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Mediterranean coastal pine forest stands: understorey distinctiveness or species cauldron?

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8

9 Abstract

A common perception of plantation forests is that they constitute an "ecological desert" and thus 10 they are often disdained by scholars. Distribution patterns of understorey assemblages of coastal 11 pine stands on sand dunes are still little known, despite such forests being widespread along the 12 Mediterranean coastline, particularly in Italy. The purpose of this study of 167 plots along Italian 13 coastlines was to analyse whether similar communities and specific species pools occur in different 14 pine forest types dominated by Pinus halepensis, P. pinea or P. pinaster. Multivariate analysis was 15 used, considering the effects of sea-inland gradient and pine canopy cover. The results indicated 16 that pine forests do not consist of specific vegetation assemblages, suggesting the idea of a "floristic 17 cauldron". Understorey distinctiveness of Mediterranean coastal pine stands is limited, with 18 psammophilous species of coastal dunes occurring mostly in P. halepensis stands, and forest 19 species mainly being linked to P. pinea stands. Thus species-specific management is not required 20 21 for different pine forest types. Like on non-forested dunes, plant species mainly followed the seainland gradient, maintaining the natural zonation of coastal vegetation. Moreover, higher pine 22 canopy cover affected both herbaceous and woody natural dune vegetation in a negative way. These 23 24 findings should not be disregarded since they have implications for management planning and

conservation and because understoreys of Mediterranean pine plantations, with a species reservoirof unknown value, have often hitherto been overlooked.

27

28 Keywords: canopy cover; dunes; Italy; plant species assemblage; sea-inland gradient; vegetation.

29

30 **1 Introduction**

Forest stands with dominance of stone pine, Aleppo pine and maritime pine (P. pinea L., Pinus 31 32 halepensis Mill., P. pinaster Aiton) occur along almost all the low sandy coast of the Italian peninsula (Biondi et al., 2009), extending from the first back dune to the last settled innermost dune 33 34 environment. Italian coastal pine forest vegetation, autochthonous or derived from ancient 35 plantations, consists mainly of P. pinea or P. halepensis (Biondi & Blasi, 2015). Historically, the primary role of coastal pine forest stands was as a shelterbelt to protect cropland from salty sea 36 spray and for production of pine nuts, timber and resin. These forests were mainly planted in the 37 38 second half of the twentieth century or later and were maintained by humans for coastal defence against impermeabilisation waterproofing and soil erosion (Calama et al., 2003). They are well 39 known for their rehabilitation capacity in Mediterranean dunal environments (see Bellarosa et al., 40 1996), as well as for recreational uses (Cutini et al., 2013) and carbon storage and sequestration 41 services (Drius et al., 2016). Planted forests that were established a long time ago are more likely to 42 43 be a habitat for biodiversity, although a common perception is that they are "ecological deserts" (Brockerhoff et al., 2008). In fact, at European level they are included in "Coastal brown dunes 44 covered with natural or almost natural thermophilous pines" (EUNIS Habitat Type Code B1.74) and 45 since the 92/43 EEC Directive Habitat in priority habitat 2270 ("Wooded dunes with P. pinea 46 and/or P. pinaster"), also including P. halepensis forests (Biondi et al., 2009). However, in recent 47 decades, environmental concern about sandy coasts and associated pine forest have increased, 48

49 mainly due to direct and indirect effects of human activities, such as coastal erosion (Raddi et al.,

50 2009), salinisation of groundwater (Antonellini & Mollema, 2010; Zanchi & Cecchi 2010),

51 trampling (Santoro et al., 2012) and urban sprawl (Reina-Rodríguez & Soriano, 2008; Malavasi et

al., 2013). These threats have been particularly intense in the last 50 years in countries bordering the
Mediterranean Sea (Curr et al., 2000).

