Metagenomic Mining for Esterases in the Microbial Community of Los Rueldos Acid Mine Drainage Formation

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Acid mine drainage (AMD) systems are extremely acidic and are metal-rich formations inhabited by relatively low-complexity communities of acidophiles whose enzymes remain mostly uncharacterized. Indeed, enzymes from only a few AMD sites have been studied. The low number of available cultured representatives and genome sequences of acidophiles inhabiting AMDs makes it difficult to assess the potential of these environments for enzyme bioprospecting. In this study, using naïve and in silico metagenomic approaches, we retrieved 16 esterases from the α/β-hydrolase fold superfamily with the closest match from uncultured acidophilic Acidobacteria, Actinobacteria (Acidithrix, Acidimicrobium, and Ferrimicrobium), Acidiphilium, and other Proteobacteria inhabiting the Los Rueldos site, which is a unique AMD formation in northwestern Spain with a pH of ∼2. Within this set, only two polypeptides showed high homology (99.4%), while for the rest, the pairwise identities ranged between 4 and 44.9%, suggesting that the diversity of active polypeptides was dominated not by a particular type of protein or highly similar clusters of proteins, but by diverse non-redundant sequences. The enzymes exhibited amino acid sequence identities ranging from 39 to 99% relative to homologous proteins in public databases, including those from other AMDs, thus indicating the potential novelty of proteins associated with a specialized acidophilic community. Ten of the 16 hydrolases were successfully expressed in Escherichia coli. The pH for optimal activity ranged from 7.0 to 9.0, with the enzymes retaining 33–68% of their activities at pH 5.5, which was consistent with the relative frequencies of acid residues (from 54 to 67%). The enzymes were the most active at 30–65°C, retaining 20–61% of their activity under the thermal conditions characterizing Los Rueldos (13.8 ± 0.6°C). The analysis of the substrate specificity revealed the capacity of six hydrolases to efficiently degrade (up to 1,652 ± 75 U/g at pH 8.0 and 30°C) acrylic- and terephthalic-like [including bis(2-hydroxyethyl)-terephthalate,
INTRODUCTION

In biotechnology, there is high interest in finding enzymes with new or improved properties (Pellis et al., 2017; Ferrer et al., 2019). This interest is especially increased in relation to enzymes from extremophiles, which are microorganisms that have evolved to thrive in extreme environments (Baveja et al., 2016), as they can efficiently operate under multiple conditions requested by industry. One example application is the eco-friendly bioconversion of cellullosic biomass by extremozymes, which produces green products and has less substrate loss (Thapa et al., 2020). Furthermore, plastic disposal is one of the major problems currently faced by the environment, as enormous quantities of synthetic plastics are non-degradable. Researchers are constantly exploring new ways to degrade plastics, and one of these ways involves using enzymes from microorganisms or microbial communities, including some that inhabit extreme environments (Nchedo Ariole and George-West, 2020).

Acid mine drainage (AMD) systems deserve special attention as a source of extremozymes. AMDs are extremely acidic runoff formations that originate from the microbial oxidation of pyrite and other sulfide minerals, which results in the production of sulfuric acid and metal-rich solutions (Méndez-García et al., 2015; Johnson and Quatrini, 2020). AMD systems are common in our planet, although only a limited number of them have been microbiologically characterized (Méndez-García et al., 2015; Johnson and Quatrini, 2020). Although it has recently been demonstrated that some of these AMD formations, such as the Los Ruedos mercury mine in northwestern Spain (Méndez-García et al., 2014), appear to be populated by a great diversity of prokaryotes, the majority of them are inhabited by a restricted set of acidophilic bacteria and archaea (Dopson et al., 2004; Golyshina, 2011; Méndez-García et al., 2015; Chen et al., 2016; Johnson and Quatrini, 2020), whose variety and abundance depend on geochemical constraints (Méndez-García et al., 2014, 2015; Huang et al., 2016).

Major bacterial lineages detected in AMD systems include the phyla Proteobacteria (Acidithiobacillus, Acidiphilum, Acidocella, Acidocaldus, Acidomonas, Acidiphaga, “Ferrovum,” Acidibacter, and Metallibacterium spp.), Nitrospira (Leptospirillum spp. such as Leptospirillum ferrooxidans, Leptospirillum ferriphilum, and “Leptospirillum ferrodiazotrophum”), Actinobacteria, Firmicutes (Sulfohalobacillus spp., and Alicyclobacillus spp.), Acidobacteria, Saccharibacteria (TM7) and other candidate phyla radiation (CPR) organisms. Archaea include the phyla Eurarcheaeota (Ferroplasma spp. such as Ferroplasma acidiphilum and “Ferroplasma acidarmanus,” Acidiplasma cupricumulans, and Unicellularplasma divulgatum), Thaumarchaeota, and the Candidate divisions “Micrarchaeota” and “Parvarchaeota” (Golyshina et al., 2000, 2009, 2016; Dopson et al., 2004; Baker et al., 2006, 2010; Golyshina, 2011; Chen et al., 2016, 2018; Gavrilo et al., 2019; Korzhenkov et al., 2019). These microorganisms are expected to be reservoirs of enzymes selected to resist acidic harsh conditions (at least regarding extracellular products) (Sharma et al., 2012), some of which might be of biotechnological relevance (Gomes et al., 2003; Adrio and Demain, 2014).

In this category, esterases and lipases from the α/β-hydrolase fold superfamily are appropriate biocatalysts for use in a modern circular bioeconomy because of their abundance (at least one per genome; Ferrer et al., 2015); the extensive knowledge that has been accumulated after the analysis of the biochemical features, sequences, and structures of more than 280,638 such proteins (Bauer et al., 2020); their ease of identification (multiple available screening methods; Reyes-Duarte et al., 2012); and their outstanding properties in terms of stability, reactivity, and scalability, which make them third-choice tools for the functionalization and modification of low-reactivity hydrocarbon-like blocks, oils, and fats (Daiha et al., 2015). Genomics and metagenomics can potentially make accessible an enormous reserve of such important biocatalysts in organisms or microbial communities inhabiting any environment, including AMD systems. However, only 239 of the 280,638 sequences available at the Lipase Engineering (LED) Database (Bauer et al., 2020) have been retrieved from cultured microorganisms (listed above) and uncultured microorganisms that are inhabitants of AMD systems, including Alicyclobacillus spp., 118 in total; Sulfohalobacillus spp., 53; Acidobacteria, 34; Acidithiobacillus, 13; Leptospirillum, 9; “Ferrovum,” 5; Acidocella, 3; and Ferroplasma, Aciditrix, Acidiphilum, and Metallibacterium, with 1 each. Among these biocatalysts, only a low-pH optimum carboxylesterase from F. acidiphilum (Ohara et al., 2014) has been characterized. This limits the assessment of the biotechnological potential of acidophiles living in AMD systems, at least regarding esterases and lipases. The minimal enzyme-level information that is known about these systems is restricted to two endo-acting amylases with no similarity to any known protein and two genes conferring metal and acid resistance from the microbial community inhabiting the AMD systems of the Carnoulès (lead–zinc) mine in France (Delavat et al., 2012) and the Tinto River in southwestern Spain (Delavat et al., 2012; Guazzaroni et al., 2013), respectively.

