Metabolic Exchange with Non-Alkane-Consuming *Pseudomonas stutzeri* SLG510A3-8 Improves *n*-Alkane Biodegradation by the Alkane Degrader *Dietzia* sp. Strain DQ12-45-1b

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**ABSTRACT** Biodegradation of alkanes by microbial communities is ubiquitous in nature. Interestingly, the microbial communities with high hydrocarbon-degrading performances are sometimes composed of not only hydrocarbon degraders but also nonconsumers, but the synergistic mechanisms remain unknown. Here, we found that two bacterial strains isolated from Chinese oil fields, *Dietzia* sp. strain DQ12-45-1b and *Pseudomonas stutzeri* SLG510A3-8, had a synergistic effect on hexadecane (*C_{16}* compound) biodegradation, even though *P. stutzeri* could not utilize *C_{16}* individually. To gain a better understanding of the roles of the alkane nonconsumer *P. stutzeri* in the *C_{16}*-degrading consortium, we reconstructed a two-species stoichiometric metabolic model, jBH1908, and integrated *in silico* prediction with the following *in vitro* validation, a comparative proteomics analysis, and extracellular metabolomic detection. Metabolic interactions between *P. stutzeri* and *Dietzia* sp. were successfully revealed to have importance in efficient *C_{16}* degradation. In the process, *P. stutzeri* survived on *C_{16}* metabolic intermediates from *Dietzia* sp., including hexadecanoate, 3-hydroxybutanoate, and α-ketoglutarate. In return, *P. stutzeri* reorganized its metabolic flux distribution to feed back acetate and glutamate to *Dietzia* sp. to enhance its *C_{16}* degradation efficiency by improving *Dietzia* cell accumulation and by regulating the expression of *Dietzia* succinate dehydrogenase. By using the synergistic microbial consortium of *Dietzia* sp. and *P. stutzeri* with the addition of the *in silico*-predicted key exchanged metabolites, diesel oil was effectively disposed of in 15 days with a removal fraction of 85.54% ± 6.42%, leaving small amounts of *C_{15}* to *C_{20}* isomers. Our finding provides a novel microbial assembling mode for efficient bioremediation or chemical production in the future.

**IMPORTANCE** Many natural and synthetic microbial communities are composed of not only species whose biological properties are consistent with their corresponding communities but also ones whose chemophysical characteristics do not directly contribute to the performance of their communities. Even though the latter species are often essential to the microbial communities, their roles are unclear. Here, by investigation of an artificial two-member microbial consortium in *n*-alkane biodegradation, we showed that the microbial member without the *n*-alkane-degrading capability had a cross-feeding interaction with and metabolic regulation to the leading member for the synergistic *n*-alkane biodegradation. Our study improves the current understanding of microbial interactions. Because “assistant” microbes showed importance in communities in addition to the functional microbes, our findings also contribute to the performance of their communities.
suggest a useful “assistant-microbe” principle in the design of microbial communities for either bioremediation or chemical production.

**KEYWORDS** flux balance analysis, interspecies metabolite transfer, metabolomics, proteomics, synergistic biodegradation of \( n \)-alkanes

Bioremediation of petroleum-contaminated environments and microbial-enhanced oil recovery are two environment-friendly, cost-effective techniques to solve serious oil-related problems in current society (1, 2). Clearly, these two biotechniques have opposite aims: one for oil removal and the other for oil recovery. However, both of them preferably induce microbes or microbial consortia with the function of \( n \)-alkane degradation (3–5). Microbial communities are usually more advantageous for hydrocarbon treatment than the individual species by providing a wider substrate spectrum, higher robustness, and better adaptation to the complicated environments (6, 7). Interestingly, it has been observed that outstanding performance of petroleum alkane biodegradation could be achieved by some microbial communities composed of not only alkane consumers but also nonconsumers (8–11). What are the roles of the microbial members who do not appear to contribute to the main function of their communities? First, they might be dormant cells in the microbial communities under current conditions, acting like seeds waiting for suitable environmental conditions (12). Second, the alkane nonconsumers might have some close affiliations (13) or contact-independent interactions (14) with the alkane-degrading members. If the microbial members without the objective function of the microbial communities are active instead of dormant, metabolic exchange is considered to be a key reason for their survival in these habitats, and their activities should have neutral or positive effects on their functional partners possibly through division of labor (15). However, in some cases, either microbial dormancy or labor division patterns are not sufficient to explain some phenomena. For example, Ghazali et al. (16) screened out six bacterial isolates from oil-contaminated soil to construct bacterial consortium 1 (including three isolates) and consortium 2 (including all six isolates). They found that consortium 2 was more effective than consortium 1 for the removal of medium- and long-chain alkanes, especially \( n \)-tetradecane, in the diesel-contaminated soil, but unexpectedly the three isolates unique to consortium 2 were unable to utilize \( n \)-tetradecane or its downstream intermediates individually. Zanaroli et al. (17) isolated two microbial consortia from cow manure with remarkable diesel biodegradation activities, especially for the degradation of \( C_{10} \) to \( C_{24} \) \( n \)-alkanes. However, except for the fungus *Trametes gibbosa*, none of the isolated cultivable bacteria grew well on \( n \)-alkanes or degraded \( n \)-alkanes individually. Hence, several questions about those microbial communities remain to be answered, such as how the alkane degraders and nonconsumers cooperate with each other to achieve synergistic alkane biodegradation and what the roles of these alkane nonconsumers within the communities are. The newly developed multiple “omics” technologies (18) and mathematical modeling approaches, such as the genome-scale metabolic model (GEM) reconstruction (19), provide promising ways to better understand the microbial communities in nature and bioprocesses.

