Control of freshwater fish louse *Argulus coregoni*: a step towards an integrated management strategy

Teija Hakalahti-Sirén¹*, Viktor N. Mikheev², E. Tellervo Valtonen¹

¹Department of Biological and Environmental Science, PO Box 35 (ya), University of Jyväskylä, 40014 Jyväskylä, Finland
²A.N. Severtsov Institute of Ecology & Evolution, Russian Academy of Sciences, 33 Leninskii Prospekt, 119071 Moscow, Russia

ABSTRACT: Harmful infections by ectoparasites of the genus *Argulus* occur repeatedly in freshwater fish farming operations where the management has largely been ineffective. Preventative methods and regular monitoring are rarely applied, so that chemical interventions become necessary. According to the Integrated Pest Management (IPM) approach, a sustainable management or control program for a parasite should be based on knowledge of the ecology of the parasite along with adoption of several prevention and control methods, the application of which is dependent upon the prevailing infection level. The application of multiple management tactics is especially important because parasites can develop resistance to chemical treatments. We took a step towards sustainable management of *Argulus* populations and tested the effect of several types of treatments on survival of *A. coregoni* at different stages in its life cycle. Parasite juveniles and adults were highly sensitive to potassium permanganate treatments (0.01 g l⁻¹), which lead to 100% mortality, whereas treatments with formalin (0.6 ml l⁻¹), sodium chloride (20 g l⁻¹) or malachite-green/formalin were not effective. Mechanical treatment by shaking infected fish in a hand net was an effective means of detaching parasites from the fish, and resulted in >80% decreases in parasite numbers. Compared to eggs in control treatments, both drying over a minimum period of 24 h and formalin treatments (120 ml l⁻¹) led to significantly higher mortality of *A. coregoni* eggs. Other treatments, i.e. drying over a period of 15 h, baths in potassium permanganate (1 g l⁻¹) or sodium chloride (50 g l⁻¹), did not significantly affect the viability of eggs. Based on the present results and previously published papers, we present an initial framework showing how *A. coregoni* populations could be managed effectively.

KEY WORDS: Integrated pest management · IPM · Control · Prevention · Parasite · *Argulus coregoni* · Fish louse

INTRODUCTION

Intensification of aquaculture has lead to the development of favourable conditions for a variety of fish disease-causing agents (Rintamäki et al. 1994, Rintamäki-Kinnunen & Valtonen 1997, Suomalainen et al. 2005, Karvonen et al. 2006), crustacean ectoparasites among others (Menezes et al. 1990, Hakalahti & Valtonen 2003, Costello 2006). Parasitic crustaceans are epithelial browsers that harm the protective epithelium of the fish and evoke stress responses by attaching and feeding on fish skin or gills. The epithelial damage and other stressor-induced changes in fish physiology increase fish susceptibility to secondary diseases (see Mustafa et al. 2000, Tully & Nolan 2002, Bandilla et al. 2006). Therefore, fish parasitized by ectoparasites often have a simultaneous fungal or bacterial infection (Cusack & Cone 1986, Bakke & Harris 1998, Bandilla et al. 2006). At commercial farms, ectoparasitic infections of fish result in stunted growth, reduced survival and increased production costs (Menezes et al. 1990, Costello 2006).

Fish farmers combat crustacean ectoparasites using a variety of methods that are either laborious, expen-
sive or both, and do not always prevent economic losses. The majority of management protocols have relied on the use of chemicals either as oral medications or bath treatments (e.g. Costello 1993, 2006). For example, the in-feed medication SLICE® (emamectin benzoate; Schering-Plough Animal Health) has been used to control many crustaceans, e.g. *Lepeophtheirus salmonis* in seawater aquaculture, *Salmincola edwardii* and *Argulus coregoni* in freshwater aquaculture (Stone et al. 2000, Duston & Cusack 2002, Hakalahti et al. 2004a). Alternative methods are, for example, following farm sites between stockings (Bron et al. 1993), use of cleaner fishes (Treasurer 2002) and removal of specific life-cycle stages from the parasite populations (Gault et al. 2002, Hakalahti et al. 2004b). The emergence of reduced sensitivity to chemical treatments with increased doses applied to control parasites, however, has lead to concerns about the sustainability of the industry (e.g. Jones et al. 1992), and consequently to the adoption of the Integrated Pest Management (IPM) concept into aquaculture (e.g. Sommerville 1998, Mordue (Luntz) & Pike 2002). An IPM program is typically based on several practises including prevention of infections, monitoring infection levels, using thresholds for action and finally implementing different types of control tactics. The IPM is credited with better and longer-term control efficacy due to decreased potential for the evolution of resistance to chemicals (Hoy 1998). IPM-based practises have been developed and applied in the control of marine parasitic copepods such as *Lepeophtheirus salmonis* (Sommerville 1998, Mordue (Luntz) & Pike 2002); however, studies on management of crustaceans in freshwaters are rare.