Keywords: acidophiles, acidophilic bacteria, acid mine drainage, biodiversity, extremozymes, esterase, metagenomics, plastic
To fill this knowledge gap, we initiated a metagenomic investigation to isolate carboxylesterases from a recently discovered and microbiologically characterized AMD formation, namely, the Los Rueldos mercury mine in northwestern Spain (Méndez-García et al., 2014). By applying homology searches in metagenomic sequences and naïve screening in clone libraries with enzyme substrates, we discovered a number of such enzymes whose characteristics are reported herein. Both function- and DNA sequence-based metagenomic methods are complementary, with each having advantages and disadvantages. Bioinformatics methods allow a rapid process of enzyme searching. However, in prokaryotic genomes, >30% of genes remain annotated as “hypothetical, conserved hypothetical or with general prediction,” and large numbers of genes may have non-specific annotations (such as putative hydrolases). The analysis of biochemical functions is likely to provide a superior approach to avoid this limitation, especially when screening novel enzymes. However, only a few hundred specific enzymatic assays exist, with a limited number of them applied in a high-throughput manner for the naïve screening of metagenomics libraries.

Although the in vivo roles and expression levels of the genes encoding the hydrolases presented in this study are unknown, their sequences and results of biochemical analyses shed new light on the enzymology of the microbial inhabitants of the Los Rueldos AMD formation, which have been neglected in enzyme prospecting.

MATERIALS AND METHODS

General Experimental Procedures

The source and brand of each of the esters (purity ≥99%) used in this study was Merck Life Science S.L.U., Madrid, Spain. The oligonucleotides used for DNA amplification were synthesized by Sigma Genosys, Ltd. (Pampisford, Cambridgeshire, United Kingdom). The Escherichia coli EPI-300-T1R strain used for pCCFOS1 fosmid library construction and screening was from Epicentre Biotechnologies (Madison, WI, United States). The E. coli strain GigaSingles used for gene cloning and E. coli strain BL21 (DE3) used for gene expression were from Novagen (Darmstadt, Germany).

Sampling Site and Sample Collection

The Los Rueldos gallery is located along the northwestern slope of the Morgao Valley (2 km northeast of the town of Mieres and 20 km southeast of Oviedo, which is the capital city of Asturias in northwestern Spain; 43°15′47″N, 5°46′9″W). It is a 70 m-long gallery with 10–30 cm depths in the shallower areas and 40–70 cm depths in the deeper sections (Méndez-García et al., 2014). Microorganisms are developed along the AMD system (pH ~2), forming a bedded acidic biofilm with uppermost oxic (B1A) and lowermost anoxic (B1B) strata. The DNA samples from B1A and B1B (see below) samples collected and used in this study were the same as those in the previous work (Méndez-García et al., 2014). Briefly, samples were collected in sterile 50 ml tubes at two sampling sites determined by the presence of each different macroscopic microbial growth morphology (B1A: up to 2 cm deep; B1B: from 2 to 15 cm deep) and kept on ice until nucleic acid extraction was performed (within the following 2 h).

Nucleic Acid Extraction, Preparation of pCCFOS1 Libraries, and Naïve Screening

The DNA samples from B1A and B1B were the same as those used in a previous work (Méndez-García et al., 2014), which were obtained using the Power Soil DNA extraction kit (Cambio, Cambridge, United Kingdom) according to the manufacturer’s guidelines. Prior to clone library construction, the metagenomic DNA was concentrated by first adding 50 µl of 3 M sodium acetate solution to 50 µl DNA extract. Precipitation was conducted by the addition of 1.25 ml of ethanol and incubation at room temperature for 10 s. Precipitated DNA was pelleted by centrifugation at 20,000 g for 10 min. The resulting pellets were washed with 500 µl of 70% (v/v) ethanol twice, and the traces of ethanol were evaporated by incubation under a fume hood at room temperature for 10 min. The resulting pellets were then dissolved in 20 µl of sterile nuclease-free water. Before cloning in the large-insert pCCFOS1 fosmid libraries using the CopyControl Fosmid Library Kit (Epicentre Biotechnologies, Madison, WI, United States) and the E. coli EPI300-T1R strain, the DNA (10 µg) that was unsheared by gel electrophoresis was subjected to shearing by pipetting through a 200 µl pipette tip 100 times, following the recommendations of the supplier (Epicentre Biotechnologies, Madison, WI, United States) to reach an approximately size of 30,000 bp. Cells of each pCCFOS1 fosmid library were suspended in glycerol to a final concentration of 20% (v/v) and stored at −80°C until further use. We generated subsets of 94,000 and 81,000 clones for the B1A and B1B samples, respectively. Restriction analysis of 10 randomly selected clones from each library revealed average insert sizes of 34,000 bp (for the B1A samples) and 39,500 bp (for the B1B samples), which included nearly 3.2 Gbp of community genomes per sample. This size is within the range of the average size range of DNA inserts in positive clones found in this study (see below).

Fosmid clones were plated onto large (22.5 × 22.5 cm) Petri plates with Luria Bertani (LB) agar containing chloramphenicol (12.5 µg/ml) and induction solution (Epicentre Biotechnologies; WI, United States) at a quantity recommended by the supplier to induce a high fosmid copy number. Clones were scored by the ability to hydrolyze α-naphthyl acetate (α-NA) and tributyrin, as previously described (Reyes-Duarte et al., 2012). Positive clones presumptively containing carboxylesterases and lipases with the α/β hydrolase fold were selected, and their DNA inserts were sequenced using a MiSeq Sequencing System (Illumina, San Diego, CA, United States) with a 2 × 150-bp sequencing v2 kit at Lifesequencing S.L. (Valencia, Spain). Before sequencing, fosmid DNA was extracted from the fosmid clones containing the metagenomic segments using the QUIAGEN Large-Construct Kit (QUIAGEN, Hilden, Germany), according to the manufacturer’s protocol. Upon the completion of sequencing, the reads were quality-filtered and assembled to generate non-redundant meta-sequences, and
genes were predicted and annotated as described previously (Placido et al., 2015).

**Selection of Genes Encoding Enzymes by Homology Sequence Analysis**

The predicted protein-coding genes obtained in a previous study (Méndez-García et al., 2014) after the sequencing of DNA material from resident microbial communities in each of the samples (B1A and B1B) with a Roche 454 GS FLX Ti sequencer (Roche Applied Science, Penzberg, Germany) were used in this study. The meta-sequences are available from the National Center for Biotechnology Information (NCBI) non-redundant public database with the IDs PRJNA193663 (for B1A) and PRJNA193664 (for B1B). Protein-coding genes identified from metagenomes (sequence-based screening) and from the DNA inserts of positive clones (naïve screen) were screened (score >45; e-value <10e-3) using BLASTP and PSI-BLAST searching (Altschul et al., 1997) for enzymes of interest against the Esterases and alpha/beta-Hydrolase Enzymes and Relatives (ESTHER) and LED databases (Fischer and Pleiss, 2003; Barth et al., 2004; Bauer et al., 2020).

