Bio-weathering of granites from Eastern Dharwar Craton (India): A tango of bacterial metabolism and mineral chemistry

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Research Article

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Abstract

The bio-weathering process contributes majorly in the physical shaping of earth's surface. The weathering of mineral is coupled to the formation of new minerals and release of bio-available nutrients for flora as well as microbial communities. Granites from the Eastern Dharwar Craton (EDC) India are felsic plutonic igneous rock composed of nutritive minerals. The major elemental composition of granite is such that, it forms an ancient ecological niche for diverse microbial communities. Interdisciplinary approaches were taken to construct a more comprehensive understanding of potential functional attributes of Actinobacteria in bio-weathering of granite. The paper includes molecular characterization of the isolated strains, detecting their granite bioweathering potentials through leaching experiments where Nocardioides showed highest mineral leaching indices with iron (Fe) being the most leached element (~6372 ppm). Scanning electron microscopic imaging indicated biofilm formation and Actinobacterial hyphae colonization. Petrographic, XRD and FTIR based study shows formation of secondary minerals [kaolinite, vermiculite and smectite]. To further establish the hypothesis of soil formation and nutrition transport, long-term (360 days) microcosm was developed. Deterioration and grain size alternations with increase in clay based minerals (kaolinite, vermiculite) and total protein content was observed. A bioinformatics based functional biodiversity approach in link to global bio-weathering of rocks by the genus Actinobacteria was adopted to understand their distribution patterns and contribution to alterations of rock minerals.

Introduction

Mineral and rock weathering are the most significant biogeochemical phenomenon occurring at the interface between the earth’s endogenic and exogenic cycles. This process is an integral part of formation of soil and sediments and plays a key role in the maintenance of soil development and fertility; along with the regulation of atmospheric constitution and global climate (Uroz et al., 2009, Chikkanna and Ghosh, 2018). In addition, weathering directly affects humans by exerting an impact on water quality and durability of buildings and monuments (Mailloux et al., 2009; Brewer et al., 2018; Chikkanna and Ghosh, 2018). Transformation of bedrock through abiotic and biotic weathering processes are necessary for metabolic availability of nutrients in the earth's critical zone (Jin et al., 2010; Akob and Kusel, 2011). These processes lead to mobilization of both essential elements such as K, Na, P, Ca, Mg, Mn, Fe, Cu, Zn and pernicious elements such as Al into the terrestrial ecosystem and surface and subsurface water systems (Gadd, 2010). Moreover, in nutrient deprived environments, endolithic microbes weathering rocks, play significant role in providing nutrients to the entire food chain including primary producers (Calvaruso et al., 2006; Uroz et al., 2009). Notably, in temperate forest ecosystems which develop on acidic soils, the ectomycorrhizal fungi which lives in symbiotic association with trees, exert selective pressure on soil bacterial communities (mycorrhizosphere) which enhance primary rock weathering process through production of organic acid molecules (Van Breemen et al., 2000; Uroz et al., 2009).

Recent studies report that rock colonizing microorganisms accelerate biochemical disruption and fractionation of rock minerals through physical (hyphal growth; Fig. Graphical abstract) and chemical
processes (release of organic acids and chelating ions; Alekseeva et al., 2009; Abdulla et al., 2009; Brewer et al., 2018). However, the mechanism and rate of microbial weathering is inconsistent and dependent on certain characteristics of the rock exterior (Mapelli et al., 2012). Actinobacteria are ancient group of bacteria with inimitable ability to tolerate desiccation and to adapt and colonize in extreme environments (Ventura et al., 2007; Battistuzzi and Hedges, 2009; Abdulla et al., 2009). Their ability to decompose complex materials have contributed to the biogeochemical cycling of both essential nutrients and minerals over ancient years and thus, they might be the first bacterial phyla to have invaded the land masses (Battistuzzi et al., 2004).

Granites are abundant igneous rocks found in EDC of peninsular India, which date back to Archaean era of 3.5–2.5 Ga (Manikyamba and Kerrich, 2011). Since EDC has acted as a nutrient reservoir for large Biosphere reserves and coffee plantations of southern India, where the endolithic microbes can be the key players in sedimentation processes (Baskaran et al., 2012; Horath et al., 2009). To our knowledge, this is the first study describing the active bio-weathering of granite from EDC. The symbiotic involvement of biology and geology to understand the relationship of microbial communities in the formation of geological framework is very rare. In this report study of bio-weathering of granites from Bangalore which lies within the EDC belt, sets the stage, to understand the impact of granite minerals in nurturing the vegetation of EDC belt. The objectives of this study are to: 1) Study the role of indigenous micro-endoliths in bio-weathering of EDC- granites, 2) Understand the role of short and long term bio-leaching processes in mobilizing minor and major nutrients, and their contribution to the biogeochemical fluxes in the ecosystem. 3) Develop a comparative biogeographical model for Actinobacterial weathering of rocks.

**Materials And Methods**

**Study area and sampling procedure**

The EDC in the southern India contains abundant granite plutons with late Archaean evidence for biological weathering. The sampling site Bangalore District, Karnataka, India (N 13°01’16.15”, E 77°34’13.91”) is located at the centre of the southern Indian peninsula (Fig. 1), 934 m above sea level, only ~300 km away from Arabian sea, with a tropical climate. The average annual rainfall is 960 mm from the moisture produced in the two adjoining seas, Bay of Bengal and the Arabian Sea. It receives rainfall from both the SW and NE monsoons and the summers are more wetter than the winters (Rahul et al., 2016). According to the Köppen-Geiger climate classification, the climate in this region was considered to be Aw. The average annual temperature is ~22.9 °C or 73.1 °F and the average annual rainfall is 960 mm, with an average annual humidity of 65.16%. This site mainly forms a part of EDC which is a natural cross section of late Archaean continental nuclei. The granite samples present in various weathered conditions were identified and collected from the monoliths. The weathered samples were obtained from the surface (Fig 1. a) and non-weathered samples from 1-2 m depth from a large monolith (Fig 1. b) using sterile hammer and chisel blade, collected in separate sterile polythene bags and stored at 4 °C, prior to processing.
Petrographical and Mineralogical studies

