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Potential and realized connectivity of the seagrass *Posidonia oceanica* and their implication for conservation

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Abstract

Aim: Connectivity assessments are crucial to large-scale conservation planning, in particular for establishing and monitoring connected networks of marine protected areas (MPAs). Using biophysical modelling and genetic analyses, we assessed potential and realized connectivity among MPA populations of a benthic foundation species, the Mediterranean endemic seagrass *Posidonia oceanica*.

Location: Adriatic and Ionian seas (central Mediterranean).

Methods: We assessed potential and realized connectivity among eight *P. oceanica* meadows, mostly located in MPAs. Potential connectivity was assessed over a time horizon of 10 years via an individual-based biophysical model whose physical component relies on fine-scale spatio-temporal ocean circulation fields. Genetic assessments of realized connectivity were carried out by means of a set of 14 neutral microsatellite loci, as well as a larger dataset of 19 loci including outlier loci that did not conform to expectations under neutrality.

Results: Our findings point out a relatively high potential connectivity through longrange dispersal of floating fruits. Genetic connectivity analyses show a complex scenario with an apparent lower realized connectivity. The *P. oceanica* meadow within Torre Guaceto MPA (TOG), a well-enforced MPA within our study area, showed one of the highest levels of genotypic richness, indicative of high levels of sexual reproduction and/or recruitment of foreign genotypes. Both biophysical modelling and population genetics indicate that TOG is important to ensure the viability of the species at the local scale, and does likely play a key role as a source of propagules for the whole Adriatic area.

Main conclusions: Our results show that realized dispersal does not necessarily match with the potential for dispersal. Still, both genetic and physical connectivity analyses show good agreement in identifying hotspots of connectivity. Such information can guide management of networks of MPAs and advance conservation of marine biodiversity.

KEYWORDS

dispersal, genetic connectivity, Lagrangian, marine protected areas, propagules, seagrass

1 | INTRODUCTION

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Spatial structuring is common in the marine environment and may often favour local adaptation (Conover, Clarke, Munch, & Wagner, 2006; Palumbi, 2004; Sanford & Kelly, 2011). This is an important issue in conservation, as it supports the design of marine protected areas (MPAs) in a way that ensures seascape connectivity, so that connected networks of MPAs can effectively sustain the persistence, recovery and productivity of marine ecosystems (McCook et al., 2009). For instance, to enable recovery of protected coral populations after a disturbance within an MPA, the potential sources of replenishing larvae also need to be protected (Underwood, Smith, Van Oppen, & Gilmour, 2007). The lack of obvious physical barriers makes the marine environment an especially good case for studying adaptation in the face of gene flow. It provides an opportunity to investigate the interaction between the diversifying effects of selection and the counteracting, homogenizing effects of gene flow (Cristescu, Constantin, Bock, Caceres, & Crease, 2012; Nosil, 2009; Räsänen & Hendry, 2008). Realized connectivity, or effective gene flow, depends on the interaction between oceanographic features, species-specific life history traits affecting dispersal, habitat availability and population demography. It can be measured by genetic approaches (Galindo et al., 2010; White et al., 2010) and complemented by assessment of potential for connectivity via individual-based biophysical models (Cowen & Sponaugle, 2009; Gallego, North, & Petitgas, 2007).

The increasingly recognized importance of connectivity is also reflected in the Aichi target 11 of the Convention for Biological Diversity (CBD), aimed at implementing a "well-connected system of protected areas" by 2020. Despite the increasing awareness of the importance of connectivity for MPA design, few studies have assessed connectivity among MPAs (but see for instance Christie et al., 2010; Hogan, Thiessen, Sale, & Heath, 2012; Planes, Jones, & Thorrold, 2009). Moreover, only very few MPA design processes have incorporated connectivity into planning (among them Beger et al., 2015; Palumbi, 2003; Weeks et al., 2014). It is in fact difficult to include information about connectivity in MPA and marine spatial planning algorithms (Beger et al., 2010). A major issue is the inherent problem that connectivity assessments are usually carried out for single species (yet not exclusively, see López-Duarte et al., 2012; Magris, Treml, Pressey, & Weeks, 2015; Melià et al., 2016 for some multispecies studies). Considering the crucial role that species-specific or population-specific demographic processes play in shaping connectivity and environmentdependent dispersal processes, focusing on a single species appears to be restrictive. However, selecting umbrella species (defined here as species with an especially important role in the investigated ecosystem, e.g., ecosystem engineers) can represent a good compromise between limiting assessment efforts and emphasizing the importance of the ecosystem (Hughes & Stachowicz, 2009).

Seagrass meadows are considered one of the most highly impacted coastal ecosystems on Earth (Duarte, Dennison, Orth, & Carruthers, 2008). Habitat loss is a major threat to seagrasses, causing increase in fragmentation of populations (Marbà, Díaz-Almela, & Duarte, 2014), whose dispersal is mainly dependent on floating shoots or seeds that, at least for some species, have a low dispersal capacity (McMahon et al., 2014). Seagrasses are also important ecosystem engineers that provide crucial ecosystem services, such as reducing wave impact, stabilizing the sediment, adding oxygen to the water, providing nursery grounds and shelter for many species (including commercially important species), exporting important amounts of carbon, nitrogen and phosphorus to coastal food webs, stocking significant amounts of organic carbon and reducing exposure to bacterial pathogens (Beck et al., 2001; Costanza et al., 2014; Duffy & Stachowicz, 2006; Heck, Hays, & Orth, 2003; Lamb et al., 2017). Ensuring connectivity of such ecologically important habitat formers is thus crucial, given the major decline of seagrasses worldwide (Short et al., 2011) with important cascading effects on the associated ecosystems (Healey & Hovel, 2004; Warry, Hindell, Macreadie, Jenkins, & Connolly, 2009). Establishing networks of suitably spaced and connected MPAs is possibly the best way to maintain effective connectivity and sustain levels of gene flow that can avoid inbreeding and allow the spread of advantageous alleles.

