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## Biogeography and shape of fungal fairy rings in the Apennine mountains, Italy

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### 26 Abstract

Fungal "fairy rings" (FR) are regular bands of vegetation caused by a centrifugal expansion of fungal 27 mycelia. It is well established that FR affect both soil chemistry and microbiome, promoting plant 28 coexistence at community scale. However, nothing is known about the distribution of these peculiar 29 patterns at regional scale. Here, we report on the FR biogeography of in the Apennine mountains 30 across a 300 km latitudinal gradient, comparing twelve sites with different elevation, geomorphology, 31 and climate. From high-resolution images we gathered data concerning FR shape, colonial density, 32 and distribution. Overall, 1,163 FR structures were found across the study area. Three type of spatial 33 34 patterns could be identified, proper rings, i.e. circular vegetation structures, arcs representing circle sections, and rotors where an arc was showing narrower circular distortions at its end points. Arcs 35 36 were the most common shape followed by rings, while rotors were rare, accounting for less than 2% of all observations. Fungal colony density varied largely both among and within sites, averaging 6.7 37 38 arcs, 2.2 rings and only 0.1 rotors per ha. On average, arcs (18.8 m) were similar in size compared to rotors (18.4 m) while rings were smaller (11.7 m). Notably, arcs present a higher frequency of 39 40 occurrence on steeper slopes while rings are mostly found on flat and moderate slopes. In relation to climate, FR occur within the altitudinal range between 546 m and 2,148 m a.s.l., corresponding to 41 42 temperature between 3.4°C and 12.7°C. and rainfall between 1,100 and 1,300 mm per year. No colonies were found with an annual rainfall lower than 800 mm. Based on this study, FR are common 43 elements of the Apennine grasslands landscape contributing to the maintenance of plant and microbial 44 diversity. A better systematic identification of the fungal species involved in the FR formation is 45 necessary in relation to the different environmental gradients observed in the study area. Furthermore, 46 other investigations and modelling demonstrations are required to understand the dynamic processes 47 behind the development of the different FR forms.. 48

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50 Keywords: Plant-soil feedback, Autotoxicity, Arcs and rotors, *Agaricus arvensis*, Basidiomycota,
51 Ecosystem engineer,.

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## 53 1. Introduction

Fungal "fairy rings" (FR) are circular bands of vegetation caused by the centrifugal expansion of 54 fungi (Edwards, 1984, 1988). More than 50 species of basidiomycetes and few ascomycetes have 55 been reported to be able to create FRs (Fox, 2006; Halisky & Peterson, 1970) in different ecosystems, 56 including sand dunes (Abesha, Caetano-Anollés, & Høiland, 2003), woodlands (Dowson, Rayner, & 57 Boddy, 1989; Peter, 2006), and grasslands (Karst, Dralle, & Thompson, 2016; Yang, Li, Liu, & 58 Zhang, 2019). FRs, spreading radially in the soil as fronts of dense mats of fungal mycelia, only 59 60 become evident above ground because of the sporadic appearance of the fungal fruiting bodies or 61 because of their effect on vegetation (Gregory, 1982). Shantz and Piemeisel (1917) classified grassland FRs into three types based on the response of vegetation: Type-1 formed by an external belt 62 63 of bare soil or dead vegetation surrounding a second belt of luxuriant vegetation; Type-2 recognizable for the presence of a luxuriant belt without any dead zone; Type-3 periodically revealed only by the 64 65 presence of fungal carpophores with no detectable changes in vegetation color and floristic composition. 66

67 The occurrence of FRs in grasslands has been the subject of scientific studies since the end of the 19th century (Elliott, 1926; Evershed, 1884; Lawes, Gilbert, & Warington, 1883; Shantz & 68 Piemeisel, 1917), particularly in the context of turfs management where they are considered 69 70 destructive phenomena (Fidanza, Colbaugh, & Davis, 2000; Filer, 1965). Several attempts have been made to explain the detrimental effect of FR fungi on vegetation. The most cited hypotheses are the 71 72 local increase in soil hydrophobicity causing the dry out of the soil (Gramss, Voigt, & Bergmann, 2005), the immobilization of nutrients in the mycelium body (Fisher, 1977), the direct pathogenic 73 behavior (Fidanza et al., 2007; Terashima, Fukiharu, & Fujiie, 2004), and the release of phytotoxic 74 compounds like cyanide (Blenis, Chow, Duncan, & Knowles, 2004; Caspar & Spiteller, 2015). 75 Differently, the mechanisms triggering the formation of luxuriant vegetation in FRs were ascribed to 76 the flush of mineral nutrients after the fungal passage, the formation of vacant niches, the release into 77 78 the soil of molecules with hormone-like activity also called "fairy chemicals", or the stimulation of a 79 symbiotic microbiome with a positive effect for the plants (Choi et al., 2010; Suzuki et al., 2016; 80 Yang, Li, Zhang, Liu, & Zhang, 2018; Zotti et al., 2020).

