Photolyases and Cryptochromes in UV-resistant Bacteria from High-altitude Andean Lakes†

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ABSTRACT

“High-altitude Andean Lakes” (HAAL) are pristine environments harboring poly-extremophilic microbes that show combined adaptations to physical and chemical stress such as large daily ambient thermal amplitude, extreme solar radiation levels, intense dryness, alkalinity, high concentrations of arsenic (up to 200 ppm) and dissolved salts. In this work, we compared the UV resistance profiles, pigment content and photoreactivation abilities of three UV-resistant bacteria isolated from distinct niches from HAALs, that is Acinetobacter sp. Ver3 (water, Lake Verde, 4400 m), Exiguobacterium sp. S17 (stromatolite, Lake Soompapa, 3570 m) and Nesterenkonia sp. Act20 (soil, Lake Soompapa, 3570 m). UV resistance ability of HAAL’s strains indicate a clear adaptation to high radiation exposure encountered in their original habitat, which can be explained by genetic and physiological mechanisms named as the UV-resistome. Thus, the UV-resistome depends on the expression of a diverse set of genes devoted to evading or repairing the damage it provoked direct or indirectly. As pigment extraction and photoreactive assays indicate the presence of photoactive molecules, we characterized more in detail proteins with homology to photolyases/cryptochromes members (CPF). Phylogenetic analyses, sequence comparison and 3D modeling with bona fide CPF members were used to prove the presence of functional domains and key residues in the novel proteins.

INTRODUCTION

The Central Andes region displays unexplored ecosystems of shallow lakes and salt flats at altitudes between 3000 and 6000 m called as “High-altitude Andean Lakes” (HAAL). These pristine environments are a rich source of poly-extremophilic microbes that shows combined adaptations to physical and chemical stress, that is large daily ambient thermal amplitude, extreme solar radiation levels, intense dryness, alkalinity, high concentrations of arsenic (up to 200 ppm) and dissolved salts (1,2). Of particular importance is the high solar irradiance in the HAAL (1,2) with values of UV-B flux at mid-day up to 10 W m⁻² and a monthly average of daily insolation of 6.6 kWh m⁻² d⁻¹; these parameters are among the highest in the world (3) (NASAOMI/Aura Online serverhttp://gdata1.sc.ives.gasc.nasa.gov). The radiation values on HAALs are 165% higher than the reported values for lakes at sea level (1,2,4–9), and the erythemal index extends above to 15, which is equivalent to 5.57 MED h⁻¹ and to ca. 1200 J m⁻² h⁻¹. The low latitude and high altitude at HAAL is responsible for a thin ozone layer and a clear sky that favor intense UV radiation, a pattern which is clearly observed in lakes such as the one on the summit of the Licancabur volcano (5917 m) and Laguna Blanca (4340 m) (Bolivia) (8,9).

According to the Comité Internationale de l’Eclairage (CIE), the ultraviolet part of the electromagnetic spectrum is subdivided into ultraviolet A (UV-A) (315–400 nm), UV-B (280–315 nm) and UV-C (<280 nm) (CIE, 2011). As solar UV-C spectrum is filtered by the atmospheric gases O₂, H₂O, CO₂ and mainly O₃, UV range affecting life on Earth falls only between 280 and 400 nm. The UV-A spectrum constitutes 95% of UV-spectrum that reaches the earth causing cell damage indirectly by causing photooxidation of compounds and generating reactive oxygen species (10,11). In turn, UV-B represents <5% of incident solar radiation; besides triggering oxidative damage, UV-B also is able to excite and ionize biologically relevant molecules, such as lipids, proteins and DNA, causing the formation of different types of photoproducts (Pho) (12). DNA damage produced by UV-B irradiation reproduced by dimerization or adduct formation between adjacent pyrimidines bases (Thymine or Cytosine) on one DNA strand. The dimerization products called as cyclobutane pyrimidine dimers (CPDs) constitute 70–80% of total photoproducts, while the adducts accounted for the rest and are termed into ultraviolet A (UV-A) (315–400 nm), UV-B (280–315 nm) and UV-C (<280 nm) (CIE, 2011). As solar UV-C spectrum is filtered by the atmospheric gases O₂, H₂O, CO₂ and mainly O₃, UV range affecting life on Earth falls only between 280 and 400 nm. The UV-A spectrum constitutes 95% of UV-spectrum that reaches the earth causing cell damage indirectly by causing photooxidation of compounds and generating reactive oxygen species (10,11). In turn, UV-B represents <5% of incident solar radiation; besides triggering oxidative damage, UV-B also is able to excite and ionize biologically relevant molecules, such as lipids, proteins and DNA, causing the formation of different types of photoproducts (Pho) (12). DNA damage produced by UV-B irradiation reproduced by dimerization or adduct formation between adjacent pyrimidines bases (Thymine or Cytosine) on one DNA strand. The dimerization products called as cyclobutane pyrimidine dimers (CPDs) constitute 70–80% of total photoproducts, while the adducts accounted for the rest and are termed.

The World Meteorological Organization has established one UV Index Unit at 25 mW m⁻² or 90 J m⁻² h⁻¹. The erythemal dose Minim (MED) is defined as the unit of weighted UV-B radiant energy that produces a barely perceptible erythema on the skin (clear skin) and is equivalent to 210 J m⁻². 1 MED/h is defined as the ratio (irradiance itself) with which UV-B radiation affects some surface.
6-4[pyrimidine-2'-one] pyrimidines or (6-4)photoproducts (6-4 PPs) (12,13). Photoproducts (6-4) can be transformed to “Dewar photoisomers” through UV-A, UV-B and UV-C radiation. The biological effects are very variable depending on the type and extent of mutation produced.

Indigenous microorganisms originally isolated from irradiated environment are in general well adapted to high UV exposure (14–20). This is also the case for HAAL’s strains that display intrinsically high UV resistance and are now considered novel models for studying adaptive responses and mechanisms of light sensing and UV-triggered mechanisms (1,2,21). Genes encoding photoreceptors were identified in these strains, and a CPD class I photolyase from Acinetobacter sp. Ver3 was functionally characterized (22). These results call for a deeper screening in strains from HAAL and request a more detailed characterization of HAAL’s novel bacterial photoreceptors that already were coined extremo-enzymes.

Among many mechanisms that bacteria have developed to counter high UV radiation, DNA repair systems are essential as they enable cells to cope with fatal DNA damage. These mechanisms are usually classified in dark repair (DR) and photorepair (PR) (23,24). This last mechanism—also called photoreactivation—allows cells to repair UV-induced damage in their DNA upon adequate light irradiation for a certain time by using an enzyme called photolyase activated by light in the UV-A and blue-light range (25).

