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Phenotypic and genotypic characterization of a marine bacterium isolated from Sundarbans, Bangladesh

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ABSTRACT

The mangrove ecosystem is a largely unexplored source for bacteria with potentiality to produce biologically active secondary metabolites. Consequently, we set our research study to isolate, characterize and screen a bacterium producing bioactive compounds from marine soil of largest mangrove forest of the world, Sundarbans, Bangladesh. A total number of 39 marine bacterial colonies were isolated and purified. Representative bioactive isolate, ANAM-5 was characterized using phenotypic and genotypic procedures since it exhibited highest antibacterial activity against a series of test organisms. The 16S rDNA sequence of the strain ANAM-5 showed close similarity with only one bacterium *Planctomyces brasiliensis*, complete genome (99.64%). But the cultural, morphological, physiological and biochemical characteristics of the strain and *P. brasiliensis* were completely different. So, it may not be logical to assign the strain ANAM-5 to the genus *Planctomyces*. Thus, to confirm the taxonomic position of the strain and to have a correct solution from this taxonomic problem, further studies is needed in respect to the DNA relatedness studies, fatty acid profiling, DNA-DNA hybridization, matrix-assisted laser desorption/ionization time-of-

flight (MALDI-TOF), metabolic finger profiling using BIOLOG, ribotyping, small subunit (SSU) sequences, cell wall composition and other characterizations.

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1. Introduction

Antibiotic resistance is now recognized as a global health problem. Although many pre-existing antibiotics have been modified to yield new derivatives, bacteria have the potential to mutate known resistance mechanisms to combat these molecules (Knowles, 1997; Levy, 1998). The situation is exacerbated by the fact that no novel antibiotics have been discovered for last 20 years. For this constant increasing of bacterial resistance, it is now desperate need to search for new sources of antibiotics.

Nature is always boundless source for novel microorganisms producing bioactive compounds having antagonistic activity against pathogenic organisms. Among the bioactive metabolites producers of commercially important drugs, bacteria have proven to be a prolific source with a surprisingly small group of taxa accounting for the vast majority of compounds discovered till date (Bull, 2004). Among bacteria, actinomycetes particularly the genus *Streptomyces* are of special interest because it able to produce wide range of bioactive secondary metabolites, such as antibiotics, antitumor agents, immunosuppressive agents, antifungal, neurotogenic, anticancer, antialgal, antimalarial, anti-inflammatory and enzymes (Ravikumar et al., 2011).

But, presently, the chance of isolating a novel actinomycete strain from a terrestrial habitat, which would produce new biologically active metabolites, has reduced (Solanki et al., 2008). So, now much attention has been focused on screening of microorganism from diverse environments such as desert biomes and marine ecosystems. Marine environment harboring a vast variety of organisms differing in their physiology and adaptation capacity. The immense diversity of this habitat along with its underexploitation is the fundamental reason for attracting researchers towards it for discovering novel metabolite producers (Solanki et al., 2008). It is estimated that less than 1% of potentially useful chemicals from marine environment has been screened so far, with microbial products representing approximately 1% of the total number (Neidleman and Laskin, 2000). As marine environmental conditions are extremely different from terrestrial ones, it is summarized that marine bacteria have different characteristics from those of terrestrial counterparts and therefore might produce different types of bioactive compounds (Imada et al., 2007).

Marine bacteria are efficient producers of new secondary metabolites that show a range of biological activities including antibacterial, antifungal, anticancer, insecticidal and enzyme inhibition. Bioactive compounds from marine bacteria possess distinct chemical structures that may form the basis for synthesis of new drugs that could be used to combat resistant pathogens (Solanki et al., 2008). So, the exploration of microbial secondary metabolites from marine environment has led to the discovery of hundreds of biologically active compounds. Marine bacteria occur on the marine soil, sediments, water and also other biomass (mangrove) and substrates.

In order to explore the diversity, antimicrobial activity of marine bacteria from mangrove environments, in the present investigation, an effort was made to screen marine soils from Sundarbans, Bangladesh which are a large unscreened and diverse ecosystem for the isolation of potent antibiotic producing marine bacteria. This research study revealed that the diversity of the Sundarbans as a rich source of new and potential marine bacteria species producing bioactive secondary molecules.

