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# Sex-related spatial segregation along environmental gradients in the dioecious

## conifer, Taxus baccata

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## 1 Abstract

2 Sex-related differences in dioecious woody plants commonly result in spatial segregation of the sexes and a male-biased sex ratio, leading to strongly structured populations. We report results 3 from more than 1000 precisely georeferenced individuals of a threatened European dioecious tree 4 5 species, *Taxus baccata* in central Italy. We hypothesized differences between males and females in both spatial distribution and growth pattern. We used point pattern analysis (PPA) to evaluate 6 the spatial distribution of trees as influenced by dioecy and tree size, and dendrochronological 7 methods to study growth pattern differences between males and females. We found a dominance 8 of female trees giving a SR of 0.79, which increased along a gradient of elevation and slope. At 9 10 the microsite level, female *Taxus* trees were clumped to a scale of 30 m while male trees were 11 randomly distributed. We observed a significant effect of both sex and decade in the growth pattern of *Taxus* trees, where annual radial growth of females was 8.3% greater than that of 12 13 males. Female trees are more commonly found in wetter sites and males are widely but sparsely distributed, causing the absence of a clear spatial segregation of the sexes. Observed differences 14 in spatial and growth patterns of males and females have the potential to influence the climate 15 change response of T. baccata and other dioecious tree species, given that range expansion 16 depends upon the availability of seed-bearing, female trees that tend to be clumped in climatically 17 18 favorable sites.

19

## 20 Keywords

Dioecy; Tree-ring; Point pattern analysis; Spatial Segregation of the Sexes; Multi-scale approach

## 24 Introduction

25 Ecological differences between males and females in populations of dioecious plants have been demonstrated by several studies (e.g. Bierzychudek and Eckhart 1988). Sex ratio (SR), defined as 26 the proportion of male to female trees, strongly influences the reproductive success and 27 conservation status of dioecious species (Grant and Mitton 1979). Particularly important for 28 reproductive success is the spatial distribution of the sexes that is often influenced by 29 environmental heterogeneity (Hultine et al. 2007). In harsh environments with limited resources, 30 microhabitat differences are responsible for the spatial segregation of the sexes (SSS -31 Bierzychudek and Eckhart 1988). The underlying hypothesis of SSS is that females require more 32 33 resources for reproduction than males, leading to the prediction that females are more common in high-resource microsites and males in low-resource microsites (Hultine et al. 2007). The 34 reproductive biology that causes SSS can thus contribute to the vulnerability of dioecious trees to 35 global change because females, requiring a greater reproductive effort, typically show slower 36 growth (de Jong and van der Meijden 2004; Iszkulo et al. 2009). On the poorer sites where males 37 dominate, lower seed availability can contribute to increased potential for regeneration failure in 38 the more resource limited portions of the species range, possibly accelerating species decline on 39 the -trailing edgell in a climate change context (Hampe and Petit 2005). However, if SSS due to 40 41 niche partitioning is adaptive, then climatic changes could diversify resource availability and a 42 subsequent concentration of females on wetter sites.

The occurrence of male and female plants on different sites, and thus their spatial pattern,
can be used as an index of gender-environmental covariance and environmental heterogeneity
(Iglesias and Bell 1989). The influences of dioecy on the spatial pattern of tree populations have
rarely been studied (Gibson and Menges 1994; Nanami et al. 1999; Hultine et al. 2007), because

the majority of studies on dioecious species have been conducted without considering male and
female trees separately (e.g. Garcia et al. 2005; Martínez et al. 2010).

Spatial point pattern analysis (PPA - Moloney 1993; Wiegand et al. 2007) can be used to analyze 49 spatial association patterns in plant communities (Ripley 1981; Stoyan and Stoyan 1994). These 50 51 techniques quantify how neighborhood density changes with distance, thus providing a scaling approach that is valuable for describing spatial heterogeneity in community ecology (Levin 52 1992). The contribution of dioecy to the spatial structure of tree communities can be measured by 53 performing a bivariate analysis that shows at different scales if there is significant attraction or 54 55 repulsion between males and females of the same species. 56 T. baccata is a dioecious tree species that plays an important role for the biodiversity of