Photolyases (Phr) together with the structurally closely related cryptochromes (Cry) form a divergent family of photoactive proteins present in all three biological domains of life, called as cryptochrome/photolyase family (CPF). This includes monomeric flavoproteins of 53–66 kDa, containing between 450 and 620 amino acids, and a noncovalently bound flavin adenine dinucleotide (FAD) as cofactor in a 1:1 ratio. Some members of this protein family also can act as an antenna pigment, such as deazaflavin, lumazin or methenylethyadruyrofolute derivatives (26–28). Cry and Phr proteins share a large percentage of sequence identity, and their 3D-structures are very similar, probably due to their origin from a common ancestor (29).

Photolyases are classified according to the type of repaired photoproduct, that is cyclobutane pyrimidine dimer (CPD-Phr) photolyases or (6-4) pyrimidine–pyrimidine photolyases (6-4-Phr). Cryptochromes (Cry) have no photolyase activity and function as signaling molecules regulating diverse biological responses such as entrainment of circadian rhythms in plants and animals (30,31). According to Oberpichler 2011 (32), seven groups can be distinguished within the CPF: CPD class I; CPD class II; CPD class III; (6-4) photolyase and animal Cry; plant Cry; Cry-DASH; FeS-BCP (including the minor clusterCry-Pro). The (6-4) photolyases are very similar to animal Cry; the Plant Cry is the sister group of CPD class III, which is more often found in bacteria. Recently, Cry-DASH was described to be able to bind flavin adenine dinucleotide that can be photoreduced by blue light; also, CryDASH has been reported to binds single-stranded DNA with very high affinity (Kd \( \approx 10^{-8} \) M), and double-stranded DNA and single-stranded RNA with far lower affinity (Kd \( \approx 10^{-6} \) M). Cry-DASH was found to repair CPDs specifically in single-stranded DNA (ssDNA) (33,34), so these proteins should be coined photolyases. Other studies showed that the CPF must be expanded by an additional class called as FeS bacterial cryptochromes and photolyases (FeS-BCPs), in which members carry a 4Fe4S cluster (32).

In this work, we compare the UV resistance profiles, photoreactivation abilities and pigment content of three UV-resistant bacteria originally isolated from distinct niches from HAALs, that is Acinetobacter sp. Ver3 (22,35) (water, Lake Verde; 4400 m), Exiguobacterium sp. S17 (stromatolite, Lake Socoma, 3570 m) (22) and Nesterenkonia sp. Act20 (21) (soil, Lake Socoma, 3570 m) for which genomes sequences are available (22,36,37). Considering the high UV resistance phenotype of HAAL’s strains, these genomes were screened for the presence and diversity of sequences with homology to CPFs members. In addition, 3D modeling and a phylogenetic comparison with other known bacterial CPF sequences were performed.

**MATERIALS AND METHODS**

**Strains and culture conditions.** UV-resistant strains used in this study were previously isolated from different HAALs at the Andean Puna in Argentina (1,2), belonging to the LIMLIA-PROBIMI Extremophlic Strain Collection.

*Acinetobacter* sp. Ver3 was isolated from the Andean Lake Verde (4400 m), *Exiguobacterium* sp. S17 was isolated from a Lake Socoma stromatolite (3570 m) (17,35,36), and *Nesterenkonia* sp. Act20 was isolated from soil around Lake Socoma (3570 m) (21). Bacterial strains from DSMZ Bacterial Culture Collection were used as UV-sensitive controls that was previously reported (19,21,38) (Fig. 1), and they were Acinetobacter johnsonii DSM 6963, Acinetobacter baumannii DSM 30007, Acinetobacter lwoffi DSM 2463, Exiguobacterium auranticum DSM 6208 and Nesterenkonia halotolerans DSM 15474.

*Acinetobacter* spp. and *Exiguobacterium* spp. were grown in Luria–Bertani (LB) broth, supplemented with 2% agar for solid medium when applicable. *Nesterenkonia* strains were grown in “H” medium (a medium modified for halophiles, containing NaCl 15 g L \(-1\), KCl 3 g L \(-1\), MgSO\(_4\) 5 g L \(-1\), sodium citrate 3 g L \(-1\)) added with 2% agar for solid medium when applicable.

For electron microscopy, aliquots of a stationary state culture of each strain were fixed overnight at 4°C in Karnovsky’s fixative, comprising formaldehyde (8% v/v), glutaraldehyde (16% v/v) and phosphate-buffered saline (PBS; 0.2 M, pH 7.4). The fixed samples were washed three times with phosphate buffer for 10 min. Later, they were fixed with 2% v/v osmium tetroxide overnight. The samples were dehydrated successively with increasing alcohol concentrations (30%, 50%, 70%, 90%, and 100%) for 10 min each and finally maintained in acetone for 24 h. The final dehydration was carried out with the critical point technique. Samples were mounted on scanning electron microscopy sample stubs and gold coated. Specimens were observed under vacuum using a Zeiss Supra 55VP (CarlZeissNTS GmbH, Germany) scanning electron microscope.

**UV-B resistance assays.** For a quick and qualitative determination of level of resistance to different UV-B irradiation doses, the assays were performed on agar media. Briefly, cultures were collected at OD\(_{300}\) nm of 0.6 and subjected to serial dilutions. Aliquots of 5 µL were then loaded onto medium agar plates (the petri plates were covered with acetate film to block out UV-C and, once inoculated, were immediately exposed to UV-B lamps (Milbert Lourmat VL-4, maximum intensity at 312 nm) as light source (see lamp spectra, Appendix S1) covered by acetate sheet (see spectra, Appendix S1). As Gram-negative strains are much more UV-sensitive than Gram-positive strains (18,19,38), different duration and irradiation doses were applied depending on the strain. *Acinetobacter* strains were exposed to 15, 30, 60, 120, 180 and 240 min of UV-B irradiation at 2.98 W m \(-2\), while *Exiguobacterium* and *Nesterenkonia* strains were exposed to 5.4 W m \(-2\) during different times (0, 60, 90 and 120 min). Then, the strains were incubated in the dark to prevent photoreactivation, for 24, 48 and 72 h at 30°C. Microbial growth was recorded with three positive signs (+++ when it was similar to the growth in the control, with two positive signs (++) when it was slightly different from the growth in the control, with one positive sign (+) when the growth was too low to allow colony counting and with a negative sign (−) when no growth at all was observed. For plotting the data in a graphic, three, two or one sign was taken as 3, 2 or 1 unit of growth for each dilution and then summed up together to reach a unique value. Negative signs were counted as null unit of growth.