2. Materials and methods

2.1. Isolation and selection of marine bacteria

From different locations of Sundarbans, Bangladesh (08-16 August, 2010), totally nine marine soil samples were collected carefully from the various depth of the earth, ranging from layers just beneath the upper surface to 1.5 meter depth. By using spread plate technique (Bernard, 2007), isolation of the marine bacteria from these soil samples were done. A total of 39 strains [ANAM-1 to ANAM-48] were isolated and purified as pure culture from these soil samples. All of these purified isolates were preserved on yeast-extract-glucose-agar slants at 40C. Then

by using streak plating technique (Alcamo and Pomerville, 2004) on yeast-extract-glucose-agar medium, all of these pure isolates were preliminary screened for antibacterial activity. The isolates ANAM-5, a brownish gray colored microorganism was chosen for further characterization since it showed highest antibacterial activity in preliminary antibacterial screening, visibility of the metabolite as well as excellent growth properties.

2.2. Morphological characterization of the selected marine bacterium

Morphology is one of the major phenotypic characteristics to identify and distinguish bacteria species. By using the method described by Shirling and Gottlieb (Shirling and Gottlieb, 1966), morphological characteristics of the strain ANAM-5 was determined. Microscopic characterization was done by cover slip culture method. In this method, three to four sterile cover slips were inserted aseptically into the sterile solidified agar medium in a petri dish at an angle of 45° to 60° by using a sterile forceps and the strain was inoculated by streak plating method, at the free space of the plate. The hyphae of the strain grew and spread on the cover slip and produced spores during incubation period (at 30.5°C for 8 days). After this incubation period, the cover slips were then taken out smoothly by using sterile forceps, mounted on slides and observed under microscope, using lactophenol cotton blue (Korzybski et al., 1967). Colonies was identified on the basis of their colony morphology and color (Shirling and Gottlieb, 1966). The characteristics of the spore-bearing hyphae and spore chains were determined by direct microscopic examination of the culture surface on opened dishes. To establish the presence or absence of chains of spores 200x – 700 x magnifications were used.

2.3. Cultural characterization of the selected marine bacterium

The cultural characteristic of the strain was also determined according to the method described by Shirling and Gottlieb (Shirling and Gottlieb, 1966). Cultural characteristics are best made on a variety of standard cultivation media such as *International Streptomyces Project* (ISP) 2 (yeast-extract-malt extract agar); ISP 3 (oatmeal agar); ISP 4 (inorganic salts-starch agar); and ISP 5 (glycerol-asparagine agar), ISP 1 (Trypton–yeast extract agar), ISP 7 (Tyrosine agar) and yeast-extract–glucose agar (YEGA). A National Bureau of Standards Color Chart was used to determine the color of the substrate mycelia, aerial mycelia, and spore mass and pigment production (Zhao et al., 2006).

2.4. Biochemical and physiological characterization of the selected marine bacterium

The biochemical and physiological characteristics of the strain were determined as described by Shirling and Gottlieb (Shirling and Gottlieb, 1966). The growth of the strain ANAM-5 was determined on different carbon sources like D-glucose, sucrose, raffinose, D (+) xylose, D (-) mannitol, fumarate, lactate, malate and pyruvate. Here, basal mineral salts agar (ISP medium 9) and different carbon sources were used for the medium of the test. The optimization of nitrogen sources such as yeast extract, peptone, L-glutamic acid, L-glutamine, gluconate, casein, beef extract, creatinine, sodium nitrate and L-asparagine was carried out by adding nitrogen source (0.2%) to the basal medium containing glucose (3%). The pH of the medium was adjusted using hydrochloric acid (1M) and sodium hydroxide (1M). For the optimization of temperature, after sterilizing and inoculating with spores the flasks were incubated at different temperatures (25-45°C).

2.5. Phylogenetic characterization of the selected marine bacterium

Phylogenetic studies were also performed. Thus, for the preparation of genomic DNA, the isolated colony from the agar plate was dispersed in 500µl of saline-EDTA buffer (NaCl 150 Mm; EDTA 10 Mm; Ph 8.2) and incubated for 1h at 37°C. Then, 10 µl of lysozyme solution (5mg/ml), 5 µl of proteinase K solution (15mg/ml) and 10 µl SDS solution (25%) were added and incubated for 30 minutes at 55°C. Lysate was extracted, purified and dissolved in nano pure sterile water (60 µl). The 16S rDNA was amplified by the PCR in reaction mixture containing KOD FX buffer with 200 Mm Dntp, 100 ng genomic DNA and 0.5 µg forward (5'-AGAGTTTGATCCTGGCTCAG-3') and 0.5 µg reverse (5'-GGTACCTGTACGACTT-3') primers. Thermal cycle was performed with a model 22331 eppendorf (Germany). The samples were subjected to an initial denaturing step consisting of 2 minutes 98°C, after which 2U of Taq polymerase was added to each sample at 90°C. The thermal profile used was 30 cycle consisting of 1 min annealing at 52°C, 2 min extension at 72°C and 1 min denaturation at 94°C. A final extension step consisting of 4 min at 65°C. PCR amplicants were detected by agarose gel electrophoresis and visualized in Dolphin WaveTech. The band was excised and isolated using TAE buffer. Purified by phenol chloroform extraction followed by ethanol precipitation and dissolved in water. The resultant PCR solution was used in sequencing reaction with

