単細胞および多細胞の紅藻類(紅色植物門)のポリアミン構成

誌名	日本微生物資源学会誌
ISSN	13424041
著者名	浜名,康栄
	小林,正樹
	古地,壯光
	林,秀謙
	新津,勝
発行元	日本微生物資源学会
巻/号	34巻2号
掲載ページ	p. 83-91
発行年月	2018年12月

農林水産省 農林水産技術会議事務局筑波産学連携支援センター

Tsukuba Business-Academia Cooperation Support Center, Agriculture, Forestry and Fisheries Research Council Secretariat



Polyamine distribution profiles in unicellular and multicellular red algae (phylum Rhodophyta)
—Detection of 1,6-diaminohexane, aminobutylcadaverine, canavalmine and aminopropylcanavalmine—

Koei Hamana^{1)*}, Masaki Kobayashi²⁾, Takemitsu Furuchi²⁾, Hidenori Hayashi¹⁾ and Masaru Niitsu²⁾

 ¹⁾Faculty of Engineering, Maebashi Institute of Technology 460-1 Kamisadori-machi, Maebashi, Gunma 371-0816, Japan
 ²⁾Faculty of Pharmacy and Pharmaceutical Sciences, Josai University 1-1 Keyakidai, Sakado, Saitama 350-0295, Japan

To consider the phylogenetic significance of cellular polyamine distribution profiles in multicellular red-algal evolution, polyamines procured through acid extraction from 27 additional red algae species (29 strains) including 18 species (19 strains) of marine multicellular species (seaweeds) were newly analyzed by HPLC and HPGC-MS, and compared with 21 previously analyzed red-algal polyamine profiles. In the unicellular thermoacidophilic order Cyanidiales, Cyanidium and Cyanidioschyzon contained putrescine, spermidine and spermine, and Galdieria contained norspermidine and norspermine in addition to the three polyamines. Freshwater/marine unicellular Porphyridium contained putrescine and spermidine, furthermore, spermine was found in some Porphyridium species. In the marine unicellular orders Dixoniellales, Rhodellales and Stylonematales, the presence of putrescine, spermidine, norspermidine, spermine and norspermine was detected. Of the five polyamines, Bulboplastis containing thermospermine instead of spermine, has a novel algal polyamine profile. In the freshwater macroalgal orders Thorelales, Batrachospermales and Compsopongonales, the presence of putrescine, spermidine, homospermidine and spermine was abundant. In red seaweeds belonging to seven orders of the class Bangiophyceae or Florideophyceae, 1,3-diaminopropane, putrescine, cadaverine, spermidine, norspermidine, homospermidine, spermine, norspermine and thermospermine were widely distributed. 1,6-Diaminohexane was distributed within 12 species, aminobutylcadaverine was found in Gelidium and Hypnea, and canavalmine and aminopropylcanavalmine were detected in Meristotheca, as unusual algal polyamines. The distribution of norspermidine and norspermine was particular in marine unicellular species. Homospermidine was a special polyamine member for freshwater multicellular species. I,6-Diaminohexane and penta-amines were found in seaweeds in addition to common red-algal polyamines. The polyamine profiles of red seaweeds distinguished them from those of green seaweeds (ulvophytes) and brown seaweeds.

Key words: aminobutylcadaverine, aminopropylcanavalmine, canavalmine, diaminohexane, polyamine, red algae, Rhodophyta

INTRODUCTION

As a contribution to advances in cellular polyamine biosynthetic diversity in the evolution of algae (Fuell *et al.*, 2010; Minguet *et al.*, 2008; Shaw *et al.*, 2010; Schweikert & Burritt, 2015), we have analyzed polyamines of various algae which evolved after primary, secondary and tertiary symbiotic processes (Hamana, 2008; Hamana & Matsuzaki, 1982, 1985; Hamana & Niitsu, 2006; Hamana *et al.*, 1990, 2004a, 2004b, 2013, 2016b, 2017, 2018). The four algal phyla Glaucophyta, Rhodophyta, Chlorophyta and Streptophyta have plastid(s) as a result of the primary endosymbiosis of a phototrophic cyanobacterium, which evolved independently without secondary and tertiary endosymbiosis, and are located together in the Archaeplastida (unranked). Glaucophytes, such as *Cyanophora* and *Glaucocystis* species, evolved in a primitive unicellular form, which contained putrescine and spermidine alone as their cellular poly-

^{*}Corresponding author

E-mail: koeihamana@gmail.com

Accepted: September 25, 2018

amines (Hamana & Matsuzaki, 1985; Hamana & Niitsu, 2006). Unicellular green algae evolved into colony forming green algae and multicellular ulvophytes (green seaweeds) within the phylum Chlorophyta. Multicellular charophytes belonging to the phylum Streptophyta transited to the ancestor of early land plants. It has been shown that the distribution profiles of norspermidine, homospermidine, norspermine, spermine, thermospermine and penta-amines are significant within multicellular green algal evolution (Hamana *et al.*, 2004b, 2013, 2018).

Although the polyamines of 21 red algae located in the two subphyla Cyanidiophytina and Rhodophytina of the phylum Rhodophyta (Yoon *et al.*, 2006) (Table 1) have been analyzed, thermospermine and penta-amines have never been detected, however, one red seaweed was included in the previous studies (Hamana & Matsuzaki, 1982, 1985; Hamana & Niitsu, 2006; Hamana *et al.*, 1990), as cited in Table 1.

