Mating structure of the blue and red shrimp, *Aristeus antennatus* (Risso, 1816) characterized by relatedness analysis

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Understanding life history variation and strategies is crucial for stock assessment and fisheries management due to the direct effects on population dynamics, effective population size, sex-ratios, levels of inbreeding, and relatedness among individuals. *Aristeus antennatus* (En — Blue and red shrimp; Fr — Crevette rouge; Sp — Gamba rosada) is one of the most exploited demersal resources in the Western Mediterranean Sea. However, information regarding the mating system and mate choice preferences remains largely unknown. Advances in molecular genetic markers and methods of inferring biological relationships among individuals have facilitated new insights into the reproductive dynamics of the species in the wild. Here, we used microsatellite markers to examine the *A. antennatus* mating system and putative mate choice preferences. Our results provided clear evidence of polyandry and polygyny. Relatedness analyses, together with *Fst* and DAPC values showed females exhibited a mating bias towards unrelated males. Mating males were inferred from spermatophores and suggested males were sympatric with females and were also from other spawning grounds. Our findings provided the first description of the reproductive behavior of blue and red shrimp.

Mating systems have numerous implications for the ecology and evolution of species due to the effects on population dynamics, effective population size (*N*ₑ), levels of inbreeding, and relatedness among individuals¹₂. It is crucial to elucidate life history variation and reproductive strategies in exploited marine species for accurate stock assessment and sustainable fisheries management³ and references therein. A variety of basic mating patterns are described for crustaceans updated in⁴, including exploited marine species. These studies have often focused on reproductive morphological traits, size and age at first maturity, and fecundity or spawning frequency³⁵. Less attention has been devoted to social interactions, mating systems, mate choice, or sex-biased dispersal³⁵. In addition, the reproductive biology of deepest sea taxa is poorly elucidated because the nature of the habitat imposes technical difficulties in observing mating behaviour⁸. Moreover, studies under captivity conditions might alter mating behaviors or kinship structure⁷.

The mating system might be simply described as the number of mates that individuals have during the breeding season⁸. Besides this basic definition, Asakura⁸ described mating systems in terms of the genetic relationships between mating individuals. Barbosa *et al.*⁹ reported that the reproductive strategy was toward improving offspring production and viability. Research has long demonstrated inbreeding caused by the mating of related individuals results in lower offspring survival and an overall population decline updated in¹⁰. Dispersal and mate choice reduce mating between closely related individuals¹¹, but mate choice based on kin recognition is the typical pattern in invertebrate species. For example, females of the German cockroach (*Blattella germanica* Linnaeus, 1767) mate preferentially with distantly related males to increase their reproductive success¹². However, Loyau *et al.*¹³ reported inconsistent results in *Drosophila melanogaster* (Meigen, 1830), where a reduction of offspring viability was observed when mating occurred between unrelated individuals. Such outbreeding depression resulted from breaking apart co-adapted beneficial gene complexes or local genetic adaptations¹¹. Weir *et al.* reviewed in¹⁴ defined relatedness between individuals as the probability of a set of genes shared between individuals are identical-by-descent (IBD). Measuring relatedness requires known pedigrees, but

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often pedigree information is not available in wild populations\(^\text{14}\). Recent advances in molecular markers facilitate probability estimates of shared ancestry from observed genotypes of a set of loci, and several Maximum Likelihood (MLE) and Method of Moment (MoM) estimators have been developed\(^\text{15}\). The ideal situation is when allele frequencies are known from the ancestral population\(^\text{14}\). Nevertheless, this is again an unrealistic situation for most wild populations and biased estimates are therefore obtained\(^\text{14,16}\). In any case, methods inferring relatedness between individuals within and among populations have thoroughly facilitated our understanding of wild populations\(^\text{16}\). Relatedness estimates serve to address several important conservation questions, e.g. mating behaviors, genetic structure, or dispersal dynamics\(^\text{14}\). For example, Stow et al.\(^\text{17}\) demonstrated relatedness due to isolation-by-distance, high female philopatry to their natal nest, and polygynandrous mating in the Australian allodapine bee (Exoneura nigrescens Friese, 1899). Kümerlu and Keller\(^\text{18}\) conducted relatedness analyses and showed limited queen and free adult worker dispersal in a population of the ant Formica exsecta (Nylander, 1846)\(^\text{18}\). In marine crustacean species, few studies of mating preferences related to kinship have contributed to our understanding of population dynamics\(^\text{18}\), but studies in open thelycum (absence of seminal receptacles) shrimp species not suitable to laboratory conditions have not been published.

