

ORIGINAL ARTICLE

Mito-nuclear discordance in the degree of population differentiation in a marine goby

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An increasing number of phylogeographic studies on marine species shows discordant patterns in the degree of population differentiation between nuclear and mitochondrial markers. To understand better which factors have the potential to cause these patterns of discordance in marine organisms, a population genetic study was realized on the sand goby *Pomatoschistus minutus* (Pallas 1770; Gobiidae, Teleostei). Sand gobies from eight European locations were genotyped at eight microsatellite markers. Microsatellites confirmed the global phylogeographical pattern of *P. minutus* observed with mitochondrial DNA (mtDNA) markers and nuclear allozyme markers. Three groups consistent with the mitochondrial lineages were defined (the Mediterranean, Iberian and North Atlantic groups) and indications of a recent founder event in the northern Baltic Sea were found. Nevertheless, differences

in the degree of population differentiation between the nuclear and mitochondrial markers were large (global F_{ST} -values for microsatellites = 0.0121; for allozymes = 0.00831; for mtDNA = 0.4293). Selection, sex-biased dispersal, homoplasy and a high effective population size are generally accepted as explanations for this mitonuclear discrepancy in the degree of population differentiation. In this study, selection on mtDNA and microsatellites, male-biased dispersal and homoplasy on microsatellite markers are unlikely to be a main cause for this discrepancy. The most likely reason for the discordant pattern is a recent demographical expansion of the sand goby, resulting in high effective population sizes slowing down the differentiation of nuclear DNA.

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Introduction

Traditionally, the genetic structure of marine organisms has been thought to be homogeneous because of the lack of obvious barriers to gene flow in the 'open' marine environment. In the last decade, however, an increasing number of population genetic studies have described distinct genetic structuring for several marine species on large and small geographical scales (Hauser and Carvalho, 2008). Those observed population genetic structures reflect both historical and contemporary processes (Balloux and Lugon-Moulin, 2002). Geographical and climatic factors acting during the Pleistocene glaciations (1800–11.5 ka) are the major factors responsible for the present genetic structure of most extant marine species (Hewitt, 2000). Heterogeneity in the marine environment owing to the influence of climate, hydrodynamics and topography, together with biological traits, such as sex-dependent migration, phyloptry and assortative mating, which may counteract gene flow, enhance genetic structuring (Ruzzante *et al.*, 1998).

These recent insights are mainly due to the increased popularity of polymorphic microsatellite markers in

marine population and landscape genetics (Jørgensen *et al.*, 2005). Microsatellites have proven for many species to be more powerful for resolving population structure than mitochondrial DNA (mtDNA) and allozyme markers (De Innocentiis *et al.*, 2004; Nesbø *et al.*, 2000). However, this is not applicable to all marine organisms (Lukoschek *et al.*, 2008). There is almost certainly a publication bias for marine studies detecting microscale population structures with microsatellites (Hauser and Carvalho, 2008). An increasing number of studies have shown a large difference in the order of magnitude for the population divergence between nuclear and mitochondrial markers. Among those, mtDNA shows a higher sensitivity to resolve the phylogeographical and population genetic structure (Peijnenburg *et al.*, 2006; Lukoschek *et al.*, 2008). This discordance in resolution among markers may result from the differential effects of genetic drift, mutation and migration on a marker class, or may result from selection or sex-biased dispersal (Buonaccorsi *et al.*, 2001). More research is required to study how common and important those factors are in the marine environment.

The sand goby *Pomatoschistus minutus* (Pallas 1770; Gobiidae, Teleostei) is a small marine demersal fish common in shallow waters along European coasts (Miller, 1986) (Figure 1). Although earlier analyses with allozyme markers showed very low population differentiation values for *P. minutus* (Stefanni *et al.*, 2003), a recent study revealed that the sand goby is highly structured at the mtDNA cytochrome *b* (cyt *b*) locus

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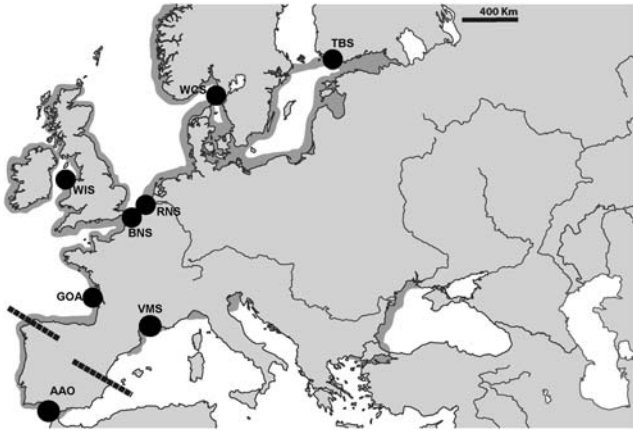


Figure 1 Geographical distribution of the eight sampling locations of sand goby *P. minutus*. The shaded area represents the distribution range of *P. minutus* according to Miller (1986). The two dashed lines represent the main phylogeographical barriers for the sand goby based on the mtDNA analysis of Larmuseau *et al.* (2009b) that differentiate the three isolated and differently evolving sets of populations. See Table 1 for sample codes.

(Larmuseau *et al.*, 2009b). On the basis of mtDNA, middle Pleistocene glaciations yielded three isolated and different evolving sets of sand goby populations. Reciprocal monophyly was observed between a Mediterranean Sea (MS) and an Atlantic Ocean (AO) Clade. The AO-Clade contains two evolutionary significant units: the Iberian Peninsula (IB) Group and the North Atlantic Group. For the North Atlantic Group, there is evidence for geographic sorting of the ancestral mtDNA haplotypes with recent independent radiations in the Baltic Sea, Irish Sea, North Sea and the Bay of Biscay (Larmuseau *et al.*, 2009b). Analyses with allozyme markers revealed weak divergence between Mediterranean and Atlantic populations for *P. minutus* and no differentiation elsewhere, which was attributed to high contemporary gene flow throughout its whole distributional range (Stefanni *et al.*, 2003). However, the biology and morphology of sand gobies indicate reduced levels of contemporary gene flow (Miller, 1986). Moreover, the mitochondrial phylogeography invalidates the interpretation of the allozyme data because of the high F_{ST} values between sand goby populations and the lack of common haplotypes between the three evolutionary significant units (Larmuseau *et al.*, 2009b). The reasons for this discordant pattern can be surveyed by using another set of nuclear markers, such as DNA microsatellites. In contrast to allozymes, DNA microsatellite markers promise a higher resolution for population differentiation in the sand goby because of a higher mutation rate (Chistiakov *et al.*, 2006). Moreover, Pampoulie *et al.* (2004) detected genetic structure on a microscale for *P. minutus* in the southern North Sea using microsatellites, whereas Gysels *et al.* (2004b) did not in the same region using allozyme markers.