**Photoreactivation assays.** To evaluate the UV-B photorepair ability, selected strains were grown in LB medium at 30°C with shaking (2.3 g). Cells were harvested in the midexponential phase by centrifugation at 10 700 g for 20 min at 4°C. The cell pellets were washed twice in 0.9%
NaCl and were kept under starvation conditions in the same solution for 12 h at 4°C. A 20 mL portion of each cell suspension (at OD600 0.6) was transferred to 45 mL sterile quartz tubes, and these were covered with acetate film to block out UV-C and then exposed to UV-B radiation at an irradiance of 5.4 W m⁻², until microbial growth decreased to ca. 50% (CFUs) of the strain under study with shaking (50 rpm) at 15°C. Controls were incubated under the same conditions, but in the dark. After UV-B exposure, 100 µL of the cell culture was removed from the tubes and microbial growth was assessed upon growth in solid medium (CFUs) after 48 h of incubation at 30°C in the dark to prevent photoreactivation. Aliquots from UV-B exposed cell suspensions were subjected to photorepair (PR) or dark repair (DR) conditions. Photorepair was allowed by incubating the suspensions under photosynthetically active radiation (PAR) during 120 min (18 Wm⁻²), using OSRAM L18 W/77 lamps with continuous shaking (0.13 g) at 15°C, while dark repair was evaluated under the same experimental conditions but without PAR illumination. After each treatment, 100 µL of cells was removed from the tubes and the number of CFUs was determined after 48 h of incubation at 30°C under dark conditions to prevent photoreactivation. Incubation in the dark was accomplished by covering the quartz tubes or agar plates with aluminum foil.

Statistical analysis. Each experiment consisted of three biological replicas, error bars are the standard deviation (O.2). The means and the standards error of both, UV resistance profiles and photorepair ability assays, were assessed through Microsoft Excel statistical functions of three biological repetitions of each experiment.

Pigments extraction. Pellets of 150 mL of Act20, S17 and Ver3 cultures grown for 3 days were harvested through centrifugation at 10 700 g for 10 min, washed twice with physiological solution (PS), and centrifuged again. Pigments from the pellets were first extracted with aliquots of a mix of 2 mL methanol and 1 mL acetone (2:1), mixed with a vortex and left at rest for 2 min. Pellets were centrifuged at 10 700 g pm, and the supernatant was collected with a Pasteur pipette, filtered and transferred to a clean tube. The procedure was repeated twice. For the complete extraction of the pigments, the procedure was repeated with 2 mL acetone, 2 mL methanol and 2 mL hexane, one solvent at a time. The mixed solution was concentrated with nitrogen at low pressure and room temperature. Then, the sample was washed with 3 mL of PS and 3 mL of diethyl ether (DEE). The diethyl ether phase was collected and, in order to release carotenoids from conjugated groups (i.e. glycosides), a step of saponification was performed by adding 1:1 10% methanolic sodium hydroxide and incubation for 6 h protected from the light. The solution was dried with nitrogen as above and washed with PS and pigments re-extracted with DEE. The pigment absorption spectra were recorded with Elisa Reader spectrophotometer (Multiskan Go, Thermo Scientific) from 200 to 700 nm using DEE as blank all-trans-Astaxanthin, Canthaxanthin and β-Carotene (all of them from SIGMA).

Sequence comparison and phylogenetic analyses. Genomes of Acinetobacter sp. Ver3 and Exiguobacterium sp. S17 are available at the NCBI database under accession numbers GCA_000632455.1 and GCA_000411915.1, respectively (39,40). Using the BLAST tool and sequences of CPF family described in previous works, we found putative candidates encoding photoreceptors in the HAAL isolates (Ver3: Act20; S17). The sequences of the two photoreceptors found in Act20 genome (unpublished) were individually uploaded in Genbank (41), with accession numbers MG800787 and MG800788. RAST annotated genomes for all three strains were inspected searching for genes related to the UV-resistome (see Appendix S2).

Figure 1. (A) Geographic position of the High-altitude Andean Lakes in the Central Andes, South America. (B–D) Map localization and landscapes of L. Socompa and L. Verde. Macroscopic and microscopic view of HAAL’s strains: (E) Acinetobacter sp. Ver3; (F) Exiguobacterium sp. S17; (G) Nesterenkonia sp. Act20.
Phylogenetic tree was constructed using 45 bacterial sequences of the CPF family retrieved from the NCBI and Uniprot-KB databases in which experimental work proved their function as photolyase or cryptochrome (Table S1). A larger phylogenetic tree was constructed using 186 CPF homologues bacterial sequences, including the small dataset of known functions of the first tree together with sequences with high identity retrieved from the NCBI and Uniprot-KB databases (Table S2). Multiple alignment was performed with ClustalX, and main tree was generated with neighbor-joining (NJ) method using MEGA7 program (36) and the larger phylogenetic tree using FastME 2.0 (42), with 1000 bootstrap replicates. Subsequent processing and visualization were performed using iTOL (http://itol.embl.de/). Eight sequences were selected from the main set in order to trace the evolution of the family using Maximum Likelihood (ML) and Maximum Parsimony (MP). All sequences used in both analyses have been functionally and/or phylogenetically characterized in previous works; for six of them, crystal structures are available. Two trees were generated through the MEGA7 program (33): the first one was made with the ML method, with 100 bootstrap replicates, and the second tree was obtained with the MP method again with 100 bootstrap replicates. The alignment of the protein sequences and key amino acids was analyzed using the Jalview 2.9.0b2 software. The levels of amino acid conservation in photoreceptors were drawn using WebLogo (http://weblogo.berkeley.edu/logo.cgi).

Three-dimensional modeling. A comparative modeling technique was used to obtain three-dimensional models of HAAL’s extremophotoreceptors. First, PSI-BLAST was used to search the database for modeling templates (43) in the Protein Data Bank (PDB, www.pdb.org). The structures employed as models are listed in Table 2. The sequences were then aligned manually to adjust the models, and finally, 100 3D models were obtained per protein, using the software Modeller 9.14 (44). The models with the lowest score were selected according to DOPE score (Discrete Optimized Protein Energy), which is used to evaluate the energy of the model generated by Modeller 9.14. This selection identifies the lower DOPE score with lower free energy associated with the model and is taken as a criterion for the best model for the protein of interest.

Energetic and structural analyses were carried out to assay the quality of the chosen models. In the beginning, we obtained the DOPE profile (with modeler by Phytont) for the sequence of amino acids in the 3D structure, both for the model and for the template, for each CPF member. Then, we generated a graph with the two DOPE profiles for the selected model and the template employed in order to determine the similarity between model and template. The Ramachandran graphics were also generated in the MolProbity server (45), which provides an easy way to evaluate the distribution of the torsion angles of a protein structure, and also provides an overview of the allowed and not allowed regions, which serves as an important indicator of the quality of the three-dimensional structure of a protein. Also, the Z-scores and the energy were determined according to the position in the sequence of the models through the ProSA server (46), which allows determination of the quality of the general model and comparison of its score with the score of proteins crystallized by X-rays or NMR of similar sizes. The energy analysis along the sequence allows detecting regions with positive energy that are usually problematic or carry errors.