Pre-Mix and primer. The reaction mixture was applied to thermal cycles in 30 cycles consisting of 10 sec at 98°C, 30 sec at 47°C and 2 min at 68°C. DNA was purified and by using Prep-A-Gene Kit (Bio-Rad). The liquid was evaporated at reduced temperature and using vacuum pressure. The dried DNA was dissolved in 20µl of polyacrylamide sequencing gel and sequenced by Applied Biosystems automated DNA sequencer. The resulting sequences were analyzed with DDBJ/EMBL/GenBank database using analysis softwares- BLAST and BIBI (Bioinformatics Bacterial Identification Tool) (Clarridge, 2004). BIBI software is available online (<http://umr5558-sud-str1.univ-lyon1.fr/lebibi/lebibiexecx.cgi>).

3. Results and discussion

Marine environment has been proven as an outstanding and attractive resource for innovating new and potent bioactives compounds producing microorganisms. Marine microbes are particularly attractive because they have the high potency required for bioactive compounds to be effective in the marine environment, due to the diluting effect of sea water (Valli et al., 2012). This present study was designed with an aim to discover a new marine bacterium strain from marine environment which is capable of producing antibacterial secondary metabolites. Considering this, the isolate ANAM-5, a brownish gray colored microorganism was chosen for further characterization due to the highest preliminary antibacterial activity (Figure 1), visibility of the metabolite as well as excellent growth properties of the strain.

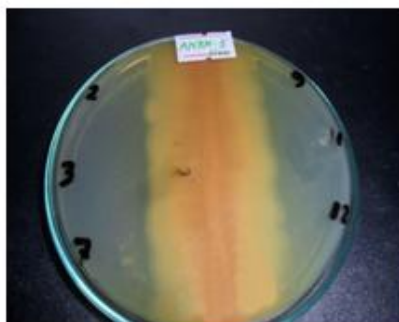


Fig. 1. Antibacterial activity of ANAM-5 isolate. The perpendicular lines marked with numerical digits are the test organism- 2. *Streptococcus agalactiae*, 3. *Bacillus cereus*, 7. *Pseudomonas aeruginosa*, 9. *Escherichia coli*, 11. *Shigella dysenteriae* and 12. *Shigella sonnei*.

The isolate ANAM-5 grew moderately at first day and after 2nd day white colonies were appeared, gradually it turned into thick and velvety cream colonies on to 3rd days. After the 3rd day ANAM-5 was started to release soluble yellow pigment through the medium. It became brownish gray (Figure 2) after the 4th day and gave deep gray colonies on 5th day. As the time passed, the aerial mycelium became more powdery as well as deep yellowish due to sporulation of the microorganism.

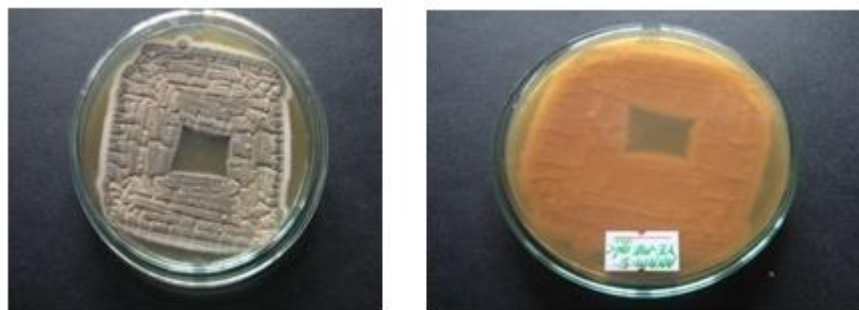


Fig. 2. Gray and brown colonies of ANAM-5.

