

Ecology and life history of *Meta bourneti* from Monte Albo (Sardinia, Italy) (#29230)

1

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
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Ecology and life history of *Meta bourneti* from Monte Albo (Sardinia, Italy)

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
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Underground environments and related biodiversity are still relatively understudied. Even widespread cave-dwelling species show a considerable paucity of information regarding their ecology and life traits. This is the case of one of the most common cave predators occurring in Europe and in the whole Mediterranean basin: the orb-web spider *Meta bourneti*. Although the congeneric *M. menardi* represented the model species in several studies, *M. bourneti* was considered very little and available information regarding this species is founded on observations performed on a handful of populations. Therefore, further studies are required to produce a more complete species overview. In this study I analyzed data on *M. bourneti* spiders collected in caves of Monte Albo (Sardinia, Italy) throughout a full year. I used binomial Generalized Linear Mixed Models to analyze spider occupancy inside cave environments. The same data were also analyzed with Generalized Linear Models, an approach which allows weighting of species absence based on its detection probability. Generalized Linear Mixed Models were used to analyze spider abundance. Analyses on *Meta bourneti* occupancy and abundance were repeated three times: for all individuals and for adults and juveniles separately. Finally, Linear Mixed Models were used to detect possible divergences in underground spatial use between adult and juvenile spiders. Although widespread on the whole mountain, *M. bourneti* generally showed low density and low detection probability; most of the individuals observed were juveniles. The spiders generally occupied cave sectors with high ceilings and deep enough to show particular microclimatic features; adults tended to occupy less illuminated areas than juveniles, while the latter were more frequently found in sectors showing high humidity. The abundance of *M. bourneti* was strongly related to high humidity and the presence of two troglophile species (*Hydromantes flavus* and *Oxychilus oppressus*); morphological sector features promoting predators' avoidance positively influenced the abundance of juveniles. However, when adults only were considered, no

significant relationships were found. Adults and juvenile spiders did not differ in spatial distribution inside the caves studied, but a seasonal distribution of the species along cave walls was observed. Microclimate appears to be one of the most important features affecting both presence and abundance of *M. bourneti* in underground environments. Individuals tended to occupy a lower height during hot seasons, probably looking for more suitable microclimatic conditions. This study represents a further  useful to better comprehend the ecology of these widespread cave-dwelling spiders.

Ecology and life history of *Meta bourneti* from Monte Albo (Sardinia, Italy)

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
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13 Abstract

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INTRODUCTION

Underground environments (from shallow cracks and burrows to the deepest karst systems) are peculiar habitats showing a characteristic combination of environmental features: they generally show little or no light, high air humidity and a relatively stable temperature resembling the mean annual temperature occurring in outdoor surrounding areas (Culver & Pipan, 2009; Smithson, 1991). Underground microclimate is generally shaped by the influence of external climate which, through openings connecting underground environments with outer ones, spread in and contribute to creating different microhabitats (Badino, 2004; Badino, 2010; Campbell Grant, Lowe & Fagan, 2007; Lunghi, Manenti & Ficetola, 2015). The most evident result of such influence is the formation of three different macro-ecological zones (Culver & Pipan, 2009). The zone adjacent to the connection with the outdoor is the most affected by external influences; indeed, the microclimate of this area generally resembles the environmental conditions occurring in surrounding outdoor areas. Then, there is the so-called twilight zone, where external influences are weaker and incoming light is generally low. Finally, there is the deep zone, where incoming light is absent and microclimatic features are the most stable.

Underground environments house a rich biodiversity (especially in animal species) which can show unique and peculiar adaptations to the different ecological zones (Romero, 2011). A species' degree of adaptation to cave life represents the ~~base of~~ ^{basis for} the general classification used in distinguishing between different groups of cave-dwelling ~~species~~ ^{organisms} (Novak et al., 2012; Pavan, 1944; Sket, 2008). The most specialized are called "troglobites", species closely connected to the deep areas of underground environments that evolved specific adaptations, such as depigmentation, anophthalmia, elongation of appendages, and reduction in metabolic rates (Aspiras et al., 2012; Bilandžija et al., 2013; Biswas, 2009; Hervant, Mathieu & Durand, 2000).