In this study, we focus on the Mediterranean endemic seagrass Posidonia oceanica, which has experienced severe habitat loss and population fragmentation over the last decades to centuries (Marbà et al., 2014; Short et al., 2011). Our research integrates connectivity assessments based on numerical simulations of the movement of sexual propagules based on oceanographic fields forced with atmospheric data and genetic analyses: the combination of these two independent approaches provides complementary information about potential and realized connectivity of P. oceanica at regional levels. We sampled eight sites of P. oceanica mostly located in MPAs encompassing five countries in the Adriatic and Ionian seas. Previous studies in the area focused mainly on mobile species (for instance, Schiavina, Marino, Zane, & Melià, 2014 on the Mediterranean shore crab, Boissin et al., 2016 on the Black scorpionfish and Carreras et al., 2017 on the peacock wrasse), showing either a N-S (crab) or a W-E discontinuity (scorpionfish), or a mixture of both (wrasse). Here we assess a foundation species and aim to determine the extent to which the selected Adriatic and Ionian populations of P. oceanica may be connected (based on neutral genetic markers and Lagrangian simulations)-given current environmental conditions and demographic processes affecting the different populations. Specifically, we address the following questions: (1) What is the level of potential connectivity, based on biophysical modelling? (2) What is the level of realized connectivity, based on genetic differentiation and assignment test? (3) How do the potential for connectivity and realized connectivity compare? Finally, we discuss our findings in the context of regional conservation management, giving important insights in the definition of management plans and MPA network design that extend beyond our case study.

2 | METHODS

2.1 | Sampling

We collected individuals of *P. oceanica* at eight sites in the Adriatic and lonian seas in five different countries during spring 2013 (Figure 1). Most populations were sampled within MPAs (see Table 1), in sites at distances from each other varying between 65 and 605 km. At each location, we

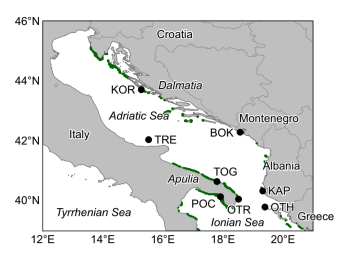


FIGURE 1 Sampling locations of *Posidonia oceanica* in the Adriatic and Ionian seas. The shading shows the suitable habitat produced by the MediSeH project (Giannoulaki et al., 2013), including a buffer to increase visibility. Location acronyms are TRE for Tremiti (MPA was established in 1989), TOG for Torre Guaceto (MPA was established in 1991), OTR for Otranto (this site is in a potential area for a future MPA), POC for Porto Cesareo (MPA was established in 1997) (all located in Italy), OTH for Othonoi in Greece (no MPA), KAP for Karaburun Peninsula in Albania (MPA was established in 2010), BOK for Boka Kotorska Bay in Montenegro (no MPA) and KOR for Kornati in Croatia (MPA was established in 1980). [Colour figure can be viewed at wileyonlinelibrary.com]

sampled *ca.* 50 individuals (spaced 5–8 m apart, a standard distance for this species) and according to "random walk" (Arnaud-Haond, Duarte, Alberto, & Serrão, 2007; Arnaud-Haond, Migliaccio, et al., 2007). This sampling strategy is a good compromise between avoiding the sampling of clonal replicates and assessing local genetic structure of a meadow by covering an extent of 250–400 m of the meadow.

2.2 | Potential (oceanographic) connectivity

2.2.1 | Potential connectivity by means of biophysical simulations

We investigated potential connectivity between sites where genetic sampling was carried out using Lagrangian oceanographic simulations. The individual-based biophysical model used here has been developed by Melià et al. (2016) and it is fully described there. The physical component of the model relies on fine-scale ocean reanalysis (in both the spatial and temporal sense) produced by the Adriatic Forecasting System, which assimilates satellite-based Earth observations and accounts for atmospheric forcing by the European Centre for Medium Range Weather Forecast (ECMWF) at 1/45° (*ca.* 13 km) and tidal signal (details at http://oceanlab.cmcc.it/afs). The ocean circulation fields are generated with the Adriatic Regional Model AREG (Oddo, Pinardi, Zavatarelli, & Coluccelli, 2006) at a daily temporal resolution, over a regular grid with a horizontal resolution of 1/45° (*ca.* 2.2 km) and 31 vertical sigma layers. The geographical domain encompasses the whole Adriatic Sea and extends southwards into the Ionian Sea down

