Diversity of cultivable bacteria from deep-sea sediments of the Colombian Caribbean and their potential in bioremediation

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Abstract The diversity of deep-sea cultivable bacteria was studied in seven sediment samples of the Colombian Caribbean. Three hundred and fifty two marine bacteria were isolated according to its distinct morphological character on the solid media, then DNA sequences of the 16S rRNA were amplified to identify the isolated strains. The identified bacterial were arranged in three phylogenetic groups, Firmicutes, Proteobacteria, and Actinobacteria, with 34 different OTUs defined at ≥ 97% of similarity and 70 OTUs at ≥ 98.65%, being the 51% Firmicutes, 34% Proteobacteria and 15% Actinobacteria. Bacillus and Fictibacillus were the dominant genera in Firmicutes, Halomonas and Pseudomonas in Proteobacteria and Streptomyces and Micromonospora in Actinobacteria. In addition, the strains were tested for biosurfactants and lipolytic enzymes production, with 120 biosurfactant producing strains (mainly Firmicutes) and, 56 lipolytic enzymes producing strains (Proteobacteria). This report contributes to the understanding of the diversity of the marine deep-sea cultivable bacteria from the Colombian Caribbean, and their potential application as bioremediation agents.

Keywords Bioremediation · Diversity of cultivable bacteria · Deep-sea sediments · Colombian Caribbean

Introduction

The ecological importance of bacteria in deep-sea sediments has been recognized since 1957, when Zobell and Morita (1957) isolated marine piezophiles. In deep-sea ecosystems, microbial communities contribute to global carbon recycling and represent Earth’s major ecological research frontier (Danovaro et al. 2014). Specifically, deep-sea sediments are substrates for bacterial growth because of its vast matrix of organic molecules mainly composed by carbon (C), nitrogen (N), and phosphorus (P) (Jamieson et al. 2013). However, the availability of these organic pools is highly variable geographically and depends on their origin, degradation rates, and utilization by benthic consumers (Li et al. 1999).
The Caribbean Sea is a semi-enclosed basin of the western Atlantic Ocean, it is bounded by the coasts of Central and South America on two sides and by the Antilles island chain and the arc of the Greater and Lesser Antilles. It has an average depth of 2400 m being a 75% more deeper than 1800 m approximately (Hernández-Ávila 2014). The Caribbean seafloor include typical abyssal soft-sediment extensions, numerous small canyons, and wide (10 km) and low-relief channels. It is divided into five deep-sea areas: the Grenada, Venezuelan, Colombian, and Yucatan Basins and the Cayman Trough, separated from each other by underwater ridges and sills (Miloslavich et al. 2010).

With the development of environmental DNA technology (Olsen et al. 1986; Pace et al. 1986), it was possible to study an unexplored diversity making the study of deep-sea environments a challenge in the field of ecology and evolution of microbial communities. In this sense, DNA sequences supply the tools to study the communities, especially gene encoding rRNAs providing a basis to estimate phylogenetic diversity and generate taxonomic inventories of microbial populations (Kai et al. 2017; Rappé and Giovannoni 2003). Additionally, the analysis of 16S rRNA gene libraries have shown that there is a substantial diversity of bacteria in deep-sea sediments, both within and between the phyla (Fry et al. 2008), with a high variation of beta-diversity, reflecting the physical, chemical and biological contrasts in deep-sea ecosystems (Zinger et al. 2011).

Worldwide, significant efforts have been made to cultivate marine microorganisms from deep-sea environments under controlled laboratory conditions, obtaining a low representation of these. Nevertheless, the few representatives available are highly valuable for discovering information about unusual types of metabolism, pressure and temperature adaptations, and growth under “extreme” conditions (Orcutt et al. 2011); also, marine microorganisms can be a source of unique metabolites for many industrial and environmental (bioremediation) applications.

Petroleum hydrocarbons have been considered one of the main pollutants of the environment, because during the process of exploration, collection and transportation of oil, leaks can arise that lead to the contamination of adjacent lands and bodies of water (Patowary et al. 2018). This problem has been reflected in Colombia, where oil spills have occurred due to accidental circumstances or attacks by groups outside the law, causing environmental damage, affecting ecosystem services and generating social and economic impacts due to the affeaction of activities such as fishing, consumption of drinking water, the cultivation of primordial species and the affectionation of marine ecosystems (Velásquez-Arias 2017).

