

Interference competition between alien invasive gammaridean species

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Abstract The relative abundances of gammaridean species in the river Rhine have profoundly changed since the invasion of *Dikerogammarus villosus* in 1994/1995. This study tested whether these changes in gammaridean dominance could have been determined by interspecific competition and unequal mortality, for example by intraguild predation (IGP). Single and two species tests have been carried out in aquariums provided with all substrata present in the main channel of the Rhine. Changes in substratum choice, increased swimming activity and increased mortality of a species were used as indicators of interspecific competition during interaction between gammaridean species. Interspecific competition and mortality between the most abundant invasive gammaridean species in the Rhine, viz. *Gammarus tigrinus*, *Echinogammarus ischnus* and

Dikerogammarus villosus were tested. In single-species experiments, *G. tigrinus* and *D. villosus* showed similar preferences for a stony substratum, whereas *E. ischnus* mostly occupied the water column. The two-species aquarium experiments indicated direct interference competition for substratum and unequal mortality between *G. tigrinus* and *D. villosus*, with *D. villosus* being the stronger competitor. Competitive stress was influenced by population density, was size-dependent and varied between the different types of substratum due to substratum choice. *G. tigrinus* did not show any behaviour indicative of interference competition in the presence of *E. ischnus*, and neither did *E. ischnus* or *D. villosus* in the presence of any of the other gammarideans. Swimming in the water layer may already enable *E. ischnus* to minimise its encounters with the stone-dwelling *D. villosus* and *G. tigrinus*. To maximise the encounters between *E. ischnus* and *D. villosus*, a fish (*Lepomis gibbosus*) was added to occupy the water layer during the aquarium experiments. *E. ischnus* showed a higher mortality in the presence of both *D. villosus* and fish, probably due to increased stress, as shelter opportunities to escape the predators had been minimised. The study shows that interference competition between gammaridean species can explain the replacement of the North American invader *G. tigrinus* by *D. villosus* in the river Rhine. *E. ischnus* and *D. villosus* both Ponto-Caspian invaders did not show interference competition in our experiments and co-exist in the Rhine.

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Introduction

Each river basin traditionally harbours a characteristic fauna, with species that remain endemic due to biogeographical barriers isolating the populations in the watersheds. Lifting these barriers by constructing waterways that link river systems leads to mixing of species and to new interactions between invasive and native species in the food web (Bij de Vaate et al. 2002; Van der Velde et al. 2002). Gammarideans have expanded their distribution all over Europe, enabled by the high connectivity level of Europe's large rivers and by intentional introductions, thus increasing their encounters (Bij de Vaate et al. 2002).

Invasions by species related to native species, as has occurred with gammarideans, can have a profound effect on a food web, as closely related species often occupy similar niches. These species can coexist under various circumstances, if resources are sufficiently available, population densities remain low (Van Overdijk et al. 2003; Van Riel et al. 2006a), the species are equally strong competitors or one of them is able to adjust its niche to avoid the other species in space or time. Limited availability of shared resources, which is most likely to occur during the population growth peak of an invasion, can result in interspecific competition. This can alter the functioning of food webs (Van der Velde et al. 2006) as it can have a huge impact on community structure.

Competition is most likely when species overlap completely in their needs for limited resources, and may result in species being replaced (Reynoldson and Bellamy 1970). The outcome of interspecific competition is not only determined by the limiting resource itself but also influenced by behavioural, physiological and morphological traits of the species involved, in relation to environmental conditions and anthropogenic disturbance (Dick et al. 1993; Carlton 1996; Moyle and Light 1996; Williamson and Fitter 1996a, b; Wisheu 1998; Van der Velde et al. 2002, 2006; Wijnhoven et al. 2003).

Competition mechanisms vary, as species can interact in different ways. Examples are pre-emptive competition, in which individuals occupy a unit of space and inhibit its occupation by others, and consumptive competition, in which a shared resource is being consumed. More direct competition is shown in interference competition, in which species harm one another by aggressive interaction while actively defending a unit of space (territorial competition), or by competitive interactions upon encounters (encounter competition) (Schoener 1983). In this paper, we use the term competition more broadly, defining it as a situation in which species use overlapping resources and may thus compete, regardless of differences in tactics of resource acquisition. An extreme form of encounter competition is intraguild predation (IGP), i.e., the killing and eating of species that use similar resources and are thus potential competitors (Polis et al. 1989), which has frequently been observed for gammarideans (Dick 1996; Dick et al. 1993; MacNeil et al. 2003). IGP may affect the distribution, population size, stability and resilience of species, reduce potential competition and promote the occurrence of alternative stable states (Polis et al. 1989; Dick and Platvoet 2000; Dick et al. 2002). Species suffering from interspecific competition may develop ways to escape this stress, for instance by avoiding encounters by reproducing at different times or by shifting their spatial niche.

Gammarideans stressed by interspecific competition increase their activity levels in terms of swimming, and seeking shelter (Garvey et al. 1994), and may eventually change their habitat utilisation. Competition for shelter can be size-mediated, giving the larger species an advantage in the competition for the larger shelter places and thus driving the competitively weaker species to smaller shelter areas. Competitively weaker species may be indirectly eliminated through predation by top predators (Garvey et al. 1994). Changes and replacements in gammaridean communities due to invasions by relatives have been frequently recorded (Pinkster et al. 1992; Dick and Platvoet 2000; Bollache et al. 2004; Jazdzewski et al. 2004; Meyer et al. 2005) and may have been determined by interspecific competition and IGP (Schoener 1983; MacNeil and Prenter 2000).

To the most successful invaders in the river Rhine belong amphipod crustaceans, which became dominant

in numbers and biomass within a very short time and have a huge impact on the Rhine food web (Van der Velde et al. 2000, 2002; Haas et al. 2002; Leuven et al. 2009). The indigenous *Gammarus pulex* strongly decreased in numbers after the North-American species *Gammarus tigrinus* became abundant in 1982 and remained abundant for decades till the appearance of the Ponto-Caspian *Dikerogammarus villosus* (Van der Velde et al. 2000). In 1987, the Ponto-Caspian *Chelicorophium curvispinum* entered the Rhine through the Mittelland Canal in Germany and became dominant, covering the hard substrata with its muddy tubes. *Echinogammarus ischnus* also made its way through the Mittelland Canal to the Rhine in 1989. Since 1992, the Rhine has been connected to the Danube by the Main-Danube canal, which facilitates invasions of the Rhine by Ponto-Caspian species (Bij de Vaate et al. 2002). The most recent mass invader, *Dikerogammarus villosus* is currently the largest amphipod species in the Rhine and is a strong, omnivorous predator (Dick and Platvoet 2000; Van der Velde et al. 2000, 2002; Dick et al. 2002; Van Riel et al. 2006b). Other alien or native gammarideans inhabiting the Rhine have remained present in low numbers, as is the case with *E. ischnus* and *D. haemobaphes*, both Ponto-Caspian invaders, or are found occasionally at times of high discharge, probably after being flushed out of the tributaries, as is the case with the native *G. pulex*, the probably early Ponto-Caspian invader *G. roeseli* and the North American *Crangonyx pseudogracilis* (Bij de Vaate and Klink 1995; Bij de Vaate et al. 2006).

