Diversity and biocide susceptibility of fungal assemblages dwelling in the Art Gallery of Magura Cave, Bulgaria

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Abstract: Magura Cave, north-western Bulgaria, possesses valuable rock-art paintings made with bat guano and dated from the period between the Eneolithic and Bronze Ages. Since 2008, the Art Gallery is closed to the general public in order to protect the paintings from vandalism, microclimatic changes caused by visitors and artificial illumination, and the consequent growth of fungi and phototrophs. Nevertheless, some tourist visits are allowed under the supervision of cave managers. This study provides the first scientific report on cultivable fungal assemblages dwelling different substrata in the Art Gallery. A total of 78 strains, belonging to 37 OTUs (Ascomycota 81%, Zygomycota 14%, Basidiomycota 5%), were isolated in the study. This fungal diversity was clearly dominated by Penicillium (50% of strains) and Aspergillus (13%). The most relevant visible fungal colonies were detected in sediments rich in bat guano, where, besides Penicillium, other guanophilic fungi such as Mucor, Mortierella, Trichosporon and Trichoderma were dominant. Conversely, scarce fungi were detected on rock surface of painted walls. Based on the biocide susceptibility assay, octylisothiazolinone (OIT) and benzalkonium chloride (BAC) were effective inhibiting the in vitro growth of dominant fungal species in Magura Cave, when applied at concentrations ranged from 100 to 1,000 mg/L. These data provide a valuable knowledge about Magura fungi, and exemplify a type of preliminary test that may be conducted before planning any biocide treatment. However, considering the irreversible effects of biocides on the ecological balance in caves, and the low fungal contamination in painted walls of Magura Cave, there is no reason to use conventional biocides in this cave. Further studies, monitoring microbial communities and microclimatic parameters, should be conducted to improve the knowledge on microbial ecology in Magura Cave and possible human impacts, as well as to allow the early detection of potential microbial outbreaks.

Keywords: fungi, cultivable microorganisms, rock-art caves, bat guano, biocides


INTRODUCTION

Magura Cave, located in Bulgaria, contains more than 700 prehistoric paintings depicting a variety of anthropomorphic figures, animals and signs that resemble letters from an unknown alphabet. These multi-layered paintings were created using bat guano, similar to those from Baylovo Cave in Bulgaria (Stoytchev, 2005) and Grotta dei Cervi in Italy (Groth et al., 2001), and dated from the period between the Eneolithic and Bronze Ages (Stoytchev, 1994). In 1984, Magura Cave was placed on the Tentative List for consideration as World Heritage Site by the United Nations Educational, Scientific and Cultural Organization (UNESCO, Tentative list 45). In 2008, sections of the cave containing the rock art (the Art Gallery) were closed to the public in order to protect the paintings from deterioration due to (i) vandalism, (ii) microclimate warming caused by visitors and artificial lighting installed in 2001, and (iii) the consequent growth of phototrophs and fungi on the walls and sediments (Arcá, 2014). In particular, the main concern was the presence of fungal colonies on paintings in the Art Gallery (Stavreva, 2012). Despite
the closure of the Art Gallery, some guided tourist visits are allowed under the supervision of cave managers. Human presence in caves always leads to alternations of the microclimate, the biogeochemical cycles, and the balance of organic matter. These parameters have enormous impact on native microbial populations. Some fast growing heterotrophic microorganisms may be favored by the new conditions, and cause microbial outbreaks. However, the presence and role of microorganisms in caves is a topic that is often ignored in cave management unless any microbial crisis arises (Saiz-Jimenez, 2012). Furthermore, the decision of closing a cave to the public for conservation reasons is always a controversial topic. Sometimes economic interests prevail over conservation leading to deterioration of cultural heritage (Saiz-Jimenez et al., 2011). In rock-art caves, when visits cannot be prevented, cave management should include ways to control organic matter inputs by visitors, and periodic monitoring of microclimate and microorganisms present in all cave compartments. Microbial ecological studies can provide useful information for early detection of dangerous microbial outbreaks (Saiz-Jimenez, 2012, 2013). Knowledge about the most relevant microorganisms, such as their colonization patterns, dispersion mechanisms, and potential adverse effects on human health and rock art (if present), can help cave managers to adopt conservation measures.

A variety of studies have demonstrated that the use of biocides in caves is a dangerous choice to control microbial outbreaks (Boston et al., 2006; Saiz-Jimenez, 2013). Boston et al. (2006) discussed the effect of different cleaning chemicals traditionally used in caves, such as chlorine bleach, soaps and solvents, on microbial communities and mineral formations. Mulec and Kosi (2009) reviewed the pros and cons of physical, chemical, and biological methods to control the phototrophic biofilms associated with artificial illumination of caves, demonstrating that all evaluated control methods have relevant weaknesses. Biocides completely change the original microbial communities, altering the natural ecological balance of the cave, and subsequent recolonizations can provoke further microbial outbreaks. In addition, when biocides are applied without the appropriate previous efficacy tests against the specific cave microbiota, the ecological effects of such treatments are totally unpredictable. Although there is no clear report about biocide treatments in Magura Cave to date, some authors mentioned that a small fraction of the paintings was treated in 1983 by Aneta Slavova, from the National Institute for Cultural Monuments, with an unknown solution which prevented fungal growth on the treated surfaces (Stavreva, 2012; Arcà, 2014).

The presence of bats is a critical challenge for conservation of rock-art caves. Bat colonies significantly affect the microbial diversity in caves due to input of organic matter through deposition of guano and carcasses, and its influence on bacterial and fungal dispersion (Vanderwolf et al., 2013; Kokurewicz et al., 2016; Ogórek et al., 2016). Bat-inhabited caves can be reservoirs of human pathogenic fungi such as *Histoplasma capsulatum*, causative agent of histoplasmosis (Cano & Hajjeh, 2001), which is frequently isolated from warm caves in tropical areas. In addition, the sudden and catastrophic appearance of the White Nose Syndrome (WNS), caused by *Pseudogymnoascus destructans*, in cave-dwelling North American bats in 2006 has stimulated interest in cave fungi in the last few years (Blebert et al., 2009; Lorch et al., 2013). A huge bat colony of more than 2,000 individuals spends the winter in Magura Cave. A total of eight bat species with priority conservation status in Europe have been identified in Magura Cave (Bulgarian Biodiversity Foundation, “See More: FOR the Bats” project).

