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Extremely High Methane Concentration in Bottom Water and Cored Sediments from Offshore Southwestern Taiwan

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ABSTRACT

It has been found that Bottom Simulating Reflections (BSRs), which infer the existence of potential gas hydrates underneath seafloor sediments, are widely distributed in offshore southwestern Taiwan. Fluids and gases derived from dissociation of gas hydrates, which are typically methane enriched, affect the composition of seawater and sediments near venting areas. Hence, methane concentration of seawater and sediments become useful proxies for exploration of potential gas hydrates in a given area. We systematically collected bottom waters and sedimentary core samples for dissolved and pore-space gas analyses through five cruises: ORI-697, ORI-718, ORII-1207, ORII-1230, and ORI-732 from 2003 to 2005 in this study. Some sites with extremely high methane concentrations have been found in offshore southwestern Taiwan, e.g., sites G23 of ORI-697, N8 of ORI-718, and G96 of ORI-732. The methane concentrations of cored sediments display an increasing trend with depth. Furthermore, the down-core profiles of methane and sulfate reveal very shallow depths of sulfate methane interface (SMI) at some sites in this study. It implies sulfate reduction being mainly driven by the process of anaerobic methane oxidation (AMO) in sediments; thus indicating that there is a methane-enriched venting source, which may be the product of dissociation of gas hydrates in this area.

(Key words: Methane, Gas hydrate, Sulfate methane interface, Anaerobic methane oxidation)

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1. INTRODUCTION

Gas hydrates, which are a kind of non-stoichiometric clathrate, are natural occurring solids composed of water molecules forming a rigid lattice of cages with most of the cages each containing a molecule of natural gas, mainly methane. These are metastable crystal products at low temperature and high-pressure. If the temperature and pressure conditions are above the gas hydrate stability boundary, then the protogenic gas hydrates dissociate into water and methane. Gas hydrates are widely distributed in the permafrost of polar regions and the strata of continental deep-sea areas (Kvenvolden 1998; Pellenbarg and Max 2000; Sloan 1998; Milkov 2004) and are considered future potential energy resources.

There are conditions under which gas hydrates beneath the ocean floor may become unstable: firstly, if there is a general change in temperature and pressure conditions at the seafloor; and secondly, if warm fluids begin moving through highly permeable faults (Pecher 2002). When dissociation occurs, the frozen sediments become liquefied releasing water and dissociated gases and thereby increasing pore pressure. These high-pressure fluids from gas hydrate dissociation permeate upward through the seafloor. Here, there is a strong potential for faults and porous layers to provide the main migration pathways for fluids in gas hydrate bearing layers (Pecher 2002; Wood et al. 2002).

The fluids dissociated from gas hydrates may erupt through craters forming mud volcanoes. This may explain the presence of submarine mud volcanoes in offshore southwestern Taiwan (e.g., Chou et al. 2000; Chiu et al. 2006), which have been considered to be related to on-land mud volcanoes (Yang et al. 2003, 2004) by Yeh (2003). Moreover, these methane-rich fluids influence the composition of pore water and bottom seawater around these gas-expelling craters.

Local scientists have done many geophysical investigations on gas hydrates in offshore southwestern Taiwan. Based on seismic reflection profiles, they found many Bottom Simulating Reflections (BSRs) and concluded that gas hydrates are widely distributed under the seaf-loor of depths of 700 to 3000 m in this region (Chi et al. 1998; Chow et al. 2000; Schnürle et al. 1999, 2002, 2006; Liu et al. 2006). The study presents the first systematic survey based on gas geochemical techniques performed in the gas hydrate area in offshore SW Taiwan. We collected bottom waters and cored sediments for fluid and gas geochemistry study during five cruises (ORI-697; ORI-718; ORI-732; ORII-1207; ORII-1230) from 2003 to 2005. The sampling sites are shown in Fig. 1.

In comparison with gas hydrate studies in other areas (Kvenvolden and Lorenson 2000; Lorenson and Collett 2000), our results show unusually high methane concentration in bottom water and sediment pore space. This phenomenon can be explained by venting methane, which could be due to the release of methane from gas hydrate dissociation migrating upward.

2. ANALYTICAL METHODS AND PROCEDURES

Bottom water and sediment samples were collected for analyzing dissolved methane concentration in seawater and pore spaces. Sediment samples were collected by both gravity (30 - 100 cm of length) and piston (300 - 500 cm of length) core samplers. Bottom seawater samples are

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southwestern Taiwan showing sampling sites investigated in this study. The region of Fig.1c is shown in Fig. 1. (a) Bathymetric map of offshore Taiwan. Box shows region of Fig.1b. (b) Bathymetric map of offshore the box. (c) Enlarged map shows the sampling sites for the detailed survey, which piston cores were taken in this study. The depth of the contours is shown in meter.

represented by core top waters. The core top waters were collected in plastic bottles and then transferred immediately to low permeability glass bottles. After being given an ultrasonic bath at 60°C for 10 minutes, the dissolved gas of the bottom water was transferred to a 7-ml Vacutainer using a gas tight syringe. The extracted gases were analyzed using gas chromatography.