**Gene Expression and Protein Purification**

The experimental procedures used for the cloning, expression, and purification of selected proteins (either from naïve or homology sequence screening) in the Ek/LIC 46 vector and *E. coli* strain BL21 (DE3) were performed as described previously (Alcaide et al., 2015). The primers used for amplification are listed in Supplementary Material. All proteins studied here were N-terminally His6-tagged, and the soluble His-tagged proteins were produced and purified at room temperature after binding to a nickel–nitrilotriacetic acid (Ni–NTA) His-Bind resin (from Merck Life Science S.L.U., Madrid, Spain) as described previously (Giunta et al., 2020), with slight modifications (the expression culture was scaled up to 1 L using 50 ml pre-inoculum). The purity was assessed as >98% using sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS–PAGE; Supplementary Figure 1) in a Bio-Rad Mini Protein system (Laemmli, 1970). Protein concentrations were determined according to the Bradford method with bovine serum albumin as the standard (Bradford, 1976). A total of approximately 0.8–37 mg of purified recombinant proteins was obtained from each 1 L culture on average, as follows: Est_A1 (6.4 mg/L), Est_A2 (25 mg/L), Est_A3 (13 mg/L), Est_A4 (37 mg/L), Est_A5 (41 mg/L), Est_A6 (7 mg/L), Est_A7 (0.8 mg/L), Est_A8 (19 mg/L), Est_B1 (1.0 mg/L), and Est_B2 (32 mg/L).

**Enzyme Assays**

The hydrolysis of 2-naphthyl acrylate (ref. 577189), tri(propylene glycol) diacrylate (ref. 246832), dibenzyl terephthalate (ref. PH000126), and bis(2-hydroxyethyl)-terephthalate (BHET; ref. 465151) (all from Merck Life Science S.L.U., Madrid, Spain) was assessed using a pH indicator assay in 384-well plates (ref. 781162, Greiner Bio-One GmbH, Kremsmünster, Austria) at 30°C and pH 8.0 in a Synergy HT Multi-Mode Microplate Reader in continuous mode at 550 nm over 24 h [extinction coefficient (ε) of phenol red, 8,450 M⁻¹ cm⁻¹]. The acid produced after ester bond cleavage by the hydrolytic enzyme induced a color change in the pH indicator that was measured spectrophotometrically at 550 nm. The experimental conditions were as detailed previously (Giunta et al., 2020), with the absence of activity defined as at least a twofold background signal. For $V_{\text{max}}$ determination, (protein): 270 μg/ml; (ester): 20 mM; reaction volume: 44 μl; T: 30°C; and pH: 8.0. Activity was calculated by determining the absorbance per minute from the generated slopes and applying the following equation:

$$\text{Rate} = \frac{\Delta \text{Abs}_{\text{min}}}{8450 \text{M}^{-1} \text{cm}^{-1}} \cdot \frac{1}{0.4 \text{ cm}} \cdot \frac{10^6 \mu \text{M}}{1 \text{ M}} \cdot \frac{0.000044 \text{ L}}{1 \text{ mg prot.}}$$

The activity toward the model esters $p$-nitrophenyl acetate (pNPC2), propionate (pNPC3), butyrate (pNPC4), octanoate (pNPC8), decanoate (pNPC10), and decanoate (pNPC12) was assessed in 50 mM Britton and Robinson (BR) buffer at pH 8.0 and 30°C by monitoring the production of 4-nitrophenol at 348 nm (pH-independent isosbestic point, $\varepsilon = 4147 \text{ M}^{-1} \text{ cm}^{-1}$) for over 5 min and determining the absorbance per minute from the generated slopes (Santiago et al., 2018). The reactions were performed at 30°C in 96-well plates (ref. 655801, Greiner Bio-One GmbH, Kremsmünster, Austria) and contained 0.09 to 3 μg proteins and 0.8 mM esters in a total volume of 200 μl. The effect of pH on the activity was determined in 50 mM BR buffer at pH 4.0–12.0, as described previously. Similar assay conditions were used to assay the effects of temperature on the ester hydrolysis of pNPC3, but in this case, the reactions were performed in 50 mM BR buffer pH 7.0. Note that the BR buffer consists of a mixture of 0.04 M H$_3$BO$_3$, 0.04 M H$_3$PO$_4$, and 0.04 M CH$_3$COOH that has been titrated to the desired pH with 0.2 M NaOH. All values were determined in triplicate and were corrected for non-enzymatic transformation. In all cases, the activity was calculated by determining the absorbance per minute from the generated slopes and applying the following equation:

$$\text{Rate} = \frac{\Delta \text{Abs}_{\text{min}}}{4147 \text{ M}^{-1} \text{cm}^{-1}} \cdot \frac{1}{0.4 \text{ cm}} \cdot \frac{10^6 \mu \text{M}}{1 \text{ M}} \cdot \frac{0.000044 \text{ L}}{1 \text{ mg protein}}$$

Poly(propylene glycol) diacrylate (ref. 455024, Merck Life Science S.L.U., Madrid, Spain) and poly(DL-lactide) with an average molecular weight 2,000 (ref. AP224, PolySciTech, Akina, IN, United States) were assayed as described previously (Hajighasemi et al., 2018). The hydrolysis of polyethylene terephthalate (PET) films (prepared as reported by Bollinger et al., 2020) and particles, which were prepared using PET from a bottle (from a local shop - Granini brand), as described previously (Pütz, 2006), was evaluated at 30°C and pH 8.0 with 270 μg protein/ml and 2 mg/ml plastic material, as previously reported (Bollinger et al., 2020).
The effect of the inhibitors mercaptoethanol (ref. M7154) and iodoacetamide (ref. I1149), which were both from Merck Life Science S.L.U., Madrid, Spain, was tested as follows. A mixture containing the purified enzymes (final concentration of 1 mg/ml) in 190 µl of 40 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (HEPES) at pH 7.0 and the inhibitors (final concentration, 1–10 mM) was incubated for 5 min to 24 h at 30–45°C. The reaction was initiated by adding pNPC₃ (0.8 mM, final concentration), and the activity was measured for over 5 min as described above and compared to control samples without inhibitors.

### Circular Dichroism to Estimate Thermal Denaturation

Circular dichroism (CD) spectra were acquired between 190 and 270 nm with a Jasco J-720 spectropolarimeter equipped with a Peltier temperature controller in a 0.1-mm cell at 25°C. The spectra were analyzed, and denaturation temperature (Tₜₜ) values were determined at 220 nm between 10 and 85°C at a rate of 30°C per hour in 40 mM HEPES buffer at pH 7.0. CD measurements were performed at pH 7.0 and not at the optimal pH (8.5–9.0) to ensure protein stability. A protein concentration of 0.5 mg/ml were performed at pH 7.0 and not at the optimal pH (8.5–9.0) to ensure protein stability. A protein concentration of 0.5 mg/ml was used. Tₜₜ (and the standard deviation of the linear fit) was calculated by fitting the ellipticity (mdeg) at 220 nm at each of the different temperatures using a 5-parameter sigmoid fit with SigmaPlot 13.0.

### Codes and Accession Numbers

The sequences were named based on the code “Est,” which refers to Esterase, followed by a letter indicating the origin of the sample, as follows: Est₄ₐ, esterase from the uppermostoxic B1A strata; and Est₈, esterase from the lowermost anoxic B1B sediment attached strata. The final number (subscript) is an arbitrary number representing the number of enzymes per site. Sequences encoding enzymes were deposited under the BioProject IDs PRJNA193663 (for B1A) and PRJNA193664 (for B1B) in the NCBI public database, with the accession numbers detailed in Table 1.