The flora of coastal pine forests has rarely been studied because artificial forests are not highly 54 55 esteemed from a conservation viewpoint (Brockerhoff et al., 2008). Although pines are the most common plantation species worldwide (about 20% of total plantation area) (FAO, 2001), pine 56 forests have been considered "second choice" in plant community research compared with other 57 58 forest types. However, pine-climate interactions (Mazza et al., 2011, 2014; Cutini et al., 2015), fire practices and their effects (Fernandes & Botelho, 2004; Rigolot, 2004; Fernandes et al., 2008) and 59 dendrochronology are widely debated in the scientific literature (Calama et al., 2003; Calama & 60 Montero, 2005). Although they are threatened like other dunal plant communities, the understoreys 61 of these forests and the processes that drive them are still largely unknown and the importance of 62 63 plantation forests for biodiversity conservation goals is a controversial issue.

Our primary objective in this study was to understand whether pine forests with different dominant 64 pine species can harbour a particular understorey species pool or should rather be considered a 65 "species cauldron". We assumed that overstorey composition and structure influence understorey 66 plant communities through modification of resources including light and soil (Messier et al. 1998, 67 Légaré et al., 2001), with some species having special affinities for a particular overstorey type 68 (Bartels & Chen, 2010). We analysed also the effects of sea-inland gradient and pine canopy cover 69 70 on the underlying layers. Major questions were: i) Does vegetation associated with a pine forest 71 type form distinct communities? ii) Are some plant species more likely to occur in one pine forest type than others? iii) How do distance to coastline and pine canopy cover affect pine understorey 72 assemblages? 73

- 74 In this way, this work aims to contribute to the question whether species-specific management is
- required for forests dominated by different pine species and their understoreys, as a guide for

vegetation management in Mediterranean coastal areas.

77

78 2 Methods

79 2.1 Pine forest-types

80 For data collection we considered the EU Directive definition of Habitat 2270 "Wooded dunes with

81 *P. pinea* and/or *P. pinaster*" that also includes *P. halepensis* forests (Biondi et al., 2009). We then

used data of three different pine forest-types dominated by: i) Stone pine (*Pinus pinea* L.); ii)

83 Aleppo pine (*Pinus halepensis* Mill.); iii) Maritime pine (*Pinus pinaster* Aiton).

84 *Pinus pinea* is scattered throughout the Mediterranean basin, mainly in coastal areas. In Italy this

⁸⁵ "umbrella-shaped" tree is the icon of Italian coastal forests (Mazza et al., 2014) and it is reported as

native to Liguria, Tuscany, Molise and the major islands, i.e. Sicily and Sardinia (Conti et al.,

87 2005), although there is little evidence of its nativeness (Abad Viñas et al., 2016a). *Pinus halepensis*

is largely present in the Western sector of the Mediterranean basin and it is the most widely

89 distributed Mediterranean pine. In Italy the species is recognized as native to all regions except the

90 Alps (Conti et al., 2005), although it has been widely planted (Mauri et al., 2016). *P. pinaster* has a

- 91 western Mediterranean distribution; this medium-size pine mainly occurs in north and central Italy.
- 92 Its presumed native distribution includes Liguria, Tuscany, Lazio and the major islands (Conti et al.,
- 93 2005).

94

95 2.2 Study area

96 The study area included pine forests established on dunes along the coasts of the Italian peninsular. 97 The forests occur in six admistrative regions on the coasts of the Tyrrhenian (Toscana, Lazio and 98 Campania) and Adriatic seas (Emilia-Romagna, Molise and Puglia), including much of the 99 distribution of pine stands in the Italian peninsular. The sites ranged about from 0 to 5 m a.s.l. and 100 phyto-climate depends mostly on latitude, ranging from Temperate to Mediterranean, passing throw 101 transition zones (Blasi & Michetti, 2007). The study area comprises mainly calcareous sediments, 102 although aeolian deposits also occur from time to time (Geoportale Nazionale, 2015).

