Subterranean Biology 27:31–51 (2018) doi: 10.3897/subtbiol.27.28909 http://subtbiol.pensoft.net





Day-night and seasonal variations of a subterranean invertebrate community in the twilight zone

Stefano Mammola¹, Marco Isaia¹

I Department of Life Sciences and Systems Biology, University of Turin, Via Accademia Albertina 13, 10123, Turin, Italy

Corresponding author: Stefano Mammola (stefanomammola@gmail.com)

Academic editor: O. Moldovan | Received 6 August 2018 | Accepted 10 September 2018 | Published 19 September 2018

http://zoobank.org/C968063A-65C8-4BEB-A90B-E96E7EC006FB

Citation: Mammola S, Isaia M (2018) Day–night and seasonal variations of a subterranean invertebrate community in the twilight zone. Subterranean Biology 27: 31–51. https://doi.org/10.3897/subtbiol.27.28909

Abstract

Being characterized by the absence of light and a reduced environmental cyclicity, the subterranean domain is generally regarded as temporally stable. Yet, in the proximity of cave entrances (twilight zones), patterns of sunlight and darkness can be detected within the 24-hour day—night cycle. In parallel, changes in the abiotic and biotic conditions are expected; however, these patterns have been rarely explored in animal communities dwelling in the twilight zone. We performed a biological investigation in a small abandoned mine in the Western Alps, monitoring it once per season, both during the day and at night. At each survey, we collected data on the spatial distribution of the resident species, their activity patterns, and the main microclimatic parameters. We observed significant daily variations in the environmental conditions during winter and spring, namely higher temperature, relative humidity and availability of trophic resources at night. In conjunction with these disparate nocturnal conditions, the abundance of troglophile species was also higher, as well as the activity patterns of one of the most frequent species inhabiting the entrance area — the orb-weaver spider *Meta menardi*. We further documented temporal changes in the composition of the parietal community, due to species using the mine as a diurnal, nocturnal or overwintering shelter. Overall, our results suggest that the communities of the twilight zone are not temporally stable and we highlight the importance of taking into account not only their seasonal, but also their daily variations.

Keywords

arthropods, seasonality, disphotic zone, spatial dynamics, day night, cave cricket, cave spiders, activity patterns, mine

Introduction

Light availability plays a crucial ecological role for organisms on the earth surface (e.g., Panda et al. 2002, Bradshaw and Holzapfel 2010, Gaston et al. 2013, Tierney et al. 2017). Alternatively, the absence of light, at least from a biological point of view (see Badino 2000), represents the most crucial factor conditioning the ecology of subsurface habitats and the evolution of subterranean inhabitants (Pipan and Culver 2012, Battiston and Marzotto 2015, Culver and Pipan 2015, Konec et al. 2015, Fišer et al. 2016, Tierney et al. 2017). As a direct consequence of the lack of solar irradiation, primary phototrophic producers are absent in the deep parts of the caves, with the exception of plant roots growing from the soil into sub-superficial subterranean habitats (Gibert and Deharveng 2002). The permanent darkness below the ground also determines the absence of daily dark-light cycles and a reduced influence of the environmental cyclicity (Poulson and White 1969, Howarth 1980, 1983), two factors which seemingly triggered the reduction in the circadian components of activity of numerous cave-limited species over their evolutionary history (Trajano and Menna-Barreto 1995, Hervant et al. 2000, Trajano et al. 2005, Moran et al. 2014, Espinasa et al. 2016). As such, the subterranean domain is generally regarded as temporally stable.

However, evidences have accumulated testifying that subterranean habitats are not entirely aseasonal (Poulson and Culver 1969, Tobin et al. 2013), and that their inhabitants are not entirely arhythmic (Friedrich 2013, Abhilash et al. 2017). Most subterranean habitats are indeed characterized by a constant flux of invaders and migrants (Chapman 1993, Culver and Sket 2002, Culver and Pipan 2009, Romero 2009, and there can be a temporal variability in the presence and distribution of vertebrates (e.g., Reichard et al. 2009, Krofel 2010, Biswas 2014, Baker 2015, Lunghi et al. 2015, 2017, Ineich and Bourgoin 2016, Balogová et al. 2017, Lunghi et al. 2018) and invertebrates (e.g., Crouau-Roy et al. 1992, Carchini et al. 1994, Gnaspini et al. 2003, Novak et al. 2004, Papi and Pipan 2011, Tobin et al. 2013, Mammola and Isaia 2014, Bento et al. 2016, Mammola et al. 2015, 2016a, 2017, Bichuette et al. 2017, Lunghi et al. 2017, Plăiașu et al. 2017) found in caves. For instance, invertebrates may move in search of food in larger cave chambers, and move back to the more stable fissures connected with caves in response to any physiological stress (Juberthie 1969, Chapman 1985). Besides, numerous species commonly exploit subterranean habitats in certain phases of their life cycle, e.g. as seasonal shelters (Novak et al. 2010a, Chelini et al. 2011, Lipovšek et al. 2016, Balogová et al. 2017) or as breeding sites (Diesel et al. 1995, Glover et al. 2008, Barzaghi et al. 2017). As a direct consequence, there are seasonal turnovers in the species composition in a typical subterranean community.

In several subterranean habitats, there are transitionary photic zones such as cave entrances, where changes in light availability can be detected during the day (but see Mejía-Ortíz et al. 2018 for a peculiar example of a cave with almost no dark zone). In these habitats, theoretically, variations in the composition of the animal community and the abundance and activity of individual species should occur within the 24-hour day–night cycle. Daily spatial and temporal dynamics have been seldom documented

in cave-dwelling species (e.g. Weber et al. 1995, Taylor et al. 2005, Polseela et al. 2011, Havird et al. 2013). In this respect, certain species of bats represent classical examples, resting inside the cave during the day and leaving it at night to feed (e.g., Rodríguez-Durán and Lewis 1987, Agosta et al. 2005, Berková and Zukal 2006). Similarly, other organisms such as nocturnal species taking shelter in caves during the day display analogous activity patterns. For example, Ineich and Bourgoin (2016) recently reported about the peculiar case of a nocturnal Madagascar gekkonid, *Paroedura tanjaka* Nussbaum & Raxworthy, 2000 (Squamata: Gekkonidae), which exploit the cave habitat during daytime to find food and as a reproductive site, meanwhile avoiding most epigean predators. However, to the best of our knowledge, such daily dynamics have never been explored at the level of the entire invertebrate community inhabiting the twilight zone.

We performed a pilot study in a small subterranean site in the Western Italian Alps, in order to unravel the existence of diurnal–nocturnal and seasonal dynamics in the abundance and patterns of activity of resident species. We hypothesized that i) there are variations in the environmental conditions at the twilight zone (e.g. microclimate, trophic resources) both seasonally and within a day–night cycle. We further hypothesized that ii) in parallel to these daily and seasonal environmental variations, there are changes in the species composition and in the abundance of the resident species. Finally, we hypothesized that iii) there are different activity patterns in the resident species during day- and night-time.

Methods

Study site

The study was performed in an abandoned mine near the hamlet of Seinera, municipality of Bruzolo, Susa Valley, Cottian Alps, Italy [entrance at 1007 m asl; 7.201E, 45.136N (WGS84 reference system)]. We chose an artificial site rather than a natural cave due to its linear shape and low ceiling, allowing us to measure the environmental parameters with high accuracy and to access resident animals more easily (Mammola 2018). The Seinera mine represents a typical subterranean habitat at the epigean/hypogean ecotone (*sensu* Prous et al. 2004). It has a horizontal length of only 22 m and the passage cross-section of about 2.5 × 2 m (Figure 1). The mine is in a mixed deciduous forest of chestnuts, oaks, maples and hornbeams, and opens in micaschist rocks used for talc extraction during the first half of the XX century.