In this study, we constructed a simple microbial consortium consisting of an alkane-degrading bacterial strain, *Dietzia* sp. strain DQ12-45-1b, and a potential non-alkane-degrading bacterial strain, *Pseudomonas stutzeri* SLG510A3-8, both of which were isolated from Chinese oil fields, to investigate the communication mechanism in the synthetic microbial consortium via GEM and omics analyses. The genera *Dietzia* and *Pseudomonas* were widely detected in oil fields (20–22). *Dietzia* species are Gram-positive bacteria that are widely distributed in diverse environments, and some *Dietzia* members have been shown to have the potential for hydrocarbon biodegradation (23). Among them, *Dietzia* sp. DQ12-45-1b was one of the few strains that were reportedly able to utilize a broad spectrum of \( n \)-alkanes (\( C_6 \) to \( C_{40} \)), and this property was attributed to the collaboration of a CYP153 alkane hydroxylase and an integral membrane alkane monoxygenase-rubredoxin fusion protein, AlkW1, in the strain (24–26).
P. stutzeri, a Gram-negative bacterium, was seldom reported to have the capability to degrade hydrocarbons. For instance, P. stutzeri SLG510A3-8 lacked essential alkane degradation genes such as alkane hydroxylase genes (27), indicating its incompetence for growth on n-alkanes. However, in line with the fact that P. stutzeri is one of the most ubiquitous and abundant species in hydrocarbon-related environments (28–30), its strains are considered to interact with the hydrocarbon degraders when they are exposed to hydrocarbons and to play important roles in their microbial communities. Here, to investigate the microbial interaction between the hydrocarbon-degrading microbes and nonconsumers when they are exposed to hydrocarbons as the sole carbon source, we reconstructed a two-species GEM of Dietzia sp. DQ12-45-1b and P. stutzeri SLG510A3-8 and analyzed the metabolic flux distribution between the two "compartments" in the stoichiometric model using the flux balance analysis (FBA) algorithm. Afterward, we did in vitro cultivation, comparative proteomics analysis, and extracellular metabolomics analysis based on the in silico prediction of the metabolic exchange between the two strains on hexadecane (C₁₆) to understand the mechanism of communication between the two strains. Our finding on the mutualistic interaction between the leading and supporting members reveal a novel assemblage strategy to construct synthetic microbial consortia with high-performing functions.

RESULTS

**C₁₆ degradation by the synthetic microbial consortium.** Bacterial strains Dietzia sp. strain DQ12-45-1b and P. stutzeri SLG510A3-8 were aerobically cocultivated on C₁₆. When P. stutzeri was exposed to C₁₆ individually, the percentage of C₁₆ removed by P. stutzeri was low in a 12-day culture period (5.21% ± 2.88%) (Fig. 1a), and the P. stutzeri cell density decreased to a very low level (Fig. 1b). This indicated that even if P. stutzeri could adhere to a small amount of C₁₆, it was unable to utilize C₁₆, which was consistent with the fact that no alkane-oxidizing gene was identified in its genome. For Dietzia sp., C₁₆ was easily available, since Dietzia sp. cells grew well on C₁₆ (Fig. 1c), with 79.24% ± 6.71% of the total C₁₆ being removed in 12 days in its monoculture (Fig. 1a). Nevertheless, the percentage of C₁₆ removal by the microbial consortium (92.94% ± 1.67%) was significantly higher than those by each individual strain (P < 0.05). On the other hand, both P. stutzeri and Dietzia sp. in the coculture system grew much better than those in monocultures, even though the cell density of P. stutzeri was an order of magnitude less than that of Dietzia sp. (Fig. 1b and c). By analyzing the time courses of Dietzia cell density and residual C₁₆ data, it was seen that the specific C₁₆ removal rate of Dietzia sp. at its highest specific growth rate was significantly higher in the coculture system (1.38 ± 0.10 pg day⁻¹ CFU⁻¹) than in its monoculture system (0.71 ± 0.16 pg day⁻¹ CFU⁻¹) (P < 0.05). Based on the efficient performance in C₁₆ degradation by the coculture of P. stutzeri and Dietzia sp., a synergistic effect on C₁₆ biodegradation through metabolic communication was pro-
posed for the coculture system, in which *P. stutzeri* might survive on the by-products of *Dietzia* sp. under the C\(_{16}\) exposure condition and, in return, secretory metabolites of *P. stutzeri* are able to induce a higher C\(_{16}\) recovery efficiency of the *Dietzia* strain.

**Reconstruction and characterization of the two-species model.** To elucidate the interspecific metabolic communication between *P. stutzeri* and *Dietzia* sp., two-species metabolic network reconstruction and consortium-level flux balance analysis (FBA) were carried out. First, we reconstructed the single-species GEM of *Dietzia* sp. DQ12-45-1b, namely, *iBH925* (see File S2 in the supplemental material), and the GEM of *P. stutzeri* SLGS10A3-8, namely, *iBH983* (see File S3 in the supplemental material). The accuracy of these two models was validated by the consistency between *in vitro* experimental data and *in silico* prediction of growth rates and available carbon source spectra (see File S1 in the supplemental material). Afterward, *iBH925* and *iBH983* were reconciled to unify the metabolites and reaction formats, followed by network integration. The resulted model for the microbial consortium, namely, *iBH1908*, contained 1,908 genes, 1,273 metabolites, and 2,670 reactions (see File S4 in the supplemental material). In *iBH1908*, the metabolic networks of *Dietzia* sp. and *P. stutzeri* were connected to each other through 68 shared metabolites (see Table S1 in File S1). Details of the modeling and analyzing processes of *iBH925*, *iBH983*, and *iBH1908* are described in File S1.