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to the 39°N parallel. The bathymetry is based on the U.S. Navy 1/60° bathymetric database DBDB1. Being performed at large scales, such reanalyses cannot account for very local and/or extreme factors (such as tidal currents and erratic, but strong winds). However, we expect that this limitation does not strongly affect our results on connectivity. In fact, though strong winds (Ruiz-Montova et al., 2012) can affect movement of floating fruits (Grech et al., 2016), this effect is minor (McMahon et al., 2014), and expected to be modest in the Adriatic considering local wind speed (Katalinić, Ćorak, & Parunov, 2014) and limited tidal currents (Poulain, 2013). The biological component of the model (see Melià et al., 2016 for a more detailed description) accounts for the key traits affecting *P. oceanica* dispersal by sexual propagules: P. oceanica produces positively buoyant fruits, which are released between January and April (Balestri & Cinelli, 2003; Buia & Mazzella, 1991) and float in the upper layers of the water column for about 28 days before dehiscence and consequent release of the sinking seed (Serra et al., 2010). Lagrangian particles-passively guided within their motion according to the oceanographic fields-were released at a density of 2,000 particles per km² from areas of suitable habitat around the eight sampling locations, within a radius of 12.5 km. The suitable habitat was derived from the suitability model for P. oceanica produced by the MediSeH project (Giannoulaki et al., 2013) on the basis of the most up-to-date information on the distribution of seagrass meadows in the Mediterranean Basin. The Lagrangian simulations covered the period from 2003 to 2013 and a total of 5×10^6 particles were released. Each particle was assigned a fixed depth between 0 and 1 m below the surface and its trajectory was stepped forward for 28 days using a fourth-order Runge-Kutta integration scheme characterized by a 6-min time step, a linear convex combination in space and a linear interpolation in time of the current velocity field.

Potential connectivity between sites was measured in terms of intensity and persistence (sensu Melià et al., 2016). Connectivity intensity was calculated as the average (over the simulation period) number of particles released from a source site and reaching the suitable area of a destination site. Connectivity persistence, expressing the continuity of a connection throughout the years, was calculated as the stabilization coefficient (i.e., the reciprocal of the coefficient of variation) of connectivity intensity. Each site can then be characterized by its retaining strength (defining as *retainer* of Lagrangian particles a place where released propagules successfully remain in situ), source strength (defining as *source* a place from where released propagules successfully reach other sites) or sink strength (defining as *sink* a place to where propagules released from other sites tend to successfully settle). Other details on modelling explorations of potential connectivity are described in Melià et al. (2016).

2.3 | Realized (genetic) connectivity

2.3.1 | DNA extraction and microsatellite amplification

We extracted DNA from *ca.* 20 mg of silica-gel-dried tissue in 96well plates using the NucleoSpin[®] 96 Plant II kit (Macherey-Nagel) **TABLE 1** Genetic diversity of *Posidonia oceanica* at the eight locations in the Adriatic and Ionian seas. The 374 individuals from eight populations and five countries were assessed with 20

samples successfully amplified at all loci (N_g); the number of multilocus genotypes (MLG); genotypic richness (R); the mean number of alleles per locus (N_g); allelic richness standardized to 27

genotypes (A ₂ ; significant <i>F</i> ve	genotypes (A_{27}) ; observed heterozygosity (H_O) ; expected heterozyg significant F values. The parameters marked with * were calculated	zygosity (H _o); ∈ ers marked witl	expected heter h * were calcula	genotypes (A_{27}); observed heterozygosity (H_0); expected heterozygosity (H_E); the fixation index (F); and the percentage of polymorphic loci in the population (%p). Figures in bold indicate significant F values. The parameters marked with * were calculated after the removal of the chloroplastic locus Poc-trm	xation i al of th	index (F); and th oplastic l	e percent ocus Poc	trn	orphic loci in th	le population (%p). Figures in	bold indicate	
Country	Population	Latitude	Longitude	Location info	z	'	MLG	R	N a	A ₂₇	н _° *	H _E *	۲*	d%
Italy	Otranto (OTR)	40.109233	18.519217	Potential area for future MPA	48	45	27	0.59	1.80 (0.16)	1.8 (0)	0.45 (0.10)	0.26 (0.05)	-0.58 (0.11)	65%
	Porto Cesareo (POC)	40.195250	17.917950	MPA established in 1997	48	43	41	0.95	2.60 (0.32)	2.39 (0.11)	0.53 (0.10)	0.34 (0.06)	-0.44 (0.09)	80%
	Torre Guaceto (TOG)	40.716650	17.800050	MPA established in 1991	48	42	42	1	2.70 (0.40)	2.61 (0.07)	0.58 (0.08)	0.42 (0.06)	- 0.41 (0.09)	80%
	Tremiti (TRE)	42.138583	15.523950	MPA established in 1989	48	46	31	0.67	2.25 (0.25)	2.22 (0.05)	0.49 (0.10)	0.30 (0.05)	- 0.49 (0.09)	75%
Albania	Karaburun Peninsula (KAP)	40.392800	19.324967	MPA established in 2010	38	37	37	4	2.55 (0.35)	2.51 (0.03)	0.56 (0.10)	0.38 (0.06)	- 0.44 (0.11)	80%
Croatia	Kornati (KOR)	43.792250	15.281483	MPA established in 1980	48	44	33	0.74	2.30 (0.19)	2.26 (0.04)	0.42 (0.09)	0.29 (0.05)	-0.32 (0.09)	95%
Greece	Othonoi (OTH)	39.836017	19.397767	No MPA	48	44	34	0.78	2.70 (0.40)	2.68 (0.03)	0.55 (0.09)	0.37 (0.05)	-0.43 (0.09)	85%
Montenegro	Boka Kotorska Bay (BOK)	42.387533	18.569633	No MPA	48	45	33	0.73	2.40 (0.25)	2.29 (0.08)	0.52 (0.10)	0.31 (0.05)	-0.50 (0.09)	80%