Sampling design

We used a sampling-square methodology to monitor the study site, as it has been shown to be an effective, non-invasive means for investigating the annual dynamics of

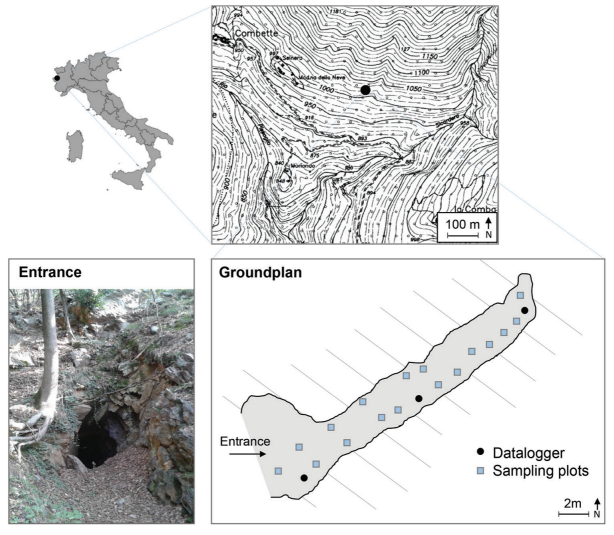


Figure 1. Map of the study area and groundplan of the Seinera mine, with indication of sampling plots and dataloggers.

invertebrates in both artificial and natural subterranean environments (Bourne 1976, Mammola and Isaia 2014, 2016).

Twenty-two sampling plots of 1×1 m were positioned from the entrance up to the end of the mine (Figure 1). We randomly distributed the sampling plots among the mine walls and roof (i.e. parietal habitats; Moseley 2009a). The mine floor was not investigated, because it was impossible to obtain a reliable visual census of the organism due to the presence of leaf litter and debris, and because the floor is not an elective microhabitat for the species considered in our analysis (Mammola and Isaia 2014, Mammola et al. 2016a; see section 'activity pattern').

We visited the study area once per season, between November 2016 and June 2017. In each season, the day of the sampling session was arbitrarily chosen to correspond to the day of new Moon closest to the solstice (summer, winter) or the equinox (autumn, spring). During each sampling session, we conducted one visit around 12:00 am, and we repeated the monitoring approximately four hours after sunset.

During each visit, we censused individuals of each invertebrate species occurring at each sampling plot. Species were identified in the field up to the lowest recognized

taxonomic rank, given the reliability of their in-field identification and our taxonomical expertise. We grouped each species into one of the most common ecological categories (troglobiont, troglophile, trogloxene, accidental organisms) used in subterranean biology (Schiner-Racovitza's categories; Sket 2008, Trajano and Carvalho 2017). Each taxon was attributed to one of these categories on the basis of the ecological information available in literature on its natural history (Lana 2001, Smithers 2005a, Isaia et al. 2011, Allegrucci et al. 2014, Mammola and Isaia 2014, Mammola et al. 2016a, 2016b, 2017). We used such information to calculate the abundance of troglophiles and trogloxenes for each sampling plot – no troglobionts were present in the study site.

We characterized the annual climatic conditions in correspondence of the entrance, 10 m inside and 20 m inside by three EL-USB-2 dataloggers (Lascar Electronics; accuracy of ± 0.5 °C for temperature and ±2.0% for relative humidity), placed at 1 meter height on the wall and programmed to record temperature and relative humidity every 12 h for the whole sampling period – one measurement at 12:00 am and one at 12:00 pm. During each survey, at the centre of each sampling plot we also measured: i) illuminance (lux; lx) using a photometric probe LP 471 Phot (Delta OHM S.r.l.; accuracy of 0.2%) pointed toward the entrance; ii) air flow velocity (WS; m/s) by a Testo 425 telescopic flow velocity/temperature probe (Testo SE and Co, KGaA; accuracy of ±0.03 m/s); and iii) temperature (T; °C) and relative humidity (RH; %) by EL-USB-2 dataloggers (Lascar Electronics; accuracy of ±0.5 °C for temperature and ±2.0 % for relative humidity).

Activity patterns

In order to obtain an estimation of the activity patterns of the three troglophile species inhabiting the mine, during each survey we recorded the diurnal and/or nocturnal movements of the spiders *Meta menardi* (Latreille, 1804) (Araneae: Tetragnathidae) and *Pimoa graphitica* Mammola, Hormiga & Isaia, 2016 (Araneae: Pimoidae), and the cave-dwelling cricket *Dolichopoda azami* Saulcy, 1893 (Orthoptera: Rhaphidophoridae). These species were chosen for this analysis owing to i) their high identification reliability in the field (Mammola and Isaia 2014, Mammola et al. 2016a); ii) their high abundance within the study site (pers. obs.); and iii) their large body size – adults of centimetric length, making a visual monitoring in the field possible.

During each diurnal and nocturnal sampling session, we monitored the activity of each individual of the three species occurring within the sampling plots. The activity was expressed as the number of seconds while the animal was moving, using a stoppable chronometer during observation sessions of one minute. We considered any movement of the animals, with or without spatial displacements. During each session, we set the LED light of our speleological helmet to the red spectrum, in order to minimize disturbance to the animal – in accordance with the general chromatic visual spectrum of most arthropods (Briscoe and Chittka 2001). For each plot and each species, we calculated the total species activity, as the sum of the individual activities divided by the number of individuals within the plot.

Statistical analysis

All statistical analyses were performed in R (R development team 2017). Differences in the environmental conditions at the twilight zone between day and night and across seasons were evaluated graphically (*Graphics* and *Lattice* R packages; R development team 2017, Sarkar 2008) and by means of standard statistical metrics. Wherever appropriate, we tested statistical differences by means of factorial linear regressions (ANOVA) or generalized linear regressions (generalized linear models; GLMs).

To analyse the day–night differences in the abundance, and seasonality of the abundance of trogloxenes and troglophiles, we used a mixed-design analysis of variance with Poisson distributed data (generalized linear mixed models; GLMMs). For the activity pattern of the three species, we used linear mixed models (LMM). GLMM and LMM were fitted with the R packages *lme4* (Bates et al. 2014) and *nlme* (Pinheiro et al. 2014), respectively. Regression-type analyses were conducted following the general protocol of Zuur and Ieno (2016).

Counts of trogloxenes, counts of troglophiles and total activity values for each plot represented the dependent variables. Environmental features (distance from the entrance, temperature, humidity, airflow and illuminance) and their relative interaction with the sampling period, either diurnal (day–night) or seasonal (winter, spring, autumn, summer), represented the independent covariates (i.e., explanatory variables). In order to capture potential non-linear trends in the response of the dependent variables, we allowed up to second order polynomial for the continuous independent variables, when appropriate. The mixed procedure accounted for multiple observations from the same sampling plot, by specifying the sampling plot within the seasonal sampling session as a random-intercept nested structure.

Prior to fitting the models, we explored the datasets following the standard protocol for data exploration proposed by Zuur et al. (2010). Indeed, the inclusion of outliers and highly correlated predictors in a regression analysis may lead to incorrect results (type I and II statistical errors). Thus, we used Cleveland dotplots to assess the presence of outliers in dependent and independent variables. We investigated multicollinearity among continuous covariates via Pearson correlation tests (r) and variance inflation factors values (VIFs), setting the threshold for collinearity at r > |0.7| and VIF> 3.0. The collinearity between continuous and categorical variables was graphically evaluated with boxplots.