In an earlier paper interspecific competition between native and invader gammaridean species occurring in the Rhine is described (Van Riel et al. 2007). The present study investigated various situations of interspecific competition between the three most abundant invasive gammaridean species occurring in the Rhine, to answer the following questions:

- (a) Could interspecific competition have determined the varying success of the successive invading gammarideans *Gammarus tigrinus*, *Echinogammarus ischnus* and *Dikerogammarus villosus*?
- (b) Are interactions between invasive gammarideans determined by interspecific competition, as indicated by mortality, swimming behaviour and substratum choice shifts?

- (c) Could the presence of a fish predator change the above-mentioned indicators of competition between gammaridean species?
- (d) Is it likely that the limited habitat heterogeneity in the Rhine's main channel influence interspecific competition?

Materials and methods

Study site characteristics and invasion history

The Rhine is a large river ecosystem under various forms of anthropogenic stress, such as water pollution and salination (Van der Velde et al. 1990, 2006; Admiraal et al. 1993; Bij de Vaate et al. 2006). The floodplain is embanked with dikes and the main channel has been canalised, reducing its heterogeneity to two main biotopes, i.e., sandy streambeds and stony riverbanks and groynes (Admiraal et al. 1993). Water quality has improved lately by sanitation, but rehabilitation of the native fauna seems to be inhibited by the large numbers of invasive species present in the Rhine (Van der Velde et al. 2000, 2002).

Field studies

Biomonitoring has been carried out in the river Waal, a Rhine distributary, in the vicinity of the Dutch town of Nijmegen (5°48'E; 51°52'N), by sampling during the years 1992, 1993 and 2001. Macroinvertebrates were sampled monthly by sampling six groyne stones from a water depth of 2 m by means of a polyp-grab operated with a hydraulic crane from a ship. Macroinvertebrates were cautiously brushed off the stones, collected and preserved in 70% ethanol, subsequently sorted by species, and counted in the laboratory using a stereomicroscope. The total surface area of the stones was calculated from the areas of all sides of the stones, in order to quantify the densities of macroinvertebrate species per square metre of substratum.

Aquarium experiments

Aquarium experiments in the laboratory were used to study possible IGP and interspecific competition for

substratum between the exotic gammaridean species *D. villosus*, *G. tigrinus* and *E. ischnus*. Interspecific predation, shifts in substratum choice and altered swimming behaviour after the addition of another species were assumed to indicate interspecific competition or IGP. The substrata used in these experiments reflected those occurring in the Rhine and provided the species with opportunities for shelter to various degrees. A series of experiments was carried out to study different situations of competition and to find out under what conditions gammarideans show interspecific competition and whether interspecific competition could be intensified by increasing densities or by introducing a top predator such as a fish. Although several fish species occurring in the Rhine have been identified as feeding on gammarideans (Kelleher et al. 1998, 2000) they could not be held in our aquaria for several reasons. Therefore, in this study, the pumpkinseed sunfish (*Lepomis gibbosus*) was used as the top predator because of its small size, its predatory behaviour with respect to macroinvertebrates including gammarideans (Van Kleef et al. 2008) and because this fish could be easily obtained from the aquarium shop. It is not occurring in the Rhine but functions in our experiments as a representative of any predatory fish.

Adult *D. villosus* (DV) and *E. ischnus* (EIS) were collected from stone substrata in the river Waal near Nijmegen (5°48'E, 51°51'N). *G. tigrinus* (GT) was collected from Lake IJsselmeer (5°32'E, 52°35'N). All specimens were kept separately, in aerated basins (40 × 40 × 50 cm) at 15°C, with a 9/15 h dark/light regime, before being released into the experimental aquaria. The gammarideans were fed dead frozen chironomids during captivity.

The experiments were carried out in a climate room at 15°C with a 9/15 h dark/light regime (2 × 36 W/840TLD lamps). Aquaria (25 × 25 × 30 cm) were filled with Rhine water and aerated. Four different types of Rhine substratum, i.e., one groyne stone (Ø 14.4 cm ± 1.5), pebbles (Ø 5.4 cm ± 1.2), gravel (Ø 2.5 cm ± 1.0) and sand, were put into cups (Ø 11.5, 6.5 cm height), which were randomly placed in each aquarium. In single-species tests, 50 adult individuals of the same species (50 GT ($n = 10$), 50 EIS ($n = 14$) or 50 DV ($n = 14$)) were allowed to choose between substrata. After 24 h, the cups with substrata were collected and the gammarideans inside each cup were counted and their body lengths

measured. The gammarideans swimming in the water column were counted and measured as well and are referred to as 'free' in the figures. The substratum containing the largest number of specimens was considered to be the preferred substratum. In addition, the numbers of specimens that had died (undamaged dead individuals) or had been consumed (only remnants of individuals found) were counted. The experiment was repeated for three combinations: *G. tigrinus* with *D. villosus*, *E. ischnus* with *D. villosus* and *G. tigrinus* with *E. ischnus*. In these two-species experiments, specimens of *G. tigrinus* or *E. ischnus* were allowed to hide for 2 h before *D. villosus* were added. The following nine density combinations were tested: 50 GT + 25 DV ($n = 10$), 25 GT + 25 DV ($n = 10$), 25 GT + 50 DV ($n = 10$), 25 GT + 25 EIS ($n = 10$), 50 GT + 50 EIS ($n = 2$), 25 EIS + 25 DV ($n = 14$), 50 EIS + 1 fish ($n = 14$), 50 DV + 1 fish ($n = 14$), 25 EIS + 25 DV + 1 fish ($n = 12$). Each experiment was repeated four times, using new individuals from the stock populations.

Echinogammarus ischnus (mean length = 1.07 cm, SD = 0.22) used in the single-species experiments and in the two-species experiments with *D. villosus* had been collected in spring (April 2001) and were significantly smaller ($P < 0.000$, t -test) than the *E. ischnus* (mean length = 0.81 cm, SD = 0.25) used in the two-species experiments with *G. tigrinus*, which had been collected in summer (August 2001). The mean body length of *G. tigrinus* used in the different experiments was similar (length = 0.71 cm, SD = 0.20) ($P > 0.05$, t -test), just as for *D. villosus* (mean length 1.15 cm, SD = 0.23) ($P > 0.05$, t -test).