*Arsteus antennatus* (Risso, 1816) (En — Blue and red shrimp; Fr — Crevette rouge; Sp — Gamba rosada) is one of the most exploited demersal fisheries resources in the Western Mediterranean Sea\(^\text{20}\). However, information regarding the mating system and mate choice preferences remain largely unknown. The species inhabits the Mediterranean Sea at depths ranging from 80 to 2800 m\(^\text{21}\), with sex and size spatial and temporal segregation\(^\text{22}\). Adult females are primarily distributed at shallower depths between 400 and 1000 m throughout the year, while adult males and juveniles inhabit deeper zones below 1000 m\(^\text{23}\). In the Western Mediterranean Sea, reproduction occurs from early spring to summer, when mature males and females form spawning aggregations at depths between 600–800 m\(^\text{23}\) and references therein. The female–biased sex ratio in these aggregations (females may reach up 80% of catches)\(^\text{24}\), suggests some degree of polygyny in *A. antennatus*. During mating, males place spermatophores in the females’ external genitalia (thelycum) with specialized enlarged and folded endopods of the first pair of pleopods, which constitute the copulatory organ called the petasma\(^\text{24}\). Females possess no seminal receptacles; hence, spermatophores attached in the thelycum are the only way to store sperm\(^\text{24}\). In other decapod species, spermatophores are lost at each molt\(^\text{25}\), so the high molt activity observed in *A. antennatus* females from May to September suggests mating might occur several times each reproductive season\(^\text{26}\). After mating, the shrimp shoal breaks up and males return to submarine canyons, while females remain in shallower waters carrying one or more spermatophores in the thelycum\(^\text{27}\). More details on how fertilization occurs remain unknown, but undoubtedly it happens in open waters and parental care is absent\(^\text{27}\), which complicates mating system determination.

In this study, we examine the mating system dynamics and genetic diversity of *A. antennatus* using microsatellite loci. In particular, we assessed multiple mating rates, male origin of multiple spermatophores attached to female thelycum, and putative male choice preferences. These data will provide the first insights into mating systems and reproductive behaviors of blue and red shrimp, which is crucial to understand how exploitation influences the resistance of this species to catastrophic decline and subsequent resilience.

### Materials and Methods

#### Sampling and DNA extraction.

In August 2015, a total of 111 samples of mature *A. antennatus* adults (59♂: 52♀) were collected while on board the *Nova Gasela* trawling vessel at the spawning ground of Palamós (600 m depth), Spain (41°53′04″N and 3°23′77″E). Mature males were identified based on rostrum shorter than 12 mm in length\(^\text{28}\) and the presence of petasma fusion\(^\text{24}\). Similarly, we followed Sardà and Demestre\(^\text{29}\) to identify adult females, and juveniles inhabit deeper zones below 1000 m\(^\text{23}\). In the Western Mediterranean Sea, reproduction occurs from early spring to summer, when mature males and females form spawning aggregations at depths between 600–800 m\(^\text{23}\) and references therein. The female–biased sex ratio in these aggregations (females may reach up 80% of catches)\(^\text{24}\), suggests some degree of polygyny in *A. antennatus*. During mating, males place spermatophores in the females’ external genitalia (thelycum) with specialized enlarged and folded endopods of the first pair of pleopods, which constitute the copulatory organ called the petasma\(^\text{24}\). Females possess no seminal receptacles; hence, spermatophores attached in the thelycum are the only way to store sperm\(^\text{24}\). In other decapod species, spermatophores are lost at each molt\(^\text{25}\), so the high molt activity observed in *A. antennatus* females from May to September suggests mating might occur several times each reproductive season\(^\text{26}\). After mating, the shrimp shoal breaks up and males return to submarine canyons, while females remain in shallower waters carrying one or more spermatophores in the thelycum\(^\text{27}\). More details on how fertilization occurs remain unknown, but undoubtedly it happens in open waters and parental care is absent\(^\text{27}\), which complicates mating system determination.

