INTRODUCTION

Speleothems of different shapes and colours, varying from black to orange to white are typical of many caves. The fluorescence of calcite speleothems is associated with humic and fulvic acids that are incorporated within the calcite crystal structure (Brennan & White, 2013). These acids originate from overlying soils and are carried into caves by percolation water (McGarry & Baker, 2000) on a seasonal cycle (Ban et al., 2008). Spectroscopic analysis of speleothems of known age appears to provide a suitable tool to facilitate future study of changes in soil and vegetation over time (Brennan & White, 2013).

Various biological mats occur alongside speleothems as a part of the subsurface habitat and decoration. They colonize rock, speleothems and sediment surfaces as yellow, white, pink, tan and gold-coloured spots, known among cave explorers as “sparkles” or “cave gold” for yellow-gold spots and as “cave silver” for those seen through condensed water droplets (Mulec, 2008). These subaerial biofilms, developed on solid mineral surfaces, are widespread under various climate and radiation conditions (Gorbushina, 2007). Especially in karst caves, the biofilms are observed in locations related to intense water condensation or seepages, and in the vicinity of underground rivers and cave entrances. Microbes colonizing surfaces in caves are responsible for changing the integrity of substrata, for example the deterioration of Palaeolithic paintings in Altamira Cave, Spain (Saiz-Jimenez et al., 2011). In these microbial mats Proteobacteria and Actinobacteria commonly dominate, followed by Bacteroidetes, Gemmatimonadetes, Firmicutes, Planctomycetes, Nitrospirae, Verrucomicrobia and Chloroflexi (Portillo et al., 2008; Portillo et al., 2009; Pašić et al., 2010). Similar dominant groups have been identified in lava tubes, and similarities were established among the extensive bacterial diversity of Terceira island in the Azores and the Big Island...
of Hawaii, two locations that are widely separated geographically (Dattagupta, 2014). In lava caves (Northup et al., 2011), but also in karst caves (Cuevza et al., 2012; Keiner et al., 2013), some microbial mats are associated with mineral formations.

Circular golden-yellowish colonies, which are also found as confluent growth, are the microbial mats most easily observed during cave exploration, other than those that are linked to lampenflora. Lampenflora is the term most commonly used to describe a community of organisms, mainly phototrophic, that develops near artificial light sources in show caves (Mulec, 2012). Such yellow mats in karst caves are periodically or constantly covered with a water film and some of them display strong adhesion to the substrata. A different fauna, especially troglobitic animals such as moths, flies and harvestmen, is observed at places with yellow mats close to cave entrances. The objective of the study was to undertake spectroscopic evaluation of yellow-pigmented subaerial biofilms and the applicability of MALDI-TOF (Matrix-Assisted Laser Desorption/Ionization Time-Of-Flight) technique for identification of cave bacteria on these samples, using the commercially available mass spectra database. Human colour perception of yellow subaerial biofilm in caves was considered further in relation to the spectroscopic characteristics of the pigment and lamps that are used to illuminate the caves.

**MATERIAL AND METHODS**

**Caves**

Yellow subaerial biofilm was sampled in three caves in Classical Karst area of southwestern Slovenia, where it was well developed and accessible to sample: Dimnice (No. 737 in the Cave Register of the Karst Research Institute ZRC SAZU and Speleological Association of Slovenia), Križna jama (Cave Register No. 66), and Sveta jama (Cave Register No. 1,158). All three caves have developed passages at different levels, with notable air flows in some galleries, and they are open to the public. Vegetation above these caves comprises shrub and forest. Temperature at the sampling sites was measured using a portable Kestrel 4,500 PocketWeather Tracker (USA).