The bacterial assemblages inhabiting the Art Gallery of Magura Cave were recently studied by Tomova et al. (2013a, b), and Ivanova et al. (2013). However, despite the relevance of fungi in cave environments and their well-known role in rock-art deterioration, no fungal studies have been conducted to date in Magura Cave. Among other reasons, the need of a fungal study in this cave is supported by the abundance of bat guano in the whole cavity, as well as the fact that fungal colonization of paintings was reported as one of the key motivations for closing the Art Gallery to the public in 2008. Considering this lack of knowledge, the goals of this study are (i) to characterize the fungal assemblages that colonize different substrata in the Art Gallery of Magura Cave by cultivation as well as culture-independent analyses, and (ii) to evaluate the susceptibility of the most abundant cultivable fungi to different biocides.

**MATERIALS AND METHODS**

**Sampling**

Magura Cave is located near Rabisha village in the Vidin district at north-western Bulgaria. The cave has an area of 28,600 m², 2.5 km of underground passages, and its entrance is situated 375 m above sea level (43°43′41.16″N, 22°34′54.84″E). This study focused on the Art Gallery (Fig. 1a), placed in a left branch of the cave which can be accessed through a narrow corridor around 200 m from the entrance. The mean air temperature and relative humidity in this gallery is 12°C and ~94%, respectively.

Surveys were carried out 2011-2015, and samples were collected from 15 different points along the Art Gallery (Fig. 1a). A total of 35 samples were analyzed, 24 from the rock surface close or on the paintings (hereafter “painting samples”, Fig. 1b-d), and 11 from cave sediments rich in bat guano (hereafter “sediment samples”, Fig. 1e-g) which frequently showed growth of fungal mycelia. Painting samples were aseptically collected using cotton swabs, and sediment samples were collected in 50 ml Falcon tubes using sterile scalpels. All samples were kept on ice during transport to the laboratory.

**Isolation of fungi**

Swabs of painting samples were transferred to 15 ml Falcon tubes containing 2 ml of sterile saline solution (0.9% NaCl). Similarly, 2 g of sediment samples were
transferred to 15 ml Falcon tubes containing 10 ml of sterile saline solution. Afterwards all samples were homogenized in the orbital shaker at 150 rpm for 30 min. Isolation of fungi was carried out by plating in triplicate 100 µl aliquots of the resulting suspensions, and three decimal serial dilutions from them, on the following culture media – potato dextrose agar (PDA), malt extract agar (MEA),Sabouraud’s glucose agar (SGA) and dichloran rose bengal chloramphenicol agar (DRBC). Culture plates were incubated at 25°C in darkness for one month, checked every two days during the first two weeks, and once per week later. This standard incubation temperature, higher than air temperature in this cave, was chosen in order to (i) save time of analysis, because the majority of fungi grow quicker at 25°C than 14°C, and (ii) compare the results to most of fungal cave studies, with similar culture conditions. Based on colony morphology, the most abundant cultivable fungi, as well as some further representative strains, were isolated in pure cultures on the same culture media. Based on colony morphology, the most abundant cultivable fungi, as well as some further representative strains, were isolated in pure cultures on the same culture media. Afterwards, selected isolates were grown on MEA slants and stored at 4°C until subsequent analyses.

**Molecular identification**

Molecular identification of all fungal strains was performed by PCR and sequencing of the rDNA internal transcribed spacers (ITS) or a fragment of the 18S rRNA gene (18S). The 18S marker was alternatively analyzed for some fungal strains in which the ITS analysis failed. Genomic DNA of each strain was extracted from its biomass grown on MEA. Biomass was collected and transferred to a 2 ml Eppendorf tube containing 500 µl TNE buffer...
(10 mM TrisHCl, 100 mM NaCl, 1 mM EDTA; pH 8) and glass beads of three different diameters (0.5, 2, and 5 mm). The mixture was shaken at 4.5 ms-1 for 1 min in a Fast Prep RiboLyser cell disrupter (Thermo Hybaid GmbH, Ulm, Germany). The DNA was purified through phenol/chloroform extraction and isopropanol precipitation. The extracted DNA was resuspended in 100 µl sterile ultrapure water, and its DNA concentration was quantified using NanoDrop 2000C (Thermo Fisher Scientific, Wilmington, DE, USA), following the manufacturer’s instructions.

The corresponding molecular marker was analyzed by conventional PCR for each strain, the whole ITS region using the primers ITS1 and ITS4 (White et al., 1990) or a fragment of 18S using the fungal primers 0817F (Borman & Hartin, 2000) and 1750-3’ (Gargas & Taylor, 1992). PCR reactions were performed in a BioRad C1000 Thermal Cycler (BioRad, Hercules, CA, USA). Cycling parameters for both markers were 95°C for 5 min, followed by 35 cycles of 95°C for 30 s, 50°C for 30 s, and 72°C for 1 min (ITS) or 2 min (18S), with a final extension at 72°C for 10 min. Reactions were performed in duplicate, and negative controls (containing no DNA) were included in each PCR trial. All PCR products were checked by electrophoresis in 1.5% (w/v) agarose gels stained with GelRed dye (Genaxxon Bioscience GmbH, Ulm, Germany) and visualized under UV light. Positive amplification products were purified by using a QIAquick PCR Purification Kit (Qiagen GmBH, Hilden, Germany) and sequenced in duplicate by Macrogen Europe Company (Amsterdam, The Netherlands) with the same primer sets used for PCR. After edition, final sequences were submitted to the European Nucleotide Archive (ENA, EMBL-EBI) under the accession numbers LT623959 – LT623995. Identification of strains was based on comparison of their rDNA sequences with the GenBank database using the BLAST algorithm from the NCBI.

**Phylogenetic relationships**

Based on previous molecular identification, representative strains of the different fungal taxa isolated from Magura Cave were selected (Table 1). Phylogenetic relationships between such representative strains and the closest related sequences of reference strains, extracted from GenBank, were estimated. The ITS sequences were aligned using MUSCLE web service (EMBL-EBI), and the phylogenetic analyses were conducted using MEGA 5.2 with the Neighbor-Joining method. Gaps were treated as missing data, the Kimura 2 parameter substitution model was used, and bootstrap values were generated using 10,000 replicates. Additionally, the resulting topology was compared with results from other treeing algorithms, including the Maximum-Likelihood and Maximum-Parsimony methods.