### Three-Dimensional Modeling

The models of the protein structures were predicted with AlphaFold 2.1.0 (Jumper et al., 2021).

### RESULTS

#### Enzyme Selection and Divergence at the Sequence Level

Microbial communities inhabiting two distinct compartments within Los Rueldos AMD formation were screened for sequences encoding carboxylesterases or lipases. For that, we used two complementary metagenomics approaches, namely, naïve and homology sequence screens. First, a total of approximately 81,000 pCCFO51 clones from each clone library (equivalent to 2.8 Gbp for B1A and 3.20 Gbp for B1B) were screened for esterase/lipase activity using plate-based screen with αNA and tributyrin as model substrates. We identified a total of 10 positive clones being active against both substrates in the B1A clone library, whereas no positives were found in the B1B library. The fosmids with insert lengths ranging from 16,545 to 41,280 bp were fully sequenced, from which 10 genes (one per positive clone), encoding presumptive esterases/lipases, were identified. In addition, we searched the predicted protein-coding genes obtained through next-generation sequencing for sequences encoding esterases and lipases by BLAST search against the ESTHER and LED databases. A total of 6 full-length sequences (B1A: 2; B1B: 4), with accession numbers EQD63018.1, EQD66234.1, EQD71191.1, EQD52136.1, EQD55146.1, and EQD26916.1, encoding potential enzymes were identified. Taken together, a total of 16 genes encoding hydrolases from the α/β-hydrolase fold superfamily, specifically, 12 from B1A (Est₄₁ to Est₄₁₂) and 4 from B1B (Est₅₁ to Est₅₁₂), were identified (Table 1). As determined by Matcher (EMBOSS package), the pairwise amino acid sequence identity for 14 of the 16 α/β hydrolases ranged from 4.0 to 44.9%. This, together with the fact that only 2 out of 16 polypeptides were highly similar (Est₄₅ and Est₅₆) differ in only 2 amino acids: arginine 152 and alanine 179 in Est₄₅, cysteine 152 and threonine 179 in Est₅₆, suggests a large divergence at the sequence level within the enzymes examined, and that the diversity of polypeptides was not dominated by a particular type of protein or highly similar clusters of proteins, but rather by diverse non-redundant sequences. Note that only 1 of 10 sequences selected after naïve screens was found in the metagenomic data generated after direct DNA sequencing (Est₄₁₁, which is 99% identical to GenBank accession no. EQD37671.1 from the Los Rueldos metagenome) (Méndez-García et al., 2014). This demonstrates that both types of screens (naïve and in silico) are complementary tools for enzyme discovery. However, deeper metagenomic sequencing could potentially detect all enzymes isolated by naïve screens.

The deduced molecular mass and estimated isoelectric point (pI) values ranged from 23.19 to 101.53 kDa and from 4.62 to 10.04, respectively. Putative proteins exhibited a maximum amino acid sequence identity ranging from 39 to 100% to putative esterases/lipases in public databases (Table 1). It is worth mentioning that Est₄₃, Est₅₆, and Est₅₃ are related to presumptive esterase/lipase-like subfamily proteins of the Serine-Glycine-Asparagine-Histidine (SGNH) hydrolases, Est₄₁₁ to presumptive glycoside-hydrolase family GH114 (N-terminal domain) and CE₄_Peaₐ_like hydrolases (C-terminal domain), and Est₅₁₂ to presumptive sialate O-acetylerases. A further TBLASTX search against metagenomics proteins deposited in databases revealed no similarity of 10 proteins with homologous AMD metagenome proteins. In contrast, five proteins (Est₄₃ to Est₅₆, and Est₅₁₂) do share from 27 to 54% homology to three proteins from the Carnoules arsenic-contaminated mine drainage (GenBank: CB107622.1, CB197521.1, and CB100527.1). This finding suggests that esterases/lipases from microbial communities from the Los Rueldos site are distantly related to proteins from other known homologous proteins from AMD formations with metagenome sequences available. It also reflects the large undiscovered pool of enzymes from bacterial species populating the Los Rueldos site.
**Primary Structure Analysis**

Based on the comparison of the primary structures, 14 families of sequence-related esterases and lipases have been reported (Arpigny and Jaeger, 1999; Rao et al., 2013). Sequence analysis categorized 13 enzymes from Los Ruedos into some of these known subfamilies (Figure 1) with most structurally similar homologs as follows: EstA8 [27%]; best hit in Protein Data Bank (PDB) 3DOH_A and EstB3 (41%; SOMB_A) to Family I; EstA1 (41%; 3V9A_A), EstA4 (41%; 4YPV_A), EstA5 (49%; 4YPV_A), and EstA6 (49%; 4YPV_A) to Family IV; EstA7 (53%; 4YPV_A) and EstB2 (41%; 1AUO_A) to Family VI; EstA3 (37%; 2OGT_A) to Family VII (EstA2); and EstA10 (41%; 4IVK_A) to beta-lactamase like Family VIII. EstA3 (43%; PDB code 3P94_A), EstA8 (44%; PDB code 3P94_A), and EstB3 (26%; PDB code 3KV_N_X) belong to Family II GDSL, but the structural alignment also confirms that they contain a domain that displays the characteristic α-β-α globular fold of the SGNH hydrolase family. In addition, EstB3 also contains a passenger domain providing the driving force for passenger translocation (Van den Berg, 2010). Three of the sequences could not be assigned to these subfamilies. First, EstA1 contains a 300 amino acid long N-terminal domain most similar to glycose-hydrolase family 114 and a 616 amino acid long C-terminal domain most similar to the carbohydrate esterase 4 (CE4) superfamily that includes chitin deacetylases (EC 3.5.1.41), N-acetylglucosamine deacetylases (EC 3.5.1.-), and acetylxylan esterases (EC 3.1.1.72), which catalyze the N- or O-deacetylation of substrates such as acetylated chitin, peptidoglycan, and acetylated xylan. Its N-terminal domain is most structurally similar (26%) to that of the glycosidase 2AAM_A and its C-terminal domain is structurally similar to the polysaccharide deacetylase from *Bacillus cereus* (4HD5). Second, EstA12 is associated with acetylxylan esterases (EC 3.1.1.72), with most similar (21%) structural homolog in PDB being 1ZMB_A. Third, EstB1 shows homology to small serine alpha/beta-hydrolase/acyl-peptidase (58%; 2FUK_A). The tentative amino acids participating in the typical catalytic triad of esterases and lipases are summarized in Table 1. Together, the analysis of the primary sequence suggests that the diversity of esterases was not dominated by a particular type of protein or a highly similar cluster of proteins, but rather by diverse non-redundant sequences belonging to different microbial groups and distinct esterase/lipase subfamilies.

