ORIGINAL ARTICLE

Synthesis of poly- and oligo(hydroxyalkanoate)s by deep-sea bacteria, *Colwellia* spp., *Moritella* spp., and *Shewanella* spp

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The production of poly(hydroxyalkanoate)s (PHAs) as ecofriendly bioplastics by various deep-sea bacteria (4 types of *Colwellia* spp., 11 types of *Moritella* spp., and 18 types of *Shewanella* spp.) from glucose, fructose, gluconate, or from one of the several plant oils as the sole source of carbon was examined at atmospheric pressure. Some of the deep-sea bacteria successfully accumulated PHAs that had a wide range of molecular weights and contained 3-hydroxybutyrate, 3-hydroxyvalerate, and the other hydroxyalkanoate units. Furthermore, with a plant oil as its sole source of carbon, *Shewanella surugensis* produced low-molecular weight oligomeric PHAs. These results provide important and basic information regarding the production of PHAs by deep-sea bacteria and on the diversity of PHA synthase enzymes in nature. *Polymer Journal* (2013) **45**, 1094–1100; doi:10.1038/pj.2013.25; published online 6 March 2013

Keywords: biosynthesis; deep-sea bacteria; poly(hydroxyalkanoate).

INTRODUCTION

Deep-sea bacteria that grow preferentially or exclusively at high hydrostatic pressures and low temperatures have been discovered in various ocean trenches, such as the Mariana Trench. Because of their unique requirements in terms of pressure, temperature, and nutrients, deep-sea bacteria have recently been identified as important bioresources that are capable of providing novel chemicals, enzymes, or genes. It has been reported that the aerobic and anaerobic pathways of fatty-acid biosynthesis in deep-sea bacteria are mutually exclusive of one another and that they differ from the comparable pathways in other organisms.^{1,2} Additionally, the biosynthesis of polyunsaturated fatty-acids (PUFAs) such as (5Z,8Z,11Z,14Z,17Z)-icosa-5,8,11,14, 17-pentaenoic acid (EPA) or (4Z,7Z,10Z,13Z,16Z,19Z)-docosa-4,7,10,13,16,19-hexaenoic acid (DHA) by fermentation with deepsea bacteria has been an attractive process because of the importance of these compounds to human health.³ Bacteria of the genus Moritella biosynthesize DHA, whereas those of the genera Colwellia and Shewanella produce EPA.⁴ Bacteria of the genus Colwellia are strictly psychrophilic (active below 20 °C), and they produce extracellular polymers relevant to biofilm formation and cryoprotection.⁵ Also, enzymes from Colwellia species can degrade polymeric compounds, indicating that Colwellia species might provide a novel method for assimilating polymeric biomass.⁶ Bacteria of the genus Shewanella are deep-sea bacteria that are

capable of biosynthesizing hydrocarbons, such as polyolefins.⁷ These various metabolic pathways for fatty-acid biosynthesis suggest that it should be possible to produce new biochemicals and biopolymers through fermentations with deep-sea bacteria.

Poly(hydroxyalkanoate)s (PHAs) are biodegradable polyesters that are biosynthesized by a variety of bacteria as intracellular materials for the storage of carbon, and their thermal, mechanical, and biological characteristics have been studied extensively.^{8–10} The mechanical properties of PHAs can be improved by means of drawing techniques, resulting in a tensile strength of as high as 1.32 GPa, an elongation-to-the-break of 31%, and a Young's modulus for the drawn PHA fiber of 8.1 GPa.¹¹ These improved mechanical properties of PHAs are comparable with those of industrial samples of polyethylene and polypropylene, and therefore PHAs are an attractive group of materials with respect to their environmental impact, carbon dioxide emissions, and mechanical strength.

PHA is synthesized via glycosylation or β -oxidation pathway (Figure 1). The genes for the synthesis of PHAs reside within an operon that includes a β -ketothiolase (*phaA*), an acetoacetyl-coenzyme A (CoA) reductase (*phaB*), and a synthase (*phaC*).¹² The thiolase and reductase contribute to synthesis of the monomeric substrate 3-hydroxybutyryl CoA, and the synthase polymerizes this monomer to form PHAs.¹² With respect to the synthesis of PHAs by the β -oxidation pathway, *phaJ* and *fabG* genes encoding (*R*)-specific

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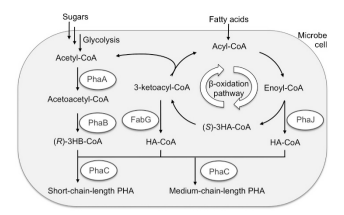


Figure 1 Pathway for the biosynthesis of PHA from sugars and fatty acids. PhaA, β -ketothiolase; PhaB, acetoacetyl-coenzyme A reductase; PhaC, PHA synthase; FabG, 3-ketoacyl acyl carrier protein (ACP) reductase; PhaJ, enoyl-coenzyme A hydratase. A full color version of this figure is available at *Polymer Journal* online.