As shown in Table 1, three new freshwater unicellular thermoacidophilic red algae belonging to the order Cyanidiales (class Cyanidiophyceae) have been isolated in Japan and Italy. Four freshwater/marine unicellular red algae belonging to the order Porphyridales (class Poryphyridiophyceae) and two new marine unicellular red algae belonging to the order Dixoniellales, Rhodellales (class Rhodellophyceae) or Stylonematales (class Stylonematophyceae) have been collected in Japan. Two new freshwater multicellular (macro) red algae belonging to the order Thoreales, Batrachospermales (class Florideophyceae) or Compsopogonales (class Compsopogonophyceae) have been identified as endangered red algae in Japan. On the other hand, many marine multicellular (macro) red algae belonging to the order Bangiales (class Bangiophyceae), Creamiales, Halymeniales, Gelididales, Gracilariales or Gigartinales (class Florideophyceae) are considered edible seaweed in Japan.

Recently, the subphylum Rhodophytina was divided into the two subphyla Proteorhodophytina (consisting of the classes Poryphyridiophyceae, Compsopogonophyceae, Rhodellophyceae and Stylonematophyceae) and Eurhodophytina (consisting of the classes Bangiophyceae and Florideophyceae) (Muñoz-Gómez *et al.*, 2017) (Table 1).

To reveal the phylogenetic, morphologic and growth-related environmental significance of cellular polyamine profiles in red-algal evolution, and furthermore, to compare the polyamine profiles of red seaweeds with those of green seaweeds and brown seaweeds, a polyamine analysis of an additional 29 red-algal samples including 19 red seaweeds (as shown in Table 1) was performed in the present study.

MATERIALS AND METHODS

The red algal strains supplied from the Microbial Culture Collection at the National Institute for Environmental Studies (MCC-NIES) and the Biological Resource Center, National Institute of Technology and Evaluation (NBRC) were cultivated phototrophically in the light (10-14 h/24 h) at 20-25°C (at 40°C for Cyanidium and Galdieria) using 1-10 l of liquid medium designed by MCC-NIES (http://mcc.nies.go.jp, 2017) and NBRC (https:// www.nite.go.jp/en/nbrc, 2017). The cultures of the non-axenic unicellular Porphyridium aerugineum NIES-1957 and Bulboplastis apyrenoidosa NIES-2742 contained some bacterial contaminants, and so the red algae were purified by sonication in an ultrasonic washing machine following differential centrifugation. The dry and wet red-algal seaweeds "Susabinori" (Pyropia yezoensis), "Narawasusabinori" (P. yezoensis form. narawaensis), "Kaigara-amanori" (P. teruipeddalis), "Uppuruinori" (P. pseudolinearis), "Egonori" (Campylaephora hypnaeoides), "Matsunori" (Polyopes affinis), "Makusa" ("Tengusa") (Gelidium elegans), "Ogonori" (Gracilaria vermiculophylla), "Yumigata-ogonori" (G. arcuata), "Fukuro-funori" (Gloiopeltis furcate), "Ma-funori" (G. tenax), "Aka-tosakanori" (Meristotheca papulosa), "Akaba-ginnansou" (Mazzaella japonica), "Kotoji-tsunomata" (Chondrus elates), "Tsunomata" (C. ocellatus), "Ibaranori" (Hypnea charoides), "Isodantsu" (Caulacanthus ustulatus) and "Dulse" (Palmaria palmata) were purchased from food markets in the habitats in Japan or Korea (Table 1).

Wet red algae harvested at the stationary phase in our laboratory (1-10 g wet weight) and the red seaweeds rehydrated and washed by an ultrasonic washing machine (100-200 g wet weight), were homogenized in 5% perchloric acid (PCA). The PCA extract was subjected to a column containing a cation-exchange resin, Dowex 50WX8 (3 cm I.D. ×1 cm or 1 cm I.D. ×3 cm), to concentrate the polyamines. The polyamines were eluted with 6M HCl and then analyzed with by high-performance liquid chromatography (HPLC) on a Hitachi L6000 using a column

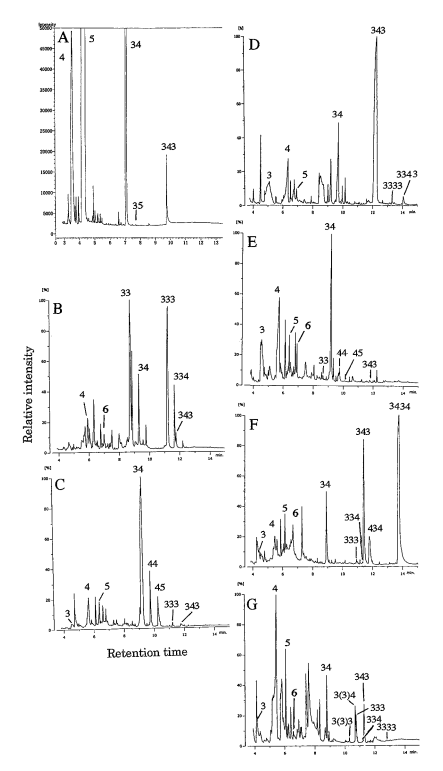


Fig. 1 HPGC analysis of the concentrated polyamine fraction of *Porphyridium purpureum* NIES-2138 "Chinorimo" grown in the presence of 1 mM cadaverine (A), *Bulboplastis apyrenoidosa* NIES-2742 No. 3 (B), *Gelidium crinale* "Makusa" ("Tengusa") (Kumamoto) (C), *Gloiopeltis tenax* "Ma-funori" (D), *Hypnea charoides* "Ibaranori" (E), *Meristotheca papulosa* "Aka-tosakanori" No. 2 (F), and *Mazzaella japonica* "Akaba-ginnansou" No. 1 (G) on SHIMADZU GC-17A (A) or JEOL JMS-700 (B-G). Several non-polyamine peaks assigned by MS spectra were detected in the chromatograms. Abbreviations for polyamines are shown in Table 1.