Petrographic study of the samples was carried out in polished thin sections under a Petrological Microscope-Olympus BX51 to understand the mineral assemblages. The mineral composition of respective minerals was obtained by using JEOL JXA-8230 Electron Probe Micro Analyser (EPMA) housed at Advanced Facility for Microscopy and Microanalysis (AFMM), IISc, Bangalore. The probe (beam) current during analysis was 12 nA and acceleration voltage 15 kV. Natural and synthetic silicates and oxides were used for calibration. Standard ZAF correction procedure was used for data reduction. The rock thin sections were carbon coated prior to analysis.

Genotypic and phenotypic characterization

Enrichment of Actinobacterial strains

The collected rock samples were aseptically cut into small chips and crushed into fine powder using sterile mortar and pestle. The *Actinobacteria* isolation was carried out as previously described (Abdulla et al., 2008), using Starch Casein Agar (composition: Soluble starch 10 g/l; Casein 0.30 g/l; KNO₃ 2 g/l; MgSO₄.7H₂O 0.05 g/l; K₂HPO₄ 2 g/l; NaCl and CaCO₃ 0.02 g/l; FeSO₄.7H₂O 0.01 g/l; Agar 18 g/l) and Actinobacteria Isolation Agar (Sigma-Aldrich/Hi-Media). The plates were prepared in triplicates for each sample and were incubated in inverted condition at 30 °C for 3-5 days.

Biochemical characterization

The biochemical characterization of the isolates was detected by oxidation of 1% p-amino dimethylaniline oxalate and catalase activity using 3% (v/v) H₂O₂ solution. The carbon substrate utilization pattern of the isolates was performed with Carb Kit (HiMedia) following the manufacturer's instructions (Table 1).

Taxonomic characterization

Taxonomic characterization of the isolated strains were done using single bacterial colony by colony-polymerase chain reaction (PCR) based amplification of 16S rRNA gene using eubacterial primer Fc27 (5′-AGAGTTTGATCCTGGCTCAG-3′) and Rc1492 (5′-TACGGCTACCTTGTTACGACTT-3′; Lane, 1991). The PCR reaction mixtures and amplification conditions were carried out as previously described (Chikkanna et al., 2018). The PCR reactions (Veriti, AB) were performed in triplicates pooled down and purified using Gel Purification Kit (Qiagen, Hilden, Germany) as per manufacturer's instructions. The purified product was then sequenced in both directions using same primers in an ABI Prism 3730 Genetic Analyzer based on Big Dye Terminator chemistry (Applied Biosystems, California, United States). Sequences were checked in Bellerophon (Huber et al., 2004) for chimera and had been submitted to GenBank database. The accession numbers are MH726216 for 16S rRNA gene sequence of *Nocardioides salarius* strain BLR1 and MH726217 for *Nocardioides bigeumensis* strain BLR2.

Nucleotide sequence accession number
The GenBank/EMBL/DDBJ accession number for the 16S rRNA gene of *Nocardioides salarius* strain BLR1 is MH726216 and for *Nocardioides bigeumensis* strain BLR2 is MH726217.

**Phylogenetic analysis**

The sequences were then compared with the nucleotide databases (GenBank/EMBL/DDBJ) using NCBI BLAST program (Camacho et al., 2009). The top ten similar 16S rRNA gene sequences in the match results were selected for each isolate and used for phylogenetic representation. The sequences were aligned using Clustal Omega tool (Sievers et al., 2011). The software MEGA (Version 5.0) was used for phylogenetic analysis (Tamura et al., 2007). The phylogenetic representation for the sequences was carried out using Neighbour-joining (NJ) method (Saitou and Nei, 1987) based on Kimura 2 parameter model (Kimura, 1980) for constructing the tree. Bootstrap test of 1000 replicates was conducted to get best topology (Felsenstein, 1985). To root the phylogenetic tree, the 16S rRNA gene sequence of *Methanobrevibacter smithii* strain B181 (Acc. No. U55235) was used as an outgroup.

**DNA G + C content**

To estimate the G + C content, DNA was degraded into nucleosides, enzymatically (Mesbah et al., 1989). The degraded product was then separated in HPLC (Shimadzu) using an analytical column (Vydac 201 SP54, C18, 5 mm; 250 × 4.6 mm) equipped with a guard column (201 GD54H; Vydac; Tomaoka and Komagata, 1984) and the non-methylated lambda phage DNA (Sigma) was used as reference.

**Phospholipo Fatty acid (PLFA) characterization**

Approximately 50 mg of the freeze-dried cell pellet of the strain BLR-1 in log-phase was used to extract membrane fatty acids and fractionate them using short phase extraction column as described earlier by Ghosh et al., 2018. The methylated PLFAs were identified and quantified by gas chromatography-mass spectrometry (GC-MS; Ghosh et al., 2018). Deuterated eicosanoic acid methyl ester (d-EAME) was used as an internal standard to a spike of 10 mg/l.

**Granite bio-weathering experiments**

Granite weathering experiments for three different analyses were carried out using modified starch casein broth in triplicates for the measurements of leached out elemental concentrations and Scanning Electron Microscopy (SEM) analysis. The experiments were carried out in 500 ml Erlenmeyer culture flasks each containing 250 ml of the modified media with 10 g of granite rock powder and pH was adjusted to 7 with 10 mM NaOH. The media used for SEM analysis contained 1 cm rock chip (Pre-weighed) instead of rock powder. Those were then inoculated with previously isolated Actinobacterial strain cultures and incubated at 30 °C in a rotary shaker at 150 rpm for 40 days. The pH was monitored every 5th day using pH 213 Microprocessor (Hanna). The cells were stained with crystal violet and observed using biological microscope for mycelia growth.