RESULTS AND DISCUSSION

UV-B resistance profile

UV resistance profiles of the studied strains and their respective controls indicated that HAAL’s poly-extremophiles were much more resistant to radiation than phylogenetically related bacteria used as control strains (Fig. 2A–C). Acinetobacter sp. Ver3 (herein, Ver3) was the most tolerant strain within the Acinetobacter spp. strains tested, being able to grow up to a UV-B dose of 42 kJ m$^{-2}$ (240 min), while A. johnsonii DSM 6963 (AJ), A. baumannii DSM 30007 (AB) and A. lwoffi DSM 2463 (AL) showed a clear inhibition even at lower exposure times (Fig. 2A). These results agreed with previous findings for the Acinetobacter sp. Strains (19,38).

The poly-extremophile actinobacterium Nesterenkonia sp. Act20 (21) (herein, Act20) was able to endure higher UV-B irradiation (UV-B dose: 68 kJ m$^{-2}$) than Ver3 and Exiguobacterium sp. S17 (herein, S17). Act20 showed a relative survival of 70% after 68 kJ UV-B irradiation, whereas N. halotolerans DMS 15474 (NH) showed a relative survival of only 20% (Fig. 2C). S17 was able to grow up to the maximum dose tested (120 m, UV-B dose: 38.8 kJ m$^{-2}$), while E. aurantiacum (EA) was completely inhibited after 90 min of exposure (Fig. 2B).

The superior UV resistance ability of HAAL strains indicates a clear adaptation to high radiation exposure encountered in their original habitat (1), which otherwise have selected a multifaceted system of shared genetic and physiological mechanisms called as UV-resistome (22). Thus, the UV-resistome depends on the expression of a diverse set of genes devoted to evade or repair the damage provoked direct or indirectly (2). Following this assumption, we compared Ver3, Act20 and S17 gene determinants involved in the following subsystems: (1) UV avoidance and protection strategies; (2) Stress sensors with their corresponding response regulators; (3) Damage tolerance and oxidative stress response; and (4) DNA damage repair (Appendix S2).

From the three strains, Act20 genome showed the most diversified response to UV as all subsystems proposed were found; this in agreement with its higher resistance profile to UV.

From all the subsystems found, we herein characterize more in detail photoreactivation abilities (included in the subsystem of DNA damage repair) and pigment content (included in the subsystem of UV avoidance and protection strategies) in the selected HAAL strains.

Photorepair ability

To evaluate whether photoreactivation is involved in the UV resistance mechanism of HAAL’s strains, their photorepair ability was tested in comparison with the sensitive controls (Fig. 2D–F). Selected strains were first exposed to UV radiation: 6.4 kJ m$^{-2}$ for Ver3, 9.7 kJ m$^{-2}$ for S17 and 12.9 kJ m$^{-2}$ for Act20. In these conditions, Ver3 and AL demonstrated a relative survival of 40%, in contrast to AJ and AB that showed a relative survival of only 2% and 10%, respectively (Fig. 2D). Aliquots of the UV-B challenged cells were then subjected to photorepair (PR) or dark repair (DR). After PR treatment, all strains increased their survival substantially. Ver3 showed the highest photoreactivation ability among all Acinetobacter strains, reaching values of up to 80%, followed by AL(50%), AJ(20%) and AB (<10%).

The resistance/photorepair profile was quite different for S17 and EA; after UV-B treatment, S17 was able to grow until a 64%, and then, it recovered with values of 77% for DR and 87% for PR, while UV-B exposed EA showed a 7% survival without recovering its viability after both, DR or PR (Fig. 2E).

Act20 and NH showed a 52% and 40% of survival after UV-B exposure, respectively. When exposed to the repairing treatments, they completely recovered under PR and showed 67% (NH) and 79% (Act20) of viability after DR (Fig. 2F). This was in line with our expectations since both Nesterenkonia strains were isolated from arid environments with high UV exposure (21,47,48).

Irrespective of the strain tested, photorepair was the most effective treatment to increase survival after UV-B. This phenomenon may be explained by the synergic effect of photolyase activity together with all repair mechanisms independent of light.
activation such as NER (nucleotide excision repair). In fact, it was suggested that photolyases can participate in light-independent repair (49,50); *E. coli* and yeast photolyase activities enhanced dark survival via specific stimulation of excision repair (49,51). Sancar et al. (50) have shown that binding of photolyase to pyrimidine dimers enhances the recognition of dimers by the excision nuclease.

Similar UV resistance profiles were found for other microbes or microbial communities reported in the HAAL; Ordonez et al. (36) isolated and characterized HAAL’s UV-B–resistant bacteria from six Andean lakes belonging to proteobacteria, Cytophaga-Flavobacterium-Bacteroides group, Firmicutes, Actinobacteria and *Bacillus*. Fernandez Zenoff et al. (52) studied the UV-B resistance of the culturable bacterial community from high-altitude wetlands; among all strains, the actinobacterium A5 was able to survive doses of 3201 kJ m⁻² under an irradiance of 4.94 W m⁻². Flores et al. showed (17) that Andean lakes bacterial communities were well adapted to high UV-B exposure and in many cases UV-B even stimulated their growth.

**Pigment content**

Light may damage biological systems by causing the excitation of photosensitive molecules (i.e. a sensitizer) which could initiate potentially harmful photochemical reactions with the surrounding molecules to generate highly reactive free radicals (53). Among the resources that microorganisms display to protect themselves from light, carotenoids are effective photo-protective compounds widespread in prokaryotes and eukaryote organisms (54). Carotenoids are diversified pigments ranging from red to yellow and produced by a wide variety of bacteria, algae, fungi and plants. These poly-isoprenoid compounds can be divided into two main groups: (1) carotenes or hydrocarbon carotenoids, which are composed of carbon and hydrogen atoms, and (2) xanthophylls that are oxygenated hydrocarbon derivatives that contain at least one oxygen function such as hydroxyl, keto, epoxy, methoxy or carboxylic acid groups (55). The protective action of carotenoids against photodynamic effects is related to their range of light absorption and their antioxidant properties in both photosynthetic and nonphotosynthetic organisms (48). Thus, in order to determine the involvement of carotenoids on the UV-B resistance profiles of the tested strains, pigment extractions were performed followed by a spectrometric analysis (Fig. 3).