Polyamines of red algae

Table 1 Concentration of polyamines of red algae

		meen		1011	01	poi	., ui					gae								
	Rhodophyta	Ref.	Dap		Cad					APCad		NSpm	Spm	TSpm	Can				AP4NSpd	
Subphylum Cyanidiophytina Freshwater unicellular thermoacidoph	hilic (micro) red algae		3	4	5	6	33	34	44	35	45	333	343	334	434	3333	3343	3434	3(3)3	3(3)4
Order Cyanidiales (Class Cyanidiophyceae)	NEO 01072	1000		1.00									0.05							
Cyanidium caldarium (RK-1) IAM R-11 (=N (KS-1) (Kusatsu spa, Ja		1990 1990	:	1.28 0.85		:	:	1.24 1.11		-	-	-	0.85 0.32					•	-	
MBIC 10236 (=NBRC		2006		1.30	-	-	2	0.77	-		-	2	0.20	-				-	-	-
(Italy 86) NIES-551			-	0.70	-	-	-	0.84		-	-		0.40	-	-	-	-	-		· -
Cyanidioschyzon melorae NIES-1332 (=NIE	S-3377) "Shianidioshizon"	2006	-	1.57	-	-	-	0.40	•		-	•	0.13	-	-	-	-	-	•	-
Galdieria sulphuraria IAM M-8		1985	-	1.72	-	-	0.08	3.09	-	-	-	0.59	0.82	*	-	-	-	-	-	-
Galdieria partita NBRC 102759	dashutias and Subshulum Eurhodonhutin	a)	-	0.20	-	-	0.12	1.02	-	-	-	0.40	0.42	0.02	-	-	-	-	-	
Subphylum Rhodophytina (Subphylum Proteorho Freshwater/marine unicellular (micro Order Porphyridales (Class Porphyridiophyceae) red algae	aj																		
Porphyridium sordidum MBIC 10454 (=NBR		2006	-	0.55	-	-		0.77			-	-	0.18	-			-			
Porphyridium purpureum (P. cruentum) 1A		1985			0.02	-		0.53	0.04	-	-	-	0.10	*	-	-	-		-	-
	IES-2138 (=IAM R-1)		-	0.52	0.01	-	-	1.06	-	-	-	-	0.52	-	-	-	-	-	-	-
	+ Cad		-	0.40	>2.0	-	-	1.16	-	0.01	-	-	0.31	-		-	-	-	-	-
	M R-3	2006	-	0.60	-	-		0.20	-	-	-	-	0.02	-	•	-	-	-	-	-
	1ES-2140 (=IAM R-3)		-	0.53	-	-	-	1.05	-	-	-	-	0.34	-	-	-	•	•	•	•
Porphyridium aerugineum NIES-1957			-	0.65	-	-	-	0.30	-	-	-	-	-	-	-	-	-	-	-	-
Porphyridium sp. NIES-1032	+ Cad		-	0.20 0.08	1.50	-	-	0.30 0.87	·	-	-	-	- 0.08	-	-	-	-	-	-	-
Marine unicellular (micro) red algae			-	0.08	-	-	·	0.07			•	-	0.00	-	-	-	-	-	-	-
Order Dixoniellales (Class Rhodellophyceae)																				
Dixoniella grisea MBIC 10460 (=NBRC 102	(778)	2006	0.02	0.35		-	0.40	0.04	-	-		-	0.02	-	-	-	-		-	-
Bulhoplastis apyrenoidosa NIES-2742	No.1		0.01	0.40	-	-	0.75	0.80		-	-	1.00	0.01	0.45	-	-	-	-		
	No.2		-	0.20	-	0.02	0.97	0.34	-		-	1.06	0.01	0.35	-	-	-	-	-	-
	No.3		·	0.20	-	0.05	1.06	0.50	-		-	00.1	0.10	0.45	-	-	-	-	-	-
Order Rhodellales (Class Rhodellophyceae)				-																
Rhodella maculata MBIC 10824 (=NBRC 10)2933)	2006		0.02	-	-	0.02	0.65	-	-	-	0.01	0.01	•	-	-	-	-	-	-
Rhodella sp. NIES-1036		2006	-	0.20 0.05	-	-	0.07	0.35 0.30	-	-	-	-	0.19 0.02	-	·	-	-	-	-	-
Corynoplastis japonica NIES-2662 Order Stylonematales (Class Stylonematophyce	199		-	0.05	-	-	0.15	0.50	î	-	-	-	0.02	-	•	-	•	•	-	-
Rhodosorus sp. MBIC 10854 (=NBRC 10294		2006		0.27			0.02	0.15				0.10	0.30		-	-			-	-
Freshwater multicellular (macro) red		2000		0			0.02	0110				0110	0100							
Order Thoreales (Class Florideophyceae)																				
Nemalionopsis tortuosa NIES-1467	"Okichi-mozuku"	2006	-	0.02	-	-	-	0.42	0.30	-	-		0.40	-	-	-	-		-	-
Thorea gaudichaudii NIES-1473	"Shima-chisuzinori"	2006	-	0.02	-	-	-	0.37	0.50	-	-	-	0.22	-	-	•	•	-	-	-
Thorea okadae NIES-1516	"Chisuzinori"	2006	-	0.02	-	-	0.10	0.51	0.45		-	-	0.22	-	-	-	-	•	•	-
Thorea hispida NIES-1572	"Futo-chisuzinori"		•	1.50	-	-	0.05	0.50	0.20	•	•	-	0.02	-	-	-	-	-	-	-
Order Batrachospermales (Class Florideophyce							0.00	0.75	0.00				0.00							