2-enoyl-CoA hydratase and 3-ketoacyl-acyl carrier protein reductase are known to be responsible for converting 3-enoyl-CoA and 3-ketoacyl-CoA into (R)-3-hydroxyacyl-CoA (Figure 1).^{13,14} PhaC is the key enzyme for the synthesis of high-performance PHAs, because its substrate specificity and specific activity affect the monomeric compositions of the resulting PHA copolymers as final products.¹⁵

In attempts to synthesize new types of PHA, numerous researchers have sought examples of PhaC that show substrate specificities that differ from those of known types of PhaC. However, no attempts have been made to produce PHAs by using deep-sea bacteria as sources of new PhaCs. Also, the biosynthesis pathway of fatty acids in deep-sea bacteria differ from the comparable pathways in soil bacteria, implying a potential to synthesize new monomers via β -oxidation pathway.^{1,2} We therefore selected and characterized a number of deep-sea strains, namely, 4 *Colwellia* spp., 11 *Moritella* spp., and 18 *Shewanella* spp., and we examined their production of PHAs from various carbon sources. The findings of this study are expected to provide important and basic information regarding the production of PHAs by deep-sea bacteria, in addition to improving our knowledge of novel PhaCs.

EXPERIMENTAL PROCEDURES

Cultivation of deep-sea bacteria

Colwellia spp. and *Shewanella* spp. were isolated and stored by the Japan Agency for Marine-Earth Science and Technology. *Moritella* spp. were obtained from RIKEN BioResource Center. All the strains that we used in this study are listed in Supplementary Table S1. These strains were plated on marine agar (Difco; Becton, Dickinson and Co., Franklin Lakes, NJ, USA) and incubated at the optimal temperature for 72 h. The optimal temperatures for the various *Moritella* spp., *Colwellia* spp., and *Shewanella* spp. are listed in Supplementary Table S1.

PHA production

The production of PHAs by 11 types of *Moritella* spp., 4 types of *Colwellia* spp., and 18 types of *Shewanella* spp. was examined on a small scale (4 ml) to screen for PHA-producing strains. Strains were precultured in a marine broth medium (Difco) under aerobic conditions in 10-ml tubes with constant shaking at 140 rpm for 48 h. The PHA-production phase was carried out in MM (mineral medium) medium containing glucose, fructose, gluconate (sodium gluconate), or soybean oil $(20 \text{ g} 1^{-1})$ as the sole source of carbon. Fermentations were performed at the optimal temperature with constant shaking at 140 rpm. for 72 h under aerobic conditions.¹⁶ The MM medium

contained 2.80 g1⁻¹ of KH₂PO₄, 3.32 g1⁻¹ Na₂HPO₄, 0.54 g1⁻¹ (NH₂)₂CO, 0.25 gl^- MgSO₄ · 7H₂O, 19.45 gl⁻¹ NaCl, and 1.0 mll⁻¹ trace elements, consisting of 0.22 g1⁻¹ CoCl₂ · 6H₂O, 9.70 g1⁻¹1⁻¹ FeCl₃, 7.80 g1⁻¹ CaCl₂, $0.12 \text{ gl}^{-1} \text{ NiCl}_2 \cdot 6\text{H}_2\text{O}, 0.11 \text{ gl}^{-1} \text{ CrCl}_3 \cdot 6\text{H}_2\text{O}, \text{ and } 0.16 \text{ gl}^{-1} \text{ CuSO}_4 \cdot 5\text{H}_2\text{O}.$ For larger-scale PHA production, strains were precultured in a marine broth medium (Difco) under aerobic conditions in a 10-ml tube with constant shaking at 140 rpm for 48 h. PHA production was carried out in MM medium supplemented with $20 g l^{-1}$ of glucose, fructose, gluconate (sodium gluconate), jatropha oil, or soybean oil as the sole source of carbon in a 500-ml shaking flask containing 100 ml of medium at the optimal temperature with constant shaking at 140 r.p.m. for 96 h under aerobic conditions.¹⁶ After cultivation, the collected cells were washed with MilliQ water and then ethanol to remove residual carbon sources and salts and then lyophilized. The lyophilized cells were weighed, and the accumulated PHAs were extracted from the dried cells with chloroform for 72 h at 70 °C, and then purified by precipitation with hexane. To determine their PHA content, the dry cells (before PHA extraction) were subjected to methanolysis with a solution consisting of 1.7 ml of methanol, 0.3 ml of 98% sulfuric acid, and 2.0 ml of chloroform at 100 °C for 140 min to convert the constituents into their methyl esters. The addition of 1.0 ml of water to the reaction mixture induced phase separation. The lower chloroform layer was subjected to gas chromatographic (GC) analysis on a Shimadzu GC-17A system equipped with Neutra Bond-1 capillary column $(30\,\text{m}\times0.25\,\text{mm})$ and a flame-ionization detector (Shimadzu Scientific Instruments, Kyoto, Japan). The PHA content was quantitatively determined through detection of the methyl esters of (R)-3-hydroxybutyrate (3HB) and (R)-3-hydroxyvalerate (3HV) units by GC and from the weight of the dry cells.