**Microcosm setup**
Two sacrificial microcosms were set up as Set A: by mixing 5 kg sterilized and dried block of granite rock with 50 mL actively growing (Optical density 1; in Actinobacteria media mentioned above), mixed culture of the isolated Actinobacterial strains, and Set B (control set): mixing 5 kg sterilized dry block of granite rock with 50 mL sterile Actinobacteria media). Both the sets were sealed and incubated for 360 days in aphotic condition at 10 °C (to mimic soil zone B; Graphical abstract) and 90% relative humidity. The rock in set A became brittle, whereas the rock in set B remain unaltered is powdered for further analysis. After the incubation period, the texture (grain size), mineral composition and leached elements were measure as described above. The total protein content was measured using Bradford's assay (Bradford 1976; Halvorson and Gonzalez, 2006), which can be also interpreted as Bradford reactive soil protein (BRSP), however, the source of the protein is only the microbial hypae.

**Chemical analyses of the rock samples**

i). **X-Ray Diffraction (XRD) based mineralogical analyses:** The mineral composition of a) initial weathered section of the granite sample from bio-weathering experiment (40 days incubation) and the post microcosm incubation (360 days incubation) were determined by using XRD based irradiation of samples (Rigaku, Smart lab powder X-Ray Diffractometer). The source used during analysis was CuKa radiation, acceleration voltage of 40 kV/30 mA with D/Tex Ultra detector. The samples were finely powdered, centrifuged and < 2m particles were used to prepare the oriented slides in three sets: air dried (air-dried oriented aggregates (AO), solvated with ethylene glycol at 60 °C for 16 h (AO+EG), and heated at 500 °C for 2 h (AO+T) (Moore and Reynolds, 1989).

ii) **Fourier Transformation Infrared Spectroscopy based mineralogical analysis:** For a comparative analysis of the mineral characteristics due to long term microcosm based bio-weathering, the powdered particles were also characterized using a PerkinElmer Spectrum Two FT-IR Spectrometer.

iii) **Inductively Coupled Plasma Mass Spectrometry (ICP-MS) based leaching quantifications:** The leached-out elements were quantified using ICP-MS (Thermo Scientific X-Series II equipped with Quadrupole mass analyser and Nickel sample and skimmer cones). For this, 1 g of each sample from bio-weathering experiment was crushed using sterile agate mortar and pestle. The sample dissolution using acids was carried out as previously described (Banerjee et al., 2016). For elemental concentration determination, ~25 mg of each sample was dissolved using a mixture of 1ml of 1N HF and 1ml of 1N HNO₃ at 120 °C. The acid mixture was evaporated after 24 h of reaction and a mixture of 1ml of 1N HNO₃ and 1ml of 1N HCl was added to the samples. After another 24 h of incubation at 120 °C, the acids were evaporated, and the residue was dissolved in 2 ml of 1N HNO₃ and transferred to 125 ml pre-cleaned HDPE bottles. The internal standards were used as previously described (Banerjee et al., 2016).

To determine the concentration of the dissolved major and minor elements, highly sensitive method of ICP-MS with No-CCT mode was used. USGS rock standards BHVO-2 (Hawaiian basalt) and BCR-2 (Columbia River basalt) were used as calibration standards. The USGS rock standard AGV-2 (Andesite)
was analysed to test the accuracy of the data. All the USGS standards were processed the same way as the rock samples. The detection limits for all elements of the ICP-MS were within 0.1 mg/l.

**Bio-weathering evidences from Scanning Electron Microscopy (SEM)**

The weathered rock material was examined by SEM for the presence of Actinobacterial hyphae colonization. The chips were removed from the flasks using sterile forceps, air dried and coated with gold. They were examined using SEM JSM IT-300 instrument with accelerating voltage of 15 kV and 7-11 mm working distance. The weathered fragments were also examined using ESEM.

**Biogeographic diversity of weathering Actinobacteria**

The diversity studies were carried out by targeting previously studied Actinobacterial strains involved in environmental bio-weathering process (Supplementary Table 1). The 16S rRNA gene sequences were retrieved from the nucleotide databases (GenBank/EMBL/DDBJ). Clustal Omega tool was used to align the sequences (Sievers et al., 2011). Phylogenetic tree was constructed using similar parameters described in the above section 2.6. The diversity parameters were calculated using DOTUR Program.

**Statistical analysis of biogeography metadata**

To understand the bacterial diversity and for taxonomical analysis, the generated sequences were grouped into Operational Taxonomic Units (OTU) based on 3% cut-off at nucleotide level for 16S rRNA gene sequences using DOTUR (Schloss and Handelsman, 2005). The Shannon-Wiener (H) and Simpson indices were calculated for measurement of biodiversity of the Actinobacterial genera in environmental bio-weathering process. Phylotype frequency curves were prepared to determine the frequency of OTUs across the Actinobacterial genera. Rarefaction analysis was undertaken in DOTUR to understand the sampling biasness.

**Results**

**Petrology and mineral chemistry studies**

The coarse-grained interlocking plagioclase-K-feldspar-quartz-biotite (Pl-Kfs-Qtz-Bt) assemblage is typical of granitic rocks (Fig. 2A). Textural alterations with Actinobacterial intergrowth within plagioclase grain indicate weathering initiation (Fig 2B). The representative microprobe point (WDS) analyses of minerals are listed in Table 2. Plagioclase contains Na$_2$O which is 12.87 wt% indicating plagioclase feldspar is albite rich. In case of K-feldspar, K$_2$O is 13.33 wt%. There are other K-bearing minerals like Bt (K$_2$O – 10.89 wt% and FeO – 26.85 wt%) present in the sample. The mineral chemistry of biotite suggests that it is Fe rich annite.