In the last decade, several studies also revealed the impact of FRs at ecosystem level. Type-1 FRs, by killing the established vegetation, create gaps that are filled by short-lived plants, thus enhancing species coexistence and plant diversity at community level (Bonanomi, Mingo, Incerti, Mazzoleni, & Allegrezza, 2012). More recently, the extensive use of next generation sequencing provided the means to study the impact of FRs on soil microbiome in terms of bacteria and fungi composition (Oh, Fong, Park, & Lim, 2016; Xing et al., 2018). Recently, Zotti et al. (2020) revealed that the passage of Type-1 FRs, caused by *Agaricus avernsis*, causes profound changes in the microbiota composition with an overall increase in taxa diversity. In fact, the passage of the FRs wipe out the established bacterial and fungal communities, promoting the recolonization of disturbance related taxa. Taken together, available evidence indicates that FRs fungi are crucial for the maintenance of plant and microbial diversity in grasslands acting as ecosystem engineers in grasslands.

Most of the previous studies investigated the impact of FRs at community scale, focusing on 93 soil chemistry (Edwards, 1984), plant communities (Yang et al., 2019), and soil microbes (Espeland, 94 95 Caesar, Sainju, Lartey, & Gaskin, 2013). However, very little is known about the factors controlling the distribution of FRs at regional scale. Recently, Karst and coauthors (Karst et al., 2016), using 96 high-resolution aerial imagery, documented that in United States the FRs colonies can develop in 97 different shapes, i.e. complete ring, arc-shaped, and rotor. The same paper reported, on mathematical 98 99 simulations reproducing the observed spatial patterns without investigating the link between ecological factors and geographical distribution of the observed different shapes of FRs. In this work, 100 101 we attempt to describe the biogeography of fairy rings at regional scale and propose a mechanistic explanation for the induction of the FR types. To this aim, we carried out a study on twelve sites in 102 103 the Italian Apennine mountains, ranging across high geomorphological, climatic and latitudinal variability, of ecosystems in which FRs exist. This allowed us to set the biological limits of 104 occurrence of such peculiar phenomenon. 105

We collected, from high-resolution images, data concerning FR shape, colonial density and distribution. Thereafter, we analyzed the relationships between fungal colonies data with selected climatic and geomorphological variables to explore the factors that affect FR distribution. The specific objectives of our study were:

- 110
- to assess the abundance and relative occurrences of different FR types, i.e. rings, arcs, and rotors;
- 112 (ii) to explore the association between geomorphology and FR types;
- 113 (iii) to reveal the distribution of FR in relation to temperature and rainfall;
- (iv) to define a new hypothesis on the causal process explaining the occurrence of different
   FR types

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117 2. Material and Methods

(i)

- 118
- 119 *2.1 Study area*

The Apennines is a ~1,000 km long mountain chain that span in latitude from 38°N to 44°N accounting 261 peaks above 2,000 m a.s.l. and The Apennine rock is mainly composed by limestone substrata, with arenaceous-pelitic flysch present only in the Laga groups (Bonanomi et al., 2020). The climate is a mountain variant of the Mediterranean type, with mean annual temperature that range from 0 °C to 8 °C in January and from 20 °C to 28 °C in the hottest month, July. Total annual precipitation also largely varies, from less than 600 and more than 3,000 mm, with abundant winter snowfall above 1000 m a.s.l.

The current treeline, dominated by the deciduous tree *F. sylvatica*, occur at elevation that range from ~1,000 m a.s.l. to ~2,100 m a.s.l. (Bonanomi et al. 2020). Because of past intensively human exploitation for agricultural purposes (i.e. cultivation, livestock, wood, charcoal production), the current treeline stay several hundred meters below its potential climatic limit that in Central and Southern Italy is above 2,000 m a.s.l. Consequently, grassland under ~2,000 m a.s.l. are considered secondary i.e. created by past human activity, with only high-elevation grassland being primary or of natural origin (ref\_marina).