In this study, two hypotheses are tested: (a) is the phylogeographical structure of the sand goby based on data of the mtDNA *cyt b* locus (Larmuseau *et al.*, 2009b) confirmed by nuclear microsatellite markers, and (b) do microsatellite markers in comparison with mitochondrial markers show a superior resolution to reveal the phylogeographical and population genetic structure?

Materials and methods

Sampling and species/sex identification

A total of 696 *P. minutus* individuals were caught at eight locations along the European coast between September 2002 and January 2007 (Table 1 and Figure 1). Many of the samples have already been included in a previous mtDNA study (Larmuseau *et al.*, 2009b). Locations BNS and VMS were sampled twice to perform an estimation of effective population size. All new samples were identified as *P. minutus* morphologically, based on the dermal head papillae (Miller, 1986) and pigmentation pattern (Hamerlynck, 1990), and genetically according to a molecular tool described by Larmuseau *et al.* (2008). The sex of each sand goby was determined by the shape of the urogenital papilla as drawn by Rodrigues *et al.* (2006).

Microsatellite genotyping

Genomic DNA was extracted from fin clips stored in 100% ethanol using the NucleoSpin Extraction Kit (Machery-Nagel GmbH, Düren, Germany). Each individual was genotyped at nine multiplexed microsatellite loci (Pmin03, Pmin04, Pmin09, Pmin16-2, Pmin20, Pmin29, Pmin31, Pmin35 and Pmin38) (Larmuseau *et al.*, 2007) on an ABI 3130 automated capillary DNA sequencer (Applied Biosystems, Foster City, CA, USA). Several methods were used to mitigate genotyping errors in the data set (Supplementary materials); marker Pmin20 was excluded for further statistical analyses.

Genetic diversity

We used genotype and allele frequencies of the microsatellite loci to obtain standard estimates of genetic diversity within and between sample sites. Genetic variation in each population was measured by calculating the mean number of alleles per locus, the observed (H_O) and unbiased expected (H_E) heterozygosities and the F_{IS} . The deviation from the Hardy–Weinberg equilibrium was assessed with GENETIX v.4.05 (Belkhir *et al.*, 2004). Allelic richness, which corrects the number of alleles for sample size, was assessed using FSTAT v.2.9.3.2 (Goudet, 2001). Exact tests of linkage disequilibrium between pairs of loci were calculated at each location, each region and across all samples using GENEPOP v.3.4. (Raymond and Rousset, 1995).

Patterns of population subdivision

Different methods were used to reveal the population substructure of *P. minutus*. First, a factorial correspondence analysis (FCA) of individual multilocus genotypes was performed using GENETIX to reveal the portion of the hyperspace of all genotypes occupied by each group of individuals. Second, population differentiation was quantified in GENETIX using the standardized allelic variation F_{ST} , estimated as θ (Weir and Cockerham, 1984), and in SPAGeDi 1.2 g (Hardy and Vekemans, 2002) using an analog of F_{ST} for microsatellites R_{ST} , estimated as \bar{n} (Slatkin, 1995). F_{ST} -linked pairwise genetic distances were calculated according to Cavalli-Sforza and Edwards (1967) (D_{CE}) with GENETIX, and R_{ST} -linked pairwise genetic distances calculated according to Goldstein *et al.* (1995) ($d\mu^2$) were obtained with SPAGeDi. F_{ST} value and D_{CE} value were tested for significance against 10^4

Table 1 Overview of the 10 samples of *P. minutus* collected at eight different locations

Code	Area	Country	Location	Date	Coordinates	N
TBS	Northern Baltic Sea	Finland	Tvärminne	Jul/06	59°50'N–23°12'E	96
WCS	Skagerrak	Sweden	Bökevik Bay, Skaftö island	Jun/06	58°14'N–11°26'E	88
RNS	North Sea	The Netherlands	Renesse	Nov/05	51°44'N–03°47'E	48
BNS1	North Sea	Belgium	Oostduinkerke	Nov/06	51°08'N–02°40'E	47
BNS2	North Sea	Belgium	Oostduinkerke	Sep/02	51°08'N–02°40'E	95
WIS	Irish Sea	Wales (UK)	Llanfairfechan	Nov/06	53°59'N–03°59'W	45
GOA	Bay of Biscay	France	Gironde estuary	Aug/06	45°36'N–01°01'W	40
AAO	Iberian Peninsula	Spain	Guadalquivir river estuary	Nov/06	36°58'N–06°10'W	95
VMS1	Mediterranean Sea	France	Vaccarès lagoon	Jan/06	43°32'N–04°35'E	94
VMS2	Mediterranean Sea	France	Vaccarès lagoon	Jan/07	43°32'N–04°35'E	48

Locations BNS and VMS were sampled twice in order to perform an estimation of effective population size. *N* indicates sample size.