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**Microsatellite loci.** Genetic diversity at thirteen polymorphic microsatellite loci developed for *A. antennatus*\(^\text{30}\) were analyzed using one singleplex and three multiplex PCR panels delineated to avoid primer-dimer and hairpin formation (assessed *in silico* with AUTODIMER v1.0 program\(^\text{30}\)). For each locus, the forward primer was labeled with one fluorescent dye (PET, 6-FAM, NED or VIC) and loci with overlapping allelic ranges were tagged with different dyes. Multiplex PCRs were conducted in a 10 µL final volume containing 1X GoTag\(^\text{31}\) G2 Hot Start Colorless Master Mix (Promega Corporation), primer multiplex mix at primer-specific concentrations (Table 1), 0 or 1.5 mM MgCl\(_2\) solution (adults and spermatophores, respectively), and ~40 ng of template DNA. The Aa1255 locus was analyzed in a singleplex panel performed in a 15 µL total reaction mixture containing 1X NH\(_4\) Reaction Buffer, 1.5 or 4.6 mM MgCl\(_2\) solution (adults and spermatophores, respectively), 1 µm of dNTP, 0.2 µm of each primer (forward and reverse), 0.375 units BIOTaq\TM DNA Polymerase (Bioline), and ~40 ng of template DNA. All PCR reactions proceeded as follows: initial denaturation step of 2 min at 94°C, followed by 30 (multiplex) or 35 (singleplex) cycles of 30 s at 94°C; annealing for 1 min 30 s at 50°C or 60°C; and elongation for 1 min 30 s at 72°C; with a final extension of 30 min at 60°C (Table 1). One µL of PCR product was mixed with 0.15 µL of GeneScan\TM 500LIZ Size Standard (Applied Biosystems) and 10 µL of highly deionized (Hi-Di) formamide (Applied Biosystems) and denatured for 3 min at 95°C. Resulting amplicons were run on an ABI PRISM 3130 Genetic Analyzer (Applied Biosystems) and genotypes scored using GENEIOUS R7.1.34.
Characteristics of the singleplex and three multiplex PCR reactions used to amplify 13 microsatellite loci in *A. antennatus*. SP, singleplex; MP1, multiplex 1; MP2, multiplex 2; MP3, multiplex 3; *PCR* final volume primer concentration.

| Locus  | Annecling temperature (°C) | Fluorescent dye 5′-forward | Allele size range | Primer concentration (µM)* | GenBank accession number |
|--------|-----------------------------|----------------------------|------------------|-----------------------------|-------------------------|
| SP     |                             |                            |                  |                             |                         |
| Aa1255 | 50                          | VIC                        | 115-163          | 0.2                         | KU195295                |
| MP 1   |                             |                            |                  |                             |                         |
| Aa138  | 50                          | 6-FAM                      | 187-249          | 0.1                         | KU195272                |
| Aa496b | 50                          | PET                        | 412-418          | 0.05                        | KU195279                |
| Aa956  | 50                          | VIC                        | 193-217          | 0.1                         | KU195289                |
| MP 2   |                             |                            |                  |                             |                         |
| Aa123  | 60                          | 6-FAM                      | 425-435          | 0.1                         | KU195269                |
| Aa667  | 60                          | NED                        | 241-265          | 0.1                         | KU195282                |
| Aa681  | 60                          | VIC                        | 224-322          | 0.1                         | KU195283                |
| Aa751  | 60                          | PET                        | 225-235          | 0.1                         | KU195285                |
| Aa1444 | 60                          | PET                        | 180-202          | 0.4                         | KU195297                |
| MP 3   |                             |                            |                  |                             |                         |
| Aa421  | 60                          | PET                        | 169-241          | 0.1                         | KU195278                |
| Aa818  | 60                          | 6-FAM                      | 160-210          | 0.2                         | KU195287                |
| Aa1061 | 60                          | VIC                        | 143-211          | 0.2                         | KU195290                |
| Aa1195 | 60                          | NED                        | 189-204          | 0.2                         | KU195293                |

Table 1. Characteristics of the singleplex and three multiplex PCR reactions used to amplify 13 microsatellite loci in *A. antennatus*. SP, singleplex; MP1, multiplex 1; MP2, multiplex 2; MP3, multiplex 3; *PCR* final volume primer concentration.