following a modified protocol optimized for a Biomek FX robotic station (Tomasello et al., 2009). We amplified 22 microsatellites (Alberto et al., 2003; Arranz et al., 2013; Procaccini & Waycott, 1998) and ran PCRs as in Jahnke et al. (2015). See Tables S1 and S2 for details on primer sequences and PCR concentrations. Three loci were subsequently removed for most analyses, resulting in a dataset of 19 loci. We only used samples that were successfully genotyped at all loci for further analyses.

2.3.2 | Scoring and data quality checks

We scored the fragments by hand or using GeneMapper[®] (Life technologies) and rechecked scoring by eye for each individual. We used Microchecker (van Oosterhout, Hutchinson, Wills, & Shipley, 2004) to detect potential scoring errors and we revisited, and adjusted if necessary, loci with possible stuttering problems. We identified clones using GenClone (Arnaud-Haond & Belkhir, 2007) and removed duplicate multilocus genotypes (MLGs) before further analyses. Specifically, only one MLG for each clone was retained if the probability that the repeated genotypes do not originate from distinct sexual reproductive events, considering possible departures from Hardy-Weinberg equilibrium (HWE), was smaller than 0.05. After removal of (significant) clones, we used MicroDrop (Wang & Rosenberg, 2012) to detect null alleles. We tested for linkage disequilibrium (LD) and HWE at each locus and across all loci in each population with GENEPOP 4.2 (Raymond & Rousset, 1995), using 100 batches and 1,000 iterations per batch and applying Bonferroni corrections. Finally, we calculated the probability of identity (PI) in GENALEX 6.5 (Peakall & Smouse, 2012) to get an indication of the power of the marker set at each location, and we used POWSIM 4.1 (Ryman & Palm, 2006) to evaluate whether the sets of microsatellites have enough power to detect population structure among locations. We used the actual allele frequencies based on unique MLGs to simulate drift to F_{ert} levels of 0, 0.001, 0.01 and 0.1 using an effective population size (N_{e}) of 500 and a varying number of generations t (0–100) with 200 replicates and 100,000 batches.

2.3.3 | Outlier tests

We used Lositan (Antao, Lopes, Lopes, Beja-Pereira, & Luikart, 2008) and BayeScan (Foll & Gaggiotti, 2008) to test whether any of the used microsatellite markers do not behave according to expectations under neutrality. In Lositan, we ran the simulations for 50,000 iterations, with a 95% confidence interval, using the options for neutral mean F_{ST} , force mean F_{ST} , a subsample size of 40, the infinite allele model and eight populations based on the sampling sites. In BayeScan, we used default settings, which results in the same probability threshold as used for Lositan. We used the R script provided by Foll and Gaggiotti (2008) in R 3.2.2 (R Development Core Team 2014) to analyse whether any loci deviate significantly from expectation under neutrality and for plotting the posterior distribution. The two methods differ in the approach to identify outliers. While Lositan identifies outliers with higher than neutral heterozygosity conditioned on

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 F_{ST} (Antao et al., 2008), BayeScan uses posterior distributions generated by MCMC to identify whether a model including selection is more likely than a model without selection for each locus (Foll & Gaggiotti, 2008). We only considered as outliers those detected by both methods.

2.3.4 | Genotypic and genetic diversity and structure

We performed MLG identification for each population separately and repeated the analysis combining all populations to investigate clone sharing among populations in GenClone (Arnaud-Haond & Belkhir, 2007). Based on MLG identification, we calculated genotypic richness for each population according to Dorken and Eckert (2001). After removal of clone mates, we used GENALEx 6.5 (Peakall & Smouse, 2012) to calculate the number of alleles per locus, polymorphism and heterozygosity. We calculated allelic richness standardized to the minimum number of genotypes present in the dataset (27 MLGs at OTR) using the STANDARICH package in R 3.2.2 (http://www.ccmar.ualg.pt/maree/ software.php?soft=sarich). We used STRUCTURE (Pritchard, Stephens, & Donnelly, 2000) for K 2 to 8, to identify potential population structure based on neutral loci, all loci and loci putatively under selection. We assumed population admixture and correlated allele frequencies, but also performed runs with no admixture and independent allele frequencies. We used a burn-in of 100,000 and subsequent 1,000,000 steps, checking for run convergence. We identified the most likely number of populations based on delta-K with STRUCTURE HARVESTER (Earl & von Holdt, 2012) and used CLUMPAK (Kopelman, Mayzel, Jakobsson, Rosenberg, & Mayrose, 2015) to generate graphs. We also used ADEGENET (Jombart, 2008) in R 3.3.2 to perform a discriminant analysis of principal components (DAPC) (Jombart, Devillard, & Balloux, 2010) with the number of principal components set to 15, following alpha score indication. In order to validate these two approaches, we also performed an AMOVA with 999 permutations in GENALEX 6.5 (Peakall & Smouse, 2012).