random permutations of the data in GENETIX. The significance for R_{ST} value and $d\mu^2$ value were tested in SPAGeDi also against 10^4 random permutations. A sequential Bonferroni test (Rice, 1989) and the false discovery rate control (Verhoeven *et al.*, 2005) were applied to correct significance levels for multiple testing. Standardized genetic differentiation measures were obtained by dividing F_{ST} measures by the maximum values for F_{ST} (Hedrick, 2005; Meirmans, 2006), calculated using the pragmatic recoding approach suggested by Meirmans (2006). To assess the influence of stepwise-like mutations versus drift on genetic differentiation, we performed a permutation test available in the software SPAGeDi. Allele size at each locus was randomly permuted among allelic states (2000 mutations) to simulate a distribution of R_{ST} values (pR_{ST}) and 95% confidence intervals (CIs) under the null hypothesis that differences in allele sizes do not contribute to population differentiation (Hardy *et al.*, 2003). Third, the classical multidimensional scaling analyses based on the two types of genetic distances were obtained using the Vegan package in R (Oksanen *et al.*, 2007). Ordination plots with a stress value below 0.20 provide interpretable information concerning intersite relationships (Clarke, 1993). Fourth, to analyze the effect of geographical distance on genetic distance, the Mantel test in GENETIX (Belkhir *et al.*, 2004), which computes the correlation between distance matrices by means of a permutation procedure (Mantel, 1967; Smouse *et al.*, 1986), was used. Geographical distances were obtained as the shortest coastal distances between sites using the electronic atlas EN-CARTA (Microsoft 2001). Both types of pairwise genetic distances, D_{CE} and $d\mu^2$, were used for the Mantel test. Fifth, a Bayesian clustering analysis was realized for the microsatellite data using the program STRUCTURE v.2.2 (Pritchard *et al.*, 2000). This approach, which estimates the number of independent genetic clusters in the data set, does not require *a priori* information about population structure, and thus provides an estimate of genetic structure independent of the origin of samples. We used the no-admixture algorithm without prior population information and used 10 000 runs as burn-in and 100 000 runs for each of three Markov chains. The no-admixture algorithm was used on the basis of detection of the lack in gene flow between and within marine systems for *P. minutus*, at the mtDNA level (Larmuseau *et al.*, 2009b). Moreover, this model is appropriate for studying fully discrete populations and is often more powerful than the admixture model at detecting subtle structure (see

manual STRUCTURE v.2.2). For each simulation of $k=1-10$ (no. of clusters), we used 10 replicates. The uncorrelated allele frequency model with parameter λ set to 1 was used as the model assuming that the allele frequencies in each population are independent. This is the original model used in Pritchard *et al.* (2000). We selected the most likely number of clusters given the data by choosing the number of clusters where we observed the largest difference in log likelihoods (ΔK) (Evanno *et al.*, 2005). Finally, the overall pattern of population genetic structure among regions was assessed with a Bayesian approach developed by Ciofi *et al.* (1999) and implemented in the program 2MOD by MA Beaumont (<http://www.rubic.rdg.ac.uk/~mab/software.html>). Two models are evaluated: (1) the gene flow model, which assumes that gene frequencies within populations are caused by a balance between genetic drift and gene flow, and (2) the drift model, in which allele frequencies within populations are evolving purely through drift. The Markov chain Monte Carlo search was carried out twice using 10^5 iterations. The first 10% of the runs were discarded to remove the effects of initial starting parameters. In each case, the probability of a model was estimated from the number of times it appeared during the simulation.

Comparison between microsatellite and mitochondrial data

MtDNA *cyt b* data were available for eight out of ten sand goby samples (TBS, WCS, RNS, BNS1, WIS, GOA, AAO and VMS1) (Larmuseau *et al.*, 2009b). On the other hand, the allozyme analyses by Gysels (2003) and by Stefanni *et al.* (2003) were restricted to samples from other locations. Therefore, it was only possible to compare statistically the results of microsatellites and mtDNA *cyt b* data. Two statistical methods were used to compare the degree of population differentiation between the types of genetic distances calculated for the microsatellite data (D_{CE} and $d\mu^2$) and the genetic distances of Tamura and Nei (1993) calculated for the mtDNA data. Data on the mtDNA *cyt b* were available for the populations. First, the pairwise F_{ST} matrices were correlated by using simple Mantel procedures (Mantel, 1967) in the Vegan library in R (Oksanen *et al.*, 2007). Permutations ($n=10\,000$) were used to evaluate statistical significance. Then, two-dimensional multidimensional scaling analysis ordinations of the two types of genetic marker were compared by Procrustes analysis

(Gower, 1975) using R software. Procrustes analysis is searching for the best match between two configurations of points in a multivariate Euclidean space using rotation, translation, reflection and dilation of one configuration. The criterion used to assess the best fit is the minimization of the sum of squares between the differences for each observation (m^2). The significance of the result, an optimal superposition of one configuration on the other (reference), is obtained through a permutation test (PROTEST) (Jackson, 1995). PROTEST is using $R = \sqrt{1 - m^2}$ as a test statistic, which can be interpreted as a correlation.

Impact of selection, sex-biased dispersal and effective population size on the degree of population divergence estimates

To verify if the variation on the microsatellites can be influenced by selection, two different approaches were considered. First, all microsatellite flanking regions were compared with sequences in GenBank by means of the BLAST program (<http://www.ncbi.nlm.nih.gov/BLAST/>) (Altschul *et al.*, 1990) to verify if the microsatellites potentially are located within the described functional regions of the genome. Second, potential outlier microsatellites were identified by using the selection detection workbenches LOSITAN (Antao *et al.*, 2008) and BayeScan (Foll and Gaggiotti, 2008). Analyses are performed for LOSITAN with 10 000 simulations for IAM and SMM, both with the options 'Neutral' mean F_{ST} and force mean F_{ST} . For BayeScan, 10 000 iterations were conducted with a thinning interval of 20 and with 10 pilot runs.

To detect differences in migration rates between females and males, deviation from the Hardy–Weinberg equilibrium (F_{IS}), differentiation among populations (F_{ST}), relatedness (r) (Queller and Goodnight, 1989), mean assignment index (mAI_C) and variance of the assignment index (vAI_C) were quantified separately for both sexes over all populations (Goudet *et al.*, 2002). Statistical significance of differences in these within-population indices was determined with 10 000 permutations using the randomization method implemented in FSTAT (Goudet, 2001).

Effective population sizes were estimated using different point methods, which do not require samples spaced over at least one temporal interval, and temporal methods, where samples are taken from the same location at two or more points in time separated by a specified number of generations (one or more). Two different point estimation methods were estimated for the 10 samples. The program NeEstimator v.1.3 (Peel *et al.*, 2004) was used for the Heterozygote Excess method, which examines the excess of heterozygotes in the sample compared with the proportion predicted under the Hardy–Weinberg equilibrium (Luikart and Cornuet, 1999). For the estimation of effective population size on the basis of linkage disequilibrium data, the program LDNe v.1.31 (Waples and Do, 2008) was used because it implements a recently developed bias correction (Waples, 2006). Finally, N_e was estimated using two temporal methods in NeEstimator v.1.3: (i) based on the Moment Approach (Waples, 1989) and (ii) on the pseudo-likelihood method (MLNE) developed by Wang and Whitlock (2003). These two methods require at least two temporally spaced samples, and therefore this analysis was only possible for VMS and BNS (Table 1).

