Distinct Community Assembly Processes of Abundant and Rare Soil Bacteria in Coastal Wetlands along an Inundation Gradient

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ABSTRACT Microbial communities commonly consist of a large number of rare taxa (RT) and few abundant taxa (AT), and it is important to identify the differences of the community assembly processes between RT and AT in response to environmental changes. However, the community assembly processes governing AT and RT in coastal wetland soils along an inundation gradient remain elusive. Here, an in situ mesocosm, with continuous inundation gradients and native mangrove *Kandelia obovata* or exotic cordgrass *Spartina alterniflora*, was established to determine the patterns and driving factors of community turnover and assembly processes of AT and RT. We found that RT exhibited a remarkably lower turnover rate than AT, and the niche breadth of RT was significantly narrower than that of AT. In comparison with AT, RT presented stronger phylogenetic signals for ecological preferences across environmental gradients. Null model analyses revealed that RT were more phylogenetically clustered and primarily governed by homogeneous selection, while AT were more overdispersed and dominated by dispersal limitation. Soil water content was the most decisive factor for community turnover and assembly processes of both AT and RT. Structural equation modeling analysis showed that RT were strongly associated with *K. obovata* biomass rather than *S. alterniflora* biomass, suggesting a strong relationship between RT and the growth of mangrove *K. obovata*. Overall, our study revealed distinct assembly processes of soil AT and RT communities in coastal wetlands, which is crucial for mechanistic understanding of the establishment and maintenance of soil microbial diversity in coastal wetlands under conditions of global environmental changes.

IMPORTANCE Coastal wetlands are one of the important ecosystems that play a crucial role in the regulation of climate change. Rare taxa (RT) exist in one habitat along with abundant taxa (AT). In this study, we found that RT exhibited narrower niche breadth and stronger phylogenetic signals than AT. Null model analyses showed that RT were more phylogenetically clustered and primarily governed by homogeneous selection, while AT were more overdispersed and dominated by dispersal limitation. Revealing the differences in the community assembly processes between AT and RT in coastal wetlands is critical to understand the establishment and maintenance of soil microbial diversity in coastal wetlands with regard to environmental changes.

KEYWORDS rare bacteria, abundant bacteria, assembly processes, plant biomass, coastal wetlands

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One of the central objectives of microbial ecology is to determine the relative contributions of deterministic processes (e.g., variable selection) and stochastic processes (e.g., dispersal limitation) in microbial community assembly (1, 2). Understanding the fundamental mechanisms of the establishment and maintenance of microbial communities is crucial for mechanistic understanding of the establishment and maintenance of soil microbial diversity in coastal wetlands under conditions of global environmental changes.
diversity is crucial for unraveling the relationships between microbial communities and ecosystem functions and for deciphering the responses and feedbacks of microbial communities to environmental changes (2–4).

Microbial communities typically exhibit a skewed distribution of species abundance, with a small number of abundant taxa (AT) and a large number of rare taxa (RT) coexisting in one habitat (5, 6). Previous studies have demonstrated the crucial role of RT in governing the functions of the microbiome, such as participating in biogeochemical cycles and facilitating plant growth (7–10). Determining the differences in the community assembly processes between RT and AT is critical to understand the responses of a rare biosphere to environmental changes. Recently, such differences in different ecosystems, including agricultural fields (11–13), subtropical bays (14, 15), and Tibetan Plateau grassland (16), have been widely studied. These studies showed that AT and RT generally exhibit distinct distribution patterns and assembly processes. For instance, it was found that the abundance of AT, instead of RT, is mainly limited by the dispersion process in agricultural fields across eastern China (11, 13) and in inland freshwater ecosystems in China (17). On the contrary, other studies have revealed that the abundance of RT is mostly limited by dispersion in subtropical bays (14) and in cascade reservoirs of the Jinsha River in China (18). Environmental factors strongly influence the balance between stochastic and deterministic processes of community assembly (1, 19, 20). For example, Tripathi et al. revealed that extreme soil pH could result in deterministic assembly of soil bacterial communities, while neutralized soil pH could lead to stochasticity during both short- and long-term succession stages (20). In addition, the changes of organic matters in soil also influence the stochastic and deterministic processes, which can shape soil bacterial communities in agroecosystems across subtropical China (19). These indicate that the relative dominance of assembly processes that structure AT and RT is varied in different ecosystems, which could be mediated by different environmental factors across both spatial and temporal scales. However, to date, little is known about the community assembly processes of AT and RT and their driving factors in coastal wetlands with environmental gradients.