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## 135 2.2 Study sites and survey design

To identify the study sites we capitalized from our recent survey of *F. sylvatica* distribution across all Apennine peaks where we examined 3,622 km of treeline and adjacent grasslands (Bonanomi et al., 2018). Based on this survey, we identified twelve mountains where FR were present and clearly visible from satellite images (Figures 1, 2).

A first data set build with the aims of assessing the FRs density in terms of number of fungal 140 colonies per ha, and the relative occurrence of different forms, i.e. ring, arc, rotor. To achieve this 141 aim, in each of the twelve mountains, and in grasslands where FRs were present, we randomly 142 selected five 1 ha plots. A total of 60 ha was surveyed: 5 plots for each of the 12 mountains. The 143 selected sites fulfilled the following requirements: i) grassland are not subject to extensive shrub 144 encroachment, i.e. less than 30% cover of either Juniperus communis or Pinus mugo; ii) rocks do not 145 cover more than 50% of the surface. In each plot, using Google Earth Pro<sup>™</sup> (Google, Inc. Mountain 146 View, CA, USA) images dating 2004-2018, we counted all occurring FRs and identified their forms 147 for a total number of observations of 547 FRs (Figure 2). 148

A second data set was built to assess the relationships between FRs occurrences with local geomorphology and climate. In detail, we selected additional sites located in the surrounding of the twelve mountains (Supplementary Table S1). Here, we extensively searched for all occurring FRs that was mapped in Google Earth Pro<sup>TM</sup> (Google, Inc. Mountain View, CA, USA) images dating 2004-2018. The geographical coordinate, the shape, the size (diameter in m), and the elevation of each FR were recorded for a total of 616 observations (Table 1). The vertical accuracy of the Google Earth-measured data was further checked in the field on 100 FR, corresponding to 16.2% of geolocated waypoints, distributed along the Apennines, with a Garmin Montana® 600 GPS device equipped with a barometric altimeter. For each FR the slope was measured as average slope of the transect crossing them. In addition, for each fragmented forms of FR (arcs and rotors) we assessed if the convexity was upslope or downslope.

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### 161 *2.3 Climatic variables*

162 Based on the literature concerning higher plants biogeography in Apennines (Bonanomi et al., 2018), we selected 10 variables to evaluate their ability to explain FR distribution (Table 1). Climatic data 163 164 were gathered from the global climate database WorldClim 2.0 http://www.worldclim.org; (Hijmans, Cameron, Parra, Jones, & Jarvis, 2005) at a spatial scale resolution of 1 km<sup>2</sup>. In the WorldClim 165 166 database, temperature variables are calculated as the spatial average of grid squares that include the selected point and adjusted according to their average elevation. Data were then adjusted at the 167 elevation of each study sites by using a 0.006°C m<sup>-1</sup> adiabatic lapse rate (Barry, 1992). For rainfall 168 variables, data were gathered from the Bioclimates of Italy because of the higher spatial resolution 169 170 compared with WorldClim for the Apennines. Shortly, rainfall data were obtained with a calibration process, via regression kriging, of the WorldClim (1.4), with a large data set from local 171 meteorological stations (Pesaresi, Biondi, & Casavecchia, 2017). 172

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## 174 *2.4 Colonies migration*

To assess FRs migration, we used the diachronic approach using satellite images of 2006 and 2012 available in Geoportale Nazionale (www.pcn.minambiente.it). Despite extensive searching, we found only a limited number of FR (N=10) where high-resolution images were available at both dates. For each FR we measured the diameter in 2006 and 2012 and the annual increment was calculated.

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## 180 *2.5 Data analysis*

Density in colonies per ha across the twelve mountain ranges selected were assessed by average values on five replicates in the same study site. Comparison of FRs density were made by FRs forms if were rings, arcs or rotors for the totality of the Apennine region and between each study site. With the same approach we compare data of relative abundance between the different forms and between the different study site across the Apennine. To obtain and evaluation of the more frequent diameter by which arcs, ring and rotor are observed we assess separately their distribution by applying normal fitting distribution curves. Distribution of rings, arcs and rotors were also visualized by the relation

with climatic variables including temperature and precipitation variables. Comparison of the highest 188 frequency of occurrence for rings and arcs were tested for the range of slope values observed in the 189 different study sites. To emphasize the relation between slope, elevation and diameter of rings and 190 arcs we performed bubble-plots between slope and elevation with labels proportional to the diameter 191 of the FRs sampled. We also visualized the occurrence of all the FRs sampled and divided by rings 192 and arcs by linear regression between slope and annual cumulated rainfall. The last three analysis 193 were conducted uniquely on arcs and rings because the lower contribution of rotors and the rarity of 194 this particular formations. Statistical analyses and plotting were carried out using Statistica 10 195 196 software (StatSoft, Tulsa, OK, USA).