Batrachospermum atrum NIES-1456	" Ishi-kawamozuku"	2006		0.04	-	-	0.20	0.75 0.60	0.32 0.44	-	-		0.02 0.08	-	-	-	-	-	-	-
Batrachospermum helminthosum NIES-145 Batrachospermum turfosum (B. vagum) IAN				0.12		-	-	1.15			-		0.08			-	2		-	
Chantransia macrospora NIES-2636	(I K-+ (=14125-2150) 11030-kawamozak	2000		0.05		-	0.35	0.65	0.03		-	-	0.06	-	-	-	÷.	÷	-	
Order Compsopogonales (Class Compsopogonales	ophyceae)																			
Compsopogon coeruleus NIES-1462	"Ohishisou"	2006		0.03	-	-	-	0.32	0.10		-	-	0.02	-	-	-	-	-	-	-
Compsopongonopsis japonica NIES-1463	"Ohishisou-modoki"	2006	i -	0.02	-	-	-	0.30	0.18	-	-	-	0.05	•	-	-	-	-	-	-
Marine multicellular (macro) red alga Order Bangiales (Class Bangiophyceae)	e (Red-algal seaweeds, Red seawe	eds)																		
Pyropia tenera var. tamatsuensis (Porphyra	tenera) "Ohba-asakusanori"	1982	0.02	0.01	-	-		0.01	-	-	-	-	0.12	*	-	-		-	-	
Pyropia yezoensis form. narawaensis	"Narawa-susabinori" (Ariake, Saga)		0.06	0.01	-	-	-	0.15	-	-	-		0.37	0.01	-	-		-	-	
Pyropia yezoensis	"Susabinori" (Ehime)		0.96	0.20	0.04	-	0.01	0.18	-	-	-	0.01	1.20	-	-	0.01	-	-	-	-
Pyropia tenuipedalis	"Kaigara-amanori" (Yamaguchi)		0.01	-	0.02	0.03	0.04	0.40	0.01	-	-	0.04	1.20	-	-	·	-	-	-	-
Pyropia pseudolinearis	"Uppuruinori" (Shounai, Yamagat	a)	0.05	-	-	-	0.02	0.52	0.01	-	-	0.02	1.25	-	-	-	-	•	-	-
Order Ceramiales (Class Florideophyceae) Campylaephora hypnaeoides	"Egonori" (Kashiwazaki, Niigata)	0.03	1.20	0.49	0.10	10.0	0.65	0.03	-	-	0.02	0.17	0.01	-	0.05	-	-		-
Order Halymeniales (Class Florideophyceae) Polyopes affinis	"Matsunori" (Kumamoto)	-	0.10	0.01	0.01	0.01	0.15	0.02	-	-	0.01	0.12	-		-	-	-	-	-
Order Gelidiałes (Class Florideophyceae) Gelidium elegans	"Makusa" ("Tengusa") (Kumamoto)	0.03	0.20	0.15	-		1.07	0.40		0.21	0.02	0.02	-	-	-	-	-		-
	(Izu, Shizuoka)	0.32	0.50	0.50	0.30	0.01	0.95	0.04	-	-	0.01	0.92	-	-	-	-	-	-	-
Order Gracilariales (Class Florideophyceae)			0.5		0.00		. ···		0			.	o ·•	0.00						
Gracilaria vermiculophylla	"Ogonori" (Korea		0.03				0.12			-	-	0.14	0.40	0.02	-	•	•	-	-	-
Gracilaria arcuata Order Gigartinales (Class Florideophyceae)	"Yumigata-ogonori" (Okinawa	9	0.01	0.04	-	0.20	0.01	1.05	0.03	-	-	0.01	0.95	0.04	-	-	-	-	•	-
• • • • •	"Fukuro-funori" (Erimo, Aomori	``````````````````````````````````````	0.04	0.46	0.33		0.01	0.51			_	0.05	0.80			0.01		-	-	-
Gloiopeltis furcata Gloiopeltis tenax	"Fukuro-funori" (Erimo, Aomori "Ma-funori" (Mie)			0.40		- 2	-	0.54	-	-	-		1.14		-	0.01	0.04	-	-	-
Hypnea charoides	"Ibaranori" (Okinawa				0.30		0.03	1.05	0.08	-	0.01	-	0.02		-	-	-	-	-	-
	-tosakanori" (Kouzu-shima, Tokyo) No.		0.10		0.05			0.82		-	-	0.10	1.22	-	0.08	0.02		0.20	-	-
	No.	2	0.01	0.10	0.08	0.07	-	0.50	-	-	-	10.0	0.90	0.01	0.20	-	-	1.04	-	-
Mazzaella japonica "Akab	a-ginnansou" (Hakodate, Hokkaido) No.		0.10			0.01	-	0.44	-	-	-	0.20	0.25	0.01	·	-	-	-	0.03	0.18
	No.		0.10					0.47	-	-	-	0.03	0.50	0.01	•	0.01	-	-	-	-
Chondrus elatus	"Kotoji-tsunomata" (Chiba		0.05				-	0.17	0.02	-	-	-	0.12	-	-	-	-	-	-	-
Chondrus oceliatus Caulacanthus ustulatus	"Tsunomata" (Chibi "Icordentou" (Chibi			1.05		0.01	-	0.07	-	•	-	0.01	0.03	-	-	-	-	-	•	-
Caulacanthus ustulatus Order Palmariales (Class Florideophyceae)	"Isodantsu" (Chib	1)	0.10	0.05	0.05	•	-	0.30	•	-	-	0.25	0.30	-	-	-	-	-	•	-
	"Dulos" (It-tl II-bb-14)	۰ ۱	0.02	0.02				1.25			-	-	0.57	_			-	-	-	-
Palmaria palmata	"Dulse" (Hakodate, Hokkaido	"	0.02	0.02			,				-	-	10.01	-						