By using FBA, the flux distribution in *Dietzia* sp. and *P. stutzeri* was simulated for individual growth or coculture in C\(_{16}\)-exposed minimal medium. A maximum growth rate \(\mu_{o,0718} = 2.38 \times 10^{-2} \text{ h}^{-1}\) (see Table S2 in File S1) for *Dietzia* sp. in monoculture was predicted by FBA on *Dietzia* sp. GEM *iBH925* at the C\(_{16}\) uptake rate of 7.18 \(\times 10^{-2}\) mmol g\(^{-1}\) h\(^{-1}\), which was determined by batch culture (see Fig. S1 in File S1). Similarly, with the same nutrient constraints, a maximum growth rate of 0 h\(^{-1}\) for *P. stutzeri* was predicted by FBA on *P. stutzeri* GEM *iBH983*. In contrast, FBA on the two-species GEM *iBH1908*, in which the *Dietzia* sp. growth rate was constrained to be 2.38 \(\times 10^{-2}\) h\(^{-1}\) while the C\(_{16}\) uptake rate was fixed at 8.00 \(\times 10^{-2}\) mmol g\(^{-1}\) h\(^{-1}\) based on our previous measurement (File S1), predicted a maximum growth rate of 5.60 \(\times 10^{-3}\) h\(^{-1}\) for *P. stutzeri*. Taken together, these predictions were in agreement with our previous findings that no *P. stutzeri* cell grew on C\(_{16}\) individually but its cell density gradually increased in the presence of *Dietzia* sp. (Fig. 1b). The global-level flux distribution in *iBH1908* reflected that *Dietzia* sp. provided five metabolites to *P. stutzeri*, while *P. stutzeri* secreted another 10 compounds for *Dietzia* sp. utilization (Fig. 2). *P. stutzeri* GEM *iBH983* and *Dietzia* sp. GEM *iBH925* were utilized to predict the importance of each individual interspecific metabolite. We simulated the growth rates of *P. stutzeri* on *iBH983* via FBA when the absorption rates of the five compounds transferred from *Dietzia* sp. to *P. stutzeri*, i.e., R-3-hydroxybutanoate, \(\alpha\)-ketoglutarate, formate, hexadecanoate, and glycolaldehyde, were set at the responding values in Fig. 2. Under these conditions, the maximum predicted growth rate of *P. stutzeri* increased from 0 to 3.00 \(\times 10^{-2}\) h\(^{-1}\), confirming that the metabolites clearly enhanced *P. stutzeri* growth. Afterward, the five compounds were taken out one by one in turn, and the growth rates of *P. stutzeri* were recalculated. The maximum predicted *P. stutzeri* growth rate (2.99 \(\times 10^{-2}\) h\(^{-1}\)) was almost unaffected when formate and glycolaldehyde uptake rates were both constrained to 0 mmol g\(^{-1}\) h\(^{-1}\). However, the drops were significant when each of the other three compounds was constrained, suggesting that R-3-hydroxybutanoate, \(\alpha\)-ketoglutarate, and hexadecanoate were key compounds secreted by *Dietzia* sp. to support *P. stutzeri* growth in the C\(_{16}\)-exposed minimal medium. Similarly, the 10 *P. stutzeri* secreted metabolites in Fig. 2 were tested in *iBH925*. Addition of the 10 compounds induced a cell growth rate rise from 2.38 \(\times 10^{-2}\) to 5.53 \(\times 10^{-2}\) h\(^{-1}\). Acetate and L-glutamate were found to be the key metabolites responsible for the significant rise, as the maximum predicted growth rate of *Dietzia* sp. decreased to 2.56 \(\times 10^{-2}\) h\(^{-1}\) if the two metabolites were excluded.

**In vitro validation of *in silico*-predicted key exchanged compounds.** Cells of *P. stutzeri* SLGS10A3-8 were inoculated in the minimal medium supplemented with R-
3-hydroxybutanoic acid, α-ketoglutaric acid, or hexadecanoic acid. As shown in Fig. 3a, *P. stutzeri* biomass accumulated on each of the three compounds after 24 h of incubation, of which α-ketoglutaric acid was preferred by *P. stutzeri* more than the other two compounds. The in vitro result confirmed that in silico-predicted key exchange compounds were able to be utilized by *P. stutzeri* and suggested that the three compounds might be the reason why *P. stutzeri* could survive in the coculture system when C_{16} was the sole exogenous carbon source.

**FIG 2** In silico-predicted interspecific metabolites that induced the growth of *P. stutzeri* SLG510A3-8 and *Dietzia* sp. DQ12-45-1b on C_{16} using iBH1908.

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**FIG 3** In vitro cultivation of *P. stutzeri* SLG510A3-8 and *Dietzia* sp. DQ12-45-1b on in silico-predicted key exchanged compounds. (a) Cell densities of *P. stutzeri* grown on hexadecanoic acid, α-ketoglutaric acid, R-3-hydroxybutyric acid, or no carbon source (control). (b) Time courses of *Dietzia* sp. DQ12-45-1b cell density (graph) and C_{16} removal rates (histogram) in treatments 1, 2, and 3, in which the cultures were supplemented with C_{16} alone, a mixture of C_{16} and acetate, and a mixture of C_{16} and glutamate, respectively.* P < 0.05.
proteins were high- and low-level downregulated, respectively (low-level upregulated in the coculture system, respectively, while 74 and 54 showed at least a 2-fold increase in expression values and were termed high-level upregulated proteins. On the other hand, a total of 91 and 39 upregulated proteins, while the remaining 37 proteins were designated low-level upregulated proteins. Among them, 26 showed at least a 2-fold increase in expression values and were termed high-level upregulated proteins.

Label-free proteomics and extracellular metabolomics analyses. To learn about the metabolic adjustment details of Dietzia sp. and P. stutzeri from their monoculture to the coculture states, Dietzia sp. and P. stutzeri cells were separately inoculated in each side of a two-chamber bioreactor (see Fig. S2 in File S1), cultivated aerobically on C16-exposed minimal medium, and harvested at the mid-exponential phase for proteomics analysis, with the monocultures of each strain as the controls. The results showed that a total of 1,541 Dietzia sp. proteins and 1,292 P. stutzeri proteins were detected from trypsin-digested protein mixtures of the coculture and monoculture samples, with a false-discovery rate (FDR) of <0.05. As shown in Fig. 4a, in the case of cocultivation, a total of 68 P. stutzeri proteins were significantly upregulated compared with those in P. stutzeri monoculture system (P < 0.05), among which 31 proteins showed at least a 2-fold increase in expression values and were termed high-level upregulated proteins, while the remaining 37 proteins were designated low-level upregulated proteins. On the other hand, a total of 91 and 39 P. stutzeri proteins were significantly high-level and low-level downregulated, respectively, in the coculture system (P < 0.05). Similarly, we found that 59 and 30 Dietzia proteins were high- and low-level upregulated in the coculture system, respectively, while 74 and 54 Dietzia proteins were high- and low-level downregulated, respectively (Fig. 4b). The functions of the differentially expressed proteins for the two strains were identified in NCBI and are listed in Tables S3 and S4 in File S1.

Among the differentially expressed proteins, we focused on proteins relating to the metabolism of C16 and the five important interspecific metabolites mentioned above.

| Treatment | Carbon source | Highest specific growth rate (day⁻¹) | P value | Responding time (days) | Responding specific C16 removal rate (pg CFU⁻¹ day⁻¹) | P value |
|-----------|---------------|-------------------------------------|---------|-----------------------|-----------------------------------------------------|---------|
| 1         | C16           | 0.47 ± 0.02                         | 6.99 ± 0.19 | 0.34 ± 0.04 | 0.05 | 1.11 × 10⁻² |
| 2         | C16 + sodium acetate | 0.55 ± 0.03                         | 8.50 × 10⁻² | 7.60 ± 0.25 | 0.54 ± 0.06 | 0.02 | 1.12 × 10⁻² |
| 3         | C16 + sodium glutamate | 0.76 ± 0.26                         | 0.17     | 7.19 ± 0.28 | 0.46 ± 0.05 | 0.01 | 1.27 × 10⁻² |
| 4         | C16 + R-3-hydroxybutanoic acid | 0.47 ± 0.02                         | 0.80     | 7.35 ± 0.28 | 0.37 ± 0.03 | 0.01 | 2.88 |
| 5         | C16 + α-ketoglutaric acid | 0.49 ± 0.06                         | 0.52     | 7.43 ± 0.25 | 0.35 ± 0.04 | 0.02 | 9.46 × 10⁻² |
| 6         | C16 + hexadecanoic acid | 0.82 ± 0.12                         | 4.93 × 10⁻² | 7.56 ± 0.19 | 0.37 ± 0.10 | 0.01 | 5.58 |

*P value between the highest specific growth rates of Dietzia sp. DQ12-45-1b in treatment 1 and the responding treatment.