**Morphological studies**

Morphological characteristics of representative strains (Table 1) were additionally characterized. For such purpose all fungal strains were cultivated on MEA under standard incubation conditions (25°C in darkness) for seven to ten days. If poor sporulation was achieved, longer incubation time and other nutrient media were used – e.g., PDA, carrot agar (CA) or potato carrot agar (PCA) (Atlas, 2010). In the case of *Penicillium* and *Aspergillus* strains, additional standard nutrient media were needed for the identification process – Czapek yeast autolysate agar (CYA), Blakeslee malt extract agar (BMEA), yeast extract sucrose agar (YES), and creatine sucrose agar (CREA) (Samson & Frisvad, 2004).

The morphological identification was based on macro- and micro-morphological properties (phenophytic and microscopic characters), i.e. the assessment of growth rate, colony morphology, production of exudate and soluble pigment, sporulation type, size and shape of all components of conidiophore or sporophore, type of conidiogenous cells, production of ascocarps, size and shape of ascocarps, asci, and ascospores; according to taxonomic literature and compendia (Domsch et al., 2007; de Hoog et al., 2000).

If necessary, simple physiological tests, such as Ehrlich test of color reactions and estimation of acid and base production on CREA, were also used for *Penicillium* and *Aspergillus* identification. All species were examined using a Zeiss Axio Scope.A1 microscope with Nomarski interphase contrast. Microscopic slides were prepared from cultures showing good sporulation using 60% lactic acid with fuchsin as a marker, containing the PCR products electrophoretic profiles of all fungal strains were also analyzed. DNA extraction protocol for fungal strains was detailed above. After PCR amplification of ITS marker, as previously described, for DGGE analysis a second round of PCR is needed, using the primers ITS1-GC (containing a GC clamp on its 5’ end) and ITS2 (White et al., 1990), and 1 µl of the first PCR product as DNA template. The cycling parameters in the second PCR were 95°C for 5 min, followed by 35 cycles of 95°C for 30 s, 58°C for 1 min, and 72°C for 1 min, with a final extension at 72°C for 10 min.

**Denaturing Gradient Gel Electrophoresis**

Fungal assemblages of environmental samples collected from paintings and sediments were characterized by denaturing gradient gel electrophoresis (DGGE). For comparison, the DGGE profiles of all fungal strains were also analyzed. DNA from environmental samples was extracted using the FastDNA SPIN Kit for Soil (MP Biomedicals, Solon, USA) following the manufacturer’s instructions. DNA extraction protocol for fungal strains was detailed above. After PCR amplification of ITS marker, as previously described, for DGGE analysis a second round of PCR is needed, using the primers ITS1-GC (containing a GC clamp on its 5’ end) and ITS2 (White et al., 1990), and 1 µl of the first PCR product as DNA template. The cycling parameters in the second PCR were 95°C for 5 min, followed by 35 cycles of 95°C for 30 s, 58°C for 1 min, and 72°C for 1 min, with a final extension at 72°C for 10 min.

For DGGE fingerprint of samples, 20 µl of positive PCR products (containing the ITS1 region) were analyzed. Gel electrophoresis was performed as previously described by Muyzer et al. (1993) in 0.5X TAE (20 mM Tris, 10 mM acetate, 0.5 mM Na₂EDTA; pH 7.8), with acrylamide gels (195×160×1 mm), using 10% (v/v) Rotiphorese Gel 30 (30% acrylamide, 0.8% bisacrylamide; Carl Roth GmbH, Karlsruhe, Germany) containing a gradient of denaturants (formamide and urea) of 25-45%. The gels were run in a D-Code System (BioRad) for 3.5 h in TAE 0.5X buffer at 60°C and constant voltage of 200 V. After electrophoresis, gels were stained in GelRed dye and visualized under UV light. A marker, containing the PCR products...
Table 1. Representative fungal strains isolated from Magura Cave.