The extract of *Acinetobacter* sp. Ver3 did not present peaks of absorption in the range of the visible light indicating the lack of pigments, while the smallest peak at 280 nm may be due to the absorption of the solvent used (Fig. 3). This evidence is in line with the whitish, almost transparent aspect of Ver3 cells and with the lack of carotenoids-coding genes in Ver3 genome (Appendix S2). Therefore, we suggest that pigment protection and/or ROS quenching by pigments are not involved in the high UV-B resistance of Ver3. In turn, both the yellowish extract of *Nesterenkonia* sp. Act20 and the orange extract of *Exiguobacterium* sp. S17 show a carotenoid-like absorbance spectrum (48), with maximum absorbance at 434 and 466 nm, respectively. For comparison, we have included three different pigment spectra; spectra from HAAL strains are more similar to the B-carotene control. Accordingly, genes involved in carotene biosynthesis were found in Act20 and S17 genomes (Appendix S2). Previous
works indicated that *Exiguobacterium aurantiacum* FH, a strain isolated from air, produced carotenoids (carotenes and xanthophylls) that displayed antifungal activity (56). In turn, this is the first report of carotenoid pigments produced by a *Nesterenkonia* strain.

Interestingly, S17 shows a clear absorbance in the 275–300 range, with peaks of significant intensity at 285 and 277 nm with a shoulder at 300 nm (Fig. 3). These peaks could be explained as a second pigment able to absorb light in the UV-B range, that is such as the carotenoid 15-cis-4,4'-diapophytoene of *Staphylococcus aureus* (51). An alternative explanation is that the peak at 287 nm is due to the absorbance of an intermediate compound of the synthesis of the carotenoid absorbing at 466 nm; Tao and Cheng studied the synthesis of a canthaxanthin from *Rhodococcus erythropolis* (49) and shows that the intermediate compound lacking the 4-keto group, the phytoene backbone, has a maximum absorption peak at 286 nm, while the insertion of the keto group to form 4-keto-carotene did not absorb at 286 but it maximum at 450 nm. The function of carotenoids in biological systems and their contribution to light stress has been largely studied (57). Carotenoids are considered to protect microorganisms, animals and plants from the destructive effects of activated oxygen species, such as superoxide (O_2^−), hydrogen peroxide (H_2O_2), singlet oxygen (1O_{2}) and hydroxy radical (OH), and are known to quench 1O_{2} (58). Many works reported positive effects of carotenoids and related them with UV-B protection in bacteria (55,59–61). It was mentioned that UV-B may have an indirect damage effect by promoting the formation of reactive oxygen species (ROS), which are highly toxic; likewise, UV-B can directly affect biomolecules like lipids, proteins and DNA by photolysis (62). The ROS plays the double role of being the inevitable by-product of aerobic metabolism on one hand and serving as a marker during stressful conditions on the other hand. The ROS defense mechanism consists of the antioxidant machinery which helps to mitigate the above mentioned oxidative stress-induced damages. The antioxidant machinery has two arms with the enzymatic components and nonenzymatic antioxidants. The

Figure 3. Absorption spectra of pigments, *Acinetobacter* sp. Ver3 (blue); *Exiguobacterium* sp. S17 (red); *Nesterenkonia* sp. Act20 (green).
machinery enzymatic includes superoxide dismutase (SOD), catalase (CAT), ascorbate peroxidase (APX), monodehydroascorbate reductase (MDHAR), dehydroascorbate reductase (DHAR), glutathione reductase (GR) and guaiacol peroxidase (GPX). While the nonenzymatic antioxidants comprising of acid ascorbic (AA), reduced glutathione (GSH), x-tocopherol, carotenoids, phenolics, flavonoids and amino acid cum osmolyte proline (62). Most of these proteins or metabolic pathways are presented in the annotated genome of Ver3, S17 and Act20 (Appendix S2). In addition, Acinetobacter sp. Ver3 demonstrated efficient ROS scavenging systems as a response to UV-B exposure (37,38). Further studies using functional genomics are in progress to characterize and integrate all the mechanisms of UV protection/ROS quenching in the HAAL’s strains.

Singh and Gabani (63) highlighted the importance of studying UV-resistant microbes and their metabolites as source for potential therapeutics. Some examples are as follows: (1) the mycosporine-like amino acids (MAAs) acting as a “sunscreen” for UV radiation in the wavelength range 310-365 nm (61) are found in a wide range of microorganisms including cyanobacteria and eukaryotic algae and protect against UV-induced DNA damage by preventing the formation of DNA dimers. They are already used as one of the compounds in UV sunscreens in the cosmetics industry and have been suggested for potential applications in the prevention of skin cancer induced by UV radiation such as melanoma (65). (2) Scytomonin is a secondary metabolite and an extracellular matrix (sheath) pigment synthesized by many strains of cyanobacteria (63). It is an aromatic indole alkaloid built from two identical condensation products of tryptophanyl- and tyrosyl-derived subunits linked through a carbon-carbon bond (64,65). It could act as a highly efficient protective biomolecule (sunscreen) that filters out damaging high-frequency UV rays, while at the same time allowing the transmittance of wavelengths necessary for photosynthesis (69). (3) Ectoine is a small organic molecule, present widely in aerobic, chemoheterotrophic and halophilic organisms. These organisms protect their biopolymers against dehydration caused by high solar exposure, high temperature, salt concentration and low water activity. Halophilic microorganisms synthesize ectoine from aspartate semialdehyde in three steps. Ectoine is already employed as protectant for human skin, in a number of different ways to avoid UV-A-induced cell damage; in fact, Lentzen and Schwarz in 2006 (70) could produce ectoine by a continuous fermentation of Acinetobacter sp. AN1 (WP_036475602), and (2) Act20Cry, a putative cryptochrome FeS-BCP with identity of 84% with a CPF of Exiguobacterium pavilionensis (WP_021066163); and (3) S17CryD, a putative Cry-DASH with 79% identity with a Cry-DASH from Exiguobacterium alkaliphilum (WP_034817977). Cry-DASH is a subgroup of the Cry gene family, which name indicated the taxa in which it was originally reported, that is Drosophila, Arabidopsis, Synchocystis and Homo (77). Nevertheless, it was later described to be present in nonphotosynthetic bacteria, fungi, plants and animals, including Neurospora, zebrish and Xenopus. Both structural and functional studies suggest that Cry-DASH proteins function as transcriptional repressors in Arabidopsis and Synchocystis (78,79). However, Selby and Sancar (2006) have demonstrated that the Arabidopsis At-Cry3 and other Cry-DASH exhibited single-stranded DNA photolyase activity.