Values are ± standard deviation.

In order to test if acetate and glutamate promoted C16 biodegradation by Dietzia sp. strain DQ12-45-1b, Dietzia sp. cells were cultivated in the C16-exposed minimal medium with the addition of low concentrations of sodium acetate or sodium glutamate, respectively. Higher Dietzia sp. biomass and lower residual C16 contents in the cultures supplemented with acetate and glutamate (Fig. 3b) indicated the importance of these two compounds in supporting Dietzia sp. growth and C16 biodegradation. Furthermore, the specific C16 removal rate of Dietzia sp. at its highest growth rate in the treatment containing either acetate or glutamate was significantly higher than that in the control (P < 0.05) (Table 1), which was in agreement with above in vitro batch culture result that the specific C16 removal rate of Dietzia sp. in the coculture system was significantly higher than that in the Dietzia sp. monoculture system. At the same time, we tested the effects of the other three compounds, which were in silico predicted to be very actively flowing at the interspecific region in iBH1908 (R-3-hydroxybutanoic acid, α-ketoglutaric acid, and hexadecanoic acid), on Dietzia sp. cell growth and C16 biodegradation. The specific C16 removal rates in Dietzia sp. monoculture systems with the addition of these three compounds were not significantly different from those in the monoculture system without any additives (Table 1), which was inconsistent with our in vitro data mentioned above. This result indicated that C16 biodegradation by Dietzia sp. could be enhanced through the in silico-predicted P. stutzeri secretions but not by some other compounds.

TABLE 1 C16 biodegradation kinetics of Dietzia sp. DQ12-45-1b in monocultures with C16 as the sole carbon source or with the addition of sodium acetate, sodium glutamate, R-3-hydroxybutanoic acid, α-ketoglutaric acid, or hexadecanoic acid
hexadecanoic acid, α-ketoglutaric acid, R-3-hydroxybutanoic acid, acetate, and L-glutamate). Among the differentially expressed proteins, nine proteins in Dietzia sp. and 13 proteins in P. stutzeri were involved in the metabolism of the six compounds; the proteins and their fold changes are listed in Tables S3 and S4. As shown in Fig. 5, the up- and downexpression of the 22 enzymes in the microbial consortium could be interpreted as if the five interspecific metabolites indeed transferred between Dietzia sp. and P. stutzeri in the in silico-predicted directions. Generally, the central metabolism of P. stutzeri became relatively less active in the coculture system than in its monoculture system. In contrast, the tricarboxylic acid (TCA) cycle of Dietzia sp. turned out to be more active when the cells were incubated together with P. stutzeri. This indicated that the P. stutzeri cells grown on Dietzia sp. secretions might reorganize the flux distribution to produce more acetate and L-glutamate for their partner via stripping their intracellular active metabolism unrelated to acetate and L-glutamate biosynthesis down to essentials while expressing more enzymes relating to acetate and L-glutamate biosynthesis.

In addition, culture samples taken from the two-chamber bioreactors were processed to obtain the cell-free supernatant for determination of the extracellular metabolomic composition. Overall, 43 compounds were detected in the coculture and monoculture systems (see Table S5 in File S1). Among the 43 metabolites, the in silico-predicted five key exchanged compounds were all detected in their original or derivatized forms in the supernatant from the coculture system, which was in agreement with above-described proteomic analysis (Fig. 5). Some other in silico-predicted interspecific metabolites were also identified in the metabolome profiles, such as dodecanoate, octadecanoate, and tetradecanoate, as well as their derivatized products, which was in agreement with our in silico prediction (Table 1). However, the remaining identified extracellular metabolites, such as D-lactic acid, glycolic acid, oxalate, inositol, and the derivatized product of ethanol, were out of the computational prediction. This suggested that the two-species metabolic model could be refined further for a better predictive capability.

In silico prediction of the key Dietzia enzyme responding for enhanced C_{16} biodegradation. According to our proteome data, the relative amounts of the known Dietzia alkane hydroxylases AlkW1, AlkW2, and CYP153 showed no significant difference between the monoculture and coculture systems (P ≥ 0.05) (see Table S6 in File S1), indicating that the synergistic C_{16} biodegradation did not depend on the expression level of Dietzia alkane hydroxylases. The key metabolic pathways affecting the C_{16}
FIG 5 Framework showing the regulation of expression of proteins relating to hexadecanoic acid, 3-hydroxybutanoic acid, α-ketoglutaric acid, acetate, and L-glutamate metabolic pathways in Dietzia sp. DQ12-45-1b and P. stutzeri SLG510A3-8 in the artificial consortium when they were exposed to C16. C16 was consumed by Dietzia sp. DQ12-45-1b and was converted to hexadecanoic acid and then to β-oxidation products. Meanwhile, two P. stutzeri enzymes involved in hexadecanoic acid β-oxidation, the long-chain acyl coenzyme A (acyl-CoA) synthetase and the 3-hydroxyacyl-CoA dehydrogenase, had high-level upregulation in the coculture system, indicating that hexadecanoic acid produced by Dietzia sp. was assimilated by P. stutzeri in the coculture system. The higher expression of P. stutzeri enzymes involved in hexadecanoic acid β-oxidation and the high-level downexpression of three P. stutzeri enzymes involved in acetate consumption, acetyl-CoA synthetase, acetyl-CoA C-acetyltransferase, and citrate synthase, would lead to the increased production of acetate. Coincidentally, the Dietzia sp. enzyme aldehyde dehydrogenase responding for acetate production was high-level downregulated when the cells coexisted with P. stutzeri. The lower expression of this protein could be interpreted as if Dietzia sp. indeed utilized extracellular acetate produced by its partner. If this was true, R-3-hydroxybutanoate, a downstream product of acetate, might accumulate in Dietzia sp. and then be secreted to the surrounding environment for P. stutzeri utilization.