Morphology had played a major role in distinguishing bacteria and in the characterization of bacteria species (Williams et al., 1983; Anderson and Wellington, 2001) and, in fact, it was the only characteristic used in many

early descriptions in the first few editions of Bergey's Manual. Under microscope (Figure 3), the marine bacteria ANAM-5 produce well developed substrate mycelium and aerial hyphae which are differentiated into long and extended filamentous chains with cylindrical, smooth spores occurring in chains of more than 5. Spores are observed after six days. The isolate ANAM-5 exhibited morphology of typical *Streptomyces*; the colonies were slow growing, aerobic, glabrous or chalky, folded, and with aerial and substrate mycelia (Clarridge, 2004). In addition, this isolate possessed an earthy odor and fitted to the description of genus *Streptomyces* in Bergey's Manual of Systemic Bacteriology.

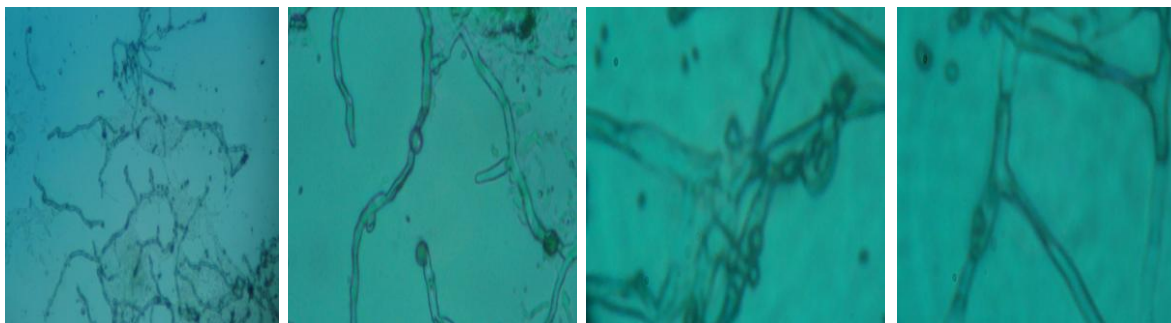


Fig. 3. Microscopic view of the ANAM-5 isolate at different magnification.

Cultural characteristics of the strain ANAM-5 in different cultural media (Table 1) showed morphology and properties of typical *Streptomyces*. The growth and pigmentation of the strain ANAM-5 was very fast in both ISP 2 and ISP 4 medium compared with both ISP 1 and ISP 5 medium where the growth of the strain was very slow. So, in ISP 2 and ISP 4 medium, the spore mass was profuse in the medium and pigmentation was very high. The growth of the strain was moderate in ISP 3, ISP 7 and YEGA medium.

The production of antibiotic by various *Streptomyces* species was greatly influenced by suitable carbon and nitrogen sources (Hobes et al., 1990). Utilization of carbon and nitrogen sources by the strain ANAM-5 is presented in Table 2. The growth as well as production of yellow pigment was good by the strain ANAM-5 in the presence of malate, sucrose and D (+) Xylose as carbon sources in the medium.

Table 1

Cultural characteristics of strain ANAM-5.

Medium	Growth	Spore mass	Aerial mycelium	Reverse mycelium	Pigmentation
Yeast-extract-malt extract agar (ISP 2)	+++	Abundant	Yellowish orange	Dark gray	+++
Oatmeal agar (ISP 3)	++	Moderate	Yellowish gray	Dark yellowish orange	++
Inorganic salt-starch agar (ISP 4)	+++	Abundant	Grayish yellow	Dark pinkish white	+++
Glycerol-asparagine agar (ISP 5)	+	Moderate	Grayish brown	Pinkish white	-
Tyrosine agar (ISP-7)	++	Moderate	Light gray	Dusky yellow	+
Trypton-yeast extract agar (ISP-1)	+	Very low	Light brown	Light yellow	-
Yeast-extract glucose agar (YEGA)	++	Moderate	Yellowish brown	Darkish pink	+

Legend: The sign '++' indicates strongly positive utilization, '+' represents positive utilization and '-' denotes negative utilization.

Utilization and metabolite production by the strain in the presence of D (+) Mannose was doubtful. In the presence of D-glucose as a sole carbon source in the medium, growth and pigmentation was moderate. We also observed that, in the presence of raffinose, pyruvate, lactate and D (-) Mannitol in the medium as carbon sources,

the growth of the strain was high but the frequency of pigmentation was moderate. In the absence of any carbon source (Negative control) the strain could not grow. Therefore, it may be concluded that, proper carbon source is essential for optimum growth of the strain and for optimum production of pigment in the medium. In case of *Streptomyces* species, with regards to carbon and nitrogen sources species specific variation may occur for cell growth and secondary metabolites production (Jonsbu et al., 2002). In this present study, we observed that the frequency of growth and production of secondary metabolites by the strain ANAM-5 was varied with the medium with different carbon sources. So, by studying the utilization of carbon sources by the strain we may conclude that the strain ANAM-5 showed the characteristics of typical *Streptomyces*.