Dimnice (45°33’48.63”N 14°2’25.65”E) has its main entrance at 567 m a.s.l., with 6,020 m of cave passages, and is formed in Upper Cretaceous limestone with notable air-streaming due to a chimney effect. During the winter incoming air at the main entrance lowers the cave temperature, primarily in the lower part of the entrance passage (Slabe, 1989; Prelovsček, 2012). Because of the lower temperature of the cave walls, condensation generally occurs close to the main entrance and is not related to percolation water. Microbial biofilms of different colours, e.g. white, yellow, purple or brown, are commonly observed at locations with traces of sediments and percolation seepages, and they are often covered with water droplets (Fig. 1B, 1D). Yellow biofilm was sampled on 8 April 2014; air temperature at the site was 6.0°C. On the day of sampling, surface air temperatures in the region were: mean air temperature 12.9°C; maximum daily temperature 19.9°C; and minimum 3.5°C. Data were collected at the Postojna Meteorological Station (45°45’56.78”N 14°11’28.85”E; Slovenian Environment Agency: http://www.meteo.si). The cave attracts approximately 3,000 visitors per year.

The explored cave passages of Križna jama (45°44’42.25”N 14°27’51.04”E), which total 8,273 m in length (entrance at 629 m a.s.l.), are developed in Jurassic limestone with lenses of dolomite (Buser et al., 1967). The cave is connected directly to the surface via a natural entrance and also indirectly by a stream that sinks in Bloško polje and flows through the cave, where it influences air temperature and humidity (Gospodarič, 1974; Slabe, 1995; Bosak et al., 2010; Prelovsček, 2012). Križna jama is an active river cave with notable air-streaming due to a chimney effect. During the winter incoming air at the main entrance lowers the cave temperature, primarily in the lower part of the entrance passage (Slabe, 1989; Prelovsček, 2012). Because of the lower temperature of the cave walls, condensation generally occurs close to the main entrance and is not related to percolation water. Microbial biofilms of different colours, e.g. white, yellow, purple or brown, are commonly observed at locations with traces of sediments and percolation seepages, and they are often covered with water droplets (Fig. 1B, 1D). Yellow biofilm was sampled on 8 April 2014; air temperature at the site was 6.0°C. On the day of sampling, surface air temperatures in the region were: mean air temperature 12.9°C; maximum daily temperature 19.9°C; and minimum 3.5°C. Data were collected at the Postojna Meteorological Station (Slovenian Environment Agency: http://www.meteo.si). Around 5,000 visitors visit the entrance part of the cave each year.

Sveta jama (45°35’23.66”N 13°51’54.14”E), with its entrance at 420 m a.s.l., has 231 m of surveyed cave passages, all formed in Palaeocene/Eocene foraminiferal limestone (Pleničar et al., 1969). This cave has no active streamways to influence the cave climate conditions. Sveta jama is the only cave in Slovenia that serves as a church, and it attracts 2,000 tourists annually. In Sveta jama, microbial biofilms of different colours are scattered throughout the cave; confluent growth (continuous bacterial growth without discrete colonies) is rarely observed. Yellow biofilms are covered with water droplets (Fig. 1C). Subaerial yellow biofilm was sampled on 18 March 2014; cave-air temperature was 9.4°C.

**Sampling and cultivation**

In all three caves yellow subaerial biofilms were sampled at sites where organic matter enters the underground system, at seeps that indicate good connectivity with the epikarst and in the vicinity of...
underground rivers or natural entrances. Using sterile equipment, biofilm was scraped from rocky surfaces, transferred in a cool box to a laboratory, weighed, serially diluted in 0.9% physiological saline and plated on culture media. To mimic the natural environment, with evidently higher organic input compared to other parts of the caves, the selected cultivation media were not primarily supportive for oligotrophic microbes. Seven different media were used to propagate microbial colonies and to estimate the cultivable part of the community: (1) Nutrient agar (NA, Fluka); (2) Malt extract agar (MEA, Fluka); (3) Sediment Agar (SA), which contained 1.0% of old cave alluvial sediment and 1.5% agar; (4) RIDA®COUNT Total Aerobic Count (R-biopharm) for heterotrophic aerobic bacteria; (5) RIDA®COUNT E. coli/Coliform (R-biopharm) for Escherichia coli and coliforms; (6) RIDA®COUNT Salmonella/Enterobacteriaceae (R-biopharm) for enterobacteria and Salmonella; and (7) RIDA®COUNT Yeast&Mold Rapid (R-biopharm) for yeasts and moulds. RIDA®COUNT plates provide a relative good proxy to monitor human impact underground (Mulec et al., 2012a; Mulec et al., 2012b).

