The Marine Endophytic Polyamine-Producing *Streptomyces mutabilis* UAE1 Isolated From Extreme Niches in the Arabian Gulf Promotes the Performance of Mangrove (*Avicennia marina*) Seedlings Under Greenhouse Conditions

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Mangrove forests in the Arabian Gulf are under continuous threat. To increase plantations of gray mangrove (*Avicennia marina*) in the United Arab Emirates (UAE), 27 endophytic actinobacterial isolates obtained from mangrove roots were in vitro pre-screened to detect the polyamines (PAs) [putrescine (Put), spermidine (Spd), and spermine (Spm)]. We also determined the abilities of the endophytic PA-producing actinobacterial isolates in enhancing the growth of *A. marina* under greenhouse conditions. Although three highly PA-producing isolates were recovered from inside mangrove root tissues, *Streptomyces mutabilis* UAE1 constantly colonized root and stem inner tissues for 12 weeks, suggesting an endophytic association between this actinobacterial isolate and mangrove seedlings. When roots were inoculated with *S. mutabilis*, mangrove growth was remarkably enhanced under gnotobiotic and greenhouse conditions. This was evident from the significant (*P* < 0.05) increases in dry weight and length of root (66.7 and 65.5%, respectively) and shoot (64.8 and 58.0%, respectively), number of branches (64.3%), total leaf area (40.2%), and photosynthetic pigments (54.5% chlorophyll *a*; 40.0% chlorophyll *b*; and 53.1% carotenoids) of mangrove compared to the PA-non-producing *Streptomyces* sp. or control treatment. Growth promotion in plants treated with *S. mutabilis* was also supported by significant (*P* < 0.05) elevations in the contents of mangrove *in planta* PAs, auxins, and cytokinins, accompanied by a decrease in abscisic acid levels. No difference was, however, detected in growth and amounts of PAs or any plant growth regulators (PGRs) in...
INTRODUCTION

Mangroves are assemblages of salt-tolerant trees and shrubs that grow in intertidal regions of tropical and subtropical coastlines (Giri et al., 2011; Tregarot et al., 2021). Mangroves are adapted to survive hostile environments because of their specialized roots, called pneumatophores, that enable gaseous exchange and growth in deep, acidic, and anaerobic soils (Howari et al., 2009). There are approximately 84 mangrove species worldwide, of which only two species can grow in the Arabian Gulf (Almahasheer, 2018; Elmahdy et al., 2020). True mangrove (Rhizophora mucronata) and gray mangrove (Avicennia marina) can be found in Iran (Zahed et al., 2010), whereas gray mangrove is the dominant species in the Arabian Gulf countries (Burt, 2014; Elmahdy et al., 2020).

Mangroves play a critical role in supporting human well-being and providing habitat for marine organisms (Tregarot et al., 2021). In addition to acting as intense carbon (C) sinks (Donato et al., 2011), mangrove forests stabilize shorelines, protect lands, and prevent erosions (Ahmed et al., 2021). Despite their importance, mangrove forests are globally under threat (Alongi, 2002); particularly Iran and the Arabian Gulf countries including Kuwait, Bahrain, Qatar, Saudi Arabia, and the United Arab Emirates (UAE; Almahasheer, 2018; Elmahdy et al., 2020). Mangrove conservation has led the UAE to become the largest mangrove-growing country in the Arabian Gulf, covering almost 48% of the overall mangroves in this region. This is the best “fruit” of a successful plantation program undertaken by the Emirate of Abu Dhabi-UAE that was initiated 20–30 years ago.

Endophytes, microorganisms that naturally reside in tissues of living plants, are potential sources of novel natural products for exploitation in agriculture, industry, and medicine (Hallmann et al., 1997; El-Tarabily et al., 2009, 2019). Bacterial endophytes confer profound impacts on their host plants through promoting growth, increasing tolerance to pathogens, and improving phytoremediation (Ma et al., 2016; Christakis et al., 2021; Faria et al., 2021). In addition, they have the ability to produce a plethora of bioactive secondary metabolites to protect plants against environmental stresses (Khare et al., 2018; Kushwaha et al., 2020).

Plant growth promoting (PGP) bacteria (PGPB) are capable to stimulate growth via an array of mechanisms that are broadly classified as direct and indirect mechanisms, known as plant growth promoters and biological control agents (BCAs), respectively (Olanrewaju et al., 2017; Moretti et al., 2020). Direct mechanisms may enhance growth through fixation of nitrogen (N), solubilization of phosphorus (P), sequestration of iron (Fe), and production of plant growth regulators (PGRs), such as auxins, cytokinins (CKs), gibberellins (GAs), ethylene (ET), and abscisic acid (ABA) (Olanrewaju et al., 2017; Gouda et al., 2018). Indirect growth promotion occurs when PGPB reduce the deleterious effects of pathogens on plant growth (Khare et al., 2018; Bektas and Kusek, 2021; Christakis et al., 2021).

In response to environmental stresses, plants produce high levels of ET, which often inhibits plant growth (Glick, 2015; Shah et al., 2019). Many PGBP may stimulate plant growth through the activity of 1-aminoacyclopropane-1-carboxylic acid (ACC) deaminase (ACCD) that lowers ACC levels and, hence, ET amounts by breaking ACC to ammonia (NH$_3$) and α-ketobutyrate (Glick, 2015; Olanrewaju et al., 2017; Acuna et al., 2019). Accumulation of polyamines (PAs) can also trigger molecular, biochemical, and physiological responses that promote stress tolerance and increase crop yield (El-Tarabily et al., 2020).

Polyamines are low-molecular-weight aliphatic nitrogenous bases containing two or more amino groups that have potent biological activities (Xu et al., 2009; Alcázar et al., 2020). Chen et al. (2019) have shown the relationship between PAs and flowering time, embryo development, and senescence. Moreover, PAs can mediate the hormonal effects or act independently as signaling molecules in response to environmental stresses (Wu et al., 2018). Exogenous applications of PAs, mainly putrescine (Put), spermidine (Spd), and spermine (Spm), protect plants against damages caused by biotic and abiotic stresses (Handa et al., 2010; Chen et al., 2019).

To conserve mangrove ecosystems in the Arabian Gulf, maintenance and restoration of microbial communities are highly recommended (Allard et al., 2020). In the last few decades, there has been a growing interest in enhancing growth and improving production in crops by using PGPB that produce PAs (Chen et al., 2019; Alcázar et al., 2020). The rhizosphere-competent strains, Actinoplanes decacensis and Streptomyces euryhalinus, are two marine PA-synthesizing PGP actinobacteria

Keywords: endophytic actinobacteria, halophyte, mangrove, plant growth promotion, polyamines
PGPA) that promote growth and seed yields of the halotolerant plant Salicornia bigelovii in the UAE (El-Tarabily et al., 2020).

In the field of marine agriculture, most research on growth promotion of mangrove has concentrated on PGBP from the rhizosphere (Bashan and Holguin, 2002; Kathiresan and Selvam, 2006; El-Tarabily and Youssef, 2010, 2011). Increasingly, endophytic bacteria have great attributes because they are less exposed to inhospitable environments of the soil and are located in living tissues where relevant activities occur (El-Tarabily et al., 2009, 2019), thus making them ideal for the next generation of PGP agents (Preyanga et al., 2021). Except of one report in 2009, 2019), thus making them ideal for the next generation of PGP agents (Preyanga et al., 2021). Except of one report in 2009, 2019), the effect of applications of endophytic bacterial or fungi to promote growth of mangrove or other marine plants in the greenhouse/nursery/field is mostly negligible.