Fish lice of the genus *Argulus* cause frequent problems for northern inland fish farming (Shimura 1983, Hakalahti & Valtonen 2003, Taylor et al. 2006). *A. foliaceus* (Linné 1758) is a common generalist species in Finland (Mikheev et al. 1998), whereas *A. coregoni* (Thorell 1865) occurs primarily on salmonids, and occasionally infects other fish genera such as cyprinids (Shimura 1983, Pasternak et al. 2004). The present study concentrated mainly on *A. coregoni*. This parasite has a direct life cycle including eggs, free-living infective larvae, and juvenile and adult stages on fish (Shimura 1981). In Finland, the larvae typically emerge in May from over-wintered *A. coregoni* eggs (Hakalahti & Valtonen 2003, 2006). Although strongly affected by water temperatures, the life cycle of *A. coregoni* is typically completed within 1.5 mo at Finnish summer temperatures (Hakalahti & Valtonen 2003, Hakalahti et al. 2006). Females deposit their eggs on hard substrata such as stones and then they die (Mikheev et al. 2001, Hakalahti et al. 2004b). The negative impact of *Argulus* spp. for a fish is a combined effect of parasitic feeding and secondary infections, either from opportunistic microbes present in the water (Singhal et al. 1990, Rahman 1996, Bandilla et al. 2006) or from pathogens spreading among host-switching parasite individuals (Cusack & Cone 1986, Bandilla et al. 2008).

The objective of the present study was to create a framework for sustainable management of *Argulus coregoni* populations using components of the IPM concept. Some fish farmers in Finland have used the in-feed medication SLICE® to reduce *Argulus* levels (Hakalahti et al. 2004a). Simulations of a recent population dynamic model of *A. coregoni*, however, show that the parasite ‘egg-bank’ consisting of eggs with variable timings of emergence over several years (see Hakalahti et al. 2004c) acts as an effective buffer against such control practises (Fenton et al. 2006). Therefore, repeated SLICE® treatments are needed to suppress an outbreak, creating a potential for resistance development. Rapid development of resistance to the insecticide lindane among exposed *Argulus* populations has already been demonstrated (Lahav et al. 1962). An alternative method comprising removal and destruction of any eggs laid on submerged artificial egg-laying substrata has been tested (Hakalahti et al. 2004b), but not yet been applied at operational farms. Thus, there is an urgent need to further develop management of *Argulus* populations (see also Taylor et al. 2006). The specific aims of the present study were to: (1) test several types of methods (chemical, biological and mechanical) to prevent and control *A. coregoni* infestations, (2) pinpoint the most vulnerable stages in the parasite life cycle against which the specific control attempts could be targeted, and (3) based on the present results and previously published work, create a framework for sustainable management of *A. coregoni* populations.

The chemicals used were sodium chloride, formalin, potassium permanganate and a malachite-green/formalin mixture, all of which are or have been common treatments against ectoparasites in Finnish fish farming (Rintamäki et al. 1994, Rintamäki-Kinnunen & Valtonen 1997). Although the use of malachite-green is banned in most European countries (from 1 October 2001 onwards in Finland), it was included owing to its traditional and most effective use as a bath treatment against ectoparasitic ciliate and fungal infections such by *Ichthyophthirius multifiliis* and *Saprolegnia* spp., respectively (Willoughby & Robert 1992, Rintamäki-Kinnunen et al. 2005a). If malachite-green/formalin has a negative effect on attached fish lice, the ban on its use may explain the increase in *Argulus* problems in the current century. In addition to chemical procedures, drying was also tested as a method to eradicate *A. coregoni* eggs. Mechanical handling of fish has

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been suggested by fish farmers to have some utility in dislodging attached ectoparasitic crustaceans. This was tested in an experiment performed under farming conditions where infected fish were shaken in hand nets.