<table>
<thead>
<tr>
<th>Representative strain*</th>
<th>Accession No.</th>
<th>Number of strains</th>
<th>Source†</th>
<th>Molecular identification‡</th>
<th>Closest related species from GenBank§</th>
<th>Similarity (%)</th>
<th>Morphological identification‡</th>
</tr>
</thead>
<tbody>
<tr>
<td>P5*</td>
<td>LT623959</td>
<td>12</td>
<td>painting / sediment</td>
<td>Penicillium sp.</td>
<td>P. cordobense, P. polonicum, P. lipoidosum, P. thomii</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>P7</td>
<td>LT623995</td>
<td>6</td>
<td>painting / sediment</td>
<td>Eurotiales**</td>
<td>Penicillium malachitum</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>P31.1</td>
<td>LT623960</td>
<td>4</td>
<td>painting</td>
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<td>P. commune, P. oxalicum, P. camemberti</td>
<td>100</td>
<td>Penicillium commune</td>
</tr>
<tr>
<td>S0-3</td>
<td>LT623961</td>
<td>1</td>
<td>sediment</td>
<td>Penicillium sp.</td>
<td>P. lanosum, P. halotolerans</td>
<td>99</td>
<td>Penicillium aurantiogriseum</td>
</tr>
<tr>
<td>S1-4</td>
<td>LT623962</td>
<td>1</td>
<td>sediment</td>
<td>Penicillium sp.</td>
<td>P. janthinellum, P. ochrochloron</td>
<td>99</td>
<td>Penicillium cf. janthinellum</td>
</tr>
<tr>
<td>2-1</td>
<td>LT623963</td>
<td>1</td>
<td>painting</td>
<td>Penicillium sp.</td>
<td>P. pannorum</td>
<td>100</td>
<td></td>
</tr>
<tr>
<td>2-2</td>
<td>LT623964</td>
<td>1</td>
<td>painting</td>
<td>Penicillium sp.</td>
<td>P. biourgeianum, P. brevicaespactum</td>
<td>100</td>
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</tr>
<tr>
<td>2-3</td>
<td>LT623965</td>
<td>1</td>
<td>painting</td>
<td>Penicillium sp.</td>
<td>P. coprophilum, P. concentricum</td>
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<td>Penicillium coprophilum</td>
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<td>LT623966</td>
<td>1</td>
<td>painting</td>
<td>Penicillium sp.</td>
<td>P. echinulatum, P. commune, P. solitum, P. caseincola</td>
<td>100</td>
<td>Penicillium solitum</td>
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<td>SF20-3</td>
<td>LT623967</td>
<td>1</td>
<td>sediment#</td>
<td>Penicillium sp.</td>
<td>P. concentricum</td>
<td>98</td>
<td></td>
</tr>
<tr>
<td>P23*</td>
<td>LT623968</td>
<td>8</td>
<td>painting / sediment</td>
<td>Aspergillus sp.</td>
<td>A. versicolor</td>
<td>100</td>
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<td>P28</td>
<td>LT623969</td>
<td>2</td>
<td>painting / sediment</td>
<td>Aspergillus sp.</td>
<td>A. versicolor</td>
<td>100</td>
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</tr>
<tr>
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<td>LT623970</td>
<td>6</td>
<td>painting / sediment</td>
<td>Simplicillium sp.</td>
<td>S. cylindrosporum, S. minatense</td>
<td>99</td>
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</tr>
<tr>
<td>5-1</td>
<td>LT623971</td>
<td>2</td>
<td>painting</td>
<td>Chaetomiaceae</td>
<td>Chaetomiom arxii</td>
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</tr>
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<td>SF15-4</td>
<td>LT623972</td>
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<td>sediment#</td>
<td>Chaetomiaceae</td>
<td>Chaetomiom piluliferum</td>
<td>99</td>
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<td>S1-8*</td>
<td>LT623973</td>
<td>2</td>
<td>sediment</td>
<td>Trichosporon sp.</td>
<td>T. porosum</td>
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<tr>
<td>SF19-2</td>
<td>LT623974</td>
<td>2</td>
<td>sediment#</td>
<td>Trichosporon sp.</td>
<td>T. jirucieci</td>
<td>98</td>
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<tr>
<td>5-2*</td>
<td>LT623975</td>
<td>3</td>
<td>painting</td>
<td>Gymnoascus sp.</td>
<td>G. reessii</td>
<td>99</td>
<td>Gymnoascus reessii</td>
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<td>S1-3</td>
<td>LT623976</td>
<td>2</td>
<td>sediment</td>
<td>Mortierella sp.</td>
<td>M. parvispora, M. jenkinii</td>
<td>99</td>
<td></td>
</tr>
<tr>
<td>S1-7</td>
<td>LT623977</td>
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<td>sediment</td>
<td>Mortierella sp.</td>
<td>M. verticillata, M. epipodidua</td>
<td>99</td>
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<td>S2-6*</td>
<td>LT623978</td>
<td>1</td>
<td>sediment</td>
<td>Mucor sp.</td>
<td>M. circinelloides</td>
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<td>Mucor circinelloides</td>
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<td>SF15-7</td>
<td>LT623979</td>
<td>1</td>
<td>sediment#</td>
<td>Mucor aligarensis</td>
<td>M. aligarensis</td>
<td>100</td>
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<tr>
<td>SF20-2</td>
<td>LT623980</td>
<td>1</td>
<td>sediment#</td>
<td>Mucor sp.</td>
<td>M. plumbeus</td>
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<tr>
<td>P16</td>
<td>LT623981</td>
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<td>painting</td>
<td>Cladosporium sp.</td>
<td>C. variabile</td>
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<tr>
<td>S1-5*</td>
<td>LT623982</td>
<td>2</td>
<td>sediment</td>
<td>Ascomycota</td>
<td>Humincola grisea, Trichocladium asperum</td>
<td>99</td>
<td>Humincola grisea var. grisea</td>
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<td>S0-9*</td>
<td>LT623983</td>
<td>1</td>
<td>sediment</td>
<td>Trichoderma sp.</td>
<td>T. koningii, T. hispanicum, T. vinide</td>
<td>100</td>
<td></td>
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<td>S1-1</td>
<td>LT623984</td>
<td>1</td>
<td>sediment</td>
<td>Trichoderma sp.</td>
<td>T. harzianum</td>
<td>100</td>
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<tr>
<td>P18*</td>
<td>LT623985</td>
<td>1</td>
<td>painting</td>
<td>Hypocreales</td>
<td>Sarocladium zeae, Sarocladium strictum</td>
<td>94</td>
<td>Acremonium zeae</td>
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<td>LT623986</td>
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<td>Exophiala sp.</td>
<td>E. salmonis</td>
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<td>S0-2*</td>
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<td>sediment</td>
<td>Bionectria sp.</td>
<td>B. rossmaniae</td>
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<td>B. rossmaniace or B. sesquisilili</td>
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<td>S1-6*</td>
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<td>T. delbrueckii</td>
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<td>LT623990</td>
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<td>sediment</td>
<td>Pochonia sp.</td>
<td>P. rubescens, P. suchlasporia</td>
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<td>S2-4*</td>
<td>LT623991</td>
<td>1</td>
<td>sediment</td>
<td>Doratomyces sp.</td>
<td>D. stemonis, D. purpureofuscus, D. castaneus</td>
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<td>sediment#</td>
<td>Debaryomycesaceae</td>
<td>Debaryomyces sp.</td>
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<td>sediment</td>
<td>Bionectriaceae</td>
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<td>Glomastix murorum</td>
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<td>P31.2*</td>
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<td>sediment</td>
<td>Pseudogymnoascus sp.</td>
<td>P. pannorum</td>
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</tbody>
</table>

*Representative strains of taxonomical groups showing identical sequences in the molecular analysis. †Strains included in the biocide susceptibility assay (Table 2). ‡Isolation source in Magura Cave, samples from the rock surface close or on the paintings collected using cotton swabs (painting), or samples from cave sediments rich in bat guano (sediment). §Sediment samples showing mycelial growth visible to the naked eye. ¶Molecular identification based on comparison of their ribosomal ITS sequences, except some strains (**) for which the 18S rDNA gene was used because the ITS primers failed, with GenBank by BLAST algorithm from NCBI. The closest relative species are detailed. ²Some strains could not be morphologically identified because they produced sterile colonies or additional studies in detail are need.
Penicillium, 2 sp. strain P9 and Mortierella 4 HPO, PO; pH 7.4), from 2 week-old MEA (2-7) after inoculation with 100 µl. suspensions was very high, developing a homogeneous min at 30 Hz. The cell concentration of all resulting MM400 mill (Retsch GmbH, Haan, Germany) for 10 beads, and subsequently homogenized in a Retsch mycelia were mixed with 10 ml PBS and 5 mm metal discs were recorded.

