Characterization of antimicrobial activity of culturable bacteria isolated from Krubera-Voronja Cave

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Abstract: In the present study we aimed to perform the first analysis of antimicrobial activity of bacteria isolated from Krubera-Voronja Cave, with the main focus on their activity against Gram-positive bacteria, including Gram-positive pathogens. Using five different media, in total 874 heterotrophic cultures were isolated from water and sediment samples collected in Krubera-Voronja Cave at a depth from 220 m to 1640 m. 14.0% of all isolates demonstrated antibacterial activity against Gram-positive and Gram-negative test microorganisms. Our results show that this percentage was not uniform: it increased with the sampling depth and was the highest in the lower part of the cave. 24 isolates were active exclusively against Gram-positive test strains *Micrococcus luteus* and *Bacillus thuringiensis*. Two isolates, namely strains 1350R2-TSA30-6 and 1410WF1-TSA30-2, were chosen for the further work because of the high and comparable activity against both Gram-positive test microorganisms. It was determined that both strains belong to the family *Bacillaceae* in phylum *Firmicutes*. The detailed bioactivity analysis of these two Gram-positive strains revealed the different mixtures of volatile compounds with antibacterial activity. The main antibacterial compounds of the strain 1350R2-TSA30-6 are pyrrolopyrazines pyrrolo[1,2-a]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)- and pyrrolo[1,2-a]pyrazine-1,4-dione, hexahydro-3-(phenylmethyl)-. The main antibacterial compound of the strain 1410WF1-TSA30-2 is 1,2-benzenedicarboxylic acid, bis(2-methylpropyl) ester. Mixtures of the volatile antimicrobial compounds of both strains were antagonistic against Gram-positive strains isolated from Krubera-Voronja Cave, and their activity against Gram-positive pathogenic bacteria substantially differed.

Keywords: antimicrobial, *Bacillus*, cave microorganisms, Krubera-Voronja Cave, volatile compound

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INTRODUCTION

Antibiotic resistance of pathogenic bacteria became one of the greatest threats to human health over the last few decades (Blair et al., 2015). A few different approaches are used to resolve this problem: chemical synthesis of novel antibiotics (Wright et al., 2014), discovery and development of novel agents capable of circumventing or neutralizing the existing resistance mechanisms (Blair et al., 2015), as well as search for new natural antimicrobials (Lamprinou et al., 2015). Considering the last-mentioned approach, caves represent one of the most attractive environments with a strong potential for the discovery of novel antimicrobials. Antimicrobial activity of different cave microorganisms against pathogenic bacteria was reported previously (Herold et al., 2005; Nakaw et al., 2009; Rajput et al., 2012; Cheeptham et al., 2013; Rule & Cheeptham, 2013; Tomova et al., 2013; Lamprinou et al., 2015); a new genus of bacteria, producing antimicrobial substances, was identified in a cave in Thailand (Nakaw et al., 2012).

Ecological conditions enabling cave microorganisms to produce antimicrobial compounds are unclear. Montano & Henderson (2013) showed that the frequency of human visitation does not affect the antimicrobial activity of cave microorganisms, while the increasing depth, and consequently increasing oligotrophy do. Energy-starved conditions in the caves prompt the complex interactions (competitive or cooperative) between different microorganisms resulting in the production of secondary metabolites,
such as pigments, siderophores, and antibiotics, which are used as informational cues (Rajput et al., 2012; Gabriel & Northup, 2013). Krubera-Voronja Cave is the Earth’s deepest cave (Sendra & Reboleira, 2012). Because of its depth, high antimicrobial activity of microorganisms living in it could be expected, but it has not been investigated so far.

Antimicrobial potential of cave actinobacteria and especially *Streptomyces* was investigated (Herold et al., 2005; Nakaew et al., 2009; Hodges et al., 2012; Rajput et al., 2012; Cheeptham et al., 2013; Rule & Cheeptham, 2013; Nimaichand et al., 2015), but the antimicrobial compounds of bacteria from other phyla – *Cyanobacteria* (Lamprinou et al., 2015), *Proteobacteria* and *Bacteroidetes* (Tomova et al., 2013) – were also of scientific interest. Our previous cultivation-independent analysis of bacterial diversity in Krubera-Voronja Cave showed that Gram-positive phyla *Actinobacteria* and *Firmicutes* were among the most numerous phyla in this cave (Kieraite-Aleksandrova et al., 2015). This indicates during the evolution the indigenous Gram-positive bacteria of Krubera-Voronja Cave could have evolved the mechanisms to compete and/or collaborate specifically with other Gram-positive bacteria. The production of antimicrobial compounds could be regarded as one of those mechanisms. Antimicrobial activity of the indigenous Gram-positive bacteria of Krubera-Voronja Cave against other Gram-positive bacteria is still unknown and should be evaluated, the rather that the search for novel antimicrobials targeting Gram-positive pathogens is of major importance. These bacteria most commonly cause severe nosocomial infections, and the resistance of these pathogens to conventionally used antimicrobials continuously increases and strengthens (Herold et al., 2005). Therefore, screening for novel narrow-spectrum antimicrobials, that would be active against Gram-positive pathogens, is vital.

In the present study we aimed to perform the first analysis of antibacterial activity of microorganisms isolated from Krubera-Voronja Cave, mainly focusing on their activity against Gram-positive bacteria including Gram-positive pathogens.