**Other Analytical studies**

The biochemical alterations of granite were detected from the mineralogical compositions. Along with the primary minerals of granite (Fig. 3), determined by XRD analysis which includes quartz (3.33 Å),
plagioclase (3.1 Å), K-feldspar (3.7 Å) and biotite (10.1 Å), secondary clay minerals like kaolinite (7.1 Å), illite (9.9 Å), chlorite (14.1 Å), vermiculite (4.77 Å) and smectite (17 Å) were also detected (Fig 4). The characteristics of these minerals were further explained using FTIR analysis (Fig 5; Supplementary Table 1).

Bio-weathering and pH changes

In the bio-weathering of granite by Actinobacterial isolates, the modified starch casein broth was used to understand the change in pH over the incubation period, in the growth media. The pH of the inoculated media decreased during the initial lag phase from 7.0 to between 5.5 and 6.0. A gradual increase in pH from 6.0 to 7.5 was observed during exponential growth phase. During the stationary phase and death phase, the pH value increased between 7.5 to 8.5 and remained there (Fig 6B).

Quantitative analyses of leaching

Dissolution experiment was carried out to better understand the mechanisms of granite dissolution by Actinobacterial isolates and to measure the concentrations of leached-out elements. The concentrations (ppm) of elements such as Na, Mg, Al, P, K, Ca, Fe and Mn were measured using ICP-MS. The maximum leached out concentrations of elements was observed in cultures of isolate BLR1 and BLR2. No leaching was observed in cultures of isolate BLR3, BLR4, BLR5 and BLR6. Iron (Fe) was the most leached element with average concentration of 6372 ppm in the culture set in comparison to 7622 ppm in granite bearing control sets. In contrast, Al and K were the least precipitated elements (Table 3; Fig 6A). The elemental concentrations in the extracts for the USGS standard AGV-2 (Andesite) were compared to the 95% confidence interval (Raczek et al., 2003), and were within the confidence limit. The biochemical and carbon substrate assimilation test of the isolates is detailed in Table 1.

Microbial alteration of granite surface

The morphology of Actinobacterial hyphae growing on the granite surface were examined by light microscopy (Fig 6d). The surface alterations and microbial growth were observed through SEM and ESEM. The SEM images (Fig 6a) of experimental granite sample section, displays altered mineral grains with rough surface; Fig 6b is showing cells forming a biofilm on weathered granite sample surface. The weathered mineral fragment with mycelia growth displayed through ESEM microphotograph (Fig 6c).

After a 360 days incubation, the microcosm experiment and control setups were compared to understand the physical, chemical and biological alterations, to understand the bio-weathering aspects clearly. The experimental setup mostly disintegrated into fine clay (<0.002 mm; 0.16% w/v), silt (0.002 to 0.05 mm; 0.69 % w/v), sand (0.05 to 2 mm; 7.23% w/v), gravels (2 -75 mm; 26.31 % w/v) and small porous rocks (7.5 -10 cm; 63.2%), whereas no change was observed in the experimental setup (Supplementary Fig. 1). The XRD data of the clay fraction showed formation of secondary minerals like kaolinite and vermiculite. The average BRSP of clay fraction was 3.67 mg/cm3, which is similar to 10-20 cm deep forest soil (Halvorson and Gonzalez, 2006).
Molecular identification and taxonomy

The 16S rRNA gene sequence of the isolate BLR1 and BLR2 showed 99% identity with *Nocardioides salarius* strain J112 (Acc. No. NR_044419), and *Nocardioides bigeumensis* strain MSL-19 (Acc. No. NR_044230) respectively, at nucleotide level and inferred as *Nocardioides salarius* strain BLR1 and *Nocardioides bigeumensis* strain BLR2 (Supplementary Table 2). The phylogenetic identification of both the strains BLR1 and BLR2 indicated that they belong to the bacterial phylum *Actinobacteria*, under the class *Actinobacteria*, order *Propionibacteriales*, family *Nocardioidaceae* (Fig 7).

GC content

The GC content of strain BLR-1 was 39.63% and strain BLR-2 was 39.21%.

PLFA analysis

The predominant membrane PLFAs of both the strain BLR-1 and BLR-2 were C14:0, C16:0, C16:1, C18:0, and C18:1. The Gram positive specific fatty acids Ci15:0, Ca15:0, Ci16:0, Ci17:0 and Ca17:0 were lesser predominant and the signature PLFAs for the phylum *Actinobacteria* 10Me16:0, 10Me19:0 (Kroppenstedt 1985), were detected in both the strains.

Actinobacterial biogeography model

Diversity study of phylum *Actinobacteria* was carried out to understand their distribution patterns and involvement in bio-weathering of rock minerals (Supplementary Fig. 2). From the large metadata set our group is working on, for this study, 99 Actinobacterial 16S rRNA were selected based on their mineral interaction properties. The distribution of Actinobacterial genera within their respective order is represented in Fig 8. In the Neighbour-joining phylogenetic tree, the identified 16S rRNA gene sequences could be classified into the primary orders *Propionibacteriales*, *Micrococcales*, *Streptomycetales*, *Pseudonocardiales*, *Corynebacteriales*, *Frankiales*, *Actinomycetales*, *Geodermatophilales* and *Rubrobacteriales*. Among them, *Propionibacteriales* was the most dominant order with 31% of the sequences, followed by *Micrococcales* (20% of sequences) and *Streptomycetales* (15% of sequences; Fig 9A). The largest clade *Propionibacteriales* consisted of a sub-clade belonging to the family *Nocardioidaceae* which included sequences from this study (Strains BLR1 and BLR2), along with published cultured and uncultured Actinobacterial genera.