Sediments were placed in a tin can of volume ca. 930 cm^3 immediately after the cores were retrieved on board. Distilled water was added into the can and a fixed space remained as the headspace. HgCl₂ was added as a bacterial inhibitor. To avoid water and gas escaping, the cover was sealed by silicone gel. After the silicone dried, the can was kept upside down and ready for analyses.

In order to release all gases into the headspace from the pore space of the sediments, the tin can was first shaken after being brought back to a shore-based laboratory and then kept at room temperature for 30 minutes. A gas-tight syringe's needle, connected to a Vacutainer, was allowed to pierce the tinplate can using a septa. Gases from the headspace were thus transferred to the Vacutainer. These extracted gases were then injected into a gas chromatograph for analysis.

Gas was introduced into a gas chromatograph (GC) (SRI 8610C) equipped with both thermal conductivity (TCD) and flame ionization (FID) detectors. Different components of the gas were separated in GC (injection temperature 30°C, held isothermal for 2 minutes, ramped to 250°C at 120°C min⁻¹). The system utilized two carrier gases, hydrogen and argon. Hydrogen was supplied by a hydrogen generator for use in the FID and in one TCD; whilst the other TCD utilized argon supplied by a cylinder tank. In general, most gas components analyzed in this system have analytical errors of less than 5% with low detection and quantification limits (Lee et al. 2005). Results of dissolved methane concentrations are reported relative to the volume of seawater from which the gases were extracted (i.e., nanoliters of gas per liter of seawater [nL L⁻¹]) and methane concentrations of cored sediments are reported relative to wet sediment from which the gases were extracted (i.e., microliters of gas per liter of wet sediment [μ L L⁻¹]).

3. RESULTS

3.1 Dissolved Methane Concentrations in Bottom Water

Results obtained from core top waters of 114 sites are listed in Table 1. Most of the samples contain methane as the dominant hydrocarbon gas. Most C_{2+} gases are below detection limits, so only the results of methane concentration are given here. The results can be arbitrarily divided into five groups: (a) Very low methane concentration (below detection limits); (b) Low methane concentration (<100 nL L⁻¹); (c) High methane concentration (100 ~ 1000 nL L⁻¹); (d) Very high methane concentration (1000 ~ 10000 nL L⁻¹); (e) Especial high methane concentration (1000 ~ 100000 nL L⁻¹); (f) Extremely high methane concentration (<1000 nL L⁻¹); The concentration distribution of the dissolved methane in bottom water in this study is compiled in Fig. 2. It shows that most sites exhibit much higher dissolved methane concentration than those in normal seawater, which is usually below 50 nL L⁻¹ (Millero 1996).

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ORI-697	CH_4	ORI-718	CH_4	ORII-1207	CH ₄ (nL/L	ORII-1230		ORI-732	CH_4
Cruise	(nL/L)	Cruise	(nL/L)	Cruise		Cruise	CH ₄ (nL/L)	Cruise	(nL/L)
G1	282	G4	3,740	G-15	384	А	19,732	G1	5,365
G5A	212	G9	4,970	G-16	441	D	4,897	G2	3,473
G6	148	G11	3,638	G-18	792	G3	21,239	G3	45,797
G10	156	G11C	822	G-20	516	G6	27.9	G4	5,494
G15	156	G12	557	G-22	876	G15	43,318	G5	81,490
G17	1,221	G14	10,838	G-24	723	G19	7,221	G7	8,667
G19	390	G16	762	G-27	749	G21	53,888	G8	6,230
G21	613	G18	2,368	G-28	809	G23	2,912	G9	64,117
G23	198,616	G22	78,731	G-29	326	G67	1,247	G10	13,364
		G24	935	G-30	458	G69	30,196	G17	53,796
		N1	6,204	G-32	705	G70	17,510	G18	21,472
		N2	9,750	G-44	308	G73	11,035	G26	140
		N4	3,793	G-45	416	G81	41,298	G27	44,743
		N5	4,543	G-47	329	G82	65,153	G28	13,194
		N6	2,450	G-48	477	G84	4,120	G31	4,186
		N8	613,268	G-50	597	G85	135	G32	274
		N9	12,702			G86	2,666	G33	8,751
		N11	10,544			G87	20,633	G36	141
						G89	58,259	G37	20,718
						G90	99.6	G96	536,32
						G91	44,383		
						G93	8,392		
						G94	11,120		
						G95	36,844		
						G96	47,551		
						G99	18,732		
						G100	25,566		
						G103	2,201		
						G105	988		
						G107	7,453		
						G108	38,559		
						G109	475		
						G110	23,921		

 Table 1. Dissolved methane concentrations measured in Offshore Southwestern

 Taiwan bottom seawaters.



