Adaptation of *Lymnaea fuscus* and *Radix balthica* to *Fasciola hepatica* through the experimental infection of several successive snail generations

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**Abstract**

**Background:** High prevalence of *Fasciola hepatica* infection (>70%) was noted during several outbreaks before the 2000s in several French farms where *Galba truncatula* is lacking. Other lymnaeids such as *Lymnaea fuscus*, *L. glabra* and/or *Radix balthica* are living in meadows around these farms but only juvenile snails can sustain complete larval development of *F. hepatica* while older snails were resistant. The low prevalence of infection (<20%) and limited cercarial production (<50 cercariae per infected snail) noted with these juveniles could not explain the high values noted in these cattle herds. As paramphistomosis due to *Calicophoron daubneyi* was not still noted in these farms, the existence of another mode of infection was hypothesized. Experimental infection of several successive generations of *L. glabra*, originating from eggs laid by their parents already infected with this parasite resulted in a progressive increase in prevalence of snail infection and the number of shed cercariae. The aim of this paper was to determine if this mode of snail infection was specific to *L. glabra*, or it might occur in other lymnaeid species such as *L. fuscus* and *R. balthica*.

**Methods:** Five successive generations of *L. fuscus* and *R. balthica* were subjected to individual bimiracidial infections in the laboratory. Resulting rediae and cercariae in the first four generations were counted after snail dissection at day 50 p.e. (20°C), while the dynamics of cercarial shedding was followed in the F5 generation.

**Results:** In the first experiment, prevalence and intensity of *F. hepatica* infection in snails progressively increased from the F1 (*R. balthica*) or F2 (*L. fuscus*) generation. In the second experiment, the prevalence of *F. hepatica* infection and the number of shed cercariae were significantly lower in *L. fuscus* and *R. balthica* (without significant differences between both lymnaeids) than in *G. truncatula*.

**Conclusion:** The *F. hepatica* infection of several successive snail generations, coming from parents infected with this parasite, resulted in a progressive increase in prevalence and intensity of snail infection. This may explain high prevalence of fasciolosis noted in several cattle-breeding farms when the common snail host of this digenean, *G. truncatula*, is lacking.

**Keywords:** Cercaria, Experimental infection, *Fasciola hepatica*, *Galba truncatula*, *Lymnaea fuscus*, Prevalence, *Radix balthica*, Redia
low prevalence of *F. hepatica* infection (generally, < 20%) and a low number of shed cercariae [5-10]. Under these conditions, it is difficult to understand the mechanism of snail infection with *F. hepatica* when *G. truncatula* is lacking in 21 cattle-breeding farms located in central France and known for high risks of animal fasciolosis (prevalence, > 70%) before the 2000s (D. Rondelaud, personal observation). As other lymnaeid species such as *L. glabra* are living in meadows present in these farms, the existence of another mode of snail infection with *F. hepatica* to explain high prevalence of bovine fasciolosis was hypothesized.

As Chipev et al. [11] in the model *G. truncatula/F. hepatica* and Southgate et al. [12] in the model *Bulinus tropicus/Schistosoma bovis* have demonstrated the role of a paramphistomid in these snail infections, a second mode of snail infection was demonstrated for French lymnaeids other than *G. truncatula*. Using a first exposure of snails with *Calocophoron daubneyi* and a second exposure with *F. hepatica* 4 hours later, successful infections of pre-adult *L. glabra* (shell height at exposure, 2.1-4 mm) with *F. hepatica*, *C. daubneyi*, or both, were reported under laboratory conditions [13,14]. This result was verified in the field because some *L. glabra* and *Lymnaea palustris* infected with *F. hepatica*, *C. daubneyi*, or both, were found in swampy meadows and wild watercress beds from central or eastern France [15-18]. However, this mode of snail infection using a paramphistomid before the penetration of *F. hepatica* miracidia did not explain all the cases of bovine fasciolosis in farms where *G. truncatula* is lacking. As paramphistomosis was not still detected before the 2000s in the livestock of seven farms (D. Rondelaud, personal observation), it was necessary to suppose a third mode of snail infection with *F. hepatica* in these farms.