**MATERIAL AND METHODS**

**Site description and sampling**

Krubera-Voronja Cave is located in the Arabica Massif (43.4184 N 40.3083 E, Western Caucasus). Sampling in Krubera-Voronja Cave was described explicitly in our previous work (Kieraite-Aleksandrova et al., 2015). It was performed on 30th July to 7th August 2012, at depths of 220 m to 1640 m. At the time of sampling, humidity ranged from 69-76% in the upper part (220-230 m in depth) of the cave to >90% in the middle (500-700 m in depth) and the lower (1215-1640 m in depth) parts of the cave. The temperature in the cave ranged from 3-4°C in the upper and middle parts to 6-8°C in the lower part of the cave.

In total, 26 samples were collected from Krubera-Voronja Cave. The collected sample materials included soil and clay from cave walls, sediment, speleothems, drinkable water from the underground camps as well as coloured spots from cave walls. All samples were collected using aseptic techniques and placed into sterile microcentrifuge tubes. The samples were transported in a cooler and stored at -20°C until microbial analysis was performed.

**Isolation of bacterial cultures**

For isolation of culturable microorganisms from water samples, 100 μL of each sample was spread over different solid microbiological media dispensed in Petri dishes. For isolation of culturable microorganisms from sediment samples, 300 mg of each sample was suspended in 700 μL of saline, and 100 μL of each suspension was spread over different solid microbiological media. Tryptic Soy agar (TSA) (Merck Millipore), Hickey-Tresner (HT) agar, Actinomyces Isolation (AI) agar, Starch Casein Nitrate agar (SCNA), and Dičo™ ISP medium 4 (ISP4) were used. The cultivation media were selected according to Cheeptham et al. (2013) and Hodges et al. (2012) as well as the previous data on the bacterial diversity in Krubera-Voronja Cave (Kieraite-Aleksandrova et al., 2015). It was previously shown that *Actinobacteria*, *Firmicutes*, and *Proteobacteria* were the dominant phyla in this cave. HT agar, AI agar, SCNA and ISP4 are usually used for the isolation and cultivation of *Actinobacteria* and TSA is routinely used for the isolation and cultivation of *Firmicutes* and *Proteobacteria*. Two different temperatures were chosen for the isolation of microorganisms from the cave samples: 4°C and 30°C. The first one was selected according to the temperature profile inside the cave (3-8°C, see above); it is suitable for the growth of psychrophiles that were believed to thrive in the cave. The second temperature is suitable for the growth of psychrotolerant microorganisms that have an optimum temperature between 20°C and 40°C, but could be metabolically active at the cave's temperature. Microorganisms were cultivated at 4°C for 7 days and at 30°C for 3 days. Only those microorganisms that stood apart in colony morphology and in ability to excrete soluble pigments were chosen for further work.

**Primary screening of microbial isolates for antibacterial activity**

In order to identify microbial isolates with antibacterial activity, a modified version of agar diffusion test (cross-streak assay) was used (Montano & Henderson, 2013). Cave microorganisms were streaked and cultivated under the same conditions and on the same solid medium which they were isolated. For example, bacterial cultures 1350R2-TSA30-6 and 1410WF1-TSA30-2 were initially isolated on TSA at 30°C. Therefore, for the primary screening for antibacterial activity, they were restreaked on TSA and cultivated at 30°C. For screening, the plates with the streaks of the cave microorganisms were covered with a layer of sterile TSA. Test microorganisms were streaked on sterile TSA perpendicularly to the already grown streaks of the cave microorganisms. Test
microorganisms were cultivated at 30°C for 24 h. The zone of growth inhibition of the test microorganisms near the cave isolate streak indicated antimicrobial activity of the cave microorganisms. No inhibition was defined as absent interruption of the target test microorganism streak where it crossed the cave isolate streak. Gram-positive (Micrococcus luteus and Bacillus thuringiensis TL8) and Gram-negative (Escherichia coli BL21(DE3) and Pseudomonas sp. VR1) bacteria served as test microorganisms.

**Genotyping of the active isolates**

The bacterial genomic DNA was extracted from fresh cell culture using the GeneJET™ Genomic DNA Purification Kit (Thermo Fisher Scientific) according to the manufacturer's instructions. BOX-PCR was performed in 50 μL of reaction mixture containing DreamTaq Green PCR Master Mix (2X) (Thermo Fisher Scientific), 0.5 μM BOXA1R primer (5’-CTA CGG CAA GGC GAC GCT GAC G-3’) and 10 ng of bacterial genomic DNA. BOX-PCR was conducted under the following conditions: initial denaturation at 95°C for 2 min followed by 29 cycles, each consisting of 95°C for 1 min, 53°C for 2 min, and 72°C for 3 min with a final extension step at 72°C for 7 min. Products of amplification were analysed by electrophoresis through 1% agarose gel.

**Phylogenetic analysis of active isolates**

Amplification of 16S rRNA genes was done according to Kuisiene et al. (2002). PCR products were cloned into E. coli DH5α using the CloneJET™ PCR Cloning Kit (Thermo Fisher Scientific). Recombinant plasmids were isolated using the GeneJET™ Plasmid Miniprep Kit (Thermo Fisher Scientific). 16S rRNA genes of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 were sequenced at the DNA Sequencing Centre (Vilnius University, Institute of Biotechnology, Lithuania); the obtained sequences were edited and sequence similarity was determined using the SEQBUILDER and MEGALIGN components of LASERGENE 6 (DNASTAR). 16S rRNA gene sequences were subjected to phylogenetic analysis using ARB (Ludwig et al., 2004). The size of the 16S rRNA gene used for the analysis was 1372 nt. The tree was rooted using the obtained sequences were edited and sequence similarity was determined using the SEQBUILDER and MEGALIGN components of LASERGENE 6 (DNASTAR). 16S rRNA gene sequences were subjected to phylogenetic analysis using ARB (Ludwig et al., 2004). The size of the 16S rRNA gene used for the analysis was 1372 nt. The tree was rooted using the 16S rRNA gene sequence of Paenibacillus polymyxa DSM 367 as an outgroup.