The responses of microorganisms toward environmental changes tend to be phylogenetically conserved (21). For example, the influence of pH in the global bacterial biogeographic distribution exhibits strong phylogenetic conservation (21, 22). Therefore, revealing the phylogenetic conservatism of microbial response traits can facilitate the prediction of their evolutionary adaptation in response to environmental changes. However, the phylogenetic distribution of microbial communities, particularly AT and RT, in coastal wetlands has not been reported.

Coastal wetlands, located in the intertidal areas across a broad range of inundation gradients, are important ecosystems with valuable functions (23–25). They experience periodic tides and are characterized by aerobic-anaerobic fluctuations and high salinity (26). In coastal wetlands, inundation frequency is one of the most important environmental factors that influence soil microbial community (27, 28) and plant growth (29, 30). Due to global environmental changes, such as sea level rise and seawater intrusion, coastal wetlands are facing prolonged flooding, which intensively impacts their ecological functions. Therefore, understanding the microbial community assembly processes along inundation gradients is important. Recent studies have reported that coastal wetlands harbor a unique soil microbial community that is fundamentally different from that in other ecosystems (31). Thus, we hypothesize that the assembly processes of AT and RT communities in coastal wetlands are different from those in other biomes. Spartina alterniflora, originally from North America, was first introduced to China in 1979 and has rapidly spread along the eastern coastlines of China over the past 4 decades (32, 33). Currently, S. alterniflora and native mangrove plant species, such as Kandelia obovata, are commonly found to coexist in most coastal wetlands in southeastern China. K. obovata is regarded as a nutrient-limited species in subtropical coastal wetlands (34), while S. alterniflora has a strong environmental adaptability and a high degree of phenotypic plasticity (35–37). In China, the invasion of S. alterniflora
into mangroves can change soil microbial community, ultimately resulting in large differences in the associated ecological functions and responses toward environmental changes (38, 39). Nevertheless, little is known about the impact of *S. alterniflora* invasion on microbial community assembly in mangroves, which compromises the understanding and prediction of the effects of *S. alterniflora* invasion on mangrove soil microbial communities.

In this study, an *in situ* mesocosm with continuous inundation gradients and planting of *K. obovata* and *S. alterniflora* monocultures was applied to evaluate the assembly processes of AT and RT. In addition, we examined the niche breadth and phylogenetic signals for the ecological preferences across environmental gradients for both AT and RT. Considering the low competition capacity, slow growth (9, 40), and the important role of RT in soil nutrient cycling and plant growth (8, 10, 41), we aimed to investigate (i) if there is any difference in the relative influence of assembly processes that govern the composition of AT and RT in a coastal wetland ecosystem along an inundation gradient, and (ii) if the relationship between RT and plant biomass varies according to plant species. Overall, our findings aid in understanding the mechanism of the establishment and maintenance of microbial diversity in coastal wetland ecosystems and in predicting the responses of soil microbial communities to global environmental changes.

**RESULTS**

**General responses of AT and RT to environmental changes.** In total, 17,634 amplicon sequence variants (ASVs) from 2,918,880 valid sequences were obtained, among which, 863 and 16,771 ASVs were classified into AT and RT, accounting for 4.89% and 95.11% of the total ASVs, respectively (see Table S1A in the supplemental material). The rarefaction curves for all of the soil samples showed a decreasing trend in the observed ASVs against the sequence numbers and eventually reached the plateau phase (see Fig. S1). Spearman’s rank correlation showed that both AT and RT exhibited a strong abundance-occupancy relationship (see Fig. S2A). We also observed strong correlations between taxa’s niche widths and their relative abundances for both AT and RT (Fig. S2B). RT contributed to 51.60% and 50.73% of the shifts of bacterial beta diversity in *K. obovata* and *S. alterniflora* soils, respectively, and the contribution of RT remained relatively stable under different contents of soil water (Fig. S2C and D). The most abundant phylum in both AT and RT was *Proteobacteria*, which tended to decrease along an increased elevation. In addition, the relative abundances of several phyla, such as *Bacteroidetes* and *Planctomycetes*, were found to be higher in RT than in AT (see Fig. S3).

We further explored the main environmental factors that influenced the soil bacterial communities. Pairwise permutational multivariate analysis of variance (PERMANOVA) revealed that elevation significantly affected both AT (*F* = 6.76, *P* < 0.001) and RT (*F* = 6.48, *P* < 0.001), while plant species only influenced RT (*F* = 2.18, *P* = 0.03) (Table S1B). This result was also supported by the variation partitioning analysis (VPA), which suggested that AT and RT were influenced by soil physicochemical properties rather than plant species (see Fig. S4). Meanwhile, Mantel tests also demonstrated that both AT and RT were strongly associated with multiple soil physicochemical properties, including soil water content, salinity, total carbon (TC), inundation, and others (Table S1C). Among them, soil water content was the most decisive factor in regulating the bacterial community structures of AT (*r* = 0.54, *P* < 0.001) and RT (*r* = 0.54, *P* < 0.001).