- 197
- 198 **3. Results**
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# 200 *3.1 Form, size, and density*

Overall, 1,163 FRs were recorded across the Apennines. Arcs were largely the most common shape 201 202 (75.1%) followed by ring (23.5%) (Figure 3). Rotors, although present, were rare accounting for only 1.4% of all observations. The relative abundance of different FR forms largely varied among 203 204 mountains, with arcs being the most common in ten sites (elencare siti) while rings only in two (Montiego and Fiegni). Rotors, instead, were found only at three sites (elencare siti). Considering all 205 study sites, we found an average density of 6.7 arcs, 2.2 rings and only 0.1 rotors per ha. Colonies 206 density varied amount mountains, with the highest values at Vettore (19.2 colonies per ha), followed 207 by four sites where the density ranged between 10 and 15 colonies per ha, and the other sites where 208 we found a density between 5 and 10 colonies (Figure 3). 209

On average, rings have smaller diameters (11.7 m) compared to arcs (18.8 m) and rotors (18.4 m). However, the size of all shapes largely varies with most of the rings falling in the dimeter classes between 5 and 15 m, with the largest ring having a diameter of 51.9 m. The majority of arcs are in the diameter classes between 10 and 20 m, but reaching the notable maximum size of 137.1 m, while the population of rotors is limited, ranging in size between 4.2 and 33.0 m (Figure 4).

- The annual average migration of the FRs was  $64.7 \pm 17.5$  cm per year, with the highest values of 151.3 cm year<sup>-1</sup> and the lowest of 40.3 cm year<sup>-1</sup>. For arcs, we always observed that the fungal colonies migrate upslope (Supplementary Figure S4).
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## 219 *3.2 Elevation, latitude and climate*

Rings, arcs and rotors showed a similar distribution in relation to elevation and climatic variables.
Arcs showed a bimodal distribution in relation to elevation, with peaks between 1,000-1,200 m a.s.l.

and 1,600-1,800 m a.s.l. and had a slightly larger elevation range than rings and rotors (Figure 5a), but no FRs were recorded below 546 m a.s.l. and above 2,148 m a.s.l. Concerning latitude, we found a significant negative relation between FR altitude and latitudinal position (Pearson r = -0.82; Supplementary Figure S1).

FR distribution was bimodal also in relation to mean annual temperature (Figure 5b). Arcs showed the largest amplitude, ranging from a minimum of 3.4 °C to a maximum of 12.7 °C of mean annual temperature. The distribution of FR to other temperature variables was similar to that observed for mean annual temperature (Supplementary Figure S2).

FR distribution in relation to annual rainfall was unimodal, slightly skewed on the left (Figure 5c). Most FRs occurs in the class of precipitation between 1,200 and 1,300 mm per year, with no cases below 813 mm and above 1,668 mm. Also in this case, arcs showed a larger ecological amplitude compared to rings and rotors. Monthly and seasonal rainfall relationships with FR distribution was similar to that of annual cumulated data (Supplementary Figure S3).

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### 236 *3.3 Slope and fairy rings*

Rings and arcs showed a different distribution in relation to mountain slope (Figure 6). Overall, rings were more common in flat and moderately sloped areas while arcs were common also on very steep slopes. In detail, 55.7% of the rings were found in areas with a slope lower than 10%, with only 2.7% occurring in steep areas (slope >30%). Instead, 57.2% of the arcs were found in areas with slopes between 10% and 30%, with 13.2% of the cases in steep slope. Notably, 99.7% of the recorded arcs showed an upslope convexity.

Moreover, we found different relationships between slope, elevation and the size of rings and arcs (Figure 7). Specifically, we found that large rings are mainly present at low elevation (<1,400 m a.s.l.) and in flat or moderately sloped areas. Instead, we found that very large arcs also occur at high elevation and in steep areas. Finally, we observed a positive correlation between slope and annual cumulated rainfall (Figure 8a). Notably, the correlation was steeper for rings than for arcs (Figure 8b).