**Diversity within and between sexes.** Genetic diversity levels within sexes (females and males) were estimated as the number of alleles per locus (*N_a*) and the observed (*H_o*) and expected (*H_e*) heterozygosity. In addition, at each sex and locus, the conformance of genotypic proportions to Hardy-Weinberg expectations (HW) was calculated using the program FSTAT v 2.9.3.235 and summarized using Wright's fixation index (*F_IS*). Loci exhibiting significant positive *F_YS* values were evaluated for potential genotyping errors due to stuttering, allele dropout, or null alleles using MICRO-CHECKER v 2.2.3 software. The Dempster et al.37 algorithm was employed to estimate the frequency of null alleles; however modifications were implemented in the FREENA software. Genetic diversity differences (*N_a, H_o, H_e, F_YS*) between sexes were statistically tested in SPSS v 2339 using the Wilcoxon signed rank test. Genetic differentiation between males and females was examined by *F_YS* values using the program FSTAT v 2.9.3.235.

Spermatophores indicated mature males effectively mating with females; therefore, we also examined these individuals for deviations from HW expectations and diversity levels, as detailed above for males and females. Diversity levels for the spermatophore group were compared with males and females and genetic differentiation from these two groups was estimated as *F_YS* values. Furthermore, discriminant analysis of principal components (DAPC) using the package ADEGENET v 1.4–2 was performed in R v. 3.3.241 to establish an additional view of differentiation among the three groups (males, females and spermatophores). DAPC does not require any specific population structure model and it is free of assumptions regarding Hardy-Weinberg or gametic equilibrium. Alleles are considered original variables with the largest between-group and smallest within-group variance.

**Paternity and relatedness in the spawning ground.** The exclusion probability of identity (*PI*) (probability that two randomly selected individuals matched their genotypes by chance) was applied to test the potential that the set of microsatellite loci was robust for individual identification using the software GIMLET v 1.3.343. The regrouping genotype option was chosen to detect multiple paternity by comparing all spermatophore genotypes. This analysis was conducted to detect males mating multiple times with either the same female or different females. Similar comparisons were used to ascertain if some spermatophores originated from sampled males.