**MATERIALS AND METHODS**

**Selection of effective chemicals for control of parasites attached to fish.** The effects of 4 different chemical treatments were tested on survival of *Argulus coregoni* under laboratory conditions. These chemicals have traditionally been employed at Finnish fish farms for managing various ectoparasites (Rintamäki et al. 1994, Rintamäki-Kinnunen & Valtonen 1997). Two concentrations of each chemical were tested; dosing choices were based on the treatment ranges used in Finnish fish farming (e.g. Rintamäki et al. 1994, Rintamäki-Kinnunen & Valtonen 1997, Rintamäki-Kinnunen et al. 2005a,b). The chemicals and their concentrations were: (1) formalin 0.6 and 0.4 ml l\(^{-1}\), (2) salt (NaCl) 20 and 15 g l\(^{-1}\), (3) potassium permanganate (KMnO\(_4\)) 1.2 and 0.9 g l\(^{-1}\), and (4) malachite green/formalin (3.7 g of malachite green mixed in 1 l formalin) 0.025 and 0.0125 ml l\(^{-1}\). Filtered lake water was used in the controls, giving 9 treatments in total. Each treatment was replicated 4 times. Each experiment was repeated with 2 different juveniles and one adult *A. coregoni* stage. Their mean body lengths, which were measured on formalin-preserved individuals (n = 360) after the experiments, were 1.1 ± 0.13 (±SD), 2.9 ± 0.40 and 7.5 ± 0.40 mm, respectively.

The experiments were performed in plastic containers each filled with 250 ml of freshly prepared chemical solution and 10 parasites. Parasites needed in the experiments were collected from rainbow trout *Oncorhynchus mykiss* (30 to 40 cm total length) at a fish farm. To collect the parasites, captured fish were anesthetized with MS-222 (3-aminobenzoic acid ethyl ester). Parasites collected were maintained and monitored in an aerated aquarium over 1 d (temperature 21 ± 1°C), and only parasites that were swimming actively and apparently in good condition were used in the experiments. At the start of each trial, parasites were randomly picked out from the holding aquarium and introduced to experimental containers using pipettes filled with a minimum amount of water (to reduce dilution of the chemicals). Parasite survival in each container was monitored under laboratory conditions (temperature 21 ± 1°C) by counting and collecting dead parasites after 15 min, 30 min, 1 h, 2 h, 4 h and 16 h from the start of the experiment. Parasite deaths were confirmed by gently prodding individuals with a needle.

Equalities of mean life times between replicate containers were tested with ANOVA; no differences were found between containers in any of the treatments (p > 0.05 in all cases). The data were checked for homogeneity of variances and for normality. Data were then analyzed with a nonparametric survival analysis whose survival functions were estimated using the Kaplan-Meier life-table method (Lee 1992).

**Minimum effective concentration and duration of potassium permanganate treatment.** Since potassium permanganate was found to be lethal to *Argulus coregoni* (see results of the experiment described in the section ‘Selection of effective chemicals for control of parasites attached to fish’), its minimum effective concentration was determined in further experiments. Treatments with potassium permanganate concentrations of 0.01, 0.005, 0.001 and 0.0001 g l\(^{-1}\) were applied. The experimental set-up was similar to the experiment described in the previous section; 4 different *A. coregoni* developmental stages were used and the number of replicate containers was 3. Parasite mortality was followed more often: after 15 min, 30 min, 1 h, 2 h, 4 h, 5 h, 7 h, 16 h and 24 h from the start of the experiment. The average body lengths of the parasites (n = 150) in the experiments were 0.8 ± 0.2 (±SD), 2.3 ± 0.3, 6.6 ± 0.5 and 10.3 ± 0.6 mm. Filtered lake water was used in controls. Since potassium permanganate concentrations of 0.001 and 0.0001 g l\(^{-1}\) did not affect parasite survival, additional concentrations of 0.004, 0.003 and 0.002 g l\(^{-1}\) were tested on parasites in size classes 6.6 and 10.3 mm. Equalities of mean life times between replicate containers were tested with ANOVA; no differences were found between the containers in any treatment (p > 0.05 in all cases). The data were checked for homogeneity of variances and for normality. Data were then analyzed with Cox regression analysis using parasite size class and potassium permanganate concentration as covariates (Lee 1992).