### 249 **4. Discussion**

Previous studies showed that fungal FR exert major effects on soil chemistry (Yang et al., 2019), microbial populations (Zotti et al., 2020), and plant diversity (Bonanomi et al., 2012). Here, we provide the first assessment of FR biogeography at regional scale. We found that rings, arcs and rotors occur with different densities and relative abundances across the Apennine mountain range, with climate and slope playing an important role in explaining the observed variability.

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# 256 *4.1. Variation in size, form and density of fairy ring*

257 FR density, size and forms varied greatly within and among sites. Rings are more common on flat areas, while arcs are more frequent in steep slopes. Rotors, on the other hand, are always very rare. 258 However, in some sites (i.e. Montiego, Ocre and Vettore) rings, arcs and rotors coexist in close 259 proximity, suggesting that local factors could explain their cooccurrence. Commonly, colonization 260 261 strategies of fungi are divided in vegetative strategies or production of spores (Dix, 2012). As demonstrated for *M. oreades* in sand dunes (Abesha et al., 2003), the initiation of FRs probably 262 263 derives from fungal spores that, after germination, spread centrifugally. The relative low density of FRs in the field coupled with the long-lived nature of these structures and the high amount of spores 264 265 produced by basidiomycetes suggest that spore establishment is a rare event (De Groot et al., 2012). In the very early phases of development, FRs are not noticeable aboveground, also after careful 266 inspection in the field, probably because the mycelium does not reach a sufficient biomass to affect 267 plants. Concerning FRs population structure, in the smallest size class (i.e. 0 to 5 m in diameter), rings 268 are far more common than arcs. On the contrary, arcs are much more common than rings in the size 269 class over 10 m in diameter. The shifting of FRs population structure, i.e. more common rings of 270 small size compared to arcs, suggest that arcs may generate from rings during their ontogenetic 271 expansion. Here, we propose two possible mode of arcs generation. In the first model, a complete and 272 regular ring brakes during its expansion due to the encounter with an obstacle of physical (i.e. a rock 273 274 or unsuitable soil patches) or biological nature (e.g. mycoparasites or another fungal front). We found several FRs with circular shape but with a missing short portion (Supplementary Figure S4a), 275 suggesting that some factors generated a rupture of the fungal front. Thereafter, as the fungal front 276 expands, the break becomes progressively larger generating the arcs. In this regard, we believe that 277 the encounter of different fungal fronts, i.e. FR coalescence, would be a common event leading to the 278 emergence of irregular shapes. However, the coalescence of two rings, while creating new shapes 279 departing in time from a regular circle, would not justify the rupture of the ring continuity and the 280 appearance of partial rings or arcs. (Supplementary Figure 4b). 281

At microscopic level the fate of fungal hyphae that belong to different genets during colonies 282 283 coalescence is largely unknown although often described as repulsion (Deacon, 2013), but at macroscopic level the rings indeed can lose their integrity (Parker-Rhodes, 1955; Stevenson & 284 285 Thompson, 1976). A similar effect resulting from rings coalescence was also presented by Carteni et al (2012) in the context of ring forming clonal plants. Accordingly, Karst et al. (2016) showed how 286 different structures can rise by the intersection of different expanding simulated colonies. The 287 "obstacle" hypothesis is consistent with the observation that large rings (>20 m in diameter) are rare 288 while arcs can often reach diameter >50 m, with some cases where the diameter are over 100 m. 289

A Fairy ring can spread at a rate of about 60 cm per year, then an arc of 50 m would be ~75 years old, and the largest observed arc (137 m) would have more than 200 years. These are not exceptional size and age for basydiomicota colonies, e.g. *Armillaria spp.* clones can cover hundreds of ha aging than 1,000 years (Bendel, Kienast, & Rigling, 2006). Such lifespan suggest that a fungal front is likely to encounter obstacles thus generating arcs from rings.

Although the "obstacle" hypothesis seems plausible and probably is operative in many cases, it does not explain the prevalence of arcs over rings in steep slope, and the case of low density of the colonies corresponding to low probability of crossing trajectories between different structures.