Characterization of PHAs

The purified PHAs were characterized with respect to their molecular weights and chemical structures. The molecular weights of the polymers were determined by using a gel-permeation chromatography system (RI-2031, PU-2086, AS-2055, C0-2065; JASCO, Tokyo, Japan) with a Shodex K-806 column (Showa Denko, Tokyo, Japan) at 40 °C. Chloroform was used as the eluent at a flow rate of 0.8 ml min⁻¹, and the concentration of the samples was approximately 1.0 mg ml⁻¹. The molecular weight was estimated by comparison to polystyrene standards. The molecular weights of the nine polystyrene standards that we used were 3 148 843, 1 074 876, 460 595, 156 528, 66 001, 28 517, 10 112, 3252, and 1319 g mol⁻¹, respectively. The PHAs were also analyzed by ¹H nuclear magnetic resonance (NMR) spectroscopy (JNM-Excalibur270; JEOL, Ltd, Tokyo, Japan) to determine their precise chemical structures and compositions.

Classification of PhaC genes

Multiple alignment of samples of PhaC genes from *Ralstonia eutropha* (*Cupriavidus necator*, YP_725940), *Aeromonas caviae* (*Aeromonas punctate*, BAA21815), *Shewanella halifaxensis* HAW-EB4 (YP_001674210.1), *Shewanella pealeana* (YP_001501610.1), and *Colwellia psychrerythraea* 34H (YP_270743.1) was achieved by using ClustalW2 software (European Bioinformatics Institute, Cambridge, UK). The amino-acid identities and similarities were calculated by means of the BLAST algorithm.¹⁷

RESULTS AND DISCUSSION

PHA productivity of deep-sea bacteria

Production of PHAs by 4 types of *Colwellia* spp., 11 types of *Moritella* spp., and 18 types of *Shewanella* spp. was examined on a small scale (4 ml) in MM medium supplemented with 20 gl^{-1} of glucose, fructose, gluconate (sodium gluconate), or with soybean oil as the sole source of carbon. The PHA contents of the bacteria were characterized by means of GC as proportions of the weights of the dry cells (Supplementary Tables S2–S4). *Colwellia* spp. other than *Colwellia* sp. strain JAMM-0421 showed no cell growth or PHA production (Supplementary Table S2). *Colwellia* sp. strain JAMM-0421 produced PHAs when glucose, fructose, or soybean oil was used as the sole source of carbon. In some cases, *Moritella* spp. showed cell growth and PHA productivity, as listed in Supplementary Table S3.

The PHA contents of *Moritella* spp. were <1% and were lower than those of Colwellia sp. strain JAMM-0421. The Shewanella spp. also showed relatively low PHA productivities (<1% PHA content), and these were independent of the type of carbon source or the strain. The results of these studies (Supplementary Tables S2-S4) did allow us to select three strains, a Colwellia sp. (JAMM-0421), a Moritella sp. (JCM21335), and a Shewanella sp., S. surugensis (JAMM-0386), for further studies because these strains produced higher PHA contents than did other species of each respective genus.