Under the order *Micrococcales*, four sub-clades were observed which were represented by the families *Brevibacteriaceae*, *Intrasporangiaceae*, *Microbacteriaceae* and *Micrococcaceae*, and the order *Corynebacteriales* consisted of two sub-clades represented by the families *Nocardiaceae* and *Mycobacteriaceae*. Whereas, the remaining clades consisted of sub-clades represented by single families as shown in (Fig 8).

Statistical analysis of Actinobacterial biogeography
Based on OTU distribution, all 99 Actinobacterial genera were grouped into 42 Operational Taxonomic Units (OTUs), with a cut-off of 3% at nucleotide level (Fig. 9B). The diversity indices for the Actinobacterial genera have been summarized in the Supplementary Table 3. The OTU richness of the Actinobacterial genera was studied by rarefaction analysis. Based on rarefaction analysis, the rarefaction curve of the 16S rRNA gene sequences showed under-saturation suggesting further sequencing effort could lead to detection of rare sequences (Fig. 9C). The Shannon (H’) index for all the Actinobacterial genera combined was 3.445, whereas the Simpson’s index for the sequences was found to be 0.0336 (Supplementary Table 3). The overall richness for all the sequences combined and measured with the Chao 1 estimator was found to be 63.111 (Supplementary Table 3). In the phylotype frequency curve based on total number of 16S rRNA gene sequences generated in this study, the dominant OTU (OTU-14) consisted of 11 sequences belonging to Streptomycetales (10 sequences) and Micrococcales (1 sequence). Likewise, ten sequences were represented by 1 common OTU (OTU-33) containing Pseudonocardiales (5 sequences); and Corynebacteriales belonging to Mycobacteriaceae (3 sequences) and Nocardiaceae (2 sequences). Additional OTUs contained 4 sequences (4 OTUs) out of which 1 common OTU represented the family Nocardioidaceae of the order Propionibacterales; 7 OTUs represented 3 sequences each and 8 OTUs with 2 sequences each. Whereas, 47% of the sequences were represented as singletons (20 OTUs) out of which 1 OTU represented the family Nocardioidaceae of the order Propionibacterales (Fig 9).

**Discussion**

Eastern Dharwar Craton (EDC) has witnessed various geological activities. The sampling site is an exposure, which is a part of EDC belt of granitic complex. Granites from EDC are significant mainly due to their Archaean age of existence and are one of the highly studied rock types due to the presence of minerals which are easily vulnerable to alterations. As apart, the mineral composition of these granite is such that, they form major nutrients to the major biosphere reserves in southern India (Jin et al. 2010; Akob and Kusel 2011).

The petrographic analysis by EPMA indicates the presence of major oxides such as SiO$_2$, Al$_2$O$_3$, K$_2$O, FeO, Na$_2$O, and CaO. The BSE (Back Scattered Electron) and XPL (Cross Polarized Light) (Fig. 2A) images of a non-weathered granite section shows lamellar twinning indicating the plagioclase grain and cross-hatched twinning of K-feldspar grain. The XPL (Cross Polarized Light) image (Fig. 2B) of a weathered granite section illustrates the microbial growth through pores and fractures causing chemical alterations of plagioclase mineral grain (arrow). The smooth surface grains are of quartz (BSE image; Fig. 2B).

The XRD data indicates presence of secondary clay minerals kaolinite and vermiculite in both the bio-weathered granite samples (short term incubation and microcosm study) along with the primary minerals plagioclase, K-feldspar and biotite (Fig. 4). The higher predominance of the secondary minerals like kaolinite, vermiculite, chlorite and smectite in the microcosm sample indicates recent weathering which in this study could be an indication of accelerated rates of microbial metabolic processes (acidification, leaching) during the experiment period (Fig. 4B). Such secondary minerals were not observed in the control sets (Fig. 3). This is a clear evidence of the microbial processes that occur in the lithosphere.
leading to formation of secondary minerals and soil from larger rock-types (Turpault et al., 2008). The laterite soil formation due to weathering of the EDC region, that sustains large biosphere reserves like the Nilgiri (Deepthy and Balakrishnan, 2005; Baskaran et al., 2012). It has been reported that the uplifting of Western Ghats with the onset of monsoonal climatic regime caused heavy rainfall (> 2000 mm) in the west coast plains of Indian peninsula and lesser rainfall (~ 500 mm) in the hinterland (Mysore plateau; Deepthy and Balakrishnan, 2005). It takes several thousand years for 1 m fresh rock to weather (Nahon, 1991). However, the temperature and precipitation can control the rate of chemical weathering (White and Blum, 1995). In this study we show that microbial reductive dissolution of the rocks can play a crucial role in pacing up the weathering and clay formation. Also as a biproduct also supports the biota sustaining on the soil to exchange nutrient through leaching and transport mechanism.

The FT-IR spectrum of the bio-weathered and garden soil samples (Fig. 5; Supplementary Table 1) showed the presence of hydroxyl stretching region of kaolinite at 3642 and 3695 cm\(^{-1}\) and as Al-OH at 912 cm\(^{-1}\). The large band near 1036 – 1000 cm\(^{-1}\) corresponds to Si-O stretching vibration and 995, 1012 cm\(^{-1}\) as Si-O stretch. These characteristic signals can be considered dioctahedral clay (Djomgoue et al., 2013). The hydroxyl stretching region of illite was detected at 3624 cm\(^{-1}\) and at 536 cm\(^{-1}\) and 754 cm\(^{-1}\) the Al(VI)-OSi, Al(IV)-OSi of illite and kaolinite were detected. However, doublet of quartz at 789 cm\(^{-1}\) and 778 cm\(^{-1}\) with Si-O-Si of quartz (in plane) at 694 cm\(^{-1}\) and out of plane Si=O; Fe–O out-of-Plane quartz at 685 cm\(^{-1}\) were detected in all the three samples.