The minimum duration of chemical treatment was determined at a potassium permanganate concentration of 0.01 g l\(^{-1}\) since this concentration was found to be the minimum concentration leading to 100% mortality of exposed *Argulus coregoni*. Parasites were exposed to potassium permanganate as in previous experiments (4 replicate containers, 10 adult parasites in each), but after 30 min or 1 h from the start of the exposure, parasites were removed from the container with a pipette and transferred to clean water containers where their survival was followed over a period of 24 h. Controls (4 replicates) were kept in clean lake water throughout.

**Mechanical treatments.** Mechanical treatments of *Argulus*-infested fish were tested as a method for detaching and removing parasites. Vigorous, brief shakings (20 to 30 s) were given to fish using a hand
net in a water tank. The treatment was performed in 2 earth-lined fish ponds containing either A. coregoni-infected rainbow trout Oncorhynchus mykiss (mean fork length 35 cm ± 2.5 SD, n = 10) or brown trout Salmo trutta (13 cm ± 0.9, n = 14). Brown trout had concurrent infections with A. coregoni and A. folicaceus. Fish in each pond were treated differently; salted water (NaCl, 0.05 g l–1) was used in the treatment of rainbow trout and lake water was used in the treatment of brown trout. Hundreds of fish in both ponds were treated and were subsequently transferred to empty ponds. The experiment was performed under farming conditions by the fish farmer, which restricted our ability to implement the experimental design.

During the treatment, groups of fish consisting of 2 to 4 individuals were captured and shaken together in a hand net. After treatment, numbers of detached parasites in the water and the numbers of parasites remaining on the fish were counted and removed. To count attached parasites, fish were narcotized with MS-222. These samples were taken in total from 10 randomly chosen rainbow trout treated in salted water and from 14 brown trout treated in lake water, respectively. Further groups of 14 and 10 randomly chosen brown trout were individually captured from the pond immediately before treatment in lake water and a day following their treatment, respectively. Captured fish were narcotized (MS-222) and the number of Argulus coregoni and A. folicaceus were counted from each fish. Fish were released back to the pond after sampling.

To evaluate the selectivity of mechanical treatment by parasite sex and size, samples of Argulus coregoni removed from rainbow trout during the experiment (mechanical treatment in salted water) were preserved in 5% formalin for further analysis. The pre-treatment sample consisted of 2 A. coregoni infrapopulations (pooled sample from 2 fish), while the post-treatment sample consisted of 10 parasite infrapopulations totalling 409 and 426 fish lice, respectively. Parasite sex and body length, which was measured from the anterior edge of the carapace to the end of the parasite abdominal lobes, were determined under a dissecting microscope (see Shimura 1983).

Selection of methods for controlling parasite eggs. Effects of various treatments under laboratory conditions were tested on hatching success of Argulus coregoni eggs. Five different types of treatments were used consisting of 3 chemical treatments and various drying periods. The treatments were: (1) potassium permanganate (KMnO₄) 1 g l⁻¹, (2) salt (NaCl) 50 g l⁻¹, (3) formalin 120 ml l⁻¹, (4) formalin 120 ml l⁻¹ applied to dried eggs (15 h), and (5) drying periods of 15, 24, 48 and 96 h. Filtered lake water was used in controls and each treatment was replicated 5 times, giving 9 treatments in total. We tested the effect of each treatment on the viability of parasite eggs collected from a commercial fish farm on 25 April 1999 (Set A). As the drying period of 15 h did not significantly affect the hatching rate of A. coregoni eggs compared to the control, additional longer drying periods (24, 48 or 96 h) were tested on parasite eggs collected on 29 May 2001 (Set B). Separate control treatments were used for each set of eggs.

During the experiments, several randomly chosen stones with attached Argulus coregoni egg clutches were exposed to each chemical (freshly prepared with lake water) for over 15 h or were allowed to dry. After the treatment periods, stones were rinsed in lake water, and egg clutches were carefully scraped off the stones. Samples of eggs (about 5 cm² area filled with eggs) were taken from each stone and transferred to plastic containers filled with 200 ml of lake water. Water in each container was changed, and hatched larvae were removed and counted daily over a period of 9 d. During the experiments with Set A and B eggs, daily water temperatures in the experimental containers were 17.5°C (±0.5) and 21°C (±0.5), respectively.