298 Our hypothesis to explain the formation of arc structures from initial smaller complete rings is based on the effect of autotoxic compounds in interaction with their leaching. Autotoxicity is a 299 phenomenon reported for microbes (Berne, Kysela, & Brun, 2010) and higher plants (Singh, Batish, 300 & Kohli, 1999) caused by localized accumulation of chemical compounds either actively produced 301 by the considered organism, or released during decomposition of its parts likelitter and fine roots. 302 303 For fungi, autotoxicity has been reported to control spore germination, and impair hyphal growth and proliferation (Bottone, Nagarsheth, & Chiu, 1998; Chitarra, Abee, Rombouts, Posthumus, & 304 Dijksterhuis, 2004). Several classes of chemicals have been considered as putative autotoxic 305 compounds, including fatty acids, alkaloids, and phenols, but none of these justifies a species 306 specificity of the effect. Recently, Mazzoleni et al. (2015) discovered the species-specific inhibitory 307 effect of extracellular DNA on plants and confirmed that the phenomenon was generally occurring 308 309 across a broad range of organisms of different kingdoms, including plants, insects, bacteria and fungi (Mazzoleni et al 2015b, Carteni et al. 2016). The inhibition by extracellular self-DNA released into 310 311 the soil by the decomposition of organic matter has unique features to provide a mechanistic explanation for the formation of FRs and also for their differentiation into different shapes. In fact, 312 DNA is soluble in water and so it is spatially distributed according to main water flows. In the FR 313 context, this logically support the theoretical framework of the formation of arcs. In flat areas, the 314 315 local accumulation of autotoxic compounds due to the mycelium turnover, would trigger, in

interaction with depletion of resources, the centrifugal hyphae growth with the development of 316 317 circular structures. A similar dynamics has been described for tussock grasses and plants capable of vegetative propagation that forms the so called hollow crowns, central dieback or monk's tonsure like 318 gaps (Bonanomi et al., 2014). In the absence of slope, the autotoxic compound is mostly leached 319 320 vertically along the soil profile, according to soil texture and hydraulic properties (Heuvelink, Burgers, Tiktak, & Van Den Berg, 2010). Under such conditions, modelling exercise demonstrate the 321 creation of ring structures and their possible coalescence into continuous waves across the landscape 322 (Cartenì et al JTB). Differently, on slopes the directional water movement creates a down slope 323 324 diffusion of the putative hydrosoluble autotoxic factor (with accumulation of the adverse effect on the lower section of the ring. If the autotoxic effect is strong enough, the lower fungal front would be 325 326 inhibited, and arcs would arise. This hypothesis is consistent with the observation that 99.7% of the arcs move upslope, with no cases of colonies moving downslope (Supplementary Figure 4c). On the 327 328 other hands, we found few rings in very steep slopes, with the few cases occurring in areas with substantial annual rainfall over 1,300 mm per years. We suggest that in theses slopes, the co-329 330 occurrence of abundant rain and rocky, porous soils may have promoted a faster vertical leaching of the autotoxic compound, thus explaining the persistence of some rings also on slopes. Theoretically, 331 332 the prevalence of vertical versus lateral water flow can be the causal factor for the maintenance of a 333 ring structure, whereas a dominance of down slope directional flow of water will cause the lower ring rupture with the creation of the arc pattern. The Vettore site (Figure 2 and Supplementary Figure 4) 334 is characterized by the co-presence of all these features, i.e. abundant annual rainfall, rocky and 335 heterogeneous soil and very steep slopes, and, consistently with our model, here arcs and rings largely 336 coexist. Further field work is ongoing to assess within FRs and arc structures the gradients of 337 concentration of the fungal DNA to observe its consistency with the proposed model of autotoxicity 338 in both flat and sloped areas. This study will use next generation sequencing, including shotgun 339 methods (Mendes, de Lima Brossi, Kuramae, & Tsai, 2015) to investigate the self-DNA hypothesis. 340

Finally, we found that rotors are rare (1.4% of all cases) and occurs only in few places at very low density (less than 0.1 colonies per ha). Karst and coauthors (Karst et al., 2016) also noticed that rotors occurs far less frequently than rings and arcs in US grasslands. Moreover, the author's simulation model suggested that soil patches with low resources could promote rotation in the advancing fungal front. Probably conditions suitable for rotors development are very special and occur infrequently in nature, in this regard fine-scale investigation of soil heterogeneity is required to test the Karst's hypothesis.