European forests (Linares 2013). T. baccata is a relict species that is declining due to poor 57 regeneration (e.g. Sanz et al. 2009), habitat fragmentation (Dubreuil et al. 2010), and reduced 58 59 genetic variation (e.g. González-Martínez et al. 2010). Other studies have assessed the population structure (Piovesan et al. 2009), structural diversity (Ruprecht et al. 2010), dendrochronological 60 potential (Moir 1999), radial growth (Cedro and Iszkulo 2011), sexual dimorphism (Iszkulo et al. 61 2009) and regeneration pattern (Devaney et al. 2014; Vessella et al. 2015) of T. baccata, but 62 sexual variation in distribution pattern and spatial associations among males and females remain 63 64 unknown. We tested the hypothesis that sex-related trait differences cause spatial segregation of the sexes at three different scales: 65

66 1. SR varies consistently with water availability as expressed by elevation and precipitation
67 gradients across the species range throughout Europe.

SR is influenced by water availability as expressed by topographic gradients (elevation
slope, continentality) within our 162-ha study area in central Italy.

70	3. Sex-specific ecological differences cause spatial segregation of the sexes (SSS) at
71	microsite scales: female trees are clustered in more productive sites, whereas males are
72	located on more stressful sites.
73	Additionally, we hypothesized that resource limitation associated with dioecy influences growth
74	patterns following sexual maturity: male trees grow more than female ones.
75	
76	Materials and methods
77	Study area
78	Macchia delle Tassinete (MdT) has been a floristic protected area since 1974 and is now a Site of
79	Community Importance (SCI) included in the Natura 2000 network (SCI IT5330013). MdT is
80	located in the vicinity of Cingoli village (43°21'N; 13°12'E) in the Marche region, central Italy.
81	The research area occupies 162 ha on predominantly north-facing slopes with elevation ranging
82	from 450 to 770 m a.s.l. The bedrock is limestone or marlstone and cambisols of the _Calcic
83	Siltic Chernozem' are the dominant soil type. Annual precipitation (1921 - 1990) averages 912
84	mm and mean annual temperature (1950 - 1990) averages 13°C (Cingoli, 631 m a.s.l.). MdT is
85	classified as a _mesoxeric Orno-Ostryetum' forest type (IPLA 2001), but tree species dominance
86	changes locally. Most frequent species are Ostrya carpinifolia Scop., Fraxinus ornus L., Quercus
87	cerris L., Fagus sylvatica L. and Taxus baccata L. Other locally abundant species are Corylus
88	avellana L., Acer opalus Mill. subsp. obtusatum (Waldst. & Kit. ex Willd.) Gams, Laburnum
89	anagyroides Medik., Sorbus torminalis (L.) Crantz, Sorbus domestica L., Acer platanoides L.,
90	Quercus pubescens Willd., and Acer campestre L. The current forest structure is the result of
91	previous coppicing on hardwood species and release of Taxus standards. The historical presence
92	of Taxus at MdT is documented by several historical monuments and documents in the Cingoli

village (Avicenna 1644). *Taxus* trees have been protected since 1500, in order to sustainably use
their valuable wood and to attract birds for hunting (Appignanesi 1982). However, the entire
MdT forest experienced clear-cut harvesting ca. 1870, for land consolidation after the Italian
unification process.

97

#### 98 *Field and laboratory methods*

In 2011, all *Taxus* trees having a DBH > 15 cm were tagged and mapped with a Trimble GeoXT
GPS receiver and a differential correction through Pathfinder Office 4.2 software was performed
(0.5-1 m estimated accuracy). For each of 1030 mapped *Taxus* trees, we recorded sex, DBH, total
height, canopy height, and four canopy radii. Sex could be determined for 951 (92.3%) of *Taxus*trees.

From a subsample of 10% of *Taxus* trees, selected according their diameter frequency distribution, we collected two orthogonal wooden cores per tree (189 cores, 110 trees) at 1.3 m height to perform a standard dendrochronological analysis. In the laboratory, all cores were glued to grooved wooden mounts and sanded to a high polish until optimal surface resolution allowed annual rings to be detected under magnification.

109

#### 110 Data analyses

111 We assessed the spatial heterogeneity of tree sex distribution adopting a multi-scale approach.

112 At the species range scale (Europe), we compared the SR of our study site with SR of other *T*.

113 *baccata* forests by using literature data (Dhar et al. 2008; Iszkulo et al. 2009; Vessella et al.