For fungal inoculation, cell suspensions were evaluated, benzalkonium chloride (BAC) and 2-octyl-3-isothiazolinone (OIT), which have a broad antimicrobial activity including against fungi. BAC, composed by mixtures of n-alkylidimethylbenzyl ammonium chlorides, belongs to the group of surface-active quaternary ammonium compounds and is active against bacteria, fungi, some viruses and protozoa. BAC biocidal action is due to disruption of intermolecular interactions causing dissociation of cellular membrane lipid bilayers, which compromises cellular permeability controls and induces leakage of cellular contents (Gilbert & Moore, 2005). OIT is an isothiazolinone derivative with broad-spectrum biocidal action used to control the growth of bacteria, fungi and algae in cooling water systems, storage tanks, emulsions and paints (Kramer et al., 2008).

Both compounds are widely used for conservation of many materials including culture heritage assets. In order to evaluate accurate concentrations of biocides, high-purity active ingredients were used for both biocides, BAC at ≥ 95.0% (Sigma-Aldrich Chemie GmbH, Munich, Germany; product number 12060) and OIT at 97.8% (non-commercial product kindly provided by Thor GmbH, Speyer, Germany). According to similar previous studies, three different biocide concentrations (100, 500 and 1,000 mg/L) were evaluated. For each strain the test was conducted on MEA plates in triplicate plus a control plate without biocide. All plates were initially inoculated with 100 µl of the fungal suspension (described below). Afterwards three sterile Whatman paper discs (5 mm in diameter) were placed on the medium and subsequently soaked with 15 µl of the corresponding biocide concentration. Plates were incubated at 25°C in darkness for 10 days and measurements of the inhibition zones around the discs were recorded.

For fungal inoculation, cell suspensions were prepared in sterile phosphate-buffered saline solution (PBS; 137 mM NaCl, 2.7 mM KCl, 10 mM Na₂HPO₄, 1.8 mM KH₂PO₄; pH 7.4), from 2 week-old MEA cultures of selected strains, following two different procedures depending on sporulation rate. For fungi with high sporulation, conidia were collected washing the colony surface with 5 ml PBS using a sterile pipet. For slow growing fungi or those with poor sporulation, mycelia were mixed with 10 ml PBS and 5 mm metal beads, and subsequently homogenized in a Retsch MM400 mill (Retsch GmbH, Haan, Germany) for 10 min at 30 Hz. The cell concentration of all resulting suspensions was very high, developing a homogeneous growth over the whole surface of culture plate in a few days (2-7) after inoculation with 100 µl.

Biocide susceptibility of fungal strains

Biocide susceptibility of major fungi isolated from Magura Cave was tested by using the agar disc-diffusion test. Two biocidal substances were evaluated, benzalkonium chloride (BAC) and 2-octyl-3-isothiazolinone (OIT), which have a broad antimicrobial activity including against fungi. BAC, composed by mixtures of n-alkylidimethylbenzyl ammonium chlorides, belongs to the group of surface-active quaternary ammonium compounds and is active against bacteria, fungi, some viruses and protozoa. BAC biocidal action is due to disruption of intermolecular interactions causing dissociation of cellular membrane lipid bilayers, which compromises cellular permeability controls and induces leakage of cellular contents (Gilbert & Moore, 2005). OIT is an isothiazolinone derivative with broad-spectrum biocidal action used to control the growth of bacteria, fungi and algae in cooling water systems, storage tanks, emulsions and paints (Kramer et al., 2008).

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RESULTS AND DISCUSSION

Fungal assemblages inhabiting Magura Art Gallery

A total of 78 strains, belonging to 37 operational taxonomic units (OTUs; whose members show identical sequences), were isolated during the study (Table 1). According to the phylogenetic study (Fig. 2 and 3), the majority of OTUs belong to the phylum Ascomycota (30 OTUs, 81%), in contrast to fewer Basidiomycota (two Trichosporon species) and Zygomycota (five species of the genera Mucor and Mortierella). Among the ascomycetes, fourteen OTUs were from class Eurotiomycetes, with the major genera Penicillium (10 OTUs, 39 strains) and Aspergillus (2 OTUs, 10 strains), and twelve OTUs were included in class Sordariomycetes, with a more diverse representation of taxa.

All OTU-representative strains were further morphologically characterized to be identified at the species level. However, only twelve strains (32%) have been identified using this approach (Table 1) due to the well-known limitations of such studies. For some studied strains, such as Aspergillus sp. P23, additional morphological, physiological and/ or phylogenetic studies would be needed for reliable identification. Other strains, such as Mortierella sp. S1-3, only produced sterile colonies which prevented their characterization. The phylogenetic trees (Fig. 2 and 3) show the final consensus identification of strains using both molecular and morphological data. Regarding DGGE characterization of Magura strains, in general the OTUs detailed in Table 1 showed distinctive patterns, like those Penicillium strains belonging to the most abundant OTU (Fig. 4a). As demonstrated in previous studies (Martin-Sanchez et al., 2012), the fungal strains in pure culture frequently develop characteristic multiband DGGE profiles, due to PCR artifacts, which complicates the interpretation of DGGE results from diverse environmental samples. In this sense, Neilson et al. (2013) concluded that DGGE provides an excellent tool for comparative community structure analysis, but such method-specific artifacts preclude its use for accurate quantitative diversity analysis.

In general, the fungal diversity found in the Art Gallery was rather low with clear dominance of a few genera. However, as well known in literature, cultivable populations do not represent the true diversity of the ecosystem. Less than 1% of the estimated microbial diversity is thought to be cultivable in laboratory conditions due to the very low growth rate of many environmental microorganisms (Amann et al., 1995). In addition, the selected culture methods, media and incubation conditions, have most likely favored these genera against the rest of microbes. In accordance with many previous cultivation-based studies in other caves (Vanderwolf et al., 2013), the fungal community in Magura Cave was clearly dominated by Penicillium and Aspergillus species (Fig. 3). Most of Penicillium strains (59%) were taxonomically placed in the subgenus Penicillium, section Fasciculata, according to the classification of Houbraken and Samson (2011) and Visagie et al. (2014a), including the species...
Fig. 2. Fungal phylogeny of the ITS region showing the taxonomical placement of representative strains from Magura Cave (in bold) and the closest reference strains, including type strains (T) and isolates from caves (C). The major taxon Eurotiiales was compressed and detailed in Fig. 3. Numbers of strains higher than one are presented in brackets. The unrooted tree was constructed using the Neighbor-Joining method applying the Kimura 2 parameter model. All positions containing gaps and missing data were eliminated. There were a total of 235 positions in the final dataset. The tree was bootstrapped 10,000 times and values above 50% are indicated at nodes. The asterisks indicate branches of the tree that were also recovered using Maximum-Likelihood and Maximum-Parsimony treeing algorithms. Bar, 0.05 substitutions per nucleotide position.