103

104 2.3 Data collection

For the purpose of the present study, field sampling in Pinus pinea stands was performed in 2014-105 2015, whereas an existing database of coastal dune vegetation (Acosta et al., 2009, Malavasi et al., 106 107 2016) was mainly used for *P. halepensis* and *P. pinaster* stands. Firstly, only plots with at least one of the pine species Pinus pinea, P. halepensis or P. pinaster were used because our interest was the 108 role of pines. Secondly, only plots on sandy soil were included in the dataset. Thus, a dataset based 109 110 on 167 plots $(2 \times 2 \text{ m})$ was defined. Each plot was assigned to a pine forest type in relation to the pine species dominant in each plot. Distance between georeferenced plot and coastline, used as a 111 proxy of sea-inland gradient, was determined remotely by GIS. The study did not involve any 112 experimental manipulations or disturbance of naturally developed relationships. Observed patterns 113 should therefore reflect long-term plant-ecological interactions. The focal species of dunal 114 115 vegetation were identified and selected using the list of diagnostic and characteristic species in the "Italian Interpretation Manual of the 92/43/EEC Directive Habitats" (Biondi et al., 2009; Biondi & 116 Blasi, 2015). Autochthonous species names are according to Conti et al. (2005), alien species names 117 118 are according to The Plant List (2013).

119

120 **2.4 Data analysis**

121 The dataset of 167 plots (94 ascribed to *Pinus pinea*, 65 to *Pinus halepensis*, and 8 to *Pinus*

122 *pinaster*) contained 269 species. Plant values were cover percentages of each species between 0 and

123 100. The complete species list is reported in Appendix A.

Evaluation of species composition distinctiveness in each forest type (with and without pine canopycovers) was performed using the following techniques:

126 1) Multi-response Permutation Procedure (MRPP), a non-parametric multivariate procedure for

testing the hypothesis of no difference in species composition between two or more groups of plots

128 chosen a priori (McCune & Mefford, 1999; McCune & Grace, 2002). A weighted mean within-

group distance in species space is calculated using Sorensen distance. MRPP consists of two

130 statistical tests: the *A* Statistic estimates within-group homogeneity and the *T* Statistic measures

between-group separability. Higher A statistic values (maximum value 1) indicate a high degree of

homogeneity within groups while a large negative T value (\leq -10.0) indicates high separability

between groups. The null hypothesis was assessed by a Monte Carlo permutation procedure with

134 999 permutations;

2) Non-Metric Multidimensional Scaling (NMDS) based on Euclidean distance, used to investigate
 community patterns without data transformations;

3) INdicator SPecies ANalysis (INSPAN), performed to find species significantly associated with
each forest type via 4999 randomization tests (Dufrené & Legendre, 1997).

139 To investigate whether sea-inland gradient and pine canopy cover significantly influenced

140 understorey species distribution in coastal pine forests, two hybrid constrained CCAs were

141 performed, with the Log (X+1)-transformed variable 'Distance'. The significance of the constrained

142 axis was tested by a Monte Carlo randomization method, using 999 permutations. To test species

143 response to sea-inland gradient and pine cover, for species with more than ten occurrences in the

dataset, including pines, we used General Additive Models (GAM) with binomial distribution, logit
link function, 2.0 df (see Yee & Mitchell, 1991; Hájková et al., 2008; Ilunga wa Ilunga et al., 2013;
Dyakov, 2016) and the quasi-distribution approach for modelling overdispersed data. GAMs were
chosen because they assume a non-linear relationship between the response and each explanatory
variable. Binomial distribution data was treated as binary. MRPP and INSPAN analyses were
performed using the software package PCORD 6.0 (McCune and Mefford 2011), whereas NMDS,
CCAs and GAMs were calculated using CANOCO v. 5.03 (ter Braak & Šmilauer 2012).

151

152 **3 Results**

Twenty-nine percent (78 out of 269) of all coastal pine stand species were focal for dune habitats 153 (Appendix A). The MRPP results (Table 1) using the data set including pine species canopy covers 154 suggested that forests dominated by different pine species host different plant species assemblages. 155 The T statistic, representing separation between pine forest types, was high, negative and highly 156 significant, while the A statistic indicated relative within-group homogeneity. However, for the data 157 set without pine species canopy covers, the results showed few differences between vegetation 158 plots, particularly when forest types were compared. Although the T statistic was negative and 159 statistically significant, within-group observed average distance was high. In addition, the A 160 statistic, a measure of within-group homogeneity, was an order of magnitude less than the value 161 obtained when pine canopy cover data was included. 162