RESULTS

Selection of effective chemicals for control of parasites attached to fish

Both concentrations of potassium permanganate tested (0.012 and 0.009 g l⁻¹) led to 100% mortality among Argulus coregoni within 16 h (Fig. 1). Mortality rates of parasites belonging to size groups 1.1 mm (log-rank test for potassium permanganate concentration 0.009 g l⁻¹: chi-squared test χ² = 75.61, p < 0.001), 2.9 mm (χ² = 75.64, p < 0.001) and 7.5 mm (χ² = 77.36, p < 0.001) were higher in potassium permanganate solutions than in controls (Fig. 1). A malachite green/formalin mixture at 0.025 ml l⁻¹ led to increased mortality rates only among parasites in size group of 2.9 mm (χ² = 10.01, p = 0.002, Fig. 1), whereas the effect of the lower concentration (0.0125 ml l⁻¹) was non-significant in this size class (χ² = 1.00, p = 0.317). Parasite survival in both sodium chloride and formalin solutions and in controls, was 100% (Fig. 1).

Minimum effective concentration and duration of potassium permanganate baths

Both potassium permanganate concentration (Cox regression: Wald = 309.99, p < 0.001) and parasite size (Wald = 95.68, p < 0.001) affected survival of Argulus
coregoni (Table 1). Parasite death rate and potassium permanganate concentration had a positive association (Fig. 2, Table 1). In general, the death rate was highest for juveniles (size class 2.3 mm); the risk of death in this size class was 3.6-fold higher than for adult parasites (size class 10.3 mm; Fig. 3, Table 1). The risk of death among exposed small parasite juveniles (size class 0.8 mm), however, was only 1.83-fold higher than for adult parasites (Fig. 3, Table 1).

All parasites that were exposed to potassium permanganate (0.01 g l–1) for over 30 min or 1 h periods of time and then transferred to clean water died within 24 h, whereas the survival of parasites in control containers kept in clean water was 100%.

**Mechanical treatments**

Eighty-one percent of Argulus coregoni were detached from rainbow trout by shaking fish (2 to 4 fish at a time) with a hand net in a salted water container. Similar treatments of brown trout in fresh water resulted in an 82.4% reduction in A. foliaceus and an 86.7% reduction in A. coregoni numbers immediately after the treatments. Note that statistical comparison between the methods is not possible, because different fish species were used. The day following the treatment (mechanical treatment in fresh water), A. foliaceus infection levels recorded on brown trout individually captured from the fish pond were lower than before the treatment (Mann-Whitney test: $U = 17.00, p = 0.002$, Fig. 4). The effect of the treatment on A. coregoni numbers on brown trout was non-significant ($U = 39.00, p = 0.051$), although a tendency towards decreased infection levels was apparent (Fig. 4).

Before the mechanical treatment (salted water) was applied, the Argulus coregoni sex ratio on rainbow trout was balanced (chi-square test: $n = 2312, \chi^2 =$

<table>
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Table 1. Argulus coregoni. Impacts of parasite size and potassium permanganate concentration on survival as assessed by Cox regression analysis. The parasite size class 10.3 mm and potassium permanganate concentration 0.01 g l–1 were used as reference categories for statistical comparisons.
0.42, df = 1, p = 0.517; Fig. 3), but became strongly male-biased after the treatment (n = 426, $\chi^2 = 31.04$, $p < 0.001$) (Fig. 5). Only 12.3% of initial number of females remained attached to fish, while 23.9% of males were left. Also, the average body lengths of female (Mann-Whitney test: $U = 11809$, $p < 0.001$) and male ($U = 21848$, $p < 0.001$) *A. coregoni* individuals attached to fish decreased as a result of the treatment. The average body lengths of attached *A. coregoni* females and males decreased from 9.1 ± 2.22 (±SD) to 8.1 ± 2.65 mm and from 9.5 ± 2.36 to 9.2 ± 2.01 mm, respectively.
Selection of methods for control of parasite eggs

Treatments affected the hatching success of Argulus coregoni eggs (Kruskal-Wallis test for Set A eggs: $\chi^2 = 26.106$, df = 5, $p < 0.001$; Kruskal-Wallis test for Set B eggs: $\chi^2 = 12.086$, df = 4, $p = 0.017$; Fig. 6). Compared to controls, a smaller proportion of eggs hatched from clutches that were treated with formalin either applied to eggs that had been allowed to dry over a period of 15 h or to moist eggs and from clutches that were allowed to dry over periods of 48 or 96 h (all pairwise comparisons: $p < 0.05$). A tendency for hatch suppression was also apparent among egg clutches allowed to dry over 24 h; however, the difference was non-significant ($p > 0.05$; Fig. 6). Drying over a period of 15 h, potassium permanganate or sodium chloride treatments did not affect the hatching success of parasite eggs (all pairwise comparisons: $p > 0.05$).