In natural habitats colonized by *L. glabra* [6] and *L. palustris* [7], experimental introduction of *F. hepatica* miracidia during four successive years had induced a progressive increase in prevalence of snail infection. This finding was at the origin of experiments carried out to demonstrate the existence of this third mode of snail infection for lymnaeids other than *G. truncatula*. The infection of five successive generations of pre-adult *L. glabra* (snail height at exposure, 4 ± 0.1 mm), originating from parents already infected with this parasite, resulted in a progressive increase in prevalence of snail infection and the number of shed cercariae [19]. In view of this last result, it was useful to determine if this mode of snail infection was specific to *L. glabra*, or it might occur in other lymnaeid species such as *L. fuscus* and *R. balthica* which were living with *G. truncatula* in swampy meadows of central France [20,21]. To verify either possibility, five successive generations of *L. fuscus* and *R. balthica* were subjected to individual bimiracidial infections in the laboratory. Resulting rediae and cercariae in the first four generations were counted after snail dissection at day 50 post-exposure (p.e.) at 20°C, while the dynamics of cercarial shedding was followed in the F5 generation.

**Methods**

**Snails and parasite**

Both snail populations were living on the commune of Thenay, department of Indre, central France. *Lymnaea fuscus* inhabited a small pond (46°35′38″ N, 1°27′10″ E), whereas *R. balthica* was colonizing a brook (47°23′22″ N, 1°17′31″ E) in the centre of Thenay. The identification of the first population as *L. fuscus* and the second as *R. balthica* was confirmed by the study of ITS-2 sequences [22,23]. Previous investigations in adults (30 per sample and per population) over the six months preceding the beginning of the experiments have demonstrated the absence of natural trematode infections in snails when they were dissected in the laboratory. One hundred adults (16–20 mm in shell height) were collected from each population and placed in 5-L and 10-L covered aquaria with 5 snails per litre of permanently-oxygenated spring water. Aquaria were subjected to constant conditions (temperature, 20°C ± 1°C; light/dark period, 12 h/12 h). Dissolved calcium concentration in spring water was 35 mg/L. Pesticide-free leaves of fresh lettuce were given as food ad libitum and spring water in aquaria was changed weekly. Egg masses laid by these parents were placed into small rearing aquaria with oxygenated spring water and finely powdered lettuce. Four-millimetre high individuals were considered as the F1 generation and were used for experiments. The protocol was similar for the F2, F3, F4 and F5 snail generations (see below). A total of 300 *L. fuscus* and 300 *R. balthica*, measuring 4 ± 0.1 mm in height and belonging to the F1, F2, F3, F4 and F5 generations (Tables 1 and 2), were used for experiments.

To compare the characteristics of *F. hepatica* infection in *L. fuscus* and *R. balthica* with those found in a common intermediate host of this parasite, a wild population of *G. truncatula* was selected. This species is the main snail host of *F. hepatica* in Europe [1-3] and may be infected at any age [24]. These snails were colonizing a road ditch at Chézeau Chrétien (46°40′27″ N, 1°21′21″ E), commune of Chitray, department of Indre, central France. As for *L. fuscus* and *R. balthica*, no trematode larval forms were found in adult snails collected over a 6-month period before the beginning of the experiments and dissected in the laboratory. One hundred snails (shell height, 4 ± 0.1 mm), belonging to the overwintering generation, were collected and acclimatized for 24 hours to laboratory temperature before being exposed to *F. hepatica* miracidia. A single generation of *G. truncatula* was used for this study.
because prevalence of *F. hepatica* infection and the number of shed cercariae were significantly lower in the F1 generation born to already infected parents than those noted for the F1 generation born to unexposed parents [25].

Several isolates of *F. hepatica* eggs were regularly collected from the gall bladders of naturally infected limousine cattle at the Limoges slaughterhouse. They were washed several times with spring water and immediately incubated in the dark at 20°C for 20 days to allow miracidial development [26].