164	Table 1. MRPH	community ?	comparison	results
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	T statistics	Sorensen distance	A statistics
Data with pine species	-62.74 (p < 0.001)	0.68	0.21

166	NMDS diagrams with plots grouped in three clusters obtained by a priori classification of forest
167	dominated by different pine species are shown in Figure 1. When pine cover data was included
168	(final stress = 0.10281), the first three axes explained 100% of community variation (axis 1
169	captured 60.1% of the variance, axis 2, 22.4%) and the biplot demonstrated clear disjunction of
170	plots dominated by different pine species (Figure 1a). In contrast, when pine canopy covers were
171	excluded (Figure 1b; final stress = 0.18482) the first three axes explained 100% of community
172	variation (axis 1 captured 38.8% of the variance, axis 2, 34.2%) and showed unclear separation of
173	plots dominated by different pine species.

174

×		

176 Figure 1. NMDS diagram of plots including (A) and excluding pine canopy cover data (B). Colors

- 177 indicate the dominant pine species. Blue diamonds are Pinus halepensis dominated plots, red stars
- are *P. pinea dominated plots and green triangles are P. pinaster dominated plots.*

179

INSPAN to detect affinity of species with the three pine forest types revealed a low percentage of 180 indicator species (10.8%) among the total number of species (Table 2; McCune & Grace, 2002). 181 Indicator species for P. pinea forest type numbered nine (3.4%) and were mainly related to forest on 182 stabilized dunes, including Brachypodium sylvaticum ssp. sylvaticum and Carex distachya. 183 Indicator species for *Pinus halepensis* forest type numbered 20 (7.4%), and were typical pioneer 184 psammophilous species, such as Ammophila arenaria ssp. australis and Medicago marina, mostly 185 associated with the forward part of the dune series. No significant indicator species for P. pinaster 186 187 forest type were detected.

189 Table 2. INSPAN diagram with dominance of Pinus halepensis, P. pinea and P. pinaster. Only

Dominant	Indicator species	IV	р
species			
Pinus pinea	Arum italicum	8.4	0.0484
(p = 0.001)	Brachypodium sylvaticum ssp. sylvaticum	13.8	0.0032
	Carex flacca	11.7	0.0072
	Carex distachya	8.5	0.0192
	Clematis flammula	9.7	0.0280
	Cyclamen repandum	7.4	0.0374
	Euphorbia peplus	10.6	0.0104
	Geranium purpureum	11.7	0.0066
	Trachynia distachya	7.4	0.0338
Pinus halepensis	Ammophila arenaria ssp. australis	9.4	0.0032
(p = 0.001)	Artemisia campestris ssp. glutinosa	10.9	0.0012

190 significant species are shown. Species are sorted in alphabetical order. *IV* = *Indicator Value*.

	Cyperus capitatus	15.6	0.0004
	Echinophora spinosa	6.2	0.0226
	Elymus farctus ssp. farctus	21.9	0.0002
	Elymus repens ssp. repens	7.8	0.0072
	Erigeron canadensis	6.2	0.0200
	Erodium laciniatum	12.5	0.0002
	Eryngium maritimum	6.2	0.0216
	Euphorbia terracina	17.2	0.0002
	Medicago marina	10.9	0.0006
	Ononis variegata	9.4	0.0034
	Phleum arenarium ssp. caesium	12.5	0.0006
	Reichardia picroides	17.1	0.0004
	Silene colorata	17.2	0.0002
	Silene vulgaris	17.7	0.0004
	Sixalix atropurpurea	20.9	0.0002
	Sporobolus virginicus	25.0	0.0002
	Verbascum niveum ssp. garganicum	12.5	0.0004
	Vulpia fasciculata	17.2	0.0002
Pinus pinaster	/	/	/

(n.s.)