Relatedness analyses were used to determine if all specimens (females, males, and mating males inferred from spermatophores) in the spawning ground were derived from the same population. We first confirmed the efficiency of seven relatedness estimators available in COANCESTRY v 1.0.1.7 software. We subsequently applied the related estimator to distinguish among low relatedness groups, which might be inherent in this species; 1,000 dyads were simulated for each of the following relationships: first cousins (*r_xy* = 0.125), second cousins (*r_xy* = 0.031), and unrelated (*r_xy* = 0). The best estimator was chosen for additional relatedness comparisons. Consequently, the distribution of observed pairwise relationship coefficients within each reproductive group (male, female, and spermatophores) was compared using COANCESTRY v 1.0.1.7 with the relationship coefficients expected for a sample of 1,000 unrelated simulated offspring generated by mating 1,000 simulated males and 1,000 simulated females from our male and female genotypes. All simulated individuals were obtained using HYBRIDLAB v 1.0 software. In addition, comparisons of relatedness between all male, female, and spermatophore pairs were used to determine if all sampled individuals derived from the same population. Finally, average relatedness in observed female-spermatophore pairs was compared to all other potential female-spermatophore pairs (excluding observed female-spermatophore pairs) and female-male pairs to determine if mate choice was associated with kinship. For each of the above comparisons, 1,000,000 replicate iterations were computed in COANCESTRY v 1.0.1.7 and a genotyping error rate of 2% was allowed.
and another three female individuals carried three. In all cases, multiple spermatophores from the same female were identified in six of fifty-two females (11.5%). Three females carried two spermatophores in their thelycum conducted an analysis with six loci (Aa138, Aa956, Aa496b, Aa123, Aa751, and Aa1195). We observed the same three groups, larger average \( A \) (\( N \)) ranged from 0.3939 in adult females to 0.4668 in mating males, genotyped from the 61 spermatophores. In all three groups, larger average \( H \) compared with \( H_s \) was detected, ranging from 0.6116 in females, 0.6166 in males, and 0.6190 in spermatophores. Significant differences were not detected for these three diversity measures between males and females, but the Wilcoxon signed rank test indicated the average \( H_0 \) (\( Z = 2.072; P = 0.038 \)) and \( H_s \) (\( Z = 2.132; P = 0.033 \)) were larger in spermatophores than in females. Several loci for each reproductive group showed significant positive \( F_{ST} \) values and \( H_e \) was often lower than \( H_s \). Following Bonferroni corrections, significant genotypic deviations from HW expectations were observed at eight loci in male and female groups and seven in spermatophores. MICRO-CHECKER v 2.2.3 revealed these significant differences were likely due to the presence of null alleles, but some biological processes also produced positive \( F_{IS} \) and \( F_{ST} \) values (e.g.: Wahlund effect from high migration rates). At loci where null alleles were detected, estimated frequencies ranged from 0.1017 to 0.8000. Results did detect significant differences between males and females (\( F_{CT} = -0.0011; P = 0.5000 \)) and males and spermatophores (\( F_{ST} = 0.0016; P = 0.5500 \)). Results did detect significant differentiation between females and spermatophores (\( F_{ST} = 0.0091; P = 0.0167 \)). Bidimensional DAPC scatterplots obtained from the 25 principal components retained more than 80% of the total genetic variance (Fig. 1a). The first component in the scatterplot depicted a distinction between females and their spermatophores. Males were located in a central position with an overlapping distribution at the left side of the central plot axis scattered with a portion of the females to the right of the main axis with some spermatophores. Spermatophores showed a somewhat bimodal distribution, scattered with females and males for the first component (Fig. 1b).

**Paternity and mating system.** Excluding locus Aa421, a combined probability of identity exclusion higher than 0.9999 was observed in all three groups; males, females, and mating males represented by their spermatophores (Table 2). Therefore, the probability that two randomly selected individuals matched their genotypes by chance was lower than 0.0001. In order to check how null alleles could influence probability of identity we conducted an analysis with six loci (Aa138, Aa956, Aa496b, Aa123, Aa751, and Aa1195). We observed the same results we previously generated, suggesting null alleles have no influence on paternity. Multiple spermatophores were identified in six of fifty-two females (11.5%). Three females carried two spermatophores in their thelycum and another three female individuals carried three. In all cases, multiple spermatophores from the same female showed different genotypes. These results indicated multiple mating in females. Identity analysis on 12 microsatellite loci resulted in identical genotypes in one pair of spermatophores from different females, which was