**Experimental protocol**

The *L. fuscus* and *R. balthica* susceptibility to *F. hepatica* miracidia was studied during five successive snail generations.

### Table 1 Several characteristics of *Fasciola hepatica* infection in four generations of *Lymnaea fuscus* and *Radix balthica* subjected to individual bimiracidial exposures, raised at 20°C and dissected at day 50 post-exposure (first experiment)

| Snail population and generation | Number of surviving snails at day 30 p.e. (%) | Number of infected snails with Immature rediae | Cercariae-containing rediae | Free cercariae | Cercarial shedding | Overall prevalence of infection (%) |
|---------------------------------|-----------------------------------------------|-----------------------------------------------|----------------------------|----------------|-------------------|-----------------------------------|
| *Lymnaea fuscus*                |                                               |                                               |                            |                |                   |                                   |
| - F1                            | 16 (32.0)                                     | -                                             | -                          | -              | -                 | -                                 |
| - F2                            | 24 (48.0)                                     | 1                                             | -                          | -              | -                 | 4.1                               |
| - F3                            | 27 (54.0)                                     | 2                                             | 2                          | 1              | -                 | 18.5                              |
| - F4                            | 29 (58.0)                                     | 2                                             | 4                          | 3              | 1                 | 34.4                              |
| *Radix balthica*                |                                               |                                               |                            |                |                   |                                   |
| - F1                            | 26 (42.0)                                     | 2                                             | 1                          | -              | -                 | 11.52                             |
| - F2                            | 27 (54.0)                                     | 2                                             | 3                          | 1              | -                 | 22.2                              |
| - F3                            | 33 (66.0)                                     | 1                                             | 5                          | 2              | 1                 | 27.2                              |
| - F4                            | 31 (62.0)                                     | 2                                             | 3                          | 7              | 3                 | 48.3                              |

*p.e.*, post-exposure.

### Table 2 Characteristics of *Fasciola hepatica* infection in *Lymnaea fuscus* (F5 generation), *Radix balthica* (F5 generation) and *Galba truncatula* subjected to individual bimiracidial exposures and raised at 20°C (second experiment)

| Population                  | *Lymnaea fuscus* | *Radix balthica* | *Galba truncatula* | Significant differences* |
|-----------------------------|------------------|------------------|--------------------|--------------------------|
| Number of snails:           |                  |                  |                    |                          |
| - at exposure               | 100              | 100              | 100                |                          |
| - at day 30 p.e. (%)        | 54 (54.0)        | 65 (65.0)        | 74 (74.0)          | $\chi^2 = 8.74, p < 0.05$ |
| Number of snails:           |                  |                  |                    |                          |
| - CS                        | 7                | 13               | 43                 |                          |
| - NCS                       | 12               | 14               | 11                 |                          |
| Prevalence of infection (%) | 35.1             | 41.5             | 72.9               | $\chi^2 = 21.99, p < 0.001$ |
| Shell growth (mm):          |                  |                  |                    |                          |
| - CS                        | 7.7 ± 1.5        | 6.0 ± 1.1        | 3.7 ± 0.9          | $F = 14.32, p < 0.001$   |
| - NCS                       | 7.3 ± 1.1        | 5.7 ± 0.8        | 3.5 ± 1.1          | $F = 16.63, p < 0.001$   |
| Prepatent period in days    | 55.7 ± 6.5       | 57.1 ± 8.3       | 43.3 ± 5.2         | $F = 4.42, p < 0.05$     |
| Patent period in days       | 11.2 ± 4.7       | 16.1 ± 5.2       | 28.8 ± 7.3         | $F = 10.21, p < 0.001$   |
| Number of shed cercariae    | 80.3 ± 35.8      | 66.8 ± 21.2      | 157.1 ± 55.2       | $F = 7.10, p < 0.01$     |
| Cadavers of NCS snails:     |                  |                  |                    |                          |
| - Free rediae               | 0.5 ± 1.5        | 0.4 ± 1.1        | 0.2 ± 0.8          |                          |
| - Free cercariae            | 13 ± 0.7         | 15 ± 0.5         | 23 ± 0.5           | $F = 4.68, p < 0.05$     |
| Percentage of shed cercariae** | 58.3             | 43.3             | 66.4               |                          |

*Mean values are given with their standard deviations for seven parameters. Each statistical comparison was performed using values recorded for the three snail species. CS, cercariae-shedding snails; F, value of ANOVA; NCS, non-cercarial shedding snails containing cercariae; NS, non-significant difference; p, probability; p.e., post-exposure; $\chi^2$, value of the $\chi^2$ test.