191

192

193 Hybrid CCA (Figure 2) with first axis constrained by sea distance significantly discriminated three

194 clusters of coastal pine forest species: typical dunal species closer to the sea, such as *Elymus farctus*

195 ssp. *farctus* and *Vulpia fasciculata*, species typical of Mediterranean maquis at intermediate

- 196 distance, such as *Phillyrea angustifolia* and *Hedera helix*, and generalist species at greater distances
- 197 from the sea, such as *Bellis perennis* and *Cerastium glomeratum*.





199

Figure 2. Hybrid constrained CCA diagram. Axis 1 captured 3.2% of the variance (p = 0.001). The 200 201 first 20 best fitting species in the ordination spaces are shown. Plant species are designated with the first three letters of their genus and species names as follows: Acacya = Acacia cyanophylla; 202 *Belper* = *Bellis perennis; Cerglo* = *Cerastium glomeratum; Cynech* = *Cynosurus echinatus;* 203 204 Dandec = Danthonia decumbens ssp. decumbens; Elyfar = Elymus farctus ssp. farctus; Erican = *Erigeron canadensis; Eupter = Euphorbia terracina; Hedhel = Hedera helix; Lotcyt = Lotus* 205 cytisoides; Lothis = Lotus hispidus; Oxa = Oxalis sp.; Phiang = Phillyrea angustifolia; Pinhal = 206 *Pinus halepensis; Pinpin = Pinus pinea; Queile = Quercus ilex ssp. ilex; Reipic = Reichardia* 207 picroides; Rhaala = Rhamnus alaternus ssp. alaternus; Smiasp = Smilax aspera; Vulfas = Vulpia 208 fasciculata. 209

Hybrid constrained CCAs with canopy cover of *P. halepensis* and *P. pinea* (Figure 3) revealed the
significant effect of pine cover on the understorey. In both CCAs, lower canopy cover was
associated with herbaceous and shrub species of natural dunal successions, whereas higher canopy
cover was associated with generalist and alien species. Hybrid constrained CCA with *P. pinaster*did not produce significant results and is therefore not shown.

215



216



219 respectively. The first 20 best fitting species in the ordination spaces are shown. Plant species are

220 *designated with the first three letters of their genus and species names as follows: A) Aspacu =*

221 Asparagus acutifolius; Belper = Bellis perennis; Brarup = Brachypodium rupestre; Cerglo =

- 222 *Cerastium glomeratum; Ciscre = Cistus creticus ssp. eriocephalus; Clefla = Clematis flammula;*
- 223 Dapgni = Daphne gnidium; Fumcap = Fumaria capreolata ssp. capreolata; Gerrob = Geranium
- *robertianum; Helsto = Helichrysum stoechas; Junoxy = Juniperus oxycedrus ssp. macrocarpa;*
- 225 Linpur = Linaria purpurea; Plalan = Plantago lanceolata; Queile = Quercus ilex ssp. ilex; Rusacu

226 = Ruscus aculeatus; Solnig = Solanum nigrum; Stemed = Stellaria media; Trirep = Trifolium 227 repens; Verarv = Veronica arvensis; Vullig = Vulpia ligustica; B) Acacya = Acacia cyanophylla; Ammare = Ammophila arenaria ssp. australis; Aspacu = Asparagus acutifolius; Cypkal = Cyperus 228 229 capitatus; Elyfar = Elymus farctus ssp. farctus; Elyrep = Elymus repens ssp. repens; Linstr = *Linum strictum; Lonimp = Lonicera implexa ssp. implexa; Lotcyt = Lotus cytisoides; Medmar =* 230 *Medicago marina; Onovar = Ononis variegata; Phiang = Phillyrea angustifolia; Phlare = Phleum* 231 arenarium ssp. caesium; Queile = Quercus ilex ssp. ilex; Rosoff = Rosmarinus officinalis; Rusacu 232 = Ruscus aculeatus; Silcol = Silene colorata.; Smiasp = Smilax aspera; Spopun = Sporobolus 233 *virginicus; Vulfas = Vulpia fasciculata.* 234 235 GAMs on pine species revealed that *P. pinea* and *P. halepensis* were highly significant (p < 0.001;

Appendix B). The curve trends (Figure 4) showed that *P. pinea* showed its greatest probability of occurrence after the first 1000 m, continuing for the rest of the sea-inland gradient. On the other hand, *P. halepensis* had the highest probability of occurrence in the first 1000 m from the coast and then disappeared. The *P. pinaster* curve is not shown in Figure 4 because it was not significant.