"trogolophile"

are

Then there are species that optionally decide to remain stable underground but still able to exit, the so-called "troglophiles"; these species can exploit different underground areas and their adaptations to cave life are reduced or even absent (*Di Russo et al., 1999; Fenolio et al., 2006; Lunghi, Manenti & Ficetola, 2017*). Finally, epigean species are randomly found in the shallowest part of underground environments are called "trogloxenes". However, this classification has turned out to be too strict (*Lunghi, Manenti & Ficetola, 2014; Romero, 2009*), as species usually thought to be accidental are indeed potential residents playing an important role for the entire ecosystem (*Lunghi et al., 2018a; Manenti, Lunghi & Ficetola, 2017; Manenti, Siesa & Ficetola, 2013*).

Despite an increasing

their

Although the undeniable increase of interest in underground ecological spaces and related biodiversity which has occurred in the last decades (see as examples *Culver & Pipan, 2014; de Freitas, 2010; Fernandes, Batalha & Bichuette, 2016; Lunghi et al., 2018e; Studier et al., 1986*), current knowledge on cave-dwelling species is still far from being considered complete. A good example is given by the troglophile orb-weaver spider *Meta bournoni*. Meta spiders are among the most common predators in cave environments (*Mammola & Isaia, 2017b; Mammola, Piano & Isaia, 2016; Manenti, Lunghi & Ficetola, 2015; Pastorelli & Laghi, 2006*). These spiders show an interesting complex life history: during their early life stages they are phototaxic and disperse in outdoor environments, while during the adult phase they become photophobic and inhabit underground environments, where they reproduce (*Chiavazzo et al., 2015; Manenti, Lunghi & Ficetola, 2015*). Meta spiders are at the apex of the underground food-chain, preying on several species using both web and active hunting (*Lunghi, Manenti & Ficetola, 2017; Mammola & Isaia, 2014; Novak et al., 2010; Pastorelli & Laghi, 2006; Smithers, 2005*). However, young spiders are in turn potential prey of other cave predators (*Lunghi et al., 2018b*). In Europe and

the Mediterranean basin area, two species of *Meta* spiders are commonly observed: *M. menardi* and *M. bourneti* (Fernández-Pérez, Castro & Prieto, 2014; Fritzén & Koponen, 2011; Mammola & Isaia, 2014; Nentwig et al., 2018). Although the former is the subject of several studies (Hörweg, Blick & Zaenker, 2012; Lunghi, Manenti & Ficetola, 2017; Mammola, Piano & Isaia, 2016; Manenti, Lunghi & Ficetola, 2015), research on *M. bourneti* is very limited (Boissin, 1973; Mammola, 2017; Mammola & Isaia, 2017a). In a recent study, Mammola and Isaia (2014) provided some of the few available data on the ecology and life history of *M. bourneti*. The authors studied the distribution and abundance of *M. menardi* and *M. bourneti* in six caves located in the north-west of Italy. Although they confirm the previously hypothesized similarities in habitat selection between the two cave-dwelling *Meta* spiders (Gasparo & Thaler, 1999), in this study it emerged that *M. bourneti* was present at warmer temperatures and showed a shift in its life cycle compared to the congeneric *M. menardi*; these findings likely result from the competition between the two species (Mammola & Isaia, 2014). However, to provide more solid knowledge on *M. bourneti* spiders, further studies involving populations from different areas are needed.

Here I provide the first information related to the ecology and life history of *M. bourneti* populations from Sardinia (Italy). This study is based on data collected by Lunghi et al. (unpublished) and aims to provide new information on the occurrence and abundance of *M. bourneti* spiders, also providing information on a possible divergence in habitat use of different age classes.

