DNA-based methods for freshwater biodiversity conservation

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Phylogeographic analysis of noble crayfish (Astacus astacus) and new insights into the distribution of crayfish plague

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- This thesis is dedicated to my grandparents -



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ABSTRACT

Worldwide one third to one half of the freshwater crayfish species are threatened with population decline or extinction. Besides habitat deterioration, pollution, and other man-made environmental changes, invasive species and pathogens are major threats to the survival of European crayfish species. Freshwater crayfish are the largest freshwater invertebrates and strongly influence the structure of food webs. The disappearance of crayfish from a water body may change the food web and could have dramatic consequences for an ecosystem.

One goal in modern species conservation strategies is the conservation of genetic diversity, since genetic diversity is an advantage for the long-term survival of a species. The main aim of my thesis was to reveal the genetic structure and to identify genetic hotspots of the endangered noble crayfish (*Astacus astacus*) throughout Europe (part 1 of my thesis). Since the most significant threat to biodiversity of European crayfish species is the crayfish plague pathogen *Aphanomyces astaci* I studied new aspects in the distribution of *A. astaci* (part two of my thesis). The results serve as a basis for future conservation programs for freshwater crayfish.

In the first part of my thesis I conducted a phylogeographic analysis of noble crayfish using mitochondrial DNA and nuclear microsatellite data. With these methods I aimed to identify its genetic hotspots and to reconstruct the recolonization history of central Europe by this species. I detected high genetic diversities in southestern Europe indicating that noble crayfish outlasted the cold climate phases during the Pleistocene in this region (*Appendix 1*). Because of the high genetic diversity found there, southeastern Europe is of particular importance for the conservation of noble crayfish. The mitochondrial DNA analysis points to a bifurcated colonization process from the eastern Black Sea basin to a) the North Sea and to b) the Baltic Sea basin (*Appendix 2*). A second independent refugium that was localized on the Western Balkans did not contribute to the colonization of central Europe. Furthermore, I found that the

natural genetic structure is dissolved, probably due to the high human impact on the distribution of noble crayfish (e.g. artificial translocation).

In the second part of this thesis using real-time PCR I identified calico crayfish (Orconectes immunis) as the fourth North American crayfish species to be carrier of the agent of the crayfish plague (Appendix 3). Furthermore I detected the crayfish plague pathogen in American spiny-cheek crayfish (Orconectes limosus) and native narrow-clawed crayfish (Astacus leptodactylus) in the lower Danube in Romania (Appendix 4). The distribution of infected spinycheek crayfish poses a threat to the native biodiversity in southeastern Europe and shows the high invasion potential of this crayfish species. Moreover, I found that even the native narrow-clawed crayfish in the Danube Delta, about 970 km downstream of the current invasion front of American crayfish, is a carrier of A. astaci (Appendix 5). This finding is of high importance, as the native species do not seem to suffer from the infection. In Appendix 6 I elucidate demonstrate that the absence of the crayfish plague agent is the most likely explanation for the coexistence of populations of European and American crayfish in central Europe. In my thesis I show that the common assumption that all North American crayfish are carrier of A. astaci and that all native crayfish species die when infected with A. astaci does not hold true.

The studies presented in my thesis reveal new aspects that are crucial for native crayfish conservation:

- 1) The genetic diversity of noble crayfish is highest in southeastern Europe where noble crayfish outlasted the last glacial maximum in at least two different refugia.
- 2) Not all American crayfish populations are carrier of A. astaci and
- 3) not all Europen crayish populations die shortly after being infected with the crayfish plague pathogen.

To conserve native crayfish species and their (genetic) diversity in the long term, further introductions of American crayfish into European waters must be avoided. However, the introduction will only decrease if the commercial trade with non-indigenous crayfish species is prohibited.

ZUSAMMENFASSUNG

Weltweit sind ein Drittel bis die Hälfte der Flusskrebsarten von Populationsrückgang oder Aussterben bedroht. Neben einer Verschlechterung der Habitate, Umweltverschmutzung und anderen vom Menschen verursachten Umweltveränderungen stellen eingeschleppte exotische Arten und Krankheitserreger eine große Bedrohung für das Überleben europäischer Flusskrebsarten dar. Flusskrebse sind die größten Wirbellosen in limnischen Systemen und haben einen entsprechend großen Einfluss auf die Struktur der Nahrungsnetze. Das Verschwinden von Flusskrebsen aus einem Gewässer kann Nahrungsnetze verändern und somit dramatische Konsequenzen für ein Ökosystem zur Folge haben.

Ein Ziel im modernen Artenschutz ist die Erhaltung der genetischen Vielfalt. Eine hohe genetische Vielfalt ist für das langfristige Überleben einer Art von Vorteil. Das Hauptziel meiner Arbeit war es, die genetische Struktur des Edelkrebses (Astacus astacus) in seinem gefährdeten europäischen Verbreitungsgebiet zu untersuchen und die besonders schützenswerten genetische 'Hotspots' zu identifizieren (Teil 1 der Dissertation). Die größte Bedrohung für die Diversität europäischer Flusskrebsarten Krebspesterreger Aphanomyces astaci dar. Daher muss die Verbreitung des Krankheitserregers bei Schutzprogrammen beachtet werden. Im zweiten Teil der Dissertation untersuchte ich neue Aspekte der Verbreitung von A. astaci. Die Ergebnisse dienen als Grundlage für zukünftige Artenschutzprogramme für Flusskrebse.

Im ersten Teil dieser Arbeit führte ich eine phylogeographische Analyse der Edelkrebse durch, um genetische 'Hotspots' zu identifizieren und die nacheiszeitliche Wiederbesiedlung Zentraleuropas durch diese Art zu rekonstruieren. Mit mitochondrialer DNA und nuklearen Mikrosatelliten-Markern ermittelte ich eine hohe genetische Vielfalt in Südosteuropa, die darauf hinweist, dass der Edelkrebs die kalten Klimaphasen des Pleistozäns in diesem Gebiet überdauerte (*Appendix 1*). Wegen der hohen genetischen Vielfalt ist Südosteuropa von besonderer Bedeutung für den Schutz des Edelkrebses. Die

mitochondriale DNA-Analyse deutet auf eine gegabelte Kolonisierung vom unteren Donaueinzugsgebiet in a) das Einzugsgebiet der Nordsee und b) das Einzugsgebiet der Ostsee hin (Kapitel 2). Ein zweites, unabhängiges Refugium, welches im westlichen Balkan lokalisiert wurde, hat vermutlich nicht zur Besiedlung Mitteleuropas beigetragen. Außerdem stellte ich fest, dass die natürliche genetische Struktur teilweise überlagert ist, wahrscheinlich aufgrund des hohen menschlichen Einflusses auf die Verbreitung des Edelkrebses (bspw. künstliche Translokation).

Im zweite Teil dieser Arbeit konnte ich mittels real-time-PCR ermitteln, dass bekannten drei nordamerikanischen Flusskrebsarten Kalikokrebse (Orconectes immunis) Träger des Krebspesterregers sind (Kapitel 3). Des Weiteren habe ich den Krebspesterrreger in der unteren Donau in Rumänien an amerikanischen Kamberkrebsen (Orconectes limosus) und europäischen Galizierkrebsen (Astacus leptodactylus) nachweisen können (Kapitel 4). Die Ausbreitung der infizierten Kamberkrebse bis in die untere Donau stellt eine große Bedrohung für die Artenvielfalt in Südosteuropa dar und zeigt das hohe Invasionspotential der Kamberkrebse. Darüber hinaus stellte ich fest, dass auch einheimische Galizierkrebse im Donaudelta, etwa 970 km hinter der aktuellen Invasionsfront des Kamberkrebses, Träger von A. astaci sind (Kapitel 5). Diese Erkenntnis ist von besonderer Bedeutung, da die einheimischen Arten offenbar nicht an der Infektion leiden. Die Untersuchung koexistierender Populationen europäischer und amerikanischer Flusskrebse ergab, dass die Abwesenheit des Krebspesterregers in diesen Populationen die wahrscheinlichste Erklärung für die erfolgreiche Koexistenz in den untersuchten Gewässern in Mitteleuropa ist (Kapitel 6).

Die Ergebnisse meiner Dissertation zeigen neue Aspekte, die von hoher Relevanz für den Schutz und Erhalt einheimischer Flusskrebsarten und deren genetischer Vielfalt sind:

Die genetische Diversität des Edelkrebses ist in Südosteuropa am höchsten.
 Dort überdauerten Edelkrebse die letzte Eiszeit in mindestens zwei unabhängigen Refugien.

- 2) Nicht alle amerikanischen Flusskrebspopulationen sind Träger der Krebspest und
- 3) nicht alle europäischen Flusskrebspopulationen sterben innerhalb kurzer Zeit an einer Infizierung mit dem Krebspesterreger.

Um einheimische Flusskrebse und deren (genetische) Vielfalt langfristig zu erhalten, dürfen keine weiteren amerikanischen Flusskrebse in der Natur ausgesetzt werden. Das unbefugte Aussetzen wird jedoch erst zurückgehen, wenn der Handel mit exotischen Flusskrebsen verboten wird.

ABBEREVIATIONS

CBD = Convention on Biological Diversity

COI = cytochrome oxidase subunit I

DNA = deoxyribonucleic acid

ESU = evolutionary significant unit

 H_D = haplotype diversity

IUCN = International Union for Conservation of Nature

MGB = minor groove binder

mtDNA = mitochondrial DNA

N = number of samples

 N_+ = number of positive samples

NICS non-indigenous crayfish species

PCR = polymerase-chain reaction

qPCR = real-time polymerase-chain reaction

rRNA = ribosomal ribonucleic acid

YBP = years before present

GENERAL INTRODUCTION

Conservation of biological diversity

In the Convention on Biological Diversity (CBD) negotiated at the United Nations Conference on Environment and Development in 1992, the conservation of biological diversity is highlighted as one of the three main goals (SCBD 1992). Biological diversity is understood as the variability among living organisms from all sources including diversity within and between species and also of ecosystems (CBD, Article 2). To maintain the genetic diversity within one species, conservation genetics aim to protect genetic variability within and between populations. The guidelines for the protection of threatened species recommend the identification of evolutionary significant units (ESU, Ryder 1986). An ESU can be defined as a population or group of populations that is sufficiently differentiated from all the other members of this species and requires an individual management or conservation strategy (Moritz 1994). An ESU may contain edemic genetic variation. In restocking programs, it is therefore recommended to use donor populations from the same ESU whenever possible to conserve the local specificity and maintain the maximum within-species diversity. Species with greater genetic diversity are more likely to be able to evolve in response to a changing environment than those with less diversity (Malcolm et al., 2007).

One of the most significant threats of biodiversity is the introduction of invasive species (McGeoch *et al.*, 2010) and diseases and parasites associated with them. Invasive species are often lacking natural predators and are competitors with the native species (Schulz *et al.*, 2006, Westman and Savolainen 2001). Although introduced species may increase the local biodiversity, they at the same time lead to a global homogenization and therefore to a loss of worldwide biodiversity (McNeely *et al.*, 2001). Compared to terrestrial ecosystems, aquatic ecosystems are particularly subject to invasive species (Dudgeon *et al.*, 2006). The connectivity of river systems facilitates the dispersal of invasive species over large distances. In addition, today artificial channels between

different rivers additionally increase the long-distance dispersal of freshwater species (Bij de Vaate *et al.*, 2002).

Otherwise, this connectivity between river systems also aided in the recolonization of the European continent by freshwater species after cold climate phases in the Pleistocene. When climate conditions were unfavorable, e.g. cold, the ranges of most European species contracted to southern Europe (Taberlet et al., 1998, Durand et al., 1999; Theissinger et al., 2012). Range contractions were followed by range expansions when the climate warmed again. The range expansion of freshwater species was most strongly influenced by the changes in landscapes through glaciers and melting water (e.g. Hänfling et al., 2009, Vonlanthen et al., 2007). This rapid northward expansion led to a reduction in genetic diversity of northern populations. In contrast, the varied topography of southern refugia allowed populations to diverge through several ice ages (Hewitt 1999). Thus, in species with a refugial area in southern Europe a genetic divergence between the ancestral populations in the South and the newly founded populations in the North of Europe can be detected as/in a decrease in genetic diversity (Reiland et al., 2002). For conservation management of a species it is important to reveal the two following aspects: 1) Where were the Pleistocene glacial refugia located? Because of their high genetic diversity, their identification is fundamental for conservation prioritization. 2) Where are possible ESUs? The reconstruction of the phylogeographical distribution aids in defining ESUs, which then serve as a basis for conservation programs. For freshwater species that often cannot migrate over terrestrial habitats, river catchments (e.g. Rhine river catchment, Danube river catchment) and sea basins (North Sea basin, Baltic Sea basin) play a significant role in subpopulational differentiation.

The freshwater crayfish

General

Freshwater crayfish are the largest freshwater invertebrates (Holdich 2002a). They occupy a variety of streams, rivers and lakes and prefer habitats with an access to shelter. Some crayfish species can have a physical impact on their environment due to their burrowing activity (Correia and Ferreira 1995). Crayfish are often key organism in food webs and have a high influence on its structure due to their omnivore behaviour (Nyström 1999). Besides on invertebrates, they feed on vertebrates like amphibians (i.e. on their eggs and larvae (Axelsson et al., 1997)) and on fish (Guan and Wiles 2002) and therefore have direct and indirect influence on the species composition of their respective ecosystems. Freshwater crayfish (Astacida) belong to the monophyletic Reptantia within the Decapoda (Scholz and Richter 1995) and are taxonomically distributed among three families, two in the northern Hemisphere (Astacidae and Cambaridae) and one in the southern Hemisphere (Parastacidae), specifically in Madagascar, southern South America and Australasia (Holdich 2002a). The Cambaridae are naturally found in the Eastern United States and in East Asia, whereas the Astacidae are distributed in the Western United States and in Europe. The diversity of crayfish in Europe is relatively low compared to that in Australia and North America. Freshwater crayfish have two centres of diversity, one in the southeastern USA and one in Victoria, Australia (Crandall et al., 2006). Today more than 460 crayfish species from North America are known (Crandall and Buhay 2011). In Europe, depending on the taxonomy, five or six species from two genera can be distinguished. Narrow-clawed crayfish (Astacus leptodactylus Eschscholz 1823), thick-clawed crayfish (Astacus pachypus Rathke 1837) and Noble crayfish (Astacus astacus Linnaeus 1758) belong to the genus Astacus and have a more northern and eastern distribution compared to stone crayfish (Austropotamobius torrentium Schrank 1803) and white-clawed crayfish (Austropotamobius pallipes Lereboullet 1858 species complex) which belong to the genus Austropotamobius. A. pallipes is considered a species complex and its taxonomic status is still under revision (Füreder et al., 2010). Schulz and Grandjean (2005) however argue that based on

genetic data this species complex consists of two separate species, *A. pallipes* and *Austropotamobius italicus*.

Today at least eight non-indigenous crayfish species (NICS) are established in the wild in Europe (Holdich *et al.*, 2009; Chucholl and Pfeiffer 2010). In the literature, a distinction is drawn between the 'Old NICS' and the 'New NICS'. The signal crayfish (*Pacifastacus leniusculus* Dana 1852), the spiny-cheek crayfish (*Orconectes limosus* Rafinesque 1817) and the red swamp crayfish (*Procambarus clarkii* Girard 1852) have been introduced before 1975 whereas the 'New NICS' have been introduced after 1980 (Holdich *et al.*, 2009). In this thesis I have investigated the crayfish plague carrier status of the 'Old NICS' spiny-cheek crayfish (Table 1 and *Appendices 3-6*) and of the 'New NICS' calico crayfish (*O. immunis* Hagen 1870) (*Appendix 3*).

Table 1 List of crayfish and crayfish pathogen species included in this thesis.

Common name	Scientific name	Conservation status	Distribution in Europe
Noble Crayfish	Astacus astacus	Vulnerable (IUCN), native	Central and eastern Europe
Narrow-clawed crayfish	Astacus leptodactylus	Least concern (IUCN), native	Native to the Ponto-Caspian regior today in most European countries
Spiny-cheek crayfish	Orconectes limosus	Invasive American species, 'Old NIC'	Western, central and eastern Europe
Calico crayfish	Orconectes immunis	Invasive American species, 'New NIC'	Upper Rhine river catchment in Germany and France
Crayfish plague	Aphanomyces astaci	Included in: 100 of the worst alien species	Throughout Europe

The noble crayfish (Astacus astacus)

While the stone crayfish and white-clawed crayfish species complex has already been subject to intensive molecular studies (e.g. Bertocchi *et al.*, 2008, Gouin *et al.*, 2006, Trontelj *et al.*, 2005), molecular studies of noble crayfish have only been focused on restricted geographical areas (Agerberg 1990, Fevolden *et al.*, 1994, Schulz 2000, Edsman *et al.*, 2002, Schulz *et al.*, 2004, Alaranta *et al.*, 2006). Therefore, a part of my thesis focuses on noble crayfish that is widely distributed across Europe. Its range extends from Russia and the Ukraine in the east, Scandinavia in the North, Greece in the South, and the United Kingdom and France in the west (Edsman *et al.*, 2010).

In the literature, different and partly contradicting taxonomic classifications of the noble crayfish (Figure 1) based on morphological criteria below species level can be found (see Smietana et al., 2006 for a review). The most widely accepted taxonomic studies (Karaman 1962, 1963) recognized the three following subspecies: A. a. colchicus, which can be found in an isolated area in the Ponto-Caspian region of the upper Rion (Caucasus); A. a. balcanicus, which populates the Western Balkans with the drainage system of the river Vardar in Macedonia and Greece and in the Lake Ohrid; the subspecies A. a. astacus with three races is distributed in the river catchment of the river Danube and the rivers that open out into the North- and Baltic Sea. Phylogenetic analysis of populations from the distribution area of the subspecies A. a. astacus is the subject of Appendices 1 and 2. Karaman (1962) named the three races of A. a. astacus according to their distribution: Northrace (A. a. a. astacus) in Elbe, Odra and probably in other rivers that open out into the North- and Baltic Sea; Danuberace (A. a. a. pretzmanni) in the upper part of the Danube and its tributary including the Drau; Southrace (A. a. a. canadziae) in the Save river catchment (tributary of the Danube in Croatia) and the lower part of the Danube (Romania, Serbia).



Figure 1 Noble crayfish (*Astacus astacus*) collected in the river catchment of the Danube in Romania (Picture: T. Schikora).

Threats to native crayfish

One-third to one-half of the crayfish species worldwide is threatened with population decline or extinction (Taylor 2002). Besides habitat deterioration, pollution, and other man-made environmental changes, invasive species and pathogens are major threats to the survival of European crayfish species. Therefore, the International Union for Conservation of Nature (IUCN) classifies noble crayfish as a vulnerable species with a decreasing population trend in the international Red List (IUCN 2012). Some national Red Lists even classify noble crayfish as an endangered species (e.g. Germany) or 'Critically Endangered' (e.g. Sweden). Noble crayfish is further included in the Bern Convention (Appendix III) and listed in the European Habitat Directive (Appendix V).

European crayfish species, in particular noble crayfish have been an object of trade, commerce and zoological studies throughout Europe for more than 2000 years (Skurdal and Taugbøl 2002). This has led to stocking of noble crayfish into numerous new localities. Translocations have been documented in central Europe

for centuries, whereas translocations are not known from southeastern Europe (see Albrecht *et al.*, 1983 for more details). This leads to an extensive mixing of crayfish on the one hand and to an overharvesting of local populations on the other hand.

Today, several pathogens (viruses, protozoans, bacteria, fungi) causing severe declines in crayfish populations are known (Edgerton et al., 2004, Longshaw 2011). The most significant threat to European crayfish species is the Oomycete Aphanomyces astaci causing the crayfish plague. A. astaci was introduced to Europe via North American (hereafter 'American') crayfish species that were stocked into European waters to replace the decreasing number of native crayfish populations (Alderman 1996). A. astaci is an invasive, crayfishspecific parasite causing mass mortalities in susceptible European crayfish populations (e.g. Kozubíková et al., 2008, Vrålstad et al., 2011). After the crayfish plague eradicated most noble crayfish populations in Sweden in 2002, only 3% of the populations were left compared to 1900 (Bohman et al., 2006). The pathogen is amplified in susceptible dying animals and spores are subsequently released into the water, usually leading to 100% mortality of the European crayfish present in the respective water body or large parts of it (OIE 2009). A. astaci is an obligate pathogen of freshwater crayfish, but can also be grown in laboratory cultures (Aldermann 1996). A co-evolution of a pathogen and the affected population may lead to a relatively stable host-parasite relationship through enhanced immune functioning by the natural host and virulence attenuation by the parasite (Edgerton and Jussila 2004). Therefore the pathogen can coexist with American crayfish species in such a balanced host-pathogen relationship. If American crayfish species are severely stressed or immunocompromised by other pathogens or environmental factors, they may succumb to the crayfish plague infection (Persson et al., 1987; Söderhäll and Cerenius 1992).



Figure 2 Narrow-clawed crayfish (*Astacus leptodactylus*) from the Romanian Danube. The dark melanisation indicates a crayfish plague infection. (Picture: L. Pârvulescu)

Probably, A. astaci was first introduced to Europe in the late 1850s. This is known as the first infection wave in which thousands of native crayfish populations were eradicated (Aldermann 1996). While the natural hosts of A. astaci, American freshwater crayfish species were not recorded during this first crayfish plague outbreak in Europe they were repeatedly introduced later (Aldermann 1996, Holdich et al., 2009). Until recently it was believed that all American crayfish are carrier of A. astaci (e.g. "This crayfish species is a permanent carrier of the parasite and there are no A. astaci-free P. leniusculus"; Cerenius et al., 2003). In **Appendix 3**, I test the New NICS calico crayfish (Orconectes immunis) for an infection with A. astaci. A positive result would identify a fourth American crayfish as transmitter and support the concern that all American crayfish species in European waters are carriers of the crayfish plague pathogen. It was further believed that native crayfish populations suffer 100% mortality when infected with the pathogen. However, recent studies have shown that a coexistence of A. astaci and European crayfish species is in some cases possible (Jussila et al., 2011; Viljamaa-Dirks et al., 2011), supposedly due to a lowered virulence of *A. astaci*. Furthermore, a coexistence could also be possible when the pathogen is absent in some populations. In *Appendices 4 and 5* I test native narrow-clawed crayfish (Figure 2) from the Romanian Danube for an infection with the crayfish plague agent. Narrow-clawed crayfish coexist with spiny-cheek crayfish (Figure 3) in the upper Romanian Danube (*Appendix 4*) and live in allopatry in the lower Danube (*Appendix 5*). In *Appendix 6* I investigate coexisting populations of noble crayfish and spiny-cheek crayfish. The prevalence of the crayfish plague has major implications on the conservation and management strategies of the threatened European crayfish species, and is critical for further reintroduction (Kozubíková *et al.*, 2009) or translocation attempts.



Figure 3 The 'Old NICS' spiny-cheek crayfish (*Orconectes limosus*) and the 'New NICS' Calico crayfish (*Orconectes immunis*) coexist in the River Rhine (Picture: B. Dahelean).

Genetic Methods

Due to the specific attributes of each molecular method, a combination of methods was necessary for this thesis to resolve the key questions. The phylogeographic analysis of noble crayfish was based on sequence variations of mitochondrial DNA (cytochrome oxidase subunit I (COI) and 16S rRNA) and length variations of microsatellites. Real-time polymerase-chain reaction (qPCR) was used for the verification of *A. astaci* in crayfish populations.

According to Avise (1987) mitochondrial DNA (mtDNA) is an ideal molecular system for phylogenetic analysis because it is distinctive, easy to isolate, has a simple genetic structure, evolves at a rapid rate and exhibits a straightforward mode of genetic transmission since mtDNA is maternally inherited. These features make mtDNA a widely used marker to reconstruct the phylogeographic history of species. It helps to uncover the ice age refuges and the colonization history of a species (Weiss 2002). When a molecular clock is known, a dated phylogenetic tree can be constructed. However, only little or no signal can be seen from sequences in bottlenecked populations that recently colonized an area (Hewitt 1999). Therefore, I additionally utilized nuclear microsatellite data to track the recent gene flow and to estimate the population diversity. The short stretches of repetitive microsatellite DNA evolve at a rapid rate when endogenous DNA polymerase in the cell makes a replication error and either mistakenly adds or removes a copy (DNA slippage). These stepwise mutations occur much more often than other types of mutation (e.g. single base substitutions in mtDNA).

In a sequence as well as in a microsatellite analysis, the first step after the DNA-extraction is the amplification of a specific DNA fragment using PCR. However, to detect small traces of DNA, a conventional PCR is not sufficient. Therefore, for the verification of *A. astaci* DNA in carrier crayfish tissues, I applied quantitative TaqMan[®] minor groove binder (MGB) qPCR. In contrast to a conventional PCR, beside the two species-specific primers, an additional species-specific probe has to anneal to the target DNA in order to get a positive signal. The TaqMan[®] MGB primer and probes bind on the minor groove of the DNA strain and are extremely sensitive to mismatches in the annealing sites of primers and

probe (Yao *et al.*, 2006). Today, this method is the most specific and sensitive method to test for the presence of *A. astaci* (Vrålstad *et al.*, 2009; Tuffs and Oidmann 2011).

OBJECTIVES

The genetic structure of a species (e.g. areas of endemism, geographical patterns in species richness) is important for conservation prioritization. Since the genetic structure is the result of historic processes, I aimed to reconstruct the post-glacial recolonization of central Europe by the endangered noble crayfish (*Appendix 2*). Furthermore, I intended to identify refugial areas that are hotspots of genetic diversity (*Appendix 1 and 2*) and should receive special attention in species conservation. To address these questions I used two genetic data sets: i) sequence data of the mitochondrial cytochrome oxidase subunit I gene (*Appendix 1 and 2*), 16S rRNA (*Appendix 2*) and ii) species-specific microsatellites (*Appendix 2*). The results serve as a basis for conservation management of European freshwater crayfish.

Since the most significant threat to European crayfish species is the crayfish plague pathogen *A. astaci*, the distribution of the disease agent has to be considered before any management action can be taken. In this thesis I aimed to expand the knowledge of the distribution of *A. astaci* using qPCR. One goal was to reveal if the American calico crayfish, which increases its range in the River Rhine rapidly since a few years, is a carrier of *A. astaci* (*Appendix 3*). On the other side, the American spiny-cheek crayfish increased its range in the lower Danube. I intended to reveal its *A. astaci* carrier status and also the status of the native narrow-clawed crayfish in the lower Danube (*Appendix 4*), as well as the status of narrow-clawed crayfish in the Danube Delta (*Appendix 5*). Furthermore, I wanted to reveal mechanisms facilating the coexistence of European and American crayfish species in the same lake.

THESIS OUTLINE

My thesis addresses the objectives with six manuscripts, compiled in the **Appendices 1-6** (Figure 4). Two manuscripts deal with the genetic structure of the European noble crayfish throughout Europe (**Appendices 1 and 2**) and four manuscripts address questions concerning the distribution of the crayfish plague in Germany, Poland and Romania (**Appendix 3 to 6**).

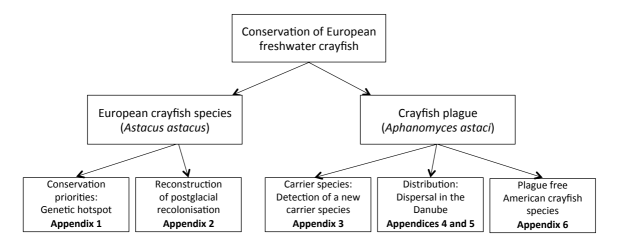


Figure 4 Overview of the different Appendices of this thesis assigned to the two topics of my PhD work: European crayfish species and crayfish plague.

PART 1 - European crayfish species (Astacus astacus)

The conservation of genetic diversity is one goal in modern conservation strategies because genetic variation represents an advantage for the long-term survival of species. To protect the within-species diversity, a good knowledge of the genetic structure of this species is indispensable. Since the genetic structure of a species is a result of both present processes and past history, the post-glacial recolonization history should be reconstructed in order to reveal the genetic structure of a species.

Appendix 1: First large-scale genetic analysis of the vulnerable noble crayfish *Astacus astacus* reveals low haplotype diversity of central European populations

In Appendix 1 I present the first study focusing on the genetic structure of noble crayfish throughout its European distribution area. I analyzed a partial sequence of the mitochondrial gene cytochrome oxidase subunit I (COI) from a large set of noble crayfish populations throughout Europe to localize genetic hotspots and to prioritize conservation effort. This study has shown that the highest haplotype diversity is located in southeastern Europe, which indicates with reasonable certainty a glacial refugium. In contrast, I showed that diversity is relatively low in central Europe, which was recolonized after the last ice age. I discovered differentiation between river catchments despite the extensive human translocation of crayfish. In this first study I revealed that more markers are needed to better understand the recolonization history of noble crayfish. Furthermore, I recognized areas from where more samples are needed in order to identify the genetic hotspots more specifically in a subsequent supplementary study.

Appendix 2: Recolonization of Europe by noble crayfish (Astacus astacus) – natural versus human-mediated processes

In the second Appendix I focus on the postglacial recolonization of central Europe by noble crayfish. For this, I improved the molecular methods by additionally using nuclear markers (microsatellites) compared to the previously performed mtDNA-based study. Additionally, 16S rRNA was used as a second mitochondrial marker besides COI. By using these methods, I was able to calculate the first calibrated tree of noble crayfish and correlate the split of lineages with climatological events. I detected a very distinct lineage in the Western Balkans that may have served as an isolated glacial refugium during the last glacial maximum. I suppose a second independent refugium in the eastern Black Sea basin from where the species recolonized central Europe through the Danube and through a second migratory route in eastern Europe.

PART 2 - Crayfish plague (Aphanomyces astaci)

For the conservation of European crayfish species, knowledge on the distribution of *A. astaci* in European waters is of great importance. This aspect correlates with the question which crayfish species are latent carrier of the disease agent.

Appendix 3: Crayfish plague agent detected in populations of the invasive North American crayfish *Orconectes immunis* (Hagen, 1870) in the Rhine River, Germany

In the third Appendix I identified calico crayfish as the fourth North American crayfish species in European waters, which is carrier of the crayfish plague pathogen. A positive verification was received from two sampling sites in the upper Rhine River where calico crayfish seems to displace its invasive predecessor spiny-cheek crayfish. This finding of the positive detection is worrying considering the fast and successful invasion process of calico crayfish. Furthermore, this outcome supports the apprehension that all American crayfish species are carriers of the crayfish plague.

Appendix 4: Invasive crayfish and crayfish plague on the move: first detection of the plague agent *Aphanomyces astaci* in the Romanian Danube

The fourth Appendix focuses on the dispersal of *A. astaci* in the Romanian Danube. The crayfish plague disease agent could be detected in American narrow-clawed crayfish as well as in European spiny-cheek crayfish. According to field observations of the first occurrences of narrow-clawed crayfish along the Danube, an invasion speed could be calculated for this species. It was estimated that the invasive narrow-clawed crayfish invade the sensitive Danube Delta area in the mid-2060s, but the crayfish plague agent may reach the delta significantly earlier.

Appendix 5: Crayfish plague pathogen detected in the Danube Delta – a potential threat to freshwater biodiversity in southeastern Europe

In the fifth Appendix I survey native narrow-clawed crayfish from the highly protected Danube Delta for an infection with *A. astaci*, with a positive test result. The delta is located 970 km downstream of the current invasion front of American

crayfish in the Danube. This result shows that the pathogen has a much wider range in this river than previously assumed. *A. astaci* seems to persist in local populations, as neither crayfish mass mortalities, nor alien crayfish species have been reported from this region. In this Appendix different possible dispersal ways of the pathogen towards the delta are discussed. However, the real transmitter/vector could not be determined.