114 2015), and used simple linear regression analysis to quantify relationships between SR and

elevation, and SR and annual precipitation. At the landscape scale, we explored SR behavior

within our study site through regression analysis along tree structural (DBH and height) andenvironmental (elevation, slope and distance from the Adriatic Sea) gradients.

At the microsite scale, univariate and bivariate Point Pattern Analyses (PPA) were applied 118 to a data set of mapped *Taxus* locations (Moeur 1993) to assess tree spatial patterns within the 119 120 study area and the spatial associations between the two sexes or two DBH classes across spatial scales from 0 - 50 m. The spatially explicit dataset that was used in the analyses comprised 419 121 males and 532 females, 784 small (DBH < 30 cm) and 56 large (DBH > 40 cm) trees. We used 122 pair-correlation functions g (r) (Stoyan and Stoyan 1994) to avoid possible misinterpretation of 123 results due to the cumulative effect typically found in other methods such as K-function analysis 124 (Ripley 1977; Perry et al. 2006). The univariate pattern of Taxus trees as a whole was contrasted 125 against the heterogeneous Poisson null model to account for first-order effects. This null model 126 differs from complete spatial randomness (CSR) in that a function  $\lambda(x, y)$  varies with location (x, 127 y) but maintains the independence of the occurrence of any point with respect to any other 128 (Wiegand and Moloney 2004). For the bivariate analyses between males vs. females and small 129 vs. large trees, we applied the random labeling null model that simply randomizes tree category 130 without changing geographic position (Wiegand and Moloney 2004). The latter is a conservative 131 method to test the significance of spatial association between two categories (e.g. males vs 132 females). The 95% confidence intervals for both univariate and bivariate analyses were computed 133 from 999 Monte Carlo simulations (Stoyan and Stoyan 1994; Wiegand and Moloney 2004) and 134 the goodness-of-fit (GoF) test for null hypothesis was performed (Diggle 2003). All analyses 135 were done only for classes with more than 15 trees (Camarero et al. 2000), applying a 1 m lag 136 distance and a maximum distance of 100 m with the grid-based software Programita (Wiegand 137 and Moloney 2004) adopting a grid size of 1 m2 and a ring width of 5 m. 138

Tree-ring widths were measured using a LINTAB semi-automatic optical system (Rinntech 139 140 2003) at a precision of 0.001 mm. All tree-ring series were first visually and then statistically checked for cross-dating and measurement errors using the CDI (Cross Dating Index) run by 141 TSAP-Win software (Rinntech 2003). Mean tree-ring chronologies for male and female Taxus 142 143 trees were compiled from selected crossdated series using the ARSTAN program (Cook and Holmes 1997). To remove interannual frequency variations each tree-ring series was standardized 144 with a spline function using a 50% frequency response. Statistical indices were calculated on a 145 common interval of 100 years (1909-2009) in order to compare male and female chronologies. 146 Tree-ring statistics included: the mean sensitivity (MS) which is the mean percent change from 147 each measured yearly ring value to the next to assess the high-frequency variation of the series 148 (Fritts 1976); the first-order autocorrelation on raw (AC) and standardized data (AC1); the 149 average correlation between series (RBAR) that was calculated for all series using the running 150 means method on overlay periods of 50 and 25 years; the Expressed Population Signal (EPS), a 151 measure of how the mean chronology derived from the sample trees represents a hypothetical 152 infinite replicated chronology (Wigley et al. 1984). Additionally, cumulative ring width and basal 153 154 area increment (BAI) curves were obtained from the mean chronologies of male and female trees. A general linear mixed-effects model (GLMM) was used to detect radial growth differences 155 between males (n = 32) and females (n = 38) over the 1920 - 2010 time period. Radial growth 156 (decadal mean) was log transformed prior to analysis, to normalize distributions and stabilize 157 variance of residual errors. Log transformed radial growth (RG-ln) was used as the response 158 variable and sex and decade were used as independent factors. Individual tree identity was 159 160 specified as a random intercept term to allow the overall growth rate to vary by tree (accounting for differences in genetics, microsite variability, etc.). GLMM was fitted by using the Linear and 161

162 Nonlinear Mixed Effects Models package (nlme 3.1-105 ver.) within the R statistical software163 (2.15.2 ver.).

164

- 165 **Results**
- 166 *Sex ratio at broad scales*
- 167 We found 419 males and 532 females, giving an overall SR of 0.79 (Table 1). Males were
- slightly larger (25.6 cm VS 25.14 cm DBH) and shorter (6.52 m VS 6.65 m height) than females,
- but these differences were not significant (t = 0.729; p = 0.47 for DBH and t = -1.02; p = 0.309
- 170 for height).