Once we fitted the initial models, including all covariates and interactions of interest, we applied model selection via backward elimination (Johnson and Omland 2004). Models were simplified by sequentially deleting covariates and/or interactions according to AICc values (Hurvich and Tsai 1989). The process was repeated until all remaining variables were statistically significant. In turn, validation plots were constructed using model residuals, and Poisson GLMMs were checked for over-dispersion.

Results

Environmental conditions

During the day, illuminance ranged from 900 lx in the vicinity of the entrance, to 0 lx at the end of the mineshaft (mean \pm sd= 22.68 \pm 103.56). Illuminance was always null at night. Airflows ranged from 0 to 0.61 m/s (mean \pm sd= 0.08 \pm 0.48). There were no daily or seasonal variations in the intensity of the airflows (ANOVA; R²= 0.08, p= 0.26 n.s.).

The mean annual temperature at the entrance, at 10 m and at 20 m inside was comparable, however, values at the entrance showed higher seasonal variability (mean±s.d. 0 m= 7.19±5.92 °C; 10 m= 8.08±2.76 °C; 20 m= 8.09±1.24 °C). Overall, the amplitude of changes and min–max ranges were progressively attenuated with increasing distance from the entrance (Figure 2). There were no significant thermal variations between day and night inside the mine (ANOVA; R^2 = 0.12, p=0.17 n.s.). At the entrance (0 m), significant variations between day and night were observed in winter and spring (Figure 2). In particular, night temperature was significantly higher in winter (LM; Winter*Night: estimated β ±SE= 0.595±0.318, p< 0.001), and significantly lower in spring (LM; Spring*Night: estimated β ±SE= -0.708±0.318, p= 0.02), with respect to diurnal temperature.

Relative humidity ranged daily and seasonally between 70% and 100% (mean±sd= 88.85 ± 5.65). Difference between day and night were more pronounced in winter and spring (Figure 3), with significantly higher humidity values at night (beta-GLM; Winter*Night, estimated $\beta\pm SE=0.550\pm0.133$, p< 0.001; Spring*Night, estimated $\beta\pm SE=0.384\pm0.141$, p= 0.006).

Animal community and abundance models

The mineshaft hosted a diversified subterranean biocoenosis, including rich populations of arthropods typical of the twilight zone of Western Alpine caves (Table 1). Over the year, we found 27 taxa within the study area. The most abundant predators [Meta menardi, Metellina merianae (Scopoli, 1763), Pimoa graphitica, Dolichopoda azami, Tegenaria cf. silvestris (Araneae: Agelenidae)] were recorded during all surveys (Table 1). Some taxa were exclusively recorded either during the day (e.g. geometrid moths) or at night (e.g. centipedes and millipedes). There were also seasonal variations in the animal community, with species found in either one [Eupolybothrus sp. (Lithobiomorpha: Lithobiidae), Callipus cf. foetidissimus (Nematophora: Callipodidae)] or more seasons [e.g. Troglohyphantes lucifer Isaia, Mammola & Pantini, 2017 (Araneae: Linyphiidae)].

Regression models were performed to identify the most important factors driving the abundance of both trogloxenes and troglophiles. Data exploration revealed that the variable temperature was collinear with the categorical variable season, and therefore

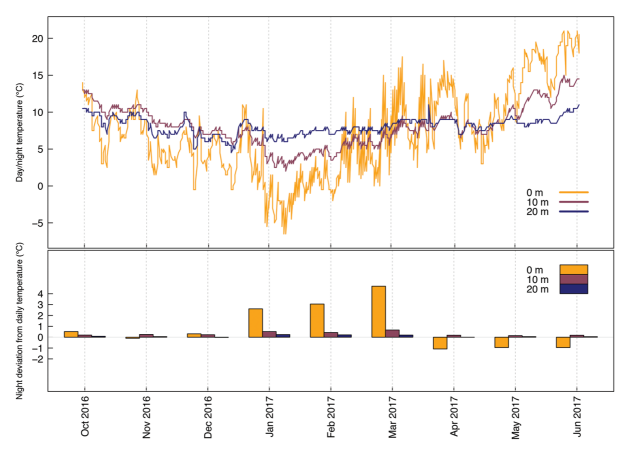


Figure 2. Temperature variation in the study area. Data refer to record of temperature and relative humidity measured every 12 h (one measurement at midday and one at midnight). Top panel: annual trends of temperatures measured at the entrance (0 m; orange line) and inside the mine (10 and 20 m; purple and blue lines, respectively). Bottom panel: mean of monthly positive and negative temperature deviations at night, with respect to the daily temperature recorded during the same period.

it was not further considered. The variable light intensity and relative humidity were collinear with the categorical variable day—night, given that illuminance was always null at night and that the relative humidity higher at night (see Figure 3). Thus, these variables were not introduced in the regression analyses.

Best AICc models and model estimated parameters are reported in Table 2. With respect to the abundance of trogloxenes, there was a significant interaction between the day–night cycle and the seasonality (Figure 4). Overall, the abundance of trogloxenes within the mineshaft was significantly higher at night, in summer and autumn. In autumn, there was the highest discrepancy between day and night, with a higher abundance at night. With respect to autumn, in the other seasons the day–night differences were significantly lower (Table 2).

Higher abundance of troglophiles were observed at night across all seasons (Table 2). Their abundance also varied seasonally with the distance from the entrance (Figure 5). In winter and spring, the highest abundance of trogophiles was predicted in the inner section of the mine, whereas during summertime closer to the entrance. In autumn, the predicted abundance of troglophiles peaked at intermediate distances (Figure 5).

Table 1. Checklist of the taxa recorded within the study site. For each taxon we report the ecological classification (Adapt; TF= Troglophile, TX= Trogloxene, AC= Accidental), the diurnal/nocturnal occurrence (D/N; D= day, N= night) and the seasonal abundance.

Species (Family)	Adapt	D/N	Autumn	Winter	Spring	Summer
ARACHNIDA: ACARINA			l.			l .
Gen. sp.	AC	D	_	_	_	1
ARACHNIDA: ARANEAE	1		ı			I.
Tegenaria cf. silvestris (Agelenidae)	TF	D/N	12	22	27	6
Amaurobius sp. (Amaurobiidae)	TX	N	_	_	2	5
Araneus diadematus Clerck, 1757 (Araneidae)	AC	D/N	7	2	4	3
Drassodes sp. (Gnaphosidae)	AC	N	_	_	_	1
Troglohyphantes lucifer Isaia et al., 2017 (Linyphiidae)	TF	D/N	_	8	3	5
Labulla thoracica (Wider, 1834) (Linyphiidae)	AC	N	_	3	6	_
Pimoa graphitica Mammola et al., 2016 (Pimoidae)	TF	D/N	39	45	22	10
Meta menardi (Latreille, 1804) (Tetragnathidae)	TF	D/N	90	100	135	74
Metellina merianae (Scopoli, 1763) (Tetragnathidae)	TF	D/N	38	32	42	60
Episinus sp. (Theridiidae)	AC	N	1	_	_	_
ARACHNIDA: OPILIONES						
Amilenus sp. (Phalangiidae)	AC	D/N	_	_	_	4
MYRIAPODA: CHILOPODA			ı			ı
Eupolybothrus sp. (Lithobiidae)	TF	N	_	_	1	_
MYRIAPODA: DIPLOPODA						
Callipus sp. (Callipodidae)	TX	N	_	_	3	1
INSECTA: DIPTERA			ı			ı
Gen. sp.	AC	D/N	569	252	223	377
Musca cf. domestica (Muscidae)	AC	N	2	_	_	_
Limonia cf. nubeculosa (Limoniidae)	TX	D/N	67	44	48	88
INSECTA: HYMENOPTERA	'		ı			ı
Gen. sp. (Formicidae)	AC	D	_	_	1	1
INSECTA: RHYNCHOTA						
Pentatoma cf. rufipes (Pentatomidae)	AC	N	_	_	_	1
INSECTA: LEPIDOPTERA						
Scoliopteryx libatrix (Linnaeus, 1758) (Noctuidae)	TX	D/N	_	6	_	_
Gen. sp. (Geometridae)	AC	D	_	_	_	2
Triphosa cf. dubitata (Geometridae)	TX	D	_	_	_	8
INSECTA: ORTHOPTERA						
Dolichopoda azami Saulcy, 1893 (Rhaphidophoridae)	TF	D/N	28	12	28	18
INSECTA: TYSANURA		1	1	1	1	
Lepisma sp. (Lepismatidae)	AC	D/N	3	2	9	1
MOLLUSCA: GASTROPODA	<u>'</u>	1	1	1	1	ı
Oxychilus sp. (Oxychilidae)	TX	D	_	1	1	_
Limax sp. (Limacidae)	AC	D/N	_	_	_	1
* * * * * * * * * * * * * * * * * * * *		1				