Statistical analysis

The numbers of surviving specimens present on the various substrata and in the water column were analyzed for substratum choice patterns. Shifts in these patterns in the two-species experiments compared to the single-species experiments were assumed to indicate interspecific interactions. Differences in substratum choice patterns of the gammaridean species under different levels of competitive stress, caused by varying densities in the aquariums, were tested using a 2-way Generalised Linear Model for Poisson distribution (SAS 8.0). The substratum choice patterns that were derived from the surviving specimens present on the substratum types were

analysed. A Poisson distribution was used in the model because the data were based on counts. A Games-Howell post hoc test (SPSS 11.5) was used to analyse any significant preference for any of the substratum types by a gammaridean species in an experiment.

The substratum choice patterns in the different aquarium experiments were clustered using Euclidean distances. The clustering was used to interpret similarities between the substratum choice patterns shown in the different aquarium experiments and not as statistical proof. Differences in body length between the gammarideans present on the various substrata within a specific treatment condition were tested in the aquarium experiments using a one-way ANOVA and a Games-Howell post hoc test on log-transformed data (SPSS 11.5). Differences in body length between individuals of species used in various aquarium experiments were tested using *t*-tests (SPSS 11.5). Intraspecific and interspecific differences in the substratum choice were tested using a paired *t*-test. Mortality rates of the specimens during the substratum choice experiments were compared using a one-way ANOVA with a Games-Howell post hoc test.

One-way ANOVA was also used to test the influence of different densities of prey and predators on intraguild predation between *D. villosus* and *G. tigrinus*. Differences between the two species in the numbers of specimens consumed were tested using a paired *t*-test.

Results

Field studies

Field observations show that the relative abundances of gammarideans in the main channel of the river Waal changed profoundly after the invasion by *D. villosus* (Fig. 1). Whereas *G. tigrinus* dominated the gammaridean community during 1992 and 1993, *D. villosus* had outnumbered *G. tigrinus* in 2001, showing similar densities as *G. tigrinus* had before. In all of the years during this study, *E. ischnus* showed low densities, but the lowest densities were found during the years when *D. villosus* was present in large numbers. During these years, water temperature was similar but chlorophyll-*a* availability decreased, as

did the abundance of *C. curvispinum*. Despite this decrease, *C. curvispinum* still dominated the macro-invertebrate fauna on the stones in terms of numbers, with population peaks of around 40,000 specimens per square metre of substratum (Fig. 1).

Aquarium studies

In single-species tests, *G. tigrinus* and *D. villosus* preferred pebbles, whereas *E. ischnus* preferred swimming in the water layer rather than occupying the stone substratum (Table 1). All species disliked sand as a substratum (Fig. 2; Table 2). Changes in substratum choice patterns of a species in the presence of other species were observed in the two-species experiments (Tables 1, 3; Fig. 3) and were influenced by substratum type and by the presence of the other gammaridean species (Table 3). Such changes in substratum choice due to the presence of another gammaridean species occurred in most cases (Table 3, significant substratum \times competition interaction). In the case of *G. tigrinus* in the presence of *E. ischnus* (2-way GLM, $P = 0.87$), *D. villosus* in the presence of *E. ischnus* (2-way GLM, $P = 0.32$) and *E. ischnus* in the presence of *D. villosus* (2-way GLM, $P = 0.89$) the data as they show no significant shift are not presented. The largest shifts in substratum choice were observed for *G. tigrinus* in the presence of higher densities of *D. villosus* and of higher densities of *E. ischnus*, for *E. ischnus* in the presence of *G. tigrinus*, and for *D. villosus* in the presence of *E. ischnus* together with a fish (Tables 1, 2; Figs. 4, 5).

In the presence of *D. villosus*, *G. tigrinus* shifted from pebbles to gravel and sand. *D. villosus* remained most abundant on pebbles once it had established itself, showing no marked shifts in substratum occupation (Tables 1, 2, Fig. 4). It was particularly the smaller specimens of *G. tigrinus* (Fig. 6) which were found hiding in gravel and digging themselves into sand for shelter (1-way ANOVA, $P < 0.001$; Games-Howell $P < 0.001$ for body length of specimens on sand and gravel). A high level of predation on *G. tigrinus* by *D. villosus* occurred when the species were put together (Table 1; Fig. 7). Predation on *G. tigrinus* was greatest when the density of *G. tigrinus* was twice that of *D. villosus*. Predation on *D. villosus* was greatest in the single-species tests (Fig. 7), indicating cannibalism.