To the best of our knowledge, no earlier studies have addressed the effect of endophytic PGPA to stimulate growth of mangrove seedlings through production of PAs under greenhouse conditions (El-Tarabily et al., 2019), the effect of applications of endophytic bacteria or fungi to promote growth of mangrove or other marine plants in the greenhouse/nursery/field is mostly negligible.

The dark grayish-black sediment used in the current study was collected from the location described in Section “Study Site and Collection of Mangrove Propagules.” The chemical characteristics of the sediment were: Electrical conductivity = 5.81 dSm$^{-1}$; pH = 8.36 (in 0.01 M CaCl$_2$); and organic C = 6.24%. Nutrients (mg kg$^{-1}$ sediment) such as available P = 8.83, total P = 85, N as nitrate = 4 and ammonium = 6.4, bicarbonate extractable potassium (K) = 241, oxalate extractable amorphous Fe = 331, and sulfate = 414 were also detected.

### Isolation of Endophytic Actinobacteria From Mangrove Roots

All microbiological media in the present study were prepared using Millipore membrane filter-sterilized full-strength seawater. Surface-sterilized mangrove propagules (section “Study Site and Collection of Mangrove Propagules”) were sown in plastic pots (23 cm diameter × 17 cm depth) containing sediment. Pots were maintained in a greenhouse (average daily photosynthetic photon flux density of 700 ± 150 µmol m$^{-2}$ s$^{-1}$; temperature of 25 ± 2°C; and relative humidity of 60 ± 5%) and watered daily with full strength seawater. Eight pots were prepared with two propagules per pot.

After 4 weeks, seedlings were collected and transferred to the laboratory in coolers for immediate processing, and roots were cut and washed. Fresh root weight was recorded prior to further processing. Roots were soaked in sterile phosphate-buffered saline (PBS) solution, pH 7.0 for 10 min (Rennie et al., 1982). Roots were then surface-disinfested as described by Sardi et al. (1992). Sterility checks were carried out for each sample to monitor the effectiveness of the disinfection procedures (Hallmann et al., 1997; Sturz et al., 1998).

Roots were macerated as described by Hallmann et al. (1997). After filtering the slurry through sterile cotton cloth, the filtrate was serially diluted (10$^{-2}$, 10$^{-3}$, and 10$^{-4}$) and aliquots (0.2 ml) were spread with a sterile glass rod over the surface of inorganic salt starch agar (ISSA; Küster, 1959) amended with 50 µg ml$^{-1}$ of each nystatin and cycloheximide (Sigma-Aldrich Chemie GmbH, Germany). Three replicated plates for each root sample dilution were dried in a laminar flow-cabinet for 15 min before incubation at 28 ± 2°C in dark for 7 days (El-Tarabily et al., 2019). Population density (PD) of endophytic actinobacteria was calculated as log$_{10}$ colony-forming units (cfu) g$^{-1}$ fresh root weight (Hallmann et al., 1997). All colonies were purified on oatmeal agar plates supplemented with 0.1% yeast extract (OMYE; ISP medium 3; Shirling and Gottlieb, 1966). Both streptomycete and non-streptomycete actinobacteria (SA and NSA, respectively) were identified based on morphological and cultural characteristics according to Cross (1989). Hyphae and spores of all isolates were stored in 20% glycerol at −70°C (Wellington and Williams, 1977).

To determine salt tolerance of isolates, SA and NSA were streaked on triplicates on ISSA medium supplemented with NaCl concentrations up to 80 g l$^{-1}$ (8%), and incubated at 28°C in dark for 7 days (Williams et al., 1972). Isolates with strong growth and heavy sporulation on ISSA supplemented with 8% NaCl indicated high salt tolerance, and were further chosen for subsequent experiments.
Qualitative and Quantitative Determination of PAs Produced by Endophytic Isolates

Endophytic actinobacterial isolates were tested for production of Put in Moeller’s decarboxylase agar medium (MDAM) amended with 2 g L⁻¹ of L-arginine-monohydrochloride (Sigma-Aldrich) and phenol red (Sigma-Aldrich) (Arena and Manca de Nadra, 2001). Plates were incubated at 28°C in dark for 2 days (El-Tarabily et al., 2020) and dark red halo found beneath and around colonies indicated Put production by the decarboxylating isolates.

Positive actinobacterial isolates obtained from the qualitative test were further evaluated for production of Put, Spd, and Spm in Moeller’s decarboxylase broth medium (MDBM) supplemented with 2 g L⁻¹ L-arginine-monohydrochloride (Arena and Manca de Nadra, 2001) by using reverse-phase high-performance liquid chromatography (HPLC; SpectraLab Scientific Inc., ON, Canada) (Marino et al., 2000). HPLC chromatograms (eight replicates per isolate) were produced by injecting 10 μl aliquot of the sample onto a 10-μm μBondapak C₁₈ column in HPLC (Waters Associates) equipped with a 254-nm UV detector (Smith and Davies, 1985).

In vitro Assessment of PGP Activities by PA-Producing Endophytic Isolates

To detect indole-3-acetic acid (IAA), flasks containing 50 ml inorganic salt starch broth (ISSB; Küster, 1959) supplied with 5 ml of 5% filter-sterilized L-tryptophan (Sigma-Aldrich) (Khalid et al., 2004) were inoculated with 2 ml of each isolate (10⁸ cfu ml⁻¹). After 7 days of incubation on a 250 rpm orbital shaker incubator in dark (El-Tarabily et al., 2019), suspensions were centrifuged (12,000 × g), and supernatants were collected and 4 ml of Salkowski reagent was added (Gordon and Weber, 1951). IAA equivalents (μg ml⁻¹) were quantitatively determined at 530 nm using spectrophotometer (UV-2101/3101 PC; Shimadzu Corporation, Analytical Instruments Division, Kyoto, Japan).

Isolates were also grown in glucose peptone broth (di Menna, 1957) supplemented with 5 ml of 5% L-tryptophan to detect IAA and indole-3-pyruvic acid (IPYA), and on medium developed by Strzelczyk and Pokoj sks-Burdziej (1984) to detect GA₃ and CKs [isopentenyl adenine (iPa), isopentenyl adenosine (iPA), and zeatin (Z)] using HPLC (Tien et al., 1979). After 10 days, extraction of PGRs from concentrated filter-sterilized cell-free broth and HPLC parameters were used to determine the concentrations of IAA, IPYA, GA₃, iPa, iPA, and Z according to Tien et al. (1979).

To screen for ACCD production, isolates were grown on ACCD agar medium (Pikovskaya, 1948) amended with bromophenol blue as an indicator in which tricalcium phosphate was replaced with insoluble rock phosphate. The disappearance of the blue color and production of clear zone beneath culture was an indicator of P-solubilization. The drop of pH and the amount of released soluble P (Murphy and Riley, 1962) in modified National Botanical Research Institute’s phosphate broth (Nautiyal, 1999) were taken as an index for the efficiency of the strains to solubilize P.

To measure nitrogenase activity and NH₃ production, acetylene-reduction assays (Dye, 1962) and Nessler's reagent (Holguin et al., 1992) were used, respectively. For siderophores production, plates of chrome azurol S agar (Schwyn and Neiland, 1987) were inoculated with isolates and incubated for 3 days at 28 ± 2°C in dark. As siderophore producers, actinobacterial isolates developed yellow-orange halo zone around the colony. Eight independent replicates for each strain were used in all these in vitro assays.