PHA production by a Colwellia sp., a Moritella sp., and S. surugensis

Three deep-sea strains, a Colwellia sp. (JAMM-0421), a Moritella sp. (JCM21335), and S. surugensis (JAMM-0036), were characterized in terms of their abilities to synthesize PHAs from glucose, fructose, gluconate, jatropha oil, or from soybean oil as the sole source of carbon in 100-ml-scale cultures at the appropriate optimal temperatures for 96 h (Table 1). The compositions of soybean oil and jatropha oil differ as listed in Supplementary Table S5, which would affect PHA productivity of the deep-sea strains. On the basis of the ¹H NMR spectra (Figure 2), Colwellia sp. (JAMM-0421) accumulated poly((R)-3-hydroxybutyrate-co-(R)-3-hydroxyvalerate) (P(3HB-co-3HV)) when glucose, fructose, jatropha oil, or soybean oil was used as the sole source of carbon. The accumulation of PHA by Colwellia sp. (JAMM-0421) fed with plant oils resulted in higher PHA contents than those produced by other deep-sea bacteria (Table 1). The monomer composition of the resulting PHA was characterized by means of ¹H NMR spectroscopy (Figure 2); this showed that the 3HV contents ranged from 2.4 to 29.0 mol%, based on the integration of the peaks corresponding to the methyl protons (3 and 7 in Figure 2). The number-average molecular weights of the P(3HB-co-3HV) synthesized by Colwellia sp. strain JAMM-0421 ranged from 170×10^3 to $340 \times$ 10^3 g mol⁻¹ (Table 1 and Figure 3b); these values are in the normal range of molecular weights for bacterial polyesters. With respect to productivity, monomeric composition, and molecular weight of PHA, Colwellia sp. strain JAMM-0421 showed no obvious differences from soil bacteria.18

Although the Moritella sp. strain JCM21335 also produced P(3HBco-3HV) with a number-average molecular weight of $120 \times$ 10³ g mol⁻¹ from fructose (Figures 3f and 4a), no PHA accumulation was detected when glucose and gluconate were used as the sole source of carbon (Table 1). Moritella sp. strain JCM21335 produced polymeric materials with number-average molecular weights of $1000-30000 \,\mathrm{g}\,\mathrm{mol}^{-1}$ from soybean oil or jatropha oil (Figures 3d and e). The ¹H NMR spectra of the polymers produced from plant oils by Moritella sp. strain JCM21335 showed that no 3HB and 3HV units were present. However, methine, methylene, and methyl protons that could be assigned to hydroxyalkanoate units were detected (Figure 4b). The methylene peaks ($\delta = 2.0-3.0$ p.p.m.) were assigned to the methylene protons of the main chains, in accord with previous report.^{18,19} Thus, the Moritella sp. strain ICM21335 produced polymers with different main chain architectures depending on the source of carbon. Generally, sugars and fatty acids are processed to PHA mainly via glycosylation and β-oxidation pathways, respectively. Based on nature of the synthesized polymers, the PhaC of the Moritella sp. appears to have different substrate specificity, especially along the main-chain direction, from that of known PhaCs, such as those of Ralstonia eutropha or Aeromonas caviae. R. eutropha and A. caviae are known as typical soil bacteria to produce PHA.12,20

Strain	Carbon source	Dry cell weight (g l ⁻¹)	PHA content (wt%)	PHA compo- sition (mol%) ^a		Number-average mole- cular weight (g mol ⁻¹)
				ЗНВ	3HV	(PDI) ^b
Colwel	lia sp. (JAM	IM-0421)				
	Glucose	1.7 ± 0.1	2.0 ± 0.1	71.0	29.0	$340 imes 10^3$ (1.5)
	Fructose	1.7 ± 0.1	2.7 ± 0.1	89.5	10.5	$300 imes 10^3$ (2.8)
	Guconate	0.6 ± 0.1	trace	100	_	$99 imes 10^3$ (1.9)
	Jatropha oil	1.4 ± 0.2	16.4 ± 1.1	96.0	4.0	170×10^3 (2.9)
	soybean oil	1.3 ± 0.1	10.9±1.6	97.6	2.4	230×10^3 (3.0)
Morite	lla sp. (JCM	21335)				
	Glucose	0.7 ± 0.2	_	_	_	_
	Fructose	1.3 ± 0.1	3.4 ± 1.0	90.5	9.5	120×10^3 (1.8)
	Gluconate	0.6 ± 0.1	0.7 ± 0.1	_	_	_
	Jatropha oil	2.4 ± 0.3	0.7 ± 0.1	ND	ND	1000–30 000
	Soybean oil	1.2 ± 0.1	1.6 ± 0.5	ND	ND	1000-30000
Shewa	nella suruge	ensis (JAMN	1-0036)			
	Gucose	0.7±0.1	_	_	_	_
	Fructose	0.1 ± 0.1	Trace	_	_	_
	Gluconate	0.1 ± 0.1	_	_	_	_
	Jatropha oil	1.9±0.3	0.4 ± 0.2	Trace	—	1.0×10^3 (1.1)
	Soybean oil	1.2±0.2	0.6±0.2	Trace	—	1.0×10^3 (1.1)

Abbreviations: 3HB, (R)-3-hydroxybutyrate (3HB) 3HV, (R)-3-hydroxyvalerate; ND, not

determined; PDI, polydispersity index; PHA, poly(hydroxyalkanoate) ^aDetermined by ¹H nuclear magnetic resonance spectroscopy.