Co- and mono-culture systems were set up to study the difference in the metabolic reactions. In mono-culture systems, P. stutzeri were low-level upregulated in the coculture system. P. stutzeri sp. enzymes (α-ketoglutarate dehydrogenase, whose function was the dehydrogenation of (S)-3-hydroxybutanoyl-CoA, was high-level downregulated in the coculture system, while a flagellin enzymealdehydedehydrogenaserespondingforacetateproductionwashigh-leveldownregulatedwhenthecellscoexistedwith P. stutzeri sp., or be secreted to the surrounding environment for P. stutzeri utilization. This inference was confirmed by the facts that an enoyl-CoA hydratase in P. stutzeri, whose function was the hydrolysis of crotonyl-CoA to the R-3-hydroxybutanoate derivative (5)-3-hydroxybutanoyl-CoA, was high-level downregulated in the coculture system, while a P. stutzeri 3-hydroxyacyl-CoA dehydrogenase, whose function was the dehydrogenation of (5)-3-hydroxybutanoyl-CoA to acetoacetyl-CoA, was high-level upregulated. In addition, if acetate was supplied by P. stutzeri to Dietzia sp., α-ketoglutaric acid, a second downstream product of acetate, might accumulate in Dietzia sp. and then be converted to L-glutamate in Dietzia sp. or be secreted to the surrounding environment for P. stutzeri utilization. This inference was confirmed by the facts that the expression of succinyl-CoA synthetase in Dietzia, which used the downstream product of α-ketoglutaric acid as the substrate, was low-level downregulated, while the expression of Dietzia glutamate synthase, which catalyzed L-glutamate production from α-ketoglutaric acid, was low-level upregulated. In P. stutzeri, the high-level downregulation of enzymes relating to L-glutamate consumption, such as glutamine synthetase and gamma-aminobutyric acid (GABA) transferase, suggested that L-glutamate might be accumulated and then be secreted to the surrounding environment for Dietzia sp. utilization. This hypothesis could explain why the glutamine synthetase and GABA transferase in Dietzia sp. were low-level upregulated in the coculture system.
constrained to 2-fold (for high-level upregulation), 1.5-fold (for low-level upregulation), 1/1.5 (for low-level downregulation), or 1/2 (for high-level downregulation) of the responding flux values in \( V_{o_{0.0718}} \). Next, the \( C_{16} \) uptake rate in \( iBH925 \) was modified to be equal to that in \( iBH1908 \) (8.00 \( \times \) 10\(^{-2} \) mmol g\(^{-1} \) h\(^{-1} \)) as mentioned above; and finally, the maximum predicted biomass growth rate \( \mu_{o_{0.08}} \) was computed via FBA again, and \( \mu_{o_{0.08}} \) was compared with \( \mu_{o_{0.0718}} \). If \( \mu_{o_{0.08}} \) with some special reaction flux constraints was similar to \( \mu_{o_{0.0718}} \), the enzymes which catalyzed the special reactions would be considered the key enzymes for the synergistic \( C_{16} \) biodegradation in the synthetic microbial consortium. It was found that the value of \( \mu_{o_{0.08}} \) (2.43 \( \times \) 10\(^{-2} \) h\(^{-1} \)) was similar to that of \( \mu_{o_{0.0718}} \) when the flux rate of \( R_{x13} \) catalyzed by succinate dehydrogenase was constrained, but the value was obviously different from that of \( \mu_{o_{0.0718}} \) when the flux rate modification was operated on any one of the remaining nine metabolic reactions. This suggested that the synergistic \( C_{16} \) biodegradation in the synthetic microbial consortium might be due to the higher-level expression of succinate dehydrogenase induced by the substrates produced from \( P. stutzeri \). The potential molecular mechanism under which the succinate dehydrogenase positively affects \( C_{16} \) biodegradation by \( Dietzia \) sp. was then examined.

Enhanced diesel biodegradation by using the synergistic microbial consortium with the addition of \textit{in silico}-predicted key exchanged compounds. Accordingly, the synergistic biodegradation capability of the microbial consortium was not limited to \( C_{15} \) but also included the synthetic \( n \)-alkane mixture of \( C_{14}, C_{16}, \) and \( C_{28} \) (see Fig. S3 in File S1), and the synergistic effect of the consortium on various \( n \)-alkanes was proved \textit{in silico} to be enhanced by the sustained supply of acetate and glutamate from \( P. stutzeri \) (see Table S7 in File S1). Here, we inferred that the microbial consortium of \( Dietzia \) sp. and \( P. stutzeri \) might have synergistic biodegradation capability on diesel oil, a specific fractional distillate of petroleum composed mainly of alkanes with chain lengths between \( C_{9} \) and \( C_{35} \) (31), and that acetate and glutamate might be able to improve the microbial consortium-based diesel biodegradation. In order to verify the inferences, \textit{Dietzia} sp. without or with the addition of sodium acetate and sodium glutamate was aerobically cultivated on the minimal medium supplemented with 10 g liter\(^{-1} \) diesel oil (treatments 3 and 4, respectively), taking the \textit{Dietzia} sp.-\textit{P. stutzeri} coculture without or with the additives as two positive controls (treatments 5 and 6, respectively) and cell-free culture and the \textit{P. stutzeri} monoculture as two negative controls (treatments 1 and 2, respectively). All the cultures were harvested on day 15 for \textit{Dietzia} sp. cell density measurement and determination of the diesel removal percentage. As shown in Fig. 6a, the cell density of \textit{Dietzia} sp. had a significantly higher increase in the coculture systems than in the \textit{Dietzia} sp. monoculture system, especially in treatment 6. As shown in Table 2, \textit{P. stutzeri} was likely incapable of diesel biodegradation, while \textit{Dietzia} sp. was effective for diesel removal. Notably, the coculture system with the addition of slight amounts of acetate and glutamate had the highest removal percentage (85.54\% ± 6.42\%), which was over 10\% higher than that of the \textit{Dietzia} sp. monoculture (75.33\% ± 3.27\%). Afterwards, we analyzed the hydrocarbon compositions of the disposed diesel oil and found that the composition profile in the coculture system with additives was much simpler than those in the cell-free culture and \textit{Dietzia} sp. monoculture (Fig. 6b), as well as those in \textit{P. stutzeri} monoculture and \textit{Dietzia} sp. monoculture with additives (see Fig. S4 in File S1). According to the composition profiles, \( n \)-alkanes and the isomers with chain lengths between \( C_{12} \) and \( C_{14} \) were completely removed from the diesel samples in treatments 3 and 6, suggesting a very effective removal of the medium-chain-length \( n \)-alkanes and their isomers by \textit{Dietzia} sp.; for the \( n \)-alkanes with chain lengths between \( C_{15} \) and \( C_{25} \), they were partially removed from the diesel sample in treatment 3 but were completely disposed of in treatment 6, leaving small amounts of \( C_{15} \) to \( C_{20} \) isomer residues. Our results suggested that the strategy using the synergistic microbial consortium plus \textit{in silico}-predicted interspecific regulators was very efficient for diesel disposal.
DISCUSSION