**Number of cercariae shed by CS snails / overall cercarial production in NCS snails.
generations via a protocol already used by Sanabria et al. [27] for F. hepatica and Vignoles et al. [28] for Fascioloides magna. F2 snails originated from eggs laid by the 50 exposed individuals of the F1 generation between weeks 2 and 5 p.e. A similar protocol was used for the F3, F4 and F5 generations. This protocol was chosen in order that these descendants have a first (F2) or multiple contacts (the F3-F5 generations) with the parasite through their infected parents.

The aim of the first experiment was to determine the aptitude of L. fuscus and R. balthica as a snail host for F. hepatica. Four groups of 50 L. fuscus each (1 group per generation from F1 to F4) were constituted. Four groups of R. balthica were also constituted according to the same protocol. Each lymnaeid was routinely exposed to two miracidia in a 35-mm Petri dish for 4 hours at 20°C (in 3.5 mL spring water). The choice of two miracidia per snail was based on the fact that a single miracidium (out of two) only developed in 75% of G. truncatula after exposure [29,30]. Each Petri dish was controlled 4 hours after the introduction of miracidia to verify their absence in spring water and thereby confirm their penetration into the lymnaeid. Snails were then raised in groups of 50 individuals in 10-L aquaria according to the same protocol used for parents. At day 50 p.e., surviving snails were dissected under a stereomicroscope to detect the presence of F. hepatica larval forms within their bodies and determine the most developed stage (immature rediae, cercariae-containing rediae, or free cercariae). The existence of cercarial shedding was also taken into account. Infected snails were then counted in relation to snail generation and each developmental stage of larval development.

As cercarial F. hepatica shedding (Table 1) was noted from F3 (R. balthica) or F4 (L. fuscus) generations, a second experiment was carried out to study cercarial shedding in the F5 generations and compare it with that occurring in G. truncatula. One hundred L. fuscus, 100 R. balthica (both belonging to the F5 generation) and 100 G. truncatula were used. Exposure of the three lymnaeids to miracidia and breeding of L. fuscus and R. balthica during the first 30 days were similar to those used in the first experiment. In contrast, the G. truncatula were raised in groups of ten individuals in 14-cm Petri dishes (60 mL spring water per dish) for 30 days according to Rondelaud et al. [31]. Snail food for G. truncatula consisted of dried lettuce leaves and dead Molinia caerulea leaves, while stems of live Fontinalis sp. ensured oxygenation of the water layer. Dissolved calcium in spring water also was 35 mg/L. Petri dishes containing G. truncatula were placed in the same air-conditioned room as the aquaria with L. fuscus or R. balthica. At day 30 p.e., each surviving snail was isolated in a 50-mm Petri dish containing 10 mL spring water, with pieces of dead grass, lettuce and spring moss. These Petri dishes were also placed at 20°C. Spring water and food, if necessary, were changed daily until snail death. When the first cercarial shedding occurred, surviving snails were subjected to a thermal shock every three days by placing their Petri dishes at 10°-13°C for 3 hours to stimulate cercarial exit [27]. Shed cercariae were then counted and removed from Petri dishes. At the death of each infected snail, its shell was measured using callipers. Cadavers of non cercarial shedding (NCS) snails that contained cercariae were routinely dissected under a stereomicroscope to count free rediae and free cercariae.