Assessment of PGP Parameters Under Gnotobiotic Conditions

To evaluate PGP abilities of isolates under gnotobiotic conditions, surface-sterilized mangrove propagules were sown in plastic pots containing sediment. Pots were maintained in a greenhouse (section “Isolation of Endophytic Actinobacteria From Mangrove Roots”) for 10 days and watered daily with full strength seawater. After 10 days, the pruned-root dip method (Musson et al., 1995) was used to introduce endophytic actinobacterial isolates inside the young seedlings. Root tips (3 mm) were trimmed using a sterilized scalpel to facilitate uptaking the actinobacterial inoculum. Young seedlings were placed in sterile plastic cups for 3 h at 25°C with their roots in direct contact with the inoculum suspension of each isolate (10⁸ cfu ml⁻¹).

Mangrove seedlings with or without the actinobacterial inoculum were then aseptically planted into glass tubes (300 × 35 mm in diameter) filled with sediment and moistened with seawater. Seedlings, serving as control treatment, were treated with autoclaved ISSB.

Seedlings were daily irrigated with full strength sterilized seawater and maintained in a growth chamber (16/8-h day of 180–200 μmol m⁻² s⁻¹ fluorescent light/dark; 25/20°C light/dark temperature cycle). Plants were harvested after 6 weeks of transplantation, washed, and separated into roots and shoots. Measurements of dry weight (DW; g) and length (cm) of shoot and root tissues were determined. Each treatment was independently replicated eight times with one seedling in each replicate.

Quantification of Internal Colonization by PA-Producing Actinobacterial Isolates

Mutants resistant to rifampicin of the three promising PA-producing isolates, along with the positive control PA-non-producing isolate, were selected on ISSA medium supplemented with rifampicin (100 µg ml⁻¹; Sigma-Aldrich) and tested...
made in PBS and spread on ISSA. After 5 days of incubation, cells were centrifuged, inoculated into 250-ml ISSB and shaken at 250 rpm on an orbital shaker incubator for 5 days. Cells were centrifuged in the Greenhouse”) experiments, 4 ml aliquots of 20% inoculum of each strain were used as an inoculum.

To assess colonization of internal root and stem tissues, the pruned-root dip method (section “Assessment of PGP Parameters Under Gnotobiotic Conditions”) was used to inoculate 10-day-old mangrove seedlings with individual isolates. Free draining pots (36-cm in diameter), filled with 14 kg of sediments collected from the area described in Section “Study Site and Collection of Mangrove Propagules,” were placed in a greenhouse (section “Isolation of Endophytic Actinobacteria From Mangrove Roots”) where seedlings were watered daily with full strength seawater to container capacity. After planting, roots and stems were sampled weekly (for 12 weeks), washed thoroughly with water, and surface-sterilized as described in Section “Study Site and Collection of Mangrove Propagules.” Root and stem samples were homogenized and treated as in Section “Isolation of Endophytic Actinobacteria From Mangrove Roots” in order to determine the PD of the four isolates on ISSA amended with rifampicin. Each treatment was replicated according to Misaghi and Donndelinger (1990). These mutants were also compared to the corresponding wild-type strains for PA production. Morphological features, growth rates, and PA production of mutants were similar to those of parental strains.

For the gnotobiotic (section “Assessment of PGP Parameters Under Gnotobiotic Conditions”) and greenhouse (section “In vivo Assessment of Growth Promotion and Endogenous PGRs in the Greenhouse”) experiments, 4 ml aliquots of 20% glycerol suspension of the four endophytes were individually inoculated into 250-ml ISSB and shaken at 250 rpm on an orbital shaker incubator for 5 days. Cells were centrifuged (12,000 × g) at 20°C for 15 min, and the pellet was suspended in 10 ml PBS and re-centrifuged (El-Tarabily et al., 2019). For each suspension, 0.1 ml of each 10^−3, 10^−4, 10^−5, and 10^−6 dilutions was made in PBS and spread on ISSA. After 5 days of incubation, a final concentration of ~10^8 cfu ml^−1 of each isolate was used as an inoculum.

TABLE 1 | In vitro production of the free PAs by selected halotolerant endophytic actinobacterial isolates.

| Isolate^a | Tolerance to NaCl (%) | Free PA^b |
|-----------|----------------------|-----------|
|           | Put                  | Spd       | Spm       |
| #1        | + 546.42 ± 17.54 a    | 168.04 ± 7.22 a | 48.24 ± 4.48 a |
| #4        | + 274.84 ± 7.68 f    | 71.58 ± 5.96 de | 13.44 ± 2.20 d  |
| #5^c      | + 0.00 ± 0.00 i     | 0.00 ± 0.00 g  | 0.00 ± 0.00 e   |
| #7        | + 159.12 ± 7.28 h    | 81.88 ± 7.40 d | 0.00 ± 0.00 e   |
| #8        | + 361.38 ± 8.92 e    | 63.10 ± 4.42 e | 24.68 ± 3.10 c  |
| #9        | + 370.16 ± 7.68 e    | 56.23 ± 3.84 e | 14.76 ± 4.06 e  |
| #12       | + 104.66 ± 17.18 f   | 0.00 ± 0.00 g  | 14.46 ± 2.80 d  |
| #14       | + 510.06 ± 8.96 b    | 141.28 ± 4.44 b | 45.38 ± 2.18 a  |
| #16       | + 198.48 ± 18.60 g   | 44.42 ± 3.12 f | 12.52 ± 2.74 d  |
| #19       | + 163.64 ± 5.56 h    | 0.00 ± 0.00 g  | 0.00 ± 0.00 e   |
| #20       | + 406.60 ± 9.68 d    | 113.13 ± 5.36 c | 34.82 ± 2.66 b  |
| #22       | + 36.42 ± 3.24 j     | 0.00 ± 0.00 g  | 0.00 ± 0.00 e   |
| #24^c     | + 543.48 ± 15.72 a   | 174.36 ± 8.16 a | 47.32 ± 3.52 a  |
| #26       | + 440.84 ± 6.32 c    | 84.33 ± 5.50 d | 23.65 ± 3.98 c  |
| #27       | + 551.58 ± 6.52 a    | 0.00 ± 0.00 g  | 0.00 ± 0.00 e   |

^aIsolates #4, #9, #12, #14, #16, #20, #22, #24, and #27 belonged to SA, while #1, #7, #8, #19, and #26 belonged to NSA, #1, #4, #8, #9, #14, #16, #20, #24, and #26 represent PA-producing isolates and were used in further experiments, while isolates #7, #12, #19, and #22, were not used in subsequent studies. ^bFree PAs (mg l^−1) were measured after 10 days of incubation at 28°C in MDMB amended with L-arginine-monoiodochloride. ^cIsolate #24 represents the endophytic PA-producing Streptomyces lutabilis UAE1, while #5 represents the endophytic PA-non-producing Streptomyces sp. 19-388. Values are means ± SE of eight independent replicates and values with the same letter within a column are not significantly (P > 0.05) different according to Fisher’s protected LSD test. PA, polyamine; Put, putrescine; Spd, spermidine; Spm, spermine; MDMB, Moeller’s decarboxylase broth medium; SA, streptomyccete actinobacteria; NSA, non-streptomyccete actinobacteria.
eight times and each replicate was determined by a single pot containing one seedling.