- 242 Figure 4. Species-response curves (GAMs) of pine species with respect to distance from sea
- 243 (probability of occurrence in relation to sea distance). Pinus halepensis and P. pinea were
- significant (p < 0.001). P. pinaster was not statistically significant.
- 245
- GAMs revealed 14 species that responded significantly to binomial distribution based on CCA with
- 247 distance (Figure 5; Appendix C). Along the sea-inland gradient, dune species *sensu strictu*, such as
- 248 *Lotus cytisoides*, showed maximum probability of occurrence close to the sea, while innerdune
- 249 species (e.g. *Phillyrea angustifolia*, *Quercus ilex* and *Brachypodium sylvaticum* ssp. *sylvaticum*)
- showed maximum response at intermediate distances, and species not linked to dune vegetation
- 251 succession, such as *Brachypodium rupestre*, were found at high distances.
- 252



254 Figure 5. Species-response curves (GAMs) with respect to sea distance for species with more than 255 10 occurrences in the dataset (probability of occurrence in relation to sea distance). Only significant models are shown. Plant species are designated with the first three letters of their genus 256 257 and species names as follows: Aspacu = Asparagus acutifolius; Brarup = Brachypodium rupestre; Brasyl = Brachypodium sylvaticum ssp. sylvaticum; Carfla = Carex flacca; Clefla = Clematis 258 *flammula*; *Euppep = Euphorbia peplus*; *Lotcyt = Lotus cytisoides*; *Phiang = Phillyrea angustifolia*; 259 *Pipmil = Piptatherum miliaceum; Queile = Quercus ilex ssp. ilex; Rhaala = Rhamnus alaternus* 260 *ssp. alaternus; Silvul = Silene vulgaris; Sixatr = Sixalix atropurpurea; Smiasp = Smilax aspera.* 261 262

GAMs to test understorey species trends in relation to P. pinea or P. halepensis canopy cover 263 showed four species in both cases that responded significantly to binomial distribution (Appendix 264 265 D). Sixalix atropurpurea and Quercus ilex showed a higher probability of occurrence at lower canopy cover of *P. pinea*, whereas species such as *Brachypodium sylvaticum* ssp. sylvaticum and *B*. 266 267 rupestre preferred higher canopy cover of P. pinea (Figure 6a). For P. halepensis a common trend 268 of focal dune species was found (Lotus cytisoides, Sporobolus virginicus), showing higher occurrence at lower canopy cover of the pine and then sharply decreasing, whereas *Piptaptherum* 269 miliaceum, which is not a focal dune species, showed a different trend (Figure 6b). 270



Figure 6. Species-response curves (GAMs) with respect to canopy of P. pinea (A) and P. halepensis
(B) on species with more than 10 occurrences in plots (probability of occurrence in relation to
canopy cover). Only significant models are shown. Plant species are designated with the first three
letters of their genus and species names as follows: Brarup = Brachypodium rupestre; Brasyl =
Brachypodium sylvaticum ssp. sylvaticum; Lotcyt = Lotus cytisoides; Pipmil = Piptatherum
miliaceum; Queile = Quercus ilex ssp. ilex; Sixatr = Sixalix atropurpurea; Spopun = Sporobolus
virginicus.