Petri plates with NA, MEA, SA and RIDA®COUNT for fungi were incubated aerobically at room temperature (~22°C), and RIDA®COUNT selective media for Escherichia coli /coliforms and enterobacteria/ Salmonella were incubated at 37°C. The inoculated plates were incubated at higher temperatures than those measured in the source caves. A previous study demonstrated that bacteria isolated from caves in Spain grew comparatively well at temperatures ranging from 13 to 45°C; however, the metabolic profile of isolates differed, which indicated an adaptive response to temperature (Laiz et al., 2003). Cultivation lasted up to 96 hours. Visible colonies were quantified in terms of Colony-Forming-Units (CFU) per g. Incubation for 4 days retrieved mostly r-strategists (fast growers) and partly K-strategists (slow growers) as the general conditions to study r- and K-strategists require 3 days to determine r-strategists and an additional 4-7 days for K-strategists at 20°C (Krišúček et al., 2005).

Characterisation of isolates

195 bacterial isolates retrieved on NA, MEA, SA, RIDA®COUNT Total Aerobic Count and RIDA®COUNT Yeast&Mold Rapid were inoculated on Blood Agar, incubated at room temperature for 24 to 48 hours and subjected to identification with MALDI-TOF using direct smear technique (Bruker Daltonik, Germany). Briefly, a smeared bacterial colony on the MALDI Steel plate was overlain with a photo-absorptive matrix – α-cyano-4-hydroxycynamic acid (CHCA) - and left to dry before analysis with Bruker MALDI Biotyper RTC software version 3.1 (Seng et al., 2009). Quality of identification was assessed according to the manufacturer (Bruker Daltonik, Germany) using the LogScore value. If the obtained LogScore value for a given isolate was below 1.700, the identification was considered unreliable, and if the value was 1.700 or higher the identification was considered reliable.

Spectroscopic analyses

Pigments from the biofilms were extracted using three different solvents, deionized water, 90% acetone and 96% ethanol. The biofilm from Sveta jama was extracted only in deionized water due to the small quantity of sampled material available. The suspension was incubated overnight in the dark (stability of pigment when exposed to light is not yet determined) and occasionally mixed vigorously. After centrifuging at 4,000 RPM for 10 min, spectroscopic analysis was performed using a Lambda 25 UV-Vis Spectrometer (Perkin-Elmer, USA).

Before fluorescence analysis was carried out using a Luminiscence Spectrometer LS 30 (Perkin-Elmer, USA), the supernatant liquid was additionally centrifuged at 14,000 RPM for 10 minutes. The excitation monochromator was set at 405 nm and the fluorescence monochromator was set to record emissions between 435 and 750 nm.

The emission spectra of lamps used to light caves during explorations were measured in the laboratory, using a Jaz spectrometer (Ocean Optics, USA; detector 200-1,000 nm). These light sources included: a wax candle; a carbide lamp; a LED (Light Emitting Diode) lamp composed of 14 individual LEDs and a...
RESULTS AND DISCUSSION

Biofilm and microbial identification

Different concentrations of cultivable microorganisms in terms of CFU/g were retrieved on selected media. The highest number of isolates was retrieved in a sample from Sveta jama. In this sample 6.5-times more colonies were retrieved on SA than on NA. The concentrations of cultivable microbes on SA and NA for the other two caves were similar. In contrast, the Sveta jama sample on RIDA COUNT plates for total heterotrophic bacteria showed the lowest concentration of cultivable microbes, $17 \times 10^3$ CFU/g (Table 1). Only a few coliform bacteria expressing β-galactosidase activity were retrieved, and there were no bacteria associated with faecal pollution, i.e., E. coli, enterobacteria. Except in one sample from Križna jama, the RIDA COUNT Yeast&Mold Rapid plates from the other two caves showed that fungi were present (Table 1). The low concentration of cultivable fungi, which are generally abundant in different cave microhabitats (Mulec et al., 2002), can be attributed to non-optimum selection of media. Yellow subaerial biofilms on solid surfaces represent live microbial biomass that serves as a nutrient source for other cave dwelling organisms. Fauna not only graze on biofilm, but also can help in microbial transmission in other parts of caves.