**Nucleotide sequence accession numbers**

16S rRNA gene sequences of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 were deposited in the GenBank under the No. KU513788 and KU513789 respectively.

**Evaluation of antimicrobial activity of active isolates by agar-well diffusion method**

Test microorganisms were cultivated on TSA medium at 30°C for 24 h and then resuspended in saline. The correlation between the optical density of bacterial suspensions at 600 nm and the number of CFU/mL after the TSA plate counts was established for test microorganisms. For the evaluation of antimicrobial activity, the suspension of the test microorganism was mixed with molten TSA to achieve final 10⁶ CFU/mL. Inoculated TSA was poured into the Petri dishes (20 mL per dish). 5 mm diameter wells were cut in the inoculated agar surface and filled with 100 μL of the sample. The zone of the growth inhibition of test microorganisms indicated antimicrobial activity of the sample and was recorded as a zone diameter (in millimeters) around the wells, minus 5-mm diameter of the well (Choma & Grzelak, 2011; Tomova et al., 2013).

**Optimization of culture growth parameters for production of antibacterial compounds**

Effects of the concentration of nutrients, growth medium pH, temperature and aeration on the culture growth were examined in order to optimize the production of antibacterial compounds. Effects of the concentration of nutrients were examined using 1xTryptic Soy broth (TSB), 0.5xTSB and highly nutritious growth medium Brain Heart Infusion broth (BHIB). To test the effect of pH, growth medium was prepared in 50 mM Tris-HCl buffer (pH 7, 8, and 9/20°C). For determination of the effect of temperature, 30°C (the isolation temperature), 16°C, 8°C (the upper temperature limit of the cave), and 4°C (the lower temperature limit of the cave) were chosen. The effect of aeration was determined at 180 rpm and 250 rpm in an orbital shaker Multitron Standard (INFORS HT).

Inoculation (5% vol/vol) was performed using bacterial cultures grown for 12 h on TSA plates at 30°C. Inocula were prepared in the respective sterile growth medium. Bacteria were cultivated in 250 mL Erlenmeyer flasks, and the samples were aseptically removed every 2 h to determine culture growth and production of antimicrobial compounds. Growth of the culture was monitored by measuring optical density at 600 nm, and antimicrobial activity of the culture was assayed in the culture supernatant by agar-well diffusion method (see above). Cells from the liquid culture were removed by centrifugation (7000×g, 20 min, 4°C).

**Extraction of antibacterial compounds by salting-out with ammonium sulphate**

Active bacterial strains were cultivated in buffered 0.5xTSB for 12 h at 30°C at 180 rpm in the orbital shaker Multitron Standard (INFORS HT). Cells were removed by centrifugation (7000×g, 20 min, 4°C), and solid ammonium sulphate was added to the supernatant to achieve 80% saturation. The precipitate was recovered by centrifugation (12000×g, 20 min, 4°C), then dissolved in 20 mM Tris-HCl buffer (pH 7). The dissolved proteins were evaluated for the antimicrobial activity using agar-well diffusion method. The dissolved proteins were also treated with recombinant proteinase K (Thermo Fisher Scientific) according to the manufacturer's instructions. Antimicrobial activity of hydrolysed proteins was also evaluated using agar-well diffusion method. Protein concentration was determined using the Pierce™ Coomassie (Bradford) Protein Assay Kit (Thermo Fisher Scientific) according to the manufacturer's instructions.
Extraction of antibacterial compounds by organic solvents

Active strains were cultivated in 100 mL of buffered 0.5xTSB for 12 h at 30°C at 180 rpm in the orbital shaker Multitron Standard (INFORS HT). Cells were removed by centrifugation (7000×g, 20 min, 4°C). Antibacterial compounds in the supernatant were extracted with equal volume of acetone, chloroform, ethyl acetate, methanol, and hexane in a separating funnel. Different supernatant samples of the same culture were separately extracted with each solvent. The organic solvent extracts were evaporated to dryness and dissolved in chloroform (Sanghvi et al., 2014).

Thin layer chromatography and bioautography

Thin layer chromatography (TLC) of chloroform extract was done on Silica gel 60 F254 plates (Merck Millipore). The antibacterial compounds were separated with ethyl acetate-methanol (3:1, vol/vol) solvent system. Chromatograms were observed under UV light and exposed to iodine vapours. Bioautography was used for the localization of antibacterial compounds in chloroform extract (Attimarad et al., 2012). TLC plate with the separated antibacterial compounds was dried, put in sterile Petri dish, and overlaid with the inoculated TSA. Inoculated TSA was prepared as described above for the agar well diffusion method. *M. luteus* was used for bioautography as the test microorganism. After incubation at 30°C for 24 h, the zone of inhibition around the spot of antibacterial compound was determined. The fractions with antibacterial activity were recovered from TLC plates, disssolved in ethyl acetate and subjected to further analysis.