Parameters studied and data analysis
The first two parameters were snail survival at day 30 p.e. and the prevalence of F. hepatica infection (calculated in relation to the number of snails surviving at day 30 p.e.). In the first experiment, prevalence took into account the numbers of snails with immature rediae only, cercariae-containing rediae, or with free cercariae. In the second experiment, prevalence was calculated using the numbers of cercariae-shedding (CS) and NCS snails. For each parameter, the differences were analyzed using a χ² test. In the second experiment, shell growth of CS and NCS snails during the experiment (calculated using the difference between shell heights at exposure and at snail death), length of the prepatent and patent periods, and the number of shed cercariae were also considered. Free rediae and free cercariae counted in NCS snail cadavers were also taken into account. Individual values recorded for these last six measurements were averaged and their standard deviations were established for each snail group. One-way analysis of variance (ANOVA) was used to establish levels of statistical significance. The different analyses were performed using Statview 5.0 software (SAS Institute Inc., Cary, NC, USA).

Results
Aptitude of Lymnaea fuscus and Radix balthica for Fasciola hepatica infection
Table 1 gives the numbers of infected snails in F1 to F4 generations of both lymnaeids (first experiment). Survival rates of L. fuscus at day 30 p.e. significantly increased (χ² = 12.66, p < 0.05) with increasing snail generation. The same finding was also noted for the number of snails containing live F. hepatica larval forms so that prevalence of infection was significantly higher (χ² = 12.59, p < 0.05) in the F4 than in the other generations. In the case of R. balthica, the differences between survival rates at day 30 p.e. were insignificant, whatever the mode of comparison. In contrast, overall prevalence of F. hepatica infection significantly increased (χ² = 10.29, p < 0.05) with increasing snail generation. Four successive generations of snails (L. fuscus) or three (R. balthica) generations, a sec-
balthica) were thus necessary to obtain the first cercarial shedding from an infected snail.

Characteristics of Fasciola hepatica infection

Table 2 gives the main characteristics of F. hepatica infection in the three lymnaeids (second experiment). Compared to G. truncatula, snail survival at day 30 p.e., prevalence of F. hepatica infection, the patent period, and the number of shed cercariae were significantly lower in L. fuscus and R. balthica, while the prepatent period was significantly longer. No significant difference between L. fuscus and R. balthica was found, whatever the above parameter considered. Compared to G. truncatula, the shell growths of CS and NCS snails during the experiment were significantly greater in L. fuscus and R. balthica. In NCS snails, free rediae and free cercariae of F. hepatica were significantly more numerous in G. truncatula than in the other two lymnaeids. If the number of shed cercariae noted in CS snails was compared to the quantity of free cercariae recorded in the bodies of NCS snails, the percentage of shed cercariae was greater in G. truncatula than in the other lymnaeids (66% vs 43-58%).

Discussion

In northern Correze, Creuse and Haute Vienne (central France), several lymnaeid species, i.e. L. fuscus, L. glabra, L. palustris and/or R. balthica, may live with G. truncatula along the same open drainage network in swampy meadows [20] or in the same wild watercress beds [32,33]. The presence of the former lymnaeids had raised a serious problem in several cattle-breeding farms known for high prevalence of fasciolosis when G. truncatula was lacking. Indeed, all the experimental infections carried out by our team with 4-mm high L. fuscus was lacking. Indeed, all the experimental infections of juvenile snails [5,8,9,34-36] were in the past 40 years were negative. On the other hand, successful infections of preadult snails and this raises again the problem of the responsible definitive host. Indeed, local cattle present in these farms were treated twice a year with anthelmintics (triclabendazole since 1990–1994) and regularly moved on new pastures for grazing. The role of local lagomorphs cannot be excluded in fasciolosis transmission because hares and rabbits were often found to be infected with F. hepatica in northern Haute Vienne [37].

The results noted for L. fuscus in the present study cannot be compared to the literature because this species had often been synonymized with L. palustris in the past [38], and experimental infections of L. fuscus with F. hepatica were only successful for juveniles measuring less than 2 mm at miracidial exposure [8,39]. Contrary to L. fuscus, several reports [3,8,40-43] described the successful infection of Radix species other than R. auricularia when juveniles were experimentally exposed to F. hepatica. But identification of these Radix species using morphological criteria was often difficult to do and was currently based on the analysis of 18S, ITS-1, ITS-2 and/or COI sequences [44-49]. As a consequence, it is difficult to compare the above results if the Radix populations concerned by these studies are not identified using DNA-based analyses. As the names of these Radix species had also changed from the 2000s [44,45], the only paper mentioning the successful infection of R. balthica (=R. peregra O.F. Müller, 1774; = R. ovata Draparnaud, 1805) and R. labiata (=R. peregra sensu Ehrmann, 1933) was reported in Belgium by Caron et al. [50,51].