Table 1. Estimations of cultivable microorganisms using different cultivation conditions and media.

<table>
<thead>
<tr>
<th>Medium</th>
<th>Cultivation temperature (°C)</th>
<th>Dimnice $\times 10^3$ CFU/g</th>
<th>Sveta jama $\times 10^3$ CFU/g</th>
<th>Križna jama $\times 10^3$ CFU/g</th>
</tr>
</thead>
<tbody>
<tr>
<td>MEA</td>
<td>RT</td>
<td>0</td>
<td>88</td>
<td>0</td>
</tr>
<tr>
<td>NA</td>
<td>RT</td>
<td>1,230</td>
<td>35,400</td>
<td>809</td>
</tr>
<tr>
<td>SA</td>
<td>RT</td>
<td>1,390</td>
<td>233,000</td>
<td>755</td>
</tr>
<tr>
<td>RIDA-Tot</td>
<td>37</td>
<td>44</td>
<td>17</td>
<td>399</td>
</tr>
<tr>
<td>RIDA-Col</td>
<td>37</td>
<td>0</td>
<td>0</td>
<td>5</td>
</tr>
<tr>
<td>RIDA-Ent</td>
<td>37</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>RIDA-Y&amp;M</td>
<td>RT</td>
<td>3</td>
<td>42</td>
<td>0</td>
</tr>
</tbody>
</table>


Altogether 195 bacterial isolates from the three caves were subjected to identification by MALDI-TOF and 149 were identified according to the commercially available mass spectra library at species and genus level. The highest number of bacteria identified at species level was in a sample from Sveta jama (66.0%), followed by Dimnice (43.5%) and Križna jama (19.9%). Different species of Pseudomonas prevailed, with the highest MALDI score value at 2.275 for Pseudomonas jessenii from Križna jama. The highest diversity at genus level was in a sample from Križna jama (Bacillus, Paenibacillus, Pseudomonas, Streptomyces and Variovorax), followed by Dimnice (Bacillus, Flavobacterium, Pseudomonas and Streptomyces) and Sveta jama where only different representatives of Pseudomonas were identified (Fig. 2). The cultivable diversity of cave subaerial biofilm from another Slovenian cave (Pajsarjeva jama) revealed that the majority belonged to Streptomyces (25%), Micrococcus (16%), Rhodococcus (10%) and Pseudomonas (9%) (Velikonja et al., 2014). Not only Streptomyces (e.g. Shirling & Gottlieb, 1966), but also Micrococcus (e.g. Brooks et al., 1980), Rhodococcus (e.g. Ichiyama et al., 1989) and Pseudomonas (e.g. Meyer, 2000) are well known pigment-producing bacterial genera. As in the current study, representatives of Bacillus and Paenibacillus were also commonly identified in Pajsarjeva jama, as well as Pseudomonas and Streptomyces. During the Pajsarjeva jama study (Velikonja et al., 2014) microbes were cultivated at 30°C for 5 weeks on nutrient-rich (glycerol-asparagine agar, peptone-yeast extract-brain heart infusion agar, 1,000-fold-diluted tryptic soy agar, starch-casein agar, malt-yeast extract agar, soil extract agar) and oligotrophic media (tap water agar). Results of culture independent studies and comparisons of environmental 16S rRNA on microbial mats from Spain, Slovenia, the Czech Republic, Portugal and Hawaii indicated that actinobacterial Pseudonocardinaceae and Gammaproteobacteria are the common dominant bacterial groups (Porca et al., 2012). Isolated fungi were not subjected to identification by MALDI.