Fig. 2. Map of methane concentrations (in nL L⁻¹) in bottom seawaters distributed in offshore southern Taiwan.

3.2 Methane and Ethane Concentrations of Cored Sediments

Methane is the main component of hydrocarbon gases, ethane is the minor component and C_{3+} gases usually are below detection limits. The results are consistent with recent work of Oung et al. (2006) in this area. Table 2 shows both methane and ethane concentrations of all

cored samples of each cruise. Note that the reported concentrations are considered to be the minimum values of the samples due to possible gas loss during core recovery, cutting, sample storage and gas transfer before analysis.

Except for a few sites, e.g., sites GD of ORI-697 and G24 of ORI-718 where the methane concentrations are very low (less than 10 μ L L⁻¹), the results show that most of the methane concentrations in the pore spaces range from ~10 to ~10000 μ L L⁻¹.

In general, most of the cored samples show less methane concentrations in the shallower depths, and methane concentration increasing with increasing depth (Fig. 3; see Table 2 for details). Extremely high methane concentrations can be found at some sites, especially sites G23 of ORI-697, G22, N8 of ORI-718, and G96 of ORI-732 cruises. In contrast, methane concentrations at some sites do not change much with depth and maintain similar values $(10^2 \sim 10^3 \ \mu L \ L^{-1})$ (Fig. 3), e.g., sites: G1, G3, G6, GA of ORI-697, and N4, N13, G4, G14 of ORI-718. It is worth noting that methane profiles at some sites show a saw-tooth shape, e.g., sites G19 of ORI-697 and N11, N9 of ORI-718, which indicate possible gas leaks at some depths or multiple gas sources underneath some sites.

Note that the gravity core samples, of three cruises, ORII-1207, ORII-1230 (except G96) and ORI-732 (except G96), show low methane concentrations ($< 10^3 \ \mu L L^{-1}$) at all sites. The length of cored sediments from gravity core samplers usually is less than 2 m, so the change of methane concentrations with depth is unknown for those sites.

4. DISCUSSIONS

4.1 Very Shallow Depth of Sulfate Methane Interface

Sulfate is involved in two chemical reactions induced by microbial activity that deplete sulfate in pore waters of the sulfate reduction zone within anoxic marine sediments. In the first process, sulfate, which is a kind of oxidant, reacts with sedimentary organic matter. The net reaction of the first process is shown as equation (1):

$$2(CH_2O) + SO_4^{2-} \rightarrow 2HCO_3^{-} + H_2S \quad . \tag{1}$$

In the second process, sulfate reacts with methane called anaerobic methane oxidation (AMO). The net reaction of the second process can be shown as equation (2):

$$CH_4 + SO_4^{2-} \rightarrow HCO_3^{-} + HS^{-} + H_2O$$
 (2)

By combining the profiles of interstitial sulfate concentration and the methane concentration of pore spaces in sediments from a gas hydrate area, it is typical to observe sulfate depleting and methane increasing dramatically at a particular depth; this is called the Sulfate Methane Interface (SMI). If methane fluxes from sediments are high, sulfate reduction rates will be fast. Furthermore, fast sulfate reduction rates induce shallower depth of the SMI. Therefore, the profiles of interstitial sulfate concentration and the methane concentration of pore space in sediments can provide an indicator for further estimation of methane fluxes from buried sediments if there are gas hydrates beneath the sea floor (Borowski et al. 1996, 1999).