| Sample | Aa138 | Aa1255 | Aa956 | Aa496b | Aa123 | Aa681 | Aa667 | Aa1444 | Aa751 | Aa818 | Aa1061 | Aa1195 | Aa421 | All loci |
|--------|-------|--------|-------|--------|-------|-------|-------|--------|-------|-------|--------|--------|-------|---------|
| Females | \( N_0 \) | 18 | 10 | 6 | 2 | 5 | 15 | 4 | 8 | 3 | 5 | 9 | 4 | 6 | 7.3080 |
| \( H_0 \) | 0.7885 | 0.4444 | 0.5192 | 0.3962 | 0.4706 | 0.4510 | 0.3269 | 0.4286 | 0.1346 | 0.2745 | 0.3725 | 0.5962 | 0.2000 | 0.3959 |
| \( H_e \) | 0.9097 | 0.7818 | 0.5907 | 0.8924 | 0.6202 | 0.6992 | 0.6667 | 0.7366 | 0.2285 | 0.6524 | 0.7063 | 0.5913 | 0.7247 | 0.6116 |
| HWE | \( 0.0164 \) | \( 0.0000^* \) | \( 0.1372 \) | \( 1.0000 \) | \( 0.0476 \) | \( 0.0000^* \) | \( 0.0000^* \) | \( 0.0000^* \) | \( 0.0002^* \) | \( 0.0000^* \) | \( 0.0000^* \) | \( 0.0000^* \) | \( 0.0000^* \) |
| Nu | 0.0632 | 0.1878 | 0.0148 | 0.0000 | 0.0800 | 0.1617 | 0.1981 | 0.1735 | 0.0941 | 0.2333 | 0.1958 | 0.0000 | 0.3004 |
| \( F_s \) | 0.1333 | 0.4315 | 0.1210 | -0.4048 | 0.2412 | 0.3550 | 0.5096 | 0.4182 | 0.4109 | 0.5792 | 0.4725 | -0.0083 | 0.7240 | 0.3559 |
| \( P_l \) | 0.8220 | 0.9145 | 0.7825 | 0.1705 | 0.7896 | 0.8814 | 0.8101 | 0.8870 | 0.3863 | 0.8259 | 0.8587 | 0.7650 | 0.8781 | 0.9999 |

Table 2. Summary statistics of genetic diversity in females, males, and spermatophores of A. antennatus. \( N_0 \), Number of alleles; \( H_0 \), observed heterozygosity; \( H_e \), expected heterozygosity; HWE, P, conformance to Hardy-Weinberg equilibrium; Nu, null allele frequency; \( F_s \), Wright’s fixation index; \( P_l \), Probability of Identity; *Significant departure from HWE; †Null alleles detected; ● see text.
consistent with the above estimated exclusion probability of identity, also supporting multiple mating in males. A genetic identity match was not found between spermatophores and males.

All relatedness coefficients produced small and similar estimates between simulated first and second cousin pairs, and more importantly, these values were not clearly differentiated from the values obtained between unrelated pairs of individuals (Table S1). A more robust relatedness estimator was not available for our data set. The triadic likelihood estimator was selected for all further analyses because it typically outperforms other estimators in natural populations, where most dyads are unrelated or only loosely related. The pairwise relatedness in our groups ranged from 0.0528 in males to 0.0657 in spermatophores (Table 3). The average relatedness among

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**Table 3.** Bootstrap test to compare relatedness ± standard error (SE) between groups using TrioML estimator. NS, no significant difference.

| Group 1                        | Mean group 1 ± SE | Group 2                  | Mean group 2 ± SE | P     |
|-------------------------------|-------------------|--------------------------|-------------------|-------|
| Females                       | 0.0584 ± 0.0024   | Unrelated                | 0.0591 ± 0.0001   | NS    |
| Males                         | 0.0528 ± 0.0021   | Unrelated                | 0.0591 ± 0.0001   | NS    |
| Spermatophores                | 0.0657 ± 0.0023   | Unrelated                | 0.0591 ± 0.0001   | P < 0.01 |
| Females                       | 0.0584 ± 0.0024   | Males                    | 0.0528 ± 0.0021   | NS    |
| Males                         | 0.0528 ± 0.0021   | Spermatophores           | 0.0657 ± 0.0023   | P < 0.01 |
| Females-spermatophore         | 0.0368 ± 0.0070   | All other female-spermatophore pairs | 0.0534 ± 0.0016 | NS |
| Females- spermatophore        | 0.0368 ± 0.0070   | All female-male pairs    | 0.0565 ± 0.0016   | 0.05 < P < 0.1 |

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**Figure 1.** Discriminant Analysis of Principal Components (DAPC) among females, males, and spermatophores. (a) Genetic differentiation among females, males, and spermatophores by DAPC. Groups are displayed by different colours and inertia ellipses. Dots represent individuals. The relative portions of the variance captured by first and second axes are illustrated in the DA eigenvalues plot (bottom right corner). The PCA eigenvalues plot in the bottom left corner shows the portion of variance retained when maintaining 25 PCA axes. (b) Density plot of individuals along the first discriminant function from the DAPC for females, males, and spermatophores.