It has been reported that the fraction of hydrocarbon-degrading microorganisms is as high as 1% to 10% of the total population in hydrocarbon-polluted environments.

TABLE 2 Diesel removal in treatments 2 to 6 after a 15-day disposal

| Treatment | Bacterium and additives                  | Diesel removal (%) |
|-----------|----------------------------------------|--------------------|
| 2         | P. stutzeri                             | 1.82 ± 0.09        |
| 3         | Dietzia sp.                             | 73.53 ± 3.27       |
| 4         | Dietzia sp. + additives                 | 76.37 ± 4.13       |
| 5         | Dietzia sp. + P. stutzeri              | 83.56 ± 5.66       |
| 6         | Dietzia sp. + P. stutzeri + additives   | 85.54 ± 6.42       |

Values are ± standard deviation.

FIG 6 Diesel biodegradation using Dietzia sp. DQ12-45-1b, P. stutzeri SLG510A3-8, or their consortium with or without the addition of sodium acetate and sodium glutamate. (a) Dietzia cell densities on day 0 (white bars) and day 15 (black bars) in the minimal medium supplemented with 10 g liter⁻¹ diesel (treatment 3), with the mixture of sodium acetate and sodium glutamate (treatment 4), with P. stutzeri (treatment 5), and with both the additive mixture and P. stutzeri (treatment 6). Letters above the bars represent significant differences. (b) TIC-GC/MS chromatograms of diesel oil samples after a 15-day disposal in the cell-free culture (treatment 1), treatment 3, and treatment 6.
which was usually less than 0.1% in unpolluted environments (32, 33). However, the functions of the remaining 90% to 99% of the microbial populations in these hydrocarbon-related ecosystems were unusually ignored. To reveal a potential contribution of these remaining microbial populations, a microbial consortium consisting of an alkane degrader, *Dietzia* sp. DQ12-45-1b, and a potential alkane nonconsumer, *P. stutzeri* SLG510A3-8, was investigated. Surprisingly, the microbial consortium was found to have a greater C₁₆ removal rate than *Dietzia* sp. alone. The synergistic effect was not limited only to C₁₆ degradation but also included degradation of synthetic and natural n-alkane mixtures. This finding was in agreement with cases of bioaugmentation with fungal-bacterial consortia, in which the bacterial members acted as the pollutant degraders while the fungal members did not (34, 35). However, the synergistic interaction between our two bacterial strains should be different from that in these cases, since fungi were referred to as the physical vectors for bacterial transport in these consortia because of use of their hyphae. Accordingly, *Dietzia* sp. was realized to convert C₁₆ into not only its biomass components but also some metabolic by-products available to *P. stutzeri*. In return, *P. stutzeri* fed back some other organic compounds to support the further growth of *Dietzia* sp. and to regulate the expression of some *Dietzia* enzymes, such as succinate dehydrogenase, for improving its C₁₆ uptake.

Succinate dehydrogenase catalyzes the electron transfer from succinate to quinone as shown in the following chemical reaction: succinate + quinone → fumarate + quinol. Succinate dehydrogenase is a central metabolic enzyme coupling the bacterial growth process controlled by the TCA cycle with the energy production resulting from the aerobic respiratory chain. Mammalian cells with succinate dehydrogenase deficiency are diagnosed to have syndromic disease (36). For microbials, there are many reports showing that this enzyme is effective for microbial adaptation under different conditions. Hartman et al. reported that succinate dehydrogenase was a governor of *Mycobacterium tuberculosis* cellular respiration in the adaptation to low-oxygen environments (37). Kiefler et al. significantly increased the carbon conversion efficiency of the industrial obligate aerobic bacterium *Gluconobacter oxydans* from its cytoplasmic oxidized compounds to biomass by knocking in a heterologous gene cluster expressing succinate dehydrogenase and the responding flavinylation factor (38). Considering that petroleum hydrocarbons cause oxidative stress to bacteria during their biodegradation (5), it is inferred that the upregulation of *Dietzia* sp. succinate reductase induced by the *P. stutzeri* secretion was used for the oxidative stress adaptation of *Dietzia* sp. during its C₁₆ biodegradation, and then the degradation rates in the adaptive *Dietzia* sp. were enhanced. Further studies using molecular techniques on *Dietzia* sp. DQ12-45-1b will show whether succinate dehydrogenase regulates C₁₆ biodegradation through oxidative stress adaptation.

Deeply understanding the metabolic interactions among members in microbial consortia with special functions helps to develop efficient microbial consortia or communities in industrial applications. Until now, almost all success stories about the rational design of microbial communities for efficient microbiological processes are based on the following two strategies. The first one is the aggregation method, in which every member could accomplish the tasks of its synthetic microbial consortium but with underperforming efficiencies (39, 40). The second strategy is the “division-of-labor” approach, in which the individual constituents perform complementary roles (41–43). Our study provides a new strategy: the boosting method, in which some members without the desired biological functions promote positive performances of other members who have the desired biological functions or stepwise functions. The microbiological functions of the communities consisting of the supporting and the leading members could be even further enhanced by strengthening their interactions. Taking the consortium of *Dietzia* sp. and *P. stutzeri* as an example, although its synergistic rate of biodegradation of diesel oil was as high as 83.56% ± 5.66%, we found that the cleanup percentage could be further improved by adding slight amounts of two exogenous compounds closely related to the synergistic n-alkane biodegradation (Table 2; see Fig. S3 in File S1 in the supplemental material).
In summary, in our study *Dietzia* sp. DQ12-45-1b and *P. stutzeri* SLGS10A3-8 were found to have a synergistic effect on C\textsubscript{16} biodegradation, even though *P. stutzeri* could not assimilate C\textsubscript{16}. By integrating *in silico* prediction and *in vitro* validation, it was found that *Dietzia* sp. and *P. stutzeri* had a cross-feeding interaction when they were coexposed to C\textsubscript{16}, in which *Dietzia* sp. was the nutrient supplier of *P. stutzeri* while *P. stutzeri* was the n-alkane degradation regulator donator to *Dietzia* sp. Our finding of the synergy mechanism between the two species provides a novel microbial community assemblage strategy for enhancing functions of microbiological processes and helps to deeply understand microbial behaviors in natural ecosystems in the future.