Gas chromatography-mass spectrometry (GC-MS) analysis of antibacterial compounds

Gas chromatographic (GC) analysis of the fractions with antibacterial activity recovered from TLC plates was performed on a PerkinElmer Clarus 580 series gas chromatograph coupled to a PerkinElmer Clarus 560 S mass spectrometer (PerkinElmer, Shelton, USA). The GC system was equipped with Elite-5MS capillary column (30 m × 0.25 mm id, 0.25 µm film thickness) coated with methylpolysiloxane (5% phenyl). Helium was employed as a carrier gas with a constant flow of 1 mL/min. GC conditions were as follows: the oven temperature was programmed: 40°C for 1 min, from 40 to 250°C at 3°C/min and held at 250°C for 8 min; the injector temperature was held at 250°C. Injection was performed in pulsed splitless mode (pulsed to 4 mL/min until 1.5 min, split 10:1) open at 1.55 min). The capillary column was connected to the ion source of the mass spectrometer by means of the transfer line maintained at 280°C. The electron ionization ion source conditions were: electron energy 70 eV and temperature 180°C. GC-MS in full scan mode was used. The analyses were carried out with a filament multiplier delay of 3 min, and the acquisition was performed in the range of m/z 33 - 600. The qualitative identification of different compounds was performed by comparing their mass spectra with those stored in the NIST (The National Institute of Standards and Technology) library. The relative amounts of individual compounds were expressed as percentages of the peak area relative to the total peak area. GC-MS identified compounds were compared with the bacterial organic volatile compounds included in the Microbial Volatile Organic Compounds (mVOC) database (Lemfack et al., 2014) as well as with those in the SuperScent database (Dunkel et al., 2009).

Evaluation of antibacterial activity against pathogens and microorganisms of the cave

For evaluation of antibacterial activity against pathogens and microorganisms of the cave, the active fractions, recovered from TLC plates and dissolved in ethyl acetate, were used.

Antibacterial activity was determined against Gram-positive pathogenic bacteria: *Bacillus cereus* ATCC 10876 (Firmicutes), *Enterococcus faecalis* ATCC 29212 (Firmicutes), *Listeria monocytogenes* ATCC 7644 (Firmicutes), *Staphylococcus aureus* subsp. *aureus* ATCC 25923 (Firmicutes), and *Rhodococcus equi* ATCC 6939 (Actinobacteria). Agar-well diffusion method was used for determination of the minimal inhibitory concentration (MIC). The inocula of the pathogenic bacteria were prepared in saline and matched 0.5 McFarland standard. 19 mL of melted and cooled down TSA medium was inoculated with 1 mL of such suspension and poured into Petri dish. The final concentration of cells in TSA was 10⁶ CFU/mL. 5 mm diameter wells were cut in the inoculated agar surface and filled with 100 µL of the sample. Six different concentrations of antimicrobial compounds were used for determination of MIC: 4.66 mg/mL (the undiluted sample), 2.33 mg/mL (1:2 dilution), 1.165 mg/mL (1:3 dilution), 0.5825 µg/mL (1:7 dilution), 291.25 µg/mL (1:15 dilution), and 145.625 µg/mL (1:31 dilution).

In the current work antibacterial activity of the active fractions was also examined against bacteria isolated from Krubera-Voronja Cave: Gram-positive strain 28TSA30-8, *Brevibacillus* sp. 1410WF1-HT30-5, *Paenibacillus* sp. 23TSA30-6, *Paenibacillus* sp. 28ISP30-5, and *Streptomyces* sp. 1350R2-SCNA30. Modified agar-well diffusion method was used for this test. Sterile TSA (20 mL) was poured into the Petri dish, and the well was cut in agar in the centre of the Petri dish. Bacterial cultures were streaked from the edge of the well to the border of the Petri dish. Then the well was filled with 100 µL of the sample containing antimicrobial compounds at a concentration of 1,165 mg/mL. Growth inhibition indicated the antibacterial activity of the tested compounds.

RESULTS

Isolation and antibacterial activity of microorganisms from samples of Krubera-Voronja Cave

In total, 874 heterotrophic cultures were isolated using five different media: TSA, HT agar, Al agar, SCNA, and ISP4. 108 isolates were from water samples, and 766 – from sediment samples collected in Krubera-Voronja Cave at the depth from 220 m
to 1640 m (Kieraite-Aleksandrova et al., 2015). The largest number of cultures was isolated on TSA (93 isolates at 4°C and 141 - at 30°C) and HT agar (207 isolates at 30°C). The smallest number of isolates was obtained using AI agar – 138 at 30°C. In general, temperature of 30°C was more suitable for the growth of heterotrophs – 714 cultures were obtained at this temperature, while only 160 cultures – at 4°C. Only TSA and ISP4 were suitable for growth at the latter temperature.

In order to examine antibacterial activity of the isolated cultures, primary screening using agar diffusion test was performed. Out of 874 cultures, 122 isolates (14.0%) demonstrated antibacterial activity. 108 active isolates were obtained from sediment samples, and 14 – from water samples. Most (63.9%) of the active isolates inhibited growth of a single test microorganism, while four isolates possessed a broad activity spectrum, being antagonistic against all test microorganisms. Only 9.3% of isolates from the upper part of the cave (sampling at 220-230 m in depth) showed antimicrobial activity, while 11.1% and 14.1% of all isolates from the middle (sampling at 500-700 m in depth) and lower (sampling at 1215-1640 m in depth) parts of Krubera-Voronja Cave, respectively, inhibited growth of at least one test strain.

Our previous cultivation-independent analysis showed that Gram-positive phyla Actinobacteria and Firmicutes were among the dominant phyla in Krubera-Voronja Cave (Kieraite-Aleksandrova et al., 2015). Therefore, we particularly focused on the isolates that exhibited antagonism only against Gram-positive but not against Gram-negative test microorganisms. In total, 31 isolates inhibited growth of two different test strains, and 24 of them were active only against Gram-positive ones. Activity of most of these 24 isolates against both Gram-positive test strains was very weak; somewhat stronger activity was observed only for seven cave isolates. Two cave isolates, 1350R2-TSA30-6 and 1410WF1-TSA30-2, were chosen for further experiments because of their high and comparable activity against both Gram-positive test strains – M. luteus from phylum Actinobacteria and B. thuringiensis TL8 from phylum Firmicutes. Both 1350R2-TSA30-6 and 1410WF1-TSA30-2 were isolated on TSA at 30°C from samples 1350R2 and 1410WF1 that have been collected in the lower part (1350 m and 1410 m depth) of Krubera-Voronja Cave. Sample 1350R2 was the sample of white clay from the rarely visited branch of the cave, and sample 1410WF1 was the sample of water from the underground camp (Kieraite-Aleksandrova et al., 2015).