Appendix 6: Absence of the crayfish plague pathogen (*Aphanomyces astaci*) facilitates coexistence of European and American crayfish in central Europe

In the sixth Appendix I study coexisting populations of introduced spiny-cheek crayfish and native noble crayfish. I resolve the question whether the coexistence resulted from reduced virulence in local *A. astaci* strains, increased resistance in susceptible crayfish species, or a complete absence of the pathogen in the American crayfish populations. A negative result of 523 crayfish tested for an infection with *A. astaci* is a very strong indication that the coexistence is possible in the absence of the pathogen in these populations. Exposure experiments confirmed these results.

SUMMARY OF RESULTS

In this chapter I will briefly summarize the most important results of my thesis. In the following discussion I will get back to these results. For more detiails on the results see the Appendices.

European crayfish species (Astacus astacus)

In the first part using sequence analyses and microsatellite analyses I revealed high genetic diversities (high number of haplotypes and number of alleles) of noble crayfish in southeastern Europe (*Appendix 1 and 2*). The haplotype diversity (H_D) is highest in the Black Sea basin (H_D = 0.851, Table 2). In contrast the genetic diversities are low in central Europe. The lowest haplotype diversity was detected in the Baltic Sea basin (H_D = 0.276). I detected the highest number of differentiated haplotypes/alleles in the Western Balkans (Adriatic Sea basin and south-western Black Sea basin; red Star in Figure 6). Six haplotypes from the Western Balkans (Hap41 – Hap46) were highly distinct from all other haplotypes (Figure 2 in *Appendix 2*). The calibrated Bayesian phylogenetic tree revealed that the Western Balkan haplotypes (Lineage 4 in Figure 5) were separated from all other haplotypes about 710,890 years before present (YBP), whereas all other differentiations occurred within the last 450,000 YBP. The microsatellite analysis supported the differentiation of the Western Balkan samples (Figure 5 in *Appendix 2*).

Table 2 Results from a combined 350 base pair (bp) fragment of the COI sequence and a 476 bp fragment of the 16S rRNA sequences for the analyzed sea basins. The number of sequenced individuals (N), the number of haplotypes (H_N) and the haplotype diversity (H_D) is given (also see *Appendix 2*).

	Sites	N	H _N	H _D
Baltic Sea	14	66	5	0.276
North Sea	59	249	13	0.316
Black Sea	82	185	28	0.851
Adriatic Sea	5	12	5	0.576

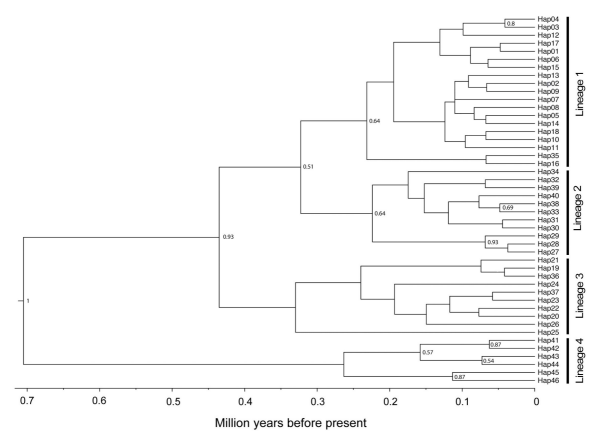


Figure 5 Phylogenetic tree generated using Bayesian Markov Chain Monte Carlo analysis implemented in Bayesian Evolutionary Analysis Sampling Trees (BEAST; Drummond *et al.*, 2012) by using a combined fragment of the COI sequence and a fragment of the 16S rRNA sequence. The axis shows the time scale in million years before present. Shown are the posterior probability values above 0.5 (see also *Appendix 2*).

In total, eight haplotypes were shared between different sea basins and between different river catchments. For example the haplotype Hap41 from the differentiated lineage 4 was shared between the Danube river catchment in the Western Balkans and the Elbe river catchment (North Sea basin).

The probability values/Bayes factors for the model selection calculated with Migrate-n version 3.3.2 (Beerli and Felsenstein 2001) using mtDNA sequence data preferred model 2b (see *Appendix 2*). This model implies a 'bifurcal' colonization route for noble crayfish from the eastern Black Sea basin (Romania, Bulgaria) to a) the North Sea basin and to b) the Baltic Sea basin (see schematic arrows in Figure 6).

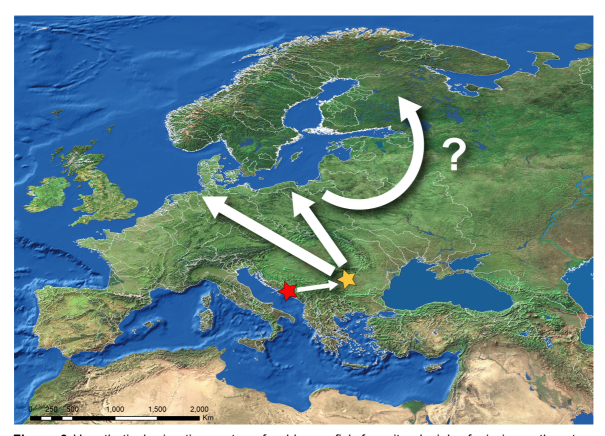


Figure 6 Hypothetical migration routes of noble crayfish from its glacial refugia in southeastern Europe into northern and central Europe are shown schematically. Arrows indicate the directions of gene-flow. Hotspots of genetic diversity are indicated with stars. Red star: Western Balkan, yellow star: eastern Black Sea basin. The '?' indicates that the route to Finland should be considered with caution because of the low sample size of Finnish samples (N = 10) (see also **Appendix 2**).

Crayfish plague (Aphanomyces astaci)

In *Appendix 3* using qPCR I tested the New NICS calico crayfish from the river Rhine in Germany positive for an infection with *A. astaci*. From the sampling site Germersheim in the main river 60% of the samples were positive (N = 50; N_{+} = 23) and from Bühl (a side channel of the Rhine) 81% were positive (N = 32; N_{+} = 26). With this positive verification I identified calico crayfish as the fourth American crayfish species to be carrier of the crayfish plague agent (*Appendix 3*).

Furthermore I detected the crayfish plague pathogen in American spiny-cheek crayfish and in native narrow-clawed crayfish in the lower Danube in Romania (*Appendix 4*). The pathogen could be detected in 41% of the tested native species (N = 49; $N_+ = 20$) and in 32% of the tested NICS (N = 71; $N_+ = 23$). Moreover, I detected that native narrow-clawed crayfish in the Danube Delta at

two sampling sites about 970 km downstream of the current invasion front of American crayfish in the Danube is a carrier of *A. astaci* (*Appendix 5*). In total, 30% of the samples from the Chilia Channel (N = 37; N_+ = 11) and 67% of the samples from the Merhei Lake (N = 3; N_+ = 2) were tested positive.

On the other side, from a total of 523 tested crayfish (490 spiny-cheek crayfish, 33 noble crayfish) from six coexisting populations of European and American crayfish species, no *A. astaci* positive individuals were found. With an assumed lowest possible *A. astaci* prevalence of 10% in a carrier population, there is a 98% probability of disease absence in five of the nine tested coexisting populations. Exposure experiments confirm these results: no abnormal mortality or behavioural changes were seen in noble crayfish kept together with American crayfish from the coexisting populations.

GENERAL DISCUSSION

In the following discussion I will first address the two parts of my thesis (native crayfish and crayfish plague) separately and after that I will discuss these aspects that are inextricably linked together combined.

Conservation considerations - native crayfish

The results of this thesis indicate that special attention in species conservation of noble crayfish should be put on the Danube river catchment in southeastern Europe, where the highest genetic diversity was detected (Appendix 1, Schrimpf et al., 2011). Noble crayfish outlasted cold climate phases in southeastern Europe while the climate conditions in central and northern Europe were unfavorable for crayfish species. The same applied to many other freshwater species (e.g. Hänfling et al., 2009; Weiss et al., 2002). In particular, I detected that noble crayfish outlasted the last glacial maximum in at least two different refugia in southeastern Europe: one was located in the Western Balkans and one in the eastern Black Sea basin (see scheme in Figure 6). Noble crayfish from the Western Balkans are particularly diverse and highly differentiated. While the Western Balkan haplotypes (Lineage 4 in Figure 5) were separated from all other haplotypes already about 711,000 YBP, all other haplotypes differentiated within the last 450,000 YBP. Therefore, I assume that the Western Balkans served as an older glacial refuge (*Appendix 2*, Schrimpf et al., unpublished data). Further, the high diversity in the eastern Black Sea basin and the similarity to central European haplotypes indicates a second refuge in the Danube river catchment from where the noble crayfish recolonized central Europe. It has to be mentioned, that the data from the eastern Black Ses basin is restriced to Romania and Bulgaria. However, it is very well possible that the glacial refuge was even further south, e.g. in Greece, and Romania and Bulgaria were recolonized from a more southern refuge before noble crayfish recolonized central and northern Europe from the eastern Black Sea basin. I revealed that the recolonization of the North Sea basin was possible along the Danube and the recolonization of the Baltic Sea basin along a second eastern corridor. The sample size of Finnish samples however was too small (N = 10) to definitely reveal the colonization route of Finland. In an additional study more samples from eastern and northern Europe should be analyzed in order to reveal the eastern corridor more specific and to reconstruct the (re-)colonization of northern Europe.

In this thesis I have detected that the genetic structure of noble crayfish is partly dissolved (Appendices 1 and 2). This altered genetic structure was probably caused by the high anthropogenic influence on the noble crayfish dispersal (e.g. translocation) throughout Europe (Skurdal and Taugbøl, 2002). The haplotype Hap41 from the differentiated Lineage 4 that was detected in the Western Balkans and in the Elbe river catchment (Appendix 2), was probably introduced to the Elbe river catchment artificially and led to a 'genetic contamination of the local population (Largiadèr et al., 2000). Long-distance translocation of noble crayfish needs to be prohibited to conserve the remaining (natural) genetic structure of the species. However, the disrupted genetic structure might explain why partly different and contradictory classifications of the noble crayfish subspecies based on morphological criteria can be found in the literature (see Smietana et al., 2006 for a review). When noble crayfish are collected for morphological determination from a specific sampling site, it is very well possible that this site was stocked artificially and that researchers did therefore not always use autochthonous crayfish to define morphological characteristics of noble crayfish for a certain region.

The geographical assignment of the three races of *A. a. astacus* based on morphological characters (Karaman 1962, 1963, see also the Introduction of my thesis) cannot be confirmed with the genetic data of individuals from the same regions (*Appendices 1 and 2*). It was not possible to reveal if the discrepancy between morphological and genetic differentiation can be explained by a difference between morphological and genetic characters (e.g. morphological and genetic characters evolve at a different rate) or by artificial translocations. Presumably, it is a combination of different aspects.

However, as long as the taxonomic status of a species is not resolved, one cannot be sure which taxonomic unit is being translocated: a different species, a subspecies, a race or a genetic variation. It is oftentimes impossible to define the

taxonomic level, since the threshold for any taxonomic level (e.g. species, subspecies, race) is difficult to define. The existence of different species concepts additionally complicates the delimitation of a species/subspecies (de Queiroz 2007). A species under the 'Biological species concept' (sensu Mayr 1942, a species is defined based on its ability to inbreed) could be considered a subspecies under the 'Phylogenetic species concept' (Cracraft 1983) or vice versa. Therefore, any introduction of individuals from a distant region needs to be prohibited and the conservation of local population should be promoted in order to protect local adaptions and endemic haplotypes/alleles. This way the greatest possible genetic variation of a species can be conserved.

The taxonomic status of European crayfish species is not fully resolved (Holdich *et al.*, 2006). One example is the white-clawed crayfish case (*A. pallipes* species complex), whose taxonomic order is still under revision. While in the past A. pallipes was considered a 'species', it is today considered a 'species complex' (Holdich *et al.*, 2006). In the future the *A. pallipes* 'species complex' might be split into two separate species (*A. pallipes* and *A. italicus*), as proposed by Grandjean *et al.*, (2002). With regard to noble crayfish, we cannot exclude, that populations today known as noble crayfish throughout Europe will in future be classified as more than one species. Only if between-catchment translocation of noble crayfish is stopped and translocation for conservation purposes will be conducted within ESUs (e.g. river catchments or even smaller units, like side arms of a river), the conservation of the species integrity is possible. In summary, there is urgent need for a close collaboration between taxonomists and molecular biologists as well as conservation biologists to reveal the taxonomic status of European crayfish species as a basis for species conservation.

Unfortunately, the region with the highest genetic diversity of noble crayfish coincides with the region that is under current threat of disease carrying American crayfish species (*Appendices 4 and 5*, Hudina *et al.*, 2009, Pârvulescu *et al.*, 2009). Therefore, before any conservation measures of native crayfish can be conducted the distribution of the crayfish plague pathogen has to be considered.

Conservation considerations - crayfish plague

Until recently it was believed that a) all (North) American crayfish are carrier of A. astaci (e.g. "This crayfish species is a permanent carrier of the parasite and there are no A. astaci-free P. leniusculus"; Cerenius et al., 2003) and that b) native crayfish populations suffer 100% mortality when infected with A. astaci (Table 3). With the positive verification of calico crayfish as carrier of A. astaci (Appendix 3, Schrimpf et al., 2013) I support the assumption a) that all American crayfish spesies are indeed potential carriers. However, I discovered potential plague-free spiny-cheek crayfish populations and therefore I show that not all populations of American crayfish species in Europe are necessarily infected with A. astaci (Appendix 6, Schrimpf et al., in press). These results demonstrate that the disease transmission risk varies substantially between different American crayfish populations. The existence of plague-free American crayfish populations has been assumed by several authors (e.g. Westman and Savolainen, 2001, Jussila and Mannonen 2004, Skov et al., 2011) but never before been studied thoroughly. In addition, I contradict the assumption b) that all European crayfish suffer 100% mortality when infected with A. astaci. With the positive verification of narrow-clawed crayfish as carrier of the agent of the crayfish plague (Appendices 4 and 5, Pârvulescu et al., 2012, Schrimpf et al., 2012), I show that latent infected native crayfish populations do exist. This result is supported by recent molecular studies, which also indicate that persistent infections of A. astaci occur in native European crayfish (Jussila et al., 2011; Viljamaa-Dirks et al., 2011; Kokko et al., 2012, Svoboda et al., 2012).

Table 3 The common assumption and the outcome of this thesis regarding the infection status of North American crayfish populations (NACP) and the reaction of European crayfish population (ECP) to an infection. \checkmark = correct, \checkmark = wrong.

	Commo	on assumption	This	s thesis
	All populations are infected	All populations die shortly after infection	All populations are infected	All populations die shortly after infection
NACP	✓	X	×	×
ECS	X	✓	X	×

These findings demonstrate that even after many years of visual observations, one cannot know if *A. astaci* is present or absent in a crayfish population. The wrong assumption of the plague's absence can have fatal consequences for other native crayfish populations because the pathogen must not be equally lethal to all populations. It is possible that some local populations evolved a resistance against this pathogen, but other recently infected populations without previous 'selection opportunity' could be susceptible and might be completely destroyed. Therefore, before native crayfish individuals are being translocated to a new locality, molecular analyses need to be done to reveal the carrier status of the donor population. On the other side, European crayfish populations are not necessarily lost, in case American crayfish occur in the same water. The native crayfish can be transferred to waters that are free of American crayfish species. Furthermore, it has to be considered that any translocation also involves the risk of spreading other diseases and associated non-target species 'accidentally' (Edgerton and Jussila 2004).

American crayfish species were initially introduced into Europe to replace native crayfish that suffered high mortalities due to environmental changes (Alderman et al., 1990). Back then, the replacement of one species by another one, which occupies a similar ecological niche, was not believed to cause any negative ecological consequences. However, today we know that there are differences in the dietary habits between different crayfish species and that their physical impact on the environment varies (see Holdich 2002b for a summary of different species). Thus, the replacement of one crayfish species by another can have fatal consequences on the environment. For this reason, nowadays the introduction of American crayfish species into the wild in Europe is forbidden and the western European native species (A. astacus, A. pallipes, A. torrentium) are listed as protected in Appendix III of the Bern Convention. Species listed under Appendix III are those who are in need of protection and any exploition of wild fauna shall be regulated in order to keep the populations out of danger (Council of Europe 2009).

Synthesis of discussion

Protection of the native biodiversity is the major aim of conservation biology (Frankham et al. 2004). In the last decades many conservation projects have been conducted that consisted in restoring the habitat, (re-)introducing native crayfish species (Taugbøl and Peay 2004, Dümpelmann 2011) or eradicating NICS in the wild (Sandodden and Johnsen 2010). However, while there are numerous reports on successful conservation measures, many introductions have failed due to new crayfish plague outbreaks. The extinction of local populations can have fatal consequences for the environment. Due to their trophic activities as omnivores crayfish play a key role in many freshwater ecosystems (Nyström 1999), and their loss may change the food-web with drastic impacts on local species composition. As long as infected crayfish species are subject to aquarium trade, NICS and the crayfish plague pathogen will repeatedly be introduced into the wild. Many people promote the introduction of exotic species because trading these species promises increased profit (McNeely 2001). The persons involved in the introduction often don't consider the negative impact these species might have on the environment. Usually it's the general public and future generations who have to pay the price.

Germany is the main importer of exotic crayfish species in Europe (Chucholl 2012). The majority of the 120 NICS, which are available in German aquarium trade originate from America and is therefore suspected to be carrier of the crayfish plague. Contrary, in Norway the import of live crayfish is banned and until recently Norway was free of NICS (Vrålstad *et al.*, 2011). However, few crayfish plague outbreaks and the recent findings of NICS in Norway (Johnsen *et al.*, 2007, Vrålstad *et al.*, 2011) are believed to have resulted from introductions of NICS and transmissions of *A. astaci* from lakes in Sweden close to the Norwegian border (Taugbøl 2004). Boats or fishing gear used on both sides of the border might have served as vectors for the *A. astaci* transmission. This example highlights that all European countries have a responsibility not only for their own environment but also for that of their neighbouring countries because non-indigenous species do not adhere to national boundaries (European Commission 2008).

CONCLUSION

The results of my studies show that a high degree of genetic diversity of noble crayfish can be conserved when the genetic hotspots in southeastern Europe receive special attention in conservation management (Appendices 1 and 2). Genetic diversity and endemic haplotypes/alleles can be protected when species introduction for conservation purposes is being conducted only within ESUs (river catchments) and translocations between river catchments are strictly avoided. However, the greatest threat to native crayfish biodiversity stems from the introduction of NICS infected with crayfish plague (Lodge et al., 2000). Because the disease agent may devastate conservation measures, its distribution has to be considered before any management action can be taken. In my thesis I could verify A. astaci in several populations in Poland, Germany and Romaia in native and American crayfish species (Appendices 3 - 6). While I could show that plague-free American crayfish populations do exist (Appendix 6) and native Euroipean crayfish can be infected without indication of illness (Appendices 3 and 4), these findings however constitute exceptions. Moreover, the results show that even native species may serve as a transmitter between water bodies even if they don't show signs of infection. Therefore, a health test should be conducted before native crayfish species are translocated.

As long as infected NICS can be purchased by anybody via pet stores or internet, they will repeatedly be introduced into the wild and eradicate genetic resources of native crayfish. NICS and the crayfish plague continue to be the greatest threat to the survival of native crayfish. Therefore, crayfish conservation needs to address the threat posed by NICS with a high priority. A change in legislation, which prohibits the trade with infected invasive species will increase the long-term survival of native crayfish species and is therefore urgently needed.

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APPENDIX

Appendix 1

Schrimpf A, Schulz HK, Theissinger K, Pârvulescu L, Schulz R. (2011) First large-scale genetic analysis of the vulnerable noble crayfish *Astacus astacus* reveals low haplotype diversity of central European populations. Knowledge and Management of Aquatic Ecosystems. 401: 35.

Appendix 2

Schrimpf A, Dahlem J, Maguire I, Pârvulescu L, Schulz HK, Theissinger K and Schulz R (unpublished) Recolonisation of Europe by noble crayfish (*Astacus astacus*) – natural versus human-mediated processes.

Appendix 3

Schrimpf A, Chucholl C, Schmidt T, Schulz R (2013) Crayfish plague agent detected in populations of the invasive North American crayfish *Orconectes immunis* (Hagen, 1870) in the Rhine River, Germany. Aquatic Invasions, 8.

Appendix 4

Pârvulescu L, Schrimpf A, Kozubíková E, Resino SC, Vrålstad T, Petrusek A, Schulz R (2012) Invasive crayfish and crayfish plague on the move: first detection of the plague agent *Aphanomyces astaci* in the Romanian Danube. Diseases of Aquatic Organisms, 98: 85-94.

Appendix 5

Schrimpf A, Pârvulescu L, Copila-Ciocianu D, Petrusek A, Schulz R (2012) Crayfish plague pathogen detected in the Danube Delta – a potential threat to freshwater biodiversity in southeastern Europe. Aquatic Invasions, 7(4): 503–510.

Appendix 6

Schrimpf A, Maiwald T, Vrålstad T, Schulz HK, Smietána P, Schulz R (in press) Absence of the crayfish plague pathogen (*Aphanomyces astaci*) facilitates coexistence of European and American crayfish in central Europe. Freshwater Biology.

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The first large-scale genetic analysis of the vulnerable noble crayfish Astacus astacus reveals low haplotype diversity in central European populations

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ABSTRACT

Major global changes (e.g., human impact or climatic cycles) have a severe impact on the distribution and diversity of species such as the vulnerable European noble crayfish Astacus astacus. This is the first large-scale study regarding haplotype diversity of A. astacus in central and southeastern Europe. We analyzed a partial sequence of the mitochondrial gene cytochrome oxidase subunit I from 416 specimens of 92 crayfish stocks of three European river basins (Black Sea, North Sea and Baltic Sea). Twenty-two haplotypes were identified, and one common haplotype was found, across the whole study area. We detected differences in the genetic diversity between major river catchments (Φ_{ST} : 0.03481 to 0.20387). The high haplotype diversity ($H_D = 0.794 \pm 0.024$) and high number of private haplotypes suggests a glacial refuge in the Balkan area. The very low haplotype diversity in central Europe ($H_D = 0.299 \pm 0.038$ and $H_D = 0.163$ ± 0.058) could be a result of human translocation and/or founder effects due to postglacial recolonization. Nevertheless, the high frequency of private haplotypes in all major catchment areas indicates a differentiation of noble crayfish populations throughout Europe despite the extensive human translocation of crayfish. The results of this study support the establishment of conservation management plans for this vulnerable species.

Key words: mtCOI sequences • haplotype diversity • human translocationconservation management • evolutionary significant units (ESU)

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RÉSUMÉ

Une première analyse génétique à grande échelle des populations vulnérables de l'écrevisse *Astacus astacus* révèle une faible diversité des haplotypes dans les populations du centre de l'Europe

D'importants changements globaux (par exemple, l'impact humain ou les cycles climatiques) ont un grave impact sur la distribution et la diversité des espèces telle que l'écrevisse à pattes rouges européenne Astacus astacus. C'est la première étude à grande échelle en matière de diversité des haplotypes d'A. astacus en Europe centrale et du sud-est. Nous avons analysé une séquence partielle du gène mitochondrial de la cytochrome oxydase, sous-unité I de 416 spécimens de 92 stocks d'écrevisses de trois bassins hydrographiques européens (Mer Noire, Mer du Nord et Mer Baltique). Vingt-deux haplotypes ont été identifiés, et un haplotype commun a été trouvé dans toute la zone d'étude. Nous avons détecté des différences dans la diversité génétique entre les principaux bassins versants (ΦST: 0,03481 à 0,20387). La forte diversité des haplotypes ($HD = 0.794 \pm 0.024$) et le nombre élevé d'haplotypes suggèrent un refuge glaciaire dans la région des Balkans. La très faible diversité des haplotypes en Europe centrale $(0,299 \pm HD = 0,038 \text{ et } 0,163 \pm HD = 0,058)$ pourrait être une conséquence de la translocation de l'homme et / ou des effets fondateurs en raison de la re-colonisation postglaciaire. Néanmoins, la fréquence élevée des haplotypes particuliers dans tous les grands bassins versants indique une différenciation des populations d'écrevisses à pattes rouges à travers l'Europe, malgré les translocations importantes de l'écrevisse. Les résultats de cette étude viennent appuyer la mise en place de plans de conservation pour cette espèce vulnérable.

Mots-clés: séquences mtCOI • diversité des haplotypes translocation • gestion de la conservation • evolutionary significant units (ESU)

INTRODUCTION

Genetic diversity among and within species is the basis of evolution and is important for the resilience of native stocks to future changes in the environment (Östergren, 2006). The convention on biological diversity (CBD) highlighted the importance of genetic diversity within species as one of the three levels of biological diversity (CBD, adopted 1992). Therefore, the identification of genetically differentiated populations is crucial for the conservation and management of a species or the regional strains within a species. The guidelines for the protection of threatened species recommend the identification of evolutionary significant units (ESU, Ryder, 1986). An ESU may be defined as a population or group of populations that is sufficiently differentiated and requires a separate management or conservation strategy (Moritz, 1994). Defining management units can aid in selecting a management program among the various populations so that the greatest overall diversity is sustained by the conservation plan (Crozier, 1997). Nevertheless, conservation managers typically do not use genetic evidence (i.e., the identification of ESUs) to support their adopted measures (Pullin et al., 2004). The IUCN red list of threatened species cites Astacus astacus as vulnerable with a decreasing population status (Edsman et al., 2010). A. astacus is widely distributed across Europe. Its range extends from Russia and the Ukraine in the east, Scandinavia in the north, Greece in the south, and the United Kingdom and France in the west (Edsman et al., 2010). Based on morphological criteria, different and partly contradictory classifications of the noble crayfish subspecies can be found in the literature (see Smietana et al., 2006 for a review). Here, we focus on the taxonomic studies of Karaman (1962) and Albrecht (1983). They described three subspecies with distinct geographical distributions: (1) A. a. colchicus is endemic to an isolated area in the Ponto-Caspian region of the upper Rion (Caucasus). (2) A. a. balcanicus inhabits the western Balkans in the drainage system of the Vardar river in Macedonia and Greece, as well as Lake Ohrid. (3) A. a. astacus inhabits the Danube river and its tributaries, as well as the rivers that open out into the North and Baltic Seas. The fragmented nature of A. astacus' typical freshwater habitats (streams, rivers and lakes with sufficient water quality and an excess of shelter) coupled with a low migratory potential

reduces the possibility of reestablishing local populations that have gone extinct (Abrahamsson, 1971; Strayer and Dudgeon, 2010). In addition to environmental contributions to the decreasing number of stocks (i.e., water pollution or channelization of streams), the oomycete Aphanomyces astaci is decreasing A. astacus' numbers (Schikora, 1906). This invasive pathogen was brought to Europe via American crayfish species (e.g., Orconectes limosus) in the midnineteenth century (Alderman et al., 1990) and caused massive die-offs in native crayfish stocks (Alderman, 1996). During the Middle Ages, the noble crayfish became a commercial object throughout Europe, and it has been sold as a delicacy ever since (Schulz et al., 2004). The translocation of individuals and whole stocks (Skurdal and Taugbøl, 2002), sometimes over large distances (Koutrakis et al., 2007), resulted in the foundation of new stocks (Albrecht, 1983) and in the mixing of indigenous and non-indigenous populations. Over the last decades, the number of stocking events that disregard the genetic structure within and between populations (Souty-Grosset and Reynolds, 2009) and cross-basin translocations in response to rapidly declining stocks have led to a contamination of local stocks (Largiadèr et al., 2000). This has led to repeated calls for modern conservation programs that consider the genetic origin of the stocking material (e.g., Schulz et al., 2004; Bertocchi et al., 2008; Souty-Grosset and Reynolds, 2009; Strayer and Dudgeon, 2010). Previous studies on noble crayfish taxonomy and biogeography are based solely on morphological characteristics (Bott, 1950, 1972; Karaman, 1962, 1963; Albrecht, 1983), while molecular studies have only focused on restricted geographical areas (Agerberg, 1990; Fevolden et al., 1994; Schulz, 2000; Edsman et al., 2002; Schulz et al., 2004; Alaranta et al., 2006). This study is the first molecular-based study of A. astacus covering a large portion of its distribution range, including river catchments of the North and Baltic Seas in central Europe and the Black Sea in southeastern Europe. The main aim of this study was to determine the degree of genetic structure in noble crayfish populations across three major catchments areas. In particular, we focused on asking to what extent this structure is the consequence of natural migration due to past climatic fluctuations during the Pleistocene or recent man-induced translocations. In the latter case, we would expect a lower genetic structure due

to shared haplotypes between geographically disconnected populations. An anthropogenic influence could have caused a homogenization of haplotypes throughout Europe. The identification of inter-basin diversity could result in the conservation of catchment-specific gene pools (*i.e.*, the identification of ESUs) to protect the present-day genetic diversity (Weiss *et al.*, 2002). Finally, the results of this study are discussed in the context of conservation management plans for this vulnerable species.

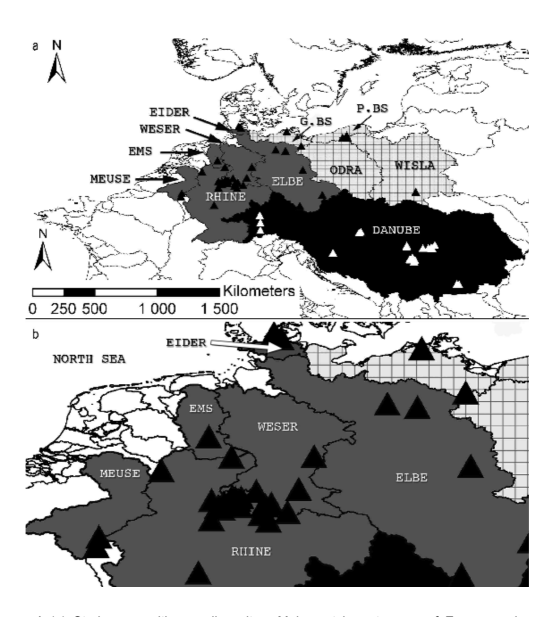


Figure 1 (a) Study area with sampling sites. Major catchment areas of European rivers are indicated by a checked pattern for the Baltic Sea catchment area; by dark grey for the North Sea catchment area; and by black for the Black Sea catchment area. G. BS = German Baltic Sea estuary, P. BS = Polish Baltic Sea estuary. (b) Larger scale map of the North Sea major catchment.

MATERIALS AND METHODS

Sampling

A total of 416 crayfish specimens from 92 sampling locations (Figure 1,Tables I and II) were collected either by hand or with traps within catchments in the North, Baltic, and Black Seas. Immediately after the lower part of one pereopod (propodus and dactylus) was taken, specimens were released at the place where they were caught. Appendages usually regenerate after a few molts. The samples were stored in 96% ethanol until DNA extraction. DNA was extracted from the muscle tissue using a standardized protocol ('rapid isolation of Mammalian DNA', Sambrook and Russel, 2001).