**MATERIALS AND METHODS**

**Bacterial strains and growth conditions.** *Dietzia* sp. strain DQ12-45-1b and *Pseudomonas stutzeri* SLGS10A3-8 were isolated in our laboratory from oil fields in China and deposited in the China General Microbiological Culture Collection Center (CGMCC) (Beijing, China) under accession numbers 1.10709 and 1.15316, respectively. After the two strains were grown in GPY (10 g liter\textsuperscript{−1} glucose, 5 g liter\textsuperscript{−1} yeast extract, 10 g liter\textsuperscript{−1} tryptone) and LB (10 g liter\textsuperscript{−1} NaCl, 5 g liter\textsuperscript{−1} yeast extract, 10 g liter\textsuperscript{−1} tryptone) media, respectively, at 150 rpm and 30°C until the mid-exponential phase, they were centrifuged and washed with the minimal medium (44) three times to make inoculating seed suspensions. Afterward, the two clean seeds were separately inoculated into two flasks or mixed in one flask, each containing 100 ml minimal medium supplemented with 0.7734 g liter\textsuperscript{−1} C\textsubscript{16}. The initial cell density of each strain was an optical density at 600 nm (OD\textsubscript{600}) of 0.1. The culture without inoculum was taken as the control. As described in File S1 in the supplemental material, cocultivation of the two strains was also done in minimal medium supplemented with a synthetic n-alkane mixture consisting of C\textsubscript{14}, C\textsubscript{16}, and C\textsubscript{28} or with diesel oil. Since the colonies of *Dietzia* sp. were orange and smooth, being significantly different from those of *P. stutzeri*, which were brown and wrinkled, samples were taken every 1 to 2 days for cell density counting by CFU until the early stationary phase. The collected data were processed using nonlinear curve fitting followed by differentiation in Origin8.5 (OriginLab Co., MA, USA) to obtain the time courses of specific growth rates.

**Analysis of C\textsubscript{16} removal rates.** At the end of the cultivation period, the residual C\textsubscript{16} in each flask was extracted using 20 ml n-hexane in a separating funnel and then was analyzed using a 5975C gas chromatograph (GC) coupled with a 7890A mass spectrometer (MS) detector (Agilent Technologies, Santa Clara, CA, USA). Helium was applied as the carrier gas at a flow rate of 25 ml min\textsuperscript{−1}. The sample injection volume was 20 ml. One sample was injected at that temperature for 5 min. The external standard method was utilized to obtain the C\textsubscript{16} concentration data were processed in Origin8.5 to obtain the time courses of specific C\textsubscript{16} removal rates. The C\textsubscript{16} removal rate was calculated as the difference in the percentage of the residual C\textsubscript{16} concentration between the experimental and the control treatments in 0.7734 g liter\textsuperscript{−1} C\textsubscript{16}.

**Reconstruction and constraint-based analysis of the two-species metabolic model.** The single-species GEMs of *Dietzia* sp. DQ12-45-1b and *P. stutzeri* SLGS10A3-8 were reconstructed as described in File S1 and then were modified by unifying the format of reactions and metabolites before being integrated to form the two-species model following the approach of Klettgord and Segrè (45). In the two-species model, the metabolites were localized into a total of five compartments as follows: (i) [d] = cytoplasm of *Dietzia* sp., (ii) [s] = extracellular space of *Dietzia* sp., (iii) [p] = cytoplasm of *P. stutzeri*, (iv) [e] = extracellular space of *P. stutzeri*, and (v) [w] = environment shared by the two strains. FBA was used to calculate the metabolic flux distribution in the reconstructed GEMs, in which the biomass reactions accounting for all known biomass constituents and their fractional contributions to the overall cellular biomass were used as the objective to calculate the flux distribution solution in the feasible flux space for the maximal growth rate. System constraints were defined with the upper bound for each metabolic reaction as 1,000 and the lower bounds as –1,000 and 0 for the reversible and irreversible reactions, respectively, except for the exchange reactions. The lower bounds of exchange reactions relating to oxygen and the medium compounds were set to −1,000, while the remaining ones were set to 0. All calculations were performed under the linear programming algorithm using the COBRA toolbox (46) on the MATLAB platform (The MathWorks Inc., Natick, MA, USA).

**In vitro validation of *in silico*-predicted key interspecific compounds.** Seeds of *Dietzia* sp. DQ12-45-1b were inoculated into 100 ml minimal medium supplemented with 0.7734 g liter\textsuperscript{−1} C\textsubscript{16} alone or with the addition of 0.0446 g liter\textsuperscript{−1} sodium acetate, 0.0368 g liter\textsuperscript{−1} sodium glutamate, 0.0283 g liter\textsuperscript{−1} R-3-hydroxybutyric acid, 0.0318 g liter\textsuperscript{−1} α-ketoglutaric acid, or 0.0174 g liter\textsuperscript{−1} hexadecanoic acid at an initial OD\textsubscript{600} of 0.1. The carbon molar contents of the five additives in the experimental treatments were equal to 2.0% of the carbon amount in 0.7734 g liter\textsuperscript{−1} C\textsubscript{16}. The pH in each treatment was adjusted to 8.0 before medium sterilization. All the cultures were incubated at 150 rpm and 30°C. Samples were taken for CFU counting and measurement of the residual C\textsubscript{16} amount until the early stationary phase. To obtain the time courses of the specific growth rates and the specific C\textsubscript{16} removal rates in each treatment, the collected data were processed in Origin8.5 as described above.