^bPolydispersity index (determined by gel-permeation chromatography).

S. surugensis accumulated oligomeric PHAs when a plant oil was used as the sole source of carbon (Table 1). The ¹H NMR spectra showed that the structure of the oligomers synthesized by S. surugensis was similar to that of the polymer synthesized by Moritella sp. (Figures 4b and c). However, in the ¹H NMR spectra of the polymer synthesized by S. surugensis, the peaks at around $\delta = 2.55$ p.p.m., which were assigned to methylene protons of the main chain, differed from those in the corresponding spectrum of the PHA from the Moritella sp. The molecular weight of the polymer synthesized by S. surugensis was markedly lower and its polydispersity index was markedly narrower than the corresponding values for the polymer from the Moritella sp. (Figure 3c). Because of the low molecular weight of the oligomeric product, we expected to be able to characterize the chain-end structures of the polymer by ¹H NMR spectroscopy; however, no peaks originating from the chain ends could be assigned.^{21,22} This might be because the synthesized polymers were blends of oligomers with various chemical structures, rather than pure copolymers. An influence of the PhaC on the molecular weight of the synthesized PHA has previously been reported for a series of mutant PhaCs from A. caviae and Pseudomonas sp. 61-3.23,24 These studies found that a difference of one order of magnitude in the molecular weight of PHAs could be induced by mutation of the PhaC. Alteration of the function of the PhaC cannot, therefore, completely explain the observed reduction in the molecular weight of the PHA to that of the oligomer synthesized

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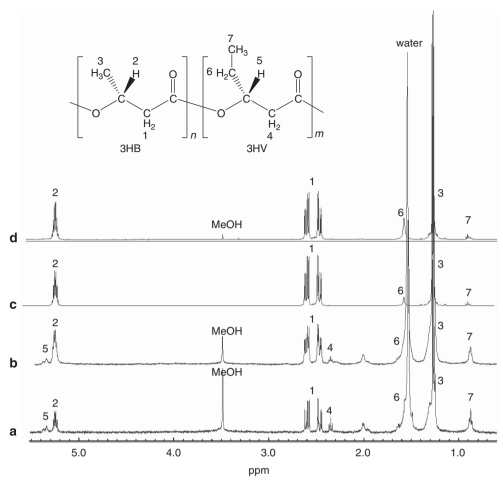


Figure 2 ¹H NMR spectra of samples of PHA synthesized by Colwellia sp. from (a) glucose, (b) fructose, (c) soybean oil, or (d) jatropha oil.

by *S. surugensis* with plant oils as the sole source of carbon (~1000 g mol⁻¹). The metabolic pathway of the strain, especially its β -oxidation pathway, must also affect the molecular weight of PHA. No previous method has been established for the biosynthesis of oligomeric PHAs, which might be useful as raw materials for cosmetics or as pharmaceutical products.²⁵ The synthesis of the oligomeric polyester from plant oils by *S. surugensis* therefore has potential as a new method for producing oligomeric PHAs for practical use.

The major difference among the three genera is found in their biosynthesis behavior for PUFAs, such as EPA or DHA.²⁶ As explained in the introduction, organisms of the genus Moritella generally biosynthesize DHA, whereas those of the genera Colwellia and Shewanella produce EPA.^{1,27} Both bacterial PUFAs are synthesized by polyketide synthase system, and DHA is synthesized via EPA.4,26 Although we could not evaluate how differences in the biosynthesis of PUFAs contributed to PHA synthesis in the current study, deep-sea bacteria with different pathways for PUFA synthesis did show differences in their productivities of PHAs. Under some culture conditions, cell growth of the strains was insufficient to permit evaluation of their PHA productivities (see Supplementary Tables S2-S4). Furthermore, the synthesized polymers could not be adequately characterized to permit determination of their chemical and chain-end structures. To determine these details of PHA productivity of the deep-sea strains in future experiments, we plan to perform large-scale cultivations (for example, 10-liter-scale fermentations)

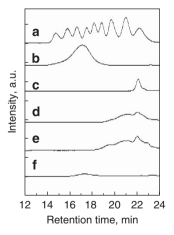


Figure 3 Gel-permeation chromatographic retention times of (a) polystyrene standards (molecular weights: 3148,843, 1074876, 460595, 156528, 66001, 28517, 10112, 3252, and 1319g mol⁻¹) and of PHA samples synthesized from various substrates by deep-sea strains: (b) from soybean oil by *Colwellia* sp., (c) from soybean oil by *Shewanella surugensis*, (d) from soybean oil by *Moritella* sp., (e) from jatropha oil, and (f) from fructose by *Moritella* sp.

under aerobic and anaerobic conditions and also to construct transgenic *R. eutropha* harboring samples of PhaCs originating from deep-sea bacteria to evaluate their substrate specificities.