2.3.5 | Genetic connectivity

We performed assignment tests in GENECLASS2 (Piry et al., 2004) using the exclusion method, because this method does not require an exhaustive sampling with every possible population of origin included in the dataset (Berry, Tocher, & Sarre, 2004; Underwood et al., 2007). This analysis was based on the dataset of neutral loci. We calculated the probability that an individual belongs to the population from which it was sampled with a partially Bayesian criterion (Rannala & Mountain, 1997) and compared the likelihood of exclusion of an individual to a distribution of likelihoods of 1,000,000 simulated genotypes in order to define a statistical threshold (Paetkau, Slade, Burden, & Estoup, 2004; Underwood et al., 2007) with a type I error of 0.05. We excluded an individual from its sampling site when the probability for exclusion was higher than 95% and we assigned the individual to another sampled population when the probability for inclusion in it was higher than 10% (Underwood et al., 2007). Otherwise, we assumed that the individual under study did not originate from the population where it was sampled, but originated most likely from an unsampled source population.

2.3.6 | Realized connectivity: isolation by (geographical) distance

We measured geographical distances between sampling locations using the shortest path over the sea without crossing land using Google Earth and used ARLEQUIN 3.5 (Excoffier & Lischer, 2010) to calculate pairwise Weir & Cockerham F_{ST} among populations and significance levels. We also calculated the unbiased estimator of Jost's D, D_{EST} (Jost, 2008) using the DIVERSITY package (Keenan, McGinnity, Cross, Crozier, & Prodöhl, 2013) in R 3.2.2 with 1,000 bootstrap replicates to test for the significance of pairwise comparisons. The two methods are to a certain degree complementary for assessing population differentiation: F_{ST} measures deviation from panmixia and is calculated based on allele frequencies; D measures deviation from complete differentiation and is based on the effective number of alleles (Whitlock, 2011; Meirmans & Hedrick, 2011). We tested for isolation by distance (IBD) for the two genetic distances separately using three datasets that contained all 19 diploid loci, only neutral loci and only outliers. We also calculated Slatkin's R_{sT} in SPAGEDI 1.4 (Hardy & Vekemans, 2002) and used 10,000 permutations to test whether $R_{\rm ST}$ is significantly higher than the permuted value pR_{sT} , which would indicate that the mutation rate exceeds the migration rate (Hardy, Charbonnel, Fréville, & Heuertz, 2003).

3 | RESULTS

3.1 | Potential (oceanographic) connectivity

The Apulian region was identified as the area with the highest potential connectivity (as obtained via Lagrangian simulations), in terms of both intensity (Figure 2, top panels) and persistence (Figure 2, bottom panels). The three Apulian sites OTR, TOG and POC (population acronyms as in caption of Figure 1) are the strongest retainers and sinks. However, while OTR and TOG are also the strongest sources, particles originating from POC do not reach any of the study sites. These three locations are connected by the current flowing southwards along the Adriatic coasts of Apulia and then turning around Salento towards the Gulf of Taranto. Particles released from TOG and OTR can potentially (yet through less intense and persistent connections) cross the Adriatic Sea and reach BOK and (only for TOG) KAP. There are also directional connections, driven by the southern Adriatic gyre, between the eastern and the western side of the Adriatic, with particles flowing from BOK to OTR and, through a less intense and persistent connection, from KOR to TRE. OTH acts in our modelling experiments only as a source of particles, and no particles reach this location from any other. TRE is a quite strong and constant source of particles for TOG and, to a lesser extent, OTR and KOR. KOR is a strong retainer and supplies particles to TRE. KAP is a good source, subsidizing Apulian sites (TOG, OTR, POC) via the southern Adriatic gyre. Particles released from OTH, instead, are not able to enter into the Adriatic Sea, but reach the two southernmost Italian sites (OTR and POC).

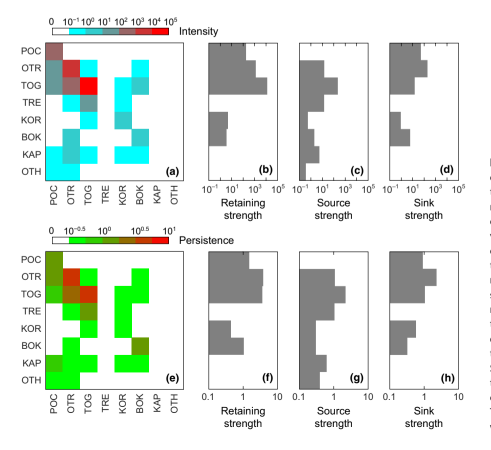


FIGURE 2 Oceanographic connectivity of eight Posidonia oceanica populations in the Adriatic and Ionian seas. Connectivity matrices (leftmost panels) show potential connectivity among sites, estimated via Lagrangian simulations, in terms of (a) intensity and (e) persistence (see text for details). Histograms show retention (b and f), source (c and g) and sink (d and h) strength of each site, as resulting by summing up the values of the corresponding matrices along the diagonal, the remaining row cells and the remaining column cells, respectively. Supplying populations are shown in the rows, receiving populations in the columns. Site acronyms as in Figure 1 and Table 1. [Colour figure can be viewed at wileyonlinelibrary.com]