Seeds of *P. stutzeri* SLGS10A3-8 were inoculated into 100 ml minimal medium supplemented with 0.872 g liter\textsuperscript{−1} hexadecanoic acid, 1.590 g liter\textsuperscript{−1} α-ketoglutaric acid, or 1.416 g liter\textsuperscript{−1} R-3-hydroxybutyric acid at an initial OD\textsubscript{600} of 0.1. The carbon molar content in each of the three compounds was equal to that in 0.7734 g liter\textsuperscript{−1} C\textsubscript{16}. The culture without any carbon source was taken...
as the control. All the cultures were incubated at 150 rpm and 30°C. Samples were taken for CFU counting after 24 h of cultivation.

**Label-free proteome identification and quantification.** Cells for proteomics analysis were cultivated in a 300-ml two-chamber bioreactor. As shown in Fig. S2 in File S1, the two chambers in the bioreactor were separated by a cellulose acetate filter (pore size = 0.2 μm; Sartorius Stedim Biotech GmbH, Göttingen, Germany). Seeds of *Dietzia* sp. DQ12-45-1b and *P. stutzeri* SLGS10A3-8 were inoculated into different chambers, each of which contained 100 ml minimal medium supplemented with 0.7734 g liter⁻¹ C₁₆. The initial cell density for each strain was an OD₆₀₀ of 0.1. Meanwhile, we developed the *P. stutzeri* monoculture control by inoculating *P. stutzeri* seeds into another bioreactor with an initial OD₆₀₀ of 0.1 in each side and did the same for the *Dietzia* sp. monoculture control. All cultures were kept at 30°C with aeration. Bacterial cells were harvested at day 8 for proteome extraction. Details of proteomic sample preparation are described in File S1. The resulting tryptic peptide samples were loaded into an LCT-Q/Orbitrap-MS (Thermo Fisher Scientific, San Jose, CA, USA). The mobile phase was comprised of solvent A (97% H₂O, 3% acetonitrile, 1% formic acid) and solvent B (99.9% acetonitrile and 0.1% formic acid). The detection of peptide spectra was as described by Liu et al. (47). The proteome qualification was performed using the Mascot software (Matrix Science, Boston, MA, USA) against genome databases for the two strains, while the proteome quantification was based on comparison of the spectrometric counts for selective proteolytic peptides (48).

**Extracellular metabolic profiling.** Cell-free cultures of *Dietzia* sp. DQ12-45-1b, *P. stutzeri* SLGS10A3-8, and their mixture were obtained by centrifugation of 2-ml samples at 4°C and 10,000 rpm for 10 min to remove C₁₆-exposed cells. For desalination, 10 ml methanol was added to each cell-free culture, vortexed for 5 min, and kept at 4°C for 5 h. The mixtures were then centrifuged at 20,000 rpm and 4°C for 10 min, and the supernatant was lyophilized (Freeze-Dryer; Boyikang, Beijing, China) for 4 h. The lyophilized samples were derivatized as described by Mujahid et al. (49). GC-MS analysis of the derivatized samples was performed on a GCMS-QP2010 (Shimadzu, Japan). One microliter of derivatized sample was injected into an RTX-SMS column (30 m by 0.25 mm by 0.25 μm) with helium as the carrier gas at a constant flow of 1 ml min⁻¹. The inlet temperature was set at 250°C. The oven temperature was held at 80°C for 2 min, ramped to 300°C at 20°C min⁻¹, and then held at 300°C for 5 min. Electron impact ionization with an ionization energy of 70 eV and transfer line temperature of 220°C was used. Mass spectra were recorded at 50 to 700 m/z for 5 to 19 min. Metabolites were identified from the GC-MS data based on mass spectral comparison to the standard NIST library 2.0 (National Institute of Standards and Technology, 2008) and Wiley 9 (Wiley-VCH Verlag GmbH & Co. KgaA, Weinheim, Germany), with a match threshold of >80 (with a maximum match equal to 100). Relative metabolite abundances were calculated from peak areas (unique mass) of identified metabolites from GC-MS data and then were amended using the peak area of methyl tetracanoate to minimize instrumental errors.

**In vitro cultivation of the two-species consortium with the addition of predicted exchanged compounds for diesel removal efficiency test.** Seeds of *Dietzia* sp. DQ12-45-1b were inoculated into the minimal medium containing 10 g liter⁻¹ diesel oil with or without the addition of *P. stutzeri* SLGS10A3-8 and exogenous addition of 0.045 g liter⁻¹ sodium acetate and 0.037 g liter⁻¹ sodium glutamate. The culture without inoculum and the one inoculated with *P. stutzeri* SLGS10A3-8 alone were taken as the two control treatments. Every treatment was performed in triplicates. All the cultures were incubated at 150 rpm and 30°C. Samples were taken weekly until the stationary phase or at the end of the culture period for *Dietzia* sp. cell density counting with CFU. At the end of the culture period, the oil removal rates were tested using the weight difference assay as described in EPA method 3500B, and diesel composition profiling was tested using GC-MS. Samples for diesel composition profiling were prepared as described previously. During the GC-MS processing, the oven temperature was set to 50°C for 1 min and then increased to 260°C at a rate of 10°C min⁻¹ and held at that temperature for 5 min, after which it was sequentially increased to 280°C at 10°C min⁻¹ and held at that temperature for 0.5 min.

**Statistical analysis.** All *in vitro* experiments were performed in triplicates and subjected to statistical analysis. The standard deviations were calculated in Microsoft Excel to represent the errors for the triplicate data. The *P* values between two triplicates were also calculated in Microsoft Excel, and a *P* value of <0.05 was considered statistically significant. For the proteomics analysis, proteins with a |log₂(fold change)| of >1 were defined as high-level regulated proteins, and those with a |log₂(fold change)| of <1 were defined as low-level regulated proteins. The significantly changed proteins were annotated according to the KEGG database to identify metabolic pathways in each strain that were influenced by the other one.

**Data availability.** The DNA sequencing data are deposited at GenBank under accession numbers CP046567.1, CP046568.1, and CP046569.1 for *Dietzia* sp. DQ12-45-1b and CP011854.1 for *P. stutzeri* SLGS10A3-8. The mass spectrometry proteomics data reported in this paper have been deposited in the open-access database ProteomeXchange Consortium via the iProX partner repository (50) with the data set identifier PXD016754.

### SUPPLEMENTAL MATERIAL

Supplemental material is available online only.

**SUPPLEMENTAL FILE 1,** PDF file, 1 MB.

**SUPPLEMENTAL FILE 2,** XLSX file, 0.3 MB.

**SUPPLEMENTAL FILE 3,** XLSX file, 0.3 MB.

**SUPPLEMENTAL FILE 4,** XLSX file, 0.5 MB.
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