Table 2

Utilization of carbon and nitrogen sources by the strain ANAM-5.

Carbon source	Utilization	Pigmentation	Nitrogen source	Utilization	Pigmentation
D-Glucose (Positivecontrol)	+	+	Yeast extract	++	++
No carbon (Negativecontrol)	-	-	Peptone	++	+
Raffinose	++	+	Creatinine	++	+
Pyruvate	++	+	Gluconate	++	++
Malate	++	++	L-glutamic acid	+	+
Sucrose	++	++	L-asparagine	±	-
Lactate	++	+	Beef extract	++	+
D (-) Mannitol	++	+	L-glutamine	++	++
D (+) Mannose	±	-	Sodium nitrate	-	-
D (+) Xylose	++	++	Casein	++	++

Legend: The sign '++' indicates strongly positive utilization, '+' represents positive utilization, '±' indicates doubtful utilization and '-' denotes negative utilization.

The growth of the strain as well as metabolite production was very good in the medium supplemented with yeast extract, gluconate, L-glutamine and casein as nitrogen sources. In the presence of peptone, creatinine, L-glutamic acid and beef extract as nitrogen sources, the growth was high but metabolites production of the strain were moderate. The growth and production of pigment was moderate for the strain in the medium containing L-glutamic acid. The growth of the strain was doubtful and metabolite production was negative in the medium containing L-asparagine. In the presence of sodium nitrate as nitrogen source in the medium, growth and metabolite production was negative. So, it may be concluded that, the strain used different inorganic and organic nitrogen sources differently and exhibited differences in metabolites production.

For, various *Streptomyces* species, different types of carbon and nitrogen sources were found to have significant effect on growth and secondary metabolites production (Mansour et al., 1996). Considering this, the strain ANAM-5 showed similar characteristics with *Streptomyces* species. The growth of the strain was very high and production of high levels of antibiotic when culture medium incubated at 29°C and pH 6.1. Extreme pH and temperature were unfavorable for antibiotic production for the strain. So, the morphological, cultural, biochemical and physiological properties of the strain ANAM-5 exhibited consistency with its assignment to the genus *Streptomyces*.

For confirmation and identification of the strain ANAM-5 at the species level, incongruities were observed. The 16S rDNA sequence of strain ANAM-5 was determined and compared to those deposited in GenBank using the BLAST and BIBI. Species identification was based on maximum score, identity and coverage values. The length of the sequence was 1131 bp. It was observed that the sequence was actually similar to only one sequence (Table 3).

According to BLAST, The 16S rDNA sequence of strain ANAM-5 exhibited close similarity (99.64%) with *Planctomyces brasiliensis*, complete genome. *Planctomyces brasiliensis* is a member of the order Planctomycetales was first isolated in 1982 in water from a salt pit, Lagoa Vermelha, Brazil. According to Drancourt et al (Drancourt et al., 2000), >99% and >97% sequence similarities were used as the cut-offs for species and genus identification, respectively, depending on the bacterial genus under investigation. On the other hand, Janda and Abbott in their recommended guidelines suggested that a minimum of >99%, and ideally >99.5%, sequence similarity be used as

the criteria for species identification (Janda and Abbott, 2007). They also proposed that other properties, e.g. phenotype, should be considered in final species identification. So, before assignment of a strain in a definite genus and species based on 16S rDNA sequencing there must be comparative study based on phenotypic characteristics between the respective strain and most closely related bacteria.

Table 3

Sequence producing significant alignment with the 16S rDNA sequence of ANAM-5.