DISCUSSION

Modern fish culture is generally conducted under high stocking densities, a favourable environment for transmission and reproduction of parasites. High abundance of parasites inevitably leads to an arms-race between farmers and pathogen; farmers seek new control methods while treatments applied are selecting for tolerance and resistance among parasites (e.g. Lahav et al. 1962, Jones et al. 1992). Instead of using a limited set of chemical treatments against parasites, developing and implementing IPM programs would be a move towards more sustainable and cost-effective fish farming. According to principles of the IPM concept, a parasite control program should be based on an understanding of parasite ecology, and should rest on several pillars including (1) prevention of infections, (2) monitoring infection levels, (3) using thresholds for action, and finally (4) implementing various control tactics.

Prevention is fundamental to IPM and reduces the likelihood of a parasite becoming a problem. Fish culture practices are likely to have a key role in reducing and preventing problems caused by fish lice. Most importantly, destruction of parasite eggs in fish ponds before restocking of fish breaks the life cycle of the parasite by eliminating or reducing the infestations (see below). Prevention of contacts between farmed and wild fish is also an important preventative practice against fish lice, as it is known that parasitic infections of farmed fish are most typically derived from wild fish (Valtonen & Koskivaara 1994). An additional important
source of infection can be adjacent fish ponds that are infested with fish lice. Thus, self-contained water circulation in each pond may help to prevent the transmission of the parasite between farming sites. The infection can be transmitted via free-swimming louse stages that may live several days without a host (Hakalahti et al. 2005, Bandilla et al. 2008). Some other methods that have the potential to decrease the growth potential of *Argulus* populations include good fish husbandry leading to less stressed fish (see review on host-parasite interactions by Walker et al. 2004, Bandilla et al. 2005) and alteration of physico-chemical conditions of the farming system. Along with other aquatic parasites (see Chubb 1977), fish lice are greatly affected by abiotic factors in their environment. Several authors have demonstrated the importance of habitat requirements of different fish louse species also differ from each other; *A. coregoni* occurs in large lakes and rivers, mainly on salmonids, whereas *A. foliaceus* is a common generalist in eutrophic, slow-flowing or still waters (Kennedy 1974, Shimura 1983, Taylor et al. 2006).

A subsequent (though not separate) step in the implementation of an IPM program is the monitoring of parasite infection levels. In the case of *Argulus coregoni*, monitoring might involve direct and indirect means: field sampling to make counts of lice on individual fish or assessment of damage caused by the lice, and the use of trapping devices. As the transmission of *A. coregoni* to fish is strongly temperature dependent and only takes place during the ice-free period, the monitoring period is relatively short, but should start before the temperature exceeds 10°C in spring (Hakalahti & Valtonen 2003, Hakalahti et al. 2006). Habitat requirements of different fish louse species also differ from each other; *A. coregoni* occurs in large lakes and rivers, mainly on salmonids, whereas *A. foliaceus* is a common generalist in eutrophic, slow-flowing or still waters (Kennedy 1974, Shimura 1983, Taylor et al. 2006).

**Paraconsistent control of** *A. coregoni* **monitoring might involve direct and indirect means:** field sampling to make counts of lice on individual fish or assessment of damage caused by the lice, and the use of trapping devices. As the transmission of *A. coregoni* to fish is strongly temperature dependent and only takes place during the ice-free period, the monitoring period is relatively short, but should start before the temperature exceeds 10°C in spring (Hakalahti & Valtonen 2003, Hakalahti et al. 2006). We also recommend monitoring of parasite egg numbers, which can be undertaken by placing removable artificial egg laying substrata near the bottom of water ponds (Hakalahti et al. 2004b; see also below). Complete control of *A. coregoni* is neither necessary for high yields nor appropriate for IPM. Fish can tolerate a certain level of louse infestation without appreciable effects on vigor (Bandilla et al. 2006). However, in contradiction, there is a link between *Argulus* infections and susceptibility to certain harmful microbial diseases (e.g. Cusack & Cone 1986, Singhal et al. 1990, Bandilla et al. 2006) leading to a reduced acceptable parasite infection levels on fish. The economic threshold for fish louse infestations still needs to be determined, and is likely to vary between different types of culture systems. Rather than waiting for the damage to become apparent, application of control measures should supplement preventative measures when the predefined threshold for action has been reached. A control strategy may be based on a combination of chemical, mechanical, biological and cultural treatments.