ORI-697 cruise	CH ₄ (μL/L)	C_2H_6 (µL/L)		ORI-718 cruise	CH_4 (uL/L)	C_2H_6 (μ L/L)		ORI-718 cruise	CH ₄ (μL/L)	C_2H_6 (µL/L)	
G1 20-30	540	- (µD/D)	Р	G4 31-40	375	8.79	Р	N8 31-40	117	21.6	Р
G1 120-130	492	_		G4 81-90	745	10.8		N8 81-90	1301	8.53	
G1 220-230	663	_		G7 31-40	595	10.6	Р	N8 131-140	94,602	34.7	
		-	Р		867	8.96	P		· · · ·	40.5	
G2 20-30	392	-	Р	G9 31-40			Г	N8 181-190	· ·		г
G2 120-130	4,964	-	D	G9 81-90	805	11.4		N9 31-40	8.75	2.76	F
G3 70-80	442	-	Р	G9 131-140	375	10.8		N9 81-90	215	20.4	
G3 170-180	294 423	-		G9 181-190	395 363	10.8 8.79		N9 131-140	37.7 4.01	14.3 2.26	
G3 270-280 G3 370-380	423 759	-		G9 231-240 G9 281-290				N9 181-190		65.1	
		25.4	р		3,668	11.7	D	N9 231-240	6,457		
G5A 70-80	166	2.05	Р	G11 31-40	384	8.71	Р	N9 281-290	10.5	0.09	
G5A 120-130		-		G11 79-88	1,138	30.6	D	N11 31-40	7.01	5.85	F
G5A 170-180		-		G11c 31-40	392	8.91	P	N11 81-90	22.1	19.3	
G5A 220-230		4.08		G14 31-40	-	2.57	Р	N11 131-140	6,722	5.96	
G5A 270-280		21.5	ъ	G14 81-90	71.4	1.75		N11 181-190	12.4	2.44	
G6 70-80	305	-	Р	G14 131-140	246	2.54	D	N11 231-240	29,261	2.66	
G6 120-130	328	20.4		G22 31-40	35.4	9.26	Р	N11 281-290	12.6	3.95	
G6 170-180	542	42.3		G22 81-90	65.9	-		N11 331-340	12.3	0.40	
G6 220-230	761	47.8	р	G22 131-140	310	17.9		N11 381-340	11.7	0.63	
G10 20-30	864	-	Р	G22 181-190	760	51.0		N13 31-40	349	8.71	F
G10 70-80	5,283	-	Р	G22 231-240	1,427 4,143	4.19		N13 81-90	366	11.3	
G15 20-30	283	-	P	G22 280-290	· ·	16.5		N13 131-140	598	9.65	
G15 70-80	507	-		G22 331-340	61,638			N13 181-190	434	6.94	
G15 120-130	1,143	5.56		G22 381-390	71,287			N13 231-240	382	10.7	
G15 220-230	4,539	74.5		G22 430-440	68,490		D	N13 281-290	389	11.6	
G15 250-260	5,742	81.4	D	G24 31-40	-	2.35	Р	N13 331-340	473	9.33	
G17 20-30	5,145	3.85	Р	G24 81-90	3.02	1.14		N13 381-390	369	9.13	
G17 70-80	24,699	167	Р	G24 131-140	-	4.24 0.50		N13 431-440	373	9.54	
G19 20-30	321	-	Р	G24 181-190 G24 231-240	5.31			N13 481-490	347	-	
G19 70-80	179 2,878	- 26.2		G24 231-240 G24 281-290	4.18 -	0.64 -		N13 531-540	376	9.91	
G19 120-130 G19 170-180	363	20.2 8.18		G24 281-290 G24 381-390	- 5.42	- 2.68					
G19 170-180 G21 70-80	378	0.10 -	Р	G24 431-440	0.14	0.08					
G21 170-80 G21 170-180	220	-	1	N1 31-40	6.54	1.42	Р				
G21 170-180 G21 220-230	425	_		N1 81-90	24.5	4.14	1				
G21 220-230 G21 270-280	20,703	_		N1 131-140	74.3	1.52					
G23 20-30	515	140	Р	N1 181-190	850	6.96					
G23 70-80	370	244		N1 441-445 450-455	2,023	1.96	lost 445-450cm				
		109		N2 81-90	7.64	3.58	P				
G23 170-180	109,412			N2 131-140	4.17	0.83					
G23 220-230	41,038	5.34		N4 31-40	377	10.5	Р				
G23 270-280	108,190			N4 81-90	380	9.84					
G23 320-330	133,354			N4 131-140	372	-					
G23 370-380	113,477			N4 191-190	393	8.82					
G23 420-430		-		N4 231-240	395	11.8					
GA 20-30	330	-	Р	N4 281-290	299	7.42					
GA 120-130	297	-	P	N4 331-340	421	10.7					
GA 220-230	281	-		N4 381-390	475	11.0					
GC 70-80	512	-		N4 431-440	518	11.3					
GC 220-230	1,203	-		N4 481-490	624	11.8					
GC 320-330	3,722	46.0		N4 531-540	938	13.0					
GD 70-80	-	36.1	Р	N6 31-40	9.11	0.83	Р				
GD 170-180	-	-		N6 81-90	7.56	3.73					
GD 270-280	835	8.46		N6 131-140	7.45	4.06					
				N6 181-190	24.1	0.51					
				N6 231-240	23.9	2.90					
				N6 281-290	762	4.79					
				N6 331-340	14,651	1.05					
				N6 381-390	64,263	0.75					
				N6 431-440	74,908	0.95					

Table 2. Methane and ethane concentrations of cored samples in this study.