Laboratory procedures

A 350 base pair (bp) fragment of the mitochondrial cytochrome oxidase subunit I (COI) was sequenced using the primer pair ASTCOI (forward primer: 5'-5'-GCGGGGATAGTAGGAACCTC-3'; reverse primer: ATTTACCGCCCCTAAAATCG-3'; Schrimpf and Schulz, in prep.). Polymerase chain reactions (PCR) were performed in a total volume of 25 µL containing 0.625u GoTaq DNA Polymerase (Promega, Mannheim, Germany), 1x Colorless GoTag[®] Flexi Buffer (Promega, Mannheim, Germany), 2 mM MgCl₂ (Promega, Mannheim, Germany), 0.24 mM of each dNTP (Fermentas, St. Leon-Rot, Germany), 0.2 μM of each primer and 20 ng of template DNA. The PCR reaction was performed using a Primus 96 Cycler (Peqlab Biotechnologie GmbH, Erlangen, Germany) under the following conditions: an initial 2 min step at 95 °C, followed by 35 cycles with 45 s at 95 °C, 45 s at 50 °C, 1 min at 72 °C, and a final extension of 5 min at 72 °C. PCR products were stored at 10 °C. PCR products were partly sequenced on a 3730 DNA Analyzer eight capillary sequencer (Applied Biosystems, MA, USA) by the company SeqIT (Kaiserslautern, Germany). The remaining PCR product was purified with the AMPure XP system (Beckman Coulter, Krefeld, Germany). The fragments were amplified in a sequencing-reaction using the primer ASTCOIF and the CEQ DTCS-Quick Start kit (Beckman Coulter, Krefeld, Germany). After ethanol precipitation, sequencing was performed on a Beckman Coulter CEQ 8000 eight capillary sequencer. Each

template was sequenced in the forward direction, and 50% of samples were randomly chosen for sequencing in the reverse direction. The sequences were edited and aligned with the Sequencher 3.0 software (gene codes corporation, Ann Arbor, Michigan). The sequences were checked manually for base pair ambiguities, nuclear copies of mitochondrial derived genes, stop codons, and high levels of divergence, as recommended by Buhay (2009). All haplotypes were submitted to GenBank.

Statistical Analyses

A median joining (MJ) network (Bandelt et al., 1999) was constructed using the software network 4.510 (www.Fluxus-Engineering.com, 2009) to identify haplotypes and to determine the phylogenetic relationships between haplotypes, which allows us to identify geographically localized haplogroups. The network was also calculated with TCS 1.21 (Clement et al., 2000) software to confirm the shape of the network. Haplotype diversity (H_D) and the number of polymorphic sites (S) per river catchment were calculated using the software ARLEQUIN 3.5.1 (Excoffier et al., 2005). The calculation of haplotype diversity is based on the frequency of a haplotype and the sample size (Nei, 1987). To detect genetic differentiation among major catchment areas, an analysis of the molecular variance (AMOVA, Excoffier et al., 1992), based on three hierarchical groups corresponding to the major catchment areas, and the pairwise Φ_{ST} -values (Weir and Cockerham, 1984) were calculated using the program ARLEQUIN 3.5.1. We further examined population genetic differentiation using exact tests of population differentiation (ETPD, Raymond and Rousset, 1995), as implemented in ARLEQUIN 3.5.1. In addition, H_D , S, and pairwise Φ_{ST} -values (Weir and Cockerham, 1984) were calculated for sampling sites in the North Sea catchment area. Here, the Ems population has been excluded from the calculation of the pairwise Φ_{ST} -values due to the low sample size (N = 5).

RESULTS

COI sequences were analyzed for 416 specimens from 96 sample sites across three major catchment areas in Europe (Tables I and II, Figure 1). There were 20 variable sites within the 350-bp fragment (0.06%). Only four of these

substitutions resulted in an amino acid change in four haplotypes (H04, H10, H21). In total, 23 unique haplotypes were detected (GenBank accession numbers: JN254659-JN254681), 17 of which were endemic to a specific river catchment. The overall haplotype diversity was 41.9%. The median joining (MJ) network is shown in Figure 2. The shape was confirmed with the TCS software. The greatest observed distance between haplotypes was 9 bp. For the Black Sea catchment, 10 out of 16 haplotypes were endemic to this region. Of the 11 North Sea haplotypes, five were endemic to the catchment area, and two out of three haplotypes were exclusively found in the rivers discharging into the Baltic Sea. Five haplotypes (H05, H12, H17, H18, H20) were shared between populations in the Black and North Seas. One haplotype (H01), representing the center of the MJ network, was found frequently in all regions, resulting in a star-like topology of the network. A list of all haplotypes for each sampling site is given in Table II.

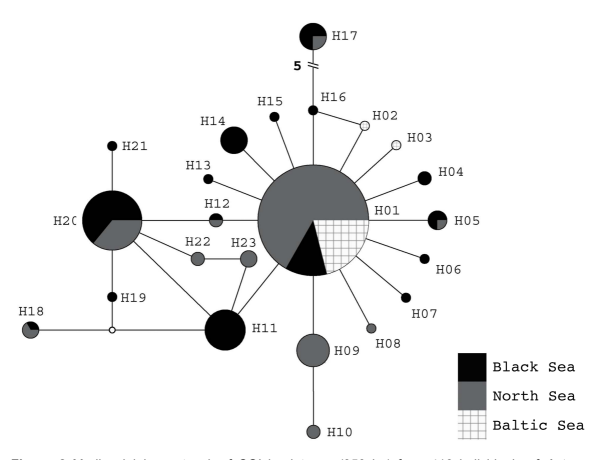


Figure 2 Median joining network of COI haplotypes (350 bp) from 416 individuals of *Astacus astacus*. The size of the circles is proportional to the frequency of the haplotypes. Median vectors are indicated as white dots. The number of base pair (bp) changes are given; no number = 1 bp change.

Table I Locality information of analyzed *Astacus astacus* populations. The originating countries and river catchments, the number of populations (Pop) and the number of sequenced individuals (N), the number of haplotypes (HN), the haplotype diversity (HD) and the number of polymorphic sites (S) are given. Country Codes: AUT = Austria; BEL = Belgium: BUL = Bulgaria; CZE = Czech Republic; GER = Germany; CRO = Croatia; HUN = Hungary; POL = Poland; ROU = Romania.

Major catchment area	Countries	River catchments	Pop	N	H _N	H _D	S
Black Sea	AUT, BUL, CRO, GER, HUN, ROU	Danube	30	106	16	0.794±0.024	17
North Sea	CZE, GER, BEL	Eider, Elbe, Rhine, Meuse, Weser	50	241	11	0.299±0.038	14
Baltic Sea	GER, POL	Uecker, Odra, Wisla, Wiprza	16	69	3	0.163±0.058	2

H_D-values (Table I) were lower for the rivers in central Europe (North Sea: $H_D = 0.299 \pm 0.038$ and Baltic Sea: $H_D = 0.163 \pm 0.058$) than for the Black Sea catchment ($H_D = 0.794 \pm 0.024$). Pairwise comparisons between river catchments revealed the lowest ΦST-value between the North and the Baltic Seas (ΦST = 0.035), and the highest value between the Black and Baltic Seas (Φ ST = 0.204, Table V). The ΦST-value between the North Sea and the Black Sea was 0.161. All values were highly significant (p < 0.001). The results of the AMOVA based on the three hierarchical groups (the major catchment areas) are given in Table III and reveal highly significant genetic differentiation between the major catchment areas (Φ ST = 0.290, p < 0.001) and among river catchments within the major catchment areas (FSC = 0.252, p < 0.001). H_D -values for river catchments that open into the North Sea ranged between 0.000 (Weser) and 0.400 (Ems), although these minimum and maximum values could be due to low sample sizes (Weser: N = 12; Ems: N = 5; see Table IV). HD for the Rhine (N = 161) was 0.227. Pairwise Φ ST-values for the North Sea major catchment area (Table VI) revealed very low genetic differentiation between sampled rivers. However, the Eider exhibited a high differentiation compared to all of the other rivers of the North Sea catchment area (Φ ST-value between 0.575 and 0.876; p < 0.001).

Table II Sampling locations and haplotypes of analyzed *Astacus astacus* and the number of the most common haplotype (H01) and the number of all other haplotypes. WIE = Wieprza, WIS = Wisla, DA = Danube, RH = Rhine, G. BS = German Baltic Sea estuary, P. BS = Polish Baltic Sea estuary, BAS = Baltic Sea, BS = Black Sea, NS = North Sea. n.a. = not available. For country codes, see Table I.

Region	[N°]	[E°]	Country	River district	Estuary	N	H01	Haplotype
Barthe	54.333	12.833	GER	G. BS	BAS	5		H02(5)
Tonkuhle	54.333	12.833	GER	G. BS	BAS	3	3	
Lake close to Miastko	54.017	16.983	PO	P. BS	BAS	10	10	
Lake close to Miastko	54.017	16.983	PO	P. BS	BAS	7	7	
Lake close to Miastko	54.017	16.983	PO	P. BS	BAS	8	8	
Rakowe Duze	53.915	16.796	PO	Odra	BAS	5	5	
Koppelsee	53.307	13.855	GER	Uecker	BAS	3	3	
Seki	54.065	17.095	PO	WIE	BAS	5	5	
Bez Nazwu	54.067	17.083	PO	WIE	BAS	4	4	
Rosko	53.903	17.143	PO	WIS	BAS	5	4	H03(1)
Czarne	50.038	21.993	PO	WIS	BAS	6	6	
Tomczyna	50.036	22.004	PO	WIS	BAS	8	8	
Freundsheimer Weiher	47.304	10.954	Α	DA	BS	3	2	H18(1)
Razdvec	43.463	24.908	BU	DA	BS	6	4	H13(1)
Beli Osam	12 858	24.655	BU	DA	BS	3		H15(1) H14(3)
Deli Osaili								H14(4)
Gorna Trape	43.444	25.073	BU	DA	BS	6	1	H16(1)
								H05(2)
Farm Augsburg	48.359	10.906	GER	DA	BS	14	9	H06(1)
								H20(2)
Wielenbach	47.841	10.952	GER	DA	BS	2	2	
Vukovina	45.693	16.109	HR	DA	BS	5		H17(5)
Kádárta	47.120	17.962	HU	DA	BS	2	2	
Pét	47.184	18.096	HU	DA	BS	1		H07(1)
Clocotici	45.241	21.838	RO	DA	BS	8		H20(7)
								H12(1)
Bradisoru de Jos		21.768	RO	DA	BS	1		H11(1)
Carasova		21.952	RO	DA	BS	1		H11(1)
Carasova		21.943	RO	DA	BS	1		H11(1)
Garliste		21.873	RO	DA	BS	5		H11(5)
Anina		21.887	RO	DA	BS	2	2	
Ocna de Fier		21.759	RO	DA	BS	4	1	H11(3)
Forotic		21.574	RO	DA	BS	6		H11(6)
Carnecea		21.636	RO	DA	BS	1		H11(1)
Hartagani		22.925	RO	DA	BS	1		H21(1)
Balsa	46.019		RO	DA	BS	1	1	
Baita		22.884	RO	DA	BS	2	2	
Bacaia	46.01	23.171	RO	DA	BS	2	2	
Balsa	46.04	23.07	RO	DA	BS	2	1	H20(1)

Region	[N°]	[E°]	Country	River district	Estuary	N	H01	Haplotype
Tamasesti	46.022	22.507	RO	DA	BS	3	3	
Almasu Mic de Munte	46.052	23.072	RO	DA	BS	3	2	H19(1)
Cladova	46.151	21.357	RO	DA	BS	4	2	H04(2)
Araneag	46.21	21.357	RO	DA	BS	2	1	H20(1)
								H12(1)
Nadas	46.206	21.359	RO	DA	BS	14		H20(12) H11(1)
Conop	46.122	21.359	RO	DA	BS	1		H20(1)
Farm Oeversee	54.688	9.422	GER	Eider	NS	5		H09(4) H10(1)
Langsee	54.575	9.593	GER	Eider	NS	9		H09(8) H10(1)
Svetlohor	48.977	13.736	CZ	Elbe	NS	7	7	
U sudu	49.822	15.358	CZ	Elbe	NS	6	4	H18(2)
Kramata	49.043	13.758	CZ	Elbe	NS	7	7	
Stepenitz	53.037	12.042	GER	Elbe	NS	5	5	
Schwarze Elster	51.582	13.97	GER	Elbe	NS	6	6	
Jäglitz	52.96	12.741	GER	Elbe	NS	7	5	H17(2)
Gut Rietberg	51.802	8.435	GER	Ems	NS	5	4	H20(1)
Florenville	49.700	5.307	BE	Meuse	NS	5	5	
Libramont	49.927	5.371	BE	Meuse	NS	4	4	
n.a.	n.a.	n.a.	BE	Meuse	NS	2	1	H12(1)
Fischbach	49.087	7.674	GER	RH	NS	6	3	H23(3)
Klausbach	52.272	7.910	GER	RH	NS	7	3	H20(4)
Mahlscheid	50.778	7.972	GER	RH	NS	8	8	
Zoo Zajak, Petstore	51.488	6.810	GER	RH	NS	4	4	
Aar	50.675	8.487	GER	RH	NS	5	5	
Allna	50.791	8.591	GER	RH	NS	4	4	
Ambach	50.702	8.275	GER	RH	NS	5	5	
Mühlgraben Caldern	50.842	8.658	GER	RH	NS	3	2	H05(1)
Dautphe	50.838	8.525	GER	RH	NS	3	3	
TA Donsbach	50.708	8.242	GER	RH	NS	5	5	
Eichelbach	50.459	9.123	GER	RH	NS	5	5	
Fohnbach	50.636	8.622	GER	RH	NS	5	5	
Gansbach	50.804	8.409	GER	RH	NS	5	5	
TA Geierstein/ Roth	50.641	8.226	GER	RH	NS	5	5	
Giebelsbach	50.997	9.012	GER	RH	NS	1	1	
Kallenbach	50.582	8.226	GER	RH	NS	10	10	
Lasterbach/ TA Mabüll	50.591	8.142	GER	RH	NS	5	5	
Merzkrebse	50.336	7.976	GER	RH	NS	9	9	
Meerbach	50.706	8.482	GER	RH	NS	1	1	
Madenmühlen	50.624	8.143	GER	RH	NS	5	5	
Mademühlen 2	50.624	8.143	GER	RH	NS	2	2	
Nanzenbach	50.780	8.489	GER	RH	NS	1	1	
Perf	50.873	8.459	GER	RH	NS	5	5	
TA Hartmann/ Rehbach	50.624	8.120	GER	RH	NS	5	5	

Region	[N°]	[E°]	Country	River district	Estuary	N	H01	Haplotype
Salzbach	50.753	8.539	GER	RH	NS	5	5	_
TA bei Spielberg	50.304	9.298	GER	RH	NS	5	2	H20(3)
Steinbruch Rot/ Schönbach	50.654	9.219	GER	RH	NS	5	5	
Stippbach	50.704	9.359	GER	RH	NS	5	5	
Pollichia Woog	49.103	7.719	GER	RH	NS	2		H20(2)
Pollichia Woog2	49.103	7.719	GER	RH	NS	2	1	H20(1)
Schlettenbachtal, Fischbach	49.088	7.711	GER	RH	NS	1		H20(1)
								H08(1)
Wolfsägertal	49.106	7.701	GER	RH	NS	7	2	H20(2) H22(2)
Meisertalweiher	49.335	7.769	GER	RH	NS	5	5	
Waldteich Irrschelde	50.763	8.399	GER	RH	NS	5	5	
Waldteich bei Wallenfels	50.774	8.438	GER	RH	NS	5	5	
Clausthal-Zellerfeld*	51.809	10.352	GER	Weser	NS	1	1	
Breitweiher/Rhön	50.507	9.730	GER	Weser	NS	2	2	
Ocherbach	50.673	9.185	GER	Weser	NS	5	5	
Urff	51.041	9.991	GER	Weser	NS	4	4	

^{*} This sequence was obtained from a preserved specimen of the Senckenberg Museum in Frankfurt (SMF 13095).

Table III Results of the analysis of molecular variance (AMOVA). Shown are the percentage of the total variance (% Var), fixation indices (F) and their significance (***: p < 0.001)based on 1000 random permutations.

Variance components	d.f.	% Var	F-statistics
Among major catchment areas	2	5.06	Φ _{CT} =0.051
Among river catchments within major catchment areas	9	23.92	Φ _{SC} =0.252***
Within river catchments	406	71.02	Φ_{ST} =0.290***

Table IV Diversity values for the North Sea sampling area. The number of sequenced individuals (N), the number of haplotypes (H_N) , the haplotype diversity (H_D) and the number of polymorphic sites (S) for seven rivers flowing into the North Sea are given.

River catchment	N	H _N	H _D	S
Elbe	38	3	0.199±0.084	8
Ems	5	2	0.400±0.237	2
Meuse	11	2	0.182±0.144	1
Rhine	161	6	0.227±0.043	5
Weser	12	1	0.000±0.000	0
Eider	14	2	0.264±0.136	1

Table V Pairwise $\sqrt{s_T}$ -values between the three major catchment areas and their significance (*: p < 0.05, ***: p < 0.001).

	Black Sea	North Sea	Baltic Sea
Black Sea	0		
North Sea	0.16131***	0	
Baltic Sea	0.20387***	0.03481***	0

Table VI Pairwise $\sqrt{\text{sr}}$ -values between five rivers flowing into the North Sea and their significance (*: p < 0.05, ***: p < 0.001).

	Elbe	Meuse	Rhine	Weser	Eider
Elbe	0				
Meuse	-0.011	0			
Rhine	0.042*	-0.018	0		
Weser	-0.016	0.008	0.007	0	
Eider	0.575***	0.816***	0.699***	0.876***	0

DISCUSSION

This is the first large-scale molecular study of Astacus astacus. The main aim was to determine the degree and origin (natural migration versus artificial translocation) of the genetic structure in noble crayfish populations across three major catchment areas. We detected very low haplotype diversities in noble crayfish populations in the North Sea and Baltic Sea catchments, with the exception of the Eider at the very northern end of Germany. The observed significant structure (Φsτ-values between 0.575 and 0.876) between the Eider and all other North Sea river catchments may be explained by an isolation of the Eider population. The Eider exhibits an east to west current, which is in contrast to the primarily south-north running rivers of southern Germany. This might result in a disconnection of the Eider from the other southern river systems, where the catchment areas of streams partly overlap and gene flow seems more likely. As a peripheral source with restricted gene flow, the Eider population contributes to the overall genetic diversity within A. astacus. To discover the origin of the private haplotypes solely found within this river catchment (indigenous or non-indigenous), additional populations nearby should be analyzed. Noble crayfish populations within the central European major catchment areas are significantly less differentiated than within the Black Sea catchment (Table I). A high haplotype

diversity and a high number of private haplotypes were detected for the Black Sea populations, suggesting a glacial refuge of the noble crayfish in the Balkan area. During the Pleistocene, severe climatic fluctuations in central Europe resulted in repeated range expansions and regressions of many taxa. The distribution of freshwater species was especially influenced by the fluctuating glaciers and varying water levels, which resulted in landscape changes and modified river systems (Hänfling et al., 2009). The recolonization of previously glaciated habitats (northern Germany) could have caused successive bottlenecks (Taberlet et al., 1998), which could explain the loss of genetic diversity and the observed dominance of haplotype H01 in the North Sea and Baltic Sea major catchments. A decrease in genetic diversity from southern to northern populations has been observed for many freshwater species, for example, the freshwater fish Vimba vimba (Hänfling et al., 2009) or the mayfly Ameletus inopinatus (Taubmann et al., 2011). The prevalence of haplotype H01 in the entire study area (Figure 1) may indicate a recolonization event from the Balkan area to central Europe. The fact that the North Sea and Baltic Sea major catchments only share one haplotype and show some degree of differentiation (Tables II and V) maybe interpreted as a result of two routes of recolonization from a common refuge. One migratory pathway into central Europe may have been the Danube drainage system after deglaciation, as presumed by Schulz and Grandjean (2005). The Danube represents a widely used corridor of postglacial expansion into central Europe for many freshwater species for which the possibility of a large-scale terrestrial dispersal has been excluded (e.g., Durand et al., 1999; Weiss et al., 2002; Hänfling et al., 2009). A second recolonization route, which would explain the divergence of the North Sea and Baltic Sea major catchments, cannot be confirmed due to the lack of sample sites from eastern Europe. The occurrence of noble crayfish in the Baltic Sea major catchment may also be the result of human translocation, although no evidence exists to support this theory.

Our study indicates that the original distribution pattern of the noble crayfish has changed significantly due to anthropogenic influence (Skurdal and Taugbøl, 2002), resulting in a mixture of indigenous and non-indigenous populations (Largiadèr *et al.*, 2000). Endemic haplotypes of the North Sea and Baltic Sea

catchments could be the result of artificial stocking, or using stock from a river catchment not included in our study (e.g., from Russia). However, presumably no stocking of native crayfish has been conducted in the Balkan States and the Ponto-Caspian region (Albrecht, 1983). We can therefore assume a natural origin of the studied populations from the Balkans (Maguire, 2009). The high frequency of private haplotypes in all major catchment areas, as well as the relatively high genetic differentiation between the catchment areas (ФST-values between 0.035 and 0.204), indicates a differentiation of populations throughout Europe despite the extensive human translocation of noble crayfish. Nevertheless, the artificial translocation of individuals can be observed in the disjunct distribution of some shared haplotypes between different drainage areas (Figure 1). Haplotype H20 could be an example of artificial distribution across central Europe. This haplotype was detected in the lower section of the Danube (Black Sea catchment), in the river catchments of the Rhine and Ems, and in individuals from a crayfish farm in Augsburg, Southern Germany (Table VII). This farm was founded by mixing four donor populations of the upper section of the Danube catchment area (Keller, pers. com.). Because restocking programs have been frequently conducted throughout Germany over the last few decades, in particular using stocking materials from the above-mentioned farm in Augsburg (Keller, 1999; Souty-Grosset and Reynolds, 2009), it is conceivable that specimens carrying haplotype H20 may originate from the Danube area and were artificially spread over central Europe. Human impact is also assumed to affect the (haplotype-) distribution of other European crayfish species, such as Austropotamobius pallipes (e.g., Grandjean et al., 2001; Diéguez-Uribeondo et al., 2008) or Austropotamobius italicus (Cataudella et al., 2010). Until today, the natural migratory routes of these species could not be fully inferred. However, the authors revealed geographical differences in the haplotype distribution and diversity, which may influence the conservation management strategies for these species.

With regard to *A. astacus*, the high frequency (74%) of private haplotypes in the Black Sea catchment highlights the importance of the Balkans as a reservoir for intraspecific genetic diversity. In addition to the topographical complexity, which allows for the isolation of single populations on a small geographical scale,

the Balkans were relatively unaffected by geological events during the glacial cycles. Thus, biodiversity is more conserved in the Balkans compared to central Europe for many species (Trontelj et al., 2005; Hänfling et al., 2009). Therefore, we strongly suggest a definition of several distinct ESUs along the Black Sea major catchment area. To define such ESUs, more populations of the Black Sea catchment area have to be analyzed because there might be much higher levels of genetic diversity and more detectable, endemic haplotyes to be found, for example, in the Upper Kolpa drainage similar to the recently detected distinct clades of Austropotamobius torrentium (Tronteli et al., 2005). So far, we suggest avoiding the intermixing of noble crayfish populations between the tributaries of the Danube. This is especially important for countries that plan to found crayfish farms for potential future restocking programs, e.g., in Bulgaria (Zaikov and Hubenova, 2007). We strongly emphasize that haplotype identification of the donor population should be conducted before a breeding stock is used for restocking. Additionally, the waters to be restocked should be carefully selected to avoid the contamination of local stocks. If restocking programs continue to translocate individuals with no regard for their population's genetic structure, the natural genetic make-up will further dissolve (Souty-Grosset et al., 2003), which is accompanied by a reduction in intraspecific diversity. Conservation strategies, therefore, need to manage populations of A. astacus as distinct ESUs and give the highest priority to the populations with high genetic diversity and unique haplotypes. Mitochondrial DNA is a widely used marker to reconstruct the phylogeographic history of species. Here, we show that the analysis of partial COI-sequences helps to understand the genetic structure of noble crayfish. However, the variation in noble crayfish COI-sequences is relatively low and the resolution is limited. Additional DNA-sequences should be included in the analysis to increase the resolution. Finally, a microsatellite analysis should be conducted to estimate genetic diversity within populations to give special attention to diverse populations and to further identify artificially stocked and naturally dispersed populations. However, as long as we cannot reject the hypothesis that private central European haplotypes developed due to local adaption, those populations that harbor endemic haplotypes are suggested to be candidates for special management plans to prevent the loss of unique haplotypes and to protect overall

genetic diversity. In conclusion, the results of this study suggest a glacial refuge in the Balkan area and a postglacial re-colonization of central Europe. Despite human translocations, which were revealed by the disjunct distribution of some haplotypes, a differentiation of noble crayfish populations in all major catchment areas supports the establishment of distinct ESUs to protect its present-day genetic diversity in Europe.

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Noble crayfish (Astacus astacus) with migratory background – natural versus human-mediated processes

Short title: European recolonisation of noble crayfish

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ABSTRACT

Conservation strategies aim to maintain the genetic diversity and the genetic integrity of a species. The phylogeographic history of a species can aid in defining areas of conservation priority. For freshwater species, the historical river structure plays a significant role in explaining the genetic differentiation and the population structure. Here, we aimed to reconstruct the post-glacial recolonization of central Europe of the endangered noble crayfish (*Astacus astacus*) and intended to identify refugial areas that are hotspots of genetic diversity. To address these questions, we analysed a fragment of the mitochondrial cytochrome oxidase subunit I (COI), and the 16S rRNA from 540 noble crayfish specimens from 158 sampling sites distributed throughout five European sea basins. Additionally, we conducted a microsatellite analysis of 289 individuals from 22 out of the 158 above-mentioned sites.

The haplotype diversity was highest in the Black Sea basin (H_D = 0.851, 28 haplotypes) and lowest in the Baltic Sea basin (H_D = 0.276, four haplotypes). The microsatellite data supported these results. Both markers revealed a particular high differentiation between populations from the Western Balkans to the remaining Black Sea populations. Western Balkan haplotypes diverged already around 710,890 years before present, whereas remaining differentiations occurred within the last 450,000 years before present. With migration modeling we detected that the North Sea basin and the Baltic Sea basin were colonized independently via different colonization paths from the eastern Black Sea basin,

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while the Western Balkans did not contribute to the colonization. The results suggest the existence of at least two refugial areas in southeastern Europe. To conserve the maximum genetic diversity, the focus of conservation prioritization for noble crayfish should be set in southeastern Europe. We further propose that each river catchment forms a separate management unit.

Key words: mtDNA sequences • microsatellite analysis • refugial area • postglacial colonization • migration model • conservation management

INTRODUCTION

The Convention on Biological Diversity (CBD) highlighted the preservation of biological diversity as one of its three main goals (SCBD 1992). According to the CBD, biological diversity includes the diversity of ecosystems, between species and within species (genetic diversity) (CBD Article 2). A high genetic diversity increases the chance of long-term survival because species with greater diversity are more likely to be able to survive in response to a changing environment compared to those with lower diversity (Malcolm *et al.*, 2007). Apart from the maintenance of the genetic diversity, the preservation of the native species' integrity (e.g. local specificity) is one of the major goals in conservation genetics (Souty-Grosset *et al.*, 2003). To protect the integrity and the within-species diversity a detailed knowledge of the phylogenetic structure of a species is indispensable. Present-day genetic structure is a result of climatic variation and associated geomorphological and hydrographic conditions in the past.

Geological events in Europe

Severe climatic oscillations and accompanying geomorphological and hydrographic conditions during the Pleistocene period resulted in great changes in species distribution areas characterised by contractions and expansions of their geographical ranges (Hewitt 1996). Southern Europe, especially the Iberian Peninsula, Apennine and the Balkans, served as a refuges during cold periods for many species (Hewitt 1999). During subsequent warm periods, a repeated northward expansion took place from refuges involving the leading edge (Hewitt 1996). Such a colonisation process of a limited number of individuals implies

successive founder effects that may lead to a loss of genetic diversity in the northern populations of cold-intolerant taxa (Taberlet *et al.*, 1998). The colonisation is followed by repeated extinctions of northern populations during cold climate phases (Hewitt 1996). In contrast, the diverse topography of southern refuges allows populations to diverge through several ice ages (Hewitt 1999).

The present distribution of freshwater organisms that often cannot migrate between river catchments has been most strongly influenced by the changes in landscapes caused by glaciers and melting water (e.g. Hänfling *et al.*, 2009; Vonlanthen *et al.*, 2007). The retreat of the glaciers formed the migration routes for freshwater species after the Last Glacial Maximum (LGM) that lasted from approx. 24,000 – 12,000 years before present (YBP). However, ephemeral contacts between river systems and changes in the flow direction of upper river systems complicate the reconstruction of the migration routes.

Over the LGM, freshwater crayfish were absent in central and northern Europe. They outlasted the glacial cycles on the Iberian Peninsula, the Italian Peninsula, the Balkan Peninsula or in the Ponto-Caspian region (Albrecht 1983). For noble crayfish (*Astacus astacus*), who probably survived the last glacial on the Balkan Peninsula (Albrecht 1983) under temperate conditions, a north- and westward spread along the Danube drainage system is most probable (Schulz and Grandjean, 2005) as it was shown for the freshwater fish species *Vimba vimba* (L, 1758) (Hänfling *et al.*, 2009), *Leuciscus cephalus* (L, 1758) (Durand *et al.*, 1999) or the European grayling *Thymallus thymallus* (L, 1758) (Weiss *et al.*, 2002). The recolonisation route reconstruction facilitates the identification of management units or evolutionary significant units (ESU) and is therefore relevant for conservation.

Noble crayfish - threats

The IUCN Red List of Threatened Species lists the noble crayfish as vulnerable with a decreasing population status (IUCN 2010). Besides environmental changes like water pollution or channelisation of streams, native crayfish species are highly threatened by invasive species. In particular, North American crayfish species pose a high threat to noble crayfish because they

potentially carry the oomycete fungus *Aphanomyces astaci* (Schikora 1906) that causes high mortalities in native crayfish populations (e.g. Alderman 1996). The high extinction rate goes along with a severe decrease in genetic diversity of European crayfish species. As the largest freshwater invertebrates and due to their trophic activities as omnivores, European crayfish play a key role in many freshwater ecosystems (Nyström 1999). The exctincion of local populations ay have drastic impacts on local biodiversity.

For more than 2000 years noble crayfish have been an object of commerce and trade (Skurdal and Taugbøl 2002) and are therefore influenced to the highest degree by human translocations (Albrecht 1983). The economic value of crayfish has led to noble crayfish stocking into numerous new localities. When the crayfish plague hit central Europe at the turn of the 19th century, noble crayfish were imported in large quantities from eastern European countries (Skurdal and Taugbøl 2002). Also today there are operating noble crayfish farms in several European countries, especially in central and northern Europe (e.g. Jussila and Mannonen 2004; Paaver and Hurt 2009) that sell crayfish as a food source, as stock for private ponds, or as donors for restocking programs in response to rapidly declining stocks. Stocking often occurs without knowing the taxonomic status of the stocking material (Souty-Grosset et al., 2003). As a consequence, the trade leads to translocations and disintegrates the natural genetic structure. In addition, the construction of channels connecting different major rivers catchments also strongly influenced the distribution of crayfish species (Albrecht 1983).

Previous study

The first molecular-based phylogeographic study of noble crayfish by Schrimpf et al. (2011) covered a large part of the distribution range and resolved the large-scale genetic structure of this species: the haplotype diversity of the cytochrome oxidase subunit I (COI) fragment in central Europe was very low compared to the high diversity in the Balkan area where a centre of range expansion was suggested. Interestingly, a strong divergence between populations from the Western Balkans (Croatia) and the eastern Black Sea basin (Romania,

Bulgaria) was found (Schrimpf *et al.*, 2011). The genetic divergence may indicate the existence of two separate glacial refuges inhabited by populations that did not exchange genes for several centuries (Hewitt 1999). However, since only five specimens were analysed from the Western Balkans a final conclusion could not be drawn. The obtained result could also be the consequence of a human-mediated noble crayfish translocation. This first phylogeographic study of noble crayfish (Schrimpf *et al.*, 2011) has also shown that one marker (COI) is not sufficient to resolve the genetic structure of noble crayfish and additional markers are needed.