According to these authors, juveniles of both species, once exposed to individual bimacridial infections with F. hepatica, could sustain complete larval development of the parasite with cercarial emergence (R. labiata) or without shedding (R. balthica).

In view of this situation mainly due to the above-mentioned taxonomic problems within the Lymnaeidae family, the results noted for L. fuscus and R. balthica can only be compared to those reported by Rondelaud et al. [19] for two populations of L. glabra infected with F. hepatica and raised according to the same protocol. In the three lymnaeids, prevalence of infection significantly increased with increasing generation of snails, even if the speed of this process over time varied with each lymnaeid species or each population in the case of L. glabra. A similar finding can also be noted for the intensity of F. hepatica infection, expressed by the number of shed cercariae in CS snails or the quantities of free rediae and free cercariae noted in dissected NCS snails. As the values of these last parameters were significantly lower in the F5 generation of L. fuscus and R. balthica (the present study) or the F7 generation of L. glabra [19] than in G. truncatula, this suggests a progressive adaptation between each lymnaeid and the parasite though the infection of several successive snail generations, as suggested by Boray [3] in his review. However, the present study does not allow us to determine the real level of L. fuscus and R. balthica susceptibility to the parasite...
because our infections were only carried out through five snail generations and the answer to this question would need the infection of a greater number of generations. Two perhaps complementary hypotheses may be proposed to comment on these changes in successive generations of infected L. fuscus and R. balthica. The first would be to admit decreased genetic variation in host resistance through selection of snails with lowest resistance and/or a series of bottlenecks during the five successive generations of snails. This first hypothesis is based on the protocol used for the two lynmaeids, with the selection of offspring coming from already infected parents. However, another hypothesis, based on a progressive decrease in the number of the snail ganglionic neurons through the successive generations of infected snails originating from already infected parents, cannot be excluded. This interpretation is supported with the report by Szymid-Adjidé et al. [52] on the numbers of neurons in the model G. truncatula/F. hepatica. According to these authors, many neurons in the dorsal lobes of cerebrod ganglia and in pedal ganglia of G. truncatula disappeared during snail infection and lost their function. If this neuronal death also exists in the corresponding ganglia of infected L. fuscus and R. balthica, the surviving neurons would gradually secrete a lower quantity of neuropeptides [53] through successive generations of infected snails. As a consequence, this decrease could induce a progressive change in the mechanisms of the snail immune system response to F. hepatica and also a progressive lifting of bottleneck exerted by the snail on larval development of the parasite.

Conclusion

The F. hepatica infection of several successive generations of snails, coming from parents infected with this parasite, resulted in a progressive increase in prevalence and intensity of snail infection. This mode of snail infection may explain high prevalence of fasciolosis noted in several cattle-breeding farms when the common snail host of this digenean, G. truncatula, was lacking.

Competing interests

The authors declare that they have no competing interests.

Authors’ contribution

DR, AT and AM designed the study, participated in field collection, performed all laboratory work and wrote main parts of the manuscript. PV made statistical analyses. GD conceived the study and contributed to the interpretation of data and writing of the manuscript. All authors have read and approved the final manuscript.

Acknowledgements

The authors gratefully thank Dr. F.F. Djukwo Teukeng for her help during experimental infections of snails and Dr. J. Cook-Moreau for revising the English text.

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Cite this article as: Rondelaud et al.: Adaptation of Lymnaea fuscus and Radix bathica to Fasciola hepatica through the experimental infection of several successive snail generations. Parasites & Vectors 2014 7:296.

doi:10.1186/1756-3305-7-296

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