Table 2. (Continued)

ORII-1207	CH_4 ($\mu L/L$)	C_2H_6 (µL/L)		ORII-1230 cruise	CH_4 (µL/L)	C_2H_6 (μ L/L)		ORI-732 cruise	CH ₄ (μL/L)	C_2H_6 (μ L/L)	
cruise G14 38-48			G	GA 40-50	<u>(µL/L)</u> 241	<u>(μL/L)</u> 8.79	G		256		G
	351	11.6	G				G	G1 40-50		8.32	G
G14 90-100	356	11.3	C	GA 90-100	235	6.66		G1 90-100	242	8.40	C
G15 40-50	405	11.0	G	GA 140-150	217	7.40	C	G3 40-50	280	8.95	G
G15 90-100	520	11.7	C	GC 40-50	280	6.08	G	G3 90-100	259	8.17	0
G16 40-50	346	12.0	G	GC 90-100	387	7.20	0	G4 40-50	288	8.48	G
G16 90-100	360	14.1	0	GD 40-50	240	7.97	G	G4 90-100	258	8.49	0
G17 40-50	362	13.1	G	GD 90-100	239	8.98		G5 40-50	311	8.53	G
G17 90-100	405	10.6		GD 140-150	238	7.55	~	G5 90-100	275	8.91	
G18 40-50	401	12.4	G	G3 92-102	222	7.56	G	G5 140-150	263	7.90	
G18 90-100	334	11.8		G3 140-150	292	7.29	-	G5 190-200	269	8.31	
G21 40-50	406	10.6	G	G4 90-100	233	5.45	G	G7 40-50	267	8.23	G
G21 90-100	466	11.0		G21 40-50	245	6.83	G	G7 90-100	355	8.95	
G23 40-50	490	13.7	G	G21 90-100	252	5.93		G8 40-50	252	7.87	G
G23 90-100	345	11.2		G21 140-150	220	6.91		G8 90-100	231	8.12	
G24 40-50	340	-	G	G69 40-50	272	8.33	G	G9 40-50	286	8.31	G
G24 90-100	366	12.8		G69 90-100	239	7.45		G9 90-100	20.7	1.64	
G27 40-50	426	13.6	G	G70 90-100	314	7.47	G	G9 137-147	242	6.98	
G27 90-100	446	13.6		G72 40-50	232	7.09	G	G10 40-50	250	8.23	G
G30 40-50	370	12.8	G	G72 90-100	236	6.67		G10 90-100	277	8.88	
G30 90-100	438	12.8		G73 40-50	235	7.16	G	G17 40-50	231	10.3	G
G31 40-50	408	8.25	G	G81 40-50	236	6.55	G	G18 40-50	297	8.25	G
G46 40-50	504	12.0	G	G81 90-100	246	-		G26 40-50	263	8.21	G
G47 40-50	415	12.1	G	G82 40-50	173	21.6	G	G26 90-100	241	8.90	
G47 90-100	366	12.0		G82 90-100	245	7.71		G26 140-150	233	8.57	
G49 40-50	486	12.0	G	G85 40-50	233	6.67	G	G27 40-50	266	8.42	G
G50 40-50	404	14.4	Ğ	G85 90-100	239	7.35	0	G27 90-100	303	7.89	0
000 10 00		1	0	G85 140-149	272	8.18		G28 40-50	288	8.23	G
				G86 40-50	239	5.08	G	G28 90-100	239	8.07	0
				G86 87-97	152	4.49	0	G28 140-150	261	8.32	
				G87 40-50	253	7.50	G	G20 140-150 G31 40-50	255	8.28	G
				G87 90-100	250	7.60	U	G31 90-100	270	7.99	U
				G88 40-50	288	9.23	G	G33 40-50	241	8.41	G
				G88 90-100	288 254	9.23 8.64	U	G36 40-50	359	8.60	G
					234 244	8.04 7.53	C		290	8.00	U
				G89 40-50			G	G36 90-100			р
				G89 90-100	243	6.42	0	G96 34-43	1,277	8.05	Р
				G90 40-50	228	12.6	G	G96 84-93	2,408	9.11	
				G91 40-50	140	3.56	G	G96 134-143	,	11.8	
				G91 90-100	315	5.97	~	G96 184-193	·	25.9	
				G94 40-50	236	6.43	G	G96 234-243	12,677	38.7	
				G95 40-50	186	5.53	G	G96 284-293	,	30.2	
				G95 90-100	265	5.62		G96 334-343	· · ·	29.6	
				G96 41-50	16,325	46.0	G	G96 384-393		34.4	
				G96 90-100	30,730	40.3		G96 434-443	40,585	31.2	
				G99 40-50	233	9.12	G				
				G99 90-100	245	6.07					
				G103 40-50	242	7.86	G				
				G103 90-100	304	7.88					
				G103 140-150	266	6.50					
				G105 40-50	233	7.31	G				
				G105 90-100	235	9.37	0				
				G105 90-100 G105 135-145	233 251	9.57					
							C				
				G107 40-50	232	6.08	G				
				G109 38-48	244	7.78	G				
				G111 40-50	265	6.43	G				