Aim of this study

In this study we analysed a second mtDNA marker (16S rRNA) to increase the resolution of the phylogenetic analyses. Although mtDNA is useful for studying historical lineage splits in the context of geological events, only little or no signal can be detected in bottlenecked populations that have recently colonised an area (Hewitt 1999). Therefore, we utilised nuclear microsatellite data to track the recent gene flow among populations and to estimate within population diversity. Moreover, we enhanced the number of sampling sites in southeastern Europe, especially in the Western Balkans. With this study we aim to answer three main questions that are fundamental for the conservation of the maximum genetic diversity of noble crayfish: a) Did the noble crayfish survive the last glacial in more than one refugial area in southeastern Europe? b) From which refugium did the noble crayfish recolonise central Europe? c) Was the Baltic Sea basin recolonised from the North Sea basin or from the Black Sea basin?

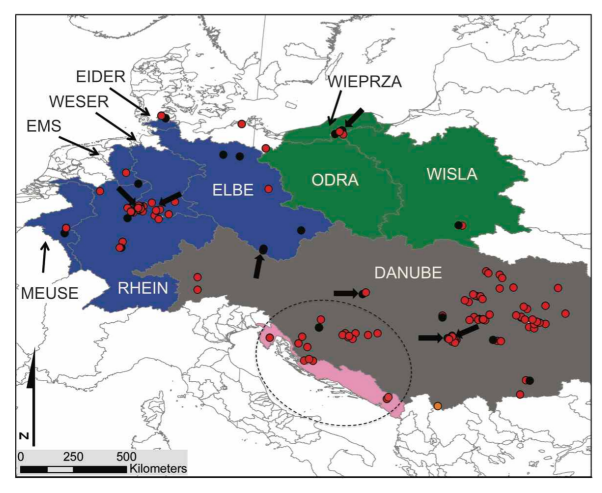


Figure 1 Study area with sampling sites. Sea basins of European rivers are colored in blue for the North Sea basin, in green for the Baltic Sea basin, in grey for the Black Sea basin and in pink for the Adriatic Sea basin. The orange circle shows the one sample from the Aegean Sea that was included in the sequence analysis. Red circles indicate populations that were sequenced and black circles indicate populations that were additionally genotyped. When the black circle is overlaid by another circle, a bold arrow point to the hidden circle. Thin arrows point to the respective river catchments. The dotted circle indicates the populations grouped as 'Western Balkans'. The two sampling sites from Finland (Valkeinen and Ylä river catchments) are not shown.

MATERIAL AND METHOD

Sampling

In total, 540 crayfish specimens from 158 sampling localities that are located within 14 river catchments (Figure 1, Table I in Supplementarly material) and from four hatcheries were collected by hand or by traps. The sampling sites were distributed within five different sea basins: 12 in the Baltic Sea basin, 59 in the North Sea basin, 87 in the Black Sea basin, four in the Adriatic Sea basin and one in the Aegean Sea basin. Immediately after the lower part of one pereopod

(propodus and dactylus) was taken specimens were released at the place where they were caught. Appendages usually regenerate after a few moults so the vitality of crayfish was not affected. Samples were stored in 70-96% Ethanol until DNA extraction. DNA was extracted from the muscle tissue using a standardized protocol ('Rapid isolation of Mammalian DNA', Sambrook and Russel, 2001).

Laboratory work

A 350 base pair (bp) fragment of cytochrome c oxidase subunit I (COI) and a 476 bp fragment of the 16S rRNA (16S), both mitochondrial DNA, were amplified using the primer pair ASTCOI (forward primer: GCGGGGATAGTAGGAACCTC; reverse primer: ATTTACCGCCCCTAAAATCG) and 16S 1471 and 16S 1472 (Crandall and Fitzpatrick 1996) respectively. The Polymerase chain reactions (PCR), the purification of the PCR products, the sequencing reaction and the sequencing were performed as described in Schrimpf et al. (2011). The annealing temperature in the PCR reaction for 16S was changed to 51° C. Sequences were edited and aligned with the SEQUENCHER 3.0 software (Gene Codes Corporation, Ann Arbor, Michigan, USA). Sequences were checked manually for base pair ambiguities, nuclear copies of mitochondrial derived genes, stop codons, and high levels of divergence, as recommended by Buhay (2009). All haplotypes were submitted to GenBank (Accessions numbers will be provided upon acceptation of the manuscript). As both COI and 16S share the same history, sequences were concatenated and treated as a single locus for the following analyses except for the calculation of the Bayesian tree.

For 289 individuals from 22 populations from 12 river catchments and the hatchery in Augsburg (Table 1 in supplementary material) a total of six microsatellite loci with a dinucleotide repeat were analysed using the species-specific primer pairs Aas2, Aas6, Aas8, Aas11, Aas766, Aas1198 (Koiv *et al.,* 2008, 2009). Forward primers were designed with a 19 bp M13-tail that was labelled during the PCR reaction using a universal fluorochrome-labelled M13 primer. PCR were carried out using a Primus 96 Cycler (Peqlab Biotechnologie GmbH, Erlangen, Germany) under the following conditions: An initial denaturation at 95°C for 2 min, followed by 35 cycles of 30 s at 95°C, 30 s at 60°C, 30 s at

72°C, and a final extension of 5 min at 72°C. The 60°C annealing was replaced by 63°C for primer Aas11 and by 57°C for primer Aas8. 2 μl PCR-product were added to 30 μl SLS (Beckman Coulter, Krefeld, Germany). The fragment analysis was performed on a Beckman Coulter CEQ 8000 eight capillary sequencer. Loci were scored using the software CEQ SYSTEM version 9 (Beckman Coulter, Krefeld, Germany). 10% of all samples were randomly chosen and repeated for estimation of the error rate (Bonin *et al.*, 2004).

Statistical analysis - Sequence analyses

The statistical analyses were calculated either for sea basins (e.g. North Sea basin) or river catchments (e.g. Rhine catchment) or for both. From the Black Sea basin we have only samples from the Danube river catchment, therefore the same samples were used for a sea basin comparison (as 'Black Sea basin') and for a river catchment comparison ('as Danube river catchment'). Genetic variation within sea basins and river catchments were measured in terms of the number of haplotypes (H_N), haplotype diversity (H_D) and number of segregating sites using DNASP v 5.10.1 (Librado and Rozas 2009). To detect genetic differentiation a hierarchical analysis of molecular variance (AMOVA, Excoffier et al., 1992) with populations from each river catchment partitioned into separate groups was performed and Φ_{ST} -values among sea basins and river catchments were estimated with ARLEQUIN v 3.11 (Excoffier and Lischer 2010). Significance was based on 1000 random permutations. To identify haplotypes and to determine the phylogenetic relationships between haplotypes a median joining (MJ) network (Bandelt et al., 1999) was constructed using the software NETWORK 4.610 (Fluxus Technology, Suffolk, UK). DNASP v 5.10.1 was used to perform a mismatch analysis, which plots the distribution of the number of differences between pairs of sequences and compares it with a fit to an unimodal Poisson distribution as expected under sudden expansion from a small population. An unimodal pattern in the graph indicates that a species underwent a recent population expansion. A multimodal, ragged pattern in the graph, however, indicates that this species has maintained a stable population size over a long period of time. The value of the calculated raggedness statistic r (Harpending et al., 1993) is lower in expanding population compared with a constant population

size. To test for deviation of sequence variation from evolutionary neutrality Tajima's *D* and Fu's F tests (Tajima 1989; Fu 1997) were carried out with ARLEQUIN v 3.11. Under a scenario of sudden expansion these values will be negative, and positive in a population that's suffered a recent decrease in population size (Excoffier *et al.*, 1992).

A phylogenetic tree was constructed using BEAST version 1.7.3 (Drummond et al., 2012). The molecular clock test was performed in MEGA v 5.05 (Tamura et al., 2011). The null hypothesis of equal evolutionary rate throughout the tree was rejected at a 5% significance level (P < 1.12). Therefore, we used a relaxed uncorrelated lognormal clock model. Lacking fossil data to calibrate a species specific molecular clock, we used a rate divergence for COI of 1.4% per million years considered as an approximate average divergence rate reported for crustaceans (Knowlton and Weight, 1998) that was recently used for the whiteclawed crayfish (Pedraza-Lara et al., 2010). Since no divergence rate is available for 16S we used an estimated molecular clock for 16S. We used a HKY substitution model (Hasegawa et al., 1985) with gamma distributed rate heterogeneity and Optimum heterogeneity parameters with empirical base frequencies, and 5 rate categories for both partitions as proposed by Treefiner (Jobb 2011). The relative fit of each model to the data was assessed using the Akaike Information Criteria (AIC, Akaike 1973). The starting tree was randomly generated with a Birth-and-Death process prior on the tree. The Markov Chain Monte Carlo (MCMC), 400 million steps long, was performed sampling every 10,000th generation. The BEAST-run was visualized and diagnosed with Tracer version 1.5 (MCMC Trace Analysis Tool, Rambaut and Drummond, 2009). TreeAnnotator version 1.7.3 (Drummond and Rambaut 2011) was used for summarising the information in a sample of trees onto a single 'target' tree whereas the first 10% of the samples were discarded as burn-in. The consensus tree was finally put into graphs with FigTree version 1.3.1 (Rambaut 2009). The cold climate phases as revealed by benthic oxygen isotope stages (δ¹⁸O, Lisiecki and Raymo 2005) were incorporated into the tree figure.

To compare the probability of different dispersal routes, we applied a model selection approach implemented in MIGRATE-N v 3.3.2 (Beerli and Felsenstein

2001). To keep computation feasible the, 152 sampling sites were pooled according to geographic patterns (sea basins) and previous results. We tested ten hypotheses that were translated into migration matrices. The first genealogy was started using a random tree. Initial theta and migrant values were generated from a F_{ST} calculation. A static heating scheme with four different temperatures was applied. We ran 1 mio generations, from which 50,000 were sampled after a burnin period of 10,000. For the log-equivalent Bayes factor (LBF) estimation we used the thermodynamic integration (T.I.) approximation using a Bezier-curve because it results in LBFs with high repeatability and little variance (Beerli and Palczewski 2010). A high T.I. value indicates a better fit of the model than a low T.I.

Statistic analysis - Microsatellite analyses

For the nuclear DNA markers the GENETIC DATA ANALYSIS (GDA) v 1.1 software (Lewis and Zaykin 2001) was used to calculate the number of private alleles (A_P) per sea basin, river catchment and population. The number of alleles (A_N) and the expected (H_E) and observed (H_O) heterozygosity for the sea basins, the river catchments and populations were performed in ARLEQUIN version 3.5.1.3 (Excoffier and Lischer 2010). The ARLEQUIN software was further used to perform an AMOVA (Excoffier et al., 1992) with populations from each river catchment partitioned into separate groups, and to calculate F_{ST}-values between river catchments and sea basins. The structure of and the variation among all genotyped individuals were determined by a factorial correspondence analysis (FCA) with the default settings in GENETIX 4.05 (Belkir et al., 1996-2004). The genetic population partitioning was evaluated using the Bayesian clustering approach in the program BAPS v 5.4 (Corander et al., 2008). We used predefined sampling units whereas all individuals of a population form one group. The program was initially run with a fixed number of clusters (K = 22). The analysis was repeated five times to determine to most likely number of K. With a probability of 1 the number of K was 15. In a second analysis, five runs, each at K = 10 to K = 20, were performed to confirm the number of clusters.

RESULTS

Sequence data

The 540 sequences of the COI and 16S genes resulted in 30 and 15 different haplotypes, respectively. COI was more diverse with H_D = 0.548 and 27 polymorphic sites, and 16S was less diverse with H_D = 0.410 and 15 polymorphic sites. The concatenated sequences with a length of 826 bp resulted in 46 different haplotypes with 42 polymorphic sites. The mean H_D for the entire sample set was 0.626. The diversity values for the sea basins suggested that populations of the Black Sea basin were highly diverse (H_D = 0.851) representing 28 haplotypes (Table 1). The Adriatic Sea basin is also characterised by a high genetic diversity (H_D = 0.576, five haplotypes), whereas it has to be noted that only 12 samples from the Adriatic Sea basin were included. The North Sea basin had a lower haplotype diversity (H_D = 0.316) and less haplotypes (13). The Baltic Sea basin had the lowest genetic diversity (H_D = 0.276, four haplotypes).

The results of the hierarchical AMOVA are shown in Table 2. Test indicated that the majority of variance was present among populations within river catchments (62.50%, p < 0.001). In total, 25.33% (p < 0.001) is attributed to variation within populations while variation was moderate between river catchments (12.16%, p < 0.05). The Φ_{ST} -values for mtDNA suggested that the sea basins were significantly different (p-values < 0.05; Table 3). The Φ_{ST} -values between the Aegean Sea basin (N = 1) on the one side and the North and Baltic Sea basins on the other side are high (Φ_{ST} = 0.789 and 0.943) but not significant. With regard to the river catchments, noble crayfish from the Eider were significantly different from all river catchments except the Valkeinen river catchment (Table 4).

Table 1 Results from the combined COI and 16S mitochondrial sequences for the analysed sea basins (BAS = Baltic Sea, NS = North Sea, BS = Black Sea, AdS = Adriatic Sea; above line) and the river catchments (below line). The Black Sea basin is also treated as a river catchment (Danube). Given are the number of sequenced individuals (N), the number of haplotypes (H_N), the haplotype diversity (H_D) (calculated with DNASP v 1.10.1) and the number of polymorphic sites (S), values for Tajima's D, Fu's F and Raggedness (r) (calculated with ARLEQUIN v 3.11). Significant values are indicated in bold (p < 0.05). The samples from the Aegean Sea basin (AeS) are excluded due to the low sample size (N = 1). Presented are the proposed groups for the calculations with MIGRATE-N (F = Finland, EBS = eastern Black Sea basin, WB = Western Balkans).

Sea basin	River catchment	Migrate	Sites	N	H _N	H _D	s	Tajima´ s D	Fu´s F	r
BAS			14	66	5	0.276	4	-1.39	-3.34	0.30
NS		NS	59	249	13	0.316	24	-1.93	-4.88	0.36
BS		EBS/ WB	82	185	28	0.851	32	-0.79	-6.29	0.03
AdS		WB	5	12	5	0.576	14	-0.52	1.58	0.23
BAS	Odra	BAS	1	5	1	0	0	0.00	0.00	0.00
BAS	Uecker	BAS	1	3	1	0	0	0.00	0.00	0.00
BAS	Wieprza	BAS	2	10	1	0	0	0.00	0.00	0.00
BAS	Wisla	BAS	3	15	3	0.362	2	-1.00	-0.92	0.19
BAS	Valkeinen	F	1	5	2	0.600	1	1.22	0.626	0.4
BAS	Ylä	F	1	5	1	0	0	0.00	0.00	0.00
NS	Eider	NS	1	9	2	0.222	1	-1.09	-0.26	0.36
NS	Elbe	NS	6	39	3	0.194	13	-1.53	3.36	0.68
NS	Ems	NS	1	5	2	0.400	3	-1.05	1.69	0.68
NS	Meuse	NS	2	9	2	0.389	1	0.16	0.48	0.20
NS	Rhine	NS	41	176	11	0.349	18	-1.82	-2.17	0.48
NS	Weser	NS	3	11	1	0	0	0.00	0.00	0.00

Table 2 Results of the analysis of molecular variance (AMOVA) calculated with ARLEQUIN v 3.11 based on mtDNA and microsatellite allele frequencies where populations are grouped according to European river catchments. The percentage of the total variance (% Var), fixation indices and their significance (*: p < 0.05 ***: p < 0.001) based on 1000 random permutations are shown. pop. = populations, Micsat = Microsatellite data.

DNA marker	Among	groups	Among po	op. within groups	Within	Within pop.		
	%	F-statistic	%	F-statistic	%	F-statistic		
mtDNA	12.16	0.122*	62.5	0.712***	25.33	0.747***		
Micsat	15.89	0.159***	29.85	0.354***	54.26	0.457***		

Table 3 Pairwise FST- and Φ ST-values between European sea basins as implemented in ARLEQUIN v 3.11. Results for microsatellites are shown above the diagonal, mtDNA data is shown below the diagonal (for abbreviation see Table 1). Significant values are indicated in bold (p < 0.05).

	BAS	NS	BS	AdS	AeS
BAS	0.000	0.365	0.283	-	-
NS	0.071	0.000	0.231	-	-
BS	0.170	0.162	0.000	-	-
AdS	0.855	0.789	0.438	0.000	-
AeS	0.943	0.789	0.162	0.291	0.000

Table 4Pairwise F_{ST} - and Φ_{ST} -values between river catchments and Finland calculated with Arlequin v 3.11. Results for microsatellite data is shown above the diagonal and mtDNA data is shown below the diagonal. Significant values are indicated in bold (p < 0.05).

	Ems	Elbe	Eider	Meu.*	Wes.*	Rhine	Uec.*	Wie.*	Odra	Wis.*	Dan.*	Val.*	Ylä
Ems	0.000	0.311	0.181	0.460	-	0.008	-	0.171	0.293	0.199	0.179	0.165	0.195
Elbe	-0.019	0.000	0.207	0.095	-	0.217	-	0.273	0.224	0.462	0.341	0.157	0.215
Eider	0.656	0.422	0.000	0.429	-	0.168	-	0.149	0.096	0.422	0.302	0.038	0.122
Meu	0.092	0.008	0.771	0.000	-	0.304	-	0.556	0.382	0.723	0.400	0.286	0.377
Wes.*	0.170	-0.014	0.910	0.156	0.000	-	-	-	-	-	-	-	-
Rhine	-0.086	0.057	0.507	0.053	0.018	0.000	-	0.163	0.249	0.188	0.201	0.127	0.169
Uec.*	-0.132	-0.166	0.845	-0.068	0.000	-0.119	0.000	-	-	-	-	-	-
Wie.*	0.149	-0.020	0.905	0.141	0.000	0.013	0.000	0.000	0.230	0.378	0.281	0.091	0.134
Odra	0.000	-0.080	0.870	0.034	0.000	-0.040	0.000	0.000	0.000	0.502	0.347	0.102	0.230
Wis.*	0.169	0.022	0.804	0.112	-0.013	0.051	-0.173	-0.019	-0.084	0.000	0.346	0.260	0.280
Dan.*	0.031	0.105	0.263	0.116	0.116	0.162	0.003	0.112	0.067	0.131	0.000	0.255	0.254
Val.*	-0.071	0.032	0.763	0.231	0.441	0.48	0.118	0.417	0.250	0.225	0.667	0.000	0.035
Ylä	0.000	-0.080	0.870	0.034	0.000	-0.040	0.000	0.000	0.000	-0.062	0.110	0.250	0.000

River catchments abbreviationed as following: Meu. - Meuse, Wes. - Weser, Uec. - Uecker, Wie. - Wieprza, Wis. - Wisla, Dan. - Danube, Val. - Valkeinen

The network showed a star-like pattern around the most common haplotype (Hap01, Figure 2) that was found in 327 samples distributed throughout the sampling area except in the Adriatic and Aegean Sea basins. Additionally, the network revealed a strong branching around two haplotypes that are common in the Black Sea basin only. A group of haplotypes from the Western Balkans (Hap41 – Hap46) formed a subsection with a distance of at least 8 bp to all other haplotypes. Four haplotypes were shared between North Sea and Baltic Sea basins, two between Adriatic Sea and Black Sea basins, one between the Baltic Sea (Finland) and the Black Sea basin and one between North Sea, Black Sea

and Baltic Sea basins. We found private haplotypes in all sea basins, seven in the North Sea basin three in the Baltic Sea basin, three in the Adriatic Sea basin, one in the Aegean Sea basin (N = 1) and 24 in the Black Sea basin, of which four were found only in the hatchery in Augsburg (Table I in Supplementarly material).

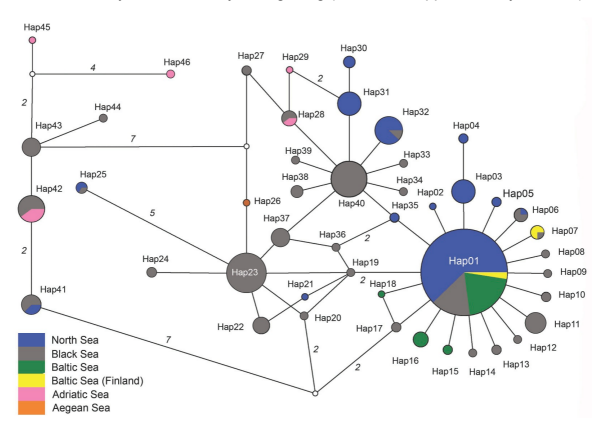


Figure 2 Median joining network constructed using NETWORK 4.610 for 826 bp of the concatenated COI- and 16S-sequences reveals genealogical relationships among 540 *Astacus astacus* samples. Each connecting branch line represents nucleotide substitutions. The number of mutational steps is given above the branch, except when it equals 1. Hollow circle represent missing haplotypes inferred from mutational changes, but absent in this data set. The size of circles is proportional to the frequency of the represented haplotype. Haplotype codes on the network correspond to samples listed in Table 1. Finnish samples correspond to the Baltic Sea basin, but were highlighted because of their high geographic differentiation to the other Baltic Sea basin samples.

The unimodal ragged pattern in the pair-wise differences graph of the mismatch analysis for the whole data set was shifted to the left of the distribution (graph not shown) and very closely matched the expected distribution for a recent population expansion (Rogers and Harpending 1992). Raggedness statistic for the whole data set was low and not significant (r = 0.088, p = 0.428) and therefore we cannot reject the hypothesis of population expansion. Mismatch distribution for

the sea basin groups revealed unimodal patterns for the North Sea and Baltic Sea basins, while multimodal and ragged distributions were detected for the Black Sea and Adriatic Sea basin (graphs are not shown). Fu's F was significant negative (p < 0.05) for the Baltic Sea and Black Sea basin and Fu's F and Tajima's D were both significant (p < 0.05) for the North Sea basin (Table 1). With regard to the river catchments, Tajima's D was significant negative for populations from the Rhine and Elbe river catchments (Tajima's D = -1.82 and -1.53, respectively; p < 0.05).

In the haplotype phylogeny inferred from the combined COI and 16S sequences haplotypes clustered into two well-supported major clades (posterior probability, pp = 1) (Figure 3). The lineage 4 contained six haplotypes from the Adriatic Sea and Black Sea basin (all Western Balkans). Haplotypes from the Black Sea basin were present in all four lineages and haplotypes from the North Sea basin in all lineages except in the lineage 4. Haplotypes from the Baltic Sea basin cluster only within one clade (lineage 1). Most splits were not significant. However, the split between lineage 4 and lineages 2 - 3 and between lineage 3 and 1 - 2 is highly supported (pp = 1 and 0.93, respectively).

For the model selection calculated with MIGRATE-N, we tested several combinations of sample groups. Due to the genetic similarity and the relatively small geographic distance between populations from the south-western Black Sea basin (Croatia) and the Adriatic Sea basin, we pooled this data (dotted circle in Figure 1). This group will be referred to as 'Western Balkans' and the remaining Black Sea basin as 'eastern Black Sea basin'. The Aegean Sea basin was excluded from this analysis due to the low sampling size. Because of the high geographic distance, we further separated Finland from the Baltic Sea basin. The remaining geographic group was North Sea basin. The probability values/ Bayes factors preferred the model 2b (T.I. = -2071.61) (Figure 4). This model implies a bifurcal colonization route from the eastern Black Sea basin to a) the North Sea basin and to b) the Baltic Sea basin. According to the favoured model the eastern Black Sea basin was colonised from the Western Balkans and the Finnish sampling sites were colonized from the Baltic Sea basin.

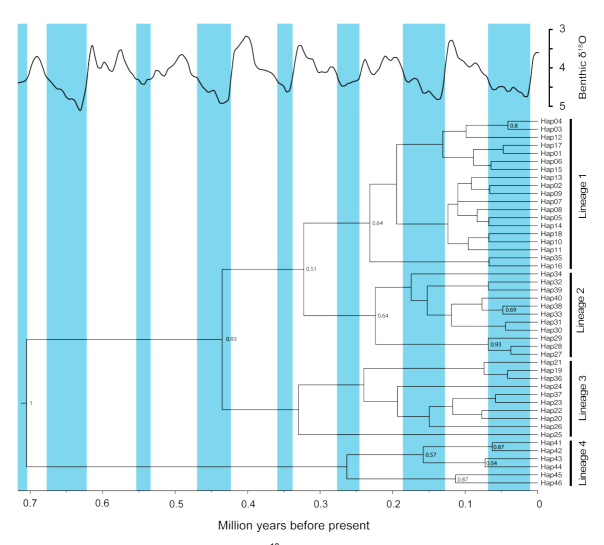


Figure 3 Benthic oxygen isotope stages (δ^{18} O; Lisiecki and Raymo, 2005; top off the figure) and haplotype phylogeny generated using Bayesian Markov Chain Monte Carlo analysis as implemented in BEAST. The calculation was based on a combined 350 bp fragment of the COlsequence and a 476 bp fragment of the 16S sequence. The axis shows the time scale in million years before present. Number at the nodes are the posterior probability values above 0.5.

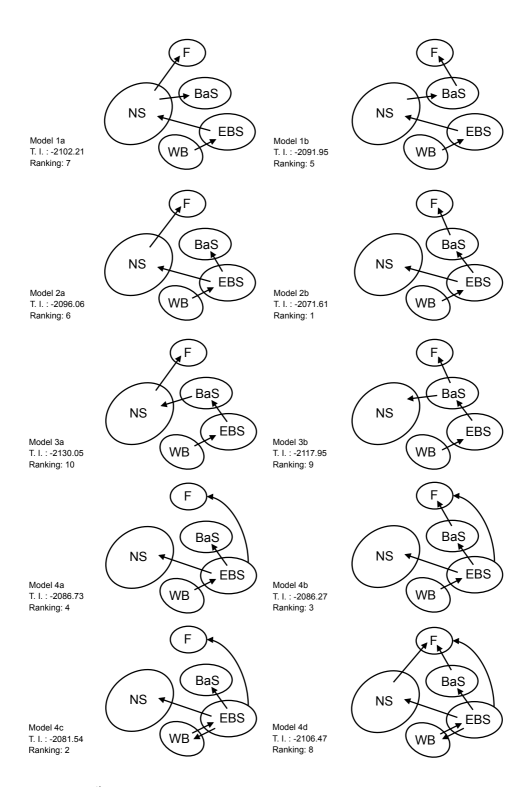


Figure 4 Results of the migration hypothesis model selection determined with MIGRATE-N v 3.3.2 between different regions according to geographic regions and genetic results. Finland (F) was separated from the Baltic Sea basin (BaS) and Croatia was separated from the Black Sea basin and pooled with samples from the Adriatic Sea basin to form the group 'WB' = Western Balkan. The remaining Black Sea basin samples form the 'EBS' = eastern Black Sea. NS = North Sea basin. Arrows indicate the directions of gene-flow. Thermodynamic Integration (T.I.) gives the fit of the model. The ranking from 1 to 10 for the models is stated. Note that only ten samples were included from Finland.

Microsatellite data

A genotyping error rate of 1.8% was estimated and should thus not bias our results. A total of 100 alleles with an average of 16.67 alleles per locus were observed. The highest H_E (0.799) and the highest number of alleles per number of samples analysed ($A_N/N = 0.92$) were detected in the Black Sea basin (Table 5). The North Sea basin ($H_E = 0.543$) is indicated by a higher heterozygosity value than the Baltic Sea basin ($H_E = 0.414$), but by a lower number of alleles compared to the number of analysed samples ($A_N/N = 0.49$) (Table 5). With regard to the individual populations, the populations from the Black Sea have higher expected heterozygosity values (mean $H_E = 0.456$) than the populations from the North Sea basin (mean $H_E = 0.320$) and Baltic Sea basin (mean $H_E = 0.231$).

In the hierarchical AMOVA tests with populations from each river catchment partitioned into separate groups the majority of variation was present within populations (54.26%, p < 0.001; Table 2), variation was moderate within river catchments (29.85%, p < 0.001) and minor among river catchments (15.89%, p < 0.001). The F_{ST} -values suggested that the sea basins were significantly differentiated (p < 0.05; Table 3). All river catchments among and within sea basins differed significantly from each other apart from the Rhine and the Ems (F_{ST} = 0.008, p < 0.05; Table 4), the Valkeinen and Eider (F_{ST} = 0.030, p < 0.05) and the Valkeinen and Wieprza.

The first two factorial components of the FCA explain 10.98% of the variance of the microsatellite data. The data points from the Black Sea basin individuals are distributed most widespread in the FCA graph (Figure 5) indicating a high genetic variation. The data points from the North Sea basin are more widespread than those from the Baltic Sea basin. There is an overlap of individuals from the North Sea and Baltic Sea basin. Data points from the Western Balkan samples clearly group apart from all other samples.

The Bayesian cluster analysis based on river catchments selected a clustering of K = 15 with a probability of 0.92 as the best model explaining the population structure (Table 5). The geographically distant Finnish populations (Baltic Sea basin) form one cluster. The hatchery in Augsburg groups in the same cluster as one population from the Rhine and one from the Ems River catchment.

One population from the Elbe, the Meuse and the Rhine form another cluster. Furthermore, two Elbe populations group in one cluster and one Wieprza population and one Eider population in another. For the remaining 11 populations (all from the Danube River catchment) each population was considered as a unique group.

Table 5 Results from the microsatellite analysis for the analysed sea basins (above line, for abbreviation see Table 1) and sampling sites (below line). The number of genotyped individuals (N), the number of private alleles (A_P) and private alleles per N (A_P/N) , the number of alleles (A_N) and alleles per N (A_N/N) and the expected (H_E) and observed heterozygosity (H_O) calculated with ARLEQUIN v 3.11 are given. The same capital letter (A-F) in the column 'Cluster' indicates populations that belong to the same cluster as evaluated using the Bayesian clustering approach in the program BAPS v 5.4. The 'x' indicates populations that form their own cluster.