MATERIALS & METHODS

Dataset

The analyzed dataset ^{contains} ~~gathers~~ information on ~~Meta~~-*bourneti* and inhabited caves from ~~the~~ Monte Albo (north-east Sardinia, Italy) (*Lunghi et al.*, ^{unpublished}). In one of the surveyed caves the presence of the species has never been detected and thus, it will not be considered in the following analyses (N of considered caves = 6). In this area the congeneric *M. menardi* is not present and thus, no potential interspecific interactions limit habitat selection of *M. bourneti* (*Mammola & Isaia, 2014*). Surveys were performed ^{seasonally} ~~seasonally~~, from autumn 2015 to summer 2016, thus covering a full year. Inner cave environments were divided horizontally into portions of 3 m (hereafter, sectors), to collect fine-scale data on both cave morphology and microclimate, as well as on the occurrence of other cave-dwelling species (*Ficetola, Pennati & Manenti, 2012; Lunghi, Manenti & Ficetola, 2017*). Within each cave sector the following abiotic data were recorded: maximum height and width, wall ^{heterogeneity} ~~heterogeneity~~, average temperature ($^{\circ}\text{C}$), humidity (%) and illuminance (lux). ~~Furthermore~~, a standardized survey method (7.5 min/sector) was used to collect data on the presence of six cave-dwelling species: *Meta bourneti*, *Hydromantes flavus*, *Metellina merianae*, *Tegenaria* sp., *Oxychilus oppressus* ^{and} *Limonia nubeculosa* (data of the latter is integrated in the present study; Table S1). These species likely interact with *Meta* spiders, as they represent both potential prey and predators (*Lunghi et al., 2018b; Manenti, Lunghi & Ficetola, 2015; Novak et al., 2010*). *Meta* spiders were also counted and ascribed to two different categories on the basis of body size (prosoma + opisthosoma): adults with fully developed pedipalps (body size $\geq 10\text{mm}$) and juveniles (body size $<10\text{ mm}$) (*Bellmann, 2011; Mammola & Isaia, 2014; Nentwig et al., 2018*). For further information on the methodology used in data collection see (*Lunghi et al., unpublished*).

Data analyses

The following analyses were performed in R (*R Core Team, 2016*) using the packages lme4, lmerTest, MuMIn, MASS, nlme, and unmarked (*Bartoń, 2016; Douglas et al., 2015; Fiske & Chandler, 2011; Kuznetsova, Brockhoff & Christensen, 2016; Pinheiro et al., 2016; Venables & Ripley, 2002*). Analyses on detection probability, species-habitat association and abundance were performed three times, one for each group studied (all individuals, adults only and juveniles only). To model species occurrence and abundance, I considered data only from surveys in which microclimatic features were recorded (cave surveys = 31, *N* of spiders = 110).

Detection probability

Cave spiders are among the species showing imperfect detection: a species is present when it is observed, but a lack of observation does not mean its true absence (*MacKenzie et al., 2006*). I estimated the detection probability of *Meta-bourneti* on the basis of twenty-seven pairs of surveys performed in all caves and in each season with a gap < 7 days (*Lunghi et al., unpublished*), a prerequisite for population closure (i.e., no immigration or emigration occurs; *MacKenzie et al., 2006*). I considered two possible covariates influencing spider detection: the depth of the cave sector (hereafter, depth) and the season. I built three models (one for each covariate and one with none) and then ranked them following the Akaike's Information Criterion (AIC); the one with the lowest AIC value was used to estimate detection probability (*Burnham & Anderson, 2002*).

Analyses on species occurrence

I used binomial Generalized Linear Mixed Models (GLMM) to assess the relationship between *Meta* spiders and the abiotic features characterizing the cave environments. The presence/absence of the spiders was used as dependent variable, while sector's morphological (height, width and wall heterogeneity) and microclimatic (temperature, humidity and illuminance) features were used as independent variables. To evaluate whether spiders' preferences change through the year, the interaction between season and each of the considered microclimatic features considered was also included as a further independent variable. Sector and cave identity were used as random factors. For each studied group, GLMMs models were built using all possible combinations of independent variables; such models were then ranked following the Akaike's Information Criterion corrected for small sample size (AICc) (Fang, 2011). The model showing the lower AICc value was considered the best model. Following the recommendations of Richards, Whittingham and Stephens (2011), models representing more complicated versions of those with a lower AIC value and nested models were not considered as candidate models. The likelihood ratio test was used to assess the significance of variables included in the best AICc models. If necessary, variables were logarithmic or square-root transformed to better fit the normal distribution (Lunghi et al., unpublished).