Table 2. For each model are shown significant variables included in the relative best AICc model. For the categorical variable Day–Night, the baseline is 'day'. For the categorical variable Season, the baseline is 'Autumn'. Variables excluded due to model selection or in the initial data exploration (collinearity analysis) are not shown. The symbol asterisk ('*') indicate an interaction.

Dependent variable	Independent variables (baseline)	Estimated β±SE	p-value
	Intercept	0.085±0.257	_
	Day-Night (Night)	1.945± 0.180	< 0.001
Abundance of external elements	Season (Winter)	1.870±0.334	< 0.001
	Season (Spring)	1.890±0.327	< 0.001
	Season (Summer)	0.534±0.352	0.130
	Season (Winter) * Day–Night (Night)	-2.212±0.210	< 0.001
	Season (Spring) * Day–Night (Night)	-2.117±0.212	< 0.001
	Season (Summer) * Day–Night (Night)	-1.250± 0.256	< 0.001
Abundance of troglophiles	Intercept	0.079±0.374	_
	Day-Night (Night)	0.243±0.078	0.002
	Distance	0.480±0.098	< 0.001
	Distance ²	-0.029±0.005	< 0.001
	Season (Winter)	0.335±0.5150	0.515
	Season (Spring)	-0.318±0.553	0.565
	Season (Summer)	1.356±0.496	0.006
	Season (Winter) * Distance	-0.358±0.130	0.005
	Season (Spring) * Distance	-0.213±0.136	0.116
	Season (Summer) * Distance	-0.380±0.136	0.005
	Season (Winter) * Distance ²	0.026±0.007	< 0.001
	Season (Spring) * Distance ²	0.018±0.007	0.001
	Season (Summer) * Distance ²	0.016±0.006	0.003
Activity of Mota manardi	Intercept	0.834±0.364	_
Activity of Meta menardi	Day-Night (Night)	1.279±0.517	0.015

Activity patterns

Overall, we recorded the activity patterns of *Pimoa graphitica* 116-times, of *Meta menardi* 399-times and of *Dolichopoda azami* 86-times. The three species showed contrasting activity patterns. The total activity of *P. graphitica* on the web and of *D. azami* on the walls were extremely reduced both during the day (*P. graphitica*, day mean activity±sd= 0.01±0.11; *D. azami*= 0.06±0.55) and at night (*P. graphitica*, night mean activity±sd= 0.80±6.71; *D. azami*= 0.17±0.58). Although overall activity was slightly higher at night in both species, the variable activity was highly zero-inflated (over 90% of observations were zero), meaning that individuals were mostly inactive. It was thus not possible to construct meaningful regression models with the available data, even using specific statistical techniques designed to deal with zero inflation (zero-inflated regression model did not converged; Zuur et al. 2012).

Conversely, *M. menardi* was in general more active, enabling to fit a robust model for this species. Model selection procedure revealed that the best model supported by observation included only the categorical variable day-night (Table 2). The total activ-

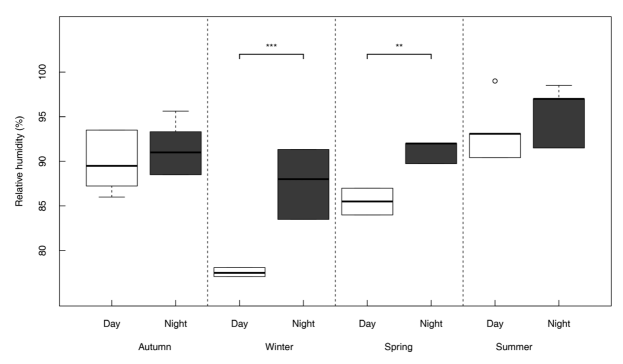


Figure 3. Boxplots showing the difference between relative humidity values during the day (white boxes) and at night (grey boxes) in the four seasons. Significant differences are highlighted by asterisks (Signif. codes: *** p<0.001, ** p<0.01).

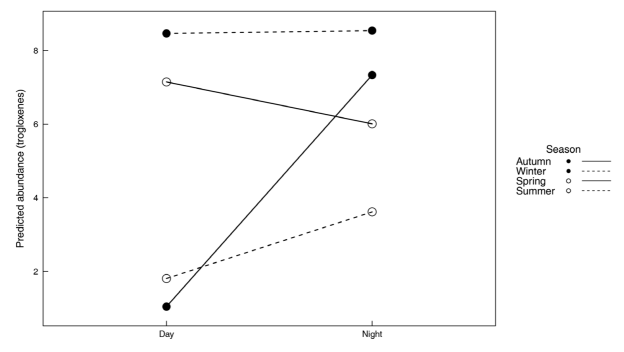


Figure 4. Interaction plot showing the effect of the interaction between seasonality and the day–night cycle on the abundance of trogloxenes.

ity of *M. menardi* was higher at night (night mean activity±sd= 2.56±5.60) than during the day (day mean activity±sd= 0.83±2.41), and this difference was statistically significant (p= 0.02). No significant differences in seasonal activity were detected, therefore the variable season was dropped during model selection.

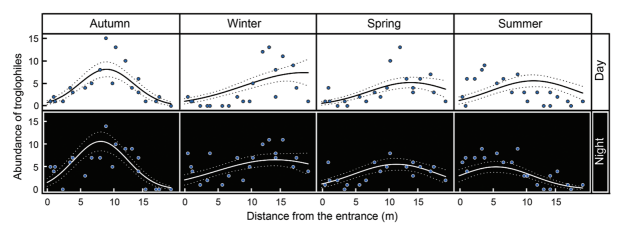


Figure 5. Predicted values (filled lines) and 95% confidence intervals (dotted lines) of the effect of distance from the main entrance in interaction with the sampling season on the abundance of troglophiles derived from the generalized linear mixed model (GLMM). Day and night trends are shown.

Discussion

Whilst the forefront on analysis of the temporal patterns in cave communities has relied on the seasonal timescale (e.g., Tobin et al. 2013, Pellegrini and Ferreira 2016, Lunghi et al. 2017, Mammola et al. 2017), this study aimed at incorporating a day/ night timescale in the analysis of abundance and activity patterns of subterranean invertebrates. We observed significant day—night variations in the abiotic conditions of the twilight zone in term of temperature, humidity and illuminance and, in parallel, variations in the composition of the community and the abundance of certain species. These variations differed across seasons, with the most significant discrepancies between day and night in winter and spring.