For light and transmission electron microscopy (TEM), root and stem samples of 6-week-old mangrove seedlings inoculated with PA-producing isolates or with the positive control PA-non-producing isolate were fixed with karlovsky’s fixative in 0.17 M phosphate buffer at pH 7.2 containing 2.5 glutaraldehyde (Sigma-Aldrich) and 2% paraformaldehyde (Sigma-Aldrich) for 24 h at 4°C. Tissues were rinsed three times in 0.17 M phosphate buffer (pH 7.2) and post-fixed with 1% aqueous osmium tetroxide (Sigma-Aldrich) and 2% paraformaldehyde (Sigma-Aldrich) for 24 h at 25°C. Tissues were dehydrated with ascending grades of ethanol (30–100%) and dipped into the propylene oxide (Sigma-Aldrich). Finally, tissue samples were infiltrated and embedded in epoxy resin (Epon 812, Agar Scientific, United Kingdom) and polymerized at 60°C in embedding oven for 24 h (Millonig, 1976).

Tissue blocks were trimmed, where semi-thin sections (1.5 µm) and ultra-thin sections (95-nm) were cut with Leica EM7 ultra micromote (Vienna, Austria). Heat-dried semi-thin sections of root and stem specimen were stained with a mixture of 1% toluidine blue and borax (Sigma-Aldrich). For light microscopic (LM) analysis, slides were observed using Olympus BH-2 (Olympus Optical Co. Ltd., Japan) LM equipped with an LM Digital Camera and Software (Jenoptik ProgRes Camera, C12plus, Germany). Ultra-thin sections (90-nm) were then collected on 200 mesh Cu grids and contrasted with 10% uranyl acetate, followed by 3% lead citrate. Finally, the grids were examined under Tecnai Spirit G2 Biotwin TEM operating at 80 kV (FEI Co., Indore, Netherlands).

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**Identification of the Most Promising Endophytic PA-Producing Actinobacterial Isolate**

The 16S rRNA gene sequencing analysis was performed by Deutsche Sammlung von Mikroorganismen und Zellkulturen GmbH (DSMZ)-Germany. Universal 16S rRNA targeting primers: 900R (5′-CGGTCAATTCATTTGAGTTT-3′); 357F (5′-TACGGGAGGCAGCAG-3′); and 800F (5′-ATTAGATACCCGTGTA-3′) (Rainey et al., 1996; Saeed et al., 2017; Kamil et al., 2018) were used. The 16S rRNA gene sequences of all representatives of this genus were retrieved from the NCBI database1 and aligned with CLUSTAL-X (Thompson et al., 1997) provided in Molecular Evolutionary Genetics Analysis 7.0 (MEGA7; Kumar et al., 2016). The phylogenetic tree was constructed using the maximum-likelihood (ML) method (Felsenstein, 1981), and consistency was investigated by 1,000 resamplings (bootstrap analysis).

Spore chain morphology and spore surface were viewed using scanning electron microscopy (SEM) manufactured by Philips XL-30 SEM (FEI Co., Netherlands).

**In vivo Assessment of Growth Promotion and Endogenous PGRs in the Greenhouse**

Free draining pots (36 cm in diameter) filled with 14 kg of mangrove sediments were collected as described in Section 1

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**TABLE 2** | In vitro production of PGRs (µg ml⁻¹), ACCD (nanomoles α-keto-butyrate mg⁻¹ protein h⁻¹), siderophores, nitrogenase enzyme, and NH₃ and effect of inoculation with RP amended Pikovskaya’s broth by halotolerant PA-producing actinobacteria isolated from mangrove roots on pH levels and concentration of available P (mg l⁻¹).

| Activity | Isolate³ |
|----------|----------|
|          | #1       | #4       | #5       | #8       | #9       | #14      | #16      | #20      | #24      | #26      |
| Production of |          |          |          |          |          |          |          |          |          |          |
| IAA equivalents | –        | 15.67 a   | –        | –        | –        | –        | –        | 8.25 b   | –        | –        | 21.25 c  |
| IAA        | –        | 37.25 a   | –        | –        | –        | –        | –        | 19.25 b  | –        | –        | 55.36 c  |
| IPVA       | –        | –        | –        | –        | –        | –        | –        | 8.24 a   | –        | –        | 15.36 b  |
| GA₃        | –        | –        | –        | –        | –        | –        | –        | –        | –        | –        | 8.14     |
| iP         | –        | 2.85 a   | –        | –        | –        | 4.36 b   | –        | 1.25 c   | –        | –        | 7.25 d   |
| iP9        | –        | 1.15 a   | –        | –        | –        | 1.30 a   | –        | 3.54 b   | –        | –        | 4.36 c   |
| Z          | –        | 2.54 a   | –        | –        | –        | –        | –        | 4.69 b   | –        | –        | 1.12 c   |
| ACCD       | –        | –        | –        | 312.25 a | –        | –        | 452.36 b | –        | –        | –        |
| Siderophores | +        | –        | +        | –        | –        | +        | –        | –        | +        | –        |
| Nitrogenase | –        | +        | –        | –        | –        | –        | –        | +        | –        | –        |
| NH₃        | –        | –        | –        | –        | –        | –        | –        | –        | +        | –        |
| Solubilization of P | +        | –        | –        | +        | +        | –        | –        | –        | –        | –        |
| P concentration (control 13.11) | 62.36 a | –        | –        | 25.25 b  | 324.3 c  | –        | –        | –        | –        | –        |
| pH (Control = 7.72) | 5.67 a  | –        | –        | 6.45 b   | 3.15 c   | –        | –        | –        | –        | –        |

³All isolates, except #5 (Streptomyces sp. 19-388), were considered endophytic PA-producing actinobacteria. Isolates #1, #6, #18, and #24 (S. mutabilis UAE1) solely produced PA but not any other PGRs, and were further tested.

Data were from eight independent replicates. Values with the same letter within a row are not significantly (P > 0.05) different according to Fisher’s protected LSD test.

PGRs, plant growth regulators; ACCD, 1-aminocyclopropane-1-carboxylic acid deaminase; NH₃, ammonia; RP, rock phosphate; PA, polyamine; P, phosphorus; IAA, indole-3-acetic acid; IPVA, indole-3-pyruvic acid; GA₃, gibberellic acid; iP, isopentenyl adenine; iP9, isopentenyl adenoside; Z, zeatin; –, not producing/effective; +, producing/effective.
“Study Site and Collection of Mangrove Propagules.” Inoculated seedlings with the promising PA-producing and the PA-non-producing endophytic isolates were prepared using the pruned-root dip method (section “Assessment of PGP Parameters Under Gnotobiotic Conditions”). A total of three treatments were carried out: (1) Negative control (seedlings inoculated with autoclaved ISSB medium; only no isolate); (2) positive control (seedlings inoculated with the PA-non-producing isolate); and (3) inoculated seedlings with the PA-producing isolate. Each treatment was replicated eight times and each replicate was determined by a single pot containing one seedling in a randomized complete block design (RCBD). Greenhouse trials were conducted in 2018 and were independently repeated in 2019.

Seedlings were placed in the greenhouse (section “Isolation of Endophytic Actinobacteria From Mangrove Roots”) and watered daily with full strength seawater to container capacity. Measurements of DW and length of roots and shoots, the number of branches, and leaf surface area (cm²) were followed monthly and recorded at 9 months post planting (mpp) of propagules. Fluorescence emission of chlorophyll (chl) a and chl b was measured at 654 and 663 nm, respectively (Holden, 1965). Carotenoids were estimated at 470 nm (Davies, 1965).