Indices of observed ASVs and Faith’s phylogenetic diversity (Faith’s PD) were used to present the taxonomic and phylogenetic diversity, respectively. Two-way analysis of variance (ANOVA) revealed that the observed ASVs of AT were mainly influenced by elevation (*F* = 5.19, *P* < 0.001), while the observed ASVs of RT were mainly affected by plant species (*F* = 17.37, *P* < 0.001) (Table S1D). In addition, Faith’s PDs of both AT and RT were influenced by either elevation or plant species, the latter of which had a more profound influence (Table S1D). The observed ASVs and Faith’s PD of RT were
significantly ($P < 0.001$) higher than those of AT, in both *K. obovata* and *S. alterniflora* soils (see Fig. S5A). In addition, the alpha diversity indices in *S. alterniflora* soil, except for the observed ASVs of AT ($P = 0.60$), were all significantly ($P < 0.05$) higher than those in *K. obovata* soil (Fig. S5A). Bacterial alpha diversity of AT and RT was strongly correlated with soil physicochemical properties (Table S1E). For AT, increasing soil water content significantly decreased the observed ASVs in both *K. obovata* ($r = -0.27, P = 0.04$) and *S. alterniflora* ($r = -0.25, P = 0.04$) soils (Fig. S5B). Similarly, the Faith’s PDs of *K. obovata* ($r = -0.69, P < 0.001$) and *S. alterniflora* ($r = -0.41, P = 0.001$) soils were significantly and negatively correlated with soil water content (Fig. S5B). However, for RT, only the Faith’s PD of *S. alterniflora* soil was significantly correlated with soil water content ($r = 0.30, P = 0.02$) (Fig. S5B).

**Distance-decay relationship and community turnover.** Principal-coordinate analysis (PCoA) revealed that AT and RT were noticeably divided along the gradients of soil water content on axes 1 and 2, with 30.63% and 27.50% interpretations on axis 1, respectively (Fig. 1A). Significant ($P < 0.001$) distance-decay relationships were also observed in AT and RT of both *K. obovata* and *S. alterniflora* soils (Fig. 1B). We further calculated the rate of community turnover ($Z$ value) (Table S1F). Standardized major axis (SMA) regression analyses revealed that the slope of distance-decay relationship in RT was significantly ($P < 0.001$) lower than that in AT, suggesting that RT had a lower rate of community turnover (Table S1F and G). In addition, the $Z$ values for both AT and RT in *S. alterniflora* soil were significantly ($P < 0.001$) higher than those in *K. obovata* soil, indicating that bacterial community in *S. alterniflora* soil was more responsive to the changes of soil water content (Fig. 1B; Table S1F and G).

The bacterial beta diversity was further partitioned into total replacement diversity (Repl) and total richness difference diversity (RichDif). We found that the dissimilarity of bacterial community compositions of AT and RT for both *S. alterniflora* and *K. obovata* soils were dominated by species replacement processes (Fig. 2). The total beta diversity (BDtotal) of RT (*K. obovata*, 0.48; *S. alterniflora*, 0.47) was higher than that of AT (*K. obovata*, 0.39; *S. alterniflora*, 0.40). RichDif accounted for a larger proportion of BDtotal (up to 71.93% in AT), especially when the difference of soil water content was small, suggesting that intensive changes of soil water content could induce species replacement (Fig. 2).

**Environmental adaption of AT and RT.** We characterized the environmental adaptations of bacterial communities by using three indices: Levin’s niche breadth index,
Orwin-Wardle resistance index, and Pagel’s lambda (\(\lambda\)) phylogenetic signal (Fig. 3 and 4). For both *K. obovata* and *S. alterniflora* soils, the niche breadth index of AT was significantly \((P < 0.001)\) higher than that of RT (Fig. 3A). However, no significant difference was found in the niche breadth indices for both AT \((P = 0.14)\) and RT \((P = 0.82)\) between *K. obovata* and *S. alterniflora* soils. In addition, we also found that the resistance index of RT was slightly higher than that of AT, though at an insignificant level \((P > 0.22)\) (Fig. 3B).

The phylogenetic signal was next examined, and we found that both AT and RT exhibited strong phylogenetic signals, indicating a significant influence of phylogenetic history on the ecological traits of the soil bacterial community (Fig. 4). RT exhibited stronger phylogenetic signals for most of the environmental variables than AT, including soil water content, TC, salinity, inundation, and others, which indicated that closely related bacterial taxa of RT exhibited more similar ecological preferences in response to environmental changes.