172 Table 1. Summary statistics (mean, median and standard error) of structural attributes and environmental variables

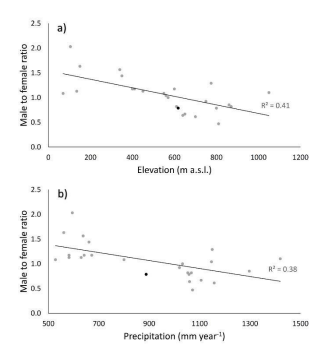
173	of <i>Taxus</i> trees at MdT study site	. DBH = diameter at breast	theight (cm); $H = I$	neight (m), $V = Volume (m3)$ .
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SEX (n)	Statistics	DBH	Н	Crown H	Crown V	Elevation	Slope	Sea Dist.
		(cm)	(m)	(m)	(m <sup>3</sup> )	(m a.s.l.)	(°)	(Km)
Males (419)	Mean	25.61	6.52	1.89	77.35	700.76	24.23	39.73
	Median	24.00	6.50	1.90	58.11	700.00	25.20	39.79
	SE	0.47	0.09	0.03	3.42	1.11	0.33	0.02
Females (532)	Mean	25.14	6.65	1.98	80.31	699.05	22.72	39.78
	Median	23.00	6.50	2.00	56.37	699.22	23.62	39.85
	SE	0.41	0.08	0.02	3.91	0.87	0.31	0.01
Undetermined (79)	Mean	25.01	6.95	1.80	99.92	640.66	21.63	39.33
	Median	22.00	6.60	1.80	62.30	681.12	24.69	39.33
	SE	1.08	0.24	0.07	10.47	8.68	1.21	0.07

At the species range scale, the MdT study site occupies an intermediate position in a topoclimatic gradient of 24 European research studies reporting the SR of *Taxus* (Fig. 1). The MdT site fits within an annual precipitation range (750 – 950 mm) that was rarely considered in previous studies, and has a lower SR than expected for its precipitation level. Across the range of *T. baccata*, the SR decreased with increasing elevation and precipitation.

179

Figure 1. Male to female ratio (SR) observed at MdT (black dot) related to elevation (a) and precipitation (b)
compared to other 25 European forests (grey dots) with *T. baccata* (Dhar et al. 2008; Iszkulo et al. 2009; Vessella et al. 2015).



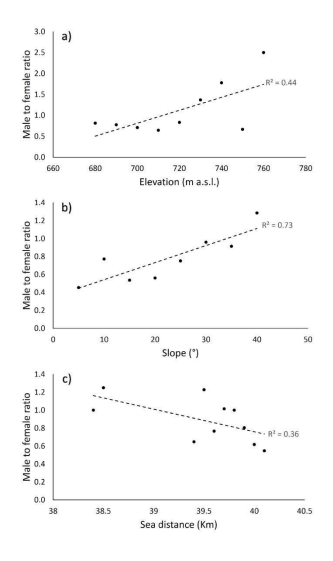
183

184

At the landscape scale, SR increased with elevation and slope steepness and decreased with
distance to the Adriatic Sea (Fig. 2). SR approximated unity in those sites located at lower
elevations and closer to the sea.

189 Figure 2. Male to female ratio (SR) observed at MdT against environmental gradients of elevation (a), slope (b) and

190 distance from the Adriatic Sea (c).