Dap (3) (abbreviations for the numbers of methylene chain units between N). 1.3-diaminopropane; Put (4), putrescine: Cad (5), cadaverine; Dath (6), 1.6-diaminohexane; NSpd (33), norspermidine; Spd (34), spermine; Can (35), aninopropylcadaverine; ABCad (45), aminobutylcadaverine; NSpm (33), norspermine; Spm (34), spermine; TSpm (34), thermospermidine; Can (43), canavalanine (aminobutylspermidine; IAPCan (333), orladopentamine; TPen (3343), thermospermidine; APCan (3434), aminopropylcadaverine; NSpm (33), norspermidine; IAM, IAM Culture Collection, Institute of Molecular and Cellular Biosciences, The University of Tokyo, Tokyo, Japan (IAM algal collections have been transferred to NBES); MBIC, Marine Biotechnology Institute for Lavironmental Studies, Tsukuba, Japan; NBIC algal collections have been transferred to NBES); MBIC, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC algal collections have been transferred to NBES); MBIC, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, algan; NBEC); MISS, National Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, MBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; NBIC, MISS, MISS, Mational Institute for Environmental Studies, Tsukuba, Japan; -, not detected (<0.005 µmol/g wet weight). ★ . Thermospermine was not analyzed by GC. Ref. are shown in the References (published year) of the present report.

of cation-exchange resin, Hitachi 2619F (=Hitachi 2720) (4 mm I.D. ×50 mm), using post-labeled fluorometry after heating with o-phthalaldehyde (Hamana et al., 2016a, 2016b). After the heptafluorobutyrization (HFB) of concentrated polyamine samples, HPGC on a SHIMADZU GC-17A and HPGCmass spectrometry (HPGC-MS) on a JEOL JMS-700, equipped with a capillary column of Inert Cap 1MS (0.32 mm I.D. \times 30 m, df 0.25 μ m) (GL Sciences) were performed at the column temperatures of $90^{\circ}C$ -($16^{\circ}C$ / min)-280°C and 120°C -(16°C / min)-280°C, respectively (Hamana et al., 2016a, 2016b; Niitsu et al., 2014, as shown in Fig. 1). Mass spectra of the HFB derivatives of all the polyamines detected in HPGC-MS were obtained for their identification (Hamana et al., 2018). Molar concentrations of cellular polyamines per gram of wet weight of the starting algae were estimated from the HPLC analysis, and two HPGC analyses were performed using authentic polyamine standards and are shown in Table 1. Polyamine extraction from seaweeds was quantitatively difficult, so the concentration values were unstable. Previous polyamine data for 21 red algae analyzed with standard GC with a packed column in 1982-2006 are cited in Table 1.

RESULTS AND DISCUSSION

Polyamines of thermoacidophilic unicellular red algae

Thermoacidophilic unicellular red algae (order Cyanidiales) constitute three genera Cyanidium, Cyanidioschyzon and Galdieria and are unique with a green colored red alga, single or few mitochondria and chloroplasts, and optimum growth at 40-50°C and pH 3-4 (Inouye, 2007; Miyagishima et al., 2017). Polyamines of a new strain of Cyanidium caldarium "Ideyukogome" and a new species of Galdieria, G. partita, were analyzed in the present study. Four strains of Cyanidium caldarium and a strain of Cyanidioschyzon melorae "Shianidioshizon" always contained putrescine (4), spermidine (34) and spermine (343) (Table 1) (Hamana & Niitsu, 2006; Hamana et al., 1990). G. partita as well as G. sulphuraria contained norspermidine (33) and norspermine (333) in addition to putrescine, spermidine and spermine (Table 1) (Hamana & Matsuzaki, 1985; Hamana et al., 1990). A small amount of thermospermine (334) in addition to spermine was possible in the Galdieria species containing norspermine as a major polyamine. Galdieria belongs to a lineage far from that of *Cyanidium* and *Cyanidioschyzon* in the order Cyanidiales (Yoon *et al.*, 2006). The higher levels of spermine, a tetra-amine, in the unicellular thermoacidophiles grown at 40–50°C compared with other unicellular red algae grown at 20–25°C seems to be related to the high optimum growth temperature of the thermoacidophiles (Hamana *et al.*, 1990). The acidophilic property of the thermoacidophiles appears to be unaffected in the cellular polyamine profiles.

On the other hand, spermine has not been found in unicellular glaucophyte algae (phylum Glaucophyta) in which putrescine and spermidine were found as the polyamine components (Hamana & Niitsu, 2006), and in the primitive unicellular green algae belonging to the class Prasinophyceae of the phylum Chlorophyta in which norspermidine and homospermidine (44) in addition to putrescine and spermidine were the major polyamines (Hamana *et al.*, 2013, 2018). Three different polyamine synthetic abilities were distributed independently in the early evolution of the three primitive unicellular algal lineages (phyla) after the primary endosymbiosis.

Polyamines of the freshwater/marine unicellular red alga *Porphyridium*

Salt tolerant P. sordidum, P. purpureum and Porphyridium sp. NIES-1032 contained putrescine (4), spermidine (34) and spermine (343). The freshwater P. aerugineum analyzed in the present study contained putrescine and spermidine but lacked in spermine. The spermine level varied within the salt tolerant Porphyridium species, however, the relation between the level and their salt tolerance is unclear. When P. purpureum NIES-2138 and P. aerugineum NIES-1957 were cultured in the presence of 1 mM cadaverine in the medium, a small amount of aminopropylcadaverine (35) was detected in the former but not in the latter, indicating that there was aminopropylation to the supplied cadaverine (5). However, this was at very low levels in the red algae. Aminobutylcadaverine (45) was not produced from the supplemented cadaverine.