Genotyping and phylogenetic analysis of active isolates

In order to determine whether isolates 1350R2-TSA30-6 and 1410WF1-TSA30-2 represent two different strains, BOX-PCR genotyping was performed. Our analysis clearly showed that these two isolates belong to two different strains (Fig. 1).

16S rRNA genes of these two strains were cloned, sequenced and analysed. The BLAST search showed that both strains belong to the family Bacillaceae in phylum Firmicutes. Both strains were assigned to the genus Bacillus (Fig. 2). 16S rRNA gene of the strain 1410WF1-TSA30-2 was most similar to those of Bacillus aerophilus, Bacillus altitudinis and Bacillus stratosphericus with 99.9% sequence similarity. Strain 1350R2-TSA30-6 clustered with Bacillus simplex, Bacillus frigoritolerans and Bacillus muralis with 99.8%, 99.6% and 99.5% sequence similarity respectively.

The effect of culture growth parameters for the production of antibacterial compounds

Optimization experiments showed that growth of a culture as well as production of antibacterial compounds depended on the concentration of nutrients in a culture medium. For the strain 1350R2-TSA30-6, the highest optical density (OD600) as well as the largest antimicrobial activity were observed in 0.5xTSB (data for BHIB not shown) (Fig. 3). For the strain 1410WF1-TSA30-2, similar maximum OD600 and antibacterial activity were obtained in TSB and 0.5xTSB, but the growth rate was higher in 0.5xTSB (Fig. 3). BHIB was the worst choice for the strain 1410WF1-TSA30-2 in terms of both culture growth and antibacterial activity (data not shown). Therefore, for both strains further optimization experiments were carried out in 0.5xTSB.

Fig. 1. BOX-PCR electrophoretic profiles. M - GeneRuler 1 kb DNA Ladder (Thermo Fisher Scientific); 1 - 1350R2-TSA30-6; 2 - 1410WF1-TSA30-2. The results of this test indicate that isolates 1350R2-TSA30-6 and 1410WF1-TSA30-2 are distinct.
Both the growth of a culture and the production of antibacterial compounds were influenced by pH of a culture medium. The best pH for the strain 1350R2-TSA30-6 was pH 7, and for the strain 1410WF1-TSA30-2 it was pH 9. Aeration did not influence the results of our experiments; highly similar results were obtained for both 180 rpm and 250 rpm. Low (16°C) temperature was not suitable for the growth of both strains – the strain 1410WF1-TSA30-2 did not grow, and OD600 for the strain 1350R2-TSA30-6 at this temperature was ~3.3 times lower than at 30°C. Consequently, the growth in the liquid culture at 8°C and 4°C was not tested.

Based on these results, both strains were further cultivated in buffered 0.5xTSB at 30°C at 180 rpm. pH of the medium was adjusted to 7 for cultivation of the strain 1350R2-TSA30-6 and to 9 for cultivation of the strain 1410WF1-TSA30-2. Growth curves and antimicrobial activity, obtained after the optimization experiments, are shown in Fig. 3. It is noteworthy that for both strains the production of antimicrobial compounds started in the exponential growth phase and continued during the growth of the culture.

**Extraction of antimicrobial compounds**

Two different approaches were used for the extraction of antimicrobial compounds from the culture supernatant – salting-out with ammonium sulphate and extraction by organic solvents.
Salted-out extracellular proteins of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 showed antibacterial activity against *M. luteus* (data not shown). The treatment with proteinase K did not diminish this activity. These experiments allowed us to suggest that antimicrobial compounds produced by both strains are not peptides or proteins.

Acetone, chloroform, ethyl acetate, methanol, and hexane were used separately for extraction of the different aqueous samples of the same culture by organic solvents. All extractions were successful; different extracts of the strain 1350R2-TSA30-6 were pyrrolizidines (Table 2, compounds No. 8-12). Their total amount was 60.3%. For four peaks (RT 46.23, 49.58, 50.19, and 58.70 min, compounds No. 8-11 in Table 2), the highest match with the library data was for pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)-. Clearly, those peaks should represent four different compounds. We could suppose that pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)- corresponds to the compound with RT 50.19 min (Table 2, compound No. 10), as its mass spectrum shows the highest match with the library data. Other three peaks should correspond to the compounds with very similar structure, as their mass spectra are quite similar with a very intense fragment peak at m/z 154 that could be attributed to hexahydro pyrrolo[1,2-α]pyrazine-1,4-dione (molecular mass 154). When harsh electron ionization is applied, the compounds undergo fragmentation and their molecular ions are not intense. Two of the compounds (with RT 49.58 and 50.19 min – Table 2, compounds No. 9 and 10 respectively) have the same molecular ion with m/z 210 indicating that they could be isomers of pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)-. A compound eluting at 46.23 min (Table 2, compound No. 8) has a molecular ion of m/z 196 and probably could be identified as pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-isopropyl)-. A compound eluting at 58.70 min (Table 2, compound No. 11) has a low intensity molecular ion peak at m/z 343. Unfortunately, very few mass spectra of related compounds can be found in the library, thus more precise identification was not possible. It should be noted that pyrrolizidines pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)- and pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(phenylmethyl)- are semivolatile cyclic dipeptides also called cyclo-