Moreover, the weathering experiments also indicate of the efficiency of the Actinobacteria to leach-out minerals from the specimens, signifying their role in ecosystem dynamics. The lowering of pH of culture media from 7.0 to 5.5 within first seven days of incubation indicates the release of organic acids such as lactic, citric, oxalic and gluconic acids. Actinobacterial strains produce such acids as a strategy to extract limiting nutrients such as P, K, Ca and S from the nutrient bearing substrate rocks through chemically altering the mineral surface (Fig. 6A; Delvasto et al., 2009). The further increase in pH during the exponential growth phase (8th to 27th day; Fig. 6B) was due to the increase in K-feldspar dissolution rate, which increases with increase in pH due to consumption of H\(^+\) ions from the solution (Roden, 2008; Uroz et al., 2009). Moreover, albite dissolves at sub-neutral pH because of surface protonation, whereas, increased density of deprotonated surface results in increased mineral dissolution above pH 7 (Roden, 2008).

The organic acids, chelating substances and pH alterations influence the dissolution rate of minerals in solution (Uroz et al., 2009). Oxido-reduction, acidolysis and chelation are important metabolic processes evidenced in microbial mineral weathering (Welch et al., 2002; Uroz et al., 2009). Actinobacteria are well known producers of commercially important extracellular enzymes such as amylase and cellulase, which catalyse the hydrolysis of simple and complex carbohydrates that result in the production of organic acids (Malisorn and Nikhome, 2014). These acids directly induce mineral dissolution through: i) They adhere to mineral surface and form complexes with metal ions thereby extracting them, ii) They catalyse the release of metals by weakening the oxygen links, iii) They affect the saturation state of solution by
creating imbalance between cations and anions and induce chelation of the metal ions (Welch et al., 2002; Hamdali et al., 2008). This is also supported by the significant reduction in rock weight during the experiment, which indicates the potential of *Actinobacteria* in mineral weathering (Supplementary Table 4).

With the mesophilic climatic condition and precipitation all through out the year the laterite, clayey soil, in this region is suggested to be the weathered granites, gneisses and migmatites (Kelly et al., 2010) the role of biological factors like endolithic bacterial groups play an important role in dissolution. The bio-weathering activity was further quantified in the ICP-MS analyses. It indicated the leaching of major nutritive cations. The increased leached-out concentration of Ca, Fe, Mg and P indicates the targeted action of Actinobacterial strains on plagioclase grains (albite) and biotite Fe(II) phyllosilicate. Since, the silicate tetrahedra in quartz is completely interlocked, no weathering occurs in quartz mineral. Biotite, a Fe(II) rich trioctahedral mica is the main Fe bearing mineral in granitic rocks and most gneisses that form important source of Fe, Al and K in soils (Shelobolina et al., 2012). Studies report that the oxidation of Fe(II) in biotite (Arocena et al., 2012; Brookshaw et al., 2014) results in its alteration to smectite vermiculite and goethite. The reduced Fe concentration indicates considerable loss of Fe from the granite specimen suggesting possible Fe(II) oxidation in biotite. Feldspars are reported to be most susceptible to microbial attack than quartz due the presence of cations that react strongly with microbially produced organic acids (Song et al., 2010). However reduced action of Actinobacterial strains on orthoclase feldspar was observed in this study. Considerable leaching was seen in Ca indicating plagioclase weathering. Also, bacterial bio-weathering of granite indicates plagioclase to be more vulnerable (Song et al., 2007).

*Actinobacteria* are the dominant microbial communities of rhizosphere. The leaching activity of major nutrients from the rock types (Fig. Graphical abstract) by Actinobacterial strains clearly infers their role in bio-availing the immobile nutrients to plants in mobile forms. Such phenomenon of rock mineral leaching carried out by physical, chemical and biological agents is a key procedure for mobilization of nutritive cations from soil to tree roots (Puente et al., 2004; Gleeson et al., 2006; Gadd, 2007; Lepleux et al., 2012).

The rhizosphere (Fig. Graphical abstract) which is the immediate region around the growing roots, is localized with chemical environment that alters the weathering rates in E (leaching zone) and C (partial weathering zone) horizons (Richter et al., 2007; Johnson et al., 2005). This is the critical zone where plants take up solutes through ionic exchange and pH reduction (as low as 3) enhancing mineral decomposition rates (Berner et al., 2004). The soil profile illustrates the blueprint of soil and topography of surrounding environment. The B horizon also called the “zone of accumulation”, receives deposits of clay and iron oxides from the above A horizon. Furthermore, in the studied region, the regular precipitation all across the year leads to a high moisture content in the interface of B and C horizons, but a drier surface. The B and C horizons harbour deep tree roots which uptake leached minerals from the microbial and chemically mediated partially decomposed rocks, which is illustrated in Fig. Graphical abstract. Trees exude H⁺ or OH⁻ to equilibrate anion-cation balance in soil. The root exudates along with organic acids (ex: oxalate, acetate) and chelating ligands from microbes acidify soil and scavenge metal ions from
metal-bearing rock minerals. Owing to the presence of tight structure of silica tetrahedra in K-feldspar, it contributes in holding the K\(^+\) ions from escaping. In albite, replacement of Si\(^{4+}\) ions by Al\(^{3+}\) ions weaken the silicon tetrahedra structure, however, in biotite mica sheets, the silica tetrahedra is squeezed between Al\(^{3+}\), Mg\(^{2+}\), Fe\(^{2+}\) and K\(^+\) ions. Hence, the cations of feldspar, anorthite, albite and biotite are released into the saprolite with chemical alteration to secondary minerals like kaolinite and vermiculite (Fig. 2). In general, the weathering of granite minerals results in the formation of secondary minerals alongside leaching such as kaolinite from plagioclase and K-feldspar Eq:1, 2 and 3; vermiculite, goethite and gibbsite from biotite alterations Eq:4.