Mathematical host-parasite population models can provide insights into the relative importance of factors influencing parasite population dynamics, and can be utilized in the development of effective and targeted control against the parasites (Hudson et al. 2002). Recently, a model based on detailed studies of the life cycle of *Argulus coregoni* was applied to simulate its population dynamics in Finnish fish farming conditions (Fenton et al. 2006). The model showed that *A. coregoni* infection level largely relies on the number of eggs in the egg-bank (see Hakalahti et al. 2004c concerning extended hatching of parasite eggs), and destroying eggs with all available means would greatly reduce the size of the parasite population (Fenton et al. 2006). The model showed that the next most effective control of *A. coregoni* may be achieved by targeting juvenile stages attached to fish (Fenton et al. 2006).

Draining and drying of egg laying sites of *Argulus* spp. has been suggested as a useful method for controlling eggs of the parasite (Bauer 1962). Some authors (Chen 1933, Hoffman 1980) recommend that drying should be accompanied by lime application. We quantified the hatching success of *A. coregoni* eggs in relation to drying period and determined a minimum period of 48 h needed to eradicate parasite eggs. A high mortality rate was also achieved by exposing the eggs directly to formalin. As complete dehydration of earth-lined ponds is time-consuming and uncertain, some fish farmers in Finland have covered the bottoms of ponds with PVC tarpaulins to facilitate drying and cleaning. Another useful method might be freezing of drained ponds during winter (T. Hakalahti-Sirén unpubl. data). In recent years, there has been increasing interest in using behaviour manipulation as a management tactic against crustacean parasites, and promising results have emerged from experiments in which artificial substrata were used to attract *A. foliaceus* and *A. coregoni* females and to trap their eggs (Gault et al. 2002, Hakalahti et al. 2004b). As the present results show, collected eggs can be effectively destroyed with minimal environmental impact simply by allowing the substrata to dry out. The eggs may also be destroyed after dislodgement with a brush or pressurized water.

Based on literature, currently used control methods against *Argulus* populations are largely based on the use of limited variety of chemicals, for example potassium permanganate, organophosphates and emamectin benzoate (e.g. Kabata 1985, Hakalahti et al. 2004a, Toksen 2006). The harmful effect of organophosphates to the environment is now widely recog-
nized, and many countries prohibit their use. In contrast, use of the in-feed treatment formulation SLICE® (in which emamectin benzoate is the active ingredient) is becoming more widespread. It has proven a very effective way to eliminate *A. coregoni* attached to fish (Hakalahti et al. 2004a), but its long-term usefulness is unknown due to a potential for tolerance development. Therefore, the medicine should only be used in response to heavy infestations after other types of controls have failed. However, of all chemicals (sodium chloride, formalin, potassium permanganate and malachite-green/formalin mixture) that were tested in the present study as means to kill parasites, only potassium permanganate was found useful.

The minimum dose of potassium permanganate which led to 100% mortality among exposed *Argulus coregoni* was 10 mg l\(^{-1}\) over 30 min. In previous studies a potassium permanganate concentration 1.5 mg l\(^{-1}\) applied as 2 subsequent treatments in 3 d was found to be effective in controlling *A. indicus* and *A. japonicus* infections on carp (Jafri & Ahmed 1994), and a concentration 0.5 mg l\(^{-1}\) resulted in a 60% decrease in the level of *A. indicus* infection (Singhal et al. 1986). In the present study, only treatments with potassium permanganate concentrations >1 mg l\(^{-1}\) significantly increased mortality of *A. coregoni*. Some studies have also recommended potassium permanganate concentrations as high as 1 g l\(^{-1}\) over 5 to 10 min for controlling of *Argulus* populations (Christensen 1989). The high variation in recommended dosing with potassium permanganate reflects its variability by water quality as an effective anti-parasite treatment, a property that also makes it a difficult and potentially deleterious treatment to use. Potassium permanganate functions as a strong oxidizing agent that reacts with any material it comes into contact with. Thus, the efficacy of the chemical is related to the amount of reducing substances (e.g. algae, detritus, inorganic reducing substances) in the water, and the amount of permanganate that is quickly reduced to manganese dioxide is called the potassium permanganate demand of the water (see Tucker 1989, Boyd 1990). The efficient use of potassium permanganate requires determination of water potassium permanganate demand before treatment. The oxidizing activity of potassium permanganate is also the primary problem for treated fish and, therefore, a careful choice of the treatment dose is essential to prevent fish mortality. It is advisable for a fish farmer to run a bioassay before treating large numbers of fish with potassium permanganate, as fish tolerance is dependent on many factors such as water pH, temperature, exposure time and the fish species (Marking & Bills 1975).