Considering a potential variation in species-habitat association over time (Lunghi, Manenti & Ficetola, 2015; Lunghi, Manenti & Ficetola, 2017) and an overall low detection probability observed for these spiders (see Results), I tested the robustness of the previous analyses using a method that allows weighting the species absence on the basis of its detection probability: the General Linear Models (GLM) (Gómez-Rodríguez et al., 2012). Unfortunately, adding random factors to this analysis is impossible, hence the cave identity was included as a fixed factor. Following the same procedure described above, for each species all possible GLMs

models were built and ranked following AICc. The significance of variables included in the best AICc model was tested using the likelihood ratio test (*Bolker et al., 2008*).

Given that for some of the groups studied the best AICc model estimating detection probability included sector depth (see *Detection probability of ~~Meta~~-bourneti*), I repeated the GLM analysis for each group including depth as a further independent variable.

Analyses of species abundance

I used GLMM to assess whether abundance of *Meta bourneti* was related to both microclimatic and biotic recorded parameters. The observed abundance of spiders was used as a dependent variable, as it represents an index of true abundance (*Barke et al., 2017*). Season, along with both microclimatic (average temperature, humidity and illuminance) and biotic (presence/absence of the five considered species) features, were used as independent variables, while sector and cave identity as random factors. The significance of variables was tested with a Likelihood ratio test.

Analyses on spatial distribution

To test whether adult and juvenile *M. bourneti* show divergences in the spatial use of underground environments, I used two Linear Mixed Models (LMM) with age class (adult/juveniles) and season as independent factors, and both sector and cave identity as random factors. The two dependent variables were the distance from the cave entrance and the height above cave floor respectively. The dataset used in this analysis is shown in Table S2.

RESULTS

Overall, a total of 182 observations of *Meta-bourneti* (64 adults and 118 juveniles) were performed within the caves studied (average \pm SE = 30.33 ± 16.49 per cave). Observations of spiders were the highest in spring (3.17 spiders/visit), followed by winter (2.92 spiders/visit), summer (2.67 spiders/visit) and autumn (1.92 spiders/visit) (Fig. 1). Of 1,538 cave surveys, spiders were observed only on 153 occasions, in most of which just one spider occupied the cave sector (129) (Table S2). Occupied cave sectors showed the following microclimatic conditions: average temperature = 14.46 ± 0.16 °C (min-max; 11.25-19.45); average humidity = 91.23 ± 0.3 % (80.6-94.3); average illuminance = 2.52 ± 1.78 lux (0-156.05). In only two cases two adults shared the same cave sector, while juveniles did this more frequently (4 times with an adult and 19 with other juveniles). Two cocoons were observed during autumn, each in a different cave. One of these was observed lying on the ground, already with numerous recently hatched spiders (Fig. 2A); during winter, spiderlings abandoned the cocoon (Fig. 2B). No further information on the second cocoon is available.

Detection probability of *Meta bourneti*

In species analysis, the model including depth as covariate was the best model (AICc = 747.93) compared to the other two (model including season, AICc = 751.36; model without covariates, AICc = 751.45); *Meta bourneti* showed an overall low detection probability (0.232). Considering adults only, the model without covariates was the best (AICc = 385.86) compared to the other two (model including depth, AICc = 385.94; model including season, AICc = 389.37); adults showed higher detection probability (0.4). Finally, for juveniles the model including

season as covariate was the best ($AICc = 557.36$) compared to the other two (model including depth, $AICc = 558.25$; model without covariates, $AICc = 559.14$); detection probability of juvenile *M. bourneti* was the lowest (0.173).