**Source Organisms of Selected Polypeptides**

A search of oligonucleotide patterns against the GOHTAM database (Ménigaud et al., 2012) and TBLASTX analysis revealed compositional similarities between the DNA fragment containing the genes for EstA1, EstA2, EstA10, and EstB4, with genomic sequences of bacteria from the phylum *Actinomycetota*. Among them, only unambiguous affiliations at lower levels could be achieved for fragments containing EstA1, EstA2, and EstA10 that may most likely belong to bacteria from the genera *Acidithrix* (EstA1 and EstA10) and *Acidimicrobium*/*Ferrimicrobium* (EstA2), both from the family *Acidimicrobiaceae* within the order *Acidimicrobiales*. Note that EstA1 and EstA10 showed 99–100% sequence identity with uncharacterized esterases and lipases (WP_052605564.1 and WP_052605292.1) from *Acidithrix ferrooxidans*, and EstA2 showed 94% sequence identity with the best hit in Protein Data Bank (PDB) 3P94_A).

**Table 1 | General sequence-based characteristics of Los Ruedos esterases.**

| Name       | Accession number | Contig bp [taxonomic origin (phylum, genus)] | Identity and best hit | pI | Putative catalytic triad |
|------------|-----------------|--------------------------------------------|-----------------------|----|--------------------------|
| EstA1 a,b  | KY010297        | 41,280 (*Actinomycetota, Acidithrix*)       | 99%; WP_052605564.1   | 4.62 | Ser146 Asp193 His270    |
| EstA2 a,b  | KY010298        | 33,407 (*Actinomycetota, Acidimicrobium*/Ferrimicrobium) | 94%; NNN14078.1       | 5.03 | Ser185 Asp131 His841    |
| EstA3 a,b  | KY010300        | 32,091 (Acidobacteria, a.a.)                | 58%; WP_041839843.1   | 5.44 | Ser183 Asp221 His394    |
| EstA4 a,b  | KY010299        | 35,459 (Proteobacteria, Acidiphilum)        | 79%; OYV70865.1       | 5.53 | Ser159 Asp254 His294    |
| EstA5 a,b  | KY010302        | 26,958 (Proteobacteria, a.a.)               | 62%; OD57651.1        | 5.43 | Ser159 Asp254 His294    |
| EstA6 a,b  | KY010302        | 39,640 (Proteobacteria, a.a.)               | 62%; OD57651.1        | 5.32 | Ser159 Asp254 His294    |
| EstA7 a,b  | EGD60018.1      | 2,621 (Proteobacteria, a.a.)               | 64%; CDU94315         | 6.29 | Ser112 Asp175 His207    |
| EstA8 a,b  | KY010301        | 38,628 (Proteobacteria, a.a.)               | 56%; WP_063675888.1   | 10.04 | Ser90 Asp2317 His240   |
| EstA9 a,b  | EGD66324.1      | 1,763 (Proteobacteria, a.a.)               | 53%; WP_055248968.1   | 7.12 | Ser120 Asp188 His220    |
| EstA10 a   | KY010303        | 16,545 (Actinobacteria, a.a.)              | 100%; WP_052605292.1  | 5.55 | Ser75 Lys75 Tyr193     |
| EstA11 a   | KY010304        | 40,600 (Proteobacteria, a.a.)              | 67%; WP_051488053     | 9.71 | Asp506 His578 His582    |
| EstA12 a   | KY010305        | 35,290 (Bacteria, a.a.)                    | 41%; WP_009508720.1   | 9.65 | Asp100 Asp311 His318    |
| EstB1 a,b  | EGD71191.1      | 2,283 (Proteobacteria, a.a.)               | 69%; WP_094623914.1   | 5.89 | Ser117 Asp165 His294    |
| EstB2 a,b  | EGD52136.1      | 2,483 (Proteobacteria, a.a.)               | 48%; WP_055790651.1   | 6.11 | Ser116 Asp164 His196    |
| EstB3 a,b  | EGD26916.1      | 13,465 (Proteobacteria, a.a.)              | 49%; WP_026633329.1   | 5.7 | Ser199 Asp313 His316    |
| EstB4 a,b  | EGD55146.1      | 13,877 (Actinobacteria, a.a.)              | 39%; WP_051823767.1   | 6.07 | Ser96 Asp202 His231     |

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a Identified by naïve screen.
b Protein expressed as active soluble form.
c Identified by sequence-based screen from metagenomic sequencing data; accession number of the sequences in NCBI being ID PRJNA193663 (for B1A) and PRJNA193664 (for B1B).
d As determined by GOHTAM database (Ménigaud et al., 2012) and TBLASTX; taxonomic assignation to phylum and genera are shown, with a.a. defining those for which ambiguous assignments below phylum level were obtained.
\* Best hit and identity as shown by TBLASTX.
1 Presumptive catalytic triad as found by 3D structure analysis and alignment with homologous proteins with solved crystal structures.
identity with an uncharacterized esterase-lipase (NNN14078.1) from Acidimicrobiaceae. EstA3 was most likely derived from an uncultured bacterium assigned to the phylum Acidobacteria with ambiguous affiliation below the phylum level. The genes for EstA4 to EstA9, EstA11, and EstB1 to EstB3 were associated with uncultured bacteria of the Proteobacteria phylum, with ambiguous affiliations at a lower taxonomic level, except for EstA4, which was most likely derived from a bacterium of the genus Acidiphilium from the family Acetobacteraceae within the order Rhodospirillales (best hit OYV70855.1 from Acidiphilium sp., 79% homology). All these bacterial groups have been detected in biofilms thriving in the Los Rueldos mine (Méndez-García et al., 2014). No clear affiliation, other than Bacteria, could be found for EstA12. Note that in some cases no clear affiliation to a taxon of source organism below the level of the phylum could be established, either because of the short fragment length or the low compositional similarity between the metagenomic fragments and the sequences of related bacterial chromosomes and plasmids do not allow proper assignations.

**Enzyme Characterization**

From the 16 sequences selected, 8 from B1A and 2 from B1B were successfully cloned, expressed, and purified as soluble active proteins when expressed in pET Ek/LIC 46 vector and *E. coli* BL21 as the host. These proteins were herein referred to as EstA1 to EstA8, EstB1, and EstB2. The remaining six (EstA9-EstA12 and EstB3-EstB4) could not be produced in soluble active form (they formed inclusion bodies) in the expression system applied herein, which consists of the use of the Isopropyl β-D-1-thiogalactopyranoside (IPTG)-inducible Ek/LIC 46 vector and *E. coli* strain BL21 (DE3) as a host, and their properties are not described herein. Refining the expression conditions, which included variations in the expression conditions (16, 30, and 37°C) and IPTG concentration (from 0.1 to 2 mM),
resulted in unsuccessful production of sufficient active protein material for characterization. Further efforts may be needed with different expression vectors, which is beyond the scope of the present study.