	Sampling site	-	Sea basin	N	A _N	A _N /N	Ap	A _p /N	HE	Ho	Cluster
Sin			BAS	62	39	0.63	4	0.11	0.414	0.156	
ı basin			NS	124	61	0.49	10	0.08	0.543	0.228	
Sea			BS	79	73	0.92	35	0.44	0.799	0.319	
	Rakowe Duze	Odra	BAS	15	16	1.07	3	0.20	0.239	0.160	х
	Bez Nazwu	Wieprza	BAS	9	13	1.45	1	0.11	0.247	0.239	Α
	Czarne	Wisla	BAS	11	11	1.00	0	0.00	0.153	0.100	Х
	Valkeinen	Kymen	BAS	13	13	1.00	0	0.00	0.226	0.161	Е
	Ylä	Kokemäenjoki	BAS	14	13	1.00	0	0.00	0.291	0.218	Е
	Florenville	Meuse	NS	15	10	0.67	0	0.00	0.197	0.046	В
	Jäglitz	Elbe	NS	7	9	1.29	0	0.00	0.147	0.074	В
	Svetlohor	Elbe	NS	10	12	1.20	0	0.00	0.216	0.147	С
	Kramata	Elbe	NS	8	11	1.37	1	0.13	0.218	0.128	С
River catchment	U sudu	Elbe	NS	13	17	1.31	0	0.00	0.377	0.244	D
ä	Ambach	Rhein	NS	18	25	1.39	0	0.00	0.491	0.456	D
zatc	Merzkrebse	Rhein	NS	5	12	2.40	0	0.00	0.296	0.200	В
er (Dauphe	Rhein	NS	19	35	1.84	6	0.32	0.531	0.368	F
.≥	Gut Riedberg	Ems	NS	15	23	1.53	0	0.00	0.503	0.344	F
	Langsee	Eider	NS	14	15	1.07	1	0.07	0.220	0.165	Α
	Clocotici	Danube	BS	8	22	2.75	7	0.88	0.511	0.494	х
	Gorna Trape	Danube	BS	22	20	0.91	1	0.05	0.387	0.349	Х
	Vukovina Lake	Danube	BS	6	16	2.67	7	1.17	0.550	0.306	Х
	Ciornovãt	Danube	BS	5	16	3.20	6	1.20	0.479	0.283	Х
	Kádárta	Danube	BS	4	10	2.51	0	0.00	0.327	0.125	х
	Dragoiestilor	Danube	BS	22	27	1.23	5	0.23	0.452	0.355	х
	Nadas	Danube	BS	12	18	1.50	1	0.08	0.487	0.353	х
	Augsburg	Hatchery		24	32	1.33	1	0.04	0.550	0.439	F

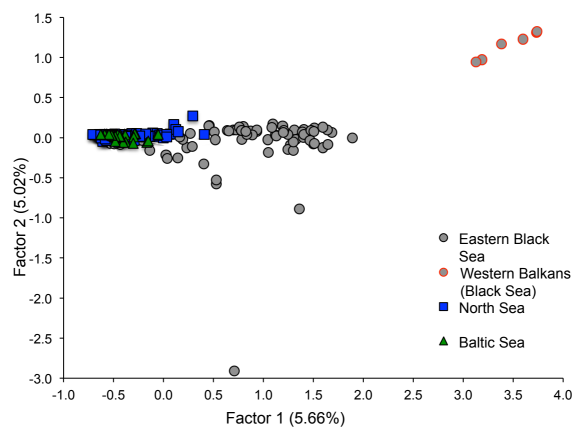


Figure 5 Factorial Correspondence Analysis over determined using GENETIX showing the degree of similarity of the 289 *Astacus astacus* individuals across six microsatellite loci based on the first two dimensions (factors). Each dot represents one individual. The distance between points reflects the degree of genetic differentiation among individuals.

DISCUSSION

Conservation strategies aim to maintain the genetic diversity and the genetic integrity of a species. The phylogeographic history of a species can aid in defining areas of conservation priority. Here, we conducted a phylogeographic analysis of noble crayfish to identify their genetic hotspots and to reconstruct the colonisation history of this native European species. Our results support the hypothesis that noble crayfish outlasted the last glacial cycles in southeastern Europe in at least two refugia. The mtDNA analysis suggests a bifurcated colonization process from the eastern Black Sea basin to a) the North Sea and b) the Baltic Sea basin.

Glacial survival in southeastern Europe

The high diversity of mtDNA and nuclear DNA indicated that southeastern Europe is the hotspot of genetic diversity of noble crayfish. In contrast, very low genetic diversities of both markers were detected in central Europe (Table 1 and 5). It is known that high levels of population differentiation characterise older populations that have already diverged before the last ice age (Hewitt 1999). Therefore, we suggest that the high genetic differentiation in southeastern Europe evolved during several glacial cycles when this area served as a glacial refuge while climate conditions were unfavourable in central and northern Europe. In particular, we observed a high genetic differentiation between the Western Balkans and the eastern Black Sea basin (Romania and Bulgaria), suggesting that noble crayfish survived in at least two glacial refuges in southeastern Europe. This long separation between both refuges is supported by the results of the Bayesian phylogenetic tree where haplotypes from the Western Balkans form a distinct lineage (Lineage 4, Figure 3) that split already around 711,000 YBP from eastern Black Sea basin haplotypes. Because of the higer differentiation of haplotyopes from the Western Balkans, it might be possible that the Western Balkans was an older glacial refuge for noble crayfish. Eventually, the Western Balkans was less affected than the eastern Balkans by adverse climate conditions. The Western Balkans is the region with the highest genetic diversity for many species (e.g. Bănărescu 2004; Sket et al., 2004). High genetic diversity in the stone crayfish (Austropotamobius torrentium) was detected by COI sequence analysis in the Kolpa / Kupa drainage along the border between Slovenia and Croatia (Trontelj et al., 2005). The high genetic diversity in the Western Balkans and in the eastern Black Sea basin as well as the high differentiation among these areas revealed by both markers supports the hypothesis of at least two refugia in southeastern Europe during the LGM.

An additional Southwest-European refugium might explain the high differentiation of haplotypes from Rhineland-Palatinate (North Sea basin; Hap30 – Hap32) to the remaining North Sea samples. It is conceivable that besides the Danube River basin there existed other refuges in Europe during the LGM. Noble crayfish were presumably present in central Europe before the last ice age, and

when the climate cooled again they could have retreated besides to a southeastern European refugium also to a Western refugium. The LGM they could have outlasted in some periodically ice-free rivers in southwestern Europe as assumed for the fish species *Cottus gobio* (L, 1758) that might have survived the LGM in today's eastern France (Vonlanthen *et al.*, 2007). However, this theory can not be supported and the differentiated haplotypes might also be a result of artificial translocation.

The divergence times in Figure 3 show that the separation of the major lineages is not related to the cold climate phases. While genetic divergence of other species took place when subpopulations were geographically isolated in different refuges (Hetitt 1999), this correlation could not be detected for noble crayfish. However, the δ^{18} O curve contains data from globally distributed sites (Lisiecki and Raymo 2005) and does not consider regional variation. Further, the divergence rate of 1.4% per million years for COI has to be considered with caution because it is not specific for freshwater crayfish species. The lack of hierarchical structure in the phylogenetic tree can be explained by a low number of base exchanges in the mtDNA caused by a founder effect of noble crayfish in Central and Northern Europe. The low genetic variation might also explain the star-like pattern around three common haplotypes in the haplotype network.

Recolonisation of central Europe

In contrast to southeastern Europe, a homogeneous gene pool dominates in the North and Baltic Sea basin (Table 1). The low genetic diversities are characteristic for a more recently colonised area (Taberlet *et al.*, 1998). In this study we have revealed a significant negative value for both Tajima's D and Fu's F for the North Sea basin (Table 1) that indicate a recent population expansion. Extinctions of noble crayfish in northern and central Europe followed by founder effects probably led to the homogenization of noble crayfish in northern and central Europe. The endemic haplotypes from the Western Balkan populations are highly differentiated from the remaining haplotypes (Figure 2 and 3). This genetic separation is also supported by the FCA analysis of the nuclear data where the Western Balkans data points grouped separately from all other

samples (Figure 5). Possibly the Alps hindered the direct spread of the Western Balkan lineages into central Europe. Additionally, as supposed for other species (Hewitt 2000) the early colonisation from another refugia prevented the expansion of noble crayfish from the Western Balkans. We rather show that the eastern Black Sea basin 'pool' (e.g. Hap 01) contributed to the post-glacial recolonisation of noble crayfish from the eastern Black Sea basin into central and northern Europe as it was demonstrated for many other freshwater species (e.g. Hänfling et al., 2009; Durand et al., 1999; Weiss et al., 2002). In the phylogenetic tree and in the MJ network haplotypes from the eastern Black Sea group together with haplotypes from the North and Baltic Sea basins. This was also confirmed by the model selection calculated with MIGRATE-N. For the calculation the Western Balkans was separated as a distinct group. The favoured model points to the bifurcated colonization route from the eastern Black Sea basin to a) the North Sea basin and to b) the Baltic Sea basin (Figure 4). For the colonisation of Finland, the model with a colonisation route from the Baltic Sea basin further to Finland was favoured. However, the colonisation from the Black Sea basin to Finland (second highest probability value) is also possible, presumably via the Don and Volga Rivers. Nevertheless, the sampling size of Finnish samples was too small (N = 10) to finally reveal this question. The F_{ST}-values (Table 3) confirm the result of the model selection. The higher differentiation between North Sea and Baltic Sea indicates a longer separation than between Black Sea and the North Sea, respectively Baltic Sea basin. However, it has to be mentioned, that from southeastern Europe besides from the Western Balkans we have only data from Romania and Bulgaria. It is very well possible that the glacial refuge was even further south, e.g. in Greece or Macedonia, and Romania and Bulgaria were recolonised after the LGM from a more southern refuge before noble crayfish recolonised central and northern Europe. This is also supported by the heterogenous distribution of haplotypes in Romania indicating that the haplotypes did not evolve in the side channels of the Danube in Romania. An artificial translocation explaining this genetic structure can be excluded (Albrecht 1983).

From a potential western refugium noble crayfish could have recolonised Rhineland-Palatinate. A survival of noble crayfish in northern German rivers

during the LGM could further explain the private haplotypes in the Eider River catchment. For example European grayling is believed to have persisted the LGM in the northern Elbe and/or Vistula River catchment (Gum *et al.*, 2005). Therefore, it might be possible that also noble crayfish were not totally absent in central Europe during the last glacial period. This scenario would contradict earlier assumptions (e.g. Albrecht 1983; Schulz and Grandjean 2005). However, these samples could also be a result of stocking events. The origin of samples from France should be subject of further investigations.

Incongruent distribution of haplotypes

Along with the influences of climatic variation, historical river structure plays a significant role in explaining the genetic differentiation and the population structure of many freshwater species (Abellán et al., 2012). The finding of the same haplotypes in the Danube and Elbe (Hap25) or in the Danube and Rhine River catchment (Hap32) could be explained by channels that were modified by periglacial melt water and have changed water flow direction between and within drainages caused by geological events (Hantke 1993). Otherwise the geographical incongruent distribution could be explained by an artificial translocation across river catchments by men. Human influence can be assumed, for example, for the distribution of the very distinct haplotype Hap41 with 10 base pairs differing from the most frequent haplotype Hap01. Hap41 was found in Croatia and in the geographically very distant Elbe River catchment. The results from the microsatellite data also indicate a disturbed distribution of noble crayfish. The cluster analysis did not assign the populations from the same river catchments into one cluster as it is expected for species with a natural genetic structure. The hatchery in Augsburg formed one cluster with a population from the Rhine (Dauphte) and the Ems River catchments (Gut Riedberg). This might be a result of noble crayfish stocking from the hatchery in Augsburg to these waters. It is well known that translocations of noble crayfish for commercial purposes have been conducted for centuries (Skurdal and Taugbøl 2002) but were intensified in the last century in response to rapidly declining crayfish stocks (Souty-Grosset et al., 2003). Former translocations have been detected with genetic methods for other crayfish species, e.g. Austropotamobius pallipes (Grandjean et al., 2002) or Austropotamobiu italicus italicus (Pedraza-Lara et al., 2010). For noble crayfish, the artificial translocation might have increased the local diversity, but destroyed the natural genetic structure of this species, especially in central Europe where the artificial influence on the noble crayfish distribution was higher than in southeastern Europe (Albrecht 1983). The translocation further decreases the overall genetic diversity of noble crayfish because local and endemic haplotypes and alleles may be lost.

According to the results from the genetic differentiation analyses (Φ_{ST} -values, F_{ST} -values), nuclear DNA and mtDNA reveal contradicting results. Contrary to the results based on mtDNA, with nuclear DNA a relatively high differentiation between the Black Sea and the North Sea as well as between the Black Sea and the Baltic Sea was observed (Table 3). The incongruence is common in phylogenetic studies because the two markers have a different mode of inheritance (Zink and Barrowclough 2008). Possible explanations are: a) The high evolutionary rate in nuclear DNA can lead to a high differentiation between some populations. At the same time it can lead to homoplasy in different alleles and result in an underestimation of genetic diversity and genetic differentiation. In this case ancestral polymorphism might not be detected anymore (Theissinger *et al.*, 2011). b) Translocation can cause a discrepancy between the two markers, especially when a translocated population is mixed with a native population.

As can be seen by the high number of missing haplotypes in the MJ network, many haplotypes are either extinct or were not included in the sampling. In particular, many haplotypes that link the eastern and Western Balkans haplotypes are missing. A more detailed analysis of southeastern European populations should be conducted and the number of populations for a microsatellite analysis should be increased in order to reveal the local hotspots of diversity and to define eastern European ESUs.

Implications for conservation

Compared to other European freshwater crayfish species where a higher number of base pair exchanges was revealed in the COI sequence (e.g. *Astacus leptodactylus*; Maguire pers. com.; *Austropotamobius torrentium* and *A. pallipes*,

Trontelj et al., 2005), noble crayfish exhibit a small genetic diversity. A good conservation management of noble crayfish is still lacking and strongly needed in whole Europe to minimize the loss of genetic diversity. This is especially of crucial importance in the Danube River catchment that is subject of a recent invasion process of American crayfish species and the crayfish plague (Hudina et al., 2009; Pârvulescu et al., 2012; Schrimpf et al., 2012). Between-catchment translocations for economic reasons and conservation programs that consist of restocking crayfish without knowledge of their taxonomic status (Souty-Grosset et al., 2003) both have negative effects on the conservation of the genetic diversity and the genetic integrity of noble crayfish. It is claimed that in restocking projects for crayfish species the donor population should be selected based on its local genetic distinctiveness. If no money for genetic analysis is available, the results of this study (inter-basin diversity) support the idea of conservation of catchmentspecific gene pools (Weiss et al., 2002) and each river catchment should be managed as a distinct ESU. In a genetic analysis an indication for an unnatural origin is given by incongruent distributed haplotypes (e.g. Hap41 in the Elbe River catchment) or significant distinct alleles in a population. When there are several options for a local donor population the ones with a high genetic diversity should be selected (Taugbøl and Peay 2004). Even if the genetic structure of noble crayfish is already partly dissolved, following these suggestions the maximum diversity of the species and the integrity of local populations will be preserved.

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SUPPLEMENTARY MATERIAL

Table I Description of the sampling locations and haplotypes of analyzed *Astacus astacus*. Given is the number of genotyped (Ms N) and sequenced (Seq N) individuals per sampling site, number of the most common haplotype (Hap01) and the number of all other haplotypes (Hap). The sampling sites are descibed by the respective sea basin (SB: BAS = Baltic Sea, NS = North Sea, BS = Black Sea; AdS = Adriatic Sea, AeS = Aegean Sea), the longitude and latitude (N°, E°), the river catchment (RC, KO = Kokemäenjoki, DA = Danube) and country (GER = Germany, PO = Poland, F =, France CZ = Czech Republic, BE = Belgium, BU = Bulgaria, A = Austria, HU = Hungary, RO = Romania, HR = Croatia, MNE = Montenegro, KOS = Kosovo). The geographical coordinates of the study sites can be obtained from the corresponding author.

Population (site)	SB	Country	RC	Ms N	Seq N	Hap01	Нар
Koppelsee Barthe	BAS BAS	GER GER	Uecker		3 5	3	Hap16(5)
Tonkuhle	BAS	GER			3	3	,
Rakowe Duze	BAS	РО	Odra	15	5	5	
Seki	BAS	PO	Wieprza		5	5	
Bez Nazwu	BAS	PO	Wieprza	9	5	5	
Czarne	BAS	PO	Wisla	11	5	5	
Rosko	BAS	РО	Wisla		5	2	Hap15(2) Hap18(1)
Tomczyna	BAS	PO	Wisla		5	5	
Lake1 close to Miastko	BAS	PO			5	5	
Lake2 close to Miastko	BAS	PO			5	5	
Lake3 close to Miastko	BAS	PO			5	5	
Valkeinen	BAS	F	Kymen	14	5	3	Hap07(2)
Ylä	BAS	F	KO	15	5	5	
Gut Rietberg	NS	GER	Ems	15	5	4	Hap32(1)
Langsee	NS	GER	Eider	14	9		Hap03(8) Hap04(1)
Stepenitz	NS	GER	Elbe		5	5	
Schwarze Elster	NS	GER	Elbe		6	6	
Jäglitz	NS	GER	Elbe	9	7	5	Hap41(2)
Svetlohor	NS	CZ	Elbe	11	8	8	
U sudu	NS	CZ	Elbe	15	6	4	Hap25(2)
Kramata	NS	CZ	Elbe	8	7	7	
Florenville	NS	BE	Meuse	15	5	5	
Libramont	NS	BE	Meuse		4	2	Hap13(2)
Aar	NS	GER	Rhine		5	5	
Allna	NS	GER	Rhine		4	3	Hap21(1)
Ambach	NS	GER	Rhine	12	5	5	
Mühlgraben Caldern	NS	GER	Rhine		3	2	Hap06(1)
Dautphe	NS	GER	Rhine	20	4	4	
Dielbach (Woog)	NS	GER	Rhine		8	3	Hap05(1) Hap31(3) Hap41(1)
Donsbach	NS	GER	Rhine		5	5	
Eichelbach	NS	GER	Rhine		5	5	
Fischbach	NS	GER	Rhine		6	3	Hap30(3)

Fohnbach NS GER Rhine	Population (site)	SB	Country	RC	Ms N	Seq N	Hap01	Нар
Gansbach NS GER Rhine 10 10 Geierstein/Roth NS GER Rhine 10 10 Waldteich, Irrschelde NS GER Rhine 5 5 Kallenbach NS GER Rhine 5 5 Lasterbach NS GER Rhine 5 5 Merzkrebse NS GER Rhine 1 1 Meerbach NS GER Rhine 5 5 Mademühlen NS GER Rhine 4 4 Mademühlen 2 NS GER Rhine 7 3 Hap32(4) Maldemühlen 2 NS GER Rhine 7 3 Hap32(4) Malzenbach NS GER Rhine 5 5 Mallscheid NS GER Rhine 5 5 Perf NS GER Rhine 1 Hap32(2) Pollichia Wo				_				_
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Wielenbach BS GER DA 2 2	Freundsheimer Weiher	BS	Α	DA				
Kádárta BS HU DA 4 3 3	Wielenbach	BS		DA		2	2	. , ,
	Kádárta	BS	HU	DA	4	3	3	

Population (site)	SB	Country	RC	Ms N	Seq N	Hap01	Нар
Pét	BS	HU	DA		1		Hap09(1)
Schiopu	BS	RO	Mures/DA		1	1	
Tebea	BS	RO	Cris/DA		1		Hap40(1)
Sohodol	BS	RO	Cris/DA		1		Hap40(1)
Racas	BS	RO	Cris/DA		1		Hap40(1)
Valea Boului	BS	RO	Cris/DA		1	1	
Barcau	BS	RO	Cris/DA		2	1	Hap40(1)
Valea Pestilor	BS	RO	Cris/DA		1	4	Hap39(1)
Valea Stoiaca	BS	RO	Cris/DA	40	2	1	Hap40(1)
Nadas	BS	RO	Cris/DA Cris/DA	13	12 1		Ham 40/4)
Derjana Clocotici	BS BS	RO RO	Cris/DA Caras/DA	8	1 8		Hap40(1) Hap36(1)
				0			Hap37(7)
Natra	BS	RO	Caras/DA		1		Hap23(1)
Comarnic	BS	RO	Caras/DA		1		Hap23(1)
Toplita	BS	RO	Caras/DA		1		Hap23(1)
Ravistea	BS	RO	Caras/DA		8		Hap23(8)
Buhui	BS	RO	Caras/DA		2	2	1140(4)
Moravita	BS	RO	Caras/DA		5	1	Hap19(1) Hap23(3)
Dognecea	BS	RO	Caras/DA		2	1	Hap23(1)
Ciornovãt	BS	RO	Caras/DA	6	7		Hap22(6) Hap23(1)
Caras	BS	RO	Caras/DA		1		Hap23(1)
Caianu	BS	RO	Caras/DA		1	1	
Bacaia	BS	RO	Caras/DA		2	2	
Balsa	BS	RO	Caras/DA		2	1	Hap40(1)
Galben	BS	RO	Jiu/DA		1		Hap40(1)
Dragoiestilor	BS	RO	Jiu/DA	22	5		Hap40(5)
Tamasesti	BS	RO	Mures/DA		4	4	
Balsa	BS	RO	Mures/DA		4	3	Hap40(1)
Cladovita	BS	RO	Mures/DA		4	2	Hap10(2)
Carpan	BS	RO	Mures/DA		1		Hap34(1)
Geoagiu	BS	RO	Mures/DA		1	1	
Conop	BS	RO	Mures/DA		3	1	Hap40(2)
Valea Crisului	BS	RO	Mures/DA		4	4	
Crisul	BS	RO	Mures/DA		3	3	
Giacas	BS	RO	Mures/DA		2	2	
Solocma	BS	RO	Mures/DA		3	3 1	
Niraj Taraia	BS BS	RO RO	Mures/DA Olt/DA		1	1	Uan22(1)
lazul	BS BS	RO	Olt/DA		1 1		Hap23(1)
Sinca	BS	RO	Olt/DA		1		Hap23(1)
Valea Adanca	BS	RO	Olt/DA		2	1	Hap23(1) Hap20(1)
Hartibaciu	BS	RO	Olt/DA		3	•	Hap23(1)
Venetia	BS	RO	Olt/DA		1		Hap23(1)
Simbrezi	BS	RO	Olt/DA		2		Hap23(2)

Population (site)	SB	Country	RC	Ms N	Seq N	Hap01	Нар
Maierus	BS	RO	Olt/DA		1		Hap24(1)
Bogata	BS	RO	Olt/DA		1		Hap24(1)
Rupea	BS	RO	Olt/DA		4		Hap23(4)
Oituz	BS	RO	Siret/DA		1		Hap23(1)
Stramba	BS	RO	Siret/DA		1		Hap40(1)
Bichigiu	BS	RO	Somes/DA		1	1	
Agrij	BS	RO	Somes/DA		1	1	
Tetisu	BS	RO	Somes/DA		1	1	
Poiana	BS	RO	Somes/DA		1	1	
Sarasau	BS	RO	Tisa/DA		1	1	
Valea Tejei	BS	RO	Tisa/DA		1	1	
Valea Holita	BS	RO	Tisa/DA		1	1	
Valea Mare	BS	RO	Tisa/DA		1	1	
Vukovina Lake	BS	HR	Sava/DA	6	5		Hap41(5)
Bačica Creek	BS	HR	Sava/DA		4		Hap42(3)
D.Y.' D' OY	DO	LID	O /D A		0		Hap44(1)
Bašnica River, Gračac	BS	HR	Sava/DA		2		Hap43(2)
Borovik Lake	BS	HR	Sava/DA		1		Hap43(1)
Dubočanka River	BS	HR	Sava/DA		1		Hap43(1)
Jaruga River, Stajničko polje	BS	HR	Sava/DA		2		Hap28(2)
Ježevo Lake, Velika Gorica	BS	HR	Sava/DA		1		Hap42(1)
Krapina River	BS	HR	Sava/DA		1		Hap43(1)
Mrežnica River, Generalski Stol	BS	HR	Sava/DA		2		Hap 27(2)
Plitvice Lakes - NP Plitvička jezera	BS	HR	Sava/DA		2		Hap28(2)
Pakra River, Kusonje	BS	HR	Sava/DA		1		Hap42(1)
Ribnjak Creek, Vladisovo	BS	HR	Sava/DA		3		Hap42(2) Hap43(1)
Trećak Creek Staro Petrovo Selo	BS	HR	Sava/DA		2		Hap42(2)
Subocka River	BS	HR	Sava/DA		1		Hap42(1)
Šumetlica Creek	BS	HR	Sava/DA		3		Hap42(3)
Rakov Creek	AdS	HR	Pazinčica		1		Hap46(1)
Zeta River	AdS	MNE	Morača River		4		Hap28(1) Hap29(1)
			-				Hap42(2)
Liverovići Lake	AdS	MNE	?		1		Hap42(1)
Paklenica River - NP Paklenica	AdS	HR	Paklenica River		2		Hap42(2)
Ričica River	AdS	HR	Ričica River		1		Hap45(1)
Unnamed stream Ferizaj (Uroševac)	AeS	KOS	?		1		Hap26(1)
Zoo Zajak	NS	GER	Hatchery/RH		5	5	
Farm in Belgium	NS	BE	Hatchery/ME		2	1	Hap35(1)
Farm Oeversee	NS	GER	Hatchery/EI		5		Hap03(4) Hap04(1)

Population (site)	SB	Country	RC	Ms N	Seq N	Hap01	Нар
Farm Augsburg	BS	GER	Hatchery/DA	24	15	8	Hap06(3) Hap07(1) Hap08(1) Hap32(2)

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Crayfish plague agent detected in populations of the invasive North American crayfish Orconectes immunis (Hagen, 1870) in the Rhine River, Germany

Anne Schrimpf¹, Christoph Chucholl², Thomas Schmidt¹, Ralf Schulz¹

ABSTRACT

Crayfish plague, caused by the parasitic oomycete Aphanomyces astaci, has driven indigenous European crayfish species to regional extinction in many parts of Europe and is among the leading threats to the remaining populations. A. astaci is known to be carried by long-established invasive crayfish species of North American origin, which are also the main vectors of the plague pathogen. In this study, we examined whether a new invasive crayfish of North American origin, the calico crayfish (Orconectes immunis), also carries A. astaci. Orconectes immunis is a recent invader of the Upper Rhine plain, where it seems to displace its invasive predecessor Orconectes limosus, which is a known carrier of the agent of the crayfish plague. Using real-time PCR, we identified the calico crayfish as the fourth invasive crayfish species to be a carrier of the crayfish plague pathogen in Europe and we confirmed the infection with A. astaci in O. limosus. These findings support the concern that all North American crayfish species in European waters are carriers of the crayfish plague pathogen. Such knowledge should prove useful for conservation efforts, management, legislation, and public education about the spread of crayfish plague and non-indigenous crayfish species.

Keywords: Crayfish plague • *Aphanomyces astaci* • calico crayfish • crustacean disease • Rhine River • invasive American crayfish species • real-time PCR • pathogen vector

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INTRODUCTION

Invasive non-indigenous species are one of the leading threats to freshwater biodiversity, besides habitat deterioration (Sala et al. 2000; Gherardi 2007). All European freshwater crayfish species (Crustacea, Decapoda, Astacidae) are highly threatened by *Aphanomyces astaci* (Schikora 1906), an invasive, crayfish-specific parasite causing crayfish plague, i.e., a fatal disease (Söderhäll and Cerenius 1999). *Aphanomyces astaci* originates from North America and was probably first introduced to Europe in the late 1850s. Its natural hosts, North-American freshwater crayfish species, were not found during the first outbreaks of the disease in Europe but were repeatedly introduced later (Alderman 1996; Holdich et al. 2009). Out of a total of more than 460 crayfish species living in North America (Crandall and Buhay 2011), today at least eight species are established in Europe in the wild (Holdich et al. 2009; Chucholl and Pfeiffer 2010).

Although all North American crayfish species are suspected to be carriers of A. astaci (OIE 2009), only three of the North American crayfish species present in the wild in Europe have been shown to be carrier of the pathogen so far: the signal crayfish [Pacifastacus leniusculus (Dana, 1852)] (Unestam and Weiss 1970), the spiny-cheek crayfish [Orconectes limosus (Rafinesque, 1817)] (Vey et al. 1983), and the red swamp crayfish [Procambarus clarkii (Girard, 1852)] (Diéguez-Uribeondo and Söderhäll 1993). These three species belong to the 'Old' non-indigenous crayfish species in Europe, i.e. have been introduced into European waters before 1975 (summarized by Holdich et al. 2009).

One 'New' non-indigenous crayfish species in Europe is the North American calico crayfish [*Orconectes immunis* (Hagen, 1870)], which was first recorded at two locations in the Upper Rhine plain in the mid-1990s (Dussling and Hoffmann 1998; Dehus et al. 1999; Gelmar et al. 2006). The pathway of introduction of *O. immunis* is unclear; both an introduction from the pet trade and as fishing bait by Canadian soldiers had been suggested (Dehus et al. 1999; Gelmar et al. 2006). However, since the calico crayfish was not known in the German pet trade prior to its establishment in the Upper Rhine plain and because this species is popular as fishing bait in North America, an introduction as fishing bait seems more likely

(see Gelmar et al. 2006; Chucholl 2012b). Since its discovery, the calico crayfish has rapidly spread upstream and downstream in the Upper Rhine plain (Gelmar et al. 2006; Chucholl 2012a) and is currently colonizing a stretch of more than 98 km, where it inhabits a wide spectrum of habitat types (summarized in Chucholl 2012a; Figure 1). Despite its presence in Europe for almost two decades, its status as carrier of *A. astaci* is still unclear.

Calico crayfish seem to replace the formerly most abundant non-indigenous crayfish species of the Rhine River, the spiny-cheek crayfish (Gelmar et al. 2006; Chucholl et al. 2008), which has been present in the Upper Rhine catchment for at least five decades (Holdich et al. 2009). In laboratory experiments, calico crayfish were shown to be superior to spiny-cheek crayfish in direct aggressive interactions and competition for shelter (Chucholl et al. 2008). Furthermore, the calico crayfish exhibits a strongly *r*-selected life history: it is rather small (at most 50 mm in carapace length), features a high fecundity (up to 500 pleopodal eggs female⁻¹), and has the fastest recorded life cycle among the crayfish species present in Central Europe, combining a high growth rate and rapid maturation (within the first summer) with short longevity (2.5 years) (Chucholl 2012a).

Knowledge about the *A. astaci* carrier status of alien crayfish populations is imperative for native crayfish conservation, risk assessment and management strategies (Peay 2009). The aim of the present study was to evaluate the status of calico crayfish populations in the Rhine catchment as carrier of *A. astaci*.

METHODS

To assess the *A. astaci* carrier status of calico crayfish in the Upper Rhine plain, we sampled two calico crayfish populations using traps and hand-held nets. One of the sampled populations was close to the currently known downstream invasion front of *O. immunis* (Germersheim, about 26 km north of Karlsruhe) and occurred in sympatry with *O. limosus*, whereas the second sampled population was located near the site of original introduction (Bühl, about 38 km south of Karlsruhe). The sampling site near the invasion front was located in the Rhine

River close to Germersheim (approx. Rhine km 390, 49°15'N, 8°25'E, see Figure 1) and the second sampling site was located with an aerial distance of about 63 km to the first sampling site at a small channel that is connected to the Rhine River near Bühl (8°04'N, 48°43'E). This site is in close proximity to the *O. immunis* occurrence reported by Dussling and Hoffmann (1998), i.e. one of the two locations where calico crayfish were first discovered in the mid-1990s. Calico crayfish is the exclusive crayfish species in this channel and previous to its establishment no crayfish had been known to occur there. In total, we collected 50 individuals of calico crayfish and 10 individuals of spiny-cheek crayfish from the Germersheim site in 2011 and 32 calico crayfish individuals from the Bühl site in 2012.

Upon capture, crayfish were identified using distinct features (Figure 2), and whole specimens were transported to the laboratory and frozen at -20°C. To evaluate the A. astaci carrier status we used the TagMan® minor groove binder (MGB) real-time PCR (gPCR) according to Vrålstad et al. (2009). This method is the most specific and sensitive method to test for the presence of the crayfish plague pathogen (Tuffs and Oidmann 2011). Before DNA-extraction, we visually checked all calico crayfish for mechanical damage and melanisation. DNA was extracted from the soft abdominal cuticle, the inner joint of two walking legs, a part of the uropods and melanised spots when present using a CTAB-method as described in Vrålstad et al. (2009). The qPCR reaction was performed on a Mastercycler[®] ep realplex S (Eppendorf) using the TaqMan[®] Environmental Master Mix to avoid PCR inhibition (Strand et al. 2011). Differing from the published PCR program, the annealing temperature was increased to 62°C and the annealing time decreased to 15 seconds to further exclude possible false positive results (T. Vrålstad, personal communication). A negative control consisting of 5 µl nuclease free water was included together with a standard series of genomic DNA from a pure A. astaci culture. Data analysis was carried out using the software Real Plex 2.2 (Eppendorf).