### Table 1. Antimicrobial activity of organic solvents’ extracts. *Micrococcus luteus* was used as the test microorganism in this agar-well diffusion assay, the width of the inhibition zone is shown in millimeters.

| Strain         | Ethyl acetate | Acetone | Hexane | Chloroform | Methanol |
|----------------|---------------|---------|--------|------------|----------|
| 1350R2-TSA30-6 | 1.5           | 3.0     | 4.0    | 3.5        | 2.0      |
| 1410WF1-TSA30-2| 1.0           | 1.5     | 1.5    | 1.0        | 1.5      |

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**Characterization of antimicrobial compounds**

Antimicrobial compounds of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 were analysed using TLC and bioautography. Both strains showed two different fractions on TLC plates, but only a single fraction of each strain showed antimicrobial activity against Gram-positive bacteria *M. luteus* (Fig. 4). Retardation factor (*R*) of the compounds with antimicrobial activity differed. *R* for the compound of the strain 1350R2-TSA30-6 was 0.85, and for the strain 1410WF1-TSA30-2 it was 0.93. The active compounds were recovered from the TLC plates and analysed by GC-MS.

GC-MS was used to determine a chemical composition of the two active TLC fractions. Main components of the active fraction of the strain 1350R2-TSA30-6 were pyrrolizidines (Table 2, compounds No. 8-12). Their total amount was 60.3%. For four peaks (RT 46.23, 49.58, 50.19, and 58.70 min, compounds No. 8-11 in Table 2), the highest match with the library data was for pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)-. Clearly, those peaks should represent four different compounds. We could suppose that pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)- corresponds to the compound with RT 50.19 min (Table 2, compound No. 10), as its mass spectrum shows the highest match with the library data. Other three peaks should correspond to the compounds with very similar structure, as their mass spectra are quite similar with a very intense fragment peak at m/z 154 that could be attributed to hexahydro pyrrolo[1,2-α]pyrazine-1,4-dione (molecular mass 154). When harsh electron ionization is applied, the compounds undergo fragmentation and their molecular ions are not intense. Two of the compounds (with RT 49.58 and 50.19 min – Table 2, compounds No. 9 and 10 respectively) have the same molecular ion with m/z 210 indicating that they could be isomers of pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)-. A compound eluting at 46.23 min (Table 2, compound No. 8) has a molecular ion of m/z 196 and probably could be identified as pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-isopropyl)-. A compound eluting at 58.70 min (Table 2, compound No. 11) has a low intensity molecular ion peak at m/z 343. Unfortunately, very few mass spectra of related compounds can be found in the library, thus more precise identification was not possible. It should be noted that pyrrolizidines pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(2-methylpropyl)- and pyrrolo[1,2-α]pyrazine-1,4-dione, hexahydro-3-(phenylmethyl)- are semivolatile cyclic dipeptides also called cyclo-

**Fig. 3. Optimization of the nutrient concentrations.** Effect on the culture growth and production of antibacterial compounds is shown. The width of inhibition zones is expressed in millimeters and shown at the sampling points. “Optimal” is the growth curve and antimicrobial activity under optimal conditions: 0.5xTSB in 50 mM Tris-HCl, pH 7, 30°C, 250 rpm for both strains. "TSB" is the growth curve and antimicrobial activity under the following conditions: 0.5xTSB, pH 7 (unbuffered medium), 30°C, 250 rpm for both strains.

**Fig. 4.** Retardation factor (*R*) of the compounds with antimicrobial activity differed. *R* for the compound of the strain 1350R2-TSA30-6 was 0.85, and for the strain 1410WF1-TSA30-2 it was 0.93. The active compounds were recovered from the TLC plates and analysed by GC-MS.
TABLE 3

| Pathogen                          | The concentration of the antimicrobial compounds in ethyl acetate, mg/mL |
|----------------------------------|-------------------------------------------------------------------------|
| Bacillus cereus ATCC 10876       | 4.660 2.330 1.165 0.583 0.291 0.146                                    |
| Enterococcus faecalis ATCC 29212 | 10.0 7.0 5.0 4.0 2.0 0                                                |
| Listeria monocytogenes ATCC 7644 | 7.0 6.5 5.0 3.5 3.0 0                                                |
| Staphylococcus aureus subsp. aureus ATCC 25923 | 3.5 1.5 0.5 0.5 0 0                                                  |
| Rhodococcus equi ATCC 6939       | 12.0 8.0 5.5 5.0 3.0 0                                                |