3.2 | Realized (genetic) connectivity

3.2.1 | MLG identification, null alleles and outliers

We identified a high number of MLGs at each location, ranging from 27 to 42 MLG per population (Table 1), resulting in 278 genets (out of 374 ramets) that were used for all further analyses. Three loci showed frequencies of null alleles above 10% in MicroDrop (Wang & Rosenberg, 2012). One of them, Poc-trn (NaF = 30.8%), is chloroplastic, that is, haploid and therefore expected to be always homozygous. The other two loci (Poc-5, NaF = 19.6% and Pooc-330, NaF = 11.1%) were removed before further analyses, while Poc-trn was retained for few descriptive statistics only (Table 1), resulting in a marker set of 19 loci. Both Lositan and BayeScan identified the same five loci to be under balancing selection (Figs S1 and S2). As the non-conformity to neutrality of these loci can affect patterns of connectivity and migration, we used three different datasets in the following analyses: (1) all diploid loci (19 markers), (2) only neutral loci (14 markers) and (3) only outlier loci (five markers under balancing selection).

3.2.2 | Linkage disequilibrium (LD), Hardy-Weinberg equilibrium (HWE) and power of the marker set

We found significant LD in 11 of 120 tests across all populations (9%) after applying Bonferroni corrections. In particular, we detected

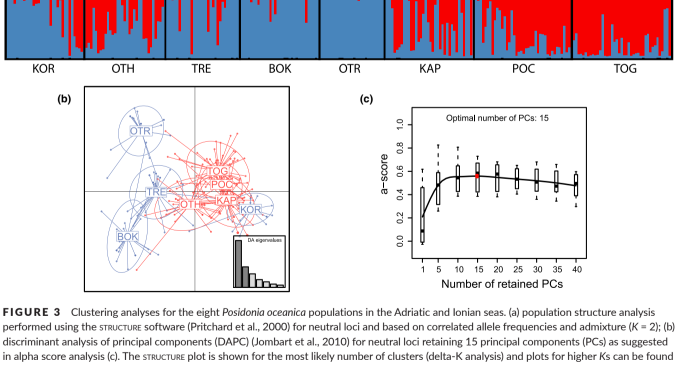
three markers to be in gametic linkage more than two times: Pooc-PCo45G11 (five times), Pooc-229 (four times) and Pooc-361 (three times). PCo45G11 is the locus with the highest number of alleles in the dataset, while the number of alleles per locus is low for the other loci (ranging from one to seven).

Seven HWE tests per population and locus were significant after Bonferroni corrections (12%). No locus deviated from HWE at more than two locations. As the HWE deviations were found to be specific to locations rather than loci and as we did not find indications of quality control problems, we retained all loci.

The probability of identity (PI) was low, ranging from 4.6×10^{-5} in OTR to 6.7×10^{-9} in TOG. The PI for sibs was higher, ranging from 5.6×10^{-3} in OTR to 1.5×10^{-4} in TOG, which are still Pl values sufficient for discerning siblings, considering the number of MLGs. Power simulations of the full marker set and the neutral marker set suggest that both sets of loci can provide a reasonably accurate picture of genetic structure, with population homogeneity rejected in 100% of the simulations when $F_{\rm ST}$ was as small as 0.01 (Table S3).

3.2.3 | Genotypic and genetic diversity and structure

Genotypic richness varied among populations, while heterozygosity was similar (Table 1). Allelic richness was generally low, ranging from 1.8 at OTR to 2.68 at OTH (Table 1). All populations had a significant excess of heterozygosity as evident by high negative F_{1S} values



in Fig. S4. Within each plot, each vertical bar represents an individual belonging to the sampling location indicated under the x-axis, clusters are colour-coded, and the y-axis of each plot shows the proportion of the genotype belonging to each cluster. The DAPC analysis was performed based on the location of sampling (as opposed to defined by the cluster analysis of DAPC) and the colour of each population represents the colour of the majority of individuals of this population in the corresponding analysis performed by the structure software. Each dot represents an individual contained into populations by a circle. Site acronyms as in Table 1 and Figure 1. [Colour figure can be viewed at wileyonlinelibrary.com]

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(Table 1). This phenomenon has been observed previously in *P. oce-anica* (Arnaud-Haond, Migliaccio, et al., 2007; Serra et al., 2010) and is most likely linked to its life history of partial clonality (Reichel, Masson, Malrieu, Arnaud-Haond, & Stoeckel, 2016). No MLGs were shared among locations. Pairwise $F_{\rm ST}$ ranged from 0.05–0.23 for the dataset of neutral loci, and all pairwise comparisons were significant (see Tables S4–S10 for values for different *F*-statistics and loci sets).