P. commune and P. solitum. Six Eurotiiales strains, with P7 as representative, were closely related to Penicillium malachitum (subgenus Aspergilloides, section Sclerotiora) based on the analysis of their 18S rRNA gene sequences (Table 1). They could not be included in the phylogenetic analysis focused on ITS region. Members of Chaetomiaceae family and genera such as Simplicitium, Trichosporon, Gymnoascus, Mortierella, Mucor, Cladosporium and Trichoderma were repeatedly isolated in the studied samples. Most of genera isolated from Magura Cave were frequently found in previous studies of air samples from indoor environments, and either in sediment or rock surface samples from other caves in Europe (Vanderwolf et al., 2013). This is logical considering the high concentration of airborne fungal spores expected in caves like Magura, populated by bat colonies and accessible to tourists, with the consequent inputs of organic matter.

Ascomycota appears to be the most abundant phylum in any natural environment irrespective of whether culture dependent or independent approaches are used. According to previous studies (Nováková et al., 2009; Docampo et al., 2011; Vanderwolf et al., 2013; Man et al., 2015), the most abundant fungal genera in caves are Penicillium, Aspergillus, Cladosporium,
Fig. 3. Eurotiales phylogeny of the ITS region showing the placement of representative *Penicillium* and *Aspergillus* strains from Magura Cave (in bold) and the closest reference strains, including type strains (T) and isolates from caves (C). *Penicillium* sections according to Houbraken and Samson (2011) are shown. Numbers of strains higher than one are presented in brackets. The tree was constructed using the Neighbor-Joining method applying the Kimura 2 parameter model, and rooted with *Gymnoascus reessii* 5-2 as outgroup. All positions containing gaps and missing data were eliminated. There were a total of 445 positions in the final dataset. The tree was bootstrapped 10,000 times and values above 50% are indicated at nodes. The asterisks indicate branches of the tree that were also recovered using Maximum-Likelihood and Maximum-Parsimony treeing algorithms. Bar, 0.02 substitutions per nucleotide position.

Fig. 4. DGGE profiles of fungal strains and environmental samples from Magura Cave. a) Strains *Penicillium* sp. P5 (= P1.1, P1.2, P4, P12, P13, P14, P15), *Simplicillium* sp. P3 (= P6), *Aspergillus* sp. P9, *Cladosporium* sp. P16 (= P17) and *Acremonium zeae* P18; marker composed by the strains P4, P9 and P16; b) Sediment samples (S) rich in bat guano, wall painting samples (W) collected from rock surface close or on the paintings, and some strains from samples W2 (*Penicillium* sp. 2-1, *Penicillium coprophilum* 2-3 and *Penicillium solitum* 2-7) and W5 (*Chaetomiaceae* 5-1 and *Gymnoascus reessii* 5-2).
Mucor, Fusarium and Trichoderma, which may be due to their ubiquitous distribution in nature. Aspergillus/ 
Penicillium was the most abundant spore type found in the air of Nerja Cave, Spain, which represented 50% 
of the total spores, followed by Cladosporium with 17% (Docampo et al., 2011). Penicillium and Aspergillus 
species grow better in warm and humid environments but can tolerate wide ranges of temperature and 
pH. They are ubiquitous molds that grow on organic matter producing abundant conidia which enhance 
their fast spreading (Visagie et al., 2014b).

Rock surface of painted walls

As described by Man et al. (2015) in Heshang Cave, 
China, differences in abundance and composition of 
fungal assemblages were detected comparing different 
substrata of Magura Cave. First, it is noteworthy to 
mention than we did not detect relevant in situ fungal 
growth on the rock surface of painted walls in the 
different sampling campaigns. Evident fungal mycelia 
were only observed in a few small areas near the 
paintings, which were especially targeted for sampling. 
This fact could explain the low number of colonies 
grown on culture media from the painting samples 
collected using swabs. For instance, cultivable fungi 
were only isolated from three (W2, W5, and W11) of 
twelve painting samples collected in the last sampling 
campaign in 2015. However, the DGGE patterns 
corresponding to nine of these samples showed 
more diverse assemblages (Fig. 4b), remarking the 
prevalence of non-cultivable fungi on this substratum. 
Despite the commented limitations of DGGE (Neilson 
et al., 2013), which hinder the comparison between 
environmental samples and isolated strains, results 
show that structure of fungal assemblages on rock 
surfaces is rather different depending on studied 
location. Samples W1 and W2, collected from 
neaby areas in the entrance of Art Gallery, showed 
similar patterns between them, and different to 
samples collected from other locations in the gallery. 
Moreover, the main members of these assemblages 
most likely correspond to non-cultivable fungi under 
the selected media and incubation conditions. 
Penicillium, Aspergillus, Simplicillium, Gymnoascus 
and Cladosporium were the most representative 
genera cultivated from these samples. The genera 
Aspergillus and Penicillium belong to the group of 
fermentative microorganisms which excrete a variety 
of organic acids that can cause biodeterioration of 
rock substratum (Sterflinger, 2000).

Bat guano-rich sediments

On the other hand, in sediment samples rich in 
bat guano, high abundance of colonies and diversity 
were found by culturing, and subsequently confirmed 
by DGGE (Fig. 4b). Although Penicillium remains the 
major group in this substratum, other genera such as 
Trichosporon, Mortierella, Mucor, Humicola and 
Trichoderma were also abundant there, while none of 
them were detected on rock surfaces. In particular, 
for those sediment samples showing mycelial growth 
visible to the naked eye (Fig. 1g), members of genera 
Mucor, Penicillium and Trichosporon, and families 
Chaetomiaceae and Debaryomycesaceae, were 
identified (Table 1). Most likely, based on morphology, 
such fungal growth was mainly caused by Mucor 
species. Jurado et al. (2010) reported similar mycelial 
growth on sediments of Castañar de Ibor Cave, Spain, 
during a fungal outbreak associated with the species 
Mucor circinelloides and Fusarium solani.