**Bacterial community assembly processes.** The phylogenetic mantel correlogram revealed significant \((P < 0.05)\) phylogenetic signals across short phylogenetic distances, indicating that the ecological traits in regulating bacterial community assembly processes were phylogenetically conserved (see Fig. S6A and B). Mantel tests demonstrated that the assembly processes were significantly \((P < 0.01)\) associated with certain environmental variables, including soil water content, salinity, TC, inundation, and others (Table S1H). Among these variables, soil water content was the most decisive factor in regulating bacterial community assembly for both AT (*K. obovata*: Mantel \(r = 0.56, P < 0.001\); *S. alterniflora*: Mantel \(r = 0.56, P < 0.001\)) and RT (*K. obovata*: Mantel \(r = 0.58, P < 0.001\); *S. alterniflora*: Mantel \(r = 0.73, P < 0.001\)). Moreover, the nearest taxon index (NTI) value of RT was significantly \((P < 0.001)\) higher than that of AT, suggesting a more phylogenetic clustering in RT (Fig. S6C). We also found that the NTI
value, except for that of AT in *K. obovata* soil (*r* = 20.07, *P* = 0.62), was significantly and negatively correlated with soil water content, indicating that the increased soil water content could reduce the level of phylogenetic clustering (Fig. 5A).

Furthermore, most of the beta nearest taxon index (betaNTI) values of AT were between 2 and 2, indicating the dominant role of stochastic processes (dispersal limitation and undominated processes) in AT community assembly (Fig. 5B). The betaNTI values of AT in *K. obovata* soil, rather than in *S. alterniflora* soil, were significantly (adjusted *R*² [Radj²] = 0.29, *P* < 0.001) and negatively correlated with soil water content (Fig. 5B). On the contrary, most of the betaNTI values of RT were less than 2, suggesting that the bacterial community assembly was mainly governed by deterministic processes (homogeneous selection) (Fig. 5B). The betaNTI values of RT in both *S. alterniflora* (*Radj²* = 0.15, *P* < 0.001) and *K. obovata* (*Radj²* = 0.05, *P* = 0.02) soils were significantly correlated with soil water content (Fig. 5B). We observed that the betaNTI was significantly (*P* < 0.001) correlated with the changes of soil water content, facilitating the transition of bacterial community assembly from homogeneous selection to stochastic processes and further to variable selection (Fig. 5C). The quantitative

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**FIG 3** Niche breadths (log₁₀ transformed) (A) and resistance indices (B) of AT and RT of *K. obovata* and *S. alterniflora* soils. Significant differences were examined by Wilcoxon rank sum test. For abbreviations, see legend of Fig. 1.

**FIG 4** Phylogenetic signals (lambda) showing the trait conservatism of AT and RT of *K. obovata* and *S. alterniflora* soils for each soil physicochemical property. For abbreviations, see legend of Fig. 1.
estimates of the relative contribution of assembly processes showed that stochastic processes were dominant in AT, while deterministic processes were dominant in RT (Fig. 5D).

Correlations between soil bacterial community and plant biomass. In order to investigate the direct and indirect effects of increasing inundation on plant biomass with simultaneous consideration of multiple factors, we used a structural equation model (SEM) to assess the potential correlations among environmental factors, bacterial community structure, and plant biomass. We found that both K. obovata and S. alterniflora displayed a hump-shaped pattern of biomass across inundation gradients (Fig. 6E). K. obovata performed the best at slightly higher elevations (with an inundation frequency between 7.73% and 13.90%), while S. alterniflora performed the best at slightly lower elevations (with an inundation frequency between 21.40% and 29.20%) (Fig. 6E). First, we identified the most important factors that influenced the biomass of K. obovata and S. alterniflora as well as the bacterial communities of AT and RT (see Fig. S7 and S8), which explained 73% and 67% of the total variation of K. obovata and S. alterniflora biomass, respectively (Fig. 7). The SEM revealed that K. obovata biomass was significantly and positively correlated with RT ($\lambda = 0.50$, $P < 0.01$), water content ($\lambda = 0.48$, $P < 0.05$), and Fe content ($\lambda = 0.18$, $P < 0.05$) while significantly and negatively correlated with inundation ($\lambda = -0.91$, $P < 0.001$) and Ca content ($\lambda = -0.26$, $P < 0.001$) (Fig. 7A). However, S. alterniflora biomass was significantly and positively correlated with salinity ($\lambda = 0.43$, $P < 0.01$) and water content ($\lambda = 0.87$, $P < 0.001$) and significantly and negatively correlated with inundation ($\lambda = -0.42$, $P < 0.01$) and Ca content ($\lambda = -0.24$, $P < 0.01$) but insignificantly ($P > 0.05$) correlated with either AT or RT (Fig. 7B). Overall, based on the total effects that were standardized from SEM,
it was suggested that *K. obovata* biomass was primarily influenced by soil water content, RT, and salinity, while *S. alterniflora* biomass was mainly driven by soil water content, inundation, and salinity (see Fig. S9).