The STRUCTURE analysis performed on the dataset of only neutral loci showed the presence of two populations clusters (K = 2, Fig. S3) as the most likely possibility. The two population clusters consist of the northern populations of KOR. TRE. BOK and the more southern OTR (Figure 3a, mostly blue populations) and the southern populations of KAP, POC, TOG and OTH (Figure 3a, mostly red populations). However, several locations (OTH and KAP in particular) show a high degree of admixture between the two clusters and when assuming higher Ks further substructuring becomes evident (Fig. S4). The STRUCTURE analysis for higher Ks (particularly K = 3-6) also shows an interesting pattern of migrants in each populations, but few admixed individuals. DAPC confirms the separation of the two clusters for all populations but KOR (Figure 3b,c); however, KOR is located close to KAP and OTH, which show a nearly 50-50 percentage of belonging to the northern and southern clusters. The DAPC also shows clearly that OTR is the most differentiated population and that all populations are differentiated from each other, which is also confirmed by the AMOVA, where the "among-populations" level explains most variance (Table S11). The STRUCTURE and DAPC analyses based on the five loci presumably under balancing selection show no detectable population structure, while the picture based on all 19 diploid loci is very similar to the analyses based on the 14 neutral loci (not shown). The identification of outlier loci is associated with high type I errors, that is, a high rate of false positive results, especially for loci that are under balancing selection (Narum & Hess 2011). The observation that results based only on neutral or on all loci are very similar suggests that loci supposedly under balancing selection may have been falsely identified.

3.2.4 | Realized (genetic) connectivity and IBD

For realized connectivity assessments, we only considered the neutral loci dataset, as dispersal should make the biggest contribution to the observed allele frequencies of neutral loci in the different populations (as opposed to selection in the other two datasets). The assignment tests (GENECLASS) show a strong population structure with only 4% of samples assigned to populations different from those of the sampling location (Table 2). TOG is identified as the most important source population, providing one individual each to TRE, KAP and BOK (Table 2). This population has the highest possible level of genotypic richness, that is, high levels of sexual recruitment. Conversely, OTR has the highest level of clonality and all sampled individuals get assigned to their own population (Table 1; Table 2). The IBD analysis did not reveal a positive correlation between neither F_{ST} nor D_{EST} (Tables S4-S9) and geographical distance for any of the three datasets (not shown). R_{ST} values were similar to F_{ST} values (see Tables S5–S11) and the permuted R_{sT} did not differ significantly from the observed value **TABLE 2** Assignment test of *Posidonia oceanica* in the eight Adriatic and Ionian populations based on the neutral microsatellite set (14 loci). For each site (acronyms as in Table 1), individuals are presented in rows according to their sampling site and classified into individuals that get assigned to their own population (self) and other sites that they get assigned to, namely Torre Guaceto (TOG), Othonoi (OTH) or unknown sources that could not be ascribed to any of the sampled populations (Unknown). The last column lists the total number and percentage of individuals that were not assigned to the population from which they were sampled

	Origin				
Population	Self	TOG	ОТН	Unknown	Total
OTR	27	-	-	-	0 (0%)
POC	39	-	-	2	2 (5%)
TOG	41	-	-	1	1 (2%)
TRE	30	1	-	-	1 (3%)
KAP	35	1	-	1	2 (5%)
KOR	31	-	1	1	2 (6%)
OTH	34	-	-	-	0 (0%)
BOK	31	1	-	1	2 (6%)
Total	241 (96%)	3	1	6	10 (4%)

(two-sided *p*-value = .69, $R_{ST} = 0.17 > pR_{ST} = 0.16$), that is, there was no indication that mutations made a high contribution to population differentiation and/or mutations do not follow a stepwise pattern.

4 | DISCUSSION

The biophysical connectivity assessments show a high potential for dispersal of P. oceanica fruits across the whole study area. The presented results on potential connectivity are robust and would neither qualitatively nor quantitatively be altered by incorporating into our biophysical model minor effects, such as movements of floating fruits caused by erratic strong winds. Realized connectivity, which can serve as an important indication for conservation policies and management, shows more complex patterns, but is apparently lower. There is high genetic structuring of the eight assessed P. oceanica sites in the Adriatic and Ionian seas, with significant pairwise population differentiation among all locations (see Tables S5-S11), and assignment tests show only a low level of recent migrants. First-generation migrants were also evident in the STRUCTURE analysis of higher Ks and the low number of admixed individuals in this analysis points to low sexual reproduction and/or non-random mating of immigrants in the assessed populations. Geographical distance was not a good predictor for genetic differentiation, but we identified two main population clusters that are in reasonable agreement with a latitudinal gradient, that is, a northern and a southern cluster (with the exception of the southern site of Otranto that groups with the northern cluster and is generally the most differentiated site). In the assignment tests, the meadow at the Torre Guaceto MPA (TOG) was identified as the most important source population. Interestingly, this result is corroborated by the

biophysical analysis, where TOG also turns out to be the most important source population. The population of TOG is within an MPA with an enforced no-take area and an enforced no-anchoring ban above the assessed *P. oceanica* meadow, and is presumably one of the most efficiently protected meadows of the evaluated sites (Guidetti et al., 2008). Our results confirm the key role played by TOG as a source of propagules, a role that was already established with physical modelling for different organisms in the whole Adriatic Basin (Melià et al., 2016; Pujolar et al., 2013). The location of this protected area was not only well chosen for achieving positive population dynamics at the local scale (Fraschetti, Guarnieri, Bevilacqua, Terlizzi, & Boero, 2013), but TOG is also very well connected to other *P. oceanica* populations in the Adriatic. We thus suggest that conservation measures for this MPA should be confirmed and possibly re-enforced.