Bioremediation is considered an effective, ecological and economic technology that allows to solve environmental problems caused by the presence of pollutants such as hydrocarbons, where microbial communities or their metabolic products are used to transform these pollutants. In less offensive compounds (Zouboulis et al. 2019). Biosurfactants are amphiphilic surface-active molecules of microbial origin produced with the aim to facilitate the accession and uptake of complex water-insoluble substrates like hydrocarbons, in this sense, they are commonly produced by hydrocarbon-degrading bacteria (Ibacache-Quiroga et al. 2013). Additionally, various studies support that hydrocarbon-degrading bacteria produce lipolytic enzymes such as lipases (Welz et al. 2021).

The aim of this study was to describe the phylogenetic diversity of the piezotolerant cultivable bacterial community from deep-sea sediments, identify the abundant and significant types of different bacteria, and to search for the potential use of these strains in the remediation of hydrocarbon contaminated environments.

Materials and methods

Sample collection and bacteria isolation

Marine deep sediments were collected using a box corer in 7 stations of the Colombian Caribbean Sea (Fig. 1), during the cruises to the blocks COL1, COL2, COL3, and COL10 conducted by the National Hydrocarbon Agency (ANH) (Vides et al. 2017). Physico-chemical parameters of the studied sediments and/or water column of the sampling sites were taken (Table S1), and the sediment samples were preserved at 4 °C until they were processed in the Marine Bioprospecting Laboratory—LabBIM at INVEMAR.

For bacteria isolation, 20 g of sediments were resuspended in 90 mL of peptone water with 1% of...
NaCl. Then, serial dilutions were made and 0.1 mL of each one were taken to cultivate by the spread plate method using different media (Table S2). The cultures were maintained at a range of temperatures between 4 and 70 °C under aerobic and anaerobic conditions during 8–30 days depending on the microbial growth observed (Table S2), an anaerobic chamber with a package of an atmosphere generation system specialized for the generation of an anaerobic atmosphere (AnaeroGenTM 2.5L- ThermoScientific) was used to maintain anaerobic conditions. Finally, the colonies that presented distinctive morphologies were selected. Axenic cultures were obtained through various subcultures in the same culture media where the strains were initially selected. A voucher and all associated information of each collected specimen was deposited at the Marine Museum of Natural History of Colombia–Makuriwa, where identification codes were assigned.

Bacteria classification and diversity

Bacteria classification was based on the analysis of 16S rRNA gene sequencing using the universal primers 27F/1492R (Lane 1991). Genomic DNA was extracted using the Invitrogen PureLink Genomic DNA kit. The DNA from all the purified isolates was used for PCR amplification of the 16S rRNA gene, then, the PCR—amplified products were subjected to sequencing by the Sanger method at National Center for Genomic Sequencing (Medellín, Colombia) reading both chains.

The sequence alignment and classification were performed in the SILVA ribosomal RNA database project (SILVA SSU 138.1) using the web resources for alignment, classification and tree services (SILVA-ACT). SILVA is a comprehensive on-line resource for quality checked and aligned ribosomal RNA sequence data (Glöckner et al. 2017; Quast et al. 2013). With more than 9 million of aligned small 16S rRNA sequences, SILVA 138.1 is an extensively curated taxonomy and non-redundant datasets provide an ideal reference for bacteria classification and sequence
alignment approaches (Quast et al. 2013). Our sequences were aligned using SINA (SILVA Incremental Aligner v1.2.11) online service provided by the SILVA ribosomal RNA project (https://www.arb-silva.de/aligner/). SINA uses a combination of k-mer sequence search and partial order alignment (POA) to maintain very high alignment accuracy of rRNA gene sequences (Pruesse et al. 2012).