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female individuals (0.0584) was not statistically different from the 1,000 simulated unrelated individuals (0.0591). Similar results were obtained for males, but the observed relatedness among spermatophores differed significantly from the simulated, indicating the spermatophores from males were more closely related than expected by chance alone (Table 3). The average relatedness among females (0.0584) was not significantly higher than the average relatedness among males (0.0528) (Table 3), but relatedness among spermatophores (0.0657) was higher than between all female pairs and male pairs (P < 0.01). In addition, the average relatedness between a female-spermatophore pair was similar to all other female-spermatophore pairs, but the value was slightly differentiated from all possible pairs involving a female and an adult male, suggesting some mating males were differentiated from the sampled males (P < 0.1) (Table 3).

Discussion

The results of our study showed a mean number of alleles per locus (9.38) (Table 2) in A. antennatus lower than the value reported in Italian blue and red shrimp samples (13.5)48. However, our results were consistent with those reported for other shrimp species, including whiteleg shrimp (Litopenaeus vannamei Boone, 1931; mean: 7.9)49 and the deep-sea hydrothermal vent shrimp (Rimicaris exoculata Williams and Rona, 1986; mean: 8.73)50. Cannas et al.48 reported higher mean H0 (0.65) with an increased number of alleles compared with our data (mean females: 0.39; mean males: 0.42). In both studies, deviations from HW genotype expectations often resulted from heterozygote deficits in several microsatellite loci, a common result in the species of Penaeoidea45–47. Several explanations for the pattern have been proposed, including stutter bands, null alleles, and biological factors, e.g. mating tactics, recent population bottlenecks, and/or subpopulation structure. In fact, Wahlund effects resulting from subpopulation structure could not be rejected, as indicated by DAPC results on sampled spermatophores (Fig. 1b). Our results provided clear evidence for the A. antennatus mating system involves multiple mating for both sexes, suggesting that the reproduction in A. antennatus involves polyandry (11%) and polygyny (1.5%). We confirmed female multiple mating by conducting an extended genotyping of spermatophores from 20 additional females collected from the same spawning ground in 2016. Each female carried more than one spermatophore in her thelycum and in each sample different male genotypes were obtained (Table S2). Multiple female mating (polyandry) is common across crustacean species, particularly in species where females do not store sperm in specific structures58, 59. Multiple mating in invertebrate females: (i) facilitates retention of genetic diversity and increased offspring fitness (e.g., the Pacific goose neck barnacle51); (ii) prevents the risks of choosing an infertile partner (e.g., polychaetes56); (iii) favors the storage of enough sperm to fertilize the entire clutch (e.g., lepidopterans57); (iv) and might prevent mate competition that causes injuries (e.g., the common rock shrimp52). In the case of A. antennatus, the benefits of multiple mating could be a compendium of the first three factors, due to the high fecundity of females (≥600,000 oocytes produced in each vitellogenesis53); the latter option is unlikely due to the sex ratio of the species (females seems to be much more abundant than males53) and the small size and without hypertrophied weaponry of the males (major chelipeds and third maxillipeds are evident in aggressively dominant males53). Nevertheless, multiple mating does not always translate into multiple paternity. For instance, females of some crustacean species show behavioral and physiological mechanisms to control male fertilization, such as selective sperm passage or spermatophore removal59, 60. In the caridean rock shrimp (Rynchocinetes typus H. Milne Edwards, 1837), females accept multiple mating to avoid injury by harassing males, but afterwards remove spermatophores from subordinate males to guarantee sperm from the highest quality males61, 62. Thus, our A. antennatus findings demonstrated multiple mating, but did not ensure multiple paternity.