Results

Genetic diversity

Mean allelic richness per location, corrected for sample size, varied between 12.088 (northern Baltic Sea) and 15.454 (MS) (mean = 13.622) (Table 2). Mean expected heterozygosity was relatively uniform among the different sampling sites with the lowest value in the northern Baltic Sea sample (0.740) and the highest in the MS samples (0.781) (mean = 0.756) (Table 2). Locations WCS and BNS2 showed a significant departure from the Hardy–Weinberg equilibrium (Table 2) (both multilocus $F_{IS} = 0.050$). Pairwise comparisons between loci revealed no significant linkage disequilibrium after sequential Bonferroni corrections.

Patterns of population subdivision

First, the graphical distribution of populations from the FCA showed that the two Mediterranean samples (VMS1 and VMS2), as well as all Atlantic samples, except the one from the IB (AAO), clustered together (Figure 2). Disregarding the AAO sample, the sample of the Bay of Biscay (GOA) was the most aberrant among the Atlantic samples. The distribution of all individuals in a FCA plot (graph not shown) shows a small overlap between the Mediterranean and Atlantic samples as well as between AAO and the other Atlantic samples.

Second, the global θ - and ρ -values across all samples, excluding the temporal samples (BNS2 and VMS2), were 0.0121 and 0.0426, respectively (both highly significant, $P < 0.001$). The global standardized multilocus F_{ST} was 0.0484. The pairwise F_{ST} values were significant after sequential Bonferroni correction and false discovery rate control for all population pairs including samples TBS, VMS and AAO, together with a limited number of population pairs between Atlantic populations (Table 3). Pairwise R_{ST} values were significant after sequential Bonferroni correction or false discovery rate control for several population pairs between VMS and Atlantic populations (Table 3). The pairwise F_{ST} and R_{ST} values between the two temporal samples were not statistically different ($P > 0.05$) for BNS (0.0038 and 0.0005, respectively) and VMS (−0.0012 and −0.0025, respectively) (Table 3). The pairwise standardized F_{ST} values between the different samples are listed in Table 4. Jackknife

Table 2 Estimates of genetic diversity of the 10 samples of *P. minutus* based on eight microsatellite markers

Population	Sample size	$H_{E \text{ n.b.}}$	H_O	F_{IS}	Allelic richness
TBS	96	0.740	0.718	0.030	12.088
WCS	88	0.760	0.722	0.050	13.211
RNS	48	0.752	0.755	−0.003	12.735
BNS1	47	0.749	0.689	0.081	12.319
BNS2	95	0.762	0.724	0.050	13.404
WIS	45	0.754	0.724	0.040	13.534
GOA	40	0.752	0.732	0.027	14.087
AAO	95	0.752	0.714	0.051	14.015
VMS1	94	0.781	0.766	0.019	15.454
VMS2	48	0.766	0.723	0.057	15.375

$H_{E \text{ n.b.}}$ is the unbiased expected heterozygosity, H_O the observed heterozygosity and F_{IS} measures deviation from Hardy–Weinberg equilibrium. Statistically significant F_{IS} values are listed in bold. See Table 1 for sample codes.

analysis revealed that locus Pmin16-2 was responsible for the largest divergence as calculated with R_{ST} , but not with F_{ST} (average pairwise $R_{ST} = 0.02974$ with locus Pmin16-2 and $R_{ST} = 0.00528$ without this locus). Random permutation of different allele sizes among allelic states at each locus revealed that estimates of R_{ST} were significantly larger than the 95% CI range of the pR_{ST} values at one single locus Pmin16-2 (Table 5), suggesting

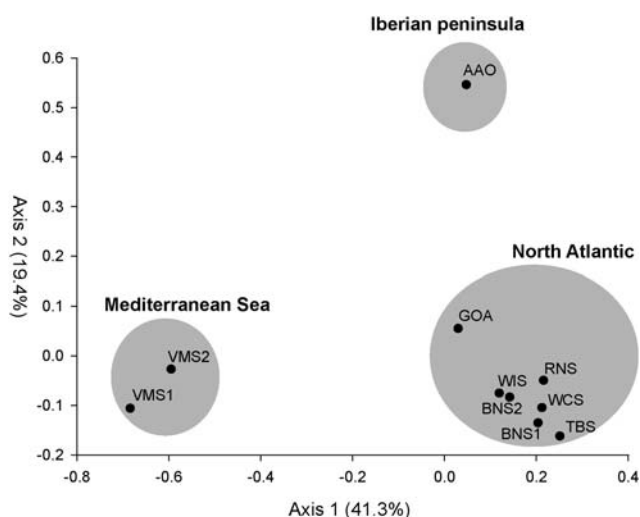


Figure 2 Factorial correspondence analysis plot based on microsatellite data of the 10 samples of *P. minutus*. See Table 1 for sample codes.

a mutational component to genetic differentiation. For the other microsatellites, neither loci allele size nor stepwise mutations strongly influence the population differentiation of sand gobies. Hardy *et al.* (2003) suggest that, in this situation, F_{ST} should be preferred over R_{ST} for estimating population differentiation.

Third, the classical multidimensional scaling analyses plots based on both types of genetic distances separated clearly the group of the two Mediterranean samples from the populations of the Atlantic (Figure 3). Both plots had a stress value below 0.20, suggesting interpretable information concerning intersite relationships.

Fourth, the global Mantel test revealed a significant isolation by distance pattern with D_{CE} ($r = 0.713$, $P < 0.05$)

Table 5 Mean single locus and multilocus pairwise estimates of F_{ST} , R_{ST} , pR_{ST} (95% distribution of central values in parentheses) between the 10 samples of sand gobies following 2000 permutations of the microsatellite alleles (Hardy *et al.*, 2003)

	F_{ST}	R_{ST}	pR_{ST} (95% range)
Multilocus	0.012	0.042*	0.011 (0.002–0.024)
Pmin09	0.008	0.014	0.008 (–0.004 to 0.029)
Pmin16-2	0.028	0.225*	0.027 (–0.002 to 0.103)
Pmin35	0.016	0.001	0.016 (–0.001 to 0.045)
Pmin38	–0.002	0.003	–0.002 (–0.004 to 0.003)
Pmin29	0.009	0.021	0.010 (–0.003 to 0.031)
Pmin31	0.005	–0.004	0.005 (–0.004 to 0.021)
Pmin03	0.006	0.006	0.006 (–0.004 to 0.026)
Pmin04	0.007	0.008	0.007 (–0.003 to 0.022)

* Indicates statistically significant values ($P < 0.01$).

