Molecular characterization of the E gene of dengue virus type 1 isolated in Guangdong province, China, in 2006

K. ZHENG¹, H.-Q. ZHOU¹, J. YAN¹, C.-W. KE^{1*}, A. MAEDA², J. MAEDA², I. TAKASHIMA³, I. KURANE⁴, H. MA⁵ AND X.-M. XIE¹

(Accepted 27 February 2008; first published online 4 April 2008)

SUMMARY

We determined the genetic relationships and origin of the dengue virus (DENV) responsible for an outbreak of dengue fever (DF) in Guangdong province, China, in 2006. Five DENV type 1 (DENV-1) isolates were obtained from human serum samples collected from DF patients during the outbreak. The nucleotide sequences of the *E* (envelope) gene were compared with those of 48 previous DENV-1 isolates: 18 from Guangdong province, one from Fujian province, one from Zhejiang province, and 28 from other countries in the South Asian region. The results suggested that four DENV-1 isolates identified in Guangdong province in 2006 might be in general circulation there, although these DENV-1 viruses may have been originally introduced into the province from other countries. In contrast, one isolate from Guangzhou city in 2006, may have been introduced by a recently imported case from Cambodia.

INTRODUCTION

Dengue fever/dengue haemorrhagic fever (DF/DHF) is a mosquito-borne disease caused by dengue virus (DENV). There are four known serotypes: dengue virus types 1, 2, 3 and 4 (DENV-1 to DENV-4) [1, 2]. DF/DHF occurs mainly in tropical or subtropical countries, e.g. in Southeast Asia, the Americas and the Caribbean. In the 1970s, there were a series of DF outbreaks in some of the southern provinces in China. The first outbreak was caused by DENV-4, the second by DENV-1, and the third by DENV-3 [3]. DF/DHF

(Email: kecw1965@yahoo.com.cn)

has been an infectious disease of particular concern in Guangdong province, China, since 1978 when a DF outbreak occurred in Foshan city [4]. The first DHF case was reported in 1985-1986, in Hainan Island, China [5, 6]. Before this epidemic in China, DF/DHF outbreaks had been reported in Thailand, Vietnam, Indonesia [7, 8], and Myanmar [9]. In the 1990s, large outbreaks of DF occurred in China, with more than 6000 cases reported [10]. In recent years, highly localized and relatively sporadic yearly outbreaks have been observed in China. The DF/DHF outbreaks in Guangdong province were thought to be related to other worldwide outbreaks in that they were caused mainly by imported cases from other countries [11–13]. Previously there had been no evidence of endemic dengue in Guangdong province; however,

¹ Center for Diseases Control and Prevention of Guangdong province, Guangzhou, PR China

² Laboratory of Prion Diseases, Graduate School of Veterinary Medicine, Hokkaido University, Sapporo, Japan

⁸ Laboratory of Veterinary Public Health, Graduate School of Veterinary Medicine, Hokkaido University, Sapporo, Japan

⁴ Department of Virology 1, National