Accession No.	Species/Strain	Total score	Query coverage (%)	E value	Max identity (%)
CP002546.1	<i>Planctomyces brasiliensis</i> , complete genome	2063	100%	0.0	99.64%

Since, only the sequence of the 16S rRNA gene may not be sufficient for the description of a new bacterial species, phenotypic and genotypic characters of newly described bacterial species should be compared with those of reference neighboring species and genera in order to identify precise phenotypic and genotypic characters that will discriminate the newly described bacterial species. Some basic and distinctive phenotypic characters should be included in the description of new taxons. These include conditions for optimum growth, the Gram staining value, colony morphology, motility, spore forming capacity and median measurements as obtained by electron microscopic examination. Biochemical profiles of the isolate, including the results of oxidase and catalase testing, antibiotic susceptibility profile and the ability to metabolize major carbohydrates, should also be useful for description of a taxon (Drancourt et al., 2000). Morphological features, such as capsules, flagella, cell size and shape are also used for the identification and classification of bacterial species (Rajendhran and Gunasekaran, 2011). When identifying bacteria, much attention is paid to how they grow on the media in order to identify their cultural characteristics, since different species can produce very different colonies (Christopher and Bruno, 2003). Each colony has characteristics that may be unique to it and this may be useful in the preliminary identification of a bacterial species.

For comparison based on phenotypic study, we examined *Planctomyces brasiliensis* and the strain ANAM-5, but we found important differences. *P. brasiliensis* differs from other bacterial taxa by several distinctive features such as internal cell compartmentalization, multiplication by forming buds directly from the spherical, ovoid or pear-shaped mother cell and a cell wall consisting of a proteinaceous layer rather than a peptidoglycan layer (Scheuner et al., 2014). It requires NaCl for growth and has a broad tolerance to salt (0.1 to 1.7 mol Na⁺/l). The cells are spherical to ovoid with a diameter of 0.7 to 1.8 μm which bear crateriform structures (Figure 4) and, in some cases, nonprosthecae appendages (Schlesner, 1989). The colonies have a dry, rough surface and a yellow to ochre pigmentation but no sporulation. It has a distinct cell cycle, with sessile mother cells forming stalks that attach to surfaces or to other stalks and motile daughter cells that bud from the mother cell. Mother cells are spherical to ovoid with stalks composed of twisted fibrils. The diameter of the mother cell is 1.1 to 1.5 μm. Multiplication occurs by budding on the distal cell pole, yielding daughter cells which are mono-trichously and polarly flagellated. Both the mother cell and bud have crateriform structures scattered over the whole of the cell surface (Schlesner, 1989). Cells have rigid stalk fibers and can form multicellular rosettes. Although it lacks peptidoglycan it has a cell envelope which consists almost entirely of protein which is rich in proline and cysteine and is stabilized to a high degree by disulfide bonds.

On the other hand, the strain ANAM-5 was aerobic, produce well developed substrate mycelium and aerial hyphae which are differentiated into long and extended filamentous chains with cylindrical, smooth spores occurring in chains of more than 5, colonies glabrous or chalky, folded, and with aerial and substrate mycelia and possessed an earthy odor. The strain ANAM-5 was gram positive in Gram staining technique and produce spore. So, it can be concluded that the strain has peptidoglycan in their cell structure but *P. brasiliensis* has no peptidoglycan layer in their cell structure.

P. brasiliensis grows chemoorganotrophically on the following carbon sources: D-cellobiose, D(+)-glucose, D(+)-galactose, maltose, melibiose, rhamnose, ribose, trehalose, N-acetyl glucosamine and glucuronate but it can not grows in the medium containing D(-) fructose, D-fucose, D(-) lyxose, α-D-melezitose, raffinose, L(-)-sorbose, D(+)-xylose, methanol, ethanol, glycerol, D(-)-mannitol, D(-)-sorbitol, acetate, fumarate, lactate, malate, pyruvate and succinate as carbon sources (Schlesner, 1989). On the other hand, the growth and metabolite production of

ANAM-5 was moderate to high in the presence of malate, sucrose, D (+) Xylose, D-glucose, raffinose, pyruvate, lactate and D (-) Mannitol in the medium as carbon sources (Table 2). But in the presence of D (+) Mannose in the medium, the utilization and metabolite production by the strain was doubtful. On the basis of utilization of carbon sources, it may be concluded that the strain ANAM-5 and *P. brasiliensis* are discriminative. *P. brasiliensis* can utilize ammonia, nitrate and N-acetyl glucosamine as a nitrogen source but it does not utilize L-glutamic acid, L-glutamine, L-serine amygdalin, gluconate or creatinine and does not require vitamins for growth (Schlesner, 1989). But, the growth and metabolite production of ANAM-5 was moderate to high in the presence of L-glutamic acid, L-glutamine, gluconate and creatinine as nitrogen sources in the medium (Table 2). In the presence of sodium nitrate as nitrogen source in the medium, there was no growth and metabolite production by the strain was observed. Based on utilization of nitrogen sources, it may be concluded that the strain ANAM-5 and *P. brasiliensis* exhibited differences.