Table 3 Pairwise F_{ST} (below diagonal) and R_{ST} (above diagonal) values of the *P. minutus* samples based on eight microsatellite markers

	TBS	WCS	RNS	BNS1	BNS2	WIS	GOA	AAO	VMS1	VMS2
TBS	—	0.017	0.012	0.003	0.023	0.017	0.044	0.018	0.118	0.118
WCS	0.007*	—	–0.002	–0.002	0.006	0.004	0.013	0.001	0.093	0.085
RNS	0.009*	0.002	—	0.004	0.014	–0.001	0.008	0.005	0.069	0.066*
BNS1	0.010*	0.003	0.001	—	0.001	0.008	0.022	0.004	0.095*	0.090
BNS2	0.007*	0.004*	0.001	0.004	—	–0.001	0.0061	0.002	0.081	0.068
WIS	0.013*	0.012*	0.007*	0.004	0.006	—	–0.004	0.001	0.055*	0.047
GOA	0.013*	0.008*	0.010*	0.004	0.005*	0.009*	—	0.002	0.042*	0.027
AAO	0.012*	0.010*	0.009*	0.012*	0.008*	0.010*	0.013*	—	0.086	0.071
VMS1	0.025*	0.017*	0.019*	0.018*	0.017*	0.015*	0.015*	0.017*	—	–0.003
VMS2	0.018*	0.011*	0.013*	0.013*	0.012*	0.014*	0.013*	0.012*	–0.001	—

Statistically significant values are listed in bold after sequential Bonferroni correction and with * after the false discovery rate control. See Table 1 for sample codes.

Table 4 Standardised pairwise F_{ST} estimates (Hedrick, 2005) based on the microsatellite data (below diagonal) and the F_{ST} estimated from mitochondrial cyt *b* sequence data (above diagonal) between the eight locations for *P. minutus*

	TBS	WCS	RNS	BNS1	BNS2	WIS	GOA	AAO	VMS1	VMS2
TBS	—	0.165	0.196	0.243	—	0.384	0.257	0.659	0.688	—
WCS	0.027	—	0.093	0.133	—	0.036	0.015	0.616	0.650	—
RNS	0.024	0.001	—	–0.049	—	0.245	0.086	0.556	0.645	—
BNS1	0.030	0.010	0.002	—	—	0.330	0.115	0.594	0.627	—
BNS2	0.024	0.020	0.002	0.019	—	—	—	—	—	—
WIS	0.032	0.047	0.018	0.019	0.021	—	0.064	0.709	0.724	—
GOA	0.050	0.042	0.031	0.014	0.030	0.034	—	0.579	0.661	—
AAO	0.045	0.043	0.042	0.054	0.037	0.038	0.055	—	0.756	—
VMS1	0.053	0.041	0.040	0.053	0.049	0.053	0.042	0.044	—	—
VMS2	0.036	0.025	0.029	0.043	0.037	0.055	0.036	0.029	–0.006	—

Mitochondrial divergence estimates were calculated as described by Larmuseau *et al.* (2009b).

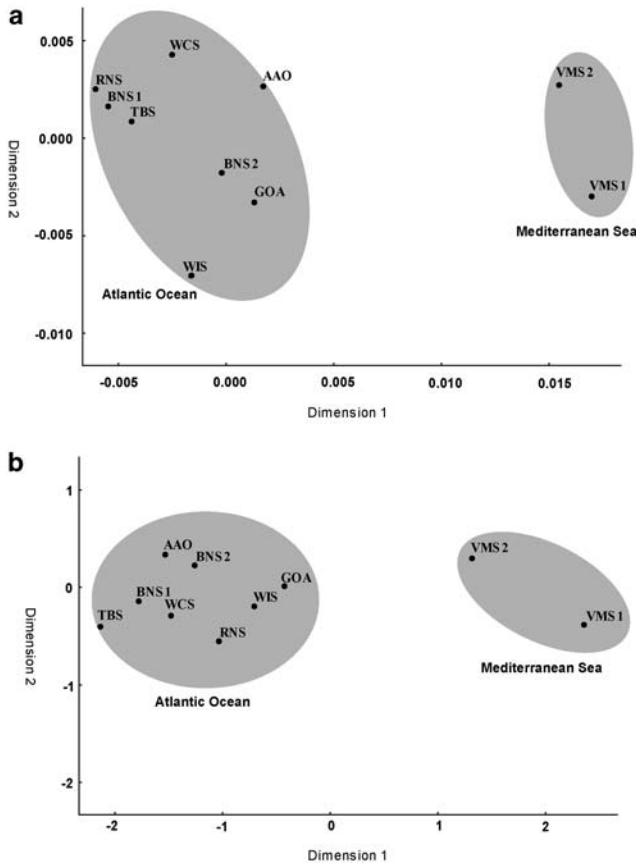


Figure 3 Classical multidimensional scaling plots of pairwise genetic distance for the microsatellite data calculated according to (a) Cavalli-Sforza and Edwards (1967) (D_{CE}) and (b) Goldstein *et al.* (1995) ($d\mu^2$) among the 10 samples of *P. minutus*. See Table 1 for sample codes.

but not with $d\mu^2$ ($r=0.166$, $P>0.05$). However, no significant isolation by distance was found for both types of genetic distances when the MS and/or IB samples were excluded from the data set.

Fifth, testing the significance of the stepwise clustering procedure performed in STRUCTURE resulted in separation of the samples into three hypothetical clusters (highest ΔK for $K=3$). The lowest proportion for a sample of membership to a particular cluster is 0.411 in cluster 1 for sample WCS and the highest proportion is 0.897 in cluster 2 for sample TBS (Table 6). All Atlantic samples (excluding TBS) had the highest assignment value for cluster 1; the northern Baltic sample (TBS) for cluster 2 and the two Mediterranean samples for cluster 3 (Table 6 and Figure 4).

Finally, the Bayesian approach used in 2MOD indicated that *P. minutus* populations were at migration-drift equilibrium. All sampled iterations revealed that a migration-drift equilibrium model was more likely than the non-equilibrium drift model.