The existence of two genetic clusters was suggested by both the set of neutral microsatellite loci and the complete set of loci, including also the five outliers, but was not necessarily confirmed by the oceanographic modelling, as most populations are predicted to supply and receive propagules to and from both northern and southern sites. For instance, in the physical modelling, the southern site of Otranto (OTR), which groups with the northern genetic cluster, has the highest probability of dispersal to two sites of the southern cluster and one (BOK) of its own cluster and is expected to receive propagules only from the southern cluster. However, this meadow has the highest levels of clonality, the lowest levels of standardized allelic richness and fixed allele frequencies with no private alleles. OTR is clearly the most differentiated of all sites in the DAPC, suggesting that postdispersal (i.e., pre- or post-settlement) processes played a role in the observed differentiation.

The levels of realized connectivity, as assessed by genetics, show a complex pattern with detectable levels of migration, but "mosaic" populations with few admixed individuals. This picture confirms the stochasticity of dispersal at small/medium spatial scales observed in other seagrasses (Kendrick et al. 2012). Possible reasons could be unsampled populations that confound the picture, pre- and postsettlement processes and non-random mating, including low levels of sexual reproduction in general. This is expected to be very pronounced for P. oceanica, as the partial clonality and longevity of clones translates into generation times that may be as long as thousands of years (Arnaud-Haond et al., 2012; Ruggiero, Turk, & Procaccini, 2002). Regional and basin scale population structuring, supported despite detectable recent migration and no IBD on most assessed spatial scales, was already described for P. oceanica over its entire distribution (Arnaud-Haond, Migliaccio, et al., 2007; Rozenfeld et al., 2008; Serra et al., 2010). As an alternative to stochastic events of long-distance dispersal, this pattern of P. oceanica population differentiation has also been proposed to stem from a stronger influence of mutation over migration at the scale of the distribution range (Arnaud-Haond et al., 2014). Under this assumption, population differentiation may be explained by historic step-by-step colonization followed by local recruitment and clonal growth, rather than contemporary gene flow. Here we used a permutation test of Slatkin's $R_{\rm ST}$ (as suggested by Hardy et al., 2003) and did not find any indication that mutations played a major role WILEY

for genetic differentiation, and we also show that oceanographic (potential) connectivity is high among the assessed populations. Potential connectivity may well be higher than realized connectivity, because of low sexual reproduction (estimates of oceanographic connectivity are based on dispersal of fruits), low settlement success after dispersal or small-scale hydrodynamics that could not be included into the oceanographic connectivity analysis. Indeed, the modelling analysis showed that two central populations, OTR and TOG, had the highest potential for acting as sources. TOG, which has high genotypic richness (indicating a high level of sexual reproduction), seems to realize this potential and supply sexual propagules to other populations. In addition, the biophysical modelling suggests that TOG can supply propagules to BOK, as also confirmed in the genetic assignment test. In contrast, OTR has a slightly lower potential for dispersal, but is genetically distinct and has a much lower genotypic richness, suggesting that this population supplies fewer sexual propagules to other meadows. This points out that the occurrence of sexual reproduction is an important parameter that may significantly influence the link between potential and realized connectivity in P. oceanica. The biophysical modelling also indicates TOG and OTR as strong retainers, a result that corroborates the outcomes of previous analyses suggesting a strong retention potential for this area (e.g., Di Franco et al., 2012; Schiavina et al., 2014): indeed, both sites have one of the highest percentages of individuals assigned to their own population in the genetic assignment test.

5 | CONCLUSIONS

Connectivity assessments are increasing rapidly in the field of conservation science (Jones et al., 2009) and information on connectivity is important to MPA network design. They can deliver information on actual dispersal rates, and identify populations that export propagules to other areas (source populations) or populations that rely on immigration for their sustenance (sink populations), as well as populations that retain their propagules locally. Moreover, linking connectivity assessments with information on the levels of genetic diversity could also be used to identify areas of high evolutionary potential (Vandergast, Bohonak, Hathaway, Boys, & Fisher, 2008). Connectivity assessments are however only one component of the MPA design process and for instance size, number, representation, replication, diversity and above all capacity are other important factors (Fernandes et al., 2009; Gill et al., 2017). In this study on connectivity, we found that the potential for dispersal was considerably higher than realized migration, but both approaches coherently identified the same optimal site, which is at the same time a strong retainer, a good source and a good sink. For species which disperse mainly by sexual propagules, yet can alternate between sexual and asexual reproduction, the amount of sexual reproduction may be a very important component to take into account when assessing connectivity. So far, the majority of connectivity assessments involving MPAs have been performed on mobile species, exclusively sexual in their reproduction (in the Adriatic, see for instance Boissin et al., 2016; Pujolar et al., 2013; Paterno et al., 2017). Our results on potential and realized connectivity indicate that II FY- Diversity and Distribution

dispersal occurs at large spatial scales (100s of km) for a sessile benthic partially clonal species and suggest that potential connectivity can be insufficient *per se* to describe population structure. Rather, postdispersal, pre-settlement and post-settlement processes have to be taken into consideration to understand discrepancies between potential and realized connectivity. Together, our findings on potential and realized connectivity, genetic structure and sexual reproduction have direct conservation application and can be used for the establishment and management of MPAs and other large-scale conservation strategies.

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BIOSKETCH

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SUPPORTING INFORMATION

Additional Supporting Information may be found online in the supporting information tab for this article.

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