**DISCUSSION**

**Distinct assembly processes of AT and RT.** Quantifying the relative contributions of deterministic and stochastic processes on community assembly is one of the central objectives of microbial ecological studies (1, 5, 42). In this study, we found that deterministic processes primarily governed RT, while stochastic processes dominated AT in coastal wetlands (Fig. 5). Our findings were consistent with previous studies which have reported the dominance of deterministic processes in RT, rather than in AT, in agricultural fields across eastern China (11, 13) and the northwestern Pacific Ocean (43). It is possible that the habitat occupancy of AT with a wide niche breadth is more likely to be limited by dispersion, whereas the distribution of RT with a narrow niche breadth might be delimited by environmental filtering. On the contrary, several studies have revealed that the assembly processes of RT were primarily limited by neutral processes, in comparison with those of AT, in subtropical bays (14) and cascade reservoirs of the Jinsha River in China (18). Such discrepancies of microbial community assembly could be partially attributed to spatial scale dependency and habitat diversity (44). Moreover, these inconsistent findings might also be explained by the variations in metabolic
activity, body size, and dispersal potential among different microbial populations in soil (45, 46).

Understanding the ecological niches and evolutionary characteristics of microbial ecological traits is important for determining the mechanism of community assembly (11, 47). Here, we further investigated the differences of environmental adaptive capabilities between AT and RT in coastal wetlands. First, we found that the niche breadth of RT was narrower than that of AT (Fig. 3). It is generally recognized that AT can efficiently utilize a wider range of resources than RT, thus becoming more abundant in the same environment (5, 11). Thus, the AT can be easily dispersed, as there are more individuals in the environment (17). On the contrary, the soil bacteria in RT were unevenly distributed in limited locations, with relatively low abundances. The restricted niche breadth of RT may be attributed to the low competitive capacity and growth rate (9, 40). Overall, the narrower niche breadth of RT than that of AT can induce increased competition among different microorganisms for similar resources,

**FIG 7** Structural equation model (SEM) revealing the direct and indirect effects of soil physicochemical properties, AT, and RT on *K. obovata* (A) and *S. alterniflora* (B) biomass. Numbers on arrows are the path coefficients and are indicative of the standardized effect size of the relationship. Arrows in black and red indicate positive and negative effects, respectively. $R^2$ means the proportion of variance explained. Solid and dashed lines indicate significant and insignificant correlations, respectively. *, $P < 0.5$; **, $P < 0.01$; †††, $P < 0.001$. For abbreviations, see legend of Fig. 1.
where environmental filtering plays a dominant role in microbial community assembly processes. Nevertheless, experimental verifications of microbial taxa in response to environmental changes are essential for environmental management, since the niche breadth analyses based on statistics might remain discrepant with the real situation.

Second, environmental adaptation could be partially reflected by phylogenetic conservation of the traits for microbial ecological preferences. Phylogenetic conservation of traits could provide projections on the evolutionary adaptation of microbial communities subject to ongoing environmental changes (21). In the present study, we found that RT exhibited stronger phylogenetic signals for most of the ecological preferences than AT, particularly for the main factors influencing bacterial community, such as soil water content, salinity, inundation, and TC (Fig. 4). In the presence of significant phylogenetic signals, environmental filtering could result in phylogenetic clustering within the community (48, 49). Accordantly, we also found that AT had a significantly lower NTI than RT, indicating that the soil bacteria in RT were more phylogenetically clustered (Fig. 5A; see also Fig. S6C in the supplemental material). In fact, previous studies have demonstrated that deterministic processes are positively associated with phylogenetic clustering in bacterial communities (20, 50).

Microbial community assembly is mediated by the balance between stochastic and deterministic processes (5, 42). We found a significantly lower rate of community turnover in RT than in AT (Fig. 1), which was consistent with prior studies based on RT in the Yangtze River basin (51) as well as in lakes and reservoirs across China (14, 17, 18). On one hand, RT can become dominant under favorable environmental conditions (7), evolve to adapt to environmental changes, and increase the community resistance to environmental disturbances (Fig. 3). On the other hand, the assembly processes of RT were dominated by homogeneous selection in comparison with the dominance of variable selection and dispersal limitation in AT assembly processes, which enabled RT with greater convergence and weaker distance-decay relationships (4, 19).