Put, Spd, and Spm were extracted from tissues of apical parts of root and shoot tissues (Flores and Galston, 1982). Quantitative determination of PAs was performed using benzoyl chloride (Sigma-Aldrich) and normal internal standard of PAs (Redmond and Tseng, 1979). Reverse-phase HPLC chromatograms were produced onto a 10-μm reverse-phase column as previously described (section “Qualitative and Quantitative Determination of PAs Produced by Endophytic Isolates”).

Endogenous auxins and ABA were extracted from mangrove root and shoot as previously described by Guinn et al. (1986), whereas GA3 and CKs were extracted from the same tissues according to Shindy and Smith (1975) and Machácková et al. (1993), respectively. HPLC parameters were applied using reverse-phase HPLC (Waters Associates) with a 254-nm UV detector. Concentrations of endogenous PGRs were obtained by comparing peak areas in unknown samples with the corresponding areas obtained in internal standard samples (Sigma-Alrich) of known concentrations.

The extraction of endogenous ACC was carried out using the method described by Lizada and Yang (1979). Derivatization of the ACC was carried out by adding phenylisothiocyanate (Sigma-Alrich), and the reverse-phase HPLC chromatograms were produced as described by Lanneluc-Sanson et al. (1986).

All extraction procedures of PAs and other PGRs were conducted on seedlings at 9 mpp of propagules. Eight replicates from independent samples were analyzed for chl a, chl b, carotenoids, PA, IAA, IPYA, ABA, GA3, CKs, and ACC.

### Statistical Analyses

All experiments were performed as RCBD. Gnotobiotic and greenhouse experiments were repeated with similar results; and data were combined and analyzed. All data were subjected to analysis of variance (ANOVA) using SAS Software version 9 (SAS Institute Inc., NC, United States). Mean values of treatments were compared using Fisher’s protected least significant difference (LSD) test at P = 0.05 levels. Actinobacterial PD was transformed into log₁₀ cfu g⁻¹ fresh root or stem weight.

### RESULTS

**In vitro Evaluation of PAs and Other PGR Activities of Endophytic Actinobacterial Isolates**

A total of 27 endophytic actinobacteria were isolated from the surface-sterilized root samples of which 23 (85.2%) isolates grew and heavily sporulated on ISSA plates containing 8% NaCl (Figure 1A), as a relevant indicator of salinity tolerance of certain isolates (Table 1). Four isolates were not chosen for further experiments because they did not grow on ISSA containing 8% NaCl (Figure 1A).

The highly salt-tolerant isolates were tested if they could produce Put in MDAM plates amended with l-arginine (Table 1). Fourteen of them, representing 51.9% of the total actinobacterial endophytes, were able to produce Put in vitro. This was clearly observed when isolates showed relatively moderate to dark red halo surrounding or beneath the colonies (Figure 1B), suggesting that these particular isolates could be considered as potential Put-producing endophytic halotolerant actinobacteria. The other nine isolates showing no or bright red halo in culture media were either non-Put producers or producers of little amount of Put and thus were excluded from subsequent tests.

To determine if they were able to produce other PAs in vitro, quantitative assays were carried out on culture extracts of the 14 isolates. The amounts of Put, Spd, and Spm synthesized by these isolates significantly (P < 0.05) varied (Table 1). According to HPLC analysis, isolates #7 and #12 produced Put and either Spd or Spm, respectively. Three isolates (#19, #22, and #27) produced Put only (Table 1) and were excluded from further studies. The remaining isolates #1, #4, #8, #9, #14, #16, #20, #24, and #26 were positive controls for determination of PAs.

| Treatment | Length (cm) | Dry weight (g) |
|-----------|-------------|----------------|
|           | Root        | Shoot          | Root          | Shoot          |
| Control   | 6.04 ± 0.43 d | 10.76 ± 0.61 a | 1.18 ± 0.11 c | 2.88 ± 0.09 d |
| #1        | 14.56 ± 0.54 b | 15.86 ± 0.24 b | 4.80 ± 0.24 a | 4.68 ± 0.34 b |
| #5        | 6.26 ± 0.19 d | 11.83 ± 0.33 c | 1.24 ± 0.14 c | 3.01 ± 0.12 d |
| #8        | 9.22 ± 0.14 c | 13.18 ± 0.46 c | 2.02 ± 0.10 b | 3.90 ± 0.20 c |
| #14       | 17.60 ± 0.28 a | 19.38 ± 0.84 a | 4.92 ± 0.16 a | 4.84 ± 0.08 b |
| #24       | 15.40 ± 0.66 ab | 20.38 ± 0.64 a | 5.03 ± 0.19 a | 6.97 ± 0.29 a |

Table 3: Effect of endophytic PA-producing actinobacterial isolates on growth parameters of mangrove (Avicennia marina) under gnotobiotic conditions.

*Isolate #5 represents the endophytic PA-non-producing Streptomyces sp. 19-388, while #24 represents the endophytic PA-producing Streptomyces mutabilis UAE1.*

Values are means ± SE of eight replicates for each treatment. Values with the same letter for each growth measurement within a column are not significantly (P > 0.05) different according to Fisher’s protected LSD test. Plants were harvested after 6 weeks.

PA, polyamine.
produced all three PAs and were further tested to synthesize other PGRs in vitro. Furthermore, three isolates (#4, #16, and #26) produced detectable levels of IAA and/or IPYA; and four (#4, #9, #16, and #26) produced different levels of iPa, iPA, and Z (Table 2). Only isolate #26 produced GA3 in the culture extracts. We also assessed the production of ACCD and nitrogenase by PGPA in vitro, as these enzymes are known for their PGP activities (El-Tarabily et al., 2019). Although isolate #20 produced both enzymes; isolates #9 and #4 produced only ACCD and nitrogenase, respectively (Table 2). Only isolates #1, #8, and #9 were able to solubilize P. Because we ought to find isolates only producing the three PAs regardless of any other PGP activities, all the above-mentioned isolates were not included in the gnotobiotic-related experiments.

Our results demonstrated that there were no detectable levels of PGRs, ACCD, and nitrogenase enzymes or P solubilization in the culture extracts of the PA-producing isolates #1, #8, #14, and #24 (Table 2 and Supplementary Figure 1). Except of isolate #8, the other four isolates produced siderophores (Table 2 and Figure 1C), suggesting that these isolates are mainly dependent on PA production.

**Evaluation of PGP Features in Mangrove Seedlings Under Gnotobiotic Conditions**

In addition to the endophytic actinobacterial isolates producing high levels of PAs (#1, #8, #14, and #24), the endophytic PA-non-producing isolate #5 (positive control) was further screened for growth promotion activities on mangrove roots under controlled-gnotobiotic conditions. The four endophytic isolates generally showed variable effects on root and shoot growth. First, we noticed that inoculation of mangrove roots with isolates #14 and #24 resulted in maximum significant (P < 0.05) increase in root and stem lengths, which was 60.8–65.7 and 44.5–47.2% higher than non-inoculated plants, respectively (Table 3). To a lesser extent, mangrove seedlings initially inoculated with isolate #1 showed increases by 58.5% in root and 32.2% in stem lengths.

Second, the same endophytic isolates #14 and #24 significantly (P < 0.05) increased DW of roots (76.0 and 76.5%, respectively) and shoots (40.5 and 58.7%, respectively) compared to non-inoculated treatments (Table 3). Isolate #1 also showed approximately similar seedling DW of roots (75.4%) and shoots (62.5%) as those inoculated with the superior isolates.