191

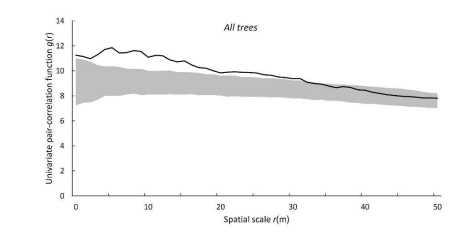
192

#### 193 *Fine scale spatial pattern*

*Taxus* trees, pooled among both sexes and all DBH classes, showed significant aggregation ( $p \le 0.01$ ) to a distance of 30 m (Fig. 3). When considered separately in univariate analyses (Fig. 4), males were randomly distributed, while females were weakly grouped ( $p \le 0.01$ ) at 15 and 35-40 m. The univariate g (r) function showed a clear tendency towards aggregation ( $p \le 0.01$ ) for small trees, whereas large trees were randomly distributed. By assessing the univariate pattern of a

```
combination between DBH and sex, we observed a similar pattern in males and females. Small
trees were weakly aggregated (p \le 0.01) while large trees were randomly distributed.
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Figure 3. Univariate analysis of the spatial pattern of 1030 *Taxus* trees at MdT. The bold line indicates the pair correlation function g(r), and shaded areas encompass the non-significant, i.e. random, distribution and represent
 points within the 2.5th and 97.5th percentile g(r) values of the 999 simulations of the null model assuming a
 heterogeneous Poisson distribution.

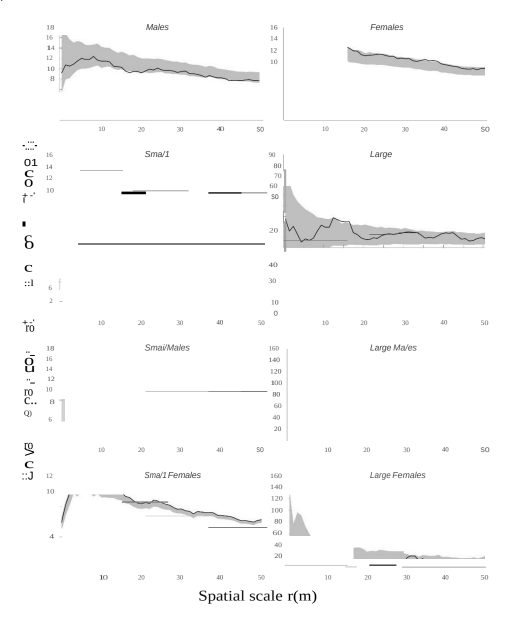


The bivariate spatial association between sexes showed a random pattern (Fig. 5).

However, significant spatial segregation ( $p \le 0.01$ ) was observed between small and large

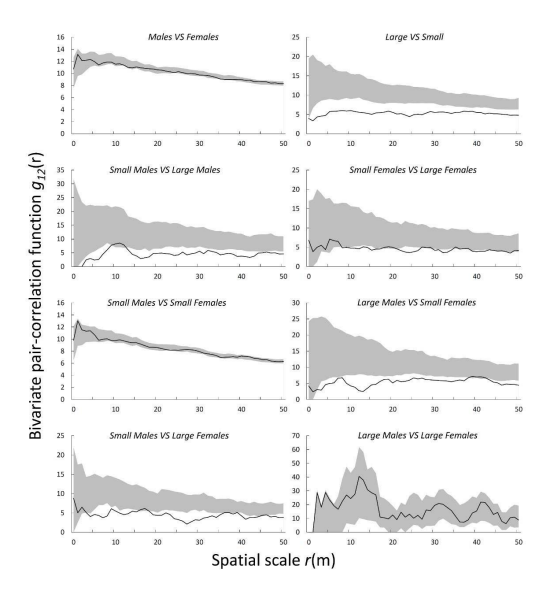
210 diameter trees, with stronger segregation for males than for females.

Figure 4. Univariate analysis of the spatial pattern of *Taxus* trees, considering sex (males and females), DBH (small and large), and their combinations (e.g. small males and large females), using the pair-correlation function g(r). Black lines indicate the pair-correlation functions g(r), and shaded areas encompass random distribution and represent points within the 2.5th and 97.5th percentile g(r) values of the 999 simulations of the null model (Random Labeling).



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- 225
- 226
- 227

Figure 5. Bivariate analysis of the spatial pattern of *Taxus* trees, considering sex (males and females) and DBH
(small and large), and their combinations (e.g. small males VS large females), using the pair-correlation function
g12(r). Black lines indicate the pair-correlation functions g12(r), and shaded areas encompass random distribution
and represent points within the 2.5th and 97.5th percentile g12(r) values of the 999 simulations of the null model
(Random Labeling).



234 *Growth pattern* 

Male and female tree-ring chronologies were well crossdated and showed similar values in most
dendrochronological descriptors (Table 2): time spans were 1879-2010 for males and 1881-2010

for females. Males and females showed similar mean ring width (0.99 mm and 1.06 mm), mean 237 sensitivity (high-frequency variability), first-order autocorrelation (low-frequency variability), 238 correlation between tree-ring series (Rbar), and adequacy of the chronologies (EPS). The two 239 BAI curves (Fig. 6a) show similar trends particularly in recent years, with a growth oscillation of 240 approximately 30 yrs caused by a coppice rotation cut on hardwood species in 1984 and the 241 resulting canopy regrowth. The periods of greatest differences between male and female growth 242 were 1907-1915, 1943-1959, and 1963-1975. Finally the cumulative ring width curves showed a 243 coincident growth pattern of male and female Taxus trees up to the 1950's and a slightly 244 increasing divergence thereafter (Fig. 6b). 245

The GLMM showed a significant effect of sex on radial growth of *Taxus* trees (p = 0.028). Annual radial growth of female trees was 8.3% greater than that of male trees (95% CI: 0.9% to 15.1%), after accounting for decade of growth and the random effects of growth variability among individuals growing in different microsites.