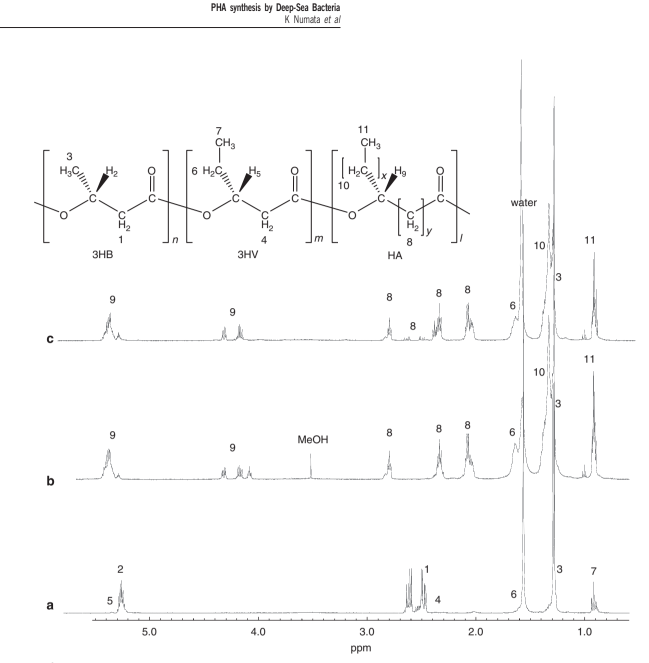


Figure 4 ¹H nuclear magnetic resonance spectra of oligomers and polymers synthesized (a) by *Moritella* sp. from fructose, (b) by *Moritella* sp. from soybean oil, and (c) by *Shewanella surugensis* from glucose. A full color version of this figure is available at *Polymer Journal* online.

PhaCs from deep-sea strains

PhaCs are key enzymes for the synthesis of PHAs and they have been classified into four classes with respect to their amino-acid sequences.¹⁵ Class I PhaCs consist of single subunits with molecular weights between 61 and 68 kDa. The PhaCs from *R. eutropha* and *A. caviae*, which have been widely studied, are known to be Class I PhaCs that exhibit different substrate specificities.^{15,20,28,29} The PhaC from *A. caviae* is an exceptional PhaC in Class I because of its low similarity (approximately 45%) to the other members of the group of Class I PhaCs and because of its substrate specificity for C₄ and C₆ monomers.^{30,31} The genomes of the deep-sea bacteria *Shewanella halifaxensis*, *S. pealeana*, and *Colwellia psychrerythraea* have been sequenced, and the appropriate PhaC sequences have been identified. On the basis of their amino-acid sequences, the PhaCs from *S. halifaxensis*, *S. pealeana*, and *C. psychrerythraea* have been classified as Class I PhaCs. To evaluate the difference between the

amino-acid sequences of PhaCs of deep-sea bacteria and those of soil bacteria, multiple alignments of the amino-acid sequences of PhaCs from S. halifaxensis, S. pealeana, and C. psychrerythraea with those from A. caviae and R. eutropha H16 were performed (Figure 5). The PhaC genes of S. halifaxensis and S. pealeana showed 92% identities in terms of their amino-acid sequences, which confirmed the PhaCs within a particular genus show high degrees of identity in terms of their amino-acid sequences. The amino-acid sequence of the PhaC from R. eutropha H16 showed 38, 42, and 43% identities to those of the PhaCs from C. psychrerythraea, S. halifaxensis, and S. pealeana, respectively. The amino-acid sequence of PhaC from A. caviae showed 40, 55, and 54% identities to those from C. psychrerythraea, S. halifaxensis, and S. pealeana, respectively. The identities between the PhaC from C. psychrerythraea and those from the two types of Shewanella were approximately 42%. Based on the yields, compositions, and molecular weights of PHA

Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	MESKSPFQDTIDNALQFGQAWMDSFGQSAQSCIVETQVEDLAQWMRSSVEH-PVNSIEQQMNWWGQQVNLFNDCIMSSLPTEKETDRRFKDPAWNEQALY MESKSPFQDAIDNAMQFGQAWMDSFGQSAQSSIVETQAEDWAQWMRSSVEH-PVNSIEQQMDWWGQQVNLFNDCIMSSLPTEKETDRRFKDPAWNEQALY MSQPSYGPLFEALAHYNDKLLAWAKAQTERTAQALLQTNLDDLGQVLEQSVEH-PVNSIEQQMDWWGQQVNLFNDCIMSSLPTEKETDRRFKAEAWSEQPIY MSSPSYGPLFEALAHYNDKLLAWAKAQTERTAQALLQTNLDDLGQVLEQ	98 108 111
Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	KYIKESYKLACNNVQSSLDNTEGLDDETRQRLSFYTRQYLNAMSPSNFVATNPEIMKLTVESKGQNLIKGLEQLQKDLAQS-ADTLNIRMTDKNAFTIGKNIATAPGKVVFKNDLFELIQ KYIKESYKLACNNIQASINNTEGLDDETRQRLSFFSRQYLNAMSPSNFVATNPEIMKLTIESKGQNLIKGLEQLQQDLEQS-VDTLNIRMTDKTAFTVGKNIATTPGKVVFKNDLFELIQ DYLKQSYLLTARHLLASVDALGUVPQKSRELRFFFRQYNAMASPSNFVATNPEIMKLTIESGGQNLIKGLEQLQQDLEQS-VDTLNIRMTDKTAFTVGKNIATTPGKVVFKNDLFELIQ NYLKQSYLLSSNMMQDTVNSLHPDDFXAEGUKFYTRQYISSYSFTNNALTNPEVGRDILESGGSLARGVENMMEDLESS-ADQLIXIRIDESSAFELGRDLAITPGKVVFKNDLYQN RFAARFYLLNARALTELADAVEA-DAKTRQRIRFAISGWVDAMSFANFLATNPEAQRLIESGGESLARGVRNMMEDLTRGKISQTDESAFEVGRNVAVTEGAVVFENEYFQLIQ : * * * * * * * * * * * * * * * * * * *	217 227 231 227
Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	YQATTEQVYKRPLLVVPFFVNKFYINDLSPERSYTQULVSQGHTVFMISWINPNADMATTDFGDYVTQGIIPALDAIEVETGEREVNGIGYČIGGTLLTAMAYLAGKRRKQRVKSATLL YQATTEQVYKRPLLVVPFFVNKFYINDLSPERSYTQULVSQGHTVFMISWNIPNAEMAATDFGDYVTQGVILALDAIEAETGEREVNGIGYČIGGTLLTAMAYLAGKRRKQRVKSATLL YSFTETVGVKRPLLVPFFVNKYYLDMEQOSLIKUAQGQTVFWISWNIPAAAQAQIDLDDVVDGVIALDGVENAATGEREVNGIGYČIGGTLALSAMAKIAARRCKQRVRSAT YKASTKVMEVPLLFIPFFINKYYLDMDEQOSLIKWAVGQGHTVFVISWINPAKTSKRFGDYMKGETASLDVIEELTKSKKVNVVGCIGGTLALALAULAARRCKQRVFG YKLEVFULVFFUNKYYLIDLGPESSLVRHVVEQGHTVFISWNIPAKAAGSTMDDYIEMAIRAEVANDISGDKINVLGFCVGGTIVSTALLAVLAARCFF ***********************************	337 347 350
Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	TTILDFGQFGELGVFINEFLISSIEAQNNARGYMDGRQMAVSFSLLRENSLYWNYYVTNYLKGESPMAFDLLHWNCDNTNITAATHNQLLRQMYLENKLKEPGGIAIDGVNIDLS TTILDFGQFGELGVFINDFLISSIEAQNNARGYMDGRQMAVSFSLLRENSLYWNYYVTNYLKGESPVAFDLLHWNCDNTNITAATHNQILRQMYLENKLKEPGGIAIDGVNIDLS TTILDFSQFGELGVFINEFLIALEAQWEAKGIMDGRQLAVSFSLLRENSLYWNYYVTNYLKGESPVAFDLLHWNDDSTNVAGK'HNSLLRLYLENQUVK-GELKINTTADIG TTLLDFEGEGIGVYLEGKULPFIKKDIKEKGLLDGRVLGLAFSMLRENSLFWSYFVNYLKKGPAFPDILHWNDDSTNVAGK'HNSLLRLYLENQUVK-GELKINTTADIG TTLLDFEGEGIGVYLEGKULPFIKKDIKEKGLLDGRVLGLAFSMLRENSLFWSYFVNYLKKGPAFPDILHWNDDSTNVAGK'HNSLKNFLKUNGAVEVEGPVFILS TTLLDFADTGILDVFVDEGHVQLREATLGGGAGAPCALLRGLELANTFSFLRPNDLVWNYVVDNYLKGNTPVPFDLLFWNGDATNLPGPWYCWYLRHTYLQNELKVPGKLTVCGVPVDLA ************************************	452 461 465
Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	KVKSPSYFLSAIEDHIAVWEGTFRGTELLNGDNTFVLAESGHIAGPMNPPNANKYGFWTNSNNKQSPLTWLAEADSHQGSWNPHWQSWIDERN-FSEKIAARSLTGNLDAPGEYV KVKSPCYFLSAIEDHIAVWEGTFRGTELLNGDNTFVLAESGHIAIINPANKYGFWTNSNNKQSPLTWLAEADSHSGSWWPHWQSWIDERN-FSDKIAARSLTGNLDAPGEYV KVKTPVLLVSAIDHIALWGGTWGGMKHGGEQFFLLAESGHIAIINPPANKYGFWTNSAEASSESWLAGATHQGGSWNPEMMOFIQNRDESEPVPARVPEGLAPAGEYV KIDTPGYFLATIADHIVLWQASYKGKALSGPTFFVLAESGHIAIINPPANKYSHWTNDALESDTAEQWEBSSTKKGSWWTDWENVUEPGS-GKQVTARKQGSKYPSIBAAFGSV SIDVPTYIYGSREDHIVPWTAAXASTALLANKLFFVLAGSGHIAGVINEVDHGKYPHWNESHWTNDALPBSPQQWLAGAIEHHGSWWPDWTAWLAGQAARNAPRPPTMAMRYRAIEPAFGXV ***	566 577 583
Shewanella halifaxensis Shewanella pealeana Aeromonas caviae Colwellia psychrerythraea Ralstonia eutropha	KQRIQDVIALKEEVHHGA 585 KQRIEDVIAPKEEVHHT 584 KVRLNPVFACPTEBDAA 594 RKRI 587 KAKA 588 :	