Additionally, we also observed that the community turnover rates for both AT and RT were significantly higher in S. alterniflora soil than in K. obovata soil (Fig. 1). This might be due to the higher variable selection and dispersal limitation in the community assembly processes of S. alterniflora soil, which increase the divergence in the microbial community composition. The above-mentioned results also suggested that the bacterial community in S. alterniflora soil is more responsive to environmental changes. Currently, S. alterniflora is rapidly spreading and occupying most of the coastal wetlands in China, which induces significant changes in the underground microbial communities and their ecological functions (33). The present study facilitates an in-depth mechanistic understanding of the responses of soil bacterial community to environmental changes in coastal wetlands.

**RT is important for mangrove plant performance.** The rare biosphere plays an important role in mediating soil nitrogen cycling (8, 10, 41), pollutant degradation (52), and plant growth (10, 53, 54). We have observed a strong association between RT and K. obovata biomass (Fig. 7), which is consistent with previous studies suggesting the importance of soil RT in regulating the aboveground biomass in alpine grassland in Qinghai-Tibetan Plateau (10). Such an association might be due to the crucial functions of RT in soil carbon cycling (55, 56), nitrogen fixation (8), and control of plant pathogens (57). Past studies have reported that RT can be more active than AT (9). Furthermore, previous studies have reported that under fluctuating conditions, such as in coastal areas experiencing periodic tides, RT could become hyperactive as a result of continuous regrowth (58). Moreover, the huge functional gene pool in RT could contribute to the outstanding metabolic potential, especially unique metabolic pathways, for the microbial communities under appropriate conditions (59). Additionally, the functionality of RT could also be enhanced by AT through microbial interactions (60). However, the current findings were mainly derived from the SEM analyses, for which the specific microbial taxa at play were unknown. Further isolation and functional
studies on these key microbial taxa by multi-omics will contribute to a deeper understanding of mangrove growth in response to environmental changes.

In contrast, a strong relationship between RT and \textit{S. alterniflora} biomass was not observed (Fig. 7), which might be due to the different plant adaptive strategies between native species \textit{K. obovata} and invasive species \textit{S. alterniflora}. \textit{S. alterniflora} is an exotic perennial C\textsubscript{4} cordgrass and has a strong environmental adaptability (37). Previous studies have demonstrated that the invasive ability of \textit{S. alterniflora} in China is mostly governed by provenance-by-environment interactions (35, 61, 62). Thus, the rapid growth and spread of \textit{S. alterniflora} are possibly attributed to its broad capacity of preadaptation and high degree of phenotypic plasticity (35, 36). Moreover, compared with \textit{K. obovata}, \textit{S. alterniflora} is more tolerant to prolonged inundation (Fig. S7E), which compromises the relationship between plant growth and soil microbial communities. However, \textit{K. obovata} biomass was significantly correlated with RT in addition to environmental factors (i.e., soil water content and inundation frequency), which might be because \textit{K. obovata} is generally regarded as a nutrient-limited species in subtropical coastal wetlands (34).

\textit{S. alterniflora} invasion in coastal wetlands of China strongly influenced the soil microbial communities and their associated functions (38, 39). \textit{S. alterniflora} can supply specific soil microorganisms with certain substrates, such as trimethylamine (63, 64). Here, we suggested that the changes of aboveground plant species were more closely associated with RT than with AT (Table S1B), due to RT’s narrower niche breadth and ecological preferences for certain habitats and resources (11, 65). With the continuous elevation of global sea level, global coastal wetlands are undergoing prolonged flooding (66), under which circumstance, \textit{S. alterniflora} could continuously and aggressively expand into most coastal wetlands (67, 68). On the contrary, native mangroves, such as \textit{K. obovata}, perform better at slightly higher elevations which are subjected to relatively less flooding. Such native mangrove habitats are fragmented and invaded by \textit{S. alterniflora} due to increased frequency and duration of flooding as well as the intensification of human activities (69, 70). Subsequently, the dominancy of \textit{S. alterniflora} could induce fundamental changes in the soil microbiome, which further influences the growth of plants (71). Moreover, it was reported that \textit{S. alterniflora} invasion could suppress \textit{K. obovata} biomass by up to 90% (29). Therefore, we suggest that protecting soil RT in coastal wetlands is crucial for maintaining and enhancing the functions of the ecosystem under the context of global environmental changes (e.g., biological invasion and deeper inundation). Future studies considering the community compositions and functions of root-associated microorganisms and plant endosymbions could facilitate a deeper understanding of the responses of plant growth to global environmental changes.