281 **4 Discussion**

Different outcomes were obtained when pine canopy cover was included in or excluded from our dataset. When tree pine canopy cover was included, floristic differences were evident between pine forest types. After pine canopy cover was excluded from analysis, the pattern disappeared and communities converged into a single cluster with less difference in plant species composition. This suggests that the different coastal pine forest types are not characterized by distinct floristic and ecologically different understoreys. Pine stands therefore do not differentiate a specific species

pool. Vascular plant species can generally colonize plantation forests regardless of canopy species 288 289 when habitat characteristics are appropriate (Carnus et al., 2006). Since all the pine forest stands studied shared substantially the same coastal dune environment, known to be a very selective 290 291 habitat for plants (Maun, 2009), the lack of floristic differentiation between them can therefore be explained by their relatively similar environmental conditions (Frelich et al., 2003). Moreover, other 292 authors have considered environmental conditions to be more important drivers of understorey 293 community composition than dominant tree species (Piwczynski et al., 2016). In the same 294 295 perspective, species composition may be highly variable and change substantially in response to environmental change in each pine forest type. Stands of different pine types, planted across the 296 297 sea-inland gradient where vegetation zonation ranges from pioneer communities of embryonic dunes to pine plantations, have been found to host a variety of species, mainly distributed according 298 299 to this gradient and often adapted to coastal environments near the sea shore (psammophilous 300 species) (Acosta et al., 2003).

Although MRPP and NMDS suggested low distinctiveness at community level, T statistics 301 302 indicated that different pine forest stands showed some differences in species assemblages, even when pine canopy cover was excluded from analysis. INSPAN supported this thesis, highlighting 303 that a small pool of species was more likely to occur in one pine forest type than another. Notably, 304 305 20 species showed significant affinity for Pinus halepensis stands, 14 of which were focal for coastal dunal habitats (Biondi et al., 2009; Biondi & Blasi, 2015) and in some cases were strictly 306 psammophilous. Among these, Sporobolus virginicus and Elvmus farctus ssp. farctus, typical of 307 embryonic Mediterranean dunes (Acosta et al., 2007; Biondi et al., 2009; 2012), showed the highest 308 indicator values for P. halepensis forest. On the other hand, in line with Biondi et al. (2009) and 309 310 Biondi & Blasi (2015), forest species were linked to P. pinea stands, where Brachypodium sylvaticum ssp. sylvaticum was the species with the highest indicator value. This may be related to 311 312 the fact that P. halepensis stands were often planted closer to the sea, whereas P. pinea stands were

planted relatively far from the coastline and therefore had a higher probability of occurrence along 313 314 most of the sea-inland gradient. In fact, P. pinea has been reported to have a wide range of distribution in coastal dune habitats, mostly growing on fixed dunes and in areas with deep water 315 316 table and relatively low salinity (Antonellini and Mollema, 2010; Angiolini et al. 2013). The spatial location of these two pine forest types, positioned slightly differently with respect to each other and 317 extending from the foredunes to the innermost dunes (paleodunes), reflects the different ecology of 318 the pine species. Indeed, P. halepensis is more pioneer, well adapted to drought (Mauri et al., 2016) 319 and able to spread by seed from plantations into adjacent natural plant communities (Higgins & 320 Richardson 1998; Lavi et al., 2005), occasionally also colonizing the inner part of the foredunes. 321 Thus, the indicator species for P. halepensis forest type were largely very specialized 322 psammophilous plants that grow exclusively on embryonic dunes, tolerating salt spray, drought and 323 unstable substrates (Garcia-Mora et al. 1999; Acosta et al., 2000). According to Maestre & Cortina 324 325 (2004), late-successional plant species are rarely observed in *P. halepensis* plantations, even after several decades. On the contrary, the species pool of P. pinea stands consisted of forest 326 327 undergrowth species, including nemoral and basically sciaphilous plants typical of evergreen oak 328 forest, typical of the natural vegetation of inner dunes and dune slack transition zones (Acosta et al., 2003). In fact, also Brockerhoff et al. (2003) found that some, especially older, pine stands showed 329 affinities with nearby native forests, allowing establishment of many native forest understorey 330 species. 331