Comparison between types of genetic distances and mtDNA

F_{ST} -values are much lower for microsatellites (global $F_{ST}=0.0121$ (CI=0.0066–0.0174) and global standardized $F_{ST}=0.0484$ (CI=0.0264–0.0696), both without the

Table 6 Summary of the assignment analysis following STRUCTURE v. 2.2 based on microsatellite data of 10 *P. minutus* samples

Population	Inferred clusters		
	1	2	3
TBS	0.059	0.897	0.044
WCS	0.411	0.388	0.201
RNS	0.514	0.295	0.191
BNS1	0.510	0.336	0.154
BNS2	0.442	0.324	0.234
WIS	0.516	0.236	0.248
GOA	0.448	0.226	0.326
AAO	0.492	0.154	0.353
VMS1	0.125	0.057	0.819
VMS2	0.160	0.079	0.762

The proportion of individuals assigned to the three hypothetical clusters is given. The highest assignment value for each population is listed in bold. See Table 1 for sample codes.

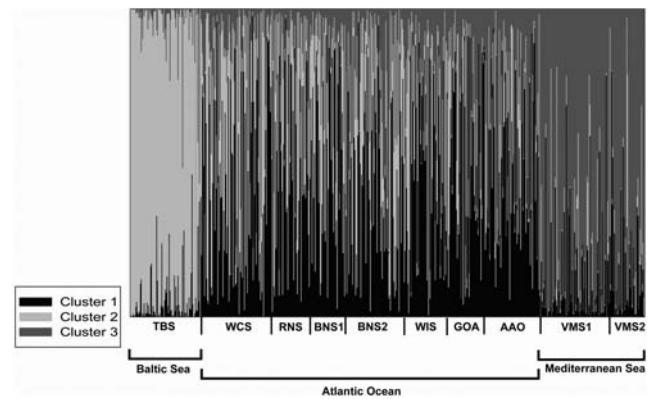


Figure 4 Results of the clustering analysis conducted in STRUCTURE 2.2 (Pritchard *et al.*, 2000) based on the microsatellite data. In the bar plot, each of the 696 individuals is represented by a vertical bar indicating its estimated proportion of membership to the three clusters ($K=3$). See Table 1 for sample codes.

temporal samples VMS2 and BNS2) than for mtDNA (global $F_{ST}=0.4293$ (CI=0.3231–0.5302)). The proportion of differences in F_{ST} between mtDNA and nuclear microsatellites is not attributed to stochastic variability among loci as there were no notable differences between different microsatellite loci. The highest global θ was found for the locus Pmin 16-2 (0.0314); the lowest θ -value was found for the locus Pmin 38 (–0.0023). Pairwise F_{ST} values between all samples based on the mtDNA data are presented together with the standardized pairwise F_{ST} estimates for microsatellites in Table 4. Each pairwise F_{ST} value for mtDNA is on average 10 times higher than the value for microsatellites. The pairwise genetic distances of the microsatellite data ($d\mu^2$ and D_{CE}) correlated well with each other (Procrustes $R=0.8339$, P -value=0.004; Mantel $R=0.8574$, P -value <0.001). Procrustes analyses and Mantel tests also showed that single-locus $d\mu^2$ and D_{CE} correlated well with the estimates based on all microsatellites. The D_{CE} pairwise genetic distances of the microsatellite data correlated well with Tamura and Nei (1993) genetic distances based on the mtDNA data (Procrustes $R=0.746$, P -value <0.001; Mantel $R=0.6804$, P -value=0.032). In contrast, the $d\mu^2$ pairwise genetic distances of the microsatellite data did not correlate

significantly with the distances based on the mtDNA data (Procrustes $R = 0.660$, P -value = 0.086; Mantel $R = 0.645$, P -value = 0.081).

Impact of selection, sex-biased dispersal and effective population size on the degree of population divergence estimates

No indication for selection for some of the microsatellite markers was observed in the analysis. The results of Lositan and BayeScan were congruent with this pattern by revealing no outlier loci. Moreover, a blast search of the flanking regions of the microsatellite loci in de GenBank database revealed that they are not linked to a functional region already sequenced for any organism available in GenBank.

A total of 352 males and 344 females were genotyped and the population genetic parameters were estimated separately for each sex. No parameter was significantly different between the sexes (Table 7), suggesting no sex-biased dispersal.

Estimates from the various point and temporal methods to estimate the N_e are not congruent with each other (Table 8). Both point methods cannot exclude the possibility that the population sizes are infinite. The two temporal methods are similar for the BNS population (~400 individuals) and for VMS (> 1000 individuals).

Discussion

Phylogeography of the sand goby

The current study with microsatellites revealed a subtle but significant genetic structure within *P. minutus*. The

highest divergence in the analysis was found between populations of the Atlantic, comprising all sand goby populations spanning from the IB to the Baltic Sea, and the MS, represented by individuals from the Vaccarès lagoon (Gulf of Lion) (Table 3 and Figures 2–4). The microsatellite marker Pmin16-2 shows a clear differentiation with a small overlap in allele size between the MS-Clade and AO-Clade, clearly indicating the historical divergence of both regions. In addition, the highest divergence noticed in the genetic analysis of allozyme markers were also found between Mediterranean and Atlantic sand gobies (Stefanni *et al.*, 2003; Larmuseau *et al.*, 2009b). The results are also congruent with the phylogenetic analysis based on mtDNA *cyt b*, which clearly showed two monophyletic clades within *P. minutus*, the Mediterranean Sea Clade (MS-Clade) and the AO-Clade (Larmuseau *et al.*, 2009b). Divergence between the two sand goby clades is most likely the result of geographic isolation caused by one of the Middle Pleistocene sea-level drops (Larmuseau *et al.*, 2009b). The Mediterranean populations showed the highest variability in the microsatellite markers (Table 2) despite the small distribution range along the north-western MS coasts and lagoons (Miller, 1986) (Figure 1). Larmuseau *et al.* (2009b) noticed the highest diversity in mtDNA in the Mediterranean samples as well. Furthermore, the highest species diversity of the ‘sand goby’ group is observed within the MS (Huyse *et al.*, 2004) supporting a Mediterranean origin of *P. minutus*, as suggested by Gysels *et al.* (2004a). Huyse *et al.* (2004) estimated the origin of the species between 1.94 and 1.18 mya (early Pleistocene). The Pleistocene glaciations were the most significant historical events during the evolutionary lifespan of most Holarctic species and are believed to have accelerated the speciation process in the present day sister taxa (Avice, 2000).