The relative level of infection by the pathogen was based on the strength of the qPCR signal. The number of observed PCR-forming units in the reaction was assigned to semi-quantitative agent levels (according to Vrålstad et al. 2009). Individuals with agent levels A_0 (no detection) and A_1 are considered uninfected, individuals with agent levels A_2 and higher are considered infected by *A. astaci* (see Table 1 caption for details).

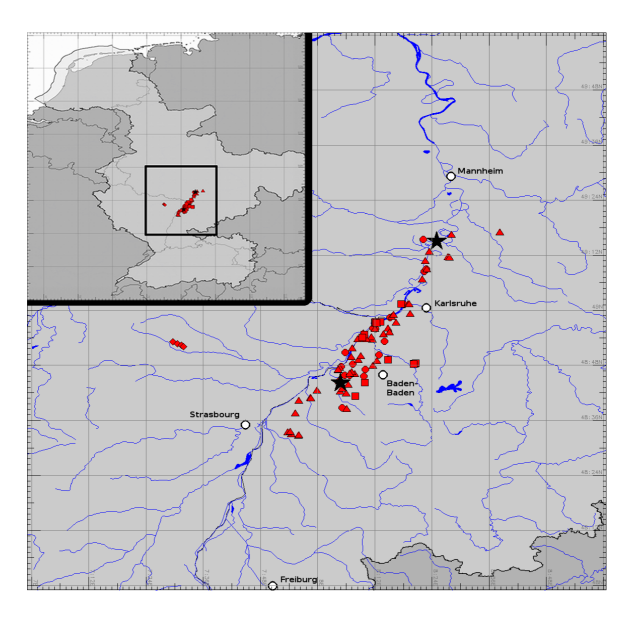


Figure 1 The known distribution of *Orconectes immunis* in Europe as summarized in Chucholl (2012) with data from Chucholl & Dehus (2011; triangles), Collas et al. (2011; diamonds) and Gelmar et al. (2006; circles), completed with unpublished data from Chucholl (squares). Black stars indicate the sampling sites of this study. The international River Basin District Rhine (data: European Environment Agency, 2011) is colored light grey. Data of waters and national borders: GADM (2012).

RESULTS

Using *A. astaci*-specific real-time PCR from a total of 92 crayfish a positive carrier status was found for about 60% of the tested crayfish (Table 1). At the sampling site Germersheim we found 23 positive detections out of 50 calico crayfish (46%) and six positive detections out of ten spiny-cheek crayfish (60%). Furthermore, *A. astaci* was detected in 26 of the 32 tested calico crayfish from Bühl (81%). Most individuals from both species and both sampling sites contained very low to moderate levels of agent DNA (A_1 to A_4). However, two calico crayfish from the Bühl site were infected at a very high level (A_6) and one spiny-cheek crayfish with mechanical damage from the Germersheim site was exceptionally high infected (A_7). Melanised areas were observed in eight calico crayfish from the Bühl site, in four calico crayfish from the Germersheim site, all of which were tested positive, and in one highly infected (A_7) spiny-cheek crayfish specimen.

Table 1 Real-time PCR detection of *Aphanomyces astaci* in investigated specimens of *Orconectes immunis* and *Orconectes limosus* from the two sampling sites from the Upper Rhine plain in Germany and percentage of specimens detected positive. For each sampling site, number of analyzed crayfish, number and proportion of infected individuals, and relative level of infection according to Vrålstad et al. (2009) are given.

Sampling site	Species	Crayfish (number)		Positives (%)	Agent level ^a							
					$\overline{A_0}$	A ₁	A ₂	A ₃	A ₄	A ₅	A ₆	A ₇
Germersheim	O. limosus	10	6	60	3	1		3	2			1
Germersheim	O. immunis	50	23	46	21	6	13	9	1			
Bühl	O. immunis	32	26	81	3	3	4	18	2		2	
	Total	92	55	60	27	10	17	30	5		2	1

^a Agent levels refer to semi-quantitative categories based on the numbers of observed PFUs (PFU_{obs}) from the *A. astaci*–specific real-time PCR. Agent level A₀: no detection; A₁: below the limit of detection (PFU_{obs} < 5); A₂: $5 \le PFU_{obs} < 50$; A₃: $50 \le PFU_{obs} < 10^3$; A₄: $10^3 \le PFU_{obs} < 10^4$; A₅: $10^4 \le PFU_{obs} < 10^5$; A₆: $10^5 \le PFU_{obs} < 10^6$; A₇: $10^6 \le PFU_{obs}$. A₀ and A₁ are both considered negative.

DISCUSSION

Using the currently most reliable molecular detection method for the agent of crayfish plague (Vrålstad et al. 2009; Tuffs and Oidtmann 2011), we have shown for the first time an A. astaci infection in calico crayfish. We found infections in this species at two sampling sites in the Upper Rhine plain with an aerial distance of about 63 km. Moreover, we confirmed the infection in spiny-cheek crayfish coexisting with calico crayfish. The relatively low agent levels (≤A₄) of most infected crayfish are typical for North American crayfish species, which show an evolved defense reaction against A. astaci that normally prevents further spread of A. astaci hyphae within their body (Cerenius et al. 2003) and results in a latent infection. Thus, North American crayfish do usually not suffer from the disease but continuously release spores into the water with elevated spore levels prior to and during molting and mortalities (Strand et al 2012). In the present study we found one spiny-cheek crayfish with a physical injury. This injury may have stressed it's immune system and thus explains the high infection status (A7) of this individual. The immune response can be visually observed in infected crayfish as melanised spots. However, melanised spots alone are not a good indicator for the infection status with A. astaci because they also appear as a reaction to physical injury. In the present study, melanisation was found in only 13 of the 29 positive tested crayfish. Therefore, the absence of melanised spots does not conclusively indicate the absence of an infection with A. astaci.

The positive verification of calico crayfish as carrier of the crayfish plague agent is worrying. Particularly, given the fast and successful spread of this 'New' non-indigenous crayfish species in the Upper Rhine plain. Moreover, the species replaces the 'Old' non-indigenous crayfish, spiny-cheek crayfish, from preferred habitats (Gelmar et al. 2006; Chucholl et al. 2008; Chucholl 2012a). Preliminary field observations suggest that calico crayfish inhabit a wider spectrum of habitats than spiny-cheek crayfish (Chucholl 2012a). Specifically, calico crayfish were found in shallow temporary backwaters adjacent to the Rhine River and brooks draining from the Schwarzwald, which are habitats from which spiny-cheek crayfish are typically absent. Calico crayfish might therefore spread *A. astaci* into habitats that were previously not colonized by spiny-cheek crayfish. However, to

date, calico crayfish have not come into contact with indigenous European crayfish species (Chucholl 2012a).

We assume that the calico crayfish will continue its fast invasion of the Rhine River and connected waterways. Artificial channels that connect the Rhine River to other large river catchment areas, such as the Danube, the Rhône, the Odra, and the Elbe, promote a fast spread of invasive aquatic species throughout Europe (Bij de Vaate et al. 2002) and will most likely also facilitate the further active spread of calico crayfish. In addition to active range expansion, secondary introductions, i.e. translocations of calico crayfish by humans, are also a potential mechanism of spread and have already occurred in Germany and possibly in France (Chucholl and Dehus 2011; Collas et al. 2011). Public and stakeholder information is therefore imperative to mitigate the risk of further secondary introductions, which are a major threat to otherwise isolated populations of indigenous European crayfish.

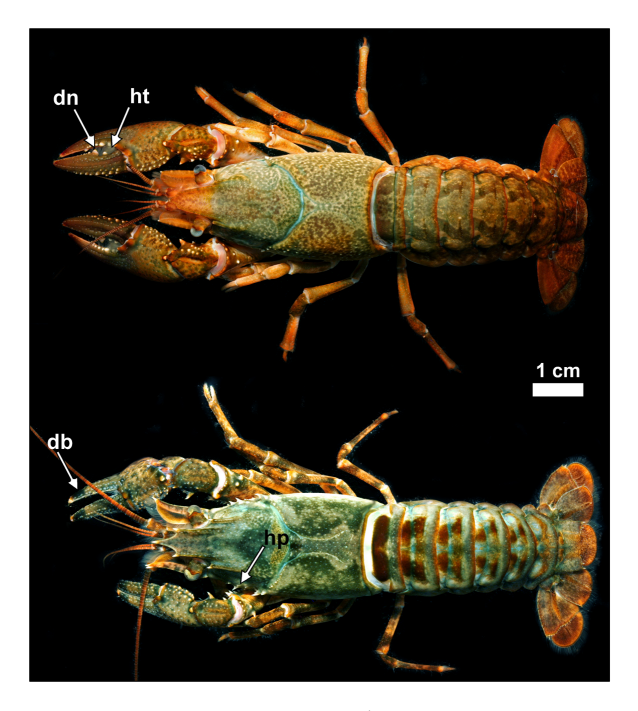


Figure 2 Orconectes immunis $\ \$ (top) and O. limosus $\ \ \$ (bottom) from the Rhine River. Arrows denote key characters to distinguish the two species (modified from Gelmar et al. 2006 and Chucholl et al. 2008): dn – distinct tooth followed by a notch on the dactylus of the chelipeds (only present in O. immunis); ht – hair tufts on the ventral side of the chelae joints of the 1st and 2nd pereiopod (only present in O. immunis); db – distinct dark bandage adjacent to the orange cheliped tips (only present in O. limosus); hp – hepatic spines (only present in O. limosus).

Finally, it is important to note that there is accumulating evidence that different North American crayfish species are carriers of different strains of A. astaci (Huang et al. 1994, Kozubíková et al. 2011) and that these strains vary in their virulence (Jussila et al. 2011; Viljamaa-Dirks et al. 2011). An important question is therefore whether calico crayfish carry a different and possibly new A. astaci strain. This question is closely linked to the question of whether a) calico crayfish were already carriers of A. astaci when they were introduced to the Rhine catchment or whether b) calico crayfish were initially uninfected and got infected later, when they came into contact with A. astaci-carrying spiny-cheek crayfish. The positive verification of *A. astaci* in the single species population close to Bühl that has existed for about 14 years is a strong indication that calico crayfish were already carrier of the agent of crayfish plague when they were introduced to the Rhine catchment. In this case, calico crayfish may carry a new and possibly more virulent strain. However, we cannot rule out the possibility of recent human influence (e.g. by fishing gear that was contaminated with A. astaci spores) or an infection via animals (e.g. predatory fish feeding on infected crayfish; Oidtmann et al. 2002). Furthermore, A. astaci originating from spiny-cheek crayfish in the Rhine River could have reached the population by gradually infecting calico crayfish that are presumably distributed in the whole channel that connects the Rhine River with the sampling site in a stepping-stone manner. In the case of the Germersheim population A. astaci can be transmitted from calico crayfish to spiny-cheek crayfish and vice versa. We hope to resolve the question of the origin of A. astaci in European calico crayfish populations in the future, when methods become available that facilitate the assignment of A. astaci strains and the detection of new strains.

The positive verification of calico crayfish as carrier of *A. astaci* adds another species to the list of highly dangerous non-indigenous species. In particular, it adds another species to the list of *A. astaci*-carrying crayfish species. The result of this study has to be implemented in native crayfish conservation strategies. A site where calico crayfish is present has to be considered as a reservoir for *A. astaci*.

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Invasive crayfish and crayfish plague on the move: first detection of the plague agent Aphanomyces astaci in the Romanian Danube

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ABSTRACT

Native European crayfish, such as Astacus leptodactylus, are threatened, among other factors, by the crayfish plague agent Aphanomyces astaci, dispersed by invasive North American crayfish. Two of these invaders, Pacifastacus leniusculus and Orconectes limosus, have extended their distribution in the River Danube catchment; the latter was detected for the first time in Romania in 2008. We monitored, at monthly intervals for over 2 yr, occurrence of native A. leptodactylus and invasive O. limosus at 6 sites on the Romanian Danube and checked for the invasive species in 4 of its tributaries. Between January 2009 and March 2011, the relative abundances of O. limosus steadily increased with time, while the native A. leptodactylus dramatically decreased in abundance. O. limosus expanded downstream at a rate of ca. 15 km yr⁻¹; in August 2011, it was already present in the upper 105 km of the Romanian Danube. An agent-specific real-time PCR analyses demonstrated the presence of A. astaci DNA in at least 32% of the analysed invasive (n = 71) and 41% of the native (n = 49) crayfish coexisting in the Danube. Furthermore, *A. astaci* was also detected in A. leptodactylus captured about 70 km downstream of the O. limosus invasion front (at the time of sampling). Assuming a steady rate of expansion, O. limosus may invade the sensitive Danube delta area in the mid-2060s, even without long-distance dispersal. The crayfish plague agent, however, may reach

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the delta substantially earlier, through dispersal downstream among populations of native crayfish.

Key words: Crayfish plague • *Aphanomyces astaci* • Quantitative real-time PCR • Molecular diagnostics • Danube • Non-indigenous crayfish • *Orconectes limosus* • Indigenous crayfish • *Astacus leptodactylus*

INTRODUCTION

Crayfish plague is a crayfish-specific disease caused by the oomycete *Aphanomyces astaci*, an obligate parasite of freshwater crayfish (Söderhäll & Cerenius 1999). While all European crayfish species are highly susceptible, infection does not usually cause disease outbreaks or death in North American crayfish species (OIE 2009) unless they are stressed (Persson & Söderhäll 1983, Cerenius et al. 1988) or exposed to extremely high concentrations of short-lived *A. astaci* zoospores (Diéguez-Uribeondo & Söderhäll 1993). It is assumed that *A. astaci* does not usually survive for more than a few weeks without a crayfish host (Söderhäll & Cerenius 1999, CEFAS 2000), and no durable stages (oospores) are known in the parasite's life cycle.

All 3 of the North American crayfish species most widely distributed in Europe - the spiny-cheek crayfish *Orconectes limosus* (introduced in 1890), the signal crayfish *Pacifastacus leniusculus*, and the red swamp crayfish *Procambarus clarkii* (both introduced in the 1950–60s) - act as carriers of the crayfish plague pathogen (e.g. Vey et al. 1983). They are, therefore, regional reservoirs of the disease in invaded areas and contribute to its further dispersion into areas dominated by native European crayfish species. However, the disease itself had already started to spread across the continent by the 1860s and 1870s (Alderman 1996), before the first docu- mented introductions of American crayfish species. There were repeated disease outbreaks in the late nineteenth and early twentieth centuries across a substantial part of the continent, leading to a sharp decline in native crayfish species. Regions affected by the disease were most of central and eastern Europe (including western Russia, Belarus, Ukraine, and the lower Danube basin), southern Fennoscandia, and parts of Western Europe, particularly Germany and the eastern half of France (Alderman 1996). Most

European regions not impacted by this particular major wave of crayfish plague outbreaks became affected later in the twentieth century, after intensive introductions of *P. leniusculus* and *P. clarkii* (e.g. Alderman et al. 1990, Taugbøl et al. 1993, Diéguez-Uribeondo 2006). Due to its virulence and devastating impact on indigenous European crayfish species, *Aphanomyces astaci* has been classified among the world's 100 worst invasive alien species (Lowe et al. 2004).

The only non-indigenous crayfish species in Romania, *Orconectes limosus*, was detected in the Romanian Danube for the first time in 2008 (Pârvulescu et al. 2009), having reached the area by downstream dispersal through the Danube. The species itself has been present in Europe since 1890, apparently due to a single successful introduction (Filipová et al. 2011). However, it did not colonise the Danube until 1985, when a wild population became established in Hungary (Puky & Schád 2006), from where it dispersed both upstream (e.g. Puky 2009) and downstream. Its downstream colonisation rate over 20 yr has been estimated at 13 to 16 km yr⁻¹ (Puky & Schád 2006). *O. limosus* has also entered Croatia through the Danube from Hungary; it has entered the River Drava and is spreading upstream at a rate of ~1.5 km yr⁻¹, affecting native populations of *Astacus leptodactylus* (Maguire & Klobuçar 2003, Faller et al. 2009).

The presence of *Orconectes limosus* in Romania may dramatically affect populations of the indigenous narrow-clawed crayfish *Astacus leptodactylus*, which occurs both in the Danube itself and in other watercourses of the catchment (Băcescu 1967). Furthermore, the indigenous stone crayfish *Austropotamobius torrentium* is common in most tributaries of the Romanian Danube (Pârvulescu & Petrescu 2010). These areas are also of crucial importance from a conservation perspective. Besides carrying *Aphanomyces astaci*, other characteristics of *O. limosus*, such as high fecundity and early maturation time, contribute to its negative pressure on native crayfish populations (Hamr 2002, Kozák et al. 2006, 2007, Schulz et al. 2006). Furthermore, a recently demonstrated potential for facultative parthenogenesis in this species (Buřič et al. 2011) may also contribute to its invasive potential.

The first outbreak of the crayfish plague in the upper Danube basin was registered in Bavaria, Germany, in early 1879; following this the outbreaks quickly

(within a few months) spread to upper Austria and further downstream (Alderman 1996). In less than 1 decade, the disease had spread as far as the Black Sea. Recently, crayfish plague outbreaks in the Danube catchment were reported from the Czech Republic (Kozubíková & Petrusek 2009) and Austria (Hochwimmer et al. 2009), and populations of *Orconectes limosus* in the Hungarian Danube have been confirmed to carry the crayfish plague pathogen (Kozubíková et al. 2010). Although Romania must have been substantially affected by outbreaks of this disease in the late nineteenth century (Alderman 1996), there are only a very few later reports of crayfish mass mortalities (for example, of native noble crayfish Astacus astacus in 1934-1935 in the Olt River, a tributary of the Danube; Bačescu 1967). However, their association with crayfish plague has never been definitively proven. During the last several years, molecular diagnostic tools employing conventional PCR and real-time PCR methods for the specific detection of Aphanomyces astaci have been developed, and these have accelerated and improved the detection and identification of the crayfish plague agent (Oidtmann et al. 2006, Hochwimmer et al. 2009, Vrålstad et al. 2009). The carrier status of symptom-free North American crayfish is also evaluated using these tools (Oidtmann et al. 2006, Kozubíková et al. 2009, 2011b, Vrålstad et al. 2009, 2011, Skov et al. 2011).

Reliable information on the presence of the crayfish plague agent in the Romanian Danube and on the level of threat posed to native crayfish by the quickly spreading *Orconectes limosus* is of key importance for freshwater management and species conservation actions. In this study, we monitored the distribution dynamics of *O. limosus* and *Astacus leptodactylus* in the upper Romanian Danube and its tributaries and analysed their carrier status using the most sensitive and reliable molecular diagnostic assays available. We employed the TaqMan[®] minor groove binder (MGB) real-time PCR (Vrålstad et al. 2009). Recent results show that this method offers higher sensitivity than any other presently available, and it is the preferred method for screening populations of North American crayfish for carrier status (Tuffs & Oidtmann 2011). Furthermore, this method leads to lower error rates in the detection of *Aphanomyces astaci* than alternatives based on conventional PCR (Kozubíková et al. 2011b).

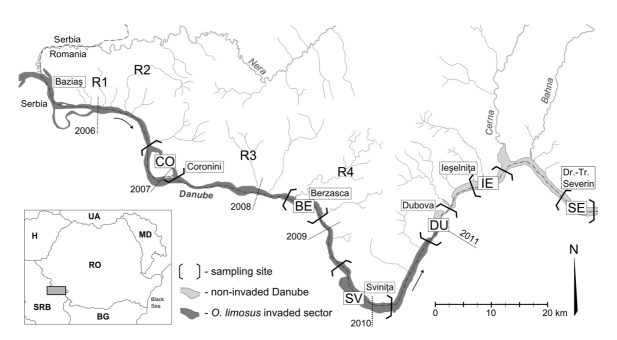


Figure 1 Sampling sites in the River Danube (CO, BE, SV, DU, IE, SE) and its tributaries (R1 to R4; for river names see 'Materials and methods') in Romania and Serbia. Estimated years indicating the rate of invasion of this species are provided; *Orconetes limosus* was recorded at the DU site in August 2011. Dr.-Tr. Severin: Droberta-Turnu Severin. International license plate codes used for countries in inset map

MATERIALS AND METHODS

Crayfish monitoring

Six sites along the eastern section of the Romanian Danube, each covering 5 km of the river's shore stretch, were surveyed monthly, using the same capture effort, from January 2009 to March 2011 in order to estimate relative crayfish abundances. One surveyed site was the location of a well-known *Orconectes limosus* invasion, near the village of Coronini (CO in Fig. 1); and another site further downstream near the village of Berzasca (BE) had more recent evidence of invasion (Pârvulescu et al. 2009). In order to estimate the actual invasion front at Sviniţa, a (SV) properly, the investigated stretch of the river was twice as long (10 km) as for the other sites. Three sites downstream of the suspected invasion area were investigated: Dubova (DU), leşelniţa (IE) and Drobeta-Turnu Severin (SE). The downstream distance to each subsequent sampling site was approximately 15 km (Fig. 1).

Crayfish were captured using 2 bottom fishing nets of 1.5 m height, 25 m width, and a mesh size of ca. 30 mm, baited with fish meat and checked twice a week for a total period of 2 wk in each month. After capture, the species and sex of each individual was determined. Four Danube tributaries, Cameniţa River (R1 in Fig. 1), Radimna River (R2), Gornea River (R3) and Berzasca River (R4), were investigated using hand sampling of the riverbed during the summer of 2010 to assess whether *Orconectes limosus* shows a tendency to move upstream and colonise smaller watercourses. These tributaries are known to be populated by the stone crayfish *Austropotamobius torrentium* in areas between 1 to 10 km upstream of the confluence with the Danube (Pârvulescu & Petrescu 2010).

Molecular diagnostics

Crayfish were sacrificed at random to determine the presence of Aphanomyces astaci in both species by freezing them in separate plastic bags at -10°C. Altogether, we analysed 71 individuals of *Orconectes limosus* and 49 of Astacus leptodactylus. The crayfish were divided and analysed separately in 3 laboratories (in Oslo, Norway; Landau, Germany; and Prague, Czech Republic). For 42 crayfish (41 O. limosus and 1 A. leptodactylus) analysed in Prague, 50 mg of crayfish tissue, including the soft ventral abdominal cuticle and a part of the tail fan (uropods and/or telson), was dissected and processed according to the methodology described in Kozubíková et al. (2009). For the remaining 78 crayfish analysed in the laboratories at Oslo and Landau (30 O. limosus and 48 A. leptodactylus), 4 different tissue segments were dissected following methodology used by Vrålstad et al. (2011) and were preserved in 96% ethanol. The tissues included (1) the soft ventral abdominal cuticle; (2) the tail fan; (3) the inner joints of 2 walking legs; and (4) black spots (if present) on the exoskeleton. The melanised spots can be caused by an immune re- action against pathogens (including A. astaci) and may therefore indicate infection (Cerenius et al. 2003). The dissection tools were thoroughly disinfected after handling each individual crayfish.

DNA from the dissected tissues was extracted according to either Vrålstad et al. (2009; for the analyses in Oslo and Landau) or Kozubíková et al. (2009; for the

analyses in Prague). An environmental control (200 µl nuclease-free milliQ water in a tube, left open during the DNA extraction procedure) and an extraction blank control were included in the subsequent real-time PCR analyses as recommended by Vrålstad et al. (2009).

A quantitative TaqMan® MGB real-time PCR (Vrålstad et al. 2009) was conducted independently in all 3 laboratories to verify the presence of Aphanomyces astaci. This method is based on the detection of a 59 bp long A. astaci-specific fragment of the most variable part of the internal transcribed spacer region (ITS1) of the nuclear rDNA. Real-time PCR reactions were performed on a Mx3005P qPCR system (Stratagene) or on a Mastercycler® ep realplex S (Eppendorf) according to Strand et al. (2011), using the TaqMan® Environmental Master Mix to avoid PCR inhibition (Strand et al. 2011), and on an iQ5 Real-Time PCR Detection System (Bio-Rad) using the Universal PCR Master Mix (Life Technologies) according to Vrålstad et al. (2009). Data analysis was carried out using MxPro software version 4.10 (Stratagene), Real Plex 2.2 (Eppendorf), and iQ5 Optical System Software (Bio-Rad), respectively. A 4-fold dilution series of genomic DNA from a pure A. astaci culture (starting concentration: 5 ng µl⁻¹) served both as a standard and positive control. Unknown samples were included as concentrated (1x) and 5x diluted templates for the Stratagene and Eppendorf PCR setup, whilst concentrated (1x) and 10x diluted templates were analysed on the Bio-Rad system. Different dilutions, used to check for potential PCR-inhibition, were used to comply with standard protocols that had been established in the different laboratories.

The agent prevalence was quantified based on PCR-forming units (PFU), according to the methodology used by Vrålstad et al. (2009), in which 1 PFU corresponds to 1 amplifiable target DNA copy. Each sample was also assigned to semi-quantitative agent levels. Samples for which a weak signal below the limit of detection (= 5 PFU) was observed were considered negative (Vrålstad et al. 2009). Absolute quantification is possible in the absence of PCR inhibition above the limit of quantification (LOQ = 50 PFU according to Vrålstad et al. 2009). Possible real-time PCR inhibition was checked for by comparison of results obtained from concentrated and diluted samples. In the absence of inhibition, with

an acceptance of 15% variation, the difference in cycle threshold (C_t) values should be 2.32 \pm 0.35 and 3.32 \pm 0.48 for the 5 and 10× dilutions, respectively, when compared to undiluted DNA, (see Kozubíková et al. 2011b for more details). However, as many samples, particularly from *Orconectes limosus*, yielded results below LOQ, a quantitative comparison could not be made. Thus, we used a 1-tailed nonparametric Wilcoxon matched pairs test to evaluate whether the undiluted samples yielded significantly higher levels of agent DNA, which would at least imply that most samples were not substantially affected by inhibition.

The identity of *Aphanomyces astaci* was further confirmed by amplifying and sequencing a 529 bp long ITS fragment (including ITS1 as well as ITS2) from 6 DNA isolates representing both host crayfish species, according to protocol used by Oidtmann et al. (2006), with primers 42 and 640 designed to be specific for *A. astaci* (but see Kozubíková et al. 2011b for discussion on their limitation). The PCR products were sequenced in the forward direction using the primer 42, and resulting sequences (GenBank accession numbers JN713915–JN713917) were aligned in Mega 5 (Tamura et al. 2011) together with those representing all presently known *A. astaci* genotype groups (As, PsI, PsII, Pc, Or) isolated from various crayfish hosts: *Astacus astacus, Pacifastacus leniusculus, Procambarus clarkii*, as well as *Orconectes limosus* (see Kozubíková et al. 2011a). The alignment was then checked visually for the presence of differences among sequences.

RESULTS

Occurrence and dynamics of Orconectes limosus and Astacus leptodactylus

The front of the downstream invasion of the River Danube by *Orconectes limosus* was, by the end of the regular monitoring (March 2011), found to be near Sviniţa, 998 km upstream from the Black Sea (Fig. 1). The species was first detected at this site in May 2010, when only the upper 5 km stretch of Site SV contained *O. limosus*, whereas no individuals of the invasive species were captured in the downstream 5 km stretch. During the 2 yr duration of the regular monitoring, *O. limosus* moved down the River Danube by about 21 km from below Site BE to Site SV (Fig. 1). However, the most recent investigation, in August 2011, has confirmed the species at Dubova (Site DU), 25 km further downstream of Sviniţa.

Altogether, 2068 individuals of *Orconectes limosus* and 4552 of *Astacus leptodactylus* were captured during the survey period (January 2009 to March 2011). The 3 downstream sites in the River Danube (DU, IE, and SE) contained only *A. leptodactylus*, of which 3006 specimens were collected during the present study. A comparison of the relative abundances of both crayfish species at the upper 3 sampling sites along the Danube (CO, BE, and SV) shows a very large decline in the native species *A. leptodactylus* at Site BE, at which the relative abundance of the invasive species was about 75% (Fig. 2). The relative abundance of *O. limosus* at the invasion front (SV) in early 2011 was about 7.2% (Fig. 2). The sex ratio of *O. limosus* varied among sites: the percentage of captured males was 60.1% at CO, and decreased further downstream to 59.4 % at BE and 47.1 % at SV.

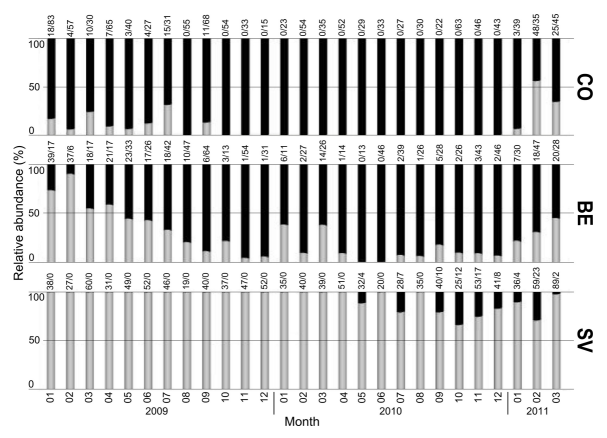


Figure 2 Astacus leptodactylus and Orconectes limosus. Relative abundance of native A. leptodactylus (grey bars) and invasive O. limosus (black bars) at 3 sites of the River Danube (see Fig. 1) over a period of 27 mo. Upper graph: CO (Coronini); middle graph: BE (Berzasca); lower graph: SV (Sviniţa; invasion front). Numbers above the bars refer to the absolute total number of A. leptodactylus and O. limosus, respectively, caught during each sampling period

The crayfish monitoring results differed between tributaries, depending mainly on the width of the tributary. In the small tributaries, Cameniţa, Radimna and Gornea (R1 to R3 in Fig. 1), each of which is less than 3 m wide at the confluence with the Danube, the invasive species was only found directly at the confluence, and only extended for some tens of metres upstream. In the Berzasca tributary (R4), with a mean width of about 7 m at the confluence, the invasive species was found up to 1400 m upstream of the tributary mouth. This was despite the fact that the time available for colonising this tributary was much shorter than for the tributaries R1 to R3 situated further upstream along the Danube. During the investigation period, a total of 161 individuals of *Orconectes limosus* were captured in the Berzasca. The distance separating the invasive species and the native *Austropotamobius torrentium* populations in the tributaries

was 1050 m at Cameniţa, 3200 m at Radimna, 4550 m at Gornea and 10800 m at Berzasca.

Molecular detection of Aphanomyces astaci

The crayfish plague agent Aphanomyces astaci was detected in 32% of the investigated Orconectes limosus specimens collected from Sites CO and BE (Table 1). PFU values in undiluted DNA samples from A. astaci-positive individuals ranged between 5 and approximately 27 000, corresponding to agent levels from A2 to A5 (Table 2), according to Vrålstad et al. (2009). Most of these samples were found to contain a low amount of the target DNA (Levels A2 and A3), and only one showed a high agent level (A5). Furthermore, A. astaci was also detected in 20 of the 49 investigated Astacus leptodactylus specimens (Table 2), even though their general state and behaviour did not indicate acute crayfish plague. Most importantly, we also detected A. astaci in 3 individuals of A. leptodactylus from Site SE, situated 68 km downstream of the O. limosus invasion front known at the time of sampling. All 6 ITS sequences obtained from isolates from 5 O. limosus from both Sites CO and BE, and 1 A. leptodactylus from BE, were invariant. They corresponded exactly to sequences from A. astaci strains of all 5 known genotype groups of the pathogen, including the one obtained from O. limosus (Kozubíková et al. 2011a). The sequencing thus corroborated the results of the real-time PCR analysis.