Our data demonstrated that males and females were not genetically differentiated, but mating males, inferred from spermatophores, were somewhat distinct from sampled males. In fact, the DAPC scatterplots showed a portion of spermatophores did not overlap with adult males (Fig. 1b) and females were clearly differentiated from their spermatophores. Females apparently bias mating towards unrelated males. Mating males inferred from spermatophores might include those sympatric with females, as well as those from other spawning grounds. Evidence for male choice in favor of genetically dissimilar partners was reported in some invertebrates, interpreted as a way of inbreeding avoidance, which improves reproductive survival and individual fitness, and increases genetic diversity of local populations63, and references therein. Liu et al.64 found that females of the cabbage beetle (Colaphellus bowringi Baly, 1865) preferred to mate with non-siblings rather than siblings, decreasing the potential for inbreeding depression. Simmons et al.65 observed similar behavior in the Australian black field cricket (Teleogryllus oceanicus Le Guillou, 1841), where females fertilized their eggs with sperm from non-sibling males rather than full-siblings. Gherardi et al.66 suggested that invertebrates make genetic decisions regarding their mating partners; however, relatively little is known about the mechanisms by which recognition is performed, especially in deep-water species, which are not amenable to laboratory conditions, as is the case for A. antennatus. One possibility for the observations made is that these species, use waterborne chemical cues (olfactory) to obtain information about conspecífics and consequently distinguish between kin and non-kin individuals57, 58. The big-clawed snapping shrimp (Alpheus hetrochaelis Say, 1818) employs this recognition mechanism to differentiate between familiar and unfamiliar individuals67. Another possibility is these taxa develop an olfactory and contact chemoreception combination mechanism, comparable to the eusocial coral-reef shrimp (Synalpheus regalis Duffy, 1996), where colony members discriminate between nest-mates and foreign conspecifics19, 68. Often, marine Penaeoidea shrimps with pure searching mating strategy have small sized males, as does A. antennatus, and use contact chemical communication to search for receptive females in reproductive aggregates21–23, 58. This strategy, for us the most plausible in A. antennatus was observed in terrestrial (e.g., mangrove tree crab38) and freshwater decapoda (e.g., freshwater atid shrimp55).

In general, when individuals of one sex do not disperse far from the natal group, they are expected to have increased relatedness on average than those that disperse71. Our results showed that the spermatophores were differentiated from sampled males and females, suggesting that they were deposited in the female thelycum by A. antennatus migrating males. In contrast, using different set of loci Cannas et al.48 analyzed the populations of
A. antennatus of three Italian collections at different depths (Sant’ Antioco, <800 m; Pesca Profonda-Sperimentale North, 1,464 m; Pesca Profonda-Sperimentale South, 1,113 m) and suggest female-biased dispersal. The different depth sampling design might explain the lack of congruency between Cannas et al. 1995 and present results (600 m). Oceanographic currents in the Mediterranean Sea are different depending water masses at distinct depths (Surface Waters, <150 m; Intermediate Waters, 150–1,000 m; Deep Waters, >1,000 m) which could facilitate different dispersal patterns 25. Contrasting results on dispersal patterns might also reflect temporal and spatial ecological variation affecting resource availability and population density 21–25,3, 24. Otherwise, the sex ratio obtained in Pesca Profonda-Sperimentale North (1:4) differed from that in the North - Western Mediterranean Sea 23,26,73, which suggested that the former locality supports a unique population dynamic.

Our research demonstrated a polyandry and polygyny mating system in the blue and red shrimp. Additional research in different natural populations will be fundamental to fully elucidate the mating system in A. antennatus. Mating tactics might vary among populations depending on population density, mate quality, mate availability or variation in environmental conditions, as Gosselin et al. 2004 reported in the American lobster (Homarus americanus). The apparent capacity of A. antennatus to select mating partners based on genetic dissimilarity provides the opportunity to pursue new research avenues and explore possible factors influencing mate choice in this species. Additionally, it will be important to include information regarding life history variation and strategies for stock assessment and effective fishery management of heavily exploited marine resources, such as A. antennatus, to ensure sustainability.

Data Availability
All data are included in the article.

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**Author Contributions**

L.P., M.V., J.L.G.M., S.H. and M.I.R. conceived the ideas and designed the study. L.P. performed experiments, analysed the data and wrote the original manuscript draft. All authors revised the manuscript for significant intellectual content and approved the final version.

**Additional Information**

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