When we tested the distance to coastline as a driving force of coastal pine forests, our results indicated that it had a significant role for pine understorey assemblages, according to its key function for the composition of natural sand dune communities (see Forey et al. 2008; Kim & Yu, 2009; Angiolini et al., 2013; Ruocco et al., 2014). Like on non-forested dunes, forest pine plant species followed the natural vegetation zonation along this gradient, with dune species, such as *Elymus farctus* ssp. *farctus* and *Lotus cytisoides*, occurring closer to sea, and forest species of

backdune evergreen oak forests, such as Quercus ilex ssp. ilex, Brachypodium sylvaticum ssp. 338 339 sylvaticum and Carex flacca, in the intermediate stands (Acosta et al., 2000). On the contrary, ruderal and generalist species such as Brachypodium rupestre, Piptapterum miliaceum, Bellis 340 perennis and Cynosurus echinatus had the highest probability of occurrence in more inland stands. 341 We observed a relationship between understorey assemblages and the sea-inland gradient with most 342 of the focal species for the natural dune zonation. Thus, these planted forests partially maintain the 343 previous assemblage of coastal vegetation, according to their inclusion in the EU priority habitat 344 2270. 345

346 Pine canopy is an important factor in Mediterranean forests, affecting plant species assemblages, 347 seedling establishment and soil fauna (McIntosh et al., 2016; Granados et al., 2016; Bonari et al., 2016; Selvi et al., 2016). Our data confirmed that P. halepensis and P. pinea canopies significantly 348 affected understorey plant communities, particularly at high coverage. Thus, plant assemblages 349 were affected similarly by the tree canopy, regardless the pine species. Typical dunal herbaceous 350 and shrub species, such as Lotus cytisoides, Ononis variegata, Sporobolus virginicus, Clematis 351 352 flammula and Juniperus oxycedrus ssp. macrocarpa, which are heliophilous, often growing on oligotrophic soils, had higher probability of occurrence with lower pine cover. Interestingly, not 353 only the herbaceous layer was affected by pine cover, as already shown by Madrigal-Gonzaléz et al. 354 355 (2010), but also shrub and tree species, including holm oak (Quercus ilex ssp. ilex), which had lowest probability of occurrence under high canopy cover. As suggested by Lemenih et al. (2004), 356 plantations with denser canopies host lower density and richness of woody species in the 357 understorey. On the other hands, Brockerhoff et al. (2003; 2008) also found that understorey 358 vegetation beneath the canopy of pine plantations may show a successional trend towards 359 360 increasing dominance of native shade-tolerant species that are typical of natural forest understoreys. In the pine forest stands considered in this study, an evident pool of forest species, including shade-361 362 tolerant species, was only found Brachypodium sylvaticum ssp. sylvaticum and Asparagus

363 *acutifolius*, while we detected an increase in the alien invasive Acacia cyanophylla and for

364 generalist species, such as *Fumaria capreolata* ssp. *capreolata*, *Brachypodium rupestre*, *Elymus*

365 *repens* ssp. *repens*, *Piptapterum miliaceum* and *Bellis perennis* at higher canopy covers.

366

367 **5** Conclusion

We studied Italian coastal pine forest stands, analysing the influence of different pine species, sea-368 369 inland gradient and pine canopy cover on their understoreys at community and species level. Our results support the concept of pine understoreys as a "floristic cauldron" without any clear floristic 370 371 differentiation between different forest types. However, minor differences in understorey assemblage were also observed and they seem related to different pine the pine species 372 characteristics, sea-inland gradient and canopy cover. In fact, coastal dune vegetation was relatively 373 374 preserved under pine canopy. This supports the idea that coastal pine forests may maintain the "valuable" Mediterranean coastal biodiversity pool. This could be related to the fact that all the pine 375 species planted were inside their original range of distribution, as well as to long established 376 377 planting. However, since stands with higher canopy covers negatively affect understorey species assemblages, forest management can make a positive contribution. Our results provide insights into 378 the role of Mediterranean coastal pinewoods and afforestations. Species-specific management does 379 not seem necessary for these forests and their understoreys, whereas other factors such as sea-inland 380 gradient and canopy cover should be taken into account. Based on our results, we could affirm that 381 382 pine stands should no longer be considered a potential threat to biodiversity. In fact, Mediterranean pinewoods do not reflect an "ecological desert" but can host many focal species and a mosaic of 383 natural habitats in the understorey, providing also ecosystem services that should be not overlooked. 384

385

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