The genetic diversity and species limits of cultured bacteria were estimated by cluster analyses in Mothur v.1.44.0 (https://www.mothur.org/wiki/Main_Page) (Schloss et al. 2009). Mothur is a microbial ecology software package focused in 16S rRNA gene sequence data analyze, using the different methods available (Schloss 2020). For the analyzes were used various levels of sequence identity between 80 and 100% to define of operational taxonomic units (OTUs), two limits were adopted, the standard limit of 97% of similarity and the recommended cut-off of 98.65% (Kim 2014; Mysara et al. 2017). Phylogenetic trees were constructed using Maximum Likelihood (ML) method, implemented in RAxML-HPC2 v8.2.10 (Stamatakis 2014), on the CIPRES Science Gateway portal v3.3 (https://www.phylo.org/) (Miller et al. 2010), under generalized time-reversible (GTR + Γ) model and 100 heuristic independent runs. For bootstrapping, we performed 1000 pseudo-replicates analyses; this was performed under auto Majority Rule Criterion (autoMRE).

The identity of the sequences was compared with reference taxonomies hosted by SILVA-ACT (RDP, EMBL-EBI/ENA and SILVA) (Cole et al. 2014; Quast et al. 2013; Toribio et al. 2017), using the least common ancestor (LCA) method and searching five of the nearest sequences with a similarity superior at 99%.

**Biosurfactants and lipolytic enzymes production screening**

Initially all the strains were put to grow in Bushnell–Hass broth (Bushnell and Hass 1941) supplemented with 1% of Diesel fuel. Bushnell–Hass is a common culture medium used as method for isolating organisms capable of utilizing hydrocarbons as carbon source. Briefly, one loop of colonies was suspended in 10 mL of medium until 0.5 McFarland scale, aliquots of 1 mL were used for inoculum of 25 mL of production medium. After 7 days of incubation at 30 °C and 140 rpm, 1 mL of the cell-free culture was obtained by centrifugation at 9520 g (4000 rpm), for 10 min at room temperature, and this cell free supernatant was used for the next tests.

The oil—spreading test was made to detect the production of extracellular biosurfactants. Briefly, 20 μL of oil crude were poured over 40 mL of distilled water in a petri dish. Then 20 μL of cell-free culture were added carefully in the center of the layer, the diameter of spreading was measured in mm. As negative control was used 20 μL of the broth without inoculum, and as positive control of displacement 20 μL of neutral detergent (Extran® MA 02), each one in different petri dishes. The assay was done by triplicate. It was considered as positive result, the spreading of the crude oil and a zone of emulsification on the surface (Walter et al. 2013).

Lipolytic activity was evaluated on staining fat emulsion agar using Nile blue A as indicator and olive oil as substrate. The cultures were incubated at 30 °C for 3–7 days and *Pseudomonas aeruginosa* ATCC 10145 was used as positive control. The observation of blue colonies surrounded by decolorate and transparent zones was an indication of positive strains for lipase production, whereas no lipolitic strains were observed as white and rose (Eisenberg 1939).

**Results**

**Sample collection and Bacteria isolation**

Three hundred and fifty-two cultivable bacteria were isolated from deep-sea sediments in seven stations E553 (1254 m), E568 (2106 m), E428 (3186 m), E466 (3328 m), E603 (3474 m), E617 (3704 m), E612 (4220 m). Initially, all the isolates were identified based on the different colony characteristics. The stations with most isolated strains were E603 (56), E617 (50), and E612 (44) which were the more farthest from the coast, followed by E428 (35), E466 (30), E553 (25), and E568 (22) (Table S3). On the other hand, physicochemical values (Temperature, salinity, dissolved oxygen) were similar in all stations.

**Bacteria classification and diversity**

The 16S rRNA gene of the cultivable bacteria isolated were sequenced with an average length of 1352 bp.
Thirty-four OTUs were defined at ≥ 97% and seventy at ≥ 98.65% of similarity (Table S3). The bacterial assemblage was arranged in three phylogenetic groups, Proteobacteria, Actinobacteria (Gram-positives with high G + C percentage), and Firmicutes (Gram-positives with low G + C percentage) (Fig. 2). The OTUs composition was 51% Firmicutes, 34% Proteobacteria and 15% Actinobacteria. Bacillus and Fictibacillus were the dominant genera in Firmicutes, Halomonas and Pseudomonas in Proteobacteria and Streptomyces and Micromonospora in Actinobacteria (Fig. 2).