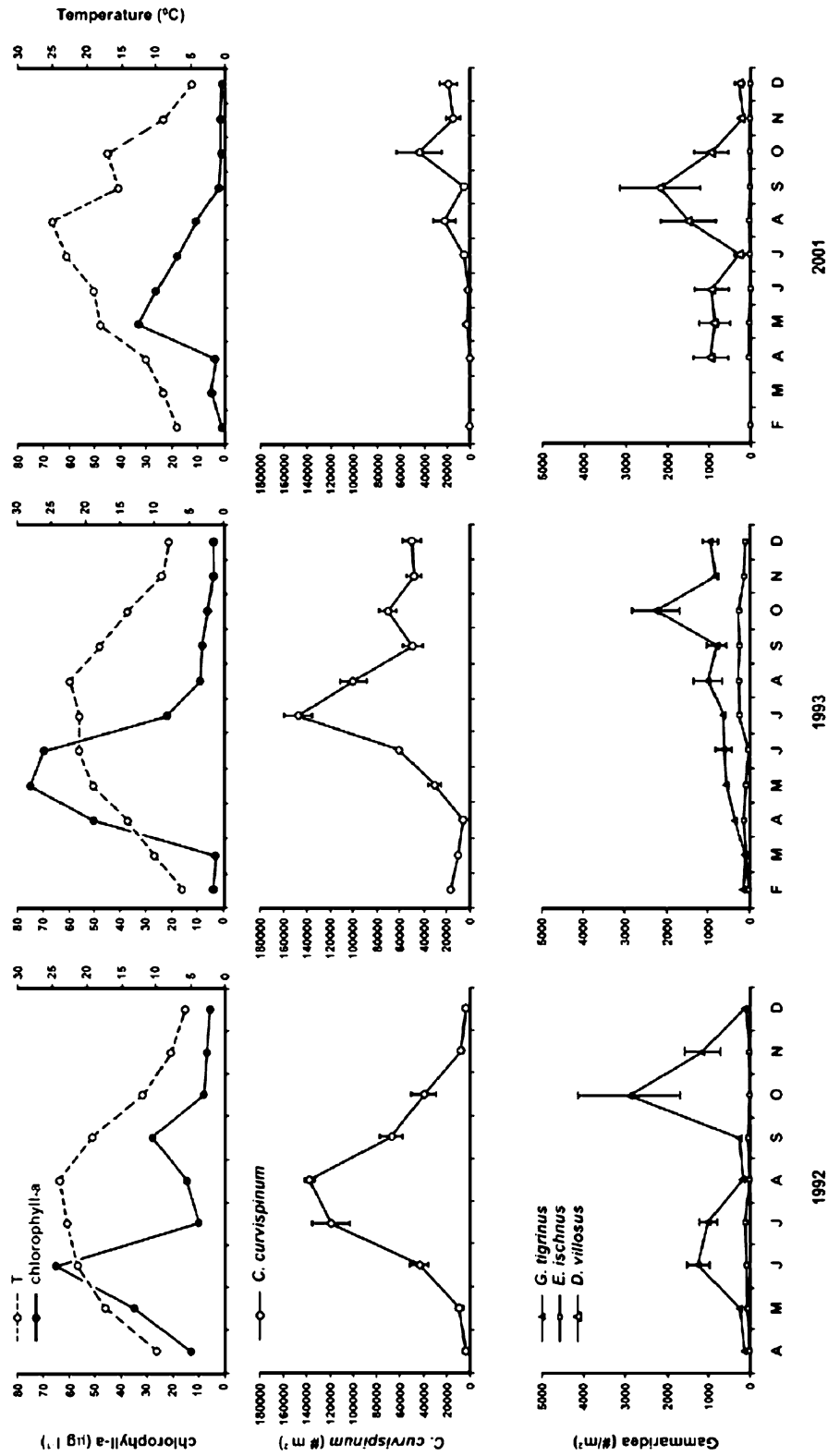


Fig. 1 Water temperatures, concentrations of chlorophyll-*a* in the water column and densities (mean \pm SEM) of *Chelicorophium curvispinum* and gammaridean species on the stone substratum of the main stream of the river Rhine during 1992, 1993 and 2001

Table 1 Substratum choice patterns and mortality (mean% \pm SEM) of *Gammarus tigrinus* (GT), *Echinogammarus ischnus* (EIS) and *Dikerogammarus villosus* (DV) in the various experiments

DV + invaders	DV alone	DV > GT	DV = GT	DV < GT				
Groyne stone	13.9 \pm 2.3	21.6 \pm 4.9	17.6 \pm 5.1	23.6 \pm 5.7				
Pebbles	32.6 \pm 3.0	44.6 \pm 6.7	35.6 \pm 8.1	28.8 \pm 5.4				
Gravel	7.1 \pm 1.3	5.2 \pm 1.5	2.4 \pm 1.3	4.0 \pm 1.3				
Sand	2.0 \pm 1.0	0.4 \pm 0.4	0.4 \pm 0.4	0.0 \pm 0.0				
Free	38.8 \pm 3.2	25.0 \pm 4.4	32.4 \pm 4.7	36.4 \pm 5.6				
Dead	5.6 \pm 1.5	1.2 \pm 0.6	4.0 \pm 1.4	1.2 \pm 0.6				
Eaten	0 \pm 0.0	2.0 \pm 0.6	7.6 \pm 3.2	6.0 \pm 2.5				
GT + invaders	GT alone	GT > DV	GT = DV	GT < DV	GT alone	GT = EIS low	GT = EIS high	
Groyne stone	4.4 \pm 1.4	3.0 \pm 1.2	2.8 \pm 1.7	3.6 \pm 1.6	4.4 \pm 1.4	7.6 \pm 2.2	22.0 \pm 5.7	
Pebbles	28.0 \pm 4.5	10.6 \pm 1.7	11.2 \pm 3.2	10.0 \pm 2.8	28.0 \pm 4.5	28.8 \pm 4.4	28.0 \pm 8.5	
Gravel	23.4 \pm 4.4	20.0 \pm 2.7	12.0 \pm 3.5	25.2 \pm 4.5	23.4 \pm 4.4	26 \pm 5.5	12.0 \pm 8.5	
Sand	10.4 \pm 1.9	15.4 \pm 4.3	9.6 \pm 2.5	17.6 \pm 6.5	10.4 \pm 1.9	9.6 \pm 3.5	7.0 \pm 1.4	
Free	22.6 \pm 2.9	24.6 \pm 3.5	29.6 \pm 5.3	10.4 \pm 2.8	22.6 \pm 2.9	22.0 \pm 4.5	21.0 \pm 7.1	
Dead	7.4 \pm 1.2	6.4 \pm 1.5	9.2 \pm 1.6	6.4 \pm 4.0	7.4 \pm 1.2	3.2 \pm 1.4	0.0	
Eaten	3.8 \pm 1.4	20.0 \pm 2.0	25.6 \pm 4.7	26.8 \pm 4.9	3.8 \pm 1.4	2.8 \pm 1.3	10.0 \pm 2.8	
EIS + invaders	EIS alone	EIS = DV	EIS + fish	EIS = DV + fish	EIS alone	EIS = GT low	EIS = GT high	
Groyne stone	5.4 \pm 0.9	5.1 \pm 1.8	10.0 \pm 2.0	5.0 \pm 2.2	5.4 \pm 0.9	10.8 \pm 1.5	13.0 \pm 15.6	
Pebbles	17.7 \pm 2.9	13.1 \pm 4.5	15.7 \pm 1.5	25.0 \pm 4.2	17.7 \pm 2.9	32.0 \pm 3.7	24.0 \pm 0.0	
Gravel	4.9 \pm 1.5	1.1 \pm 0.8	4.0 \pm 1.0	6.0 \pm 2.3	4.9 \pm 1.5	21.2 \pm 0.7	18.0 \pm 11.3	
Sand	1.4 \pm 0.6	1.4 \pm 0.8	0.5 \pm 0.5	0.0	1.4 \pm 0.6	1.6 \pm 0.7	0.0 \pm 0.0	
Free	66.4 \pm 3.7	74.0 \pm 5.4	59.8 \pm 2.0	50.3 \pm 3.3	66.4 \pm 3.7	31.2 \pm 4.5	29.0 \pm 12.7	
Dead	4.14 \pm 0.9	5.1 \pm 2.5	10.0 \pm 1.3	13.7 \pm 1.3	4.14 \pm 0.9	3.2 \pm 2.1	0.0 \pm 0.0	
Eaten	0.0	0.0	0.0	0.0	0.0	0.0	16.0 \pm 8.5	
DV + invaders	DV alone	DV = EIS	DV + fish	DV = EIS + fish	DV alone	DV > GT	DV = GT	DV < GT
Groyne stone	13.9 \pm 2.3	20.0 \pm 4.4	11.8 \pm 1.8	27.7 \pm 2.7	13.9 \pm 2.3	21.6 \pm 4.9	17.6 \pm 5.1	23.6 \pm 5.7
Pebbles	32.6 \pm 3.0	25.4 \pm 3.6	23.2 \pm 4.6	4.7 \pm 1.5	32.6 \pm 3.0	44.6 \pm 6.7	35.6 \pm 8.1	28.8 \pm 5.4
Gravel	7.1 \pm 1.3	7.4 \pm 1.8	5.7 \pm 1.9	8.0 \pm 2.1	7.1 \pm 1.3	5.2 \pm 1.5	2.4 \pm 1.3	4.0 \pm 1.3
Sand	2.0 \pm 1.0	2.0 \pm 0.9	0.3 \pm 0.2	0.0	2.0 \pm 1.0	0.4 \pm 0.4	0.4 \pm 0.4	0.0 \pm 0.0
Free	38.8 \pm 3.2	37.7 \pm 4.6	55.0 \pm 2.8	56.3 \pm 2.6	38.8 \pm 3.2	25.0 \pm 4.4	32.4 \pm 4.7	36.4 \pm 5.6
Dead	5.6 \pm 1.5	7.4 \pm 1.4	4.0 \pm 0.5	3.3 \pm 0.9	5.6 \pm 1.5	1.2 \pm 0.6	4.0 \pm 1.4	1.2 \pm 0.6
Eaten	0.0 \pm 0.0	0.0 \pm 0.0	0.0 \pm 0.0	0.0 \pm 0.0	0 \pm 0.0	2.0 \pm 0.6	7.6 \pm 3.2	6.0 \pm 2.5