Note : G = Gravity core; P = Piston Core.



Fig. 3. Methane profiles of cored samples in this study. Figs. 3a and b (1): Methane concentrations are low in the shallower depth, which increases with increasing depth ($< 10 \sim 10^5 \ \mu L L^{-1}$). Figs. 3a and b (2): Methane profiles at some sites show a saw-tooth shape ($> 10^4 \ \mu L L^{-1}$; $< 10 \ \mu L L^{-1}$). Figs. 3a and b (3): Methane concentrations do not change much with depth and maintain similar values ($0 \sim 10^3 \ \mu L L^{-1}$). Note that CH₄ concentrations have been converted from reported units ($\mu L L^{-1}$). The number in parentheses represents the methane concentration (nL L⁻¹) in bottom seawaters at that site.

Geochemical profiles for cored samples are plotted in Fig. 4 with corresponding sulfate concentrations (Lin et al. 2006) in pore water and methane concentrations in pore space. According to the depths of the SMI, the profiles can be grouped into three categories as discussed below:

- (a) **No obvious sulfate reduction trend and no discernable SMI (Fig. 4a):** The profiles show that methane concentrations are very low in pore spaces showing no changes with increasing depths and sulfates are not depleted. This may be due to two reasons: low methane fluxes, or the short length of cores. The sites where the depths of SMI cannot be determined are: G1, G2, G6, GA, GC, G10 of ORI-697, and G4, G11, G9, G24, N1, N2, N9, N11, N13 of ORI-718.
- (b) There is obvious sulfate reduction trend, and depths of the SMI can be predicted at a few meters below the core length (< 10 m) (Fig. 4b): Profiles show that methane concentrations do not change with increasing depths. Only some stations contain higher methane concentrations in the bottom part of cored samples. Due to the short lengths of cored samples, the depths of the SMI cannot be measured. However, there might be higher methane fluxes in deeper sediments that affect the sulfate profiles with increasing depth. The curves of the sulfate profiles show a steep slope. Therefore, the depths of the SMI might be less than 10 m based on sulfate gradients, e.g., sites: G3, G15, G17, G19, G21 of ORI-697, and G14, N4 of ORI-718.</p>
- (c) The sulfate reduction trend is very sharp and the SMI appears at very shallow depths (<4 m) (Fig. 4c): The profiles show that the depths of the SMI range between 100 and 400 cm, e.g., G5A, G23 (the sudden decrease of methane concentration at 225 cm might be due to leakage or sampling problems such as the introduction of air during storage) of ORI-697; G22, N6, N8 of ORI-718, and G96 of ORI-732. Note that the methane and sulfate profiles of gas hydrate bearing sediments from Blake Ridge at Ocean Drilling Program (ODP) sites 994, 995, and 997 show that the depths of the SMI are around 20 m (Dickens 2001). Compared to the results of ODP 204 sites, the sites in this study present much shallower SMI depths, especially at sites G23 of ORI-697 and N8 of ORI-718 (<100 cm). The low methane concentrations (~10 nL L⁻¹) are controlled by AMO in the sulfate reduction zone. Below 100 ~ 300 cm, methane concentrations can be up to 10⁴ nL L⁻¹ and sulfate becomes totally depleted. To sum up, the stations shown in Figs. 4b and c have very shallow depths of the SMI indicating that there should be very high methane fluxes below the sea floors in the region.