In turn, Nesterenkonia sp. Act20 presents two sequence homologues to CPF in its genome: (1) Act20Phr, a putative CPD-Class III photolyase with 83% identity with a CPD from Nesterenkonia sp. AN1 (WP_036475602), and (2) Act20Cry, a putative cryptochrome FeS-BCP with an identity of 75% with a CPF of Nesterenkonia sp. AN1 (WP_036477144). Up to now, only one crystal structure of the type CPD class III photolyases is available and was reported for Agrobacterium tumefaciens (80). Within the Nesterenkonia genus, no specific studies on photoreceptors were reported, and therefore, little is known about their biological mechanisms, which otherwise does allow these microbes to survive in their original hostile environment.

Genome screening for cryptochromes and photolyases

Considering the experimental proof of efficient photorepairing in HAAL’s strains, genomes were screened for the presence of sequences with homology to known photolyases/cryptochromes (Table 1). Each of the three genomes displays at least one classical photolyase and one cryptochrome clustering together with the so-called FeS-BCPs group. FeS-BCPs comprehend a newly characterized CPF members containing the amino acids necessary to bind the cofactors, and in addition, four conserved cysteine residues for the coordination of an iron-sulfur cluster (32).

Acinetobacter sp. Ver3 presents two CPF sequences: Ver3Phr, a photolyase class I showing 67% identity with the photolyase from Acinetobacter sp. VT 114 (WP_054581756) (73) and Ver3Cry, a putative cryptochrome displaying 95% identity with a cryptochrome from Acinetobacter sp. VT 511 (WP_048881868). The latter protein can be classified according to Geisselbrecht et al. (74) as member of the new clade Cry-Pro. These cryptochromes are usually present in proteobacteria and cyanobacteria. The cryptochrome B (CryB) from Rhodobacter sphaeroides has been characterized, and the crystal structure is already available. It is involved in the regulation of photosynthesis gene expression (74). The Cry-Pro were classified in a wider clade of proteins called as FeS-BCPs, with 350 sequences identified from bacterial organisms, and also including many human and plant pathogens such as V. cholerae and P. syringae (32). Zhang et al. (75) and Zadow et al. (76) also described photolyase (6-4) activity for a FeS-BCP sequence from A. tumefaciens and Rhodobacter sphaeroides.

The genome of Exiguobacterium sp. S17 revealed the presence of three proteins belonging to the CPF family: (1) S17Phr, a putative CPD Class I photolyase displaying 83% identity with other photolyases from Exiguobacterium mexicanum (WP_034777040); (2) S17Cry, a putative cryptochrome FeS-BCP with identity of 84% with a CPF of Exiguobacterium pavilionensis (WP_021066163); and (3) S17CryD, a putative Cry-DASH with 79% identity with a Cry-DASH from Exiguobacterium alkaliphilum (WP_034817977). Cry-DASH is a subgroup of the Cry gene family, which name indicated the taxa in which it was originally reported, that is Drosophila, Arabidopsis, Synchocystis and Homo (77). Nevertheless, it was later described to be present in nonphotosynthetic bacteria, fungi, plants and animals, including Neurospora, zebrish and Xenopus. Both structural and functional studies suggest that Cry-DASH proteins function as transcriptional repressors in Arabidopsis and Synchocystis (78,79). However, Selby and Sancar (2006) have demonstrated that the Arabidopsis At-Cry3 and other Cry-DASH exhibited single-stranded DNA photolyase activity.

Conserved domains and functional residues of photolyases and cryptochromes

Inspection of protein sequences of putative CPD photolyases Ver3Phr, S17Phr and Act20Phr showed two clear domains: pfm03441 FAD binding domain of DNA photolyase and pfm00875, specific of DNA photolyases, which binds a light harvesting cofactor. In Act20Phr, an even more conserved zone was identified within the domain pfm00875, a region cd00293 called Usp (universal stress protein family). Usp is a small cytoplasmic bacterial protein whose expression is enhanced when the cell is exposed to stress agents.
In turn, the three putative cryptochromes of the FeS-BCP clade Ver3Cry, S17Cry and Act20Cry showed a single conserved N-terminal domain: pfam04244 being member of the superfamily c804458 (deoxyribodipyrimidinid photolyase-related protein). S17CryD presents the typical Cry-DASH domain TIGR02765 together with the two domains already described as pfam03441 and pfam00875.

Multiple sequence alignments (MSA) among HAAL’s photolyases and cryptochromes in comparison with reference sequences (Figs. 4A and 5A) allow us to confirm the conserved regions and key residues involved in functional activity, that is FAD binding (Phr/Cry); it was also searched the specific amino acids such as W-triad (Phr) and specific residues for the union of other cofactors such as FeS cluster binding (Cry) which help us for the classification. A better way to evidence the levels of amino acid conservation is by drawing a sequence logo (seq-Logo) from an MSA (Figs. 4B and 5B). The produced cartoon displays statistically significant differences in position-specific symbol compositions among multiple sequence alignments (81). In addition, we obtained one-hundred models for each HAAL’s photoreceptor to assess structural similarity with homologue crystallized proteins. Models with the lowest DOPEs scores were selected for each photoreceptor (Fig. 6). The DOPE profile was calculated for both templates and models and represented in a plot as a fast way to compare their similarity. For all seven cases, the DOPE profile of the model shows the same pattern as the template. The Ramachandran plot also showed a good folding of the models (Appendices S3–S9). The Z-score and the energy profile per amino acid residue were determined as another way to evaluate the quality of the models, comparing them with crystallized structures (Table 2). For all models, the values of Z-score were similar to the template and always fitted between the maximum and minimum values.

The activity of proteins from the CPF family depends on the possibility of changing the oxidation state of the bound FAD, allowing a cycle reduced–oxidized–reduced, which is favored by the generation of an excited initial state caused by UV-A (320–400 nm) and blue light (400–500 nm). This oxidation state cycle generates the repair activity on the damaged DNA in the case of the photolyases, and for the cryptochromes, it is considered as the starting event for the signaling cascade (82). Thus, we analyzed in all HAAL’s photoreceptors the N-terminal region, responsible for the FAD binding in the MSA (Fig. 4) and 3D models (Figs. 6 and 7 and Appendices S3–S9). Overlays of the photolyases Ver3Phr, S17Phr and Act20Phr against templates show a good structural similarity that materializes in a congruent fold of the protein (Fig. 7 and Appendices S3, S5 and S8). The structures revealed an αβ domain in the N-terminal region and a second α helix domain in the C-terminal region, both connected by a long interdomain loop that envelopes the αβ domain as it is common in photolyase. The αβ domain showed a Rossman fold with five parallel β strands covered at its sides by α helices, which also is seen in other photolyases. The binding site for FAD in photolyase of E. coli (EcPhr) is composed by the fol-
in vivo (26). Furthermore, Tyr residues are also involved in ET in other photolyases: for example in the CPD class I photolyase from *Anacystis nidulans* (83), and in *Methanosarcina mazei* CPD class II photolyase, a Tyr residue is required for full photoreduction. In the *Xenopus laevis* (6–4) photolyase, the involvement of a Tyr residue in photoreduction was shown by electron paramagnetic resonance (84,85) (Weber 2002, Holub 2017). But in all cases studied, the tyrosine is coupled to tryptophans. The W-triad in Ver3Phr is W286, W363 and W310 (86); in S17Phr, it is W369, W346 and W293; in Act20Phr, it is W391, W368 and W312; and in S17CryD, it is W369, W345 and W292 (Figs. 4A and 7A–C). The 3D models confirm the presence of the W-triad, which aligned coincidently in all three Phr models (Fig. 7A).