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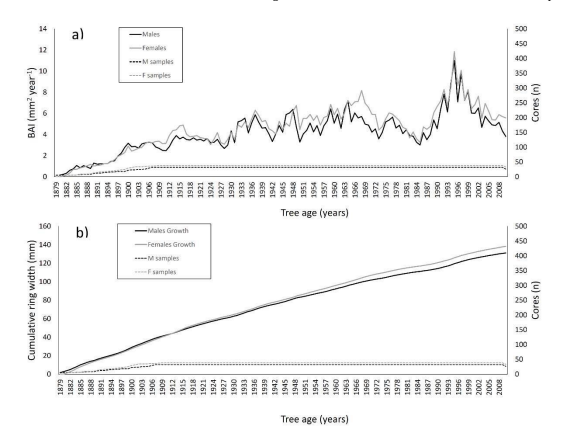
Table 2. Comparison of males and females chronologies by common statistical dendrochronology indices (MRW:
mean ring width; MI = mean index value; Rbar = average correlation between series; EPS = expressed population
signal; MS = mean sensitivity of the average chronology; AC1 = first-order autocorrelation on raw data; AC =
autocorrelation on standardized data).

Series	Time Interval	Common Interval	MRW	MI	Rbar	EPS	MS	AC1	AC
(n)	(yrs)	(yrs)	(mm)						
Males (32)	1879-2010	1909-2009	0.99	0.97	0.16	0.86	0.16	0.7	-0.1
Females (38)	1881-2010	1909-2009	1.06	0.98	0.14	0.86	0.12	0.8	-0.2

255

256

Figure 6. Basal Area Increment (a) and cumulative ring width for males (n = 32) and females (n = 38) at MdT study
site. Continuous lines indicate BAI and cumulative ring width and dotted lines indicate the number of sampled cores.



260

261

### 262 **Discussion**

263 *Sex ratio at species range and landscape scales* 

At the regional scale, our study site occupies an intermediate position along a European environmental gradient where the SR decreased with increasing elevation and precipitation (Fig. 2). We observed SRs that were female-biased, and lower than would be expected given regional levels of precipitation and elevation. Female-biased populations have also been found on wetter sites in the Alps (Hilfiker et al. 2004) and in the Andes (Nuñez et al. 2008), adding support to the hypothesis that the proportion of females increases with water availability. At the landscape-scale in our MdT study area, we observed a high SR on steeper, higher elevation sites, the opposite relationship as observed for the regional-scale comparisons. This apparent discrepancy likely arises because the sites closer to the ridge are more water-limited because of shallow, rocky soils, and thus are least favorable for female *Taxus* trees. On the contrary, at a regional scale higher elevation sites coincided with increased water availability (Iszkulo et al. 2009).

Males were less abundant than females at MdT and the difference between them in tree size (DBH and height) was not significant. These findings are partly discordant with other studies on *T. baccata* of central Europe that found larger and more abundant males on drier sites (Iszkulo et al. 2009).

SR increased with increasing diameter, but decreased with increasing height consistently with other European studies on SR of *Taxus* trees (e.g. Iszkulo et al. 2009). Taken together, these data suggest that males put more carbon into diameter growth, and females into height growth, when they are in the older growth stages. However, this could also be a matter of where they grow: possibly the females, growing on the wetter sites, have more side-pressure from competing trees and so allocate more carbon to height growth.

286

287 Influence of dioecy on spatial patterns at microsite scales

Adult *Taxus* trees (DBH > 15 cm) were strongly grouped at fine spatial scales (< 30 m) at MdT, probably representing a pattern of repeated clustering. On the contrary, other authors have found random distribution in adult *Taxus* trees (Martinez et al. 2010). Considering the two sexes separately, we observed a weak but clustered pattern in females and a random pattern in males. The observed patterns were consistent with the hypothesis of males being the more stress tolerant that can occupy a broader spectrum of suitable microsites (Hultine et al. 2007; Iszkulo et al.2009).