Successful identification of microbes by means of the MALDI-TOF technique depends upon the limited availability of mass spectra in the database. Combination of MALDI-TOF with proper identification of environmental isolates, including cave isolates based on 16S rRNA and ITS sequences, will provide a tool for rapid, simple and cost-effective identification. In addition to species identification, the questions related to pigment characteristics can be studied simultaneously.

Fluorescence of microbial biofilm and spectroscopy of yellow pigment

Synthesis of pigments among bacteria is a common phenomenon; the majority of bacteria identified by MALDI-TOF from yellow mats can produce pigments and contribute to the colouring of caves. For example, Flavobacterium can produce the yellow carotenoid pigment, zeaxanthin (Dufosse, 2006). Some Bacillus also produce pigments (Tobie, 1945) and Variovorax has a xanthomonadin biosynthetic gene cluster (Zhou et al., 2013). The human pathogen Staphylococcus aureus synthesizes triterpenoid carotenoid pigments (Marshall & Wilmoth, 1981). Many Pseudomonas are known to have versatile metabolism (Rojo, 2010) and to synthesise pigments (Meyer & Abdallah, 1978). Pseudomonas produces a yellow-green water soluble fluorescent siderophore pyoverdine, which is responsible for the capturing of iron (Meyer, 2000). Many streptomyces also produce different pigments that, in caves with Paleolithic rock paintings, contribute notably to the
biodeterioration of rock art (Groth et al., 1999). The small number of streptomycetes revealed in this study can be linked to the selection of non-optimum growth media. At the moment it is not clear whether absorption and emission spectra of the biofilm (Fig. 3A, C, and E) belong to one microbial species or more, or whether they represent a combination of more pigments.

The yellow pigment is partially water soluble. This can be observed in caves when condensed water droplets on intact subaerial biofilms adopt a yellowish hue. Biofilms from all three analysed caves displayed a similar absorption maximum, in the violet part of the light spectrum. The absorption peak from Dimnice was at 407 nm when water was used as solvent, at 410 nm in acetone and at 393 nm in ethanol (Fig. 3A). Pigment from Križna jama showed an absorption peak at 419 nm in acetone, but in water and ethanol the peak was less well expressed than in the sample from Dimnice (Fig. 3C). Pigment from a Sveta jama sample had a maximum absorption in water at 400 nm (Fig. 3E). Pigments extracted in water were further excited by light at 405 nm (violet). The emission spectra were very similar for all samples having two emission peaks. In a Dimnice sample the major peak was at 460 nm (blue) with a minor one at 600 nm (orange) (Fig. 3A). In a sample from Križna jama, the major peak was at 460 nm and the minor peak at 602 nm (Fig. 3C). Similarly, the major peak for a Sveta jama sample was at 460 nm and the minor peak at 603 nm (Fig. 3E). Additional analyses are needed to help unravel the chemical structure of the pigment(s).

Perception of yellow subaerial biofilm and its potential role

The human colour perception of cave subaerial biofilms depends both upon the quality and quantity of the light that is used for illumination and on the presence or absence of a water film that acts as a magnifying glass over the microbial mats. Different light sources are used for exploration in caves and mines; in the past carbide lamps and candles were commonly used, but LED lamps have been used more recently. To a degree carbide lamps (Fig. 3B), wax candles (Fig. 3B) and halogen lamps (Fig. 3D) have similar spectra, with a large emission extending from the nominal red edge of the visible spectrum at 700 nm further towards the infrared area that is largely responsible for heat emission. Human colour perception of objects is different in the case of LED lamps. Analysis of emission spectra of the two commonly used cool white LED lamps showed two emission maxima, for the Petzl Duo at 469 and 556 nm, and for the Scurion at 446 and 556 nm (Fig. 3D). Such LEDs emit light close to the absorption maximum of yellow biofilm, which is why such biofilms are more clearly traced in caves when they are illuminated by this type of lamp. Specifically, the relative low quantity of photons in the violet-blue part of the spectrum as against the orange-red part of the spectrum for carbide lamps and candles results in less fluorescence of yellow pigmented biofilms when these two lamps are used for lighting. How humans perceive colours depends also upon light intensity. When scotopic vision is dominant at luminance levels between of $10^{-2}$ and $10^{-6}$ Lux the luminous...
efficiency is different to that when photopic vision dominates (1-10^6 Lux). During periods of scotopic vision the human eye is most sensitive to light around 500 nm, whereas the sensitivity peak shifts to a longer wavelength, around 550 nm during photopic vision (Fig. 3F) (Judd & Wyszecki, 1975; Vos, 1978; Sharpe et al., 2005). Because of this physiological adaptation of human eyes, yellow subaerial biofilms are more vividly observed in comparison with other coloured mats in caves with generally low levels of illumination.