General reactions of conversion of primary granite minerals to secondary minerals.

\[
\text{2NaAlSi}_3\text{O}_8 + 3\text{H}_2\text{O} + 2\text{CO}_2 \rightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + \text{H}_2\text{SiO}_4 + 2\text{NaHCO}_3
\]

(Albite) (Kaolinite) (Silicic acid) (Sodium Bicarbonate) \hspace{1cm} \text{Eq 1}

\[
\text{CaAl}_{2}\text{Si}_{2}\text{O}_8 + 3\text{H}_2\text{O} + 2\text{CO}_2 \rightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + \text{Ca} (\text{HCO}_3)_2
\]

(Anorthite) (Kaolinite) (Calcium Bicarbonate) \hspace{1cm} \text{Eq 2}

\[
\text{2KAiSi}_3\text{O}_8 + 3\text{H}_2\text{O} + 2\text{CO}_2 \rightarrow \text{Al}_2\text{Si}_2\text{O}_5(\text{OH})_4 + \text{H}_2\text{SiO}_4 + 2\text{KHCO}_3
\]

(K-feldspar) (Kaolinite) (Silicic acid) (Potassium Bicarbonate) \hspace{1cm} \text{Eq 3}

\[
\text{K(MgFe}^{2+})(\text{SiAl})_10(\text{OH})_2 + \text{H}_4\text{SiO}_4 + \text{Mg}^{2+} + \text{H}_2\text{O} \rightarrow \text{Mg(MgFe}^{2+})(\text{SiAl})_10(\text{OH})_2 + \text{FeOOH}
\]

(Vermiculite) (Goethite)

\[
+ \text{Al(OH)}_3 + \text{H}^+ + \text{K}^+
\]

(Gibbsite) \hspace{1cm} \text{Eq 4}

The cations from the leached granite mobilize to upper horizons for plant uptake. Hence, this signifies the role of microbial rock weathering in mediating various biogeochemical cycles in the ecosystem.

The biogeographic Actinobacterial diversity studies signify the continuing role of \textit{Actinobacteria} in bio-weathering of different rock types. Dominance of \textit{Nocardioidaceae} family belonging to the order \textit{Propionibacteriales} was observed both in culture based and biogeographical diversity studies. The previous investigations indicated predominance of \textit{Nocardioides} signifying their potential role in bio-weathering of granitic rocks at St. Katherine, Egypt, through acid production, metal leaching and phosphate and sulphate solubilization (Abdulla, 2009). Consistent with this, we detected mineral leaching and acid production abilities by isolates belonging to \textit{Nocardioides} in bio-weathering of granites from
EDC. *Micrococcales* and *Streptomycetales* were the following abundant orders detected. Members of such genera have been recovered from volcanic environments suggesting their role in bio-weathering of volcanic rocks such as basaltic and obsidian glass and thus contribute in carbonate-silicate cycle (Cockell et al., 2013).

Interestingly, release of siderophore (Fe (III)-specific chelate) and catechol derivatives produced by *Streptomyces* sp. are suggested to enhance the dissolution of iron containing minerals such as hornblende (Liermann et al., 2000; Kalinowski et al., 2000). Actinobacterial groups belonging to the genus *Nocardioides*, *Streptomyces*, *Pseudonocardia* and *Nocardia* have been recovered from decayed ancient stone from a tomb in Egypt as well as in Palaeolithic rock arts, suggesting their involvement in biodeterioration of rock substrates (Groth et al., 1999; Abdulla et al., 2008).

A study on volcanic rocks in Iceland reported diverse bacterial communities with *Rubrobacteraceae* of the phylum *Actinobacteria* to be dominant within crystalline basalt rocks suggesting their ancient lineage in volcanic rock bio-weathering (Cockell et al., 2013). Interestingly, another study reported the predominance of *Rubrobacteraceae* within volcanic ash deposits of Kilauea volcano, Hawaii, U.S.A (Gomez-Alvarez et al., 2007). *Geodermatophilaceae* are an Actinobacterial family that are specialized in variety of biotopes, with dry and extreme biomes, predominately rock habitats (Nie et al., 2012; Trujillo et al., 2015). Colonization of such species on monuments and rocks in Mediterranean basin were investigated for their contribution to deterioration of rock materials (Urzi et al., 2001; Trujillo et al., 2015).

Most importantly the biogeographic study clearly indicates a global unavailability and ignorance of such an important environmental process, which bridges the gap between inorganic and organic global chemistry. *Geodermatophilaceae* are an Actinobacterial family that are specialized in variety of biotopes, with dry and extreme biomes, predominantly rock habitats (Nie et al., 2012; Trujillo et al., 2015). Colonization of such species on monuments and rock in Mediterranean basin were investigated for their contribution of deterioration of rock minerals (Urzi et al., 2001; Trujillo et al., 2015).

### Conclusion

Overall this study focuses on the bio-weathering of EDC granite by indigenous endolithic genus *Nocardioides*. The multidisciplinary approaches to design experiment adopted analytical techniques, confirm the enhanced mineral leaching from granite acting a rich nutrient source to the poor soil quality in the belt. Furthermore, the microscopic analyses revealed the Actinobacterial biofilm formation also leads surface alterations in granite minerals, forming colonizing spots for other group microorganisms. Moreover, through an approach to understand the influence of bio-weathering in various biogeochemical cycles, it can be inferred that microbial rock weathering mediates the leaching of essential nutritive cations that form the life-sustaining resources for plants as well as other life forms in the critical zone and thus, contribute in equilibration of ecosystem dynamics. In addition, the biogeographic Actinobacterial diversity study provides an ideal framework of their distribution patterns in ecosystem and their various bio-weathering potential. Besides, the biogeographic diversity study of Actinobacterial
genera in bio-weathering of monuments has an empirical significance in addressing attempts to restore monuments from biodeterioration.

**Declarations**

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**Competing interests**

The authors declare no competing interests.