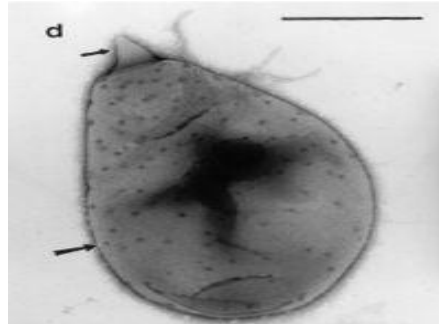


Fig. 4. *P. brasiliensis* having crateriform structures and the polar unicorn prostheca-like projection (Fuerst et al., 1997).

Although 16S rRNA sequencing has been by far the most common, reliable and convenient method of bacterial species identification (Song et al., 2003), this technique has some major limitations. The presence of mosaicism considered as one of the major limiting factor for 16S sequence based microbial identification. PCR amplification bias and cloning bias can also result in an imprecise representation of the microbial diversity. Similarly, the presence of abundant partial sequences of 16S genes in the databases results in ambiguous classification (Rajendhran and Gunasekaran, 2011).

The major difficulties and controversies in interpreting 16S rRNA gene sequence data relate to the assignment of bacterial species based on similarity search results, as no threshold values are available as for DNA–DNA hybridization (Petti, 2007). Another major limitation to its wider use is the difficulty associated with interpretation of 16S rRNA gene sequence results. The use of 16S rRNA gene sequencing for bacterial identification depends on a significant interspecies difference and a small intraspecies difference in 16S rRNA gene sequences. Therefore, one of the major limitation is that when two different bacterial species share almost the same 16S rRNA gene sequence; this technique alone would not be useful for distinguishing them confidently (Woo et al., 2009). Identification of bacterial species based on sequence analysis of the 16S rRNA gene relies on matching the obtained sequence with the existing sequence. Matching with a sequence that was incorrectly identified leads to incorrect identification (Phumudzo et al., 2013). So, 16S gene sequencing lacks taxonomic resolution at the species level for some closely related species (Janda and Abbott, 2007), subspecies, or recently diverged species (Fox et al., 1992). For this reason, *Escherichia coli* and *E. fergusonii* species, as well as the subspecies *Bartonella vinsonii subsp. arupensis* and *B. vinsonii subsp. vinsonii*, are indistinguishable, when using 16S sequence similarity comparisons (Adekambi et al., 2003).

The presence of multiple copies of the rRNA operon and intragenomic heterogeneity of the 16S genes is considered as one of the limiting factor for the use of this gene for species identification. Several recent reports have suggested the existence of divergent 16S sequences within a single organism (Rajendhran and Gunasekaran, 2011). The reported intragenomic variability of 16S was as high as 6.4% in the case of the *Thermobispora bispora* genome (Wang et al., 1997). Thus, the sequence heterogeneity within a genome can affects vigorously in the phylogenetic resolution of a particular species based on the 16S sequences. Horizontal gene transfer and recombination events have also been reported in ribosomal genes (Miller et al., 2005), which distort phylogenetic signals and thereby affect the phylogenetic classification. Horizontal transfers of 16S gene segments and the presence of mosaic like structures have been reported in *Rhizobium*, *Aeromonas*, *Bradyrhizobium*, *Streptococcus*

and actinomycetes (Eardly et al., 1996; Schouls et al., 2003). Therefore, species identification using 16S gene based probes or homology analysis of partial 16S sequences may lead to misidentification, because it may represent only a part of the mosaic like structure.

However, there are 'blind spots' within some major genera, in which 16S rDNA sequences are not sufficiently discriminative for the identification of certain species (Kwok and Chow, 2003; Kwok et al., 1999). 16S rDNA gene sequencing is highly useful in regards to bacterial classification; it has low phylogenetic power at the species level and poor discriminatory power for some genera (Gupta et al., 1983; Woo et al., 2001). In some cases, different species of bacteria with considerably different physiologies were reported to contain nearly identical 16S genes (Palys et al., 1997). For example, the 16S sequences of certain species of *Rhizobium* such as *R. galegae* were found to be more similar to *Agrobacterium* sequences than other *Rhizobium* species. On the basis of these observations, it has been proposed that the genus *Rhizobium* may be corrected as a species within the genus *Agrobacterium* (Young et al., 2001). However, this proposal was not accepted because of the ecological and genomic differences that exist between these two genera (Farrand et al., 2003). In such circumstances, species identification solely based on the 16S sequences may lead to misidentifications.