The bacteria were identified and related with isolates previously registered in the databases of taxonomic references (RDP, EMBL-EBI/ENA and SILVA). Most of the isolated belonged to the phylum Firmicutes, followed by Proteobacteria (Fig. 3A). Several interesting and recently described genera were identified, such as Piscibacillus (Tanasupawat et al. 2007), Lysinibacillus (Ahmed et al. 2007), Fictibacillus (Glaeser et al. 2013), Aquibacillus (Amoozegar et al. 2014), Domibacillus (Seiler et al. 2013), Idiomarina (Ivanova et al. 2000), Cellulosimicrobium (Schumann et al. 2001), Cobetia (Arahal et al. 2002), Paenibacillus (Krishnamurthi et al. 2009), and Plantactinospora (Qin et al. 2009) (Fig. 3b, Table S3).

Biosurfactants and lipolytic enzymes production screening

One hundred and twenty isolated strains were identified as biosurfactant producers by the oil-spreading test. The majority of the promising biosurfactant producers belong to the Firmicutes (85 strains) and the station with the most biosurfactant producing strains was the deepest E612 (4220 m) (33). On the other hand, a less number of strains (56) were identified as lipase producers, the majority belongs to the Proteobacteria (29), followed by Firmicutes (23), and the majority of the lipase producing strains were from the station E603 (3474 m) the farthest from the coast. In

![Fig. 2 Maximum likelihood (ML) tree on the GTR + G model (ln -14,587.3509), based on the nearly complete and aligned 16S rRNA gene sequences. Above the branches, bootstrap support values (> 50) are provided for each node](image)
addition, 20 strains showed capacity to produce both biosurfactants and lipases, specifically from the species Alteromonas sp., Pseudomonas sp., Micro-
monospora sp., Bacillus sp., Halomonas sp., Chromo-
halobacter sp., Cobetia sp., Fictibacillus sp., Paenibacillus sp., and Domibacillus sp. (Fig. 4, Table S4).

Discussion

The sampling stations E466, E428, E568, E553 (Fig. 1) are located in the central part of the Colombian basin, in front of the coastline between the Tayrona National Park in the Magdalena Department, covering an area of 2.500.000 ha. The approximate minimum distance to the coast is 60 mn and the maximum is 160 mn, covering depths from 2100 to 4500 mn that is the most extensive geomorphological feature within the Caribbean Large Ecosystem (Vides and Alonso 2016). The four stations are highly influenced by the delta of the Magdalena River, presenting changes in the water composition in hundreds of kilometers inside the sea contributing with a high quantity of sediments in suspension (142.6 $\times$ 10$^6$ t a$^{-1}$) with a sedimentation rate $\leq$ 1.430 mm a$^{-1}$ in the deepest areas of the delta front (Restrepo et al. 2015). This change in the water composition also might cause complex hydraulic phenomena, such as the conformation of a submarine delta composed of submarine sediment bars (Castañ-Uribe et al. 2003). On the other hand, the sampling
stations E603, E612, and E617 (Fig. 1) are located on the northeastern extreme of Colombia’s marine territory on the Caribbean Basin. It also includes a portion of the Beata Ridge, an underwater ridge of the south bed of the Caribbean Sea (Vides et al. 2018).

For the definition of OTUs, the cut-off at $\geq 98.65\%$ are biologically most representative according to the bacterial community (Fig. 3). This result agrees with the proposed by Kim (2014), who calculated this value based in the complete sequence of the 16S gene allowing a more accurate estimation of the microbial diversity. In this study, 352 cultivable bacteria were isolated by classic microbiology techniques; these strains tolerate the sudden change of pressure, so they can be cataloged as piezotolerant (Velásquez et al. 2018). The cultivation of microbes is still necessary, because it contributes to the comprehensive understanding of the detailed physiology and the complex environmental processes associated with them. Likewise, this is the first step for the microbes to be applied on environmental remediation, energy and pharmaceutical industry. In addition, these cultures can provide complete genomes and the means to verify potential metabolic ability or ecosystem function (Zhang et al. 2018).