4.2. Extremely High Methane Concentration

In general, dissolved methane concentration in average seawater is very low, about 20 nL L⁻¹. For example, in the Gulf of Mexico excluding areas where gas hydrate were found, methane concentration below 200 m is less than 50 nL L⁻¹ except whenever methane is saturated due to bacterial activity or photochemical processes (Millero 1996). In the area offshore SW Taiwan, measured methane concentrations in core top water samples are higher than general seawater background values and vary from 10^2 to 10^5 nL L⁻¹. Moreover, some extremely high methane



Fig. 4. Selected profiles of methane concentrations of pore space in cored samples and dissolved sulfate concentrations in pore water. (a) The depths of SMI cannot be determined because of the short length of cored samples; (b) The depths of SMI cannot be determined. According to the sulfate gradient, it can be predicted that the depths of SMI are very shallow (<10 m); (c) The depths of SMI can be determined and are very shallow (<4 m). The shadow in different profiles shows the possibility range of the depths of SMI. The data of sulfate concentrations are from Lin et al. (2006).</p>

concentrations (> 100000 nL L⁻¹) have been found in the study area, e.g., at sites G23 of ORI-697, N8 of ORI-718, and G96 of ORI-732. Furthermore, these values are even higher than the very high dissolved methane concentration in seawater columns of the Hydrate Ridge area (Suess et al. 1999). This indicates that there are active venting sources injecting methane into seawater in this area.

Based on the results of methane concentrations in the pore spaces of cored samples, there are extremely high methane concentrations at some sites, especially sites G23 of ORI-697, G22, N8 of ORI-718, and G96 of ORI-732. The highest methane concentration was detected at depths of 320 - 330 cm at site G23 of ORI-697 and its value is 133354 (μ L L⁻¹). This value is higher than that obtained at the gas hydrate and gas venting rich area of the Blake Ridge (Kvenvolden and Lorenson 2000; Lorenson and Collett 2000).

Generally, the value of dissolved methane concentration in bottom seawater indicates whether there is a gas source beneath the sea floor or not at that station and can be taken as an efficient method of investigating whether or not there is a methane vent on the sea floor. Comparing the results of dissolved methane concentrations in bottom seawater with methane concentrations of pore spaces in sediments (Table 1; Table 2; Fig. 2), there is good correlation at sites of ORI-697, i.e., the higher the dissolved methane concentration in bottom seawater, the higher the methane concentrations of pore spaces in cored samples and vise versa. However, no good correlations are found at other sites especially in ORII-1230 and ORI-732 cruises except sites G22, N8 of ORI-718, and G96 of ORI-732. Most of the stations contain high dissolved methane concentrations without unusually high methane concentrations in the pore spaces of cored samples ($[CH_4] < 400 \ \mu L \ L^{-1}$).

The uncorrelated results of dissolved methane concentrations in bottom seawater and methane concentrations of pore space in sediments of ORII-120 and ORI-732 cruises might be caused by the water samples. As the bottom water samples are represented by gravity core top water samples, they might be disturbed by core top sediments. Therefore, methane in the pore spaces may have been released into the core top water, making the core top water rich in highly dissolved methane. Note that the methane concentrations of pore spaces are much higher than the dissolved methane concentrations in bottom water (up to 3 orders of magnitude). Although methane concentrations of pore spaces do not have unusually high values (presume 200 μ L L⁻¹), the dissolved methane concentrations can easily reach abnormally high values (200000 nL L⁻¹) after being disturbed by methane of the pore spaces. For this reason, the methane concentrations of pore spaces in core top sediments (the average depths range from 30 - 50 cm) are treated as an indicator of methane vent on the seafloor (Table 3; Fig. 5).

Actually, the dissolved methane concentrations of core top seawater can not be used as the only indicator to estimate if there are methane vents on the seafloor. In order to study the unusual values at one station, dissolved methane concentrations at different depths of the water column (e.g., Chen and Tseng 2006; Yang et al. 2006) and methane concentration of pore spaces at different depths of cored samples need to be taken into consideration.

Dissolved methane concentrations in bottom water and methane concentrations of pore spaces in deeper sediments are extremely high at sites N8, G23 of ORI-697, G22 of ORI-718, and G96 of ORI-732. The depths of the SMI in these four sites are also very shallow. As mentioned before, the stations have very shallow SMI depths indicating that there might be