Novákova (2009) reported a comprehensive fungal 
inventory (195 taxa) from Domicia Cave system, 
Slovakia. The highest number of taxa (92) was found in 
bat guano samples, where species of genera Penicillium, Mucor, Doratomyces and Trichoderma 
were prevalent. Furthermore, Mucor species often 
formed visible colonies on bat droppings and surface of 
guano heaps from these caves. Bat guano samples from 
other Slovakian caves were recently studied by 
Ogórek et al. (2016), with Penicillium, Aspergillus, 
Mucor and Rhizopus as dominant genera. Trichosporon 
species are widespread in cave sediments, especially 
in those rich in bat guano, which highlights the 
relevance of this cave substratum as reservoir of 
potentially pathogenic fungi (Saiz-Jimenez, 2012). 
This basidiomycetous yeast-like genus has frequently 
been reported as causal agent of superficial infections, 
as well as opportunistic agent of invasive infections 
(Colombo et al., 2011). Different studies have reported 
novel Trichosporon species isolated from sediments of 
bat-inhabited caves in Japan and Slovakia (Sugita 
et al., 2005; Novákova et al., 2015). Likewise, many 
Mortierella species were usually isolated from bat 
dung samples collected in caves in Japan (Degawa 
& Gams, 2004). Species of the genera Mucor, 
Mortierella, Trichoderma and Pseudogymnoascus were 
cultured from dead bats in Berryton Cave, Canada, 
being the fast growing Zygomycetes, Mucor and 
Mortierella, particularly abundant on freshly dead 
bats (Vanderwolf et al., 2016). The results confirm 
that members of genera found in Magura Cave are 
significant constituents in many karstic caves. 
Their ecological function and possible impact on the 
paintings are not well understood yet.

The main input of organic matter in Magura Cave is 
clearly from the bat colony inhabiting this cavity 
long time ago, with the consequent accumulation 
of bat guano in sediments of the whole Art Gallery, 
and recurrent presence of visible fungal colonies 
covering these sediments. However, according to our 
observations and results, the fungal growth on rock 
surface of painted walls has been quite scarce up to 
date. As expected and demonstrated in this study, 
the current fungal community in Magura Cave is mainly 
composed by guanophilic species. Considering that 
Magura paintings were made using bat guano, this 
cave has most likely been inhabited by bats and their 
associated guanophilic microbiota since the origin of 
paintings thousands of years ago. Therefore, any 
conservation measure addressed to reduce organic 
matter inputs by controlling bat populations, and/or 
cleaning bat guano, does not make sense. Besides, 
such proposals would not be conceivable with current 
bat protection rules. The potential input of organic 
matter coming from tourists may be considered 
insignificant in comparison to bat guano contribution.
However, humans can critically affect environmental conditions in caves, e.g., increasing variables such as temperature and carbon dioxide concentration, and lead to an ecological imbalance that stimulates the growth of certain microorganisms causing microbial outbreaks.

**Fungal susceptibility to biocides**

The susceptibility of Magura fungi to two of the most commonly used biocides, BAC and OIT, was additionally evaluated by agar disc-diffusion test. A total of seventeen strains, representative of the major phylotypes dwelling in the Art Gallery, were studied (Table 2). In general, both biocides were able to inhibit the fungal growth on culture plates when applied at described concentrations (100, 500 and 1,000 mg/L). These concentrations are significantly lower than those specified in manufacturers’ instructions of commonly used commercial products, in general ranged from 1 to 25 g/L. Only Trichoderma sp. strain S0-9 was resistant to all tested BAC doses. The majority of studied strains showed much higher susceptibility to OIT in comparison to BAC (Fig. 5a-c), except for yeast species. Torulaspora sp. S1-9 showed the opposite trend (Fig. 5d-f) and Trichosporum sp. S1-8 showed similar susceptibility to both biocides. The lowest OIT concentration tested (100 mg/L) inhibited all fungi, except Trichoderma sp., causing a significant inhibition zone diameter (>16 mm) in 13 of 17 strains. In this sense, it is noteworthy to mention the high inhibition degree of some fungi, such as Penicillium sp., Exophiala sp., Doratomyces sp. and Pseudogymnoascus sp., to 500 and 1,000 mg/L OIT, which could not accurately be assessed because of their great inhibition zones (diameters > 50 mm; ++++). The most susceptible fungi to BAC were the mentioned yeasts, Trichosporum sp. and Torulaspora sp., as well as Exophiala sp., with inhibition zone diameters lower than 30 mm (Table 2). Growth inhibition by BAC was very limited (5-15 mm diameter) or non-existent in 9 of 17 evaluated species, including the prevalent Simplicillium sp., Mucor circinelloides, and Trichoderma sp.

These results provide valuable knowledge about the susceptibility of cave fungi to common biocides. According to these data, one may think that a biocide treatment based on both active substances (especially OIT), could be a useful tool to control the fungal growth in Magura Cave. However, biocide efficacy is questionable in cave ecosystems because of the presence of complex biofilms, inaccessible to biocides, and the fast inactivation of biocides through biotic and abiotic factors (Martin-Sanchez et al. 2012). In this sense, about the *in vitro* biocide assay included in this study, it should be noted that inhibition zones shown in Table 2 were recorded after a few days of incubation (ranged from 2 to 7 days, mostly 3 days), during the initial phase of fungal growth. However, afterwards, fungal colonies began to colonize the inhibition zone progressively reducing its area with the time. Hence, it is expected that efficacy of biocides will be limited to a time period after application, especially under natural cave environmental conditions.

Some cave restoration efforts in the last decades applied conventional biocides leading to detrimental effects on the complex microbial communities inhibiting caves (Saiz-Jimenez, 2013). Intense biocide treatments were applied in Lascaux Cave.

Table 2. Biocide susceptibility of fungi isolated from Magura Cave.