The hypothetical difference between marine and terrestrial microorganisms derives from salinity an important factor for microbial growth, osmotic regulation in fresh and marine water is completely different (Sivaperumal et al. 2017). Besides, the marine bacterial communities seem significantly influenced by depth and the distance from the coast. The most abundant Phylum in all stations was Firmicutes except for E603 (Fig. 3a), which is the station farthest from the coast. The majority of the species from the Firmicutes are related to endospore-forming bacteria (EFB). The ability to produce endospores and the great metabolic and physiological diversity are two characteristics of the EFB that allow their distribution in all environments (da Silva et al. 2013). Likewise, the sporulation allows EFB to tolerate different environmental conditions and constitute a seed bank that allows populations to persist over longer time scales and thereby maintain biodiversity in an environment (Cupit et al. 2019).

Of the nine families of Firmicutes, the Bacillaceae and the Planococcaceae are the largest taxa. Members of the family Bacillaceae are reported as Gram positive, rod-shaped, endospore forming and moderately halophilic some have been isolated from marine environments (Lu et al. 2001), fermented foods (Tanasupawat et al. 2007) and salt lakes (Amoozegar et al. 2014). Within this family, one of the most common genus is Bacillus. In this study, Bacillus strains were the most commonly found (Fig. 3b), specifically, the station E553 had the highest abundance of strains. This is in concordance with the fact that Bacillus strains needs more nutrition (Mondol et al. 2013) and the delta of the Magdalena River influences these stations, probably contributing with organic material. The members of Bacillus genus are known to be halotolerant, alkalitolerant and/or alkaniphilic (Glaeser et al. 2013). In addition, few members of the family Planococcaceae were isolated, they belong to the genus Paenisporosarcina. The Paenisporosarcina genus has been recently described, this genus is close to the genus Sporosarcina, its members are Gram positive rods and/or cocci, non-motile, strictly aerobic, and form round endospores (Krishnamurthi et al. 2009).

The Proteobacteria and Firmicutes phylum have been reported as the most known and readily cultivable microorganisms from the marine environment (da Silva et al. 2013; Ettoumi et al. 2010; Fuhrman & Hagström 2008; Kai et al. 2017; Orcutt et al. 2011; Velmurugan et al. 2011; Zhang et al. 2018). In this study, the Proteobacteria phylum was the second in abundance, the majority of the Proteobacteria strains were isolated in the station E603 (Figs. 1, 3a), which is farthest from the coast. Proteobacteria is known to have the largest divisions within the prokaryotes, this group is divided into five classes designated as $\alpha$, $\beta$, $\gamma$, $\delta$, and $\epsilon$, also this phylum is known to have a major proportion of Gram negative organisms (Gupta 2000). Due to its halotolerance nature a great number of marine bacteria, belong to the Alteromonas, Pseudalteromonas and Halomonas species. In this study, the most common Proteobacteria isolated species belong to the Halomonas genus, this genus was first described by Vreeland et al. (1980), they are part of the Gammaproteobacteria class, and its members are described as rod-shaped, halophilic or halotolerant, aerobic or anaerobic (Vreeland et al. 1980). Finally, the third phylum found in this study is Actinobacteria. These are aerobic, Gram positive and non-motile bacteria, with high guanosine-cytosine content in their DNA (70–80%). Actinobacteria members are commonly found in terrestrial soils, but recently some
studies reported the isolation of Actinobacteria from coastal environments like mangrove swamps and deep sea sediments, as this study (Jose et al. 2021).

There are not similar reports of bacterial diversity studies in marine sediments of the Caribbean Colombian Sea using classic microbiology techniques. Nevertheless, there is a report in the southern where it was used environmental DNA technology (Franco et al. 2021). In contrast this study found the proteobacteria phylum as the most abundant followed by bacteroidetes, and firmicutes. Besides the most abundant genus were Erythrobacter (Proteobacteria) followed by Gramella (Bacteroidete) and Pseudomonas (Proteobacteria). It is important to highlight that the cultivable diversity can’t be a reference for the diversity composition, but it is still necessary to search for cultivable microorganisms under controlled laboratory conditions to obtain valuable culture collections and to study novelty metabolic adaptations that could serve for future biotechnological applications.