In conclusion, this study demonstrates the mechanisms of AT and RT community assembly in response to environmental changes in coastal wetlands. \textit{S. alterniflora} invasion in coastal wetlands increases the sensitivity of soil microbial communities to environmental changes. Furthermore, homogeneous selection is dominant in RT, while dispersal limitation primarily governs AT. In addition, the RT community is strongly associated with \textit{K. obovata} biomass rather than \textit{S. alterniflora} biomass. These findings provide a scientific foundation for better understanding the responses of coastal wetland ecosystems to global environmental changes from the viewpoint of microbial ecology.

**MATERIALS AND METHODS**

\textit{In situ mesocosm design.} An \textit{in situ} mesocosm was designed and established in the Zhangjiang River Estuary (23°55'N, 117°26'E) in Fujian Province, China, according to previous studies (29, 72, 73). This area has a subtropical marine climate (annual mean air temperature at 21.5°C) with irregular semidiurnal tides (average range of 2 m to 3 m) (26). The mesocosm allowed us to investigate the soil microbiome along a fine-grained inundation gradient, eliminating the influences of other environmental changes and simulating the natural conditions.

In brief, three platforms, each consisting of two subplatforms (4 m in length by 2 m in height) that were placed in parallel along the direction from shoreward to seaward, were established in the intertidal...
were measured. Brie reads were discarded to avoid possible biases according to prior studies (17, 78). The SILVA database sequences were denoised by using DADA2 (version 2019.10) (77). Meanwhile, ASVs with fewer than 2 poor quality (read length of 200 bp or average quality score of $>20$) were removed. Then, the filtered sequences were denoised by using DADA2 (version 2019.10.0) (77). Meanwhile, ASVs with fewer than 2 reads were discarded to avoid possible biases according to prior studies (17, 78). The SILVA database (<https://www.arb-silva.de/>, version 132) was employed for microbial taxonomy assignment. Finally, a total of 2,918,880 sequences were obtained from all of the 120 soil samples. Each sample was rarefied to 24,324 sequences (minimum) for downstream analyses. Then, all the ASVs were classified into 6 categories based on the cutoffs as described by recent studies (65, 79, 80): always abundant taxa (AAT), those with a relative abundance of $\geq 1\%$ in all samples; conditionally abundant taxa (CAT), those with a relative abundance of $0.01\%$ in all samples and $\geq 1\%$ in some samples; always rare taxa (ART), those with a relative abundance of $<0.01\%$ in all samples; conditionally rare taxa (CRT), those with a relative abundance of $<0.01\%$ in some samples but $\geq 1\%$ in any sample; moderate taxa (MT), those with a relative abundance between 0.01% and 1% in all samples; and conditionally rare and abundant taxa (CRAT), those with a relative abundance ranging from rare (<0.01%) to abundant ($\geq 1\%$). For the comparative study of AT and RT and to avoid confusions AAT, CAT, and CRAT were jointly counted as AT, and ART and CRT were jointly counted as RT (65, 81). Here, we mainly focused on the AT and RT, as MT were not detected in this study. The general description of AT and RT is shown in Table S1A in the supplemental material.

Statistical analyses. Two-way ANOVA was used to examine the effects of elevation, plant species, and their interactions on the alpha diversity of soil bacterial communities. Spearman’s rank correlation rho ($\rho$) was used to assess the relationship between soil physicochemical properties and bacterial alpha diversity. The significant difference between any two slopes was analyzed and compared by SMA regression analysis using the SMATR package in R software (82). A significant difference in bacterial alpha diversity between any two groups was examined by Wilcoxon rank sum test. A PERMANOVA was conducted to determine the significant differences in soil bacterial communities among elevations and between plant species. In addition, a VPA was performed to assess the effects of plant species and soil physicochemical properties on bacterial community structure using the vegan package. Then, a Mantel test was used to determine the significance of the relationship between each soil physicochemical property and bacterial community structure. PCoA was performed based on the
weighted UniFrac dissimilarities along environmental gradients. The spatial turnover rate (Z value) of soil bacterial communities was determined based on the distance-decay relationship between Sørensen similarity and environmental distance (83). The contribution of RT to the shifts in bacterial community dissimilarity was estimated according to Shafe et al. (84). To assess the difference between AT and RT subpopulations, Spearman’s rank correlation ρ was used to assess the abundance-occupancy relationship of bacterial taxa between the log-transformed mean relative abundance of bacteria and the number of sites they occupied (85). The relationship between taxa’s niche width and their log-transformed mean relative abundance was also determined (86, 87). In addition, to better understand the biodiversity patterns and to explore their causes, compositional dissimilarities (beta diversity) among sites were subdivided into replacement and richness difference components (Podani family, Sørensen dissimilarities) using the adespatial package.