Preferred substrata are in italics

In the presence of *G. tigrinus*, *E. ischnus* was more abundant on the substratum than in the single-species tests, whereas *G. tigrinus* did not change its substratum choice pattern (Fig. 4; Tables 1, 2, 3). The species did not avoid each other but seemed to prefer the same substratum.

Neither *D. villosus* nor *E. ischnus* significantly changed their substratum choice in each other's

presence (Table 2; 3; Fig. 5), and mortality did not increase (Table 1; Fig. 5), making direct competition between these invaders unlikely. When a top predator such as a fish was present in an experiment with both *E. ischnus* and *D. villosus*, *E. ischnus* suffered a slightly higher mortality than *D. villosus* (*t*-test, $P = 0.18$) (Fig. 5), and both species significantly changed their substratum choice patterns (Tables 2, 3; Fig. 5).

Fig. 2 Substratum choice and mortality (mean \pm SEM) of *Gammarus tigrinus* (GT), *Dikerogammarus villosus* (DV) and *Echinogammarus ischnus* (EIS) in single species tests. Significant differences in a species' substratum choice are presented in Table 2

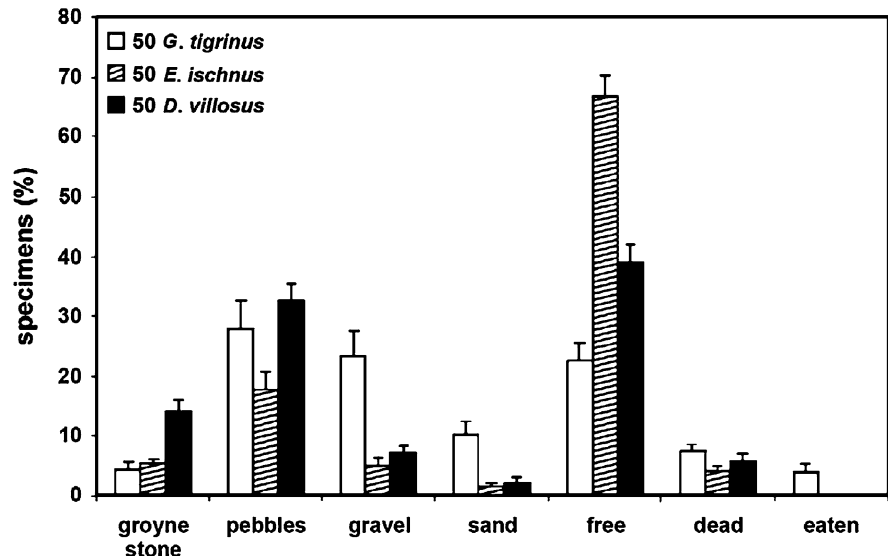


Table 2 Significant differences between substratum distribution patterns of gammarideans in aquarium studies (analysed by Games-Howell post hoc test)

Figure	Species	Experiment	Groyne stones	Pebbles	Gravel	Sand	Free
3	<i>G. tigrinus</i>	50GT	B	A	A	B	A
3	<i>E. ischnus</i>	50EIS	C	B	CD	D	A
3	<i>D. villosus</i>	50DV	B	A	B	C	A
4a	<i>G. tigrinus</i>	50GT	B	A	A	B	A
4b	<i>G. tigrinus</i>	25GT + 50DV	B	AB	A	AB	A
4a	<i>D. villosus</i>	50DV	B	A	B	C	A
4b	<i>D. villosus</i>	25DV + 50GT	A	A	B	B	A
4c	<i>G. tigrinus</i>	50GT	B	A	A	B	A
4c	<i>E. ischnus</i>	50EIS	C	B	CD	D	A
4d	<i>G. tigrinus</i>	25GT + 25EIS	B	A	A	B	A
4d	<i>E. ischnus</i>	25EIS + 25GT	C	A	C	D	B
6	<i>E. ischnus</i>	50EIS	C	B	CD	D	A
6	<i>E. ischnus</i>	25EIS + 25DV	BC	B	C	C	A
6	<i>E. ischnus</i>	50EIS + fish	BC	B	C	D	A
6	<i>E. ischnus</i>	25EIS + 25DV + fish	C	B	C	C	A
6	<i>D. villosus</i>	50DV	B	A	B	C	A
6	<i>D. villosus</i>	25DV + 25EIS	AB	A	BC	C	A
6	<i>D. villosus</i>	50DV + fish	BC	B	CD	D	A
6	<i>D. villosus</i>	25DV + 25EIS + fish	B	CD	C	D	A