In the order Porphyridiales containing a single chloroplast, although *Porphyridium purpureum* (formerly *P. cruentum*) "Chinorimo" NIES-2138 and NIES-2140 were isolated from a terrestrial hot spring in Kagoshima, Japan and land soil in the Tohoku region, Japan, respectively, they showed salt tolerance and good growth in a marine algal medium (NIES-ESM medium) that was also used for marine *P. sordidum* MBIC 10454 and *Porphyridium* sp. NIES-1032 isolated in the Pacific Ocean, Okinawa, Japan. *Porphyridium aerugineum* NIES-1957 was isolated from a land soil in Nepal and is a blue-green colored red alga likely related to the thermoacidophilic red algae belonging to the order Cyanidiales (Gantt *et al.*, 1968), and it thrives in a freshwater algal medium (NIES-C medium). These findings suggest that the marine species are more primitive phylogenetically than the freshwater species within the genus *Porphyridium*.

Polyamines of marine unicellular red algae

Marine unicellular red algae are located in the orders Dixoniellales (class Rhodellophyceae), Rhodellales (class Rhodellophyceae) and Stylonematales (class Stylonematophyceae). Norspermidine (33) (and norspermine (333)) was detected in six marine species belonging to the three orders, and their polyamine profiles were distinguished from those of *Porphyridium* species (order Prophyridiales of class Porphyridiophyceae). Newly analyzed *Corynoplastis japonica* (Yokoyama *et al.*, 2009) contained norspermidine as a major polyamine. Although few analytical data were obtained, the same distribution profile of polyamines was suggested within the classes Rhodellophyceae and Stylonematophyceae.

Brownish Bulboplastis apyrenoidosa NIES-2742 containing a single chloroplast was isolated from the mangroves of Iriomote Isl., Okinawa, Japan (Kushibiki et al., 2012) and contained a high level of norspermidine, spermidine (34), norspermine and thermospermine (334), and a low levels of spermine (343), indicating its preferential aminopropylation to the aminopropyl moieties of norspermidine and spermidine to produce norspermine and thermospermine, respectively, in the red alga. The same polyamine profile was obtained in acid extracts from the three different cultures No. 1, 2 and 3 (Table 1). This polyamine profile containing of thermospermine as a major polyamine is unique among all previously analyzed algae. 1,6-Diaminohexane was detected as a minor polyamine in two of the three Bulboplastis samples.

Polyamines of freshwater macro red algae

The freshwater multicellular red algae "Chisuzunori", "Kawamozuku" and "Ohishisou" that grow in rivers or lakes in Japan are known to be endangered. The 10 species of this group, belonging to two orders of the class Florideophyceae and an order of the class Compsopogonophyceae (Table 1), were analyzed in both previous studies and present work. They ubiquitously contained homospermidine (44) in addition to putrescine (4), spermidine (34) and spermine (343). Norspermidine (33) was found in four of the 10 species, whereas norspermine (333) was not found in any of the species analyzed. The ubiquitous occurrence of homospermidine in multicellular red algae suggests that homospermidine synthetic abilities have been coupled with multicellular evolution within the freshwater macro red algae.

Unusual polyamines of red seaweeds

Although the macro red-algal seaweeds purchased from food markets were washed in artificial seawater by sonication, the seaweeds were contaminated by other marine microorganisms. However, the unusual polyamines found in the seaweeds have never been detected in various marine bacteria. 1.6-Diaminohexane (6) was widely distributed within the red seaweeds. Although its biosynthetic pathway is unknown, this diamine is also widely distributed in brown seaweeds (Hamana et al., 2017) but not in green seaweeds (Hamana et al., 2018). On the other hand, thermospermine (334) is found in many freshwater multicellular green algae, green seaweeds and brown seaweeds as a major polyamine, and is rich in the thallic species (Hamana et al., 2017, 2018). Thermospermine was detected sporadically in the red seaweeds as a minor polyamine.

Since Gelidium elegans "Makusa" ("Tengusa") is rich in agar, acid extraction of the whole polyamine fraction from the alga and concentration of polyamines on ion-exchange resin were difficult to achieve, so the polyamines were analyzed in only one sample of four polyamine extractions from the Kumamoto sample. Polyamine analysis of the Shizuoka sample was performed for only one sample among three polyamine extractions. Aminobutylcadaverine (45) was found in the Kumamoto sample in which homospermidine (44) was at a high level, but not detected in the Shizuoka sample in which homospermidine was at a low level. Hypnea charoides "Ibaranori" contained aminobutylcadaverine (45) as a minor polyamine. Campylaephora hypnaeoides "Egonori", Gloiopeltis furcate "Fukuro-funori" and Mazzaella japonica "Akabaginnansou" were rich in cadaverine (5) but poor in homospermidine (44), however, aminobutylcadaverine was not detected in them. Aminobutylcadaverine has been found in a unicellular green alga, *Westella* sp. DBT, however, the alga was rich in cadaverine and homospermidine (Hamana *et al.*, 2018).

Two tertiary branched triamines N^{4} aminopropylnorspermidine (3(3)3) and N^{4} aminopropylspermidine (3(3)4) were detected in *Mazaella japonica* (No. 1) as a minor polyamine. Although the two branched triamines were not detected in the red-algal sample No. 2 and its detection was not replicable, the triamines have also been found in a green alga, *Chlorogonium complex* (Hamana *et al.*, 2018).

Canavalmine (434) has been detected in some green algae as a minor polyamine (Hamana *et al.*, 2013, 2018). Both aminopropylcanavalmine (3434) and canavalmine were found as major polyamines in two samples of *Meristotheca papulosa* "Akatosakanori". Although canavalmine and the aminopropylated canavalmine, aminopropylcanavalmine, have been found in some leguminous seeds (Hamana *et al.*, 1992: Matsuzaki *et al.*, 1990) and sea cucumber (Hamana *et al.*, 1991), this is the first known report of a major occurrence of canavalmine and aminopropylcanavalmine in algae.