Spider occurrence

Results of the two analyses (GLMM and GLM) were consistent, thus showing a substantial similarity in the identification of significant variables (Tables 1 and 2). The occurrence of *M. bourneti* was positively related to sector height and humidity; the best GLMM ~~model~~ also included the interaction between season and illuminance (Tables 1 and 2). The occurrence of adult spiders was negatively related to illuminance; the best GLM model also detected a positive relationship to sector height (Tables 1 and 2). The occurrence of juvenile spiders was positively related to sector height and humidity; a significant relationship with season was included in the best model of both analyses. The best GLMM ~~model~~ also included a significant relationship between season and illuminance (Tables 1 and 2).

Results of GLM including sector depth as a further independent variable were identical to those of the previous GLM analyses (Table S3).

Spider abundance

The abundance of *Meta bourneti* was related to sector humidity ($F_{1,481.38} = 6.61$, $P = 0.01$) season ($F_{3,518.3} = 3.36$, $P = 0.018$) and the presence of *Hydromantes flavus* ($F_{1,645.19} = 21.91$, $P < 0.001$) and *Oxychilus oppressus* ($F_{1,645.1} = 24.01$, $P < 0.001$). Spiders were more

abundant in cave sectors with high humidity and where *H. flavus* and *O. oppressus* were present. The abundance of adults showed no significant correlation with the variables considered. The abundance of juveniles showed a relationship to sector temperature ($F_{1,223.76} = 4.15$, $P = 0.043$), humidity ($F_{1,524.87} = 7.41$, $P = 0.007$), season ($F_{3,548.94} = 4.22$, $P = 0.006$) and the presence of ~~both~~ ^{both} *H. flavus* ($F_{1,645.47} = 25.06$, $P < 0.001$) and *O. oppressus* ($F_{1,645.38} = 31.33$, $P < 0.001$); juvenile spiders were generally more abundant in warm cave sectors showing high humidity and where *H. flavus* and *O. oppressus* were present.

Spider distribution

Distance from cave entrance did not differ by age classes ($F_{1,122} = 0.26$, $P = 0.608$) (Fig. 3A) nor between seasons ($F_{3,122} = 0.58$, $P = 0.626$). Vertical distribution of spiders (i.e., height from the cave floor) did not differ by age classes ($F_{1,113} = 0.85$, $P = 0.358$) (Fig. 3B) but a significant effect of season was detected ($F_{3,113} = 6.20$, $P < 0.001$); spiders were generally at a lower height during spring and summer.

DISCUSSION

~~Meta~~-*bournet*i spiders represent one of the top predators commonly occurring in Monte Albo caves; indeed, spiders were usually present in all underground environments ~~considered~~ ^{sampled}. The only cave of the dataset in which *M. bournet*i was never observed was located at an elevation exceeding 1000 m a.s.l. (Lunghi et al., unpublished); ~~there~~, unsuitable environmental conditions for the species likely occur there (Lunghi et al., 2018d; Mammola & Isaia, 2014). The highest number of spiders observed occurred in spring, a season in which invertebrates are generally

more active (*Bale & Hayward, 2010*). In the populations studied, the life cycle of *M. bourneti* differed slightly from what was observed in north-western Italian populations (*Mammola & Isaia, 2014*); in September, cocoons were already spun, and spiderlings started to emigrate in January. This variation in breeding phenology probably occurred because the two study areas are characterized by different climatic conditions (*Hijmans et al., 2005*). Indeed, it was recently shown that climatic conditions occurring at the surface can significantly influence the underground breeding activity of troglophile species (*Lunghi et al., 2018c*). However, the two data collections on *M. bourneti* were performed in different periods (2012-2013 in north-west Italy and 2015-2016 in Sardinia), it is therefore still unclear whether such a divergence was due to a change in local climate or to an annual fluctuation of climatic conditions.