Discussion

Invasive gammarideans have had a great impact on the macroinvertebrate community in the Rhine; the North American invader *G. tigrinus* dominated the Rhine for years and the relative abundance of gammarideans in the Rhine changed profoundly after the invasion of *D. villosus* in 1995 (Van der Velde

et al. 2000, 2002; Van Riel et al. 2006b). Within 6 years, *D. villosus* had outnumbered *G. tigrinus* and dominated the gammaridean community. Interspecific competition has proved important in determining gammaridean communities (Schoener 1983; Polis et al. 1989; Dick 1996; Petren and Case 1996; Dick and Platvoet 2000; MacNeil and Prenter 2000; Abrams 2001; Van Overdijk et al. 2003), but so have

Table 3 Influences of substratum type, interspecific relations and the interaction between interspecific relations and substratum type on the substratum choice pattern shown by

gammaridean species in the presence of a potentially competitive invader species, analysed by means of a 2-way Generalised Linear Model for Poisson distribution (SAS 8.0)

Substrate preferences of	Source	num DF	den DF	F value	Pr > F	Chi-square	Pr > ChiSq
<i>G. tigrinus</i> in the presence of <i>D. villosus</i>	Substrate type	4	180	20.84	<0.0001	83.37	<0.0001
	Competition	3	180	18.68	<0.0001	56.04	<0.0001
	Substrate × competition	12	180	3.18	0.0004	37.17	0.0001
<i>G. tigrinus</i> in the presence of <i>E. ischnus</i>	Substrate type	4	90	16.19	<0.0001	64.77	<0.0001
	Competition	1	90	14.54	0.0003	14.54	0.0001
	Substrate × competition	4	90	0.31	0.8718	1.23	0.8726
<i>D. villosus</i> in the presence of <i>G. tigrinus</i>	Substrate type	4	200	101.42	<0.0001	405.67	<0.0001
	Competition	3	200	11.62	<0.0001	34.86	<0.0001
	Substrate × competition	12	200	2.56	0.0036	30.67	0.0022
<i>D. villosus</i> in the presence of <i>E. ischnus</i>	Substrate type	4	130	63.18	<0.0001	252.7	<0.0001
	Competition	1	130	17.9	<0.0001	17.9	<0.0001
	Substrate × competition	4	130	1.17	0.3265	4.68	0.3212
<i>D. villosus</i> in the presence of fish	Substrate type	4	120	120.72	<0.0001	482.88	<0.0001
	Competition	1	120	4.99	0.0273	4.99	0.0255
	Substrate × competition	4	120	4.9	0.0011	19.6	0.0006
<i>D. villosus</i> in the presence of <i>E. ischnus</i> and fish	Substrate type	4	240	164.44	<0.0001	657.77	<0.0001
	Competition	3	240	14.03	<0.0001	42.09	<0.0001
	Substrate × competition	12	240	8.1	<0.0001	97.19	<0.0001
<i>E. ischnus</i> in the presence of <i>D. villosus</i>	Substrate type	4	130	195.25	<0.0001	780.99	<0.0001
	Competition	1	130	26	<0.0001	26	<0.0001
	Substrate × competition	4	130	2.02	0.0954	8.08	0.0887
<i>E. ischnus</i> in the presence of fish	Substrate type	4	120	131.42	<0.0001	525.68	<0.0001
	Competition	1	120	22.13	<0.0001	22.13	<0.0001
	Substrate × competition	4	120	3.06	0.0194	12.23	0.0157
<i>E. ischnus</i> in the presence of <i>D. villosus</i> and fish	Substrate type	4	240	316.35	<0.0001	1265.41	<0.0001
	Competition	3	240	13.39	<0.0001	40.16	<0.0001
	Substrate × competition	12	240	3.32	0.0002	39.87	<0.0001
<i>E. ischnus</i> in the presence of <i>G. tigrinus</i>	Substrate type	4	110	55.27	<0.0001	221.07	<0.0001
	Competition	1	110	1.96	0.164	1.96	0.1612
	Substrate × competition	4	110	14.14	<0.0001	56.57	<0.0001

factors such as water temperature and ionic content (Wijnhoven et al. 2003), chlorophyll-*a* availability (Rajagopal et al. 1999; Platvoet et al. 2006), the abundance of bulk prey species like *C. curvispinum* (Van Riel et al. 2006b) and alternative refuges, which are scarce in the main stream of the river Rhine, as macrophytes are absent and the substrata other than stones offer little shelter, resulting in frequent encounters between highly abundant invasive gammarideans with limited chances to escape competition (Van Riel et al. 2006b).

Whereas water temperatures were comparable during the 3 years this study was performed (1992, 1993 and 2001), both chlorophyll-*a* and the abundance of *C. curvispinum* had decreased in 2001, which may have resulted in a reduced food availability for gammarideans (Van der Velde et al. 2000; Van Riel et al. 2006b). Despite its decreased densities, *C. curvispinum* remained dominant among the macroinvertebrate fauna on the stones in terms of numbers, with population peaks of around 40,000 specimens per square metre of substratum (Fig. 1)

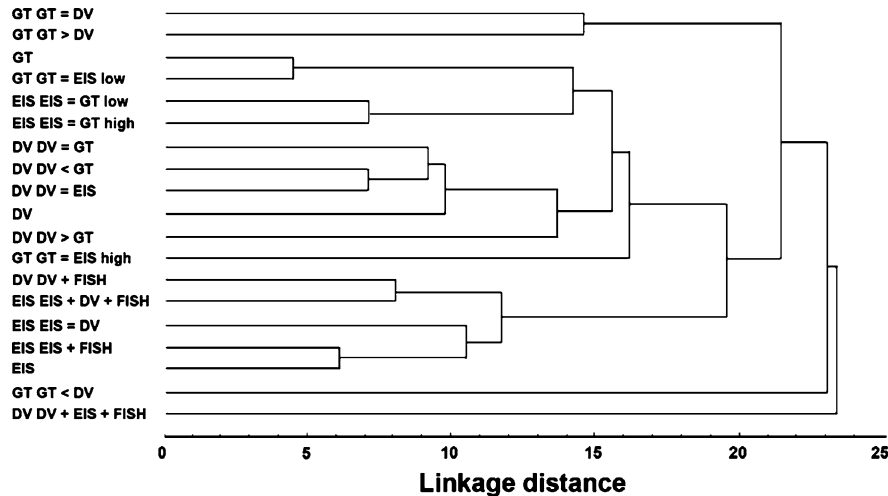


Fig. 3 Complete linkage of substratum choice patterns of the surviving specimens in the various aquarium experiments. Species are indicated by the following codes: GT = *Gammarus tigrinus*, DV = *Dikerogammarus villosus* and EIS = *Echinogammarus ischnus*. The first two letters of the codes indicate the

species which showed the choice pattern, while the following letters indicate the experimental combinations of the species and the densities at which the patterns were observed