Caldopentamine (3333) and thermopentamine (3343), probably produced from norspermine and spermine, respectively, were detected in some red seaweeds, but homocaldopentamine (3334), which is produced from thermospermine, was not detected in the red seaweeds. The three penta-amines caldopentamine, thermopentamine and homocaldopentamine have been found in green and brown seaweeds containing norspermine, spermine, thermospermine and canavalmine, whereas aminopropyl-canavalmine has not been detected (Hamana *et al.*, 2017, 2018).

Brown seaweeds belonging to the class Phaeophyceae of the phylum Heterokontophyta, evolved after the secondary symbiosis of a red alga within stramenopiles, and are unique among heterokontophytes in developing into multicellular forms with differentiated tissues. The distribution of the long linear diamine, 1,6-diaminohexane, penta-amines and hexa-amines has been observed in 12 typical brown seaweed species (Hamana *et al.*, 2017). On the other hand, the macro green algae belonging to the classes Ulvophyceae and Charophyceae were not present in 1,6-diaminohexane and hexa-amines, which contained small amounts of penta-amine (Hamana *et al.*, 2018). As shown in the present study, 1,6-diaminohexane was found in the most of red seaweeds except for *Meristotheca papulosa* "Aka-tosakanori", but no significant amounts of penta-amine and hexa-amine were detected. The polyamine profiles of red seaweeds resemble to those of green seaweeds more closely than those of brown seaweeds.

ACKNOWLEDGEMENTS

We thank NIES and NBRC for supplying the red algae and Ms. Fumi Mori of NIES for advice on unicellular red-algal cultures.

REFERENCES

- Fuell, C., Elliott, K.A., Hanfrey, C.C., Franceschetti, M. & Michael, A.J. 2010. Polyamine biosynthetic diversity in plants and algae. Plant. Physiol. Biochem. 48: 513-520.
- Gantt, E., Edwards, M.R. & Conti, S.F. 1968. Ultrastructure of *Porphyridium aerugineum* a blue-green colored rhodophytan. J. Phycol. 4: 65-71.
- Hamana, K. 2008. Cellular polyamines of phototrophs and heterotrophs belonging to the lower eukaryotic phyla Cercozoa, Euglenozoa, Heterokonta and Metamonada. J. Gen. Appl. Microbiol. 54: 135-140.
- Hamana, K. & Matsuzaki, S. 1982. Widespread occurrence of norspermidine and norspermine in eukaryotic algae. J. Biochem. 91: 1321–1328.
- Hamana, K. & Matsuzaki, S. 1985. Further study on polyamines in primitive unicellular eukayotic algae. J. Biochem. 97: 1311-1315.
- Hamana, K. & Niitsu, M. 2006. Cellular polyamines of lower eukaryotes belonging to the phyla *Glaucophyta, Rhodophyta, Cryptophyta, Haptophyta* and *Percolozoa*. J. Gen. Appl. Microbiol. **52**: 235-240.
- Hamana, K., Matsuzaki, S., Nitsu, M., Samejima, K. & Nagashima, H. 1990. Polyamines of unicellular thermoacidophilic red alga *Cyanidium caldarium*. Phytochemistry **29**: 377-380.
- Hamana, K., Niitsu, M., Samejima, K. & Matsuzaki, S. 1991. Novel tetra-amines, penta-amines and hexaamines in sea urchin, sea cucumber, sea squirt and bivalves. Comp. Biochem. Physiol. 100B: 59-62.
- Hamana, K., Matsuzaki, S., Niitsu, M. & Samejima, K. 1992. Distribution of unusual polyamines in legumes seeds. Can. J. Bot. **70**: 1984–1990.

- Hamana, K., Aizaki, T., Arai, E., Saito, A., Uchikata, K. & Ohnishi, H. 2004a. Distribution of norspermidine as a cellular polyamine within micro green algae including non-photosynthetic achlorophyllus *Polytoma, Polytomella, Prototheca* and *Helicosporidium*. J. Gen. Appl. Microbiol. 50: 289-295
- Hamana, K., Sakamoto, A., Nishina, M. & Niitsu M. 2004b. Cellular polyamine profile of the phyla Dinophyta, Apicomplexa, Ciliophora, Euglenozoa, Cercozoa and Heterokonta. J. Gen. Appl. Microbiol. 50: 297-303.
- Hamana, K., Niitsu, M. & Hayashi, H. 2013. Occurrence of homospermidine and thermospermine as a cellular polyamine in unicellular chlorophyte and multicellular charophyte green algae. J. Gen. Appl. Microbiol. 59: 313–319.
- Hamana, K., Furuchi, T., Hayashi, H. & Niitsu, M. 2016a Additional analysis of cyanobacterial polyamines —Distribution of spermidine, homospermidine, spermine, and thermospermine within the phylum *Cyanobacteria*—. Microb. Resour. Syst. **32**: 179-186.
- Hamana, K., Furuchi, T., Nakamura, T., Hayashi, H. & Niitsu, M. 2016b. Occurrence of penta-amines, hexa-amines and methylated polyamines in unicellular micro-algae belonging to the phyla *Heterokontophyta* and *Labyrinthulomycota* of the subdomain *Stramenopiles*. J. Gen. Appl. Microbiol. **62**: 320-325.
- Hamana, K., Kobayashi, M., Furuchi, T., Hayashi, H. & Niitsu, M. 2017. Polyamine analysis of brownalgal seaweeds (class Phaeophyceae) from food markets —Distribution of diaminohexane, pentaamines, and hexa-amine—. Microb. Resour. Syst. 33: 3-8.
- Hamana, K., Kobayashi, M., Furuchi, T., Hayashi, H. & Niitsu, M. 2018. Polyamine analysis of unicellular, colonial, and multicellular green algae —Detection of aminobutylcadaverine, N¹-aminopentylspermidine, N⁸-aminopentylspermidine, and penta-amines—. Microb. Resour. Syst. **34**: 73-82.
- Inouye, I. 2007. The Natural History of Algae: Perspective of Three Billion Years Evolution of Algae, Earth and Environment, second edition. Tokai University Press, Hatano, Kanagawa, Japan (in Japanese).
- Kushibiki, A., Yokoyama, A., Iwataki, M., Yokoyama, J.,