Limited genetic differentiation was observed between the Atlantic samples with microsatellites, but the FCA and the pairwise F_{ST} values suggest that the IB’s sample is different from those of all other Atlantic populations (Figure 2 and Table 3). These results are congruent with the two mitochondrial groups within the Atlantic samples: the Iberian (IB-Group) and the North Atlantic Group. This divergence is likely the result of the population decline within different refugia during glaciations in the Middle Pleistocene (Larmuseau *et al.*,

Table 7 F-statistics, relatedness (r), mean assignment (mAI_C) and variance assignment (vAI_C) for each sex based on the microsatellite data. Significance (P) was assessed using the randomization method of Goudet *et al.* (2002)

Sex	F_{IS}	F_{ST}	R	mAI_C	vAI_C
Males	0.040	0.011	0.022	-0.078	8.643
Females	0.042	0.009	0.017	0.080	9.819
P	0.890	0.380	0.378	0.478	0.255

Table 8 Estimates and the 95% confidence intervals (CIs) of the effective population sizes (N_e) of the populations of *P. minutus* using point methods (heterozygote excess and linkage disequilibrium) and temporal methods (moment based approach and MLNE) on the microsatellite data

Populations	Heterozygote excess		Linkage disequilibrium		Moments based approach		MLNE	
	Estimate	95% CI	Estimate	95% CI	Estimate	95% CI	Estimate	95% CI
TBS	Infinity	NA	724.3	324.5–infinity				
WCS	9927	NA	Infinity	78350.4–infinity				
RNS	145.7	NA	Infinity	421.2–infinity				
BNS1	15.9	NA	Infinity	-729.2–infinity				
BNS2	Infinity	NA	1159.9	245.1–infinity	470	192.5–infinity	361.9	187.4–1504
WIS	Infinity	NA	196.8	115.9–557.7				
GOA	Infinity	NA	2146.3	247.3–infinity				
AAO	Infinity	NA	547.1	297.8–2505.8				
VMS1	13.5	NA	Infinity	1014.3–infinity				
VMS2	Infinity	NA	1413.3	291.5–infinity	Infinity	113.3–infinity	1651.3	161.3–infinity

The temporal methods were only estimated for the BNS and VMS locations. See Table 1 for sample codes.

2009b). On the basis of the Bayesian assignment test and pairwise F_{ST} values, the position for the sample of the northern Baltic Sea (TBS) was remarkable (Tables 3, 6 and Figure 4). This is most likely the result of a founder event, based on the low variation on the microsatellite loci in this population (Table 2) and as suggested by Larmuseau *et al.* (2009b) based on the mtDNA data. The difference in the *cyt b* haplotype network between the southern and northern Baltic Sea samples suggested that the Baltic Sea has been colonized in two phases over a period of 8000 years, with a stronger founder effect in the north. The presence of only two common *cyt b* haplotypes in the northern samples HBS and TBS (haplotypes NA01 and NA28) with their many uniquely derived haplotypes suggested that only a few individuals founded the northern Baltic Sea (Johannesson and André, 2006). A recent study has already shown that sand gobies are locally adapted to the specific light environment of the northern Baltic Sea in comparison with the sand gobies of the North Sea and the AO (Larmuseau *et al.*, 2009a). If only a few sand gobies colonized the northern Baltic, the populations in this region will be genetically diverged from the founding population, mainly through a shift in allele frequencies. Such shifts are detectable with pairwise F_{ST} values and the STRUCTURE analysis (Table 3 and Figure 4). In contrast, individual multilocus genotypes are less sensitive for shifts in allele frequencies, and analyses based on multilocus genotypes, such as the FCA plot (Figure 2), will not be able to distinguish between different groups after a founding event.

The Mantel test and Procrustes analysis illustrated that the microsatellite results are congruent with the mtDNA phylogeographic pattern. However, those tests were not significant when the $d\mu^2$ genetic distance was used for microsatellites. This suggests that, although both measures are correlated, F_{ST} correlates better with mtDNA than R_{ST} . In theory, F_{ST} is more sensitive than R_{ST} for recent intraspecific divergence (Gaggiotti *et al.*, 1999; Balloux and Lugon-Moulin, 2002). Moreover, R_{ST} can be less accurate in reflecting population differentiation because of its higher associated variances (Balloux and Lugon-Moulin, 2002). Therefore, the number of loci screened has to increase before a consistent pattern is reached (Gaggiotti *et al.*, 1999; Balloux and Goudet, 2002). This has already been empirically observed with European grayling *Thymallus thymallus* (Koskinen *et al.*, 2004).

Overall, the results confirm the first hypothesis of this study that stated that microsatellite markers reveal a phylogeographic pattern congruent with the patterns based on mtDNA and allozyme data.

Differences in the degree of divergence estimates among nuclear and mtDNA markers

The difference in the degree of population differentiation between the nuclear and mitochondrial markers is remarkable. The mitochondrial differentiation values are more than an order of magnitude higher than the nuclear differentiation, even after standardization of the microsatellite results. The standardized measure allows comparison between loci with different levels of genetic variation (Hedrick, 2005). Therefore, several other factors can be the cause of the observed discrepancy.

First, selection may have differential effects on genetic markers. It has been suggested that balancing selection may significantly influence the distribution of allozyme diversity (De Innocentiis *et al.*, 2001). Loci experiencing balancing selection will have allele frequencies more similar than expected under neutrality, reducing the F_{ST} estimates. Allendorf and Seeb (2000) concluded that estimates of population structure produced by allozymes were generally comparable to those obtained with other nuclear markers, including microsatellites. They noted that when differences between marker classes did occur, they were usually due to one or a few exceptional loci and not all of them. Each used microsatellite marker was tested for positive or balanced selection in comparison with the other markers in the Lositan and BayeScan analysis, but no marker seemed to be an outlier. Similar to microsatellites, it is also not safe to assume *a priori* that mtDNA evolves as a strictly neutral marker (Ballard and Whitlock, 2004). Selection on mtDNA may accelerate the coalescence of lineages, and thus increase the levels of differentiation observed between populations (Peijnenburg *et al.*, 2006). The observation of very low nuclear differentiation in sand gobies while mtDNA data revealed no gene flow between the three isolated Middle Pleistocene lineages cannot be explained by selective evolution for mtDNA. The rejection of the null hypothesis in different neutrality tests for the mtDNA data of *P. minutus* was assigned to demographical expansions instead of selection (Larmuseau *et al.*, 2009b). Various demographic analyses on the mtDNA data showed that the intra-assembly genetic structure of *P. minutus* contains signatures of demographic expansion events.