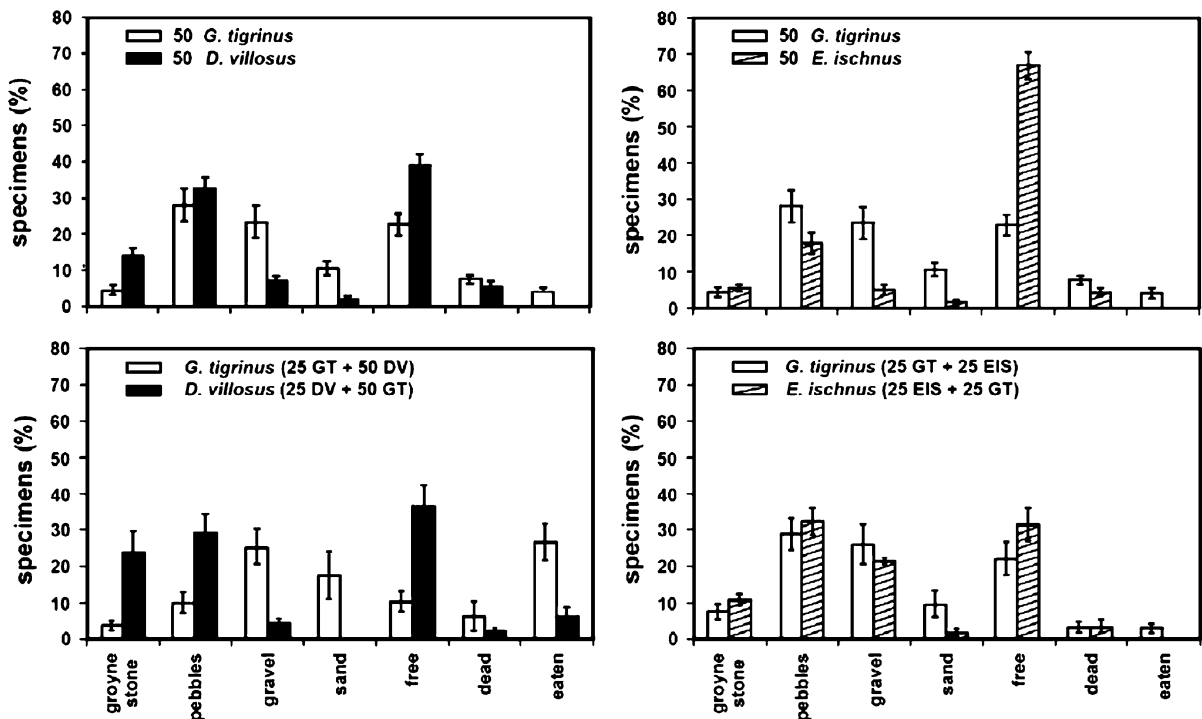


Fig. 4 Substratum occupation and mortality (mean \pm SEM) by *Gammarus tigrinus* (GT), *Dikerogammarus villosus* (DV) and *Echinogammarus ischnus* (EIS) in single and two species

tests. Significant differences in a species' substratum choice are presented in Table 2

(Van Riel et al. 2006b). If the 2001 decrease was large enough to limit *C. curvispinum* as a food resource for gammarideans, it may have intensified

competition and predation between the abundant gammarideans. Before this hypothesis can be confirmed, however, IGP and interspecific competition

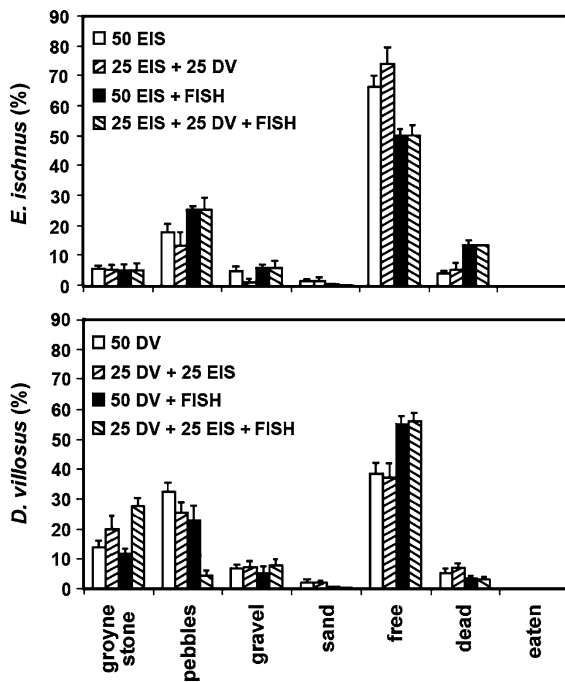


Fig. 5 Substratum occupation and mortality (mean \pm SEM) of *Dikerogammarus villosus* (DV) and *Echinogammarus ischnus* (EIS) in single and two species tests in the absence and in the presence of a predator (*Lepomis gibbosus*) in aquarium experiments. Significant differences for a specific density combination are displayed in Table 2 (Games Howell's multiple comparison test, $\alpha = 0.05$)

between invasive gammarideans in the Rhine must first be demonstrated, which was the goal of the present substratum choice experiments.

Our aquarium experiments showed shifts in substratum choice and alterations in swimming behaviour which indicated interspecific competition between gammaridean species. The substratum choices of *G. tigrinus* and *D. villosus* as tested in the single-species experiments, were found to be highly similar, probably resulting in a high encounter probability. Both species preferred mid-size stone substratum, disliked sand as a substratum, and regularly swam around in the water column. The aquarium experiments revealed that direct competition for substratum can occur between these two species, with *D. villosus* being the stronger competitor. Competitive stress was influenced by population density, and varied between the different types of substratum as a result of substratum choice. Interspecific competition seemed to take place by means of encounter competition and

predation, as *G. tigrinus* was preyed upon increasingly in the presence of *D. villosus* and shifted its substratum choice to finer types of substratum, where it hid. The smaller specimens in particular fled to the finer substrata and even dug themselves into the sand.

The third abundant invasive gammaridean species in the Rhine, *E. ischnus*, primarily occupied the water column in the aquarium experiments and was therefore less likely to encounter the other invaders frequently. No behaviour indicative of competition was found for any species in the presence of *E. ischnus*, neither for *E. ischnus* in the presence of any of the other invaders. However, when a fish occupied the water column and the stone substratum was at the same time occupied by *D. villosus*, *E. ischnus* suffered higher mortality, which could be the result of increased stress, as shelter opportunities to escape from the predators were minimised.