To further explain the patterns of beta diversity, Levin’s niche breadth indices of each bacterium were calculated by using the spa package. The Orwin-Wardle resistance indices for the soil bacterial community were further calculated (88). In addition, the phylogenetic signals were estimated to examine if the environmental preference of a bacterial ASV was related to the phylogeny, reflecting the degree of phylogenetic conservatism in response to environmental gradients (21). To obtain the potential trait, we identified the ecological preferences of each ASV via the Spearman’s correlations between the relative abundances of ASVs and each soil physicochemical property (89). Next, Page’s lambda was applied to measure the phylogenetic signals for the environmental preferences of taxa using the phytools package (90, 91). The value of lambda is between 0 and 1, with larger values indicating a stronger phylogenetic signal.

A previously developed null modeling approach, based on the assumption of phylogenetically conserved environmental preferences of microbial lineages, was used to determine the relative influences of community assembly processes (2). This approach has been shown to provide robust estimates of the relative influences of different microbial community assembly processes in a range of ecosystems (1, 2, 20, 44). To observe phylogenetic conservatism among microbial ASVs, we tested phylogenetic signals in association with habitat using the “mantel.correlog” function with 999 randomizations (2, 42, 92). The community assembly processes were evaluated by calculating the NTI and betaNTI using the “ses.mntd” function in the picante package (93) and a previously developed null modeling approach (2), respectively. NTI values, the negative values of ses.mntd output, were used to evaluate the phylogenetic community assembly at a within-community scale, in which the positive and negative values indicated clustering and overdispersion of taxa across the overall phylogeny, respectively (50). According to Stegen et al., betaNTI is the number of standard deviations that the observed beta mean nearest taxon distance (betaMNTD) is from the mean of the null distribution (2, 42). A value of betaNTI of >2 or less than –2 indicates greater than or less than the expected phylogenetic turnover, respectively. The combination matrix of betaNTI values and Bray-Curtis based Raup-Crick (RCbray) was applied to estimate the relative contributions of homogeneous selection, variable selection, dispersal limitation, homogenizing dispersal, and undominated processes in governing community assembly (42). Pairwise betaNTI of less than –2 or >2 indicated homogeneous selection or variable selection, respectively. RCbray of less than –0.95 or >0.95 indicated significant deviations from the null model expectation. |betaNTI| of <2 with RCbray of less than –0.95 or >0.95 indicated a contribution of homogenizing dispersal or dispersal limitation, respectively. Otherwise (|betaNTI| < 2 and |RCbray| < 0.95), the shifts in community composition were undominated. The significance of the relationship between betaNTI and soil physicochemical properties was assessed by Mantel tests.

An SEM was used to evaluate the direct and indirect effects of inundation changes on K. obovata and S. alterniflora biomass. Prior to SEM analysis, classification random forest analyses were conducted by using the rfPermute package to identify the most important predictors that influenced plant biomass and bacterial community structures (represented by PCoA axis 1) of AT and RT, respectively. One thousand permutation replicates were then used to construct the null distribution and calculate the P values. All selected predictors were included in the SEM as independently observed variables. Since most of the variables were not normally distributed, the probability that a path coefficient differed from zero was assessed by using the bootstrapping method (94). Meanwhile, three metrics were used to test the goodness-of-fit of SEM (95): (i) chi-square test, in which a good fit is defined as 0 ≤ χ²/df ≤ 2 and 0.05 < P ≤ 1.00, (ii) Bollen-Stine bootstrap test, in which a good fit is defined as 0.10 < Bollen-Stine bootstrap and P ≤ 1.00, and (iii) root mean square error of approximation (RMSEA), in which a good fit is defined as 0 ≤ RMSEA ≤ 0.05. Then, the standardized total effect of SEM attributes on plant biomass was calculated. SEM analyses were performed by using AMOS 21 (IBM SPSS Inc., Chicago, IL, USA). Statistical significance was determined by a P value of < 0.05 for all analyses.

Data availability. All the obtained sequences were deposited in the NCBI Sequence Read Archive (SRA) with BioProject accession number PRJNA597962.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

FIG S1, PDF file, 0.5 MB.
FIG S2, TIF file, 2.6 MB.
FIG S3, PDF file, 0.1 MB.
FIG S4, PDF file, 0.1 MB.
FIG S5, PDF file, 0.2 MB.
REFERENCES


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H. Chu and Y. Zhang designed the experiments. G.-F. Gao and D. Peng collected the samples and determined the soil and plant variables. G.-F. Gao performed the bioinformatic analyses, and H. Chu evaluated the data. G.-F. Gao wrote the original manuscript, and D. Peng, B. M. Tripathi, Y. Zhang, and H. Chu revised and approved the manuscript.