The substrate profile of all α/β hydrolases was first examined using a set of esters commonly used to characterize esterases and lipases, namely, pNP esters such as pNPC$_2$, pNPC$_3$, pNPC$_4$, pNPC$_8$, pNPC$_{10}$, and pNPC$_{12}$. All ester hydrolases preferred short-chain-length pNP-esters, particularly pNPC$_2$ (EstA$_3$), pNPC$_3$ (EstA$_2$, EstA$_4$, EstA$_5$, EstA$_6$, EstB$_1$, and EstB$_2$), and pNPC$_4$ (EstA$_1$, EstA$_7$, and EstA$_8$) (Table 2). Within all six pNP ester tested, all but one (EstA$_3$) enzyme was able to hydrolyze up to pNPC$_{12}$, albeit at a much lower level (from 62- to 3,900-fold) compared to shorter derivatives. Considering the preferred pNP esters, the maximum specific activity ranged from 3.06 ± 0.03 to 679.8 ± 9.8 U/mg. We further test the possibility that the enzymes hydrolyze substrates other than pNP esters, particularly, plastic substrates and esters relevant to plastics. Using previously described conditions (Hajighasemi et al., 2018; Bollinger et al., 2020), we did not find that any of the enzymes hydrolyzed large plastic materials such as poly(propylene glycol) diacrylate, poly(DL-lactide), amorphous PET film, and PET nanoparticles. However, by using a pH-indicator assay, we found that the enzymes were able to hydrolyze other terephthalate esters and acrylate esters. Thus, as shown in Table 3, six of the enzymes hydrolyzed esters relevant to acrylic acid plastics, e.g., 2-naphthyl acrylate and tripropylene glycol diacrylate, a commonly used material principally exploited to prepare thermally stable polymers (He et al., 2017). These substrates, herein found to be converted at a maximum rate of 3,915 ± 48 U/g, are rarely hydrolyzed by esterases and lipases. There are only two examples reported, namely, human salivary pseudocholinesterase and cholesterol esterase (Finer and Santerre, 2004; Cai et al., 2014). In addition, one of the enzymes (EstA$_8$) was capable

### TABLE 2 | Specific activity (U/mg pure protein) for each of the enzymes tested over a set of p-nitrophenyl (pNP) esters of different lengths.

| Ester          | pNPC$_2$      | pNPC$_3$      | pNPC$_4$      | pNPC$_8$      | pNPC$_{10}$   | pNPC$_{12}$   |
|----------------|---------------|---------------|---------------|---------------|---------------|---------------|
| EstA$_1$       | 76.43 ± 0.31  | 158.5 ± 2.5   | 204.8 ± 10.7  | 2.75 ± 0.14   | 0.62 ± 0.05   | 0.19 ± 0.09   |
| EstA$_2$       | 30.39 ± 0.19  | 50.56 ± 0.78  | 27.68 ± 0.83  | 1.60 ± 0.01   | 0.82 ± 0.08   | 0.16 ± 0.05   |
| EstA$_3$       | 30.40 ± 0.11  | 4.73 ± 0.05   | 1.16 ± 0.03   | n.d.$^a$      | n.d.$^a$      | n.d.$^a$      |
| EstA$_4$       | 28.39 ± 2.6   | 280.4 ± 8.7   | 196.1 ± 2.7   | 32.55 ± 2.67  | 11.19 ± 0.48  | 0.100 ± 0.03  |
| EstA$_5$       | 120.6 ± 5.3   | 665.9 ± 10.8  | 293.4 ± 14.1  | 8.11 ± 1.45   | 0.29 ± 0.02   | 0.17 ± 0.01   |
| EstA$_6$       | 520.1 ± 1.7   | 679.8 ± 9.8   | 467.7 ± 6.4   | 11.93 ± 0.06  | 0.37 ± 0.02   | 0.12 ± 0.01   |
| EstA$_7$       | 1.91 ± 0.03   | 2.18 ± 0.08   | 3.23 ± 0.06   | 0.53 ± 0.02   | 0.14 ± 0.01   | 0.02 ± 0.01   |
| EstA$_8$       | 14.07 ± 0.95  | 26.13 ± 0.71  | 70.39 ± 0.16  | 24.81 ± 0.09  | 0.95 ± 0.06   | 0.85 ± 0.02   |
| EstB$_1$       | 4.73 ± 0.33   | 15.18 ± 0.86  | 9.05 ± 0.61   | 4.25 ± 0.24   | 0.51 ± 0.04   | 0.24 ± 0.09   |
| EstB$_2$       | 2.32 ± 0.01   | 3.06 ± 0.03   | 2.17 ± 0.05   | 0.54 ± 0.04   | 0.12 ± 0.02   | 0.04 ± 0.01   |

Values calculated from triplicates at pH 8.0 and 30°C.

$^a$Not detected, activity level below detection limits under our assay conditions.

pNPC$_2$: p-nitrophenyl acetate; pNPC$_3$: propionate; pNPC$_4$: butyrate; pNPC$_8$: octanoate; pNPC$_{10}$: decanoate; pNPC$_{12}$: decanoate.

### TABLE 3 | Specific activity (U/g pure protein) for each of the enzymes able to hydrolyze a set of structurally different plastic-related esters.

| Substrate       | Structure | Specific activity (U/g pure protein) |
|-----------------|-----------|-------------------------------------|
| 2-Naphthyl acrylate | ![Structure](image) | 370.7 ± 20.3 37.7 ± 4.1 50.2 ± 0.4 1308 ± 79 n.d.$^a$ n.d.$^a$ 748.1 ± 15.9 144.5 ± 10 |
| Tri(propylene glycol) diacrylate | ![Structure](image) | 514.5 ± 36.3 4.9 ± 1.1 8.6 ± 0.7 3915 ± 48 n.d.$^a$ n.d.$^a$ 1652 ± 75 73.5 ± 2.9 |
| Dibenzyl terephthalate | ![Structure](image) | n.d.$^a$ n.d.$^a$ n.d.$^a$ n.d.$^a$ n.d.$^a$ n.d.$^a$ 432.2 ± 27.5 n.d.$^a$ |
| BHET$^b$        | ![Structure](image) | 308.3 ± 3.9 5.0 ± 1.0 n.d.$^a$ n.d.$^a$ 26.7 ± 1.7 12.3 ± 0.4 336.9 ± 3.6 91.1 ± 3.2 |

Assays were performed in triplicate with values for each of the replicates given in the table with standard deviation. Values calculated at pH 8.0 and 30°C.

$^a$Not detected, activity level below detection limits under our assay conditions.

$^b$Time course of the degradation shown in Supplementary Figure 3.
of hydrolyzing dibenzyl terephthalate (432.2 ± 27.5 U/g), an intermediate produced during chemical PET recycling with benzyl alcohol in the presence of a catalyst (Donahue et al., 2003). No esterase or lipase has been reported to date that degrades this substrate. In addition, six of the esterases (EstA1, EstA2, EstA5, EstA6, EstA8, and EstB2) efficiently hydrolyzed BHET (from 5.0 ± 1.0 to 336.9 ± 3.6 U/g), an intermediate in the degradation of PET (Yoshida et al., 2016). High performance liquid chromatography (HPLC) analysis, performed as described (Bollinger et al., 2020), confirmed the hydrolysis of BHET to mono-(2-hydroxyethyl)-terephthalic acid (MHET) and not to terephthalic acid. To conclude, the enzymes reported herein from the Los Rueldos AMD formation showed high activity for converting and recycling components of synthetic plastics, namely, acrylic- and terephthalate-based plastics, and could be of potential use in developing plastic degradation strategies.

Using gNPC3 as a substrate, the purified proteins were most active at temperatures ranging from 30 to 65°C (Figure 2). The average annual temperature in Los Rueldos is 13.8 ± 0.6°C, which varied from 10 ± 0.6°C to 17.1 ± 0.6 (Méndez-García et al., 2014), with a temperature of 17°C when samples were taken (July). At this value, the enzymes retained from 20 to 61% of the activity shown at the optimal temperature (Figure 2).