Second, studies documenting a weaker population subdivision for nuclear than maternally inherited genetic markers often attribute these discrepancies to male-biased dispersal. Sex-biased dispersal is common in nature (Cano *et al.*, 2008); however, it has only been described for a limited number of marine fishes (Consuegra and de Leaniz, 2007). No indications for sex-biased dispersal were found for *P. minutus* with microsatellite markers (Table 7). Contemporary gene flow between the Mediterranean and Atlantic basins is also unlikely because of the discontinuous distribution of the sand goby (Miller, 1986) (Figure 1). Moreover, only males have to migrate successfully to explain the pattern. However, females are expected to be the most mobile sex, especially during the spawning period when males are guarding their nest (Lindström *et al.*, 2006).

Third, various technical problems, including homoplasy, may have reduced the signal of differentiation detected by the microsatellite markers. Homoplasy occurs when different copies of a locus are identical in state, although not identical by descent. The situations where size homoplasy is most prevalent involve high mutation rates and large population sizes together with strong allele size constraints (Estoup *et al.*, 2002). Therefore, effects of homoplasy are expected to be common for microsatellites in marine fishes (O'Reilly *et al.*, 2004), which has implications for the identification of genetic structuring (Carreras-Carbonell *et al.*, 2006). Microsatellites probably suffered from higher levels of homoplasy than mtDNA because of higher mutation rates and larger effective population sizes (Balloux *et al.*, 2000). The various microsatellite markers of *P. minutus* indicate homoplasy because of the high allele numbers and the

limited size range of all markers (on average 1 allele per 2,053 bp). Single-nucleotide polymorphisms and microsatellite markers with less alleles covering a limited size range might be useful to understand the power of homoplasmy in the analysis. However, simulation studies suggest that size homoplasmy will have much less effect on estimates of population differentiation than gene migration or genetic drift (Estoup *et al.*, 2002). Therefore, it is unlikely that homoplasmy is the main cause of the observed differences.

Finally, mitochondrial markers can be more sensitive in detecting differentiation because of a lower effective population size than nuclear markers (Shaw *et al.*, 2004). Genetic drift effects are linked to effective population size (N_e), and therefore it is possible that an ecologically relevant population structure remains undetectable by using neutral markers when the N_e is high (Bentzen, 1998). Marine fish have the potential to have high N_e , and therefore recently separated large populations may appear genetically homogeneous even in the complete absence of contemporary gene flow (Hauser and Carvalho, 2008). Simulations in the study by Buonaccorsi *et al.* (2001) showed that differences in the magnitude of estimated population subdivision from nuclear and mitochondrial markers could be accounted for entirely by differences in effective population sizes and polymorphisms on F_{ST} estimates. The haploid and maternal-only inheritance of mtDNA has an effective population size of one-quarter that of nuclear DNA, making it more susceptible to effects of genetic drift (Shaw *et al.*, 2004). This explanation has been invoked for the discordant patterns in population differentiation between nuclear and mitochondrial markers for marine organisms, such as the blue marlin *Makaira nigricans* (Buonaccorsi *et al.*, 2001), Patagonian toothfish *Dissostichus eleginoides* (Appleyard *et al.*, 2002; Shaw *et al.*, 2004) and the olive sea snake *Aipysurus laevis* (Lukoschek *et al.*, 2008). Simulations in EASYPOP v.1.7 (Balloux, 2001) with specific biological information about the sand goby showed no difference with the simulations of the study by Buonaccorsi *et al.* (2001) (results not shown). Moreover, *P. minutus* is known to be one of the most abundant fish species across almost its full range (Pasquaud *et al.*, 2004; Ehrenberg *et al.*, 2005; Maes *et al.*, 2005). Therefore, high N_e most likely explains the discordant patterns between nuclear and mitochondrial data for the sand goby. Point methods to estimate the N_e of the present sampled populations cannot invalidate the null hypothesis of an infinite population size for *P. minutus* (Table 8). Nevertheless, point methods are not always reliable and are biased (Wang and Whitlock, 2003), especially when the sample size is small (<100 individuals) and below the true N_e (England *et al.*, 2006). Both temporal N_e estimates show a limited population size, especially for the BNS location (100–1000 individuals) (Table 8). Still, there are crucial differences between the two temporal methods and the assumption of a closed system without migration could not be fulfilled. Migration inside the marine system can therefore cause a strong underestimation of the N_e (Wang and Whitlock, 2003). On the other hand, the high number of alleles for microsatellites confirms the hypothesis of a high effective population size of more than one thousand individuals to maintain the high genetic variation (Ewens, 1972; Poulsen *et al.*, 2006). The Pmin20 locus was excluded from the analysis because,

after genotyping more than 90 individuals, the number of alleles almost matched the number of fish genotyped. The studies of Jones *et al.* (2001) and Pampoulie *et al.* (2004) also observed microsatellite markers with a very high number of alleles. By organizing a better sampling strategy and using straightforward N_e -estimation methods incorporating migration (Wang and Whitlock, 2003), better estimates of the N_e might confirm the expected high N_e for *P. minutus*.

Our present results do not confirm the second hypothesis, stating that microsatellite markers are more sensitive for population differentiation on a macroscale than mitochondrial and allozyme markers. The most likely reason for the discordant pattern between nuclear and mitochondrial loci is that the recent demographical expansion in *P. minutus* resulted in high effective population sizes, slowing down the differentiation at nuclear loci.

Conclusion

Marine organisms have a high potential for gene flow and population size. However, it is not known whether the observed low genetic differentiation for so many marine species reflects high effective population sizes and low gene flow, high effective population sizes and high rates of gene flow, or low effective population sizes and high rates of gene flow (Hauser and Carvalho, 2008). In this study on *P. minutus*, the scenario of high effective population sizes and low gene flow best explains the observed genetic pattern. However, this conclusion could only be reached with more than one genetic marker. Therefore, one has to be cautious when interpreting the present-day genetic structure in terms of gene flow while using one type of marker or statistical method.

Conflict of interest

The authors declare no conflict of interest.

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