The present ban to use malachite green in aquaculture does not seem to explain increased incidences of *Argulus* spp. infections since the malachite green/formalin mixture was not sufficiently effective in killing *A. coregoni*. Other chemicals previously used as bath treatments against argulids include salt (NaCl) and formalin (Kabata 1985). Fish farmers in Finland generally use 10 to 20 g l\(^{-1}\) sodium chloride solution over a period of 20 to 60 min against *Chilodonella* ciliates on fish (Rintamäki et al. 1994). According to Singhal et al. (1986), dipping of infested fish in 30 mg l\(^{-1}\) sodium chloride for 2 to 5 min is very effective treatment against *A. indicus*. However, in the present study, 20 g l\(^{-1}\) salt bath over a period of 24 h did not affect survival of *A. coregoni*. Also, formalin bath treatments (0.12 ml l\(^{-1}\)) are used extensively to control some external protozoan infections of fish (Jørgensen & Buchmann 2007). In the present study, a relatively high formalin concentration (0.6 ml l\(^{-1}\)) did not increase mortality of *A. coregoni* juveniles or adults.

Surprisingly high numbers of *Argulus coregoni* and *A. foliaceus* (>80% of original) were dislodged from fish through shaking them in hand nets in a water container. The detachment rate of *A. coregoni* depended on parasite sex and size, dislodgement rates of gravid females being especially pronounced. Detachment of females may explain the difference in dislodgement between the sexes, because remaining on a host during disturbance is likely to be risky for breeding females. In addition, many of the females were ready to lay their eggs on the bottoms of ponds (Mikheev et al. 2001, Hakalahti et al. 2004b). The observed difference by parasite sex may also be explained by weight differences; egg carrying females sank faster than males of the same body length (authors’ unpubl. data). Application of mechanical treatments, however, is not an advisable response to low fish louse infection levels as they would elicit stress reactions and scale loss in fish, increasing their susceptibility to other disease-causing agents (e.g. Pickering & Pottinger 1989). Nevertheless, on a small scale, shaking could be used in response to heavy infections, e.g. when other treatment options are limited or when done in connection with normal fish husbandry practices (fish grading, vaccination, transfer etc.) and in conformity with animal welfare guidelines. When infested fish are treated mechanically or transferred between aquaculture units, the water must be collected and sieved or treated in order to kill parasites. To our knowledge, no other biological methods (e.g. cleaner fishes; see Bannilal et al. 2008 concerning predation by fish on *A. coregoni*) besides behavior manipulation and desiccation of eggs have been applied to reduce *Argulus* numbers in aquaculture. Cultural control methods could include fallowing, i.e. maintenance of host-free seasons, which certainly would diminish the parasite population size, but not necessarily break the life cycle of the parasite.
due to diapausing eggs (see Mikheev et al. 2001, Hakalahti et al. 2004c).

Here, we present a framework for sustainable management of *Argulus coregoni* populations. Financial losses caused by the fish lice would be best tackled by implementing preventative control methods. However, to make this applicable, further investigation is required to establish factors affecting the growth potential of fish louse populations. Another essential part of managing *Argulus* populations would be the regular monitoring of louse infection levels. When the threshold level for louse infection has been reached, intervention by applying control treatments is needed to prevent extensive damage caused by the parasite. The use of multiple tactics and rotation of treatments is a preferable and, in most cases, also a possible option to prevent extensive damage caused by the parasite. Such an approach would mitigate the development of parasite tolerance towards chemical and medical treatments (e.g. SLICE®, emamectin benzoate) currently available for the control of fish lice. The present experiments showed that *A. coregoni* can be controlled effectively by draining and drying of parasite egg laying sites. If not possible, another effective means was treatment of eggs with formalin. The experiments also showed that parasite juveniles and adults were susceptible to treatments with potassium permanganate. Shaking fish in a hand net also resulted in large decreases in the numbers of mature fish lice on fish.

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