Figure 5 Multiple alignment of poly(hydroxyalkanoate) (PHA) synthases (PhaCs) originating from *Shewanella halifaxensis, Shewanella pealeana, Aeromonas caviae, Colwellia psychrerythraea,* and *Ralstonia eutropha* H16. Asterisks denote a putative catalytic triad. The arrow indicates Ala-505 for PhaC from *A. caviae.* A full color version of this figure is available at *Polymer Journal* online.

(Supplementary Tables S1 and S2–S4), differences in the degree of identity between soil bacteria and deep-sea bacteria might not be sufficient to explain the differences in the functions of their respective PhaCs.

Several groups have reported effects of mutations of PhaCs, such as PhaCs from A. caviae, Pseudomonas sp. 61-3, R. eutropha, or Allochromatium vinosum.23,24,32-37 A mutation at position Ala505 (indicated by an arrow in Figure 5) of the PhaC from A. caviae affected the composition and molecular weight of the PHA.²³ Position Ala505 is a neighboring position to an active site (His503), as indicated by the asterisk in Figure 5. The deep-sea strains shown in Figure 5 have an alanine moiety in the position corresponding to Ala505 of PhaC from A. caviae, implying that this position is not of major significance in terms of controlling the molecular weight of PHA over several orders of magnitude. This finding and the results in Table 1, taken together, indicate that PhaCs originating from deep-sea bacteria have the potential to regulate the molecular weight of PHAs and to take up new monomer units, especially when the β-oxidation pathway is used to produce monomers similar to those deep-sea bacteria. The present study also showed that PhaCs from deep-sea bacteria are useful research targets in producing novel types of PHA.

CONCLUSIONS

This is the first investigation of the production of PHAs at normal pressures by three genera of deep-sea bacteria: *Colwellia, Moritella*, and *Shewanella*. We attempted to produce PHAs by using 33 strains of deep-sea bacteria with glucose, fructose, gluconate, or with plant oils as the sole source of carbon. In several combinations of strain and carbon source, the deep-sea bacteria successfully accumulated PHAs that contained different monomer units and had different molecular weights from those of the corresponding polymers produced by soil bacteria. It is noteworthy that with plant oils as sole sources of carbon, *S. surugensis* accumulated oligomeric PHAs. Hydroxyalkanoate units were also incorporated into the PHA accumulated by a

Moritella sp. JCM21335 strain when a plant oil was used as the carbon source. This finding provides a potential method for controlling the biosynthesis of oligomeric PHAs, useful as raw materials for cosmetics or as pharmaceutical products, by using deep-sea bacteria and/or PhaCs from deep-sea bacteria. Furthermore, this new insight provides a biotechnological option for the production of various polymeric materials and chemicals, including PHA and its oligomers, by deepsea bacteria, permitting the possible efficient utilization of deep-sea bioresources.

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