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### 349 *4.2. Climate, geomorphology and fairy ring*

At a global scale climate is the primary factor controlling the distribution of living organisms, with 350 351 mean annual temperature that is considered a key parameter to explain biological distribution along latitudinal and altitudinal gradients (Bennie, Duffy, Inger, & Gaston, 2014; Körner, 2012). However, 352 353 much less is known about fungal biogeographic distribution (Tedersoo et al., 2014), and few 354 concerning FRs (Shantz & Piemeisel, 1917). In this context, our study provides a wider description 355 of FR distribution in relation to temperature and rainfall. In general, we found that FR mostly occurs within the temperature range between 4°C and 12°C, with altitudinal upper distribution likely limited 356 by low temperature. In our survey, the highest FR occur at 2,148 a.s.l. where the mean annual 357 358 temperature of ~3.4°C likely poses a climatic constraint upon the growth of FRs. In addition, at this elevation in Apennine a substantial shift in grasslands plant composition and structure occurs (ref. 359 360 Marina), with a possible reduction of the relative abundance of the plant species that the fungus use as a substrate. It will be interesting to further investigate the different fungal species altitudinal 361 362 distribution and their interactions with the plant communities. Our current knowledge of fungal species forming FR is sparse, based on either morphological and molecular methods we were capable 363 364 to identify Agaricus arvensis at the Puro-Rogedano site (Zotti et al., 2020). Here, the bi-modal distribution of FR in respect to mean annual temperature with two peaks at 5-6°C and 10-11°C 365 366 suggest that different fungal species may form FR in cold and mild climatic conditions. Indeed, 367 further studies are urgently required to assess the taxonomy of species that forms arcs and ring along the investigated altitudinal and latitudinal gradient. 368

FRs rarely occurs below 800 m, with the lowest occurrence at 546 m where the mean annual 369 temperature is ~12.7°C. Moreover, with found a strong negative correlation between altitude and 370 latitudinal occurrences of FR, meaning that in Central Italy we found arcs and ring at lower elevation 371 compared to Southern Italy. such altitudinal range and its shift with with latitude suggest that 372 temperature plays a critical role on FR distribution. First, the absence of FR below ~500 m could be 373 related to a shift of fungal colony spreading mode. We suggest that when the fungi lives in optimal 374 temperature condition, their mycelium show a "phalanx like strategy" (ref.), producing the circular 375 pattern, reflecting vigorous growth of the fungal front. Instead, at the edge of the ecological niches 376 377 where growth conditions are not optimal, e.g. at low altitude with high temperature, the fungi are unable to form a compact fungal front showing a weaker spread by single hyphae, classifiable as 378 "guerrilla strategy". Thus not producing FR structures. A second hypothesis to explain the lacks of 379 FR at lower altitudes is related to land-use in Apennine mountains. In fact, at lower elevation these 380 landscape are heavily managed with both pastures and grassland being replaced by intensive 381 agricultural system and urban areas (ref marina). This, indeed, could contribute to the rarity of FR 382 383 but cannot explain their absence in the residual grasslands present at low elevation.

Alongside temperature, precipitation also affects FRs distribution with most of the cases that 384 occurs where rainfall is between 1,100 and 1,300 mm, with no cases below 800 mm. This indicate 385 that FRs cannot proliferate in arid and semi-arid climates. On the other hands, we do not found FRs 386 where rainfall exceed 1,700 m. In the field we observed that FR do not occur in flat area subject to 387 periodical flooding, suggesting that basidiomycetes fungi cannot tolerate prolonged anoxic soil 388 conditions (Tavzes, Pohleven, & Koestler, 2001). According, we found that ring and arcs occurs both 389 in flat and sloped areas in the range between 800 and 1,400 mm of annual rainfall, whereas over the 390 1,400 threshold we mainly found FRs only where the slope exceed 20%. In general terms, many 391 392 basidiomycetes including wood degrading species are less abundant and perform poorly under anoxic condition (ref, ref). Overall, our data indicate that both shortage and excessive rainfall, below 800 393 394 mm and above 1,700 mm, respectively, could hamper FR development. However, in flat areas subject to flooding, because of poor drainage conditions, a reduction of the upper limit of annual rainfall was 395 396 recorded, corresponding in this case to 1,400 mm per year.