In addition, the binding site for the second cofactor is characteristic in some photolyases, and the amino acids are merely conserved, as shown in the sequence alignment around the positions of N108, E109, C292 and K29 (Fig. 4A). The CPD class I of Ver3 is the most similar to the CPD of *E. coli* in the amino acids sequence binding a second cofactor. The second cofactor in *E. coli* photolyase is MTHF, and glutamic acid is key for MTHF binding (87,88).

For the FeS-BCPs, the region containing the four cysteines for the sulfur part of the cluster (4Fe4S) (74) is the most conserved part (Fig. 5B); in *Rhodobacter sphaeroides*, they were identified as C346, C434, C437 and C45069. In the MSA, these residues are conserved in Ver3Cry and S17Cry but lost in Act20Cry (Fig. 5A). In coincidence, when an overlay is performed for the FeS-BCPs putative sequences (Fig. 7D–F), the four Cysteines are conserved and coincident in their spatial distribution, with the only exception of Act20Cry, which shows at the corresponding positions two proline (C381, C470), one alanine (C476) and one aspartate residue (C473). It is accepted that the cysteines are necessary to coordinate the [4Fe4S] cluster of the large subunit of primases (PriL-CTD) in archaea and eukarya and in FeS-BCPs (32,74). The absence of the pattern may indicate a loss of [4Fe4S] cluster, a novel function and/or a different structural disposition of the Act20Cry. Nevertheless, there are some examples where the clusters are bound or established for other amino acids different than cysteines (89).

Broach and jarrett (2006) (90) studied biotin synthase, an enzyme that catalyzes the addition of sulfur to dethiobiotin that contains a cluster of 4Fe4S, and in this work, they detail that the cluster is stabilized by the three cysteines, and an arginine, this last one is fundamental for maintaining the activity of the proteins. In Zu (91), they analyze and cite more than one example of proteins called Rieske [2Fe2S] clusters, where the cluster is stabilized by two histidine residues. However, there is no evidence for a cluster [FeS] without the stabilizing four cysteines in FeS-BCP cryptochromes—they are strictly conserved in crystallographic structures and widely retained in the clade—suggesting a low chance to find a cluster [FeS] coordination in the Act20Cry product. Interestingly, phylogenetic analysis indicated that the Act20Cry protein clustered in the FeS-BCP closer to the root and quite distant from the other members including Ver3Cry, S17Cry and the readily characterized FeS-BCP from *A. tumefaciens* and *R. sphaeroides* (Fig. 8).

On the other hand, the function of the FeS cluster is not very clear yet; usually, the [4Fe4S] clusters are oxidized acting as sensor for reactive oxygen species. Then, oxidative damage of the cluster could trigger disordering or structural change of the C-terminal end and alter the DNA binding, which in turn can act as a photoreactive agent, in a similar way to that proposed to the W-triad (74). This hypothesis may be sustained as FeS-BCP from *Rhodobacter sphaeroides* showed [6-4] photolyase activity.
this kind of proteins had initially been thought to be absent in prokaryotes and in most eukaryotic organisms (75, 76). Photolyase-like genes with FeS clusters were confirmed in the two strains of HAAL analyzed in this article, suggesting a possible [6-4] photolyase activity for them. The formation of the [6-4] photoproduct depends on the dose of UV radiation and increases with increasing dose reaching up to 40% of the photoproduct fraction (28). It is likely that the acquisition of [6-4] photolyases by HAAL bacteria is a compensatory adaptation to the high dose of UV radiation present in their original environment. However, in HAAL’s Acinetobacter strains the most common photoproducts repaired were the cyclobutane pyrimidine dimers (CPDs) (19). For further characterizations, assessment of a repair function for (6-4) photoproducts will be needed in these strains. Other possible functions of these cryptic photoreceptors may be to stimulate the NER system or to act as regulators enhancing photoreactivation or stress recovery upon UV-B. In previous work, a FeS-BCP has been reported to act as a cryptochrome regulating the expression of the photosynthetic apparatus in *Rhodobacter sphaeroides* (74). Although the herein selected HAAL’s bacteria are not photosynthetic, the abundance and quality of sunlight available in the Andean lakes, as well as the molecular versatility of this family of proteins, provide the microorganisms harboring them an advantage in an extreme environment where energy and nutritional resources are scarce.

**Phylogeny of bacterial photolyases and cryptochromes**

The phylogenetic analysis was designed in order to compare the HAAL’s sequences with homologue bacterial proteins with documented functional or structural information (Fig. 8A). The sequences clustered into eight groups; six of them were previously described as CPD class I photolyases (type 8-HDF), CPD class I photolyases (type MTHF), CPD class II photolyases, CPD class III photolyases, Cry-DASH and FeS-BCP-cryptochromes (34). Furthermore, two additional groups were identified herein: They include the sequences of *Thermus thermophilus* HB27 and *Streptomyces griseus*, respectively. A similar outcome
was found when ca. 100 sequences from the database were added to the analysis (Appendix S10, Table S2).

The photolyase from *Thermus thermophilus* HB27 (92) resides at a debatable phylogenetic position. In some works, it is included in the CPD class I photolyases (74,80,93,94), while in others lies in different clusters (32,95,96). It is stable at temperatures over 60°C and presents flavin mononucleotide (FMN), a highly efficient (97) second chromophore that has not been found in other bacterial CPD photolyases. The genus *Thermus* sp. usually reaches its optimum living temperature between 65 and 72°C (98), making it an excellent source of extremozymes. The closest orthologous photolyase in the databases belongs to the genus *Meiothermus* sp., which is also a thermophile with an optimal temperature range between 50 and 65°C (99). However, there is no functional proof or analysis reported that this protein intervene in the photoreactivation process.