<table>
<thead>
<tr>
<th>Strains</th>
<th>Identification</th>
<th>Benzalkonium chloride</th>
<th>2-Octyl-3-isothiazolinone</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Growth inhibition</td>
<td>Growth inhibition</td>
<td></td>
</tr>
<tr>
<td></td>
<td>100 mg/L</td>
<td>500 mg/L</td>
<td>1,000 mg/L</td>
</tr>
<tr>
<td>P14*</td>
<td>Penicillium sp.</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>2-7</td>
<td>Penicillium solitum</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>P38*</td>
<td>Aspergillus sp.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>P3</td>
<td>Simplicillium sp.</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>S1-8</td>
<td>Trichosporon sp.</td>
<td>+</td>
<td>++</td>
</tr>
<tr>
<td>5-2</td>
<td>Gymnoascus reessii</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>S2-6</td>
<td>Mucor circinelloides</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>S1-5</td>
<td>Humicola grisea var. grisea</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>S0-9</td>
<td>Trichoderma sp.</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>P18</td>
<td>Acremonium zeae</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>S0-12</td>
<td>Exophiala sp.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>S0-2</td>
<td>Bionetria rossmaniae</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>S1-6</td>
<td>Arthrinium arundinisi</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>S1-9</td>
<td>Torulaspora sp.</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>S2-2</td>
<td>Pochonia sp.</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>S2-4</td>
<td>Doratomyces sp.</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>P31.2</td>
<td>Pseudogymnoascus sp.</td>
<td>-</td>
<td>+</td>
</tr>
</tbody>
</table>

*aAll selected strains are representative ones included in Table 1, except (*) P14 and P38, whose representative strains are P5 and P23, respectively.

*bInhibition zones were measured after 3 days of incubation (25°C, darkness), except for the strains S2-6 (after 1 day), 5-2 and S2-2 (after 2 days), P3 (after 4 days), and S0-12 (after 1 week).

*Scale according to inhibition zone diameter: no inhibition (+), 5 - 15 mm (+), 16 - 30 mm (++), 31 - 50 mm (+++), and >50 mm (+++). Values of inhibition zone diameter were average of three replicates.
during several periods over years, mainly based on BAC but including other compounds such as OIT, formaldehyde, streptomycin sulphate and polymyxin sulphate. In particular, BAC concentrations used were ranged from 5 to 25 g/L, and OIT concentrations around 1 g/L (Martin-Sanchez et al., 2015). Such prolonged chemical treatments caused negative effects to the cave environment, and selected resistant microorganisms which later played a key role in the black stain outbreak (Bastian et al., 2009; Martin-Sanchez et al., 2012, 2015). Akatova et al. (2009) evaluated the efficiency of different cleaning protocols on phototrophic biofilms of Salpetre Cave (Collibató, Spain), including mechanical removal using 70% ethanol and subsequent treatment with BAC biocides. The treated areas of speleothems were monitored under white or green light. After one year, none of the treatments was fully effective to control the recolonization by phototrophic microorganisms in the stalactite illuminated with white light. Recently, Urzì et al. (2016) reported the changes of the microbial assemblages in green/greyish phototrophic biofilms from Domitilla Catacombs (Rome, Italy) caused by a biocide treatment with the same compounds, BAC (8-12 g/L) and OIT (1.4-2 g/L). In this study, such biocides had little effect on cyanobacteria, but caused significant changes on bacterial population increasing its number and diversity.

Some authors have proposed the use of hydrogen peroxide as an environmental friendly alternative to control microbial outbreaks (Faimon et al., 2003; Boston et al., 2006). Jurado et al. (2010) reported satisfactory results using this product to control the fungal outbreak in Castañar de Ibor Cave, Spain, mainly associated with *Mucor* species. They proposed carrying out mechanical removal of visible fungal colonies followed of disinfection of affected areas by hydrogen peroxide application. The main advantage of hydrogen peroxide is its degradation in contact with organic matter towards harmless molecules for the cave environment such as water and free oxygen gas.

**CONCLUSIONS**

Rock-art caves, like Magura Cave, are one of the most valuable properties of our ancient cultural heritage which attract massive tourism. For an appropriate cave management, especially in rock-art caves opened for visitors, it is essential to investigate the diversity and ecological role of microorganisms inhibiting there. This study provides the first scientific report on cultivable fungal assemblages dwelling different substrata in the Art Gallery of Magura Cave. As typical for caves, the fungal diversity in this gallery was clearly dominated by *Penicillium* and *Aspergillus* species which might due to their ubiquitous distribution in the nature. The most obvious growth of fungal mycelia in this gallery was observed in sediments rich in bat guano. These samples showed much higher number and diversity of colonies grown on culture plates than samples collected from rock surface of painted walls. Besides *Penicillium*, the dominant phylotypes in these sediments were guanophilic fungi such as *Mucor*, *Mortierella*, *Trichosporon* and *Trichoderma*. Conversely, on the rock, a few small areas near the paintings showed evident fungal mycelia, and scarce fungi were isolated from some of these samples.
Based on biocide susceptibility assay, the active compounds BAC and OIT were effective inhibiting the in vitro growth of dominant species from Magura Cave when applied at concentrations ranged from 100 to 1,000 mg/L. These data provide a valuable knowledge about Magura fungi, and exemplify a kind of preliminary test that may be conducted before planning any biocide treatment. However, considering the irreversible effects of biocides on the ecological balance in cave environments, multiple factors should carefully be evaluated before taking such critical decision in caves with rock-art paintings. As described by Martin-Sanchez et al. (2012), preliminary studies on possible advantages and disadvantages of applying biocides in each particular case are required. Any biocide treatment should be planned after testing the active compounds under different conditions, including (i) laboratory assays like in this study, but also (ii) field assays in real conditions, which are very difficult to design properly. In Magura Cave, considering the low fungal contamination in rock surfaces of painted walls, and the previously mentioned risks, there is no reason to use conventional biocides. These treatments should be avoided in order to preserve the relatively stable current balance, which of course is already altered compared to its original natural condition. Alternatively, if a more significant fungal outbreak arises, cleaning protocols combining careful mechanical removal of fungal biomass and subsequent disinfection of affected areas using hydrogen peroxide could be conducted, as long as this cleaning treatment does not lead to damages in rock-art paintings.

Human visitors can influence the quantity and diversity of microbiota in caves through of microclimate changes, organic inputs as well as lighting systems. Such human impacts only could partially be predicted by periodic microclimatic and microbiological studies. To the best of our knowledge, in Magura Cave, there was no microclimatic study up to date, and the microbiological reports are still scarce to shed light on its microbial ecology. Hence, further studies, monitoring of microbial communities and microclimatic parameters in the different cave substrata, should be conducted in order to improve the knowledge on microbial ecology in Magura Cave and to allow the early detection of potential microbial outbreaks.

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