DISCUSSION

For isolation of microorganisms from Krubera-Voronja Cave, general purpose media (TSA) as well as the growth media designed for the isolation and cultivation of actinobacteria (HT agar, AI agar, SCNA, ISP4) were used. The choice of the latter four media was based on the well-known antimicrobial activity of the members of phylum Actinobacteria (Rule & Cheeptham, 2013). Nevertheless, the percentage (14.0%) of Krubera-Voronja Cave isolates, that produce antimicrobial compounds, was quite low and comparable to some shallow caves: Backcountry Cave (6 m in depth), Spider Cave (44 m in depth), and Fort Stanton Cave (134 m in depth) – 21.1%, 19.8% and 8.6% respectively (Montano & Henderson, 2013). On the other hand, data for some other shallow caves show higher percentage of active isolates than of Krubera-Voronja Cave: 66% for the Left Hand Tunnel passage in Carlsbad Cavern (316 m in depth), 40% for Lechuguilla Cave (489 m in depth) (Montano & Henderson, 2013), 57% for Helmckens Falls Cave (Rule & Cheeptham, 2013), 73.9% for Magura Cave (56 m in depth) (Tomova et al., 2013). Our results also showed that the percentage of bacteria capable of antagonistic activity increased from the upper to the lower part of the cave. These results were in accordance with the findings of Montano & Henderson (2013) who revealed that the increasing depth influenced the antimicrobial activity of the cave isolates. To the best of our knowledge, our present study is the first report on the antimicrobial activity of bacterial isolates in Krubera-Voronja Cave.
The other focus of our research was the antibacterial activity targeting Gram-positive bacteria. It should be noted that Gram-positive pathogens are a major cause of nosocomial bacterial infections. Staphylococci, enterococci, streptococci, and *Clostridium difficile* are the most important species of clinical interest (Rossolini et al., 2014). Antibiotic resistance issues are common among Gram-positive pathogens, and the percentage of nosocomial infections caused by antibiotic-resistant Gram-positive bacteria is increasing. For example, in the United States, approximately 60% of staphylococcal infections are now caused by multidrug-resistant *S. aureus*, and these percentages continue to rise (Rice, 2006). Therefore, the development of novel antibiotics that would be active against Gram-positive pathogens is imperative. Only a small portion (24 out of 874) of bioactive isolates were antagonistic exclusively against Gram-positive bacteria, moreover – the bioactivity of the most of these 24 isolates was very weak. Two most active strains 1350R2-TSA30-6 and 1410WF1-TSA30-2, isolated from the lower part of the cave, were chosen for detailed characterization of the antimicrobials. It is of interest that these strains, although antagonistic against Gram-positive test strains *M. luteus* and *B. thuringiensis* TL8, were also Gram-positive - both strains were assigned to the genus Bacillus.

Bacteria of the family *Bacillaceae* are mainly known for their antimicrobial peptides – both ribosomally and nonribosomally synthesized peptide antibiotics. Nonribosomally synthesized antibiotic bacitracin A is produced by *Bacillus licheniformis*, lipopeptides surfactin, iturin, bacillomycin, and fengycin are produced by several strains of *Bacillus subtilis*. Well-known ribosomally synthesized peptides (*bacteriocins*) of *Bacillaceae* include subtilin and subtilosin A from *B. subtilis*; coagulin from *Bacillus coagulans*; bacthuricin F4, thuricin 17, entomocin 9, and tochinin from *B. thuringiensis*; cerecin 7 from *B. cereus*; bacillacin 490 from *B. licheniformis* (Sumi et al., 2015). On the other hand, bacilli also produce non-peptide antimicrobial compounds, such as bis (2-ethylhexyl) phthalate, synthesized by *Bacillus pumilus* (Moushumi Priya & Jayachandran, 2012); macrolide antibiotics macrolactins 1-3 produced by *Bacillus* sp. (Mondol et al., 2011); macrolides gageomacrolactins 1-3 produced by *B. subtilis* (Tareq et al., 2013). Therefore, production of both antimicrobial peptides and non-peptide antibiotics was expected for bioactive strains from Krubera-Voronja Cave.

The antimicrobials of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 differed significantly. The active compounds behaved differently in some organic solvents during extraction as well as had different *R* values in TLC. It was also noticed that antimicrobial substances of the strain 1350R2-TSA30-6 were stable, and the inhibition zone was visible constantly during the growth of the test microorganism in bioautography experiments, while antimicrobial substances of the strain 1410WF1-TSA30-2 inhibited the test microorganism only at the first stages of its growth, but disappeared through further cultivation due to evaporation of active compounds or their inactivation by the growing culture of *M. luteus* (data not shown). Activity of antimicrobials of the strain 1350R2-TSA30-6 was higher than of strain 1410WF1-TSA30-2. The common characteristics for both strains included poor production of antimicrobials in a highly rich growth medium BHIB as well as continued production of active compounds during the growth of the culture starting in the exponential growth phase. The first characteristic can be associated with the need to produce antimicrobials under starvation, but not under the nutrient-rich growth conditions (Montano & Henderson, 2013). The second feature (the time of production) indicated that antimicrobial compounds (or at least some of them) were primary metabolites (Sanchez & Demain, 2008).

Chemical compounds in the active TLC fractions of both strains differed markedly. The main antibacterial agents of the strain 1350R2-TSA30-6 were 2,5-diketopiperazine antibiotics. Most of 2,5-diketopiperazines are stable to proteolysis (Pérez-Picazo et al., 2009), and we have also observed this effect after treatment of the salted-out extracellular proteins of the strain 1350R2-TSA30-6 with proteinase K. Antitumor, antifungal, antiviral, and antibacterial activities of cyclic dipeptides are well-known (Yan et al., 2004; Pandey et al., 2010; de Carvalho & Abraham, 2012; Kumar et al., 2014). Cyclic dipeptides are usually produced by Gram-negative bacteria, but have been also isolated from Gram-positive. Among the genera of the family *Bacillaceae*, they were so far identified only in the genus *Bacillus* (de Carvalho & Abraham, 2012; Leyton et al., 2012). Because of their activity spectrum, cyclic dipeptides attract much attention as potential pharmaceuticals. Our results clearly showed that antibacterial fraction of the strain 1350R2-TSA30-6 was active against Gram-positive pathogenic bacteria *B. cereus*, *E. faecalis*, *L. monocytogenes*, *S. aureus* sp. *spp. aureus*, and *R. equi*. Cyclic dipeptides are secondary metabolites (Leyton et al., 2012), therefore, if they would be the only compounds with bioactivity, then the antimicrobial activity...
effect would appear only in the stationary phase of the culture growth. But antimicrobial activity was also detected in the exponential growth phase of the strain 1350R2-TSA30-6. We have concluded that other volatile compounds, identified in an active fraction of this strain, were responsible for this early effect. For example, antibacterial, antiviral, and antifungal activities were previously reported for 1,2-benzenedicarboxylic acid, diisooctyl ester (Rameshthangam & Ramasamy, 2007; Maruthupandian & Mohan, 2011) and 1,3-dimethyl-benzenze (El-Shouney et al., 2014). Some compounds (ethylbenzene, p-xylene) were previously identified in mixtures of bacterial volatiles (Tenorio-Salgado et al., 2013), while others (cyclohexanone) - in plant extracts (Kim et al., 2011) possessing antimicrobial activity.