It is self-evident that the most easily detectable day-night variation in the environmental conditions of the twilight zone pertain the illuminance. Most species inhabiting the twilight zone should theoretically perceive and respond to variations in light availability within the typical circadian cycle of 24 hours – although some erratic patterns of activity have been documented (Koilraj et al. 2000, Hoenen 2005, Pasquali and Sbordoni 2014). Variations in the composition of the community between day and night were indeed observed. Aside from the obvious differential daily availability of sunlight, at least in the vicinity of the entrance there were also variation in the climatic conditions, especially in respect to higher moisture content at night. A reduced relative humidity in particular, is a well-known limiting factor for the presence of the subterranean fauna (Howarth 1980, 1983, Sharratt et al. 2000). Pronounced sensitivity to saturation deficit was experimentally demonstrated in subterranean beetles (Boyer-Lefèvre 1971), crickets (Yoder et al. 2011) and spiders (Howarth 1980, Hadley et al. 1981, Mammola and Isaia 2017). The fact that subterranean species are preferentially associated with humid microhabitats (Howarth 1980, 1983), indirectly implies that, at night, the twilight zone should represent a more suitable habitat for the subterranean fauna due to the higher levels of relative humidity – at least in winter and spring.

In conjunction with these disparate diurnal–nocturnal and seasonal conditions, we observed variation in the abundance and composition of the animal community. The parietal community was primarily composed by troglophile predators, some trogloxenes plus a variety of accidental species (Table 1). It is worth noting that prior to this study, only two species were recorded in the study site, namely the spiders Pimoa graphitica (Mammola et al. 2016b) and Troglohyphantes lucifer (Isaia et al. 2017). Across the four seasons, the abundance of troglophiles was in general higher at night. There were also variations in the presence of trogloxenes in the study area, with greater nocturnal abundances during summertime and in autumn. It has been demonstrated that predators inhabiting the twilight zone have access to a range of potential prey, which mostly includes accidental organisms that move into the subterranean system from the external environment (Smithers 2005b). The temporal differences in the abundance of trogloxenes and accidental elements in the twilight zone (mostly dipterans), in turn, determines a differential availability of potential prey items for the resident predators. During summertime and autumn, when the availability of these potential prey items was higher at night, troglophiles predators (Dolichopoda azami, Meta menardi, Pimoa graphitica) displaced themselves toward the entrance of the mineshaft, most likely in order to exploit this increased availability of trophic resources. Conversely, during winter, the predicted abundance of troglophiles was higher deeper inside the mine.

We also noticed that some of the taxa used the mine either during the day or at night. For instance, the nocturnal moth [*Triphosa* cf. *dubitata* (Lepidoptera: Geometridae)] likely uses the mine as a diurnal shelter, whereas myriapods were exclusively documented at night. In contrast to moths, the latter case may not reflect a true biological pattern, but can be explained in light of a differential detectability of the species between day and night, i.e. myriapods are mostly active at night and preferably occur in sheltered and hardly accessible microhabitats during the day. Finally, a small part of the community used the mine for overwintering, e.g. the herald moth *Scoliopteryx libatrix* (Linnaeus, 1758) (Lepidoptera: Noctuidae).

Activity patterns

The high nocturnal activity that we observed in *Meta menardi* can be explained considering the parallel higher nocturnal availability of potential prey items. Indeed, a relation between the presence of *Meta* spiders and the availability of prey was previously documented (Mammola and Isaia 2014, Manenti et al. 2015). The other species considered in this study, instead, were mostly inactive. This is not surprising in the case of *Pimoa graphitica*, which is regarded as a sit-and-wait hunter (Mammola et al. 2016a, 2016b). Conversely, our results do not meet our expectation for the cave-dwelling cricket *Dolichopoda azami*. In particular, given that the recorded activity pattern was extremely low, and abundance of individuals was similar during day and night, our data seemingly did not confirm the hypothesis that *Dolichopoda* crickets leaves the cave environment at night to forage outside (see, e.g., Di Russo et al. 1991, 1994, Carchini et al. 1994, Lana 2001), at least in this specific site.

No significant variations in the activity patterns were detected in respect to the other environmental predictors considered in this study. However, given that these results were obtained in uncontrolled environment, the picture obtained of activity patterns remains preliminary. Detailed experiments performed in laboratory conditions would be useful, specifically aimed to establish rhythms of activity during controlled light/dark cycles and to evaluate the circadian clock of troglophile species inhabiting the twilight zone.

Concluding remarks

Although based on a single subterranean community, these results of this study can be used as a jumping-off point to introduce new ideas about our perception of subterranean ecosystems. We acknowledge that it may seem counterintuitive to use caves as model systems where to investigate biological cycles related to light availability. In fact "[...] the subterranean ecological theater is, by definition, dark" (Culver 2016, p. 67). Yet, in the twilight zone, variations in the availability of sunlight can be detected within the regular daily light—dark cycle. Our data evidence that, in parallel, there are daily variations in some microclimatic and trophic conditions, conditioning the animal community resident in this transitional area.

For many years, deep cave habitats have been the central models for studying the ecology and evolution of subterranean life (Culver and Pipan 2015). In recent years, however, the twilight zone is receiving increasingly attention as a model system in ecology and evolution. These transitional habitats represent ecotones characterized by clear environmental gradients (Prous et al. 2004), which can be used as ideal models for to study of transition in species assemblages (Sharipova and Abdullin 2007, Moseley 2009b, Prous et al. 2015, Mammola et al. 2017). Furthermore, non-strictly cave species inhabiting these transitional habitats offer intriguing opportunities for shedding light on the process of adaptation to subterranean environments (Yao et al. 2016) and for the study of classic ecological topics, such as competition dynamics, the niche theory and the predator-prey interactions (Novak et al. 2010b, Luštrik et al. 2011, Mammola and Isaia 2014, Mammola et al. 2016a). In this frame, our data emphasize the fact that the study of cave twilight zone communities should preferably incorporate a temporal perspective, as already suggested by other authors (e.g. Culver and Sket 2002, Bichuette et al. 2017, Lunghi et al. 2017).

Acknowledgments

This work was developed in the frame of the research project 'The dark side of climate change' funded by University of Turin and Compagnia di San Paolo (Grant Award: CSTO162355; PI: Dr. Marco Isaia). We thank Davide Giuliano for leading us the mine of Seinera, and Emanuela Palermo for fieldwork assistance. A special thanks goes to Rebecca Wilson for providing useful comments and proofreading our English, and to Rodrigo Lopes Ferreira and an anonymous referee for useful comments during the review process.