The measured root and stem lengths and weights of seedlings treated with the endophytic PA-producer (#8) were significantly (P < 0.05) higher than those non-treated or treated with the PA-non-producing isolate #5, but the measurements were significantly (P < 0.05) less than in the other tested PA-producing isolates (Table 3). Accordingly, isolate #8 was not selected for further experiments. Thus, isolates #1, #14, and #24 were tested in the greenhouse. All tested PGP features using isolate #5 were comparable to those of non-inoculated mangrove plants (Table 3).

**Endophytic Colonization of Tissues by Selected PA-Producing Actinobacterial Isolates**

In general, the potent PA-producing isolates (#1, #14, and #24) recovered at all samplings on a biweekly basis up to 12 weeks, from internal root and stem tissues of mangrove,
thus maintaining plant health as well as their endophytic nature. Except of #14, the total population of isolates increased significantly \((P < 0.05)\) in all examined weeks in root and stem tissues (Figure 2). We noticed a significant \((P < 0.05)\) increase until week 6 inside both tissues, followed by a drop in PD of isolate #14 starting week 10 in roots and 8 in stems until the end of the experiment. This indicates that isolate #14 did not sufficiently recover from tissues in week 6 onward. Based on our results, colonization of isolate #1 inside stems was limited. This was evident from the insignificant \((P > 0.05)\) mean of total population in this isolate in mangrove stems between week 2–6 and week 6–12 (Figure 2). This suggests that this isolate does not colonize stem tissues efficiently. Therefore, isolates #1 and #14 were not included in the subsequent greenhouse trials.

For isolate #24, PD dramatically increased for the tested period of up to 12 weeks of root and stem colonization (Figure 2), thus suggesting a beneficial plant–microbe interaction, i.e., plant growth promotion. From the beginning to the end of the colonization examined weeks, similar increase patterns in tissue PD of the PA-non-producing isolate #5 were observed, making it a perfect positive control for the greenhouse experiments in comparison to the finally selected PA-producing isolate #24.

We also carried out microscopy examination on the tissue-associated actinobacterial isolate #24. At 6 weeks post inoculation (wpi), spores of isolate #24 were abundantly present within parenchyma cells of cortex and xylem of mangrove-inoculated roots (Figure 3). Mycelial growth carrying the spiral spore chain of isolate #24 within cortical cells of roots and stems was also detected. Semi-thin sections revealed that the endophytic actinobacterial isolate #24 could successfully colonize intracellular spaces between root cells and intercellularly within xylem vessels, indicating its abilities to translocate between cortical and xylem conducting tissues (Supplementary Figure 2). We also figured out that many spores belonging to isolate #24 were intracellularly found in the process of germination and formation of germ tubes (Supplementary Figure 2). Not

FIGURE 3 | Colonization of mangrove tissues by the endophytic isolate #24. Light micrograph of semi-thin sections of gray mangrove (A) root and (B) stem tissues not inoculated (control; top); or inoculated with isolate #24 (treated; middle); and spiral chain of spores (close-up; bottom) of isolate #24 within treated tissues (1,000×). In (A, B), all sections were stained with 0.1% toluidine blue showing the distribution of mycelium and spiral chain of spores (red arrows) in root and stem tissues within mangrove cells. Isolate #24 represents the PA-producing Streptomyces mutabilis UAE1. PA, polyamine. Bars: 20 µm.
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**FIGURE 4** | Inter- and intra-cellular colonization of mangrove tissues by the endophytic isolate #24. Transmission electron micrograph of ultra-thin sections of a 6-week-old gray mangrove (A) non-inoculated (control; 6,000×) and (B) inoculated (treated; 6,000×) root (top) and stem (bottom) tissues with *Streptomyces mutabilis* UAE1. (C) A close-up of spiral chain of spores of *S. mutabilis* UAE1 within the root (top) and stem (bottom) tissues (20,500×). In (A–C), all sections were stained with uranyl acetate and lead citrate showing the distribution of spiral chain of spores (red arrows). Isolate #24 represents the endophytic PA-producing *Streptomyces mutabilis* UAE1. PA, polyamine. Bars: (A,B-top) 2 µm; and (B-bottom,C) 500 nm. N, nucleus.

Surprisingly, mycelial growth of isolate #5 (PA-non-producer) and straight chain of spores were located within the root and stem of mangrove at 6 wpi (Supplementary Figure 3).

Compared to control treatment (Figure 4A), TEM studies further confirmed the presence of the spiral chain of spores and substrate mycelia of isolate #24 within the cortex of roots (Figure 4B) and stems (Figure 4C). Overall, our data suggest that isolate #24 is an actinobacterial endophyte that inhabits within living root and stem tissues of mangrove.

**Classification and Identification of the PA-Producing Endophytic Actinobacterial Isolate**

The 16S rRNA gene sequence of the promising PA-producing isolate #24 (NCBI Genbank accession number: MT883496) was compared with other species of *Streptomyces* and a phylogenetic tree was generated. Sequence analysis of 1,520-bp portions of the 16S rRNA gene indicated that this strain was 100% similar to the described species *Streptomyces mutabilis* NRRL ISP-5169T (NR_044149), but only 98.8% similar to *Streptomyces rochei*, *Streptomyces enissocaeilis*, *Streptomyces plicatus*, and *Streptomyces vinaceusdrappus* (Figure 5A). The rest of actinobacterial species were less than 99% similar to the isolate-of-interest. In addition, pure cultures of isolate #24 developed light gray aerial mass color and light gray substrate mycelium with no distinctive pigment on the reverse side of cultures on ISP medium 3 (Figure 5B). The isolate formed closed spiral chains consisting of 5–10 mature, smooth-surfaced spores on the aerial hyphae (Figure 5C). Spore chains belonged to section spirales. This suggests that the PA-producing isolate #24 could be considered as *S. mutabilis* (Preobrazhenskaya and Ryabova in Gauze et al., 1957) Pridham et al. (1958) Strain UAE1.

The sequence of the 1,521-bp region of 16S rRNA gene (accession number: MT883497) of isolate #5 showed the closest similarity to that of *Streptomyces qaidamensis*, *Streptomyces chartreusis*, and *Streptomyces osmaniensis*, with 98.7, 98.2, and 98.0%, respectively (Supplementary Figure 4). Thus, this suggests that the isolate could be any of these species or a new species that has not been identified previously. Pure cultures of strain ID 19-388 displayed whitish light gray colored aerial spore mass on ISP 3 medium, and produced yellow substrate mycelium with distinctive yellow pigment on the reverse side of cultures (Supplementary Figure 4). Based on the SEM analysis, spores were cylindrical with smooth-textured surface and spore chains were long and straight with >50 spores per chain which belonged to section rectiflexiles (Supplementary Figure 4). A more reliable identification, such as DNA–DNA hybridization, of this stain is required for proper classifications. Thus, the PA-non-producing endophytic actinobacterial isolate #5, which served as a positive control treatment in gnotobiotic (section "Evaluation of PGP Features..."
FIGURE 5 | Taxonomic determination of PA-producing *Streptomyces mutabilis* UAE1, based on phylogenetic, cultural, and morphological characteristics. (A) The tree showing the phylogenetic relationships between *S. mutabilis* UAE1 (isolate #24; MT883496; 1,520 bp) and other members of *Streptomyces* spp. on the basis of 16S rRNA sequences. (B) Aerial mycelia (left) and substrate mycelia (right) growing on ISP medium 3 supplemented with yeast extract, and (C) scanning electron micrograph (5,500) of spore chains and structure of *S. mutabilis* UAE1 which belonged to section spirales. In (A), numbers at nodes indicate percentage levels of bootstrap support based on a maximum-likelihood analysis of 1,000 resampled datasets. PA, polyamine. Bar, 0.002 substitutions per site. GenBank accession numbers are given in parentheses.

in Mangrove Seedlings Under Gnotobiotic Conditions”) and greenhouse (section “Growth Promotion of Mangrove by *S. mutabilis* UAE1 in the Greenhouse”) experiments, was designated as *Streptomyces* sp. 19-388.