Many authors have studied coloured bacterial mats in lava tubes (Northup et al., 2011; Hathaway et al., 2014) and karst caves (Mulec, 2008; Pašić et al., 2010; Velikonja et al., 2014) but, nevertheless, little is yet known about their nature and role in subsurface ecosystems. In situations outside caves pigments from secondary metabolism can help to protect biofilms against lethal doses of UV irradiation, acting as “sunscreen” compounds (Gao & Garcia-Porca, 2011); this is not the case inside caves. Roles in other ecological interactions should also be considered, e.g. protection against grazers by production of antibiotics, as with *Streptomyces* (Hobbs et al., 1990), or secretion of siderophores. Siderophores are important in virulence (pathogenicity) expression and development of biofilms by different microbes (Wynn-Williams et al., 2002; Saha et al., 2013).

Fluorescence of yellow subaerial biofilm is probably not directly linked solely to the most frequent natural fluorophores, such as aromatic amino acids, flavins, vitamin A, chlorophylls and NADH (Campbell & Dwek, 1984). An absorption peak around 405 nm in the different solvents used indicates stability and potential biotechnological application, for example similar to the Green Fluorescent Protein, GFP. GFP is also very stable with tolerance to fixatives, detergents and chaotropes (Ward, 1998). Further studies are needed, not just to explain the role of pigmented subaerial biofilms in caves, but also to understand their interaction with cave surfaces as important players in changing cave micromorphology, e.g., biochemically induced etching of solid surfaces, and their potential use in biotechnology and medicine.

Caves do offer an important biotechnological pool of

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**Fig. 3.** Emission and absorption spectra: A) absorption spectra of subaerial biofilm from Dimnice after pigment extraction in deionized water (Dim), 90% acetone (Dim-Ac), 96% ethanol (Dim-Et), and a fluorescence spectrum of the extract in deionized water when excited at 405 nm (Ex = 405); B) emission spectra of a carbide lamp and a candle; C) absorption spectra of subaerial biofilm from Križna jama after pigment extraction in deionized water (Kri), 90% acetone (Kri-Ac), 96% ethanol (Kri-Et), and a fluorescence spectrum of the extract in deionized water when excited at 405 nm (Ex = 405); D) emission spectra of a halogen lamp and a Petzl Duo LED caving lamp and a Scurion LED lamp; E) absorption spectra of subaerial biofilm from Svéta jama after pigment extraction in deionized water, and a fluorescence spectrum of the extract in deionized water when excited at 405 nm (Ex = 405); F) spectral luminous efficiency function for scotopic and photopic vision (Judd & Wyszecki, 1975; Vos, 1978; Sharpe et al., 2005).
microbes, for example in Carlsbad Cavern, USA, a microbe has been isolated that can degrade hazardous benzothiazole (Barton, 2006).

CONCLUSIONS

Yellow biofilms on surfaces in caves are an important source of live microorganisms. They are well-observed in caves due to the contrast with the surrounding surfaces, and recently more evident due to fluorescence characteristic of the pigment(s) related to the usage of cool white LEDs for illumination. It is not yet clear whether the two emission peaks from yellow biofilms are associated with the same pigment; nor is the role of pigment(s) in underground habitats fully understood. Subaerial biofilms that cover surfaces in caves are probably more widespread than current data suggest, but some of them are difficult to observe when the contrast with the underlying surface is low, for example white or transparent colonies on white rock surfaces. MALDI-TOF showed considerable promise as an identification and research tool for future use in cave microbiology.

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