References

- Abhilash L, Shindey R, Sharma VK (2017) To be or not to be rhythmic? A review of studies on organisms inhabiting constant environments. Biological Rhythm Research 48(5): 677–691. https://doi.org/10.1080/09291016.2017.1345426
- Agosta SJ, Morton D, Marsh BD, Kuhn KM (2005) Nightly, seasonal, and yearly patterns of bat activity at night roosts in the central Appalachians. Journal of Mammalogy 86(6): 1210–1219. https://doi.org/10.1644/05-MAMM-A-012R1.1
- Allegrucci G, Rampini M, Di Russo C, Lana E, Cocchi S, Sbordoni V (2014) Phylogeography and systematics of the westernmost Italian *Dolichopoda* species (Orthoptera, Rhaphidophoridae). ZooKeys 437: 1–23. https://doi.org/10.3897/zookeys.437.7917
- Badino G (2000) Is it always dark in caves? International Journal of Speleology 29: 89–126.
- Baker GM (2015) Quantifying wildlife use of cave entrances using remote camera traps. Journal of Cave and Karst Studies 77(3): 200–210. https://doi.org/10.4311/2015ES0101
- Balogová M, Jelić D, Kyselová M, Uhrin M (2017) Subterranean systems provide a suitable overwintering habitat for *Salamandra salamandra*. International Journal of Speleology 46(3): 321–329. https://doi.org/10.5038/1827-806X.46.3.2026
- Barzaghi B, Ficetola GF, Pennati R, Manenti R (2017) Biphasic predators provide biomass subsidies in small freshwater habitats: A case study of spring and cave pools. Freshwater Biology 62(9): 1637–1644. https://doi.org/10.1111/fwb.12975
- Bates D, Maechler M, Bolker B, Walker S (2013) lme4: Linear mixed-e ects models using Eigen and S4. R package version 1.0-5. http://CRAN.R-project.org/package=lme4
- Battiston R, Marzotto A (2015) Evaluating and measuring biodiversity in a subterranean light-gradient. Biodiversity Journal 6(3): 709–718.
- Bento DDM, Ferreira RL, Soucza-Silva M, Bellini BC, Vasconcellos A (2016) Seasonal variations in cave invertebrate communities in the semi-arid Caatinga, Brazil. Journal of Cave and Karst Studies 78: 61–71.
- Berková H, Zukal J (2006) Flight activity of bats at the entrance of a natural cave. Acta Chiropterologica 8(1): 187–195. https://doi.org/10.3161/1733-5329(2006)8[187:FAOBAT]2.0.CO;2
- Bichuette ME, Nascimento AR, von Schimonsky DM, Gallão JE, Resende LPA, Zepon T (2017) Terrestrial fauna of the largest granitic cave from Southern Hemisphere, southeastern Brazil: A neglected habitat. Neotropical Biological Conservation 12(2): 75–90. https://doi.org/10.4013/nbc.2017.122.01
- Biswas J (2014) Occurrence and distribution of cave dwelling frogs of peninsular India. Ambient Science 1: 17–25. https://doi.org/10.21276/ambi.2014.01.2.rv02
- Bourne JD (1976) Notes préliminaires sur la distribution spatiale du *Meta menardi*, *Triphosa dubitata*, *Triphosa sabaudiata*, *Nelima aurantiaca* et *Culex pipiens* au sain d'un écosystéme cavernicole (Grotte de Scierce: Mte. Savoie). International Journal of Speleology 8: 253–267. https://doi.org/10.5038/1827-806X.8.3.2
- Boyer-Lefèvre NH (1971) Influence de la dessiccation sur trois coléoptères de la tribu des Sphodrini vivant dans milieux différents. Bulletin de la Société d'histoire naturelle de Toulouse 107: 595–605.

- Bradshaw WE, Holzapfel CM (2010) Light, time, and the physiology of biotic response to rapid climate change in animals. Annual Review of Physiology 72: 147–166. https://doi.org/10.1146/annurev-physiol-021909-135837
- Briscoe AD, Chittka L (2001) The evolution of color vision in insects. Annual Review of Entomology 46(1): 471–510.
- Carchini G, Rampini M, Sbordoni V (1994) Life cycle and population ecology of the cave cricket *Dolichopoda geniculata* (Costa) from Valmarino cave (Central Italy). International Journal of Speleology 23: 203–218. https://doi.org/10.5038/1827-806X.23.3.6
- Chapman PRJ (1985) Are the cavernicoles found in hawaiian lava tubes just visiting? Proceedings of the University of Bristol Spelaeological Society 17(2): 175–182.
- Chapman PRJ (1993) Caves and Cave Life. Harper Collins, London.
- Chelini MC, Willemart RH, Gnaspini P (2011) Caves as a winter refuge by a Neotropical harvestman (Arachnida, Opiliones). Journal of Insect Behavior 24(5): 393–398. https://doi.org/10.1007/s10905-011-9264-x
- Culver DC (2016) 23rd International Conference on Subterranean Biology. Subterranean Biology 19: 65–85. https://doi.org/10.3897/subtbiol.19.9827
- Culver DC, Pipan T (2009) The Biology of Caves and other Subterranean Habitats. Oxford University Press, Oxford.
- Culver DC, Pipan T (2015) Shifting paradigms of the evolution of cave life. Acta Carsologica 44(3): 415–425.
- Culver DC, Sket B (2002) Biological monitoring in caves. Acta Carsologica 31(1): 55-64.
- Crouau-Roy B, Crouau Y, Source CF (1992) Dynamic and temporal structure of the Troglobitic beetle *Speonomus hydrophilus* (Coleoptera: Bathysciinae). Ecography 15(1): 12–18. https://doi.org/10.1111/j.1600-0587.1992.tb00002.x
- Di Russo C, De Pasquale L, Carchini G, Sbordoni V (1991) Preliminary data on the feeding habit of Dolichopoda (Orth. Rhaphidophoridae) by means of faecal content analysis. Mémoires de Biospéologie 18: 81–84.
- Di Russo C, Carchini G, Sbordoni V (1994) Life-history variation in *Dolichopoda* cave crickets. In: Danks HV (Ed.) Insect Life-cycle Polymorphism. Netherlands: Kluwer Academic Publisher, 205–226. https://doi.org/10.1007/978-94-017-1888-2_10
- Diesel R, Bäurle G, Vogel P (1995) Cave breeding and froglet transport: a novel pattern of anuran brood care in the Jamaican frog, *Eleutherodactylus cundalli*. Copeia 2(3): 354–360. https://doi.org/10.2307/1446898
- Espinasa L, Collins E, Finocchiaro A, Kopp J, Robinson J, Rutkowski J (2016) Incipient regressive evolution of the circadian rhythms of a cave amphipod. Subterranean Biology 20: 1–13. https://doi.org/10.3897/subtbiol.20.10010
- Fišer Ž, Novak L, Luštrik R, Fišer C (2016) Light triggers habitat choice of eyeless subterranean but not of eyed surface amphipods. The Science of Nature 103(1–2): 7.
- Friedrich M (2013) Biological clocks and visual systems in cave-adapted animals at the dawn of speleogenomics. Integrative and Comparative Biology 53: 50–67. https://doi.org/10.1093/icb/ict058
- Gaston KJ, Bennie J, Davies TW, Hopkins J (2013) The ecological impacts of nighttime light pollution: a mechanistic appraisal. Biological reviews 88(4): 912–927. https://doi.org/10.1111/brv.12036