**Growth Promotion of Mangrove by *S. mutabilis* UAE1 in the Greenhouse**

In our greenhouse experiments, we inoculated mangrove seedlings with *S. mutabilis* UAE1 (isolate #24), *Streptomyces* sp. 19-388 (#5), or no isolate (control). Although both *Streptomyces* sp. inoculants did not cause negative impact on growth, inoculations with the PA-producing isolate *S. mutabilis* undoubtedly promoted growth and development of mangrove (Figure 6). This was clearly observed when *S. mutabilis* significantly (*P < 0.05*) increased DW (Figure 6B) and enhanced length (Figure 6C) in roots and shoots, compared to seedlings initially inoculated with *Streptomyces* sp. 19-388 or non-inoculated plants after 9 mpp of propagules. In comparison to *S. mutabilis*-inoculated plants, the number of branches decreased significantly by 63.5% in control treatment and 65.0% in the PA-non-producing *Streptomyces* sp. 19-388 treatment (Figure 6D). The total leaf area in plants treated with *S. mutabilis* was 43.3 and 37.1% larger than in non-inoculated and *Streptomyces* sp. 19-388-inoculated plants, respectively (Figure 6E). We did not find significant differences in any of the tested growth-related agronomic characteristics in roots inoculated with *Streptomyces* sp. 19-388 or control treatment (Figures 6A–E).

Due to the strong association between the photosynthetic pigments and growth rate of crops (El-Tarabily et al., 2019, 2020; Mathew et al., 2020), we measured the contents of chl *a*, chl *b*, and carotenoids *in planta*. Our data showed that amounts of both chlorophyll pigments as well as carotenoids were significantly (*P < 0.05*) higher in *S. mutabilis*-inoculated plants than in plants inoculated or not with *Streptomyces* sp. 19-388 after 9 mpp (Figure 6F). Thus, there was no significant difference between contents of any of the three pigments in control and *Streptomyces* sp.-inoculated seedlings. In contrast to PA-non-producing
Streptomyces sp. 19-388, PA-synthesizing S. mutabilis UAE1 significantly ($P < 0.05$) stimulated photosynthetic pigment production, and consequently this might result in growth promotion of mangrove.

**Effect of S. mutabilis on Endogenous PAs and Phytohormone Contents in Mangrove Tissues**

To figure out the mechanism associated with mangrove growth promotion by S. mutabilis UAE1, we analyzed the endogenous levels of free PAs and phytohormones in root and shoot tissues. When S. mutabilis was inoculated in roots, roots and shoots of mangrove had significantly ($P < 0.05$) higher levels of Put (Figure 7A), Spd (Figure 7B), and Spm (Figure 7C) than those grown with or without amendment of PA-non-producing Streptomyces sp. 19-388. In general, the endogenous three PA amounts in shoots were higher than in roots in all treatments (Figure 7).

Under greenhouse conditions, the concentration of phytohormones in tissues of plant inoculated with S. mutabilis UAE1 and in those inoculated with Streptomyces sp. 19-388 or not was significantly ($P < 0.05$) different (Figure 8). Inoculated seedlings with S. mutabilis were characterized by almost 41–44 and 30–33% higher concentration of IAA in roots and shoots, respectively, than those in positive (Streptomyces sp. 19-388) and negative (no isolate) control treatments (Figure 8A). Likewise, the tested IPYA showed similar patterns of increase in both tissues of seedlings treated with the PA-producing isolate (Figure 8B). There was 19–21, 38–49, and 49–55% greater contents of iPA, iPa, and Z, respectively, in tissues of seedlings originally treated with S. mutabilis than those in other treatments (Figures 8C–E). We noticed, however, that ABA concentration was decreased by approximately 2.5-fold in both
tissues of *S. mutabilis*-inoculated compared to *Streptomyces* sp. 19-388-inoculated and control plants (Figure 8F).

In our controlled-greenhouse experiments, there was no significant difference in the endogenous levels of GA3 and ACC in mangrove tissues in the three treatments (Supplementary Figure 5). The levels of all phytohormones in shoot tissues were relatively higher than in root tissues. Taken together, our data suggest that *S. mutabilis* UAE1 is a PA-producing endophytic isolate that increases PA production *in planta*, thus regulating endogenous hormonal levels to enhance growth of mangrove.

**DISCUSSION**

The present study reports—for the first time—increased growth of mangrove planted in sediments and irrigated with seawater using PA-producing endophytic PGPA, as an important application in marine agricultural activities. Our current project integrates biological diversity, ecological significance, and socio-economic impact on coastal communities through restoration of mangrove forests in the Arabian Gulf. Here, we identified and characterized *S. mutabilis* UAE1 isolated from mangrove roots, which stimulated growth of seawater-irrigated mangrove seedlings under controlled greenhouse conditions. Consistent with previous studies, *Streptomyces* is the most dominant genus among endophytic actinobacterial communities of mangrove plants (Xu et al., 2016; Li et al., 2017). Toumatia et al. (2016) have found that *S. mutabilis* IA1 is an endophyte colonizing wheat tissues possessing biocontrol and PGP properties. In another study, the performance of *S. mutabilis* showing antitubercular and antimicrobial activities against bacterial pathogens has also been reported (Mahmoud et al., 2015).

In the present study, PA production by *S. mutabilis* UAE1 was determined according to the following procedure: (i) Halotolerant actinobacterial isolates identified as endophytes were *in vitro* screened based on high production of the three PAs (Put, Spd, and Spm); (ii) isolates secreting PGRs, other than PAs, were eliminated; (iii) all PA-producing isolates were tested for their PGP abilities under gnotobiotic conditions; (iv) the sole strain colonizing both root and stem tissues of mangrove was chosen; and (v) assessing *S. mutabilis* UAE1 as a PGPA isolate on agronomic characteristics and endogenous PA and other PGR contents in mangrove seedlings irrigated with seawater. In these trials, *Streptomyces* sp. 1D 19-388, the PA-non-producing isolate #5, was used as a positive control.

In general, the implementation of PGPB that can fix N or solubilize P has a positive impact on growth of marine plants including mangrove (El-Tarabily and Youssef, 2010). Seedlings of *S. bigelovii* irrigated with seawater have shown high performance in growth and seed yields after the application of N-fixing and/or P-solubilizing bacteria originally obtained from mangrove rhizosphere (Bashan et al., 2000; Rueda-Puente et al., 2003). The P-solubilizing *Oceanobacillus picturae*, isolated from mangrove soils promoted growth of mangrove seedlings under saline conditions in the greenhouse (El-Tarabily and Youssef, 2010). Our results demonstrated that the endophyte *S. mutabilis* did not fix N, solubilize P, or produce other PGRs. Thus, production of high amount of the three PAs by this strain contributed to growth promotion of mangrove in the gnotobiotic and greenhouse experiments. This current report is the first to reveal the potential of endophytic actinobacteria isolated from root tissues to stimulate mangrove growth under greenhouse conditions by possibly stimulating PA and regulating phytohormone levels *in planta*.