We did not find spatial segregation of the sexes (SSS) at MdT, meaning that there was not 295 significant aggregation nor repulsion between males and females. This was probably because 296 female trees were clustered and more common in wetter portions of the elevation gradient, 297 whereas males were widely distributed. Homogeneity of habitats at fine scales can be another 298 possible cause of absence of spatial segregation of the sexes (Ueno et al. 2007; Schmidt 2008; 299 Iszkuło et al. 2011). However, female Taxus trees at MdT exhibited a clear tendency towards 300 clustering in wetter sites. Another possible explanation for the absence of SSS at MdT could be 301 that seed dispersal in T. baccata is primarily by birds causing a wide distribution of seeds 302 (Hubbell 1979; Murray 1988) that can in turn weaken the effect of dioecy on the spatial 303 heterogeneity of trees (Nanami et al. 1999). Birds can spread seeds across long distances, but 304 often distribute seeds in a clustered manner (e.g. burial sites for seed-caching species; below 305 favorite roosting sites). 306

Nevertheless, we found spatial segregation between smaller and larger trees, a pattern 307 already described for adult *Taxus* trees and regeneration in other European sites (Dovčiak 2002; 308 Piovesan et al. 2009; Devaney et al. 2014). As expected, small trees were aggregated for both 309 males and females, whereas large trees were randomly distributed. The spatial segregation 310 between smaller and larger trees is probably due to a negative intraspecific neighborhood effect 311 that has been previously reported for *T. baccata* (Dovčiak 2002; Piovesan et al. 2009; Devaney et 312 al. 2014) and for other species (He and Duncan 2000). Moreover, the traditional coppice system 313 with dense Taxus standards may have favored the establishment of Taxus regeneration into forest 314 canopy gaps close to adult Taxus trees. 315

316

#### 317 Influence of dioecy on growth patterns

The mean chronologies of male and female *Taxus* trees emerged as being very similar showing 318 that the overall growth patterns were analogous. However, despite wide variability among 319 individual trees, female *Taxus* trees in our study area showed a significantly higher growth rate. 320 Nonetheless, other authors have reported a radial growth reduction of female trees associated 321 with the achievement of sexual maturity in the studied population (e.g. Obeso 2002; Bañuelos 322 and Obeso 2004; Cedro and Iszkulo 2011). Female individuals may require more resources than 323 males because of higher reproduction effort (Iszkulo et al. 2009). A possible explanation of the 324 325 counter-intuitive pattern observed at MdT is that female growth can be favored by the more 326 productive sites that they occupy, as opposed to males that occur across a wide variety of microsite conditions. Where female trees of dioecious species occupy more favorable sites than 327 males, greater resource availability may compensate for greater reproductive requirements. Other 328 authors found no significant differences between sexes in Araucaria angustifolia mean radial 329 growth in South America, but they observed a higher growth rate of females in the last two 330 decades (Cattaneo et al. 2013). 331

Our results highlight that in order to relate observed spatial patterns in dioecious species 332 333 to habitat conditions, we need to consider abiotic and biotic constraints of tree establishment. Future research should investigate the spatial pattern of male and female trees along wide 334 environmental gradients, in order to better understand the role of climate and topography in the 335 SSS process. Site conditions, habitat variability, and water availability in particular are expected 336 to strongly influence the strength of segregation between sexes in dioecious species (Dawson and 337 Bliss 1989; Dudley 2006). Spatial distribution modeling of dioecious species requires accounting 338 339 for the different resource requirements and distribution patterns of male and female trees,

340	particularly in the climate change context where range expansion to favorable sites depends upon
341	the availability of seed-bearing, female trees (Tognetti et al. 2012; Hultine et al. 2013).
342	
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350	
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