- Gibert J, Deharveng L (2002) Subterranean ecosystems: a truncated functional biodiversity. Bioscience 52: 473–481. https://doi.org/10.1641/0006-3568(2002)052[0473:SEATFB]2.0.CO;2
- Glover AM, Altringham JD (2008) Cave selection and use by swarming bat species. Biological Conservation 141(6): 1493–1504.
- Gnaspini P, Santos FH, Hoenen S (2003) The occurrence of different phase angles between contrasting seasons in the activity patterns of the cave harvestman *Goniosoma spelaeum* (Arachnida, Opiliones). Biological Rhythm Research 34(1): 31–49. https://doi.org/10.1076/brhm.34.1.31.14082
- Hadley NF, Ahearn GA, Howarth FG (1981) Water and metabolic relations of cave-adapted and epigean lycosid spiders in Hawaii. Journal of Arachnology 9: 215–222.
- Havird JC, Weeks JR, Hau S, Santos RS (2013) Invasive fishes in the Hawaiian anchialine ecosystem: investigating potential predator avoidance by endemic organisms. Hydrobiologia 716(1): 189–201. https://doi.org/10.1007/s10750-013-1568-8
- Hervant F, Mathieu J, Durand JP (2000) Metabolism and circadian rhythms of the European blind cave salamander *Proteus anguinus* and a facultative cave dweller, the Pyrenean newt (*Euproctus asper*). Canadian Journal of Zoology 78(8): 427–1432.
- Hoenen S (2005) Circadian patterns in the activity of the Brazilian cave cricket *Strinatia brevipennis* (Ensifera: Phalangopsidae). European Journal of Entomology 102(4): 663–668. https://doi.org/10.14411/eje.2005.094
- Howarth FG (1980) The zoogeography of specialized cave animals: a bioclimatic model. Evolution 34: 394–406. https://doi.org/10.1111/j.1558-5646.1980.tb04827.x
- Howarth FG (1983) Ecology of cave arthropods. Annual Review of Entomology 28: 365–389. https://doi.org/10.1146/annurev.en.28.010183.002053
- Hurvich CM, Tsai CL (1989) Regression and time series model selection in small samples. Biometrika 76: 297–307. https://doi.org/10.1093/biomet/76.2.297
- Ineich I, Bourgoin T (2016) Sex and food in a cave during daytime, a possible way to escape predators for the nocturnal Madagascar gekkonid *Paroedura tanjaka* Nussbaum and Raxworthy, 2000. Herpetology Notes 9: 187–190.
- Isaia M, Paschetta M, Lana E, Pantini P, Schőnhofer AL, Christian E, Badino G (2011) Subterranean arachnids of the Western Italian Alps (Arachnida: Araneae, Opiliones, Palpigradi, Pseudoscorpiones). Museo Regionale di Scienze Naturali, Torino, 325 pp.
- Isaia M, Mammola M, Mazzuca P, Arnedo MA, Pantini P (2017) Advances in the systematics of the spider genus *Troglohyphantes* (Araneae, Linyphiidae). Systematics and Biodiversity 15: 307–326. https://doi.org/10.1080/14772000.2016.1254304
- Juberthie C (1969) Relations entre le climat, le microclimate et les *Aphaenops cerberus* dans la Grotte de Saite Catherine. Annales de Spéléologie 24: 75–104.
- Koilraj AJ, Sharma VK, Marimuthu G, Chandrashekaran MK (2000) Presence of circadian rhythms in the locomotor activity of a cave-dwelling millipede *Glyphiulus cavernicolus sulu* (Cambalidae, Spirostreptida). Chronobiology International 17(6): 757–765. https://doi.org/10.1081/CBI-100102111
- Konec M, Prevorčnik S, Sarbu SM, Verovnik R, Trontelj P (2015) Parallels between two geographically and ecologically disparate cave invasions by the same species, *Asellus aquaticus* (Isopoda, Crustacea). Journal of Evolutionary Biology 28: 864–875. https://doi.org/10.1111/jeb.12610

- Krofel M (2010) Winter observations of Eurasian Lynx (*Lynx lynx*) inspecting karst caves and dolines in the Northern Dinaric Mountains, Slovenia. Terenska Notica: 65–67.
- Lana E (2001) Biospeleologia del Piemonte. Atlante fotografico sistematico. [Biospeleology in Piedmont: Systematic photographic atlas]. 'La grafica nuova', Regione Piemonte, Torino.
- Lipovšek S, Novak T, Janžekovič F, Eiland N, Leitinger G (2016) Malpighian tubule cells in overwintering cave crickets *Troglophilus cavicola* (Kollar, 1833) and *T. neglectus* Krauss, 1879 (Rhaphidophoridae, Ensifera). PloS ONE 11 (7): e0158598
- Lunghi E, Manenti R, Ficetola GF (2015) Seasonal variation in microhabitat of salamanders: environmental variation or shift of habitat selection? PeerJ 3: e1122. https://doi.org/10.7717/peerj.1122
- Lunghi E, Manenti R, Ficetola GF (2017) Cave features, seasonality and subterranean distribution of non-obligate cave dwellers. PeerJ 5: e3169. https://doi.org/10.7717/peerj.3169
- Lunghi E, Bruni G, Ficetola GF, Manenti R (2018) Is the Italian stream frog (*Rana italica* Dubois, 1987) an opportunistic exploiter of cave twilight zone? Subterranean Biology 25: 49–60.
- Luštrik, R, Turjak M, Kralj-Fišer S, Fišer C (2011) Coexistence of surface and cave amphipods in an ecotone environment. Contributions to Zoology 80(2): 133–141.
- Mammola S (2018) Finding answers in the dark: caves as models in ecology fifty years after Poulson and White. Ecography. https://doi.org/10.1111/ecog.03905
- Mammola S, Isaia M (2014) Niche differentiation in *Meta bourneti* and *M. menardi* (Araneae, Tetragnathidae) with notes on the life history. International Journal of Speleology 43(3): 343–353.
- Mammola S, Isaia M (2016) The ecological niche of a specialized subterranean spider. Invertebrate Biology 135(1): 20–30. https://doi.org/10.1111/ivb.12113
- Mammola S, Isaia M (2017) Spiders in caves. Proceedings of the Royal Society of London B: Biological Science 284(1853): 20170193. https://doi.org/10.1098/rspb.2017.0193
- Mammola S, Piano E, Giachino PM, Isaia M (2015) Seasonal dynamics and micro-climatic preference of two Alpine endemic hypogean beetles. International Journal of Speleology 44(3): 239–249. https://doi.org/10.5038/1827-806X.44.3.3
- Mammola S, Piano E, Isaia M (2016a) Step back! Niche dynamics in cave-dwelling predators. Acta Oecologica 75: 35–42.
- Mammola S, Hormiga G, Arnedo MA, Isaia M (2016b) Unexpected diversity in the relictual European spiders of the genus *Pimoa* (Araneae, Pimoidae). Invertebrate Systematics 30(6): 566–587.
- Mammola S, Piano E, Giachino PM, Isaia M (2017) An ecological survey on invertebrate communities at the epigean/hypogean interface. Subterranean Biology 24: 27–52. https://doi.org/10.3897/subtbiol.24.21585
- Manenti R, Lunghi E, Ficetola GF (2015) The distribution of cave twilight-zone spiders depends on microclimatic features and trophic supply. Invertebrate Biology 134(3): 242–251. https://doi.org/10.1111/ivb.12092
- Mejía-Ortíz LM, Pipan T, Culver DC, Sprouse P (2018) The blurred line between photic and aphotic environments: a large Mexican cave with almost no dark zone. International Journal of Speleology 47(1): 69–80. https://doi.org/10.5038/1827-806X.47.1.2155

