence of young hedge plantations. pp. 299–303 in Desender, K., Dufrêne, M., Loreau, M., Luff, M.L. & Maelfait, J.-P. (Eds) *Carabid Beetles: Ecology and Evolution*. Kluwer Academic Publishers.
- Harper, G.L., King, R.A., Dodd, C.S., Harwood, J.D., Glen, D.M., Bruford, M.W. & Symondson, W.O.C.** (2005) Rapid screening of invertebrate predators for multiple prey DNA targets. *Molecular Ecology* **14**, 819–827.
- Hengeveld, R.** (1980a) Food specialization in ground beetles: an ecological or a phylogenetic process? (Coleoptera, Carabidae). *Netherlands Journal of Zoology* **30**, 585–594.
- Hengeveld, R.** (1980b) Qualitative and quantitative aspects of the food of ground beetles (Coleoptera, Carabidae): a review. *Netherlands Journal of Zoology*, **30**, 555–563.
- Hommay, G.** (2002) Agriolimacidae, Arionidae and Milacidae as pests in West European sunflower and maize. pp. 245–254 in Barker, G.M. (Ed.) *Molluscs as crop pests*. CABI Publishing, Wallingford, UK.
- Iglesias, J., Castillejo, J., Ester, A., Castro, R. & Lombardia, M.J.** (2002) Susceptibility of the eggs of the field slug *Deroceras reticulatum* to contact with pesticides and substances of biological origin on artificial soil. *Annales of Applied Biology* **140**, 53–59.
- Kaiser, H., Geiersberger, U., Grimm, B. & Paill, W.** (1993) Untersuchungen über die biologischen und ökologischen Voraussetzungen des Massenauftritts der Spanischen Wegschnecke. Final report. Graz, Austria.
- Keller, M., Kollmann, J. & Edwards, P.J.** (1999) Palatability of weeds from different European origins to the slugs *Deroceras reticulatum* Müller and *Arion lusitanicus* Mabilie. *Acta Oecologica* **20**, 109–118.
- Langan, A.M., Taylor, A. & Wheeler, C.P.** (2004) Effects of metaldehyde and methiocarb on feeding preferences and survival of a slug predator (*Pterostichus melanarius* (F.): Carabidae, Pterostichini). *Journal of Applied Entomology* **128**, 51–55.
- Lindroth, C.H.** (1985) The Carabidae (Coleoptera) of Fennoscandia and Denmark. Vol. 1. *Fauna Entomologica Scandinavica* **15**, 1–225.
- Lindroth, C.H.** (1986) The Carabidae (Coleoptera) of Fennoscandia and Denmark. Vol. 2. *Fauna Entomologica Scandinavica* **15**, 226–497.
- Luff, M.L.** (2007) *The Carabidae (ground beetles) of Britain and Ireland*, 2nd edn. Royal Entomological Society, St Albans, UK.
- Mair, J. & Port, G.R.** (2001) Predation on the slug *Deroceras reticulatum* by the carabid beetles *Pterostichus madidus* and *Nebria brevicollis* in the presence of alternative prey. *Agriculture and Forest Entomology* **3**, 169–174.
- McKeme, A.R., Symondson, W.O.C., Glen, D.M. & Brain, P.** (2001) Effects of slug size on predation by *Pterostichus melanarius* (Coleoptera: Carabidae). *Biocontrol Science and Technology* **11**, 81–91.
- McKeme, A.R., Symondson, W.O.C. & Glen, D.M.** (2003) Predation and prey size choice by the carabid beetle *Pterostichus melanarius* (Coleoptera: Carabidae): the dangers of extrapolating from laboratory to field. *Bulletin of Entomological Research* **93**, 227–234.
- Mendis, V.W., Bowen, I.D., Liddell, J.E. & Symondson, W.O.C.** (1996) Monoclonal antibodies against *Deroceras reticulatum* and *Arion ater* eggs for use in predation studies. pp. 99–106 in Henderson, I. (Ed.) *Slug and snail pests in agriculture*. BCPC Symposium Proceedings No. 66. Farnham, UK: British Crop Protection Council.
- Oberholzer, F. & Frank, T.** (2003) Predation by the carabid beetles *Pterostichus melanarius* and *Poecilus cupreus* on slugs and slug eggs. *Biocontrol Science and Technology* **13**, 99–110.
- Oberholzer, F., Escher, N. & Frank, T.** (2003) The potential of carabid beetles (Coleoptera) to reduce slug damage to oilseed rape in the laboratory. *European Journal of Entomology* **100**, 81–85.
- Paill, W.** (2000) Slugs as prey for larvae and imagines of *Carabus violaceus* L. (Coleoptera, Carabidae). pp. 221–22 in Brandmayr, P., Lövei, G.L., Brandmayr, T.Z., Casale, A. &



- Vigna Taglianti, A. (Eds) *Natural history and applied ecology of carabid beetles*. Sofia, Bulgaria, Pensoft Publishers.
- Pail, W.** (2004) Slug feeding in the carabid beetle *Pterostichus melanarius*: seasonality and dependence on prey size. *Journal of Molluscan Studies* **70**, 203–205.
- Pakarinen, E.** (1994) The importance of mucus as a defence against carabid beetles by the slugs *Arion fasciatus* and *Deroceras reticulatum*. *Journal of Molluscan Studies* **60**, 149–155.
- Port, C.M. & Port, G.** (1986) The biology and behaviour of slugs in relation to crop damage and control. *Agricultural Zoology Reviews* **1**, 255–299.
- Poulin, G. & O'Neil, L.G.** (1969) Observations sur les prédateurs de la limace noire, *Arion ater* (L.) (Gastéropodes, Pulmonés, Arionidés). *Phytoprotection* **50**, 1–6.