Using gNPC3 as a substrate, all enzymes showed an optimum pH for activity from neutral to slightly basic, which varied from 7.0 to 9.0 (Figure 2). This finding suggests that these proteins are most likely intracellularly produced, consistent with the absence of signal peptides in their sequences. Even though the enzymes showed a slightly basic optimum pH, all retained 33–68% of their activity at pH 5.5. Interestingly, EstA6 shows two activity peaks, one at pH 5.5 and one at pH 9.0, while EstA5, which only differs in two amino acids, has an optimum pH of 9.0 (Figure 2).

Sequence analysis revealed that EstA5 and EstA8, which have their origins in a bacterium of the phylum Proteobacteria, differ in only 2 amino acids (99.4% identity). Positions 152 and 172 are occupied by Arg and Ala in EstA5 and by Cys and Thr in EstA8, respectively. Notably, EstA5 was most active at 30°C, retaining more than 80% of the activity at temperatures from 20 to 45°C (Figure 2). The optimum temperature for activity increased up to 45°C for EstA6, which maintained more than 80% of its activity in the range from 30 to 60°C. Analysis by circular dichroism revealed that EstA5 showed a sigmoidal curve from which a temperature of denaturation of 60.4 ± 0.2°C could be obtained (Figure 3). However, the curve for EstA6 shows two transitions, one with a denaturation temperature of 48.1 ± 0.8°C and a second at 75.7 ± 0.2°C. The presence of these two phases may therefore indicate that the presence of these two amino acids may contribute to protein stability and its denaturation under distinct conditions. This result may explain the higher optimum temperature for the activity of this enzyme compared to EstA5, and the stabilization effect of Cys152 and Thr172. This difference in thermostability between EstA5 and EstA6 can probably be explained by the difference in amino acid 152, since EstA6 has a Cys that would allow it to make a possible disulfide bridge with Cys181, giving it higher thermostability than EstA5 (since EstA5 has an Arg at position 152 instead of a Cys), as shown by examination of the 3D models (Supplementary Figure 2 and Figure 4). It is plausible that this difference may also be responsible for the different pH profiles of both enzymes (Figure 2). If the disulfide bridge was present in EstA6, it could be removed by reduction or chemical modification. Activity tests revealed that both enzymes are resistant to the reducing agent beta-mercaptoethanol, with no activity lost even after 24 h of incubation at 30–45°C in the presence of a 1–10 mM inhibitor. By contrast, in the presence of the cysteine alkyllating agent iodoacetamide, both enzymes were rapidly inactivated after 5 min. Thus, we could not verify the presumptive formation of the disulfide bridge in EstA6, and further studies are needed to test this assumption.

**DISCUSSION**

The effects of environmental constraints as prime forces shaping AMD populations have only begun to be elucidated through omics studies (Méndez-García et al., 2015). These effects are also of high interest in the context of the isolation and characterization of novel enzymes, for which limited data are available. However, the difficulty of cultivating organisms inhabiting AMD sites, which is due to their longer generation times, lower biomass yields, and cultivation conditions that are not yet fully understood, requires different strategies to overcome the problems associated not only with their cultivation, but also with the isolation of enzymes. Metagenomic approaches allow the screening of enzymes from such extreme environments. However, by using these tools, we have thus far explored only a small fraction of the enormous diversity on the planet, especially that of organisms inhabiting extremely acidic environments (Ferrer et al., 2015), again indicating the importance of establishing enzyme screening programs for AMD sites. The particular characteristics of the Los Rueldos AMD site (Méndez-García et al., 2015) that make it an interesting study site include the following. First, it is populated by a larger diversity of Bacteria and Archaea compared to other AMD sites, containing a total of 39 different species. Second, it has high microbial heterogeneity in local microniches defined by its O2 concentration gradients and spatial and biofilm architecture. As an example, only 1 of 18 species inhabiting the two distinct compartments in a stratified streamer investigated herein, namely, the oxic uppermost (B1A) and anoxic lowermost (B1B) sediment-attached strata, was shared. Therefore, it is plausible that Los Rueldos may also contain a greater diversity of microbial products such as enzymes.

We have sought to address this possibility by screening for esterases and lipases from the α/β-hydrolase fold superfamily in microbial communities inhabiting the Los Rueldos AMD site. These enzymes are desired tools for biocatalysis in a variety of industrial sectors (Daiha et al., 2015; Ferrer et al., 2015). Microorganisms that can survive under low pH values similar to those in Los Rueldos (pH ~2) could be good sources of enzymes that can be used, for example, as additives in detergents, for the biobleaching of pulp and paper, in the clean-up of effluent streams from the textile processing industry, and in the degradation of plastics (Gomes et al., 2003;
FIGURE 2 | pH and thermal profiles of the purified enzymes. The data represent the relative percentages (%) of specific activity (U/g) in triplicates compared with the maximum activity using pNPC₃ as substrate. For raw data see Supplementary Table 1.
Adrio and Demain, 2014; Nchedo Ariole and George-West, 2020) and other polymers (Fütterer et al., 2004).

We used two complementary approaches for enzyme mining. A sequence-based metagenomic approach that searched for homologous enzymes in the metagenomic sequence data and function-driven screens in which expression libraries were used to identify, by using specific colorimetric substrates (Ferrer et al., 2016; Peña-García et al., 2016), clones containing enzymes of interest that could be missed in shallow metagenomics sequencing. By using both approaches, we identified 16 sequences that were potentially encoding esterases and lipases. The amino acid sequences were distantly related to sequences found in other AMD formations, which was in agreement with the distinct Los Rueldos-specific populations (Méndez-García et al., 2014). Indeed, we only observed some degree of sequence identity (27–54%) to 3 homologs from the Carnoulès (lead-zinc) mine, France (Bertin et al., 2011) in 6 of 16 sequences. In addition, the large differences among the recovered enzymes may correspond to the high population diversity that characterizes the Los Rueldos site (Méndez-García et al., 2014).

Notably, activity-based screens did not yield any active clones from the library created from the anoxic lowermost strata (B1B), while they yielded 10 active clones from the library created from the oxic uppermost strata (B1A). Thus, we searched for such enzymes in B1B by screening sequence data generated in a previous study (Méndez-García et al., 2014). It is plausible that the presence of low-O2-adapted microbial species in B1B, in contrast to the aerobic species in B1A, may account for the low efficiency of heterologous gene expression after cloning of the genetic material in the E. coli host and, possibly, the lower efficiency of the screening tests in the B1B library compared to...
that obtained for B1A. However, the fact that similar proportions of identified proteins (by naïve and sequence screening) in B1A (7 of 12) and B1B (2 of 4) could be produced as soluble active proteins when expressed in *E. coli* suggests that this may not be the only reason explaining the absence of positive clones in the B1B library. We cannot rule out that the native promoters of the partial genes from microorganisms inhabiting B1B cloned in the pCCFOS1 fosmids were inactive in *E. coli*, resulting in failed active clones on the plates. The data provided in Tables 2, 3 revealed that B1B enzymes were among the least-active enzymes among all hydrolases identified and characterized in the present study, and it is therefore also plausible that the low efficiency of the screening tests may have been due to the low activity level of enzymes from microorganisms inhabiting the anoxic B1B compartment. Additionally, it is plausible that different screening conditions (temperature, pH, inductor concentration, etc.) may be needed to detect other active proteins and that the enzymes from B1B would be more active under other assay conditions, the investigation of which is beyond the scope of the present study.