Considering these limitations, in order to increase the accuracy of 16S rDNA sequencing for identification of bacteria demands for the use of alternative methods to confirm finding. More importantly, it is necessary to interpret the results of 16S rDNA sequencing with preliminary phenotypic, biochemical and physiological test results for the confirmation upto species level. Among the alternative methods fatty acid profiling, matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF), metabolic finger profiling using BIOLOG, ribotyping, together with the computational tools employed for querying the databases that are associated with these identification tools and high-throughput genomic sequencing can use in bacterial identification (Phumudzo et al., 2013). Whole genomic DNA relatedness is considered to provide the absolute resolution in bacterial taxonomy. It is generally accepted that all taxonomic information about a bacterium is incorporated in the complete nucleotide sequence of its genome (Goris et al., 2007). Sequencing of several conserved genes (multilocus sequence analysis; MLSA) within the bacterial genome has been proposed to improve the discriminatory power (Gevers et al., 2005). The MLSA approach represents the bacterial genome to some extent and has higher discriminatory power than single locus analysis. Similarly, genomic fingerprinting method such as Rep-PCR (ERIC-, REP-, BOX-PCR), RAPD and AFLP provide more discriminatory power. However, whole genomic DNA-DNA hybridization (DDH) is considered as the "gold standard" for bacterial species identification. Organisms that show more than 70% DDH values and less than 5% difference in their melting temperature (DTm) are considered to belong to the same species (Wayne et al., 1987; Gevers et al., 2005).

Therefore, in order to prevent misinterpretation of 16S rRNA gene sequence results, they should always be interpreted in the context of basic phenotypic test results (Woo et al., 2009). Moreover, key phenotypic tests can help to distinguish between 'the closest match' by 16S rRNA gene sequencing and medically important bacterial species with similar 16S rRNA gene sequences, should be performed for differentiation of these possibilities (Woo et al., 2001; Woo et al., 2003d; Lau et al., 2002; Cloud et al., 2005). Particular attention has to be paid if there is discrepancy between 16S rRNA gene sequence and phenotypic test results. For example, 16S rRNA gene sequencing able to identify the various isolates as *Tsukamurella* to the genus level, but all *Tsukamurella* species possess >99% nucleotide identities with each other. Phenotypic tests have to be performed to distinguish among the various *Tsukamurella* species (Woo et al., 2003d).

New bacterial species are reported based on the polyphasic approach and this polyphasic approach utilizes phenotypic, genotypic and phylogenetic data of the microorganisms (Colwell, 1970; Vandamme et al., 1996; Ghosh et al., 2007). The results obtained on analyzing various phenotypic characteristics and phylogenetic studies based on 16S rDNA sequencing demonstrate that though the strain ANAM-5 is most closely related to the organism *Planctomyces brasiliensis* in phylogenetic development as well as sequence similarity but the various phenotypic characteristics of ANAM-5 like morphological, cultural, physiological and biochemical were completely different from *Planctomyces brasiliensis*. So, the consignment of the strain ANAM-5 to the genus *Planctomyces* may not be logical. Thus, further studies should be carried out in respect of the DNA relatedness studies, fatty acid profiling, DNA-DNA hybridization, small subunit (SSU) sequences, cell wall composition, matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF), metabolic finger profiling using BIOLOG, ribotyping and other characteristics to confirm the taxonomic position of the strain ANAM-5.

4. Conclusion

The marine environment represents a relatively untapped source of biologically active compounds that can be applied in pharmaceutical and cosmeceutical industries. Our findings explore the resource of a mangrove forest, Sundarbans for finding antibacterial metabolites producing marine bacterium. The strain ANAM-5 differs morphologically, culturally, physiologically and biochemically from *Planctomyces brasiliensis*. Strain ANAM-5 is phylogenetically most similar to the genus *Planctomyces*, but the phylogenetic relationship is not sufficient to permit classification of strain ANAM-5 in this genus and for the description of a new bacterial species. So, the strain ANAM-5 may not be assigned to the genus *Planctomyces*. Thus to confirm the taxonomic position and to provide absolute resolution to these taxonomic problem of the strain ANAM-5, more studies should be carried out in respect of the of the DNA relatedness studies, fatty acid profiling, DNA-DNA hybridization, small subunit (SSU) sequences, cell wall composition, matrix-assisted laser desorption/ionization time-of-flight (MALDI-TOF), metabolic finger profiling using BIOLOG, ribotyping and other characterizations.

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