Table 1 Aphanomyces astaci infecting Astacus leptodactylus and Orconectes limosus. Number of crayfish from the Romanian Danube analysed for A. astaci infection using real-time PCR and percentage of specimens detected positive

Locality	Species	No. of analysed crayfish	<i>A. astaci-</i> positive	% infected		
Coronini (CO)	A. leptodactylus	1	1			
	O. limosus	37	9	24		
Berzasca (BE)	A. leptodactylus	24	6	25		
	O. limosus	34	14	41		
Sviniţa (SV)	A. leptodactylus	21	10	48		
Severin (SE)	A. leptodactylus	3	3	100		
Total	A. leptodactylus	49	20	41		
	O. limosus	71	23	32		

Table 2 Aphanomyces astaci infecting Astacus leptodactylus and Orconectes limosus. Summary data on the real-time PCR detection of A. astaci from native and invasive crayfish from 4 sites on the River Danube (see Fig. 1). Agent levels refer to semi-quantitative categories based on the observed numbers of PCR-forming units (PFU_{obs}), according to Vrålstad et al. (2009). Agent Level 0 (A₀; no detection) and Agent Level 1 (A₁; detection below the limit of detection: PFU_{obs} < 5 PFU) are both considered negative. Remaining categories (A₂ to A₅) are considered as a positive detection of the pathogen. A₂: 5 PFU \leq PFU_{obs} < 50 PFU, A₃: 50 \leq PFU_{obs} < 10³ PFU, A₄: 10³ PFU \leq PFU_{obs} < 10⁵ PFU

Commission site	Charles	No. of crayfish analysed	Agent level						
Sampling site	Species		A ₀	A ₁	A ₂	A ₃	A ₄	A ₅	
Coronini (CO)	A. leptodactylus	1						1	
	O. limosus	37	21	7	4	4		1	
Berzasca (BE)	A. leptodactylus	24	18		2	3		1	
	O. limosus	34	13	7	10	4			
Sviniţa (SV)	A. leptodactylus	21	11		5	2	3		
Severin (SE)	A. leptodactylus	3				3			

Significantly higher levels of *Aphanomyces astaci* DNA were found in samples of undiluted template DNA in the PCR setup, in comparison to $5 \times$ diluted template samples using Environmental Master Mix (1-tailed Wilcoxon matched pairs test, n = 31, Z = 2.86, p = 0.002) as well as when compared to $10 \times$ diluted samples using Universal PCR Master Mix (n = 9; Z = 2.67; p = 0.004). However, for 1 *Astacus leptodactylus* and 2 *Orconectes limosus* individuals, *A. astaci* was detected in diluted samples only. This suggests that although inhibition was not a major problem, it occasionally affected the efficiency of detection, potentially underestimating the extent of infection.

DISCUSSION

Our study clearly showed that *Orconectes limosus* is rapidly spreading downstream in the Romanian stretch of the Danube, and that its populations are infected by the crayfish plague pathogen *Aphanomyces astaci*. We also demonstrated that *A. astaci* had been transferred to local populations of the native narrow-clawed crayfish *Astacus leptodactylus*, which strongly declined in coexistence with *O. limosus*. Furthermore, the pathogen was detected in an *A. leptodactylus* population well in advance of the main invasion front of the invasive

species. The rate of *O. limosus* invasion in the Romanian Danube identified by our study was comparable to other studies quantifying its dispersal in this river. Given that in January 2009 *O. limosus* was found in the Berzasca region (Site BE; 1015 km before the Danube flows into the Black Sea), and reached Sviniţa by May 2010 (Site SE; 998 km upstream of the Black Sea), the estimated average colonisation speed is 1.23 km mo⁻¹, or about 15 km yr⁻¹, consistent with the findings of other studies (Puky & Schád 2006, Hudina et al. 2009). In August 2011 the species reached Dubova (DU), having moved 25 km down stream of Sviniţa, at a rate of 1.56 km mo⁻¹, slightly above the average calculated colonisation speed.

The serious decline in the relative abundance of the indigenous species Astacus leptodactylus in the Coronini area (CO), and the decrease in its relative abundance in the Berzasca area (BE), suggests that the invasive species has a tendency to eliminate the indigenous populations. The population dynamics of native European crayfish susceptible to crayfish plague that are in contact with invasive crayfish populations is, amongst other aspects, strongly dependent on the presence of the pathogen. For example, coexistence and a slow displacement of the native species due to the competitive advantages of the invader has been observed for mixed populations of Orconectes limosus and Astacus astacus in which the pathogen was not detected (Schulz et al. 2006) and also for coexisting A. astacus and presumably non-infected Pacifastacus leniusculus (Westman et al. 2002). When Aphanomyces astaci is present, however, a fast elimination of whole populations of European native crayfish can be expected (Westman et al. 2002).

Our confirmation of the causative agent of the crayfish plague in the Romanian Danube highlights the threat that the presence of *Orconectes limosus* poses to the indigenous crayfish populations (Longshaw 2011). We confirmed that substantial proportions of *O. limosus* carry the crayfish plague pathogen in both stretches of the Danube we investigated, where the species is already well established (CO and BE). Since the abundance of *O. limosus* is still low at the present invasion front, no conclusions can yet be made on the presence of *Aphanomyces astaci* for this site. However, we provided real-time PCR-based evidence for the presence of *A. astaci* in native *Astacus leptodactylus* (with

melanised spots on the exoskeleton) about 70 km below the suspected invasion front of *O. limosus* at the time of sampling. Although we cannot rule out that some infected *O. limosus* could have dispersed substantially farther than suggested by our monitoring data, it is likely that *A. astaci* is spreading downstream ahead of the invasive crayfish, by gradual expansion through local *A. leptodactylus* populations. Multiple mechanisms may contribute to the pathogen dispersal, apart from movement activity of crayfish. These include, for example, passive transport of infected live or dead crayfish (or even their exuviae). Furthermore, a direct dispersal of zoospores carried for a short distance by river currents may also be a source of infection for populations of *A. leptodactylus* further downstream. Due to these mechanisms, *A. astaci* may move down the River Danube much faster than the invasion front of *O. limosus*.

Dispersal of the crayfish plague through a river with abundant populations of susceptible crayfish may be very fast, as demonstrated by the rate at which the Danube catchment was affected during the major outbreak of crayfish plague in the 1870s–1890s (Alderman 1996). It is, therefore, possible that the pathogen will reach the Danube delta within a few years. On the other hand, if the rate of expansion of *Orconectes limosus* does not change substantially, the invasive crayfish itself may reach the Danube delta in the mid-2060s. However, anthropogenic long-range dispersal (for example, by shipping) may speed up its invasion considerably. The importance of invasive crayfish for introduction and dispersal of *Aphanomyces astaci* has also recently been reported for *Procambarus clarkii* in Italy (Aquiloni et al. 2011).

In crayfish populations sensitive to crayfish plague, the pathogen prevalence usually gradually increases over time, mostly reaching 100% (OIE 2009). This typically results in complete elimination of populations impacted by the plague, including stream populations of *Austropotamobius torrentium* (e.g. Kozubíková et al. 2008). However, some evidence exists that *Astacus leptodactylus* is at least sometimes more resistant to infection than other European species (Unestam 1969) and may be able to coexist with *Aphanomyces astaci* for extended periods of time. This has been implied particularly for some Turkish lakes where the pathogen has been assumed to be present continuously for over 2 decades,

despite the absence of American crayfish species in the country (Harlioğlu 2008). Recent molecular analyses of material from one of these Turkish lakes (Svoboda et al. 2012) confirmed that *A. astaci* may indeed persist in a population of *A. leptodactylus* that retains sufficient density for commercially harvesting. It is, therefore, possible that *A. leptodactylus* populations in the Danube will not be completely eliminated after introduction of *A. astaci* but will rather serve, at least temporarily, as a potential source of its further dispersal. This is also supported by our finding of *A. leptodactylus* in winter 2011 at Site CO, in which *Orconectes limosus*, and supposedly also *A. astaci*, had been present for about 4 yr (Fig. 2).

The facts that not only American but also native species of crayfish may serve as the source of crayfish plague infection and that not all North American crayfish are carriers of the pathogen, together with observed coexistence between European and North American crayfish species (Nylund & Westman 1992, Westman et al. 2002, Hudina et al. 2009, Skov et al. 2011), highlight the importance of using reliable detection methods for testing for presence of *Aphanomyces astaci* in suspected host populations. In the present study, we successfully applied MGB real-time PCR as proposed by Vrålstad et al. (2009) and Strand et al. (2011) to evaluate infection status independently in 3 laboratories and used ITS sequencing as an independent verification of the crayfish plague agent *A. astaci*.

We mostly found relatively low PFU values and corresponding very low (A2) and low (A3) agent levels (according to Vrålstad et al. 2009) but, nevertheless, unambiguously confirmed the presence of *A. astaci* in both invasive *Orconectes limosus* and native *Astacus leptodactylus*. Our results suggest that PCR inhibition was unlikely a major problem in our analyses. Nonetheless, the proportion of infected individuals might be underestimated, as we could not analyse the whole crayfish body for *A. astaci* presence, and detections of trace amounts of pathogen DNA (under 5 PFU) were considered negative to avoid potential false positives.

The results of monitoring of Danube tributaries confirmed the observations of Petrusek et al. (2006) from the Elbe catchment, where *Orconectes limosus* rarely penetrates far into loworder watercourses unless aided by humans. However, the risk of the crayfish plague agent expanding to native crayfish populations in small

streams cannot be ignored, even in the absence of any direct encounters with *O. limosus*, mainly because uncontrolled fishing takes place regularly in this area. *Aphanomyces astaci* can be transferred from one water body to the next through fishing gear, contaminated traps, the transfer of infected animals, and probably even by predators preying on crayfish (Nylund & Westman 1992, Alderman 1996, Oidtmann et al. 2002). It has been estimated that if the European-wide distribution and abundance of native stocks continue to decline and invasive crayfish species continue rapid expansion at the present rates, all European watersheds suitable for crayfish might be inhabited by invasive species within 100 yr (Skurdal & Taugbøl 2002, Holdich et al. 2009). The example of Sweden, where only around 5% of the native crayfish populations remain since the introduction of the crayfish plague (Fjälling & Fürst 1988, Bohman et al. 2006), shows the need for increased conservation effort to protect native species as soon as the plague agent enters a country.

Since invasive crayfish are not as widespread in Romania as in most other European countries, Romanian conservation managers have the opportunity to learn from the experience of other countries and act before it is too late. Based on our data, we assume the crayfish plague agent may reach the Danube delta very soon, and no measures can stop or slow it down. Protection of Romanian native crayfish populations may be made more efficient in preservation of *Astacus astacus* and *Austropotamobius torrentium* by identifying or artificially establishing so-called 'ark-sites' (Peay 2009a). Detailed investigations, and a well-organised management plan based on sound research findings, are strongly recommended, but additional conservation measures including education of local stakeholders and communities, fishermen, and children (Peay 2009b) are needed as well. This is particularly relevant in this region, where fishing is a major occupation, and simple and inexpensive measures (such as avoidance of using the same fishing tools in other rivers as in the Danube) may prevent the disease spreading from the Danube into the tributaries.

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Crayfish plague pathogen detected in the Danube Delta – a potential threat to freshwater biodiversity in southeastern Europe

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ABSTRACT

The crayfish plague, caused by the oomycete Aphanomyces astaci, is probably the most significant reason for declines in European freshwater crayfish species. One of its hosts, the North American spiny-cheek crayfish Orconectes limosus, extends its range in the river Danube and recently reached the territory of Romania. We used highly sensitive A. astaci-specific real-time PCR to test if the native narrow-clawed crayfish Astacus leptodactylus in the highly protected Danube Delta about 970 km downstream of the current invasion front of American crayfish is a carrier of the crayfish plague. Thirteen out of 40 analysed native A. leptodactylus tested positive for the crayfish plague pathogen, infected individuals were found at both sampled localities within the Danube Delta. Therefore A. astaci has a much wider range in this river than assumed. The pathogen seems to persist in local populations, as neither crayfish mass mortalities nor alien crayfish species have been reported from the region. Aphanomyces astaci may have reached the Delta by long-range passive dispersal of infected hosts or pathogen spores, or by gradually infecting populations of native crayfish in upstream regions of the Danube in a stepping-stone manner. Alternatively, the crayfish plague may have persisted in the Danube Delta as chronic infection from an old plague wave in the 19th century. In any case, the presence of this pathogen in the lower Danube may become a threat to conservation of European crayfish and to freshwater biodiversity in many regions of southeastern Europe, at present considered 'crayfish plague-free'.

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Key words: *Aphanomyces astaci* • *Astacus leptodactylus* • Black Sea • crayfish plague • Danube Delta • freshwater biodiversity • invasive species

INTRODUCTION

The distribution of freshwater organisms is affected by varying climatic cycles and topographic features (e.g. changes in river flow) and in the last centuries to a high degree by direct and indirect human impacts (e.g. by species translocations, habitat alterations, and anthropogenic pollution). For several decades, the distribution and abundance of native European crayfish species has been strongly affected by the crayfish plague (Holdich 2002). The oomycete Aphanomyces astaci Schikora, the causative agent of this disease, had been most probably introduced from North America to Europe in the late 1850s together with some of its original hosts, North American freshwater crayfish species (Alderman 1996). This aggressive pathogen is listed among the world's 100 worst alien species (Lowe 2004) because of its devastating effects resulting in mass mortalities of whole populations of European freshwater crayfish. Since its introduction, the crayfish plague has destroyed many European crayfish populations and caused substantial losses to wild crayfish stocks as well as to valuable fisheries (Alderman 1996). Three of its original host species in Europe extended their range within the continent enormously by active migration as well as by humanmediated dispersal. The widespread presence of American crayfish populations that serve as reservoirs of the pathogen (e.g. Kozubíková et al. 2009) makes it a continuous threat to local crayfish (Holdich et al. 2009). Up to now, it has been believed that all native crayfish from Europe are highly susceptible to crayfish plague, and that infection by the plague pathogen generally leads to their death. However, a few recent studies reported that native crayfish populations may persist for several years or even decades with certain levels of infection by A. astaci (Jussila et al. 2011; Kokko et al. 2012; Svoboda et al. 2012). Despite these exceptional cases, crayfish plague poses a high risk to waters not yet affected by the disease, especially in eastern European countries where it is less widespread (Holdich et al. 2009).

In the lower Danube basin, there had been no reports of outbreaks of this disease for decades, although the whole river had been substantially affected by crayfish plague in the late 19th century (Alderman 1996). However, as in other parts of Europe, American crayfish had not been reported from the Danube basin during that first huge infection wave but colonised it only much later. The spinycheek crayfish (Orconectes limosus Rafinesque, 1817) was recorded for the first time in the river in 1985 in Hungary (Puky and Schad 2006). Since then it has spread along its course and reached Romania by 2008 (Pârvulescu et al. 2009). There it coexists and slowly displaces populations of native narrow-clawed crayfish (Astacus leptodactylus Eschscholtz, 1832) (Pârvulescu et al. 2012). Spiny-cheek crayfish populations in the Danube were repeatedly shown to host A. astaci (Kozubíková et al. 2010; Pârvulescu et al. 2012). Interestingly, the presence of this pathogen was also confirmed by molecular methods in healthylooking individuals of narrow-clawed crayfish ~70 km downstream of the presumed invasion front of spiny-cheek crayfish in the Danube (Pârvulescu et al. 2012) in August 2011. This sampling site is located ~900 km upstream of the river delta.

The downstream colonization rate of the Danube by the spiny-cheek crayfish was estimated to about 13-16 km·yr⁻¹ (Puky and Schád 2006; Pârvulescu et al. 2012). At this rate, this invasive species would reach the Danube Delta no sooner than in the 2070s. However, the presence of *A. astaci* in healthy-looking narrow-clawed crayfish (Pârvulescu et al. 2012) suggested that the crayfish plague pathogen may be steadily spreading ahead of the invasion front of American crayfish. The knowledge of the geographic distribution of the crayfish plague pathogen is important in order to prioritize conservation management of the most endangered populations. Presence and absence data of *A. astaci* is, for example, important for the concept of 'ark sites', a common component of modern management plans for endangered crayfish in Europe (Kozák et al. 2011). An ark site is an isolated refuge site where native crayfish species are not at the risk from adverse factors, including colonization by invasive crayfish species and *A. astaci* (Peay 2009). Individuals from non-infected populations that are in danger of getting infected by crayfish plague in the near future can be translocated to ark-

sites to save them from dying and to conserve the intraspecific diversity. Translocation from an infected population, on the other hand, holds the threat of further spreading the agent of the crayfish plague.

The Danube Delta is included in the UNESCO List of World Natural Heritage Sites (UNESCO 2012) because of its outstanding biodiversity. An infection of native crayfish with *A. astaci* in this highly protected area may have dramatic consequences. The elimination of native crayfish may lead to a strong cascading effect in the whole food web with unpredictable implications for other freshwater organism. To evaluate if the concept of ark sites can be applied to crayfish from the Danube Delta and to further provide a basis for conservation management measures, we tested native narrow-clawed crayfish from the Danube Delta for possible infection by *A. astaci*, using a highly specific and sensitive molecular method to detect pathogen DNA.

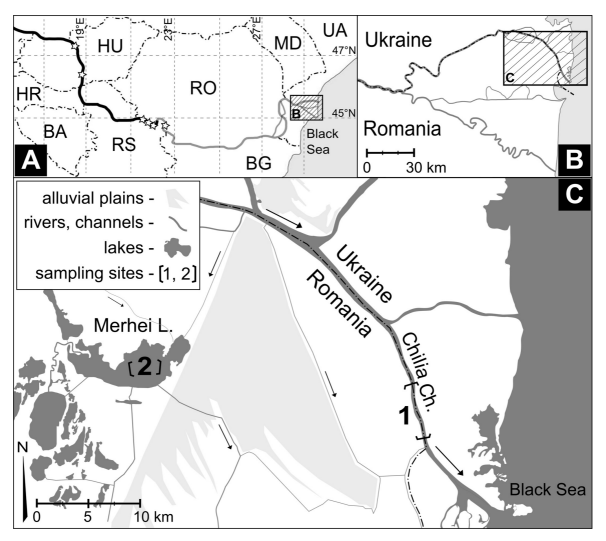


Figure 1. A: Overview of the distribution of *Orconectes limosus* (dark line) and the sites with confirmed presence of the crayfish plague pathogen (stars) in the Danube; **B**: Overview of the Danube Delta; **C**: Sampling sites in the Danube Delta (1- Chilia Channel, 2- Merhei Lake). The arrows indicate the water flow direction.

METHODS

In May 2011, we captured freshwater crayfish at two different places within the Danube Delta, in the Chilia main channel and in the Merhei Lake (Figure 1). Water exchange between the channel and the lake, which are connected by 12 kilometres of a narrow canal, is very low. Five days were spent along a 5-km stretch of the Chilia Channel (between 45.31N, 29.67E and 45.27N, 29.68E) by capturing crayfish into traps. Due to a difficult access, only one daylight capture was possible in the Merhei Lake (45.32N, 29.45E), by using fish-baited nets. Additionally, we checked a local fish market and catches of five local fishermen

around the Danube Delta for any non-native crayfish species that might be available for sale.

To avoid any potential cross-contamination by the pathogen, we used new traps and a toolbox without any previous contact with crayfish. As whole specimens of narrow-clawed crayfish cannot be collected in this protected area, only two parts of the uropods and, if present, one melanised walking leg, were sampled from each crayfish and stored in 96% ethanol. We extracted DNA from the crayfish tissues as described in Vrålstad et al. (2009). For detection of *A. astaci* DNA, we used the quantitative TaqMan® minor groove binder (MGB) real-time PCR (Vrålstad et al. 2009), the most sensitive and specific detection assay available at present (Tuffs and Oidtmann 2011). Real-time PCR reaction was performed on a Mastercycler® ep realplex S (Eppendorf) using the TaqMan® Environmental Master Mix to avoid PCR inhibition (Strand et al. 2011). We increased the annealing temperature to 62°C and decreased the annealing time to 15 seconds to further exclude possible false positive results (T. Vrålstad, unpublished data; see also Kozubíková et al. 2011b).

To increase credibility of results, the sample set was subdivided into three subsets and different people tested each subset on a different day, with similar results. Apart from using negative controls (i.e., samples containing no DNA), we ruled out potential laboratory contamination by analysing one sample subset together with presumably non-infected spiny- cheek crayfish from German populations coexisting with susceptible crayfish. All negative controls, including DNA isolates from these American crayfish, tested negative.

We carried out data analysis using the software Real Plex 2.2 (Eppendorf). The relative level of infection by the pathogen depends on the strength of the real-time PCR signal and corresponding amounts of PCR-forming units in the reaction, and is expressed as semi-quantitative agent levels (according to Vrålstad et al. 2009; see also Table 1).

RESULTS

Altogether 40 individuals of narrow-clawed crayfish were collected in the Danube Delta, 37 from the Chilia Channel and three from Merhei Lake. No other crayfish species were observed, and all investigated crayfish specimens (approximately 600 individuals) from commercial fish captures were identified as the native narrow-clawed crayfish.

Table 1. Real-time PCR detection of *Aphanomyces astaci* in investigated specimens of *Astacus leptodactylus* from the Danube Delta.

Sampling site	No. of analysed	A. astaci	Agent level ^a						
	crayfish	positive	A ₀	A ₁	A ₂	A ₃	A ₄	A ₅	A ₆
Chilia Channel	37	11 (30%)	23	3	5	3	1	1	1
Merhei Lake	3	2 (67%)		1	2				
Total	40	13 (32%)							

^aRelative levels of infection (expressed as agent level after Vrålstad et al. 2009) are given. Individuals with agent levels A_0 (no detection) and A_1 (very low real-time PCR signal, corresponding to less than 5 PFU_{obs}, i.e., PCR forming units observed in the reaction) are conservatively considered uninfected. Individuals with agent levels A_2 and over (A_2 : 5 ≤ PFU_{obs} < 50, A_3 : 50 ≤ PFU_{obs} < 10³, A_4 : 10³ ≤ PFU_{obs} < 10⁴, A_5 : 10⁴ ≤ PFU_{obs} < 10⁵; A_6 : 10⁵ ≤ PFU_{obs} < 10⁶) are considered infected by A. astaci.

Presence of *Aphanomyces astaci* was confirmed at both sampling sites. Overall, DNA of the pathogen was detected by the real-time PCR in 32% of analysed narrow-clawed crayfish specimens (Table 1): in 11 out of 37 individuals captured from the Chilia Channel, and in two out of three specimens from the Merhei Lake. Most samples contained low levels of pathogen DNA ($7 \times$ agent level A_2 , $3 \times A_3$); however, three individuals from the Chilia Channel were highly infected (agent level A_4 or more). Melanised areas were observed in the walking legs and in the telson, uropods or abdominal cuticle of four individuals from the Chilia Channel, all of which were infected by *A. astaci*.

DISCUSSION

We have demonstrated that a substantial proportion of narrow-clawed crayfish populations in at least some parts of the Danube Delta were infected by Aphanomyces astaci, although no mass mortalities of crayfish have been reported from this region for several decades. The observed levels of infection might still be underestimated, as various body parts in which A. astaci may be present could not be analysed due to species conservation legislation. However, crayfish plague detection rate in signal crayfish (Pacifastacus leniusculus Dana, 1852) was almost as high for uropods (the tissue of choice in this study) alone as for multiple body parts (Vrålstad et al. 2011). Melanised spots that were observed in four infected narrow-clawed crayfish could have been a visible indication of infection (as considered e.g. by Nylund and Westman 2000) but such spots also appear due to immune reaction to other pathogens and after a mechanical injury (Schulz et al. 2006). In a Turkish population of A. leptodactylus in which A. astaci seems to persist for over two decades, melanisation was observed on crayfish infected by the pathogen as well as on those in which real-time PCR assay did not confirm its presence (Svoboda et al. 2012). On the other hand, an absence of melanised spots is not an indication of pathogen-free crayfish. We did not notice these signs in the other nine positive as well as in negative tested narrow-clawed crayfish. These symptoms should be thus interpreted with care.

The capture success in the Merhei Lake, from which we obtained only three individuals, was low. Although no unusually high crayfish mortalities have been reported from the Danube Delta recently, we cannot rule out the possibility that a population decline already took place in this lake. However, the catching was constrained by the fact that we could only spend one day at this lake and the access to the water edge was limited.

Recent dispersal of the pathogen

As no non-native crayfish species have ever been reported from the Danube Delta, nor were they captured in the investigated areas or observed on the markets, it is unclear how *A. astaci* has reached the region. One plausible possibility is that the infection originates from American crayfish in the upstream

regions of the Danube, and has spread downstream much earlier and faster than expected by Pârvulescu et al. (2012). There are several potential processes that may have ensured long-range pathogen dispersal: (i) The plague pathogen could have been infecting native narrow-clawed crayfish that are widely distributed in the lower Danube in a stepping-stone manner, by short-range downstream dispersal of zoospores between susceptible animals. Since the survival of spores in freshwater is at least seven days at 14°C (Unestam 1969), the spores, if released, may be carried in running water for around 150 kilometres considering a current velocity of 0.5 m·sec⁻¹. (ii) Passive transport of infected live or dead crayfish (or even their exuviae) may have contributed to the pathogen dispersal since A. astaci remains viable for at least five days in a crayfish cadaver (Oidtmann et al. 2002). (iii) Other animals, particularly fish feeding on infected crayfish, may transport the pathogen from infected to healthy populations (Oidtmann et al 2002). (iv) Boats that frequently pass through the Danube could transport non-native crayfish or pathogen spores (e.g. in the ballast water) for long distances. We also cannot rule out an undetected expansion of spiny-cheek crayfish to some lower reaches of the Danube.

Relic from an old infection wave?

An alternative explanation for the origin of the present infection in the Danube Delta is that the crayfish plague pathogen could be a relic from the original infection wave that caused mass mortalities in crayfish along the Danube in the 19th century (Alderman 1996). Since then, the pathogen might have persisted, possibly as chronic infection, in local narrow-clawed crayfish populations for more than a century. This would contradict the general assumption that the pathogen is lethal to this native European crayfish species. However, there is some evidence that long-term coexistence of narrow-clawed crayfish and *A. astaci* may be possible. In particular, it has been repeatedly reported that *A. astaci* persists in some Turkish lakes inhabited by narrow-clawed crayfish since the mid-1980s (e.g., Harlioğlu 2008), and this has been supported by the recent molecular detection of *A. astaci* (Svoboda et al. 2012; Kokko et al. 2012). Similarly, coexistence of presumably even more sensitive noble crayfish *Astacus astacus* (Linnaeus, 1758) with *A. astaci* was recently reported from two

Finnish lakes in which the crayfish plague agent had probably been present for several years without a proven presence of alien crayfish (Jussila et al. 2011; Viljamaa-Dirks et al. 2011).

Several mechanisms that may promote coexistence between native European crayfish and the crayfish plague pathogen have been discussed but so far they remain unclear (Jussila et al. 2011; Viljamaa-Dirks et al. 2011; Svoboda et al. 2012). It is assumed that some native crayfish exhibit certain levels of tolerance to the infection, possibly strengthened by a strong selection pressure due to the initial crayfish plague outbreaks. Furthermore, it is possible that the pathogen has also adapted to the new hosts and lowered its virulence over the years since it is evolutionary disadvantageous from the pathogen's perspective when the host population dies out. In order to resolve this question, laboratory experiments with susceptible crayfish exposed to A. astaci should be designed and potential differences in crayfish mortality rates recorded. The settings should vary with respect to (i) the origin of *A. astaci* (regions where either latent infections of susceptible crayfish or mass mortalities are observed), (ii) the origin of crayfish hosts (populations coexisting with the pathogen, such as those in the Danube Delta or Turkey, vs. other regions), (iii) A. astaci spore concentration (which likely varies with different host densities).

It seems that different American crayfish species carry distinct pathogen strains (Huang et al. 1994; Kozubíková et al. 2011a), which might differ in their virulence to European crayfish. For instance, signal crayfish host other *A. astaci* strains than those involved in the first crayfish plague infection wave or those recently isolated from spiny-cheek crayfish (Kozubíková et al. 2011a). The signal crayfish currently extends its range to the Danube tributaries in Croatia (Hudina et al. 2009) and may reach the lower Danube in the future, possibly introducing a different *A. astaci* strain that might be more virulent to narrow-clawed crayfish. It appears that signal crayfish rapidly displaced some native crayfish populations in Croatia (Hudina et al. 2009), probably due to transmission of *A. astaci*. Although narrow-clawed crayfish in the Danube Delta seems to persist at present in the presence of *A. astaci*, this might change in case of infections by a more virulent strain of this pathogen.

Threat to freshwater biodiversity

Despite occasional reports on coexistence of populations of European crayfish with *A. astaci*, we have to assume that native crayfish species are threatened by further dispersal of the crayfish plague agent, and local stakeholders should react accordingly. Besides narrow-clawed crayfish, this applies to native noble crayfish and stone crayfish (*Austropotamobius torrentium* Schrank, 1803), distributed in the drainage area of the lower Danube (Holdich et al. 2009) and to the mesohaline thick-clawed crayfish (*Astacus pachypus* Rathke, 1837), which occurs only in small parts of the Black and Caspian seas (Holdich et al. 2006). The Southern Balkans, a glacial refuge for many freshwater species (Hewitt 1999), harbours high genetic diversity within European crayfish (Trontelj et al. 2005; Schrimpf et al. 2011).

It is possible that we have detected the infection in the Danube Delta in an early stage and that mass mortalities will follow, as described by Alderman (1996) for the 1890s. Especially when spores are dispersed in low densities and water temperature is low, mortalities may not be apparent for a long time (Alderman et al. 1987). Therefore, crayfish populations in the Danube Delta might be hit by crayfish plague outbreaks associated with high crayfish mortalities when the water temperature increases in the summer. European crayfish, as the largest freshwater invertebrates and due to their trophic activities as omnivores, play a key role in many freshwater ecosystems (Nyström 1999), and their loss may have drastic impacts on local biodiversity. This is relevant for all regions threatened by crayfish plague but might be particularly important in the wetlands of the Danube Delta.

Conservation implication

Since funding for conservation is often limited, a precise knowledge of the situation and accurate prediction of sites most at risk of becoming invaded by non-native species is fundamental to create an effective management plan (Keller et al. 2008). We presume *A. astaci* is probably present along the whole Danube main channel, therefore translocations of freshwater crayfish as well as conservation actions on crayfish (e.g. the concept of ark sites) in the Danube

have to be considered carefully. On the other hand, the side channels of the Danube could be prioritized for management actions. Native crayfish populations in the side channels that are threatened by a further dispersal of *A. astaci* and are negatively tested for an infection could be translocated to ark sites to avoid further loss of within-species diversity. Furthermore, since many invasive species, including North American crayfish and the crayfish plague pathogen carried by them, are difficult or impossible to eradicate, the best option for limiting total impacts is often to restrict spread (Keller et al. 2008). Therefore, local stakeholders, communities and fishermen need to get informed about the high risk of spreading invasive crayfish or even native crayfish infected with *A. astaci*.

An important question is whether *A. astaci* will be able to disperse from the Danube through the brackish Black Sea (with an average salinity of 18 psu) into other river basins. It remains open whether it could survive in hosts that tolerate salinities up to 21 psu (e.g. narrow-clawed crayfish, signal crayfish and spiny-cheek crayfish; Jażdżewski et al. 2005). Furthermore, the dispersal of the pathogen might be possible by ship traffic. To avoid spreading of *A. astaci* through plague-free eastern European countries by cargo ships, ballast water treatment systems should be strictly implemented.