ORI-697	pore space CH ₄ (μL/L)	bottom water CH ₄ (nL/L)	OR2-1230	pore space CH ₄ (μL/L)	bottom water CH ₄ (nL/L)
G1 20-30	540 (663)	282	GA 40-50	241	19,733
G5A 70-80	166 (22,251)	212	GD 40-50	240	4,897
G6 70-80	305 (761)	148	G3 40-50	292	21,239
G10 20-30	864 (5,283)	156	G21 40-50	245 (252)	53,888
G15 20-30	283 (5,742)	156	G69 40-50	272	30,196
G17 20-30	5,145 (24,699)	1,221	G73 40-50	235	11,035
G19 20-30	321 (2,878)	390	G81 40-50	236 (246)	41,298
G21 70-80	378 (452)	613	G82 40-50	173 (245)	65,153
G23 20-30	515 (133,354)	198,616	G85 40-50	233 (272)	135
ORI-718			G86 40-50	239	2,666
G4P 31-40	375 (745)	3,740	G87 40-50	253	20,633
G9B-P 31-40	867 (3,668)	4,970	G89 40-50	244	58,259
G11B-P 31-40	384 (1,138)	3,638	G90 40-50	228	100
G11C 31-40	392	822	G91 40-50	140 (315)	44,383
G14 81-90	71 (246)	10,838	G94 40-50	236	11,120
G22 31-40	35 (68,490)	78,731	G95 40-50	186 (265)	36,844
G24 31-40	3 (5)	935	G96 41-50	16,325 (30,730)	47,551
N1 31-40	7 (2,024)	6,204	G99 40-50	233 (245)	18,732
N2 81-90	8	9,750	G103 40-50	242 (304)	2,200
N4P 31-40	377 (938)	3,793	G105 40-50	233 (251)	988
N6 31-40	9 (74,908)	2,450	G107 40-50	232	7,453
N8 31-40	117 (121,716)	613,268	G109 38-48	244	475
N9 31-40	9 (6,457)	12,702	G111 40-50	265	12
N11 31-40	7 (29,261)	10,544	ORI-732		
OR2-1207		-	G1 40-50	256	5,365
G15 40-50	405 (520)	384	G3 40-50	280	45,797
G16 40-50	346 (360)	441	G4 40-50	288	5,494
G18 40-50	401	792	G5 40-50	311	81,490
G24 40-50	340 (366)	723	G7 40-50	267 (355)	8,667
G27 40-50	425 (446)	749	G8 40-50	252	6,230
G30 40-50	370	458	G9 40-50	286	64,117
G47 40-50	415	328	G10 40-50	250 (277)	13,364
G50 40-50	404	597	G17 40-50	231	53,796
ORI-732			G18 40-50	297	21,471
G31 40-50	255 (270)	4,186	G26 40-50	263	140
G33 40-50	241	8,751	G27 40-50	266 (303)	44,743
G36 40-50	359	141	G28 40-50	288	13,194
G96 34-43	1,277 (42,826)	536,323			

 Table 3. Comparison with dissolved methane in bottom waters and methane concentration of cored samples at the same site.

Note: The number in parentheses represents the highest methane concentration of pore-spaces at that site.

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Fig. 5. Distribution of methane concentrations (in μ L L⁻¹) of pore spaces in offshore southern Taiwan. Note that plotting methane concentrations from ORI-697 and ORI-718 are taken the data from upper 20 ~ 30 m sediments; while, those from ORII-1207, ORII-1230, and ORI-732 are from upper 40 ~ 50 m sediment.

very high methane fluxes below the sea floor, which causes the extremely high methane concentrations at these four sites. Moreover, besides the intensive BSRs (Liu et al. 2006) and the very high methane fluxes, there are high sulfate reduction rates, authigenic carbonates and pyrites existing at sites N8, G23 of ORI-697, and G22 of ORI-718 (Huang et al. 2006; Horng et al. 2006; Jiang et al. 2006; Lin et al. 2006). Therefore, beneath these sites, the upward migrating methane through shallow sediments into the seawater might be released from the dissociation of gas hydrates.

5. CONCLUSIONS

- 1. Sediments and bottom waters were systematically collected by piston cores and gravity cores during five cruises from 2003 to 2005. The results show that many sites exhibit extremely high methane concentrations in both bottom water and sediments pore space compared to other gas hydrate areas in the world.
- 2. In general, the dissolved methane concentrations in bottom seawater are very high (> 1000 nL L⁻¹). Most values are much higher than the average seawater value (< 50 nL L⁻¹), especially at site N8 of ORI-718, where the value reaches 613268 nL L⁻¹. This implies that a large amount of methane is venting from deep sediments up to the seawater in a wide area.
- 3. Methane is the main component of hydrocarbon gases in the study area, while ethane is the minor component and C_{3+} gases are usually below detection limits. Most of the cored sediments collected show a positive correlation of methane concentration in pore space with depth.
- 4. Methane concentrations at many sampling sites are higher than 100000 μ L L⁻¹, e.g., site G23 of ORI-697 and N8 of ORI-718. This infers that methane sources exist beneath the sea floor at these sites.
- 5. From methane and sulfate depth profiles, it has been found that the depths of the SMI are very shallow at many stations, which indicate the presence of very high methane fluxes. The high methane fluxes in this area might have been caused by the dissociation of gas hydrates below the sea floor. After gas hydrates dissociate, a large amount of methane is released and migrates upward to the shallow parts of sediment and the seawater. This can explain the unusually high methane concentration observed in this area.

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