Regardless of the problems associated with the screening efficiency in different environments, including extreme AMD formations such as Los Rueldos, the analysis of the optimal pH profile of 10 out of the 16 hydrolases that could be produced in active form additionally revealed that their optimal pH was in the range from 7.0 to 9.0. This finding suggests that all hydrolases are presumptively produced intracellularly by acidophiles that thrive in the acidic Los Rueldos environment with a pH of 2.0. A similar phenotype has been found for other enzymes from AMD inhabitants, such as ATP-dependent DNA ligase from *F. acidarmanus* Fer1 (Jackson et al., 2007) and ene-reductase from *Ferrovum* sp. JA12 (Scholtesse et al., 2016), with pH optima of 6.0–7.0. However, we observed that most of the enzymes showed a slightly acid-stable phenotype, retaining ~33–68% of activity at pH 5.5. It is plausible that the identification of enzymes with neutral-like pH optima is a consequence of screening tests performed at neutral pH, using a vector and host that allow mostly intracellular proteins to be produced and that presumably acid-stable enzymes could not be detected. In the future, performing naïve screens at such low pH values may help obtain additional active clones. However, while specific adaptations need to be explored in great detail, the retention of a high activity level at a slightly acidic pH might be attributed to the prevalence of acidic amino acids (negatively charged at a neutral pH) on the surfaces of these enzymes (Supplementary Figure 2), as reported for other proteins from acidophiles (Wu et al., 2020). Indeed, the relative frequencies of acidic residues in proteins in this study ranged from 67% to 54%, except for EstA (37%). As no major differences in pH profiles were observed when comparing the enzymes with the highest (EstA: 67%) and lowest (EstA: 37%) percentages of acidic residues, it is possible that other factors affect the activity and stability of the studied proteins from the Los Rueldos site. An example of this is the differences in stability against pH and temperature of EstA and EstA6, which show very different features and have only a two amino acid difference despite having the same percentage of acidic residues.

Furthermore, the biochemical properties of the esterases reported in this study revealed that all enzymes showed an activity-stability trade-off characteristic of mesophilic-adapted enzymes (from 30 to 65°C), which is a phenotype that has also been found for enzymes from other AMD inhabitants (Golyshina et al., 2006; Jackson et al., 2007; Ohara et al., 2014; Scholtesse et al., 2016). It is noticeable, however, that 5 of 10 characterized enzymes retained at least 50% of their maximal activity at temperatures as low as 12°C. The fact that the Los Rueldos site is characterized by a relatively low temperature compared to other AMD sites (Méndez-García et al., 2014) may account for this low-temperature-active phenotype. However, the lack of biochemical information on enzymes from other AMD sites does not allow us to validate this assumption.

Finally, it should be emphasized that the activity levels of the characterized enzymes (maximum for the best pNP substrates: approximately 680 to 3 U/mg, depending on the hydrolase) were in the range of other reported enzymes of different origins with esterase and lipase activities (Ferrer et al., 2015; Martinez-Martinez et al., 2018). The data suggest that the low-O2-adapted microbial species developed in the anoxic lowermost (B1B) sediment-attached strata do contain less-active enzymes than those developed in the oxic uppermost (B1A) strata under the conditions used herein. Whether this is typical in vivo or is a result of bias due to the assay conditions would require the characterization of a larger number of enzymes from both microenvironments in the Los Rueldos AMD system. The capacity of six of the enzymes from the Los Rueldos AMD formation, and thus the bacteria that contain them, for degrading acrylic- and terephthalate-like esters is noticeable. These enzymes could potentially be of use in developing plastic degradation strategies that have yet to be explored. In this context, the taxonomic distribution of top protein hits and the results of genome linguistics analysis suggested that the metagenomic fragments containing the six characterized enzymes that can potentially degrade plastic substrates most likely belong to Actinobacteria (genera related to *Acidithrix* and *Acidimicrobiurn/Ferrimicrobiurn*), Acidobacteria, and Proteobacteria (some are related to *Acidiphilium*). These are groups of acidophiles that have been largely neglected with respect to enzyme discovery.

Polyethylene terephthalate-degrading bioprospecting has shown that only a tiny fraction of carboxylic ester hydrolases can degrade PET and its intermediates BHET and MHET, and can degrade PET and its intermediates BHET and MHET, including Actinomyces (i.e., *Thermobifida, Thermomonospora*, and *Saccharomonospora*), Bacillus, Firmicutes (e.g., *Clostidiurn*), *Bacteroidetes*, *Proteobacteria* (e.g., *Pseudomonas, Enterobacteria*, and *Ideonella sakaiensis*), and fungi [*Fusarium* and *Thermomyces (Humicola)*] (for a recent example, see Yoshida et al., 2016; Danso et al., 2018; Bollinger et al., 2020; Yan et al., 2021). In this study, we found that bacteria from the genera *Acidithrix* (the host of EstA1), *Acidimicrobiurn/Ferrimicrobiurn* (the host of EstA2), and unknown genera from the phylum *Proteobacteria* (the hosts of EstA5, EstA6, EstA8, and EstB2) could degrade BHET and could
potentially degrade PET or PET oligomers under conditions yet to be explored, as no PET hydrolysis was detected under the assay conditions employed herein. Thus, the metagenomics approach applied herein expands the range of microorganisms containing enzymes supporting BHET hydrolysis and, possibly, PET depolymerization. For effective PET hydrolysis, in addition to a high degradation rate at 40–70°C, a broad range of pH stability (toward both the alkaline and acidic ranges) is one of the prerequisites of applied enzymes (Maurya et al., 2020), and these features were characteristic of some of the esterases from Los Rueldos reported herein. Further studies will reveal the catalytic efficiency and stability of hydrolases from Los Rueldos AMD systems to establish PET degradation systems or to support these systems in combination with other known PET-degrading enzymes.

It is plausible that the capacity to degrade plastic substrates comes from the adaptation of the active sites of enzymes to metabolize microbial polymeric substances that are naturally occurring in AMDs. Thus, in Los Rueldos, as in other AMDs such as the Richmond Mine at Iron Mountain (Jiao et al., 2010, 2011; Martínez-Martínez et al., 2013), the organisms present might contribute to the use/degradation of extracellular polymeric substances (EPS), which requires a broad range of enzymes (Flemming and Wingender, 2010).

**DATA AVAILABILITY STATEMENT**

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

**AUTHOR CONTRIBUTIONS**

MF and AP conceived this study. MM-M contributed to gene cloning and expression. PV, MM-M, LF-L, and MF performed biochemical data and interpreted the data. SR and VG performed the 3D modeling. CM-G and MF contributed sample processing and library construction. OG contributed the phylogenetic classification and properties. MF drafted and revised the manuscript. All authors discussed, read, approved the manuscript, and authorized its submission for publication.

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**SUPPLEMENTAL MATERIAL**

The Supplemental Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fmicb.2022.868839/full#supplementary-material

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