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Freshwater Biology (in press)

Absence of the crayfish plague pathogen (Aphanomyces coexistence of European and American astaci) facilitates crayfish in central Europe

Running head: Plague absence support crayfish coexistence

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SUMMARY

Most European crayfish species are strongly threatened, mainly as a result of the introduced crayfish plague agent, Aphanomyces astaci, transmitted by invasive North American crayfish. Long-term coexistence of American and European crayfish species is therefore regarded as almost impossible, even thought some coexisting populations have been observed.

In this study, crayfish were collected from presently coexisting populations of the introduced spiny-cheek crayfish (Orconectes limosus) and the native noble crayfish (Astacus astacus) from nine standing waters in central Europe. Our aim was to resolve whether the coexistence resulted from reduced virulence in local strains of A. astaci, increased immunity in the native crayfish, or an absence of the pathogen in these populations. We used highly sensitive A. astaci-specific real-time PCR to evaluate the crayfish latent carrier status, combined with transmission experiments to further validate the molecular results.

From the total of 523 crayfish tested (490 spiny-cheek crayfish, 33 noble crayfish), none positive for A. astaci were detected. Transmission experiments confirmed these results: No abnormal mortality or behavioural changes were seen in noble crayfish kept together with American crayfish from the coexisting populations. If we assume a very low prevalence of A. astaci of 10% in a carrier population, there is a 98% probability of disease absence in five of the nine

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coexisting populations tested. Hence, a consistent absence, or an extremely low prevalence of *A. astaci*, seems to allow the coexistence of European and American crayfish in these central European populations.

The results are important for native crayfish conservation and management, and demonstrate that disease transmission risk may vary substantially between the different populations of spiny-cheek crayfish in central Europe.

Keywords: Crayfish plague • Invasive species • Coexistence • Real-time PCR • transmission experiment

INTRODUCTION

Land use, climate and biotic exchange are the main drivers of the loss of freshwater biodiversity (Sala et al., 2000). The introduction of alien species is thought to have a higher impact on biodiversity in aquatic than in terrestrial ecosystems, and particularly in lakes (Sala et al., 2000). Crayfish are the largest, mobile freshwater invertebrates, occupying lakes, rivers and streams. They are considered keystone species with a high biological impact on their environment due to their omnivory (Holdich, 2002; Reynolds & Souty-Grosset 2012). Today, the existence of all the native European crayfish species is gravely threatened by the introduction and spread of alien species. The crayfish plague pathogen, Aphanomyces astaci (Oomycetes), carried and transmitted by North-American crayfish species is the major cause of the sharp decline in native crayfish populations in Europe. While this obligate crayfish parasite lives in a balanced host-parasite relationship with North American crayfish, it is a lethal pathogen of native European crayfish, usually leading to 100% mortality (OIE, 2012). Therefore, this aggressive pathogen is listed among the world's 100 worst alien species (Lowe et al., 2004). Aphanomyces astaci originates from North America (Unestam, 1972), but has repeatedly been brought into Europe along with introduction of American crayfish species (Alderman, 1996).

If American crayfish species are stressed or immunocompromised as a result of moulting, infection by other pathogens or an adverse environment, they too may succumb to crayfish plague (Persson *et al.*, 1987; Söderhäll & Cerenius,

1992). However, they normally cope with *A. astaci* due to evolved defence mechanisms involving melanin encapsulation of the parasite that prevents further spread within the crayfish (Unestam, 1969; Söderhall & Cerenius, 1999). Consequently, the animals can be clinically healthy, but still act as infection reservoirs of *A. astaci* (Söderhall & Cerenius, 1999; Taugbol, 2004; Holdich *et al.*, 2009; OIE, 2009). Four species of North American crayfish introduced to Europe are confirmed carriers of the disease: i.e. the spiny-cheek crayfish (*Orconectes limosus*), the signal crayfish (*Pacifastacus leniusculus*), the red swamp crayfish (*Procambarus clarkii*) (Holdich *et al.*, 2009), and the calico crayfish (*Orconectes immunis*) (Schrimpf *et al.*, 2013). *Orconectes limosus* was the first known carrier of *A. astaci* in Europe (Kozubiková *et al.*, 2011a). It was introduced to Poland and Germany around 1890, and is now wide spread in 21 countries (Holdich *et al.*, 2009).

All North American crayfish should according to the OIE (2009) be regarded as likely carriers of A. astaci. This, together with the catastrophic decline of native European crayfish (Bohman, Nordwall & Edsman, 2006, Holdich et al., 2009), suggests that the stable coexistence of American and European crayfish species is unlikely. However, examples of apparently long-term coexistence have been found ocassionally for signal crayfish and noble crayfish (Astacus astacus) in Finland (Westman & Savolainen, 2001; Westman, Savolainen & Julkunen, 2002) and Sweden (Söderbäck 1994), for spiny-cheek crayfish and native narrowclawed crayfish (Astacus leptodactylus) in Romania (Pârvulescu et al., 2012), and for spiny-cheek crayfish and noble crayfish in lakes from Poland (Schulz et al., 2006). Indications of pathogen free American crayfish populations have also been found sporadically (Matasová et al., 2011; Skov et al., 2011). Schulz et al. (2006) suggest that the displacement of noble crayfish by spiny-cheek crayfish in some lakes in Poland is caused by direct competition instead of via A. astaci infection of the European species. A replacement of noble crayfish was also proposed by Westman et al. (2002) for a long-term coexisting population of signal crayfish and noble crayfish in Finland.

The coexistence of American and European crayfish species might be possible if some European crayfish have evolved resistance against the pathogen.

In this case, the pathogen should be found in the population. The same would apply if a less virulent strain of *A. astaci* had evolved that is not lethal to European crayfish species. It has been shown that different North American crayfish species are carriers of different genotypes of *A. astaci* (Huang, Cerenius & Söderhäll, 1994; Diéguez-Uribeondo, Cerenius & Söderhäll, 1995, Kozubíková *et al.*, 2011a) and that these genotypes may vary in their virulence (Viljamaa-Dirks *et al.*, 2011). Makkonen *et al.* (2012) assume that the high genetic diversity of *A. astaci* chitinase genes may indicate its adaption to specific host species.

These new results may explain why a few populations of native European crayfish are resistant to some extent against the pathogen, as has been demonstrated recently in a population of noble crayfish in Finland where the *A. astaci* genotype group A, or As-type, was isolated in pure culture (Viljamaa-Dirks *et al.*, 2011). The As-genotype has only been isolated from noble crayfish and hence the original American host species in unknown (Huang *et al.*, 1994). The high polymorphism of the As-genotype, and coevolution with noble crayfish may explain its adaptation to native European crayfish species (Makkonen *et al.*, 2012). Other recent papers on the basis of molecular evidence suggest that persistent infections can occur in native crayfish populations (Jussila *et al.*, 2011; Viljamaa-Dirks *et al.*, 2011; Kokko *et al.*, 2012; Schrimpf *et al.*, 2012; Svoboda *et al.*, 2012). A third scenario that may allow the European species to coexist with American species could be the complete absence of the pathogen in these stocks. In that case, all tests for the presence of the pathogen should be negative.

In the present study, we aimed to test which of these three hypotheses (i.e. a) resistant native crayfish, b) reduced pathogenicity, c) no pathogen) may apply to the apparently long-term coexistence of the North American spiny-cheek crayfish and the native noble crayfish in several populations in central Europe. We used quantitative real-time PCR to clarify the *A. astaci* carrier status in the coexisting stocks of nine waters in Germany and Poland, and strengthened this molecular analysis with classic transmission experiments.

METHODS

Study Sites and Crayfish Sampling

With the help of local fishery administrations in Germany and Poland, coexisting populations of spiny-cheek crayfish and noble crayfish were localized. Between 2006 and 2011, we trapped noble crayfish and spiny-cheek crayfish, between July and September each year from five and four lakes in Poland and in Germany, respectively, where the species coexist (Table 1). The two species have coexisted in these lakes at least since 2006, when this study was started, and no exceptional mortalities of noble crayfish (indicating a plague outbreak) have been reported since then. In the five Polish lakes, the first records of the coexistence are from 2000 (Milaczewo), 2001 (Piasek, Trzcinskie), 2003 (Zielone) and 2006 (Plociowe). When possible we collected the invasive crayfish species without harm the native noble crayfish. The lakes are between approximately 5 and 50 ha surface area. We also collected spiny-cheek crayfish from two German and Polish lakes (Plociowe and Hennweiler), where no noble crayfish have been observed for at least one decade. We also collected signal crayfish from the Heimbach, a German river where native species are no longer present. These individuals where primarily included as positive controls for demonstrating the expected positive carrier status for American crayfish. Finally, we took noble crayfish from the "First Bavarian Crayfish hatchery" in Augsburg, Germany as a negative control (Table 2). This hatchery has been surveyed for mortalities of noble crayfish very carefully before and after sampling, and none has been reported for more than 20 years (Max Keller, pers. comm).

Molecular analyses

Reliable evaluation of the presence or absence of the crayfish plague pathogen is best achieved with molecular diagnostics. A molecular screening of American and European crayfish using the agent specific and quantitative TaqMan® minor groove binder (MGB) real-time PCR (qPCR; Vrålstad *et al.*, 2009) was performed in order to detect putative *A. astaci* infections. The assay has demonstrated very high sensitivity and specificity for *A. astaci*, and is recommended for evaluating the carrier status of American crayfish (Tuffs &

Oidtmann, 2011). A total of 490 spiny-cheek crayfish and 33 noble crayfish were tested from coexisting populations, and eight noble crayfish were tested from the crayfish farm in Augsburg (Tables 1 & 2). As indicated above, spiny-cheek crayfish and signal crayfish from the single species lakes (Table 2), were included in the qPCR analyses as positive controls.

Due to two different time periods of processing and qPCR analyses of the data included in this study, we will in the following refer to two sample sets that were processed separately (A and B, Tables 1 & 2). For sample set A (244 samples), only the soft abdominal cuticle was used for DNA-extraction. Here, the cuticle was cut into two halves longitudinally and the two samples tested. For sample set B (319 samples), three tissue types were included (abdominal cuticle, uropods and the inner joint of a walking leg according to Vrålstad et al., 2011). After each dissection, tools were disinfected using DNAexitus (AppliChem GmbH, Germany). The DNA was extracted according to Vrålstad et al. (2009). An extraction blank control not including tissue material was included in each DNA extraction setup, and an environmental control consisting of 200 µl nuclease free water that was left open during the extraction procedure. Both types of control were included in the further qPCR analyses. The qPCR procedure was performed according to Vrålstad et al. (2009) with some modifications. The qPCR for sample set A was run on a MJ research Chromo4, and the Universal PCR Master Mix was replaced by 1 u GoTaq DNA Polymerase (Promega, Germany), 1× Colourless Go Tag® Flexi Buffer (Promega, Germany), 2.5 mM MgCl₂ (Promega, Germany), 0.2 mM of each dNTP (Fermentas, Germany) and 2 µg BSA (Fermentas, St. Leon-Rot, Germany). For this sample set, agent levels were assigned on the basis of corresponding Ct-values (Vrålstad et al., 2009; Kozubíková et al., 2011b). Sample set B was run on a Mx3005P qPCR system (Stratagene, USA) using the TaqMan® Environmental Master Mix (Applied Biosystems, USA) to avoid PCR inhibition (Strand et al., 2011). For half of the sample set B, the annealing temperature was increased to 62°C and the annealing time decreased to 15 seconds to further increase the assay specificity (T. Vrålstad, unpubl. data). Data analysis was carried out using MxPro software ver. 4.10 (Stratagene, USA) and Opticon Monitor 2.03 (MJ Research, Canada). DNA-

extracts were tested in duplicates as concentrated (1x) and 5x diluted templates. A no-template PCR control consisting of 5 µl nuclease free water was included in each run together with a standard series of genomic DNA from a pure *A. astaci* culture. The standard series (a four-fold dilution series) included four calibrant points for each run, and was used to assign positive results to semi-quantitative agent levels according to Vrålstad *et al.* (2009).

Statistics

A statistical test was conducted in R v.2.12.2 (R Development Core Team, 2011) using the function epi.detectsize in the package epiR (Stevenson *et al.*, 2010) to estimate the probability of the absence of disease (i.e. *A. astaci*). The function computes the number of random samples (individuals) required to declare a population free from the crayfish plague agent at a certain confidence level depending on the expected prevalence of the pathogen in an infected population (or the theoretical prevalence level we want to be able to reveal), the given population size (N), test sensitivity (Se) and specificity (Sp) (Dohoo, Martin & Stryhn, 2003).

If we assume a mean population density of two individuals m⁻² for an unexploited population (Abrahamsson, 1966), the population size (including crayfish of all size classes) varies between 10⁵ individuals for a 5 ha lake and 10⁶ individuals for a 50 ha lake. We do not know if the population sizes for mixed coexisting populations vary, therefore the test was run for different hypothetical population size estimates that should apply to coexisting populations as well as to single species lakes (N = 10³; N = 10⁴; N = 10⁵; N = 10⁶). Expected prevalence (Prev = % positive individuals in a population), including the smallest prevalence we want to be able to reveal, was also tested with different estimates (high Prev = 0.8; low Prev = 0.3; very low Prev = 0.1; exceptionally low Prev = 0.01). The real-time method used has very high sensitivity (Vrålstad *et al.*, 2009; Tuffs & Oidtmann, 2011). While analysis of the uropod tissue alone detected 84% of the total of 86% positives of signal crayfish, analyses of the soft abdominal cuticle targeted only 30% (Vrålstad *et al.*, 2011). Notably, in spiny-cheek crayfish, the soft abdominal cuticle was most frequently positive (85% positives of a total of

92% infected specimen; Oidtman *et al.*, 2006). Therefore, a test sensitivity of 0.8 seems realistic for spiny-cheek crayfish, even when only the cuticle is analysed. However, to account for various sensitivity scenarios, we included three different sensitivity estimates (0.3; 0.5; 0.8) in the statistical tests. Specificity was set to 0.99, since the real-time method used has proved to be very specific and detects all known genotypes of *A. astaci* (Vrålstad *et al.*, 2009; Tuffs & Oidtmann, 2011). The test was run for two confidence levels (0.98 and 0.95).

Transmission experiments

In the transmission experiment noble European crayfish susceptible to A. astaci and potentially infected American crayfish were kept together. High mortality and behavioural change of the native species indicates a positive carrier status of the American crayfish. Spiny-cheek crayfish from German and Polish stocks were kept under controlled conditions, together with noble crayfish from the crayfish farm in Augsburg (Germany) (Table 3) for 63 and 91 days, respectively. From each of the coexisting populations, and from the single species population from Plociowe, we placed six crayfish (three spiny-cheek crayfish and three noble crayfish from the crayfish farm, the latter one known not to be infected) in a separate aquarium with a bottom area of 0.125 m² (31.25 L) to form a coexisting crayfish assemblage with a population density of 48 individuals m⁻². As soon as an animal died, it was replaced to keep the population density stable. Dead animals were frozen and kept for subsequent verification of A. astaci by molecular diagnostics. Fine gravel was used as substratum in the aguarium. No crayfish shelter was offered in order to increase stress and thereby weaken of the immune system. We used regular tap water with the following mean measured water quality values: Sodium = 3.2 mg L⁻¹, potassium = 2.1 mg L⁻¹, calcium = 27.5 mg L^{-1} , magnesium = 6.9 mg L^{-1} , aluminium = 0.020 mg L^{-1} , iron = 0.030 mg L^{-1} , ammonium = 0.07 mg L⁻¹, nitrate = 2.0 mg L⁻¹, nitrite \leq 0.05 mg L⁻¹, chloride = 6.3 mg L^{-1} , sulfate = 15.8 mg L^{-1} , total organic carbon < 1 mg L^{-1} . Water temperature $(18 - 25^{\circ}\text{C})$, oxygen content $(7.0 - 9.6 \text{ mg L}^{-1})$ and pH (7.7 - 8.4) were monitored over the whole study period and were each in the physiological tolerance range of both crayfish species and the pathogen A. astaci.

RESULTS

Molecular Diagnostics

From the total of 523 crayfish (490 spiny-cheek crayfish, 33 noble crayfish) that were tested from the nine localities with coexisting populations, the qPCR diagnostic detected no trace of *A. astaci* in any tissue sample (Table 1). About 56% of the crayfish were tested for three different tissues, while the remaining 44% were tested for the soft abdominal cuticle only. Hence, a total of 1338 crayfish tissue samples were all negative for the coexisting populations. On the contrary, positive status was confirmed in about 66% of the American crayfish tested from single species lakes and a stretch of a river (Table 2). Three out of five spiny-cheek crayfish individuals from Plociowe, where a relatively high noble crayfish mortality was observed in the transmission experiment, were positive (observed prevalence 60%), and all six individuals from Hennweiler were positive (observed prevalence 100%). Furthermore, the pathogen was found in 12 of the 21 signal crayfish tested from Heimbach (observed prevalence 57%). The noble crayfish from the crayfish farm, serving as negative controls, were indeed negative.

Table 1 Number of crayfish per coexisting population that remained negative for *Aphanomyces astaci* presence by real-time PCR. "-": no data available. The geographical coordinates of the study sites can be obtained from the corresponding author and have been made available to the chief-editor of *Freshwater Biology*.

Sampling site Country		Sample Set	A. astacus (number)	O. limosus (number)	Total	
Bielener lake*	GER	Α	13	91	104	
Heisteberger Weiher	GER	В	-	51	51	
Mastholter lake*	GER	Α	5	13	18	
Milaczcewo*	PO	Α	2	34	36	
Piasek*	PO	15xA/95xB	3	107	110	
Steinbruch Roth	GER	В	-	92	92	
Lake Stosinko wielkie	PO	В	-	30	30	
Trzcinskie*	PO	Α	8	16	24	
Zielone*	PO	34xA/24xB	2	56	58	
Total			33	490	523	

^{*} American crayfish from these coexisting populations were used in transmission experiment with farmed noble crayfish. Tough, the individual from the transmission experiment are not included in this table.

Table 2 Number of crayfish per reference population tested for *Aphanomyces astaci* presence and according agent level by *A. astaci* specific qPCR. "-": no data available.

Code ^a	Sample Set	Crayfish	Total	Positives (number)	Positives (%)	Agent level ^b				
		(number)				A0	A 1	A2	А3	A4
		A. astacus								
Cf	Α	8	8	-	-	8	0	0	0	0
		O. limosus								
PI^c	Α	5	5	3	60	2	0	1	1	1
Hw	В	6	6	6	100	0	0	0	4	2
		P. lenisuculus	;							
Не	В	21	21	12	57	9	0	3	8	1

^a Cf = Crayfish farm (Augsburg), Hw = Hennweiler, He = Heimbach (all Germany), PI = Plociowe (Poland). Hw, He and PI = single species lakes.

Transmission Experiment

The molecular results were supported by the transmission experiment. A behavioural change (lack of coordination, loss of escape reflex) in noble crayfish indicating crayfish plague infection was observed in the aquarium stocked with spiny-cheek crayfish from the single species lake Plociowe (Table 3). The proportion of dead noble crayfish was relatively high, with 0.07 dead individuals d¹. In the six aquaria housing spiny-cheek crayfish that originated from the six coexisting populations the noble crayfish mortality was lower (in three aquaria 0.02 d¹ and in the other three aquaria 0.03 d¹) (Table 3). The qPCR analysis demonstrated that noble crayfish kept in an aquarium with spiny-cheek crayfish from the sampling site Plociowe were infected by *A. astaci*, while similar tests remained negative for the other noble crayfish that had been exposed to spiny-cheek crayfish from the co-existing populations.

b Agent levels refer to semi-quantitative categories based on the numbers of observed PCR forming units (PFU_{obs}) from the *A. astaci*–specific real-time PCR. Agent Level 0 (A0): negative; (A1): below the limit of detection (PFU_{obs} < 5); A2: $5 \le PFU_{obs} < 50$; A3: $50 \le PFU_{obs} < 103$; A4: $103 \le PFU_{obs} < 104$ (according to Vrålstad *et al.*, 2009).

^c Spiny-cheek crayfish from the Plociowe site used in transmission experiment with farmed noble crayfish.

Table 3 Number of dead *Astacus astacus* during the experiment duration (63 to 91 days) in transmission experiments with *Orconectes limosus* from the studied sampling sites in Germany (GER) and Poland (PO).

Sampling site	Country	Coexistence	Dead <i>A. astacus</i> (Individuals/day)	Observed behavioral change
Bielener lake	GER	+	0.02	-
Mastholter lake	GER	+	0.03	-
Milaczewo	PO	+	0.02	-
Piasek	PO	+	0.03	-
Trzcinskie	PO	+	0.03	-
Zielone	PO	+	0.02	-
Plociowe	РО	-	0.07	+

Probability of Absence of the Pathogen

Calculations of the required number of crayfish individuals analysed to estimate the probability of absence of A. astaci are listed in Table 4. The number strongly depends on the assumed prevalence and test sensitivity, while population size affects these numbers to a lower extent, at least if N is high. With the highest tested estimates for prevalence (80% positives), sensitivity (0.8), and population size $(N = 10^6)$, positives will be detected if four individuals are tested (98% confidence level). With the lowest tested estimates for prevalence (1% positives) and test sensitivity (0.3), but the highest population size (N = 10^6), ~1300 individuals must remain negative before the pathogen can be declared absent with 98% confidence. The numbers tested in the current study vary with locations. If we assume 80% test sensitivity, comparable to results in Vrålstad et al. (2011), and 10% as the lowest probable prevalence of A. astaci positive individuals in an American crayfish population, five of the nine coexisting spinycheek crayfish and noble crayfish populations tested in this study (i.e.≥ 47 tested crayfish) (Table 1) can be declared pathogen free with 98% confidence (Table 4). Further, if A. astaci was present, but undetected, in any of the tested coexisting populations (Table 1), the prevalence would in all cases be less than 30% (i.e.≥ 14 tested crayfish required to detect one positive at 98% confidence level; Table 4).

Table 4 The required sample sizes, calculated from the epi.detectsize function in the epiR package in R, to confirm absence of disease/negative carrier status at a given confidence level (98% and 95%) given all samples are negative.

Test parameters ^a		Samples no	This study ^c			
Prevalence	Sensitivity	$N = 10^3$	$N = 10^4$	$N = 10^5$	$N = 10^6$	
0.80	8.0	4/3	4/3	4/3	4/3	9/9
0.30	8.0	14/11	14/11	14/11	14/11	9/9
0.1	8.0	44/34	47/36	47/36	47/36	5/5
0.01	8.0	278/237	453/353	484/371	487/373	-
0.80	0.5	8/6	8/6	8/6	8/6	9/9
0.30	0.5	23/18	24/18	24/18	24/18	7/7
0.1	0.5	68/54	76/58	76/58	76/58	3/3
0.01	0.5	351/310	699/548	771/592	779/597	-
0.80	0.3	14/11	14/11	14/11	14/11	8/9
0.30	0.3	39/30	41/32	42/32	42/32	5/6
0.1	0.3	107/86	126/97	128/98	128/98	0/1
0.01	0.3	421/387	1088/867	1277/982	1299/996	-

^a Test parameters in the analysis: prevalence tested from high (0.8), moderate (0.3), low (0.1) to very low (0.01). Sensitivity was tested for high (0.8), medium (0.5) and low (0.3) test sensitivity. Test specificity was set to 99%. Reduced test specificity (down to 50%) did not affect the number of individuals needed.

DISCUSSION

Pathogen free Coexisting Populations?

On the basis of tissue analyses with molecular diagnostics (qPCR) combined with traditional transmission experiments we have demonstrated a consistent lack of *A. astaci* positive crayfish individuals in nine coexisting populations of American and European crayfish species in central Europe. To our knowledge, only a few lakes harbouring coexisting American and European crayfish have been recorded (e.g. Westman & Savolainen, 2001; Söderbäck 1994; Schulz *et al.*, 2006), and this study is the first to investigate thoroughly the *A. astaci* carrier status in such populations. Many studies have shown that the introduction of American crayfish species is followed by the mass mortality of

^b The test was performed with four different hypothetical population sizes (N = 103, 104, 105, 106) for two different confidence levels (0.98 and 0.95).

^c Numbers of coexisting populations out of the nine tested in this study that are demonstrated free of *A. astaci* in spiny-cheek crayfish at given test parameters and confidence levels (0.98 and 0.95). Analysed individuals of noble crayfish are not included in the calculations to avoid assumptions of "equal prevalence" in both species.

susceptible European crayfish species due to transmission of the crayfish plague (e.g. Bohman *et al.*, 2006; Kozubíková *et al.*, 2008; Vrålstad *et al.*, 2011). Some of the key factors that could explain prolonged coexistence could therefore involve either evolved resistance in native crayfish, reduced virulence in the pathogen strain, or absence of the pathogen.

To our knowledge, there has been no demonstration of increased resistance in native European crayfish populations, although recent results do indicate that persistent infections of A. astaci in native European crayfish occur (Jussila et al., 2011; Viljamaa-Dirks et al., 2011; Kokko et al., 2012, Svoboda et al., 2012; Schrimpf et al., 2012). Whether this is due to increased crayfish immunity, or to reduced pathogen virulence, remains to be clarified in most cases. For our study, however, the absence of the pathogen is strongly supported. Only when we assume that a very low prevalence of A. astaci in the population is possible (\leq 5%), combined with low test sensitivity, the sample sizes required to confirm absence of the pathogen increase to unfeasible numbers. Hence, absence, or an extremely low prevalence of the crayfish plague pathogen in these populations stands out as the most likely factor for the observed coexistence.

It is a challenge to test sufficient individuals to exclude the possibility of a very low prevalence of *A. astaci*. As an explanation why some American crayfish populations are *A. astaci*-free it is conceivable that a few pathogen free animals actively immigrated into the coexisting populations, or were passively introduced. Alternatively, environmental or biological changes or factors could have eradicated *A. astaci* from the crayfish populations. Strand *et al.* (2012) recently demonstrated that *A. astaci* qPCR, used directly on water samples from tanks with *A. astaci* positive signal crayfish, is able to detect the pathogen spore content. They found that *A. astaci* spores were continuously released into the water from the carrier crayfish with raised spore levels before and after moulting and mortalities. Hence, screening a large number of individuals could involve analyses of water containing many crayfish. The method detects down to a single spore in the water sample (Strand *et al.*, 2011; 2012). It could therefore strengthen our hypothesis of pathogen absence if a large number of individuals from coexisting

populations were kept in a common tank over time without releasing a detectable number of *A. astaci* spores into the water.

Pathogen Prevalence

Before results from molecular screenings of American crayfish populations became available, it was generally assumed and expected that all American crayfish are carriers of *A. astaci* (Unestam, 1972; Oidtmann *et al.*, 2006, OIE, 2009). Many more recent studies have investigated the carrier status of American crayfish. However, the crayfish number and tissue types tested vary greatly, making comparisons difficult. Highly variable prevalence has been reported using the Vrålstad *et al.* (2009) qPCR method, from 0-100% in *O. limosus* and 0-37% in *P. leniusculus* (Kozubíková *et al.*, 2011b), but here only soft abdominal cuticle was analysed in a variable number of individuals per population.

Three comparable studies testing 20-40 crayfish individuals per population and three tissues per crayfish (cuticle, uropods and walking legs) also demonstrated extremely variable prevalence. In one of the studies, no positive individuals were found in populations of signal crayfish and noble crayfish coexisting in the same river in Denmark (Skov et al., 2011), while in the other studies, the observed prevalence of *A. astaci* in a signal crayfish populations were 86%, 90% and 100% for one Norwegian and two Finnish populations, respectively (Vrålstad et al., 2011; Strand et al., 2012). In the Norwegian case, the high prevalence explained the recent crayfish plague outbreaks in native noble crayfish populations in this watercourse (Vrålstad et al., 2011). In our study, the prevalence of spiny-cheek crayfish positive for A. astaci in the populations from Plociowe and Hennweiler was 60% and 100%, respectively. The lower prevalence limit for an obligate crayfish parasite like *A. astaci* remains unknown. It has been shown that the pathogen prevalence in a weakly infected population decreases from April to November, whereas the prevalence in a highly infected population remains high over several years and during different seasons (Matasová et al., 2011).

Supplementary Transmission Experiment

The classical transmission experiment conducted for all six populations tested out of the total of nine coexisting populations revealed no indication of an infection with A. astaci, and supported thereby the molecular analyses. The power of the experiment was confirmed when noble crayfish was kept with A. astaci positive spiny-cheek crayfish from a single population from Plociowe. In this aquarium, the noble crayfish mortality was about three times higher than in the stock with crayfish from coexisting populations. The increased mortality serves as a strong sign of disease transmission (Matthews & Reynolds, 1990; Oidtmann et al., 2006) from infected spiny-cheek crayfish to uninfected noble crayfish. Noble crayfish from this aquarium indeed were infected at the end of the experiment. The results of the transmission experiments therefore agree with the qPCR results. However, the long duration of transmission experiments is a disadvantage, particularly if further conservation measures depend on the confirmed infection status with A. astaci. Consequently, the fast qPCR verification of a positive case makes the time-consuming transmission experiments redundant. On the other hand, for the verification of pathogen absence, the transmission experiment turned out as a valuable supplementary test. Since a wrong negative result can have fatal consequences for native crayfish, a second independent verification is indeed strongly recommended.

Conservation and Management Considerations

Knowledge about the presence of the crayfish plague can influence conservation activities and actions (Kozubíková *et al.*, 2009). In Europe, many indigenous crayfish populations have succumbed due to this disease (e.g. Alderman, 1996). The pathogen poses a severe threat to the noble crayfish, and if a highly virulent *A. astaci* genotype infects the coexisting populations all noble crayfish will die within a few weeks. On the other hand, infection by the less virulent As-genotype might be less harmful (Viljamaa-Dirks *et al.*, 2011). An important outcome of this study is that European crayfish populations are not necessarily lost, at least not in the short run if American crayfish occur in the same water. In the long run, however, competitive exclusion of native crayfish is a likely outcome (see below). However, since predators, fishing gear and any item that has been in contact with contaminated water, may transmit the pathogen

between waterbodies (OIE, 2009), the risk of disease transmission into pathogenfree waters, and thus also into the coexisting population is great unless measures are taken to prevent infection from outside. At the same time, we cannot be 100% sure that these populations are pathogen free, as there is still a 2% probability of positive carriers in the populations (given the test assumptions made here). Hence, coexisting populations should not be considered as fully safe regarding further transmission risk originating from these populations.

Non-infected European crayfish can be translocated to refuge localities like "ark sites" to conserve intraspecific diversity (Kozák *et al.*, 2011; Peay, 2009). Such crayfish should preferably originate from single species lakes of native crayfish. However, native crayfish from well documented plague-free coexisting populations may also be considered worth translocating into ark sites depending on their genetic composition and uniqueness. This is particularly important since permanent coexistence, even in the absence of crayfish plague, seems unlikely in the long run, due to other factors such as competitive exclusion by the alien crayfish, as has been shown for coexistence between noble crayfish and spiny-cheek crayfish (Schulz *et al.*, 2006) or noble crayfish and signal crayfish (Söderbäck, 1994; Westman *et al.*, 2002).

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ERKLÄRUNG

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Landau, 16. Januar 2013

Anne Schrimpf

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English

- Schrimpf A, Maiwald T, Vrålstad T, Schulz HK, Smietána P, Schulz R (in press) Absence of the crayfish plague pathogen (*Aphanomyces astaci*) facilitates coexistence of European and American crayfish in central Europe. Freshwater Biology
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Presentations

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