Regarding the marine environment extreme conditions (low temperatures, alkaline pH, high tolerance to saline conditions, and high pressure), marine microorganisms could produce unique metabolites. In this study, the isolated strains were tested for the production of biosurfactants and lipolytic enzymes production important compounds for bioremediation applications. One hundred and twenty strains were identified as lipases producers, the majority belong to the Proteobacteria phylum (29) followed by Firmicutes (23), and just 2 members of the Actinobacteria phylum a Micromonospora sp. and a Prauserella sp. (Fig. 4 and Table S4). Besides, the majority of the biosurfactant producing strains belong to the deepest station E612 (4220 m).

Biosurfactants are amphiphilic surface-active molecules commonly produced as consequence of the presence of complex water insoluble substrates like hydrocarbons, also, they are involved in other biological functions such as microbial competition and intra or inter-species communication (Ibacache-Quiroga et al. 2013). In addition to the hydrocarbons degradation, biosurfactants could replace the surfactants used in pesticides and, at the same time, improving the quality of the soil through the improvement of the bioremediation process degrading hydrocarbons as well as heavy metals (Drakontis and Amin 2020). Firmicutes members like Bacillus sp. as well as Proteobacteria species like Pseudomonas, which have been also highly reported (Mishra et al. 2021).

Furthermore, in comparison to the enzymes derived from terrestrial fonts the enzymes from marine microorganisms has some advantages like temperature and solvent stability, various catalyzing activities, high production in less time, and they are easy to harvest (Sivaperumal et al. 2017). In this sense, 56 strains were identified as lipases producers, the majority belong to the Proteobacteria phylum (29) followed by Firmicutes (23), and just 2 members of the Actinobacteria phylum a Micromonospora sp. and a Prauserella sp. (Fig. 4 and table S4). Lipolytic enzymes like lipases are hydrolases that catalyze triglycerides to fatty acids and glycerol. In marine environments, lipase production are related to the mineralization of complex organic matter, probably as consequence of the degradation of dead plant and animals (Patanla et al. 2016), lipases are useful in many industries like detergent, pharmaceutical, food, as well as in bioremediation by the degradation of contaminants like dimethylphthalate (Mita et al. 2010) and oil (Busha 2021).

In conclusion, 352 marine cultivable bacteria were isolated from deep-sediment samples from the Colombian Caribbean finding the phylum Firmicutes as the predominant, 120 were found as biosurfactant and 56 as lipases producing strains. This study highlights the fact that a great deal remains to be learned about the bacterial diversity present in these ecosystems and its potential in the bioremediation of environments contaminated with hydrocarbons.

**Author contributors** LMB and MAM: Writing-original draft preparation, MQ, MM-G and LMJ-S: Conceptualization, Methodology, data curation, MS-A and JG-L: Supervision, reviewing.

**Funding** Thanks to the National Hydrocarbon Agency—ANH and to the Marine and Coastal Research institute “José Benito Vives de Andrés”—INVEMAR for the financial support (Contribution number CTRB-1328 - from the Evaluation and Use of Marine and Coastal Resources Program - VAR, Marine Bioprospecting Line). Thanks to the information analysis laboratory—LABSIS at Marine and Coastal Research Institute “José Benito Vives de Andrés”—INVEMAR, for making the map (Fig. 1). M.A.M. acknowledges FAPESP for the Pos-Doc fellowship (2018/11910–1) and CAPES (Finance code 001). This research was carried out within the framework of the contract (Contrato marco de acceso a recursos genéticos y sus
productos derivados 123 número RGE 0158-5) granted to the Marine and Coastal Research Institute “José Bémito Vives de Andrés” –INVEMAR by the Ministerio de Ambiente y Desarrollo Sostenible de Colombia (Minambiente).

Declarations

Conflict of interest The authors did not report any potential conflict of interest. All the authors have approved the final manuscript. The authors declare that this manuscript has not been published previously.

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