The photolyase of the genus *Streptomyces* sp., functionally characterized only in *Streptomyces griseus*, has been included in previous works as a member of CPD class I photolyases with an unidentified antenna cofactor, possibly derived from 8-HDF (8-hydroxy-7,8-didemethyl-5-deazariboavin) (100–102). The genus *Streptomyces* is known for its remarkable biological, metabolic and genetic complexity as they showed the most complex biology growing with branched mycelium, filaments with hyphae and reproducing aerial branches with chains of spores (103); in this way, a good part of its biological cycle is exposed to radiation. In addition, they are producers of the largest number of antibiotics, 8700 antibiotics estimated in 2002, compared to 2900 produced by the rest of the bacteria, and 4900 by fungi (104), as well as generators of a variety of enzymes for extracellular digestion and signaling molecules. In agreement, with its complex lifestyle and metabolism, they allocate large genomes (8-9 Mb, approximately), with the highest estimated number of genes (105). As members of the phylum Actinobacteria, *Streptomyces*

### Table 2. Z-score of the generated models and the corresponding templates.

| Strain | Photoreceptor | Models | Template |
|--------|---------------|--------|----------|
| Ver3   | Pfr CPD-I     | -10.02 | -11.47   | 1DNP     |
|       | Pfr FeS-BCP   | -10.17 | -10.88   | 4DIA     |
|       | CRY FeS-BCP   | -10.82 | -10.88   | 4DIA     |
| S17    | Pfr CPD-I     | -8.92  | -11.47   | 1DNP     |
|       | Pfr FeS-BCP   | -8.72  | -10.73   | 1NP7     |
|       | CRY FeS-BCP   | -8.97  | -11.59   | 4U63     |
| Act20  | Pfr CPD-III   | -8.97  | -10.42   | 5ZX5     |
|       | Pfr FeS-BCP   | -8.97  | -10.42   | 5ZX5     |

**Figure 6.** Models of the CPF members under study. (A) Ver3Pfr, (B) Ver3Cry, (C) S17Pfr, (D) S17CryD, (E) S17Cry, (F) Act20Pfr, (G) Act20Cry.
...display a large G-C content in this genome, mostly above 70% unlike other Gram-positive bacteria whose G-C content is usually <50%. It should be kept in mind that the higher the G-C content, the lower is the amount of total photoproduct, although an increase of photoproduct with cytosine content is observed, which is highly mutagenic compared to the photoproduct TT (53,56). These characteristics make their genomes more exposed to photo inducible mutations. Some of these traits are also found in other genera of Actinobacteria (102). It is probable that some of these factors or the combination of more than one have influenced the notable specialization of the enzyme photolyase resulting in a separated clade (Fig. 8A).

To trace the evolution of the family CPF, a phylogenetic analysis was performed using both Maximum Likelihood (ML) and Maximum Parsimony (MP) methods. They were plotted using cladograms (Fig. 8B,C) in order to identify the main steps of molecular evolution in the CPF family in bacteria. For our knowledge, this is the first analyses of CPF evolutionary process taking into account only the bacterial domain.

The main steps in the evolution of groups of bacterial photolyases are presented (Fig. 8B,C). The first photolyase described (step 1) would have resulted from the duplication of a gene with an [FeS] cluster that acquired the function of CPD repair (76). The gain of an antenna (step 2) in order to amplify the energy absorption would be the next step. This chromophore would probably have been flavin itself or a derivative (8-HDF, FMN, DMRL, FAD) for several reasons. Among others, one would consider its high photochemical efficiency, required in a primitive environment without a protective atmosphere against UV rays (100,102). The change (step 3) of the CPD function by [6-4] repair would result in the FeS-BCP class or photolyases and bacterial cryptochromes with iron–sulfur cluster. The early appearance of this function would increase the amount of photoreactivation, considering that the photoproduct [6-4] is abundant at high doses of UV irradiation (28), characteristic of the environment in which prokaryotes evolved. The loss of the cluster [FeS] (step 4) would have been the following step for the origin of the majority of the current members of the family. A long deletion (step 5) in the C-terminal α helix domain, which affects the FAD binding mechanism, may give rise to the CPD II photolyases (54,102). Photolyases CPD class III (step 6) would have originated from the recruitment of the pterinic chromophore (MTHF), which, although less efficient, is more abundant in nature (88,99,100). This class of photolyases includes an alternative photoreactive triad and the stabilization of the MTHF chromophore by π-stacking. Other recruitments of the MTHF, yet in a different and poorly conserved site would have occurred in folate CPD I (step 7) and some Cry-DASH proteins (step 8).

CONCLUDING REMARKS

In this work, we compared the UV resistance profiles, pigment content and photoreactivation abilities of three UV-resistant bacteria isolated from distinct niches from HAAL’s; out of these, the soil actinobacterium, Nesterenkonia sp. Act20, was found to be the most resistant extremophile.
UV resistance ability of HAAL’s strains indicate a clear adaptation to high radiation exposure encountered in their original habitat, which can be explained by genetic and physiological mechanisms called as UV-resistome. Thus, the UV-resistome depends on the expression of a diverse set of genes devoted to evading (pigments, wall components, biofilm formation) or repairing/coping the provoked damage (DNA repair/oxidative stress response) which are currently under further investigation by multi-omics approaches.

Two components of the UV-B resistome were further explored in this paper; pigment extraction indicates the presence of carotenoid-like compounds in S17 and Act20 cells suggesting an antioxidative defense or protective role for them. On the other hand, photoreactivation upon UV-B damage was efficient in all three strains; consequently, we found proteins with homology to photolyases/cryptochromes (CPF) in Ver3, Act20 and S17 genomes. Each one displays at least one classical photolyase and a cryptochrome clustering together with the so-called FeS-BCPs group. Phylogenetic analyses, sequence comparison and 3D modeling with bona fide CPFs were used to prove the presence of functional domains (FAD binding) and key residues (W-triad) in the novel proteins. The most intriguing protein was, however, Act20Cry for which the phylogenetic analyses indicate a close relation with BCPs carrying an iron–sulfur cluster. However, so far inspection of the sequence in detail and 3D homology models suggests no possible cluster coordination for this protein. Further overexpression, purification and spectroscopic/structural characterization of Act20Cry will be needed to solve this open question.

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SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section at the end of the article:
Appendix S1. UV-B lamp and acetate filter spectra.
Appendix S2. UV-resistome components found in HAAL strains.
Appendix S3. Modelling and quality evaluation of Ver3Phr.
Appendix S4. Modelling and quality evaluation of Ver3Cry.
Appendix S5. Modelling and quality evaluation of S17Phr.
Appendix S6. Modelling and quality evaluation of S17CryD.
Appendix S7. Modelling and quality evaluation of Act20Phr.
Appendix S8. Modelling and quality evaluation Act20Cry.
Appendix S9. Phylogenetic tree of Cryptochrome/Photolyase family members of the Bacteria domain generated using neighbor-joining methods.
Table S1. Table of the sequence used to build the main phylogeny tree.

Table S2. Table of the sequence used to build the second phylogeny tree.

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