- Moran D, Softley R, Warrant EJ (2014) Eyeless Mexican cavefish save energy by eliminating the circadian rhythm in metabolism. PLoS ONE. 9(9): e107877. https://doi.org/10.1371/journal.pone.0107877
- Moseley M (2009a) Size matters: Scalar phenomena and a proposal for an ecological definition of 'cave'. Cave and Karst Science 35(3): 89–94.
- Moseley M (2009b) Are all caves ecotones. Cave and Karst Science 36(2): 53–58.
- Novak T, Sambol J, Janžekovic F (2004) Faunal dynamics in the Železna jama cave. Acta Carsologica 33: 249–267.
- Novak T, Thirion C, Janžekovič F (2010a) Hypogean ecophase of three hymenopteran species in Central European caves. Italian Journal of Zoology 77: 469–475.
- Novak T, Tkavc T, Kuntner M, Arnett AE, Delakorda SL, Perc M, Janzekovic F (2010b) Niche partitioning in orbweaving spiders *Meta menardi* and *Metellina merianae* (Tetragnathidae). Acta Oecologica 36(6): 522–529.
- Panda S, Hogenesch JB, Kay SA (2002) Circadian rhythms from flies to human. Nature 417(6886): 329–335. https://doi.org/10.1038/417329a
- Papi F, Pipan T (2011) Ecological studies of an epikarst community in Snežna jama na planini Arto–an ice cave in north central Slovenia. Acta Carsologica 40(3): 505–513. https://doi.org/10.3986/ac.v40i3.62
- Pasquali V, Sbordoni V (2014) High variability in the expression of circadian rhythms in a cave beetle population. Biological Rhythm Research 45(6): 925–939. https://doi.org/10.1080/09291016.2014.934077
- Pellegrini TG, Ferreira RL (2016) Are inner cave communities more stable than entrance communities in Lapa Nova show cave? Subterranean Biology 20: 15–37.
- Pinheiro J, Bates D, DebRoy S, Sarkar D, R Core Team (2014) nlme: Linear and Nonlinear Mixed Effects Models. R package version 3.1-118. http://CRAN.R-project.org/package=nlme
- Pipan T, Culver DC (2012) Convergence and divergence in the subterranean realm: a reassessment. Biol J Linn Soc. 107: 1–14. https://doi.org/10.1111/j.1095-8312.2012.01964.x
- Plăiașu R, Ozgul A, Schmidt BR, Băncilă RI (2017) Estimation of apparent survival probability of the harvestman *Paranemastoma sillii sillii* (Herman, 1871) from two caves. Animal Biology 67(2): 165–176. https://doi.org/10.1163/15707563-00002529
- Polseela R, Vitta A, Nateeworanart S, Apiwathnasorn C (2011) Distribution of cave-dwelling phlebotomine sand flies and their nocturnal and diurnal activity in Phitsanulok Province, Thailand. The Southeast Asian Journal of Tropical Medicine and Public Health 42(6): 1395–404.
- Poulson TL, Culver DC (1969) Diversity in terrestrial cave communities. Ecology 50: 153–158. https://doi.org/10.2307/1934678
- Poulson TL, White WB (1969) The cave environment. Science 165: 971–981. https://doi.org/10.1126/science.165.3897.971
- Prous X, Ferreira RS, Martins RP (2004) Ecotone delimitation: Epigean-hypogean transition in cave ecosystems. Austral Ecology 29: 374–382. https://doi.org/10.1111/j.1442-9993.2004.01373.x

- Prous X, Ferreira RL, Jacobi CM (2015) The entrance as a complex ecotone in a Neotropical cave. International Journal of Speleology 44(2): 177–189. https://doi.org/10.5038/1827-806X.44.2.7
- Reichard JD, Gonzalez LE, Casey CM, Allen LC, Hristov NI, Kunz TH (2009) Evening emergence behavior and seasonal dynamics in large colonies of Brazilian free-tailed bats. Journal of Mammalogy 90(6): 1478–1486. https://doi.org/10.1644/08-MAMM-A-266R1.1
- Rodríguez-Durán A, Lewis AR (1987) Patterns of population size, diet, and activity time for a multispecies assemblage of bats at a cave in Puerto Rico. Caribbean Journal of Science 23(3–4): 352–360.
- R Development Core Team (2017) R: A Language and Environment for Statistical Computing. R Foundation for Statistical Computing.
- Romero A (2009) Cave Biology. Cambridge University Press, New York.
- Sarkar D (2008) Lattice: Multivariate Data Visualization with R. Springer, New York.
- Sharipova MY, Abdullin SR (2007) Algological investigation of cave ecotones. Siberian Journal of Ecology 14(6): 1017–1023.
- Sharratt NJ, Picker MD, Samways MJ (2000) The invertebrate fauna of the sandstone caves of the Cape Peninsula (South Africa): patterns of endemism and conservation priorities. Biodiversity and Conservation 9: 107–143. https://doi.org/10.1023/A:1008968518058
- Sket B (2008) Can we agree on an ecological classification of subterranean animals?. Journal of Natural History 42(21–22): 1549–1563. https://doi.org/10.1080/00222930801995762
- Smithers P (2005a) The early life history and dispersal of the cave spider *Meta menardi* (Latreille, 1804) (Araneae: Tetragnathidae). Bullettin of the British Arachnological Society 13(6): 213–216.
- Smithers P (2005b) The diet of the cave spider *Meta menardi* (Latreille 1804) (Araneae, Tetragnathidae). Journal of Arachnology 33: 243–246.
- Taylor SJ, Krejca JK, Denight ML (2005) Foraging range and habitat use of *Ceuthophilus secretus* (Orthoptera: Rhaphidophoridae), a key trogloxene in central Texas cave communities. American Midland Naturalist Journal 154: 97–114. https://doi.org/10.1674/0003-0031(2005)154[0097:FRAHUO]2.0.CO;2
- Tierney SM, Friedrich M, Humphreys WF, Jones TM, Warrant EJ, Wcislo WT (2017) Consequences of evolutionary transitions in changing photic environments. Austral Entomology 56(1): 23–46. https://doi.org/10.1111/aen.12264
- Tobin BW, Hutchins BT, Schwartz BF (2013) Spatial and temporal changes in invertebrate assemblage structure from the entrance to deep-cave zone of a temperate marble cave. International Journal of Speleology 42(3): 203–214. https://doi.org/10.5038/1827-806X.42.3.4
- Trajano E, Carvalho MR (2017) Towards a biologically meaningful classification of subterranean organisms: a critical analysis of the Schiner-Racovitza system from a historical perspective, difficulties of its application and implications for conservation. Subterranean Biology 22: 1–26. https://doi.org/10.3897/subtbiol.22.9759
- Trajano E, Menna-Barreto L (1995) Locomotor activity pattern of Brazilian cave catfishes under constant darkness (Siluriformes, Pimelodidae). Biological Rhythm Research 26(3): 341–353. https://doi.org/10.1080/09291019509360347

- Trajano E, Duarte L, Menna-Barreto L (2005) Locomotor activity rhythms in cave fishes from Chapada Diamantina, northeastern Brazil (Teleostei: Siluriformes). Biological Rhythm Research 36(3): 229–236. https://doi.org/10.1080/09291010400026272
- Weber F, Casale A, Bohkle C, Rusdea E (1995) Daily temperature cycles as a "Zeitgeber" of the circadian locomotion rhythm of the Carabid Beetles *Sphodropsis ghilianii*. Mémoires de Biospéologie 22: 227–234.
- Yao Z, Dong T, Zheng G, Fu J, Li S (2016) High endemism at cave entrances: a case study of spiders of the genus Uthina. Scientific Reports 6: 35757. https://doi.org/10.1038/srep35757
- Yoder JA, Benoit JB, LaCagnin MJ, Hobbs HH (2011) Increased cave dwelling reduces the ability of cave crickets to resist dehydration. Journal of Comparative Physiology B 181(5): 595–601. https://doi.org/10.1007/s00360-011-0555-5
- Zuur AF, Ieno EN (2016) A protocol for conducting and presenting results of regression type analyses. Methods in Ecology and Evolution 7: 636–645. https://doi.org/10.1111/2041-210X.12577
- Zuur AF, Ieno EN, Elphick CS (2010) A protocol for data exploration to avoid common statistical problems. Methods in Ecology and Evolution 1: 3–14. https://doi.org/10.1111/j.2041-210X.2009.00001.x
- Zuur AF, Saveliev AA, Ieno EN (2012) Zero inflated models and generalized linear mixed models with R. Highland Statistics Limited, Newburgh.
- Zuur AF, Ieno EN, Walker NJ, Savaliev AA, Smith GM (2009) Mixed effect models and extensions in ecology with R. Springer, Berlin.