GC-MS results for the strain 1410WF1-TSA30-2 markedly differed from the results for strain 1350R2-TSA30-6. The only common antimicrobial compound was 1,2-benzenedicarboxylic acid, diisooctyl ester. Cyclic dipeptides were not found, but a range of other compounds with different bioactivities were identified. Antibacterial, antiviral, antifungal, or antioxidant activities were previously reported for dibutyl phthalate (Maruthupandian & Mohan, 2011), nonadecane (Fernando et al., 2005), phenol, 2,4-bis(1,1-dimethylethyl)- (Varsha et al., 2015). Some compounds (hexadecane, octadecane) were previously identified in antimicrobial or nematicidal mixtures of bacterial volatiles (Gu et al., 2007; Karanja et al., 2010), while others (1-nonadecene, 2-methyl-ecosane) - in plant extracts with antibacterial activity (Boussaada et al., 2008). The main compounds in antibacterial mixture of volatiles of the strain 1410WF1-TSA30-2 were benzoic acid, octadecyl ester and 1,2-benzenedicarboxylic acid, bis(2-methylpropyl) ester. We could not find any information about putative antimicrobial activity of octadecyl benzoate; it was not listed among bacterial volatiles in the mVOC and SuperScent databases either. But the esters of 1,2-benzenedicarboxylic acid (bis(2-ethylhexyl) ester, bis(2-methylpropyl) ester, butyl-2-methylpropyl ester, butyl-2-ethylhexyl ester, diisooctyl ester, etc.) are frequently the main components of antibacterial and antifungal mixtures of volatiles from different sources: algae (Sivakumar, 2014), plants (Joshi et al., 2011), fungi (Nakalembbe & Kabasa, 2012), bacteria (El-Mehalawy et al., 2008). Actually these esters are believed to be responsible for antimicrobial activity of these mixtures of volatiles. Antiviral, antifungal, and antibacterial activities were demonstrated for 1,2-benzenedicarboxylic acid, bis(2-ethylhexyl) ester; while antimicrobial, antioxidant, and antitumor activities were shown for 1,2-benzenedicarboxylic acid, mono(2-ethylhexyl) ester (Kavitha et al., 2009; Karthikeyan et al., 2014). Antibacterial and antifungal activities were confirmed for 1,2-benzenedicarboxylic acid, bis(5-ethylheptyl) ester (Kavitha et al., 2009). 1,2-benzenedicarboxylic acid, bis(2-methylpropyl) ester, found in active antibacterial fraction of the strain 1410WF1-TSA30-2, had been also identified in a few other representatives of the genus Bacillus (Hao & Lu, 2007; Malliaiah & Muthamilan, 2015).

It is known that bacteria use volatile metabolites for inter- and intraspecific communications as well as in order to respond and adapt to environmental changes (Audrain et al., 2015). Microorganisms in caves form multispecies communities with complex interaction networks among their members, and bacterial volatile mixtures can participate in these interactions. For example, it had been previously shown that Bacillus sp. can start producing cyclic dipeptides when challenged by a co-culture of other Bacillus species (de Carvalho & Abraham, 2012). Our results showed that antimicrobial volatile mixtures of strains 1350R2-TSA30-6 and 1410WF1-TSA30-2 could be also involved in the competition and engagement with other bacterial species in Krubera-Voronja Cave. The strains of Gram-positive genera Paenibacillus and Brevibacillus (phylum Firmicutes, family Paenibacillaceae) were sensitive to active antimicrobial fractions of both strains isolated from the same cave. The main difference between volatiles of both investigated strains was in their antagonism against actinobacterial strain of the genus Streptomyces. It is interesting that antimicrobial compounds of the strain 1410WF1-TSA30-2 were antagonistic against bigger number of cave strains than antimicrobial compounds of the strain 1350R2-TSA30-6. This is contrary to the zero antimicrobial activity of volatiles of the strain 1410WF1-TSA30-2 against pathogenic bacteria. It was supposed that the primary role of volatiles of the strain 1410WF1-TSA30-2 is the competition with the indigenous Gram-positive microorganisms of the cave.

In conclusion, our results clearly demonstrated that while screening for novel bioactive compounds, examining the lesser-studied phyla, such as Firmicutes, rather than the commonly examined Actinobacteria is an important and promising approach. Cave actinobacteria are usually the main targets of bioactivity assays, and therefore the antimicrobial potential of other bacteria is underestimated. Detailed analysis of two Firmicutes strains, isolated from Krubera-Voronja Cave, revealed two different mixtures of compounds with antibacterial activity. The main antibacterial compounds of the strain 1350R2-TSA30-6 were pyrrolopyrazines, and the main antibacterial compound of the strain 1410WF1-TSA30-2 was bis(2-methylpropyl) ester of 1,2-benzenedicarboxylic acid. To the best of our knowledge, our study is the first report on chemical characterization of antimicrobial mixtures of Firmicutes isolated from caves.

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