- R Development Core Team,** (2008) R: A language and environment for statistical computing. R Foundation for Statistical Computing. Vienna, Austria. Available online at <http://www.R-project.org>.
- Schroeder, F.C., Gonzalez, A., Eisner, T. & Meinwald, J.** (1999) Miriamin, a defence diterpene from the eggs of a land slug (*Arion* sp.). *Proceedings of the National Academy of Sciences USA* **96**, 13620–13625.
- South, A.** (1992) *Terrestrial Slugs: Biology, Ecology and Control*. London, UK, Chapman & Hall.
- Sunderland, K.D.** (2002) Invertebrate pest control of slugs by carabids. pp. 165–214 in Holland, J.M. (Ed.) *The Agroecology of Carabid Beetles*. Andover, UK, Intercept.
- Symondson, W.O.C.** (1993) The effects of crop development upon slug distribution and control by *Abax parallelepipedus* (Coleoptera, Carabidae). *Annales of Applied Biology* **123**, 449–457.
- Symondson, W.O.C.** (1994) The potential of *Abax parallelepipedus* (Coleoptera, Carabidae) for mass breeding as a biological control agent against slugs. *Entomophaga* **39**, 323–333.
- Symondson, W.O.C.** (2002) Diagnostic techniques for determining carabid diets. pp. 111–164 in Holland, J.M. (Ed.) *The Agroecology of Carabid Beetles*. Andover, UK, Intercept.
- Symondson, W.O.C.** (2004) Coleoptera (Carabidae, Staphylinidae, Lampyridae, Drilidae and Silphidae) as predators of terrestrial gastropods. pp. 37–84 in Barker, G.M. (Ed.) *Natural Enemies of Terrestrial Molluscs*. Oxford, UK, CAB International.
- Symondson, W.O.C., Mendis, V.W. & Liddell, J.E.** (1995) Monoclonal antibodies for the identification of slugs and their eggs. *EPP0 Bulletin* **25**, 377–382.
- Symondson, W.O.C., Glen, D.M., Whiltshire, C.W., Langdon, C.J. & Liddell, J.E.** (1996) Effects of cultivation techniques and methods of straw disposal on predation by *Pterostichus melanarius* (Coleoptera: Carabidae) upon slugs (Gastropoda: Pulmonata) in an arable field. *Journal of Applied Ecology* **33**, 741–753.
- Symondson, W.O.C., Glen, D.M., Ives, A.R., Langdon, C.J. & Wiltshire, C.W.** (2002) Dynamics of the relationship between a generalist predator and slugs over five years. *Ecology* **83**, 137–147.
- Thiele, H.U.** (1977) *Carabid Beetles in their Environments*. Berlin, Germany, Springer Verlag.
- Thomas, R.S.** (2002) An immunological and behavioural study of the role of carabid beetle larvae as slug control agents in cereal crops. PhD thesis, Cardiff University, Cardiff, UK.
- Thomas, R.S., Glen, D.M. & Symondson, W.O.C.** (2008) Prey detection through olfaction by the soil-dwelling larvae of the carabid predator *Pterostichus melanarius*. *Soil Biology and Biochemistry* **40**, 207–216.
- Thomas, R.S., Harwood, J.D., Glen, D.M. & Symondson, W.O.C.** (2009) Molecular tracking of subterranean density-dependent predation by carabid larvae on slugs. *Ecological Entomology* **34**, 569–579.
- Tod, M.E.** (1973) Notes on beetle predators of mollusks. *The Entomologist* **106**, 196–201.
- Toft, S. & Bilde, T.** (2002) Carabid diets and food value. pp. 81–110 in Holland, J.M. (Ed.) *The Agroecology of Carabid Beetles*. Andover, UK, Intercept.
- Tomasgård, T.E.H.** (2005) Populasjonsdynamikk, næringspreferansar og reproduksjon/vekst hjå snilen *Arion lusitanicus* Mabille, 1868 (In Norwegian). 'Population dynamics, food preferences, reproduction and growth in the slug *Arion lusitanicus* Mabille, 1868'. Unpublished Candidata Scientiarum-thesis, University of Bergen, Bergen, Norway.
- Traugott, M.** (1998) Larval and adult species composition, phenology and life cycles of carabid beetles (Coleoptera, Carabidae) in an organic potato field. *European Journal of Soil Biology* **34**, 189–197.
- Turin, H., Penev, L., Casale, A., Arndt, E., Assman, T., Makarov, K., Mossakowski, D., Szél, G. & Weber, F.** (2003) Species accounts. pp. 151–284 in Turin, H., Penev, L. & Casale, A. (Eds) *The Genus Carabus in Europe: A Synthesis*. Sofia, Bulgaria, Pensoft Publishers.
- von Proschwitz, T.** (1996) Utbredning och spridning av spansk skogssnigel (*Arion lusitanicus* Mabille) och röd skogssnigel (*Arion rufus* L.) – en översikt av utvecklingen i Sverige (In Swedish). 'Distribution and spreading of the Iberian slug (*Arion lusitanicus* Mabille) and the red slug (*Arion rufus* L.) – an overview of the development in Sweden'. Göteborgs Naturhistoriska Museums årstryck **1996**, 27–39.
- von Proschwitz, T. & Winge, K.** (1994) Iberiaskogssnegl – en art på spridning i Norge (In Norwegian). 'The Iberian slug – a species expanding in Norway'. *Fauna* **47**, 195–203.
- Zar, J.H.** (1999) *Biostatistical Analysis, 4th edn*. New Jersey, USA, Prentice-Hall.