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## **398 5. Implications for management and future directions**

FRs are notable features of grassland ecosystems and available evidence demonstrate that these fungi 399 400 promote plant and microbe coexistence at local scale. This study, recording the presence of FR across 401 a whole Mountain range, albeit within a specific set of environmental conditions. FR occur at density of several colonies per hectare, highlight their importance for maintenance of biodiversity in 402 Mediterranean grassland. In this regard, FR fungi operate as ecosystem engineers and by killing the 403 established vegetation and microbiome, allow the periodical flourishing of life by ecosystem 404 rejuvenation. However, FR occurs only in permanent grassland and moved pastures and we observed 405 that they disappear after shrub encroachment, a comment vegetation dynamics in Apennine because 406 of land abandonment (Bonanomi, Incerti, & Allegrezza, 2013; Malandra, Vitali, Urbinati, & 407 Garbarino, 2018). Specific management plan could be applied in some sites of special values (e.g. 408 409 Puro-Rogedano, Fiegni, Montiego) to maintain summer moving and avoid J. communis establishment and encroachment. 410

Further advances in understanding FR dynamics are needed to explain the formation or arcs, ring and rotors. The combination of long-term field manipulative experiments and modeling work would help to falsify the "obstacle" and the "autotoxic" hypotheses proposed to explain the formation of arcs from rings during the ontogenetic development of fungal fronts. Finally, a substantial knowledge gap still requires the taxonomic identification of fungal species that forms FR across altitudinal and latitude gradients in Apennine.

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Variable	Unit		
Geographical			
Latitude	degree		
Elevation	m a.s.l.		
Slope‡	%		
Climatic			
Mean annual temperature (MAT)	°C		
Mean monthly temperature <sup>*</sup>	°C		
Max temperature of warmest month	$^{\circ}\mathrm{C}$		
Min temperature of coldest month	$^{\circ}\mathrm{C}$		
Temperature annual range	$^{\circ}\mathrm{C}$		
Mean temperature of warmest quarter	$^{\circ}\mathrm{C}$		
Mean temperature of coldest quarter	$^{\circ}\mathrm{C}$		
Mean cumulated annual rainfall	mm y <sup>-1</sup>		
Mean cumulated monthly rainfall <sup>†</sup>	mm mo <sup>-1</sup>		

Table 1. Variables accounted for the analysis of FR across the Apennines range.

‡ Slope measured above, across, and below the treeline.

\* Monthly mean temperature from January to December.

<sup>†</sup> Monthly cumulated rainfall from January to December.



Figure 1. The twelve study sites (black dots) selected across the Apennine Mountains range (Italy).

Figure 2. Selected satellite images of three study sites (Fiegni, Montiego, and Vettore) with photostitched examples of a ring, an arc and a rotor. Bars for scale.



**Figure 3**. Density and relative occurrence of different fungal colony types. Average of fungal colonies per ha across all study sites (a) and in each mountain (b). Relative abundance of the different shapes in either whole data set (n=547) (c) or in each site (d).



**Figure 4.** Distribution in classes of size (diameter in m) of rings (above), arcs (middle), and rotors (below). All observations for rings (n=183), arcs (n=416), and rotors (n=17) are reported. Red lines represent the best fitting normal distribution. Dashed lines show average values.



**Figure 5.** Distribution of rings, arcs, and rotors in relation to elevation (above), mean annual temperature (middle), and annual rainfall (below).



**Figure 6**. Distribution of rings (grey bars and area) and arcs (white bars and area) in relation to slope. Dashed lines refer to mode values. Red contours lines represent best fitting normal distribution.



**Figure 7.** Bubble plot showing distribution of rings (up) and arcs (down) in Apennine grasslands in relation to slope and elevation. Label size for rings (circle) and arcs (triangle) is proportional to sample diameters



![](_page_25_Figure_0.jpeg)

**Figure 8**. Linear regression between slope and annual rainfall for all FR (above) and separately for arcs and rings (below). Straight lines represent the fitted regressions. We must add some stat

**Supplementary Figure S1**. Linear regression between FR altitude and latitude across Apennines. Black circles indicate the position of each FR, straight lines represent the fitted regressions.

![](_page_26_Figure_1.jpeg)

Supplementary Figure S2. ..... Bars plot distribution with all other temperature variables

Supplementary Figure S3. ..... Bars plot distribution with all other rainfall variables

**Supplementary Figure S4**. Selected images of different FR types. A: examples of rings with missing short portion of the fungal front (yellow arrows) at Fiegni site; B: examples of FR coalescence, i.e. the encounter of two fungal fronts (yellow arrows), at Montiego site; C: a population of arcs migrating upslope at the Rogedano site. Bars for scale.

![](_page_28_Picture_1.jpeg)