West, J.A. & Hara, Y. 2012. New unicellular red alga, *Bulboplastis apyrenoidosa* gen. et sp. nov. (Rhodellophyceae, Rhodophyta) from the mangroves of Japan: Phylogenetic and ultrastructural observations. Phycol. Res. **60**: 114-122.

- Matsuzaki, S., Hamana, K., Okada, M., Niitsu, M. & Samejima, K. 1990. Aliphatic penta-amines found in *Canavalia gladiate*. Phytochemistry **29**: 1311-1312.
- Minguet, E.G., Vera-Sirera, F., Marina, A., Carbonell, J. & Blazquez, M.A. 2008. Evolutionary diversification in polyamine biosynthesis. Mol. Biol. Evol. 25: 2119– 2128.
- Miyagishima, S., Wei, J.L., Nozaki, H. & Hirooka, S. 2017. Cyanidiales: Evolution and habitats, *In* Kuroiwa, T., Miyagishima, S., Matsunaga, S., Sato, N., Nozaki, H., Tanaka, K. & Misumi, O. (eds.), *Cyanidioschyzon merolae*, p. 3–15, Springer, Singapore.
- Muñoz-Gómez, S.A., Mejia-Franco, F.G., Durnin, K., Colp, M., Grisdale, C.J., Archibald, J.M. & Slamovits, C.H. 2017. The new red algal subphylum Proteorhodophytina comprises the largest and most divergent plastid genomes known. Curr. Biol. 27: 1677-1684.
- Niitsu, M., Takao, K., Kato, Y., Nagase, E., Furuchi, T. & Hamana, K. 2014. Occurrence of N¹methylputrescine, N¹, N⁴-dimethylputrescine, N¹-methylhomospermidine and N¹, N⁹dimethylhomospermidine in tomato root. Int. J. Plant Biol. Res. 2(4): 1022.
- Schweikert, K. & Burritt, D.J. 2015. Polyamines in macroalgae: advances and future perspectives. J. Phycol. 51: 838-849.
- Shaw, F.L., Elliot, K.A., Kinch, L.N., Fuell, C., Phillips, M.A. & Michael, A.J. 2010. Evolution and multifarious horizontal transfer of an alternative biosynthetic pathway for the alternative polyamine sym-homospermidine. J. Biol. Chem. 285: 14711-14723.
- Yokoyama, A., Scott, J.L., Zuccarello, G.C., Kajikawa, M., Hara, Y. & West, J.A. 2009. Corynoplastis japonica gen. et sp. nov. and Dixoniellales ord. nov. (Rhodellophyceae, Rhodophyta) based on morphological and moleculer evidence. Phycol. Res. 57: 276-289.
- Yoon, H.S., Müller, K.M., Sheath, R.G., Ott, F.D. & Bhattacharya, D. 2006. Defining the major lineages of red algae (Rhodophyta). J. Phycol. 42: 482-492.

浜名康栄1), 小林正樹2), 古地壯光2), 林 秀謙1), 新津 勝2)

1)前橋工科大学工学部, 2)城西大学薬学部

細胞内ボリアミン構成と多細胞紅藻類の系統進化との関連性を調べるため、大型多細胞の海藻 18 種(19 株)を含む 27 種 (29 株)の紅藻類から酸抽出したボリアミンを HPLC と HPGC-MS にて追加分析し, 既分析の紅藻 21 種のポリアミン構成と 比較した.好熱好酸性単細胞の Cyanidiales 目では、*Cyanidium と Cyanidioschyzon* がプトレスシン、スペルミジン、スペル ミンを含有し、*Galdieria*は、これらに加えて、ノルスペルミジンとノルスペルミンを有していた.単細胞の淡水産および海 *産 Poryphyridium*では、プトレスシンとスペルミジンで、種によりスペルミンを含有していた. 海産単細胞性の Dixoniellales 目、Rhodellales 目、Stylonematales 目では、プトレスシン、スペルミジン、スペルミンに、ノルスペルミジンとノルスペル ミンが加わった. *Bulboplastis*はこの5ポリアミンの内のスペルミンに代わってサーモスペルミンを主要ポリアミンの一つと して含む特徴があった. 淡水産多細胞 Thoreales 目、Batrachospermales 目、Compsopogonales 目では、プトレスシン、ス ペルミジン、スペルミンにホモスペルミジンが追加される種が優勢であった. Bangiophyceae 綱と Florideophyceae 綱の7 目に属する海藻では、1.3-ジアミノプロパン、プトレスシン、カダベリン、スペルミジン、スペルミン、ノルスペルミジン、 ノルスペルミン、ホモスペルミジン、サーモスペルミンが広く分布していた. 1.6-ジアミノへキサンが海藻 12 種に検出され、 アミノブチルカダベリンが *Gelidium と Hypnea*に、カナバルミンとアミノプロピルカナバルミンが *Meristotheca* に検出され た. これら紅藻のポリアミン構成は系統進化的には異なる緑藻(アオサ藻)や褐藻のポリアミン構成とは異なっていた.