The present study demonstrated different interactions between potential competitors. The two-species aquarium experiments showed species shifting their substratum choice, IGP by *D. villosus* on *G. tigrinus*, species altering swimming behaviour and increased competitive stress in the presence of a fish predator. On the other hand, our monitoring studies on stones in the main stream of the river Waal showed *D. villosus* to be a dominant species, with other gammarideans only occurring in low numbers. Of the two most important habitats, groyne stone riverbanks and sandy streambeds, gammarideans preferred the stones (Van Riel et al. 2006a, b). Increasing the heterogeneity of the ecosystem would provide weaker competitors with refuges to escape IGP by *D. villosus* and/or *G. tigrinus* (Van Riel et al. 2004). In times of limited resources, interspecific competition for substratum and mutual predation can determine the colonisation success of a gammaridean species in the ecosystem.

Based on field observations and experimental studies the following conclusions could be drawn: (a) Interspecific competition and unequal mortality could partly have determined the varying success of the gammaridean invaders in the Rhine, (b) Shifts in mortality, swimming behaviour and substratum choice do indicate interactions between invasive gammarideans, which are determined by IGP and interspecific competition, (c) When refuges are limited, fish predators intensify interspecific competition between gammarideans, and (d) The limited

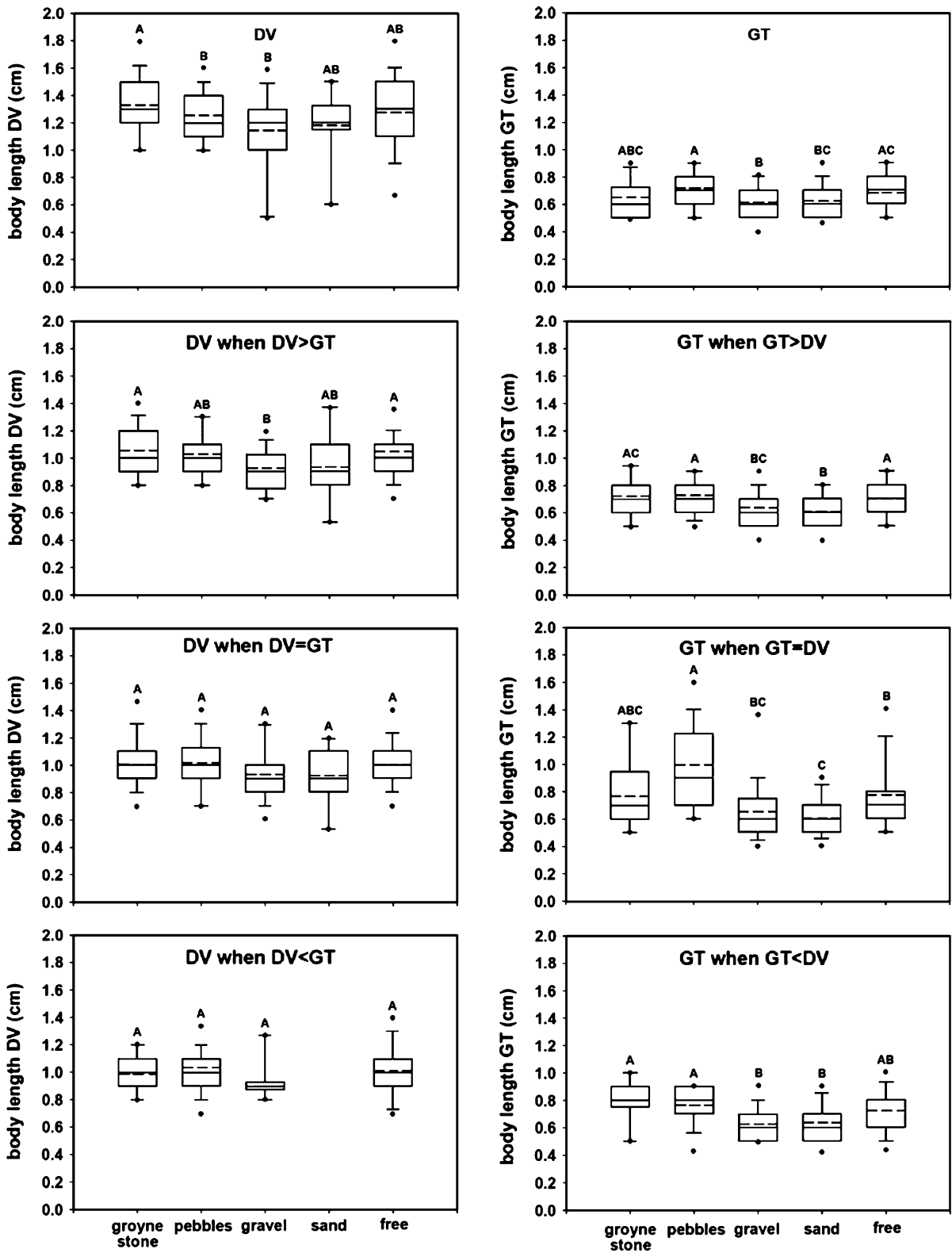


Fig. 6 Body lengths of *Dikerogammarus villosus* (DV, a–d) and *Gammarus tigrinus* (GT, e–h) specimens present on the different substrata in the aquarium experiments. Levels of interspecific competition were manipulated by varying the population densities: 50 DV or GT (a, e), 50 DV or GT + 25 of the other species (b, f), 25 DV or GT + 25 of the other species (c, g), and 25 DV or GT + 50 of the other species (d, h). Different letters indicate significant differences between the body lengths of specimens of the same species present on different substrata. Whisker-box plots represent 25–75% of the values. Bars indicate 5–95% of the values. Dots refer to 1 or 99%. Mean values are indicated by a dashed line in the boxes, whereas a solid line in a box refers to median values

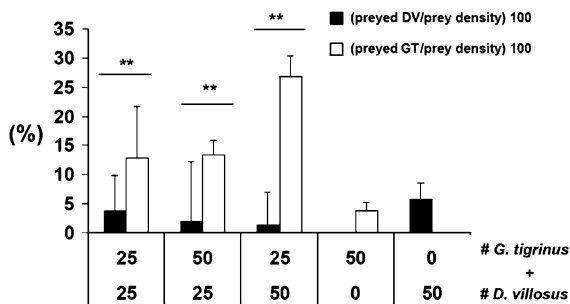


Fig. 7 Predation (mean \pm SEM) between *Gammarus tigrinus* (GT) and *Dikerogammarus villosus* (DV) when the species are present in different densities. The x-axis shows the numbers of specimens of each species present in the aquarium experiments. Different letters indicate significant differences within the same experiment. Bars marked with asterisks indicate significant differences between the different treatments. Significance levels: *** $P < 0.001$, ** $P < 0.01$, * $P < 0.05$

number of biotopes stimulates competition for stones in the Rhine's main channel. More shelter options could increase the opportunities for potentially competitive species to co-exist.

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