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Tables

Table 1. Biochemical characterization of Nocardioides salarius strain BLR1 and for Nocardioides bigeumensis strain BLR2
| Phenotypic profiles                      | Nocardioides salarius strain |
|-----------------------------------------|-----------------------------|
|                                        | BLR-1 | BLR-2 |
| Gram Character                          | +     | +     |
| Catalase activity                       | +     | +     |
| Oxidase activity                        | +     | +     |
| Starch hydrolysis Activity              | +     | +     |
| Esculin activity                        | +     | +     |
| Nitrate reductase activity              | -     | -     |
| **Carbohydrate Utilization**            |       |       |
| Lactose                                 | -     | -     |
| Xylose                                  | +     | +     |
| Maltose                                 | +     | +     |
| Fructose                                | +     | +     |
| Dextrose                                | +     | +     |
| Galactose                               | +     | +     |
| Raffinose                               | -     | -     |
| Trehalose                               | -     | -     |
| Melibiose                               | -     | -     |
| Sucrose                                 | +     | +     |
| L-Arabinose                             | +     | +     |
| Inulin                                  | +     | +     |
| Sodium Gluconate                        | -     | -     |
| Glycerol                                | +     | +     |
| Salicin                                 | -     | -     |
| Dulcitol                                | +     | +     |
| Inositol                                | +     | +     |
| Sorbitol                                | +     | +     |
| Mannitol                                | -     | -     |
| Adonitol                                | -     | -     |
| Arabitol                                | +     | +     |
| Erythritol                              | +     | +     |
| a-methyl D-manno side                   | -     | -     |
| Xylitol                                 | +     | +     |
| ONPG                                    | +     | +     |
| D-Arabinose                             | -     | -     |
| Citrate utilization                     | +     | +     |
| Malonate utilization                    | -     | -     |
| Sorbose                                 | -     | -     |

+ indicates positive result/isolate is able to utilize the C source
- indicates negative result/isolate is unable to utilize the C source
ONPG is ortho-nitrile b-D-galactopyranoside

Table 2. Representative mineral chemistry of granite sample.
Table 3. Elemental concentrations of weathered granite in contrast with initial

| Element | Na | Mg | Al | P | K | Ca | Fe | Mn |
|---------|----|----|----|---|---|----|----|----|
| ppm     | ppm| ppm| ppm| ppm| ppm| ppm| ppm| ppm|
| Initial concentration* | 28460 | 1176 | 69210 | 271.6 | 38290 | 7055 | 7622 | 236.2 |
| Average of triplicates  | 28110 | 954.7 | 69020 | 221.6 | 38030 | 6547.5 | 6372 | 205.65 |

*concentrations prior to weathering

Figures
Figure 1

A) Geological study map of peninsular of India (Left). Field photographs of granite sample (Right) B) Highly weathered granite surface, C) Non-weathered granite surface. Note: The designations employed and the presentation of the material on this map do not imply the expression of any opinion whatsoever on the part of Research Square concerning the legal status of any country, territory, city or area or of its authorities, or concerning the delimitation of its frontiers or boundaries. This map has been provided by the authors.
Figure 2

A. (Top) XPL (Cross-Polarized Light) image of non-weathered granite (2x). (Bottom) BSE image of non-weathered granite illustrating quartz, K-feldspar and plagioclase mineral assemblage (70x). B. (Top) XPL (Cross-Polarized Light) image of weathered granite (2x) showing mineral alteration and Actinobacterial intergrowth within plagioclase grain (arrow). (Bottom) BSE image of weathered granite illustrating quartz and plagioclase mineral assemblage (75x).
Figure 3

X-ray diffraction of the source rock sample indicating primary minerals.
Figure 4

X-ray diffraction of the bio-weathered samples A) Short incubation -40 days B) Long incubation- microcosm study (360 days). The samples were treated as air-dried oriented aggregates (AO), solvated with ethylene glycol at 60 °C for 16 h (AO+EG), and heated at 500 °C for 2 h (AO+T), and analysed.
Figure 5

The FTIR transmittance vs wavelength indicating corresponding species identification in the samples: unweathered granite and bio-weathered granite -microcosm study (360 days). The garden soil sample was used as a reference to identify the clay mineral species.
Figure 6

A. SEM and ESEM photomicrographs of the mineral surface. a) Surface of mineral showing alterations (2000x), b) 242 Biofilm formation on mineral surface (10,000x), c) ESEM image of weathered fragment with mycelia growth 243 (4400x). d) Light microscopy image of Nocardioides sp. BLR2 hyphae (100x). B. Leaching of elements indicated by the increase in final setup (post-experiment) relative to initial fluxes. C. Acid dynamics during the growth phase (mean; n = 3) in of bio-weathering experimental setup.
Figure 7

Neighbour joining phylogenetic tree of 16S rRNA gene sequences of the studied Nocardioides isolates (strain BLR1 and strain BLR2 in green) and their closest cultured and uncultured relatives as deposited in GenBank. Scale bar indicates 0.05 substitutions per site. The 16S rRNA gene sequence of Methanobrevibacter smithii B181 is used as an outgroup.
Figure 8

Neighbour-joining phylogenetic tree of the overall 16S rRNA gene sequences dataset for the phylum Actinobacteria, retrieved from GenBank and generated in the current study (green). Scale bar indicates 0.05 substitutions per site. 16S rRNA gene sequence of Methanobrevibacter smithii B181 is used as an outgroup.
Figure 9

A) Phylogenetic distribution of the studied sequence dataset within their respective orders. B) Rarefaction curve from DOTUR analysis of the studied sequence dataset. C) Phylotype frequency curve of the studied sequence dataset distributed among 42 phylotypes.

Supplementary Files

This is a list of supplementary files associated with this preprint. Click to download.

- FigGA.pdf
- SupplementaryData.docx