Killiny and Nehela (2020) have proposed that enhanced growth can possibly be due to the function of PAs as secondary messengers for phytohormones (e.g., auxins) within plant cells.
FIGURE 8 | Effect of the endophytic \textit{Streptomyces mutabilis} UAE1 on endogenous PGR contents of mangrove. Endogenous contents of auxins (A) IAA and (B) IPYA; cytokinins (C) iPA, (D) iPa, and (E) Z; and (F) ABA in mangrove root and shoot tissues after treatment with C, Ssp, and Sm. Mangrove seedlings were grown in an evaporative-cooled greenhouse and maintained at 30 ± 2°C. Values are means ± SE of 16 replicates for each treatment from two different independent experiments. Mean values followed by an asterisk are significantly ($P < 0.05$) different from each other according to Fisher's protected LSD test. Bars represent SE.

Endogenous contents of auxins, cytokinins, and ABA were measured at 9 months post planting the inoculated mangrove propagules. DW, dry weight; C, control (non-inoculated inorganic salt starch broth); Ssp, PA-non-producing isolate #5 (\textit{Streptomyces} sp. 19-388); Sm, PA-producing isolate #24 (\textit{S. mutabilis} UAE1); PA, polyamine; IAA, indole-3-acetic acid; IPYA, indole-3-pyruvic acid; iPA, isopentenyl adenine; iPa, isopentenyl adenosine; Z, zeatin; ABA, abscisic acid.

Growth promotion by \textit{S. mutabilis}, in the present study, was associated with the increase of endogenous PA levels, enhancement of photosynthetic pigments, and increases of IAA and IPYA in roots and shoots. Following the application of PA-producing rhizosphere-competent actinobacteria (El-Tarabily et al., 2020) or \textit{Bacillus amyloliquefaciens} (Idris et al., 2007), increases in endogenous auxins in treated plants have been detected. We also found that growth promotion by the PA-producing isolate was associated with increases in iPA, iPa, and Z levels. PAs might also play a key regulatory role in plant growth and development through regulating auxin/CKs ratio (Ikeuchi et al., 2016). Similar observations of high PAs contributing to plant growth promotion through regulation of endogenous CK levels have also been reported (Ortíz-Castro et al., 2008). Our study sheds light on the role of PAs as a plant growth promoter in marine agriculture. The effects of PAs in inducing mangrove growth can be explained as the augmentation of Put, Spd, and Spm concentrations and the balance of auxin and CKs in plant tissues.

In general, plant responses to unfavorable environment can be modulated by various plant hormones and PGRs. Synergistic and antagonistic interactions between PAs and other phytohormones, such as ABA and ET, may occur to regulate cellular processes in plants (Killiny and Nehela, 2020). Other studies of endogenous ABA deficiency have shown that ABA can limit ET production, consequently leading to shoot and root growth under water stress (Sharp and LeNoble, 2002). In \textit{S. bigelovii}, elevated PAs in roots and shoots and increased productivity of seeds have been linked to decreasing ABA levels in both tissues when plants were inoculated with the PA-producing actinobacterial isolate (El-Tarabily et al., 2020). Thus, the endogenous levels of ACC (the natural precursor of ET) remained unaffected. ACC contents in tissues of \textit{S. bigelovii} treated with another PA-producing rhizosphere competent isolate of \textit{Streptomyces tritolerans} isolated from mangrove sediments were comparable to control treatments, albeit increases in endogenous PAs and auxins (Mathew et al., 2020). Similarly, treating mangrove propagules with the PA-producing isolate \textit{S. mutabilis} not only...
FIGURE 9 | Model of the endophytic Streptomyces mutabilis UAE1 promoting growth of mangrove. The increased PA (Put, Spd, and Spm) levels produced by the endophytic S. mutabilis UAE1 induce endogenous contents of Put, Spd, and Spm in tissues of mangrove, resulting in increasing other PGRs in plant tissues. In planta ABA was reduced, resulting in stress relief in plant tissues. This ultimately leads to increase biomass and length of roots and stems, number of branches of stems, and photosynthetic pigments in leaves. Taken together, the PA-producing isolate S. mutabilis UAE1 has the ability to promote growth of mangrove seedlings. Green and red arrows represent increase/stimulation and decrease/inhibition, respectively. ?, unknown induction/suppression mechanism. PA, polyamine; Put, putrescine; Spd, spermidine, Spm, spermine; IAA, indole-3-acetic acid; IPYA, indole-3-pyruvic acid; iPA, isopentenyl adenine; iPa, isopentenyl adenoside; Z, zeatin; GA₃, gibberellic acid; ABA, abscisic acid.

increased PAs in plant tissues and enhanced growth of mangrove seedlings, but also lowered ABA contents and maintained normal levels of ACC in planta. Our data support previous findings that the regulatory role of PAs in plant growth can modulate ABA both under normal and stressful conditions (Hatmi et al., 2018; Alcázar et al., 2020). Thus, this modulation is mostly dependent on plant species and degree of tolerance to stress (Rangan et al., 2014).

Except Put, Spd, and Spm, no other PGRs were produced in culture filtrates of S. mutabilis UAE1. Thus, enhanced growth of mangrove due to high levels of PAs produced by this strain would result in increase of PA levels, regulation of endogenous auxins/CKs, and suppression of ABA in plant tissues. Accordingly, we proposed a model showing the possible mechanism of S. mutabilis UAE1 to stimulate the growth of gray mangrove under seawater irrigation and greenhouse conditions (Figure 9). We could not eliminate the possibility that the PGP activity of S. mutabilis UAE1 might be due to other mechanisms/metabolites that were not examined in this study. Elucidating regulatory mechanisms responsible for the PGP ability of this endophytic actinobacterium is on top of our priorities. Although the endophytic nature of S. mutabilis exerted beneficial effects on plant development, mobilization of this actinobacterial strain within tissues and translocation of PAs/PGRs from roots to stems could maximize the efficiency of PGP properties in plant tissues. Among other strategies, bioaccumulation (from soil to root or inside colonizing tissues) or translocation (from root to shoot tissues), PGPB promote growth and enhance phytoremediation (Ma et al., 2016; Liu et al., 2017). In our study, PAs produced by S. mutabilis UAE1 to promote growth of mangrove could be attributed to the expression of genes mediating auxins, CKs and by reducing ABA contents in cells.

This is the first study to utilize PA-producing microorganisms to enhance growth of mangrove under greenhouse conditions. These findings would in the future be adapted to natural competitive nursery and/or field environments. The endophytic actinobacterial isolate S. mutabilis UAE1, displaying all these PGP characteristics and boosting plant productivity in such saline conditions similar to those found in the UAE, can potentially be up-scaled to the field level. In the long run, this beneficial approach can be used in coastal areas of the Arabian Gulf to propagate marine plant species (e.g., mangrove) for reforestation purposes.
DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found below: The 16S rRNA gene sequences of *Streptomyces mutabilis* UAE1 and *Streptomyces* sp. 19-388 were deposited in Genbank accession numbers MT883496 and MT883497, respectively.

AUTHOR CONTRIBUTIONS

KE-T and SA conceived and designed the research, supervised the study, analyzed the data, and wrote the manuscript. AE, AH, and EG performed the microscopic experiments. SA developed the phylogenetic analyses. KE-T, GR, and ST assisted with the experiments and/or data evaluation. All authors critically revised the manuscript and approved the final version.

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