Occurrence of *M. bourneti* was generally related to cave sectors showing high humidity; this variable was observed to have the same effect on juvenile spiders, while adults showed a high occurrence in cave sectors with low light (Table 2). These particular microclimatic conditions (high humidity and low illuminance) usually occur in areas far from the connection to the surface, where external influences are weaker and the microclimate is more stable (*Culver & Pipan, 2009; Lunghi, Manenti & Ficetola, 2015*). As was pointed out for both *M. bourneti* and *M. menardi*, these spiders occupy cave areas deep enough to show suitable microclimatic conditions, but still in the proximity of sites with elevated prey abundance (*Lunghi, Manenti & Ficetola, 2017; Mammola & Isaia, 2014; Manenti, Lunghi & Ficetola, 2015*). However, the tendency of *M. bourneti* to occupy cave sectors with high ceilings is just the opposite of what was observed for *M. menardi* (*Lunghi, Manenti & Ficetola, 2017*). Considering that these two species show similar hunting strategies (*Mammola & Isaia, 2014*), the different preferences of cave sector morphology may be driven by some other ecological ~~reasons~~^{factors}. For example, in cave

sectors with high ceilings, spiders may have more surface (i.e., cave wall) to escape from potential predators present in the same cave sectors (e.g., *Hydromantes* salamanders; *Lunghi et al.*, 2018b). Indeed, sector height was particularly significant for juveniles, while for adults this variable was not included in the best AICc model (Tables 1 and 2).

Analyses of spider abundance identified both environmental and biological features as potential determinants. In cave areas with high humidity, *Meta-bourneti* showed the highest abundance. Furthermore, the presence of two of the species considered (*Hydromantes flavus* and *Oxychilus oppressus*) had a strong influence on spider abundance. While it is possible that *M. bourneti* shares the same microhabitat preference with these species (*Ficetola et al.*, in press), trophic interactions between *M. bourneti* and these two species may explain this particular association (*Lunghi et al.*, 2018b; *Mammola & Isaia*, 2014). However, results from spider abundance analyses must be carefully interpreted. The majority of observations were related to juveniles (~73%) and this may have biased the analysis performed at species level. Indeed, results from the two analyses (all spiders and juveniles only) were basically the same, while when only adults were considered, no significant variables were detected.

Distribution of spiders in underground environments did not differ by age class: all individuals showed the same horizontal and vertical distribution (Fig. 3). Two or more spiders were rarely observed inside the same cave sector, and these circumstances generally involved juveniles (Table S2). Information relating to the behavior of this species is virtually absent; hence it is possible that individuals may be territorial, at least in some populations. Considering the limited sample size analyzed here (*Lunghi et al.*, unpublished), further studies are needed to better comprehend the behavior of *Meta-bourneti* spiders. Seasonality did not affect *Meta* spider distribution along the horizontal development of the cave, but it strongly affected the vertical

distribution of all individuals; during hot seasons, spiders were found closer to the cave floor. Air circulation in cave environments is characterized by two main air layers, where the lowest has a cooler temperature (Badino, 2010). Therefore, it may be that during hot seasons the temperature of the upper layer becomes too high and spiders move toward the ground floor looking for more suitable microclimatic conditions (Lunghi, Manenti & Ficetola, 2017).

CONCLUSION

This study represents the first analysis performed on island populations of *Meta bourneti*, with the aim of adopting a more complete approach to the study of different ecological aspects of these cave-dwelling spiders. *Meta* spiders were found to be widespread in underground environments of Monte Albo, but with low densities. The species' life cycle, as well as the distribution of individuals inside caves, appears to be strongly dependent by local climatic conditions, showing some divergences from mainland Italian populations. Microclimate was one of the main features affecting both presence and abundance of *M. bourneti* in underground environments; morphological cave features promoting predators avoidance were also important for juvenile spiders. During their underground phase, spiders showed the same tendency to avoid the shallowest part of the caves, areas which likely have unsuitable microclimatic conditions. Surely enough, the vertical movement of spiders suggests a specific behavior of individuals aiming to limit exposure to unsuitable microclimatic conditions. However, further studies on populations from different geographical areas may help in providing a better overview of the ecology of these widespread cave-dwelling species.

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Table 1 (on next page)

The best five AICc models relating the presence of *Meta bourneti*.

In both GLMM and GLM analyses, the presence of the respective ~~studied~~ group (a-f) was used as dependent variable. Independent variables were: Height, Width and wall Heterogeneity^(Het) of sectors, Season of survey, average Temperature^(Temp), Humidity^(Hum) and Illuminance (Lux) recorded inside each sector. We added as further independent variables interactions (*) between season and microclimatic features (temperature, humidity, illuminance). In GLMM analyses we used both sector and cave identity as random factors; in GLM, cave identity was included as ^{an additional} ~~further~~ independent variable. The X indicate the presence of the variable into the respective AICc model; – indicate that the variable was not used in the analyses.

Independent variables included into the model										df	AICc	Δ-AICc	Weight	
Height	Width	Het	Season	Cave	Temp	Hum	Lux	Temp*S	Hum*S	Lux*S				
GLMM														
a)	Meta spiders													
X			X	—		X	X			X	12	453.2	0	0.329
X	X		X	—		X	X			X	13	454.5	1.31	0.171
X			X	—		X	X		X	X	15	454.6	1.39	0.164
X			X	—	X	X	X			X	13	454.9	1.72	0.139
X		X	X	—		X	X			X	13	455.3	2.14	0.113
b)	adults													
				—			X				4	218.7	0	0.210
X				—			X				5	218.8	0.09	0.201
X				—		X	X				6	219.1	0.36	0.176
				—		X	X				5	219.2	0.47	0.166
		X		—			X				5	219.7	0.97	0.130
c)	juveniles													
X			X	—	X	X	X			X	13	343.7	0	0.235
X			X	—		X	X			X	12	344.3	0.54	0.180
X			X	—		X	X		X	X	15	344.3	0.55	0.179
X			X	—		X			X		11	344.6	0.89	0.151
X			X	—	X	X			X		12	344.7	0.97	0.145
GLM														
d)	Meta spiders													
X			X	X		X					11	149.4	0	0.373
X	X		X	X		X					12	151	1.62	0.166
X			X	X		X	X				12	151.4	1.99	0.138
X		X	X	X		X					12	151.4	2.05	0.134
X			X	X	X	X					12	151.6	2.18	0.126
e)	Adults													
X			X	X			X				11	129.1	0	0.324
X	X		X	X			X				12	130.4	1.22	0.176
			X	X			X				10	131	1.83	0.129
X		X	X	X			X				12	131	1.88	0.127
X			X	X		X	X				12	131	1.89	0.126
f)	Juveniles													
X			X	X		X					11	105	0	0.301
X			X	X	X	X					12	105.5	0.48	0.236
X		X	X	X		X					12	106.6	1.59	0.136
X			X	X		X	X				12	107	1.92	0.115
X	X		X	X		X					12	107	1.96	0.113

Table 2 (on next page)

Parameters related to the presence of *Meta bourneti* spiders.

For each group (a-c) are shown significance of variables included in the relative best AICc model of the respective analysis. Shaded variables are those included in the best model of both GLMM and GLM analysis.

Factor	GLMM			GLM		
	<i>B</i>	χ^2	<i>P</i>	<i>B</i>	χ^2	<i>P</i>
a) <i>Meta bournetii</i>						
Season		10.25	0.016		4.97	0.174
Cave					10.86	0.054
Height	0.28	15.9	<0.001	0.27	17.35	<0.001
Humidity	13.82	14.24	<0.001	11.07	9.57	0.002
Illuminance	-1.65	0.03	0.86			
Illuminance*Season		14.96	0.002			
b) adults						
Season					1.17	0.761
Cave					7.65	0.177
Height				0.2	4.22	0.04
Illuminance	-2.63	7.75	0.005	-2.84	11.65	<0.001
c) juveniles						
Season		18.29	<0.001		9.05	0.029
Cave					12.52	0.028
Height	0.29	14.52	<0.001	0.28	13.99	<0.001
Temperature	0.33	2.61	0.106			
Humidity	16.95	16.09	<0.001	12.87	8.24	0.004
Illuminance	-1.42	0.07	0.794			
Illuminance*Season		10.58	0.014			

Figure 1

Observation of *Meta bourneti* spiders performed in Monte Albo's caves.

~~The graph reports the number of observation performed from Autumn 2015 to Summer 2016.~~

Seasonal number of observed spiders is given separating adults (blue) and juveniles

(orange)- From Autumn 2015 to Summer 2016.

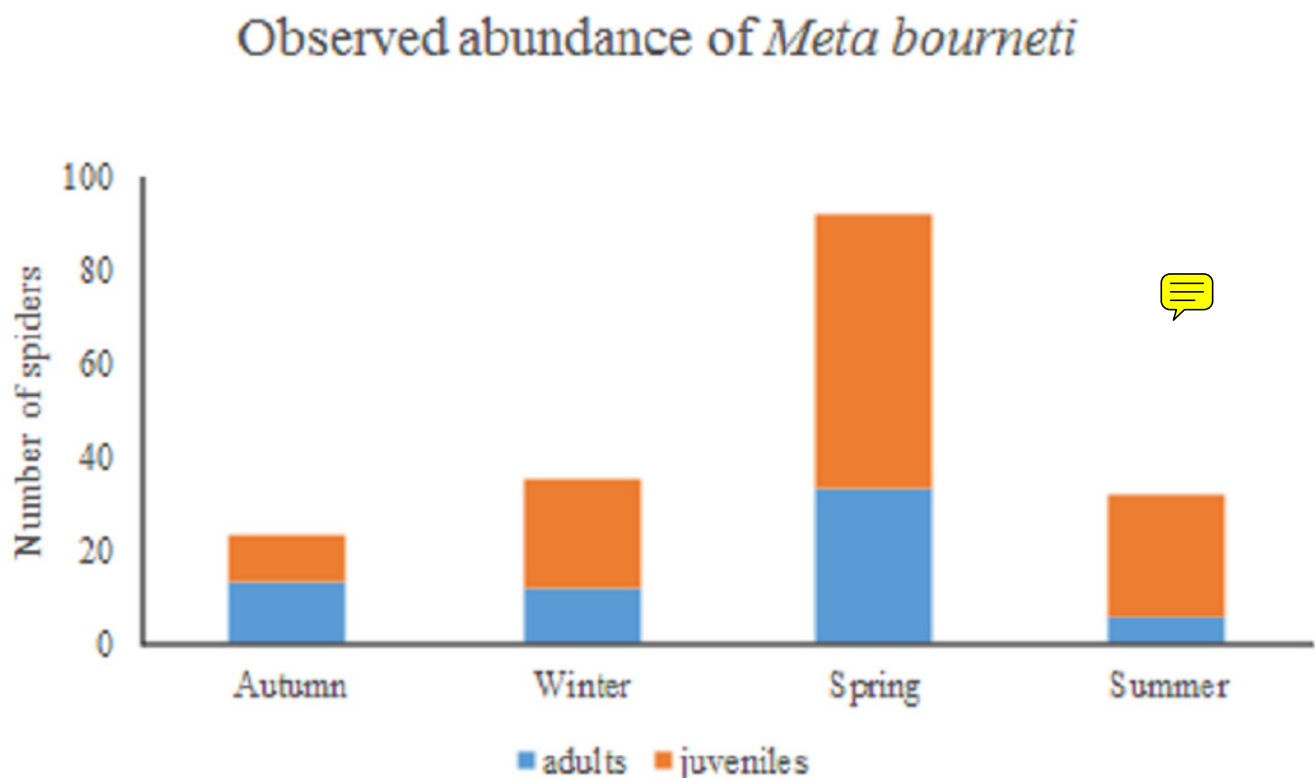


Figure 2

Spiderlings of *Meta bourneti* abandoning their cocoon.

A) A cocoon found in autumn, already laying on the cave floor; inside is possible to observe both spiderlings and opened eggs. B) The same cocoon during winter; spiderlings were abandoning the cocoon.



Figure 3

Boxplots indicating the distribution of *Meta* spiders inside caves.

Differences between adults and juveniles in the use of A) horizontal (i.e., distance from cave entrance) and B) vertical (i.e., height above cave floor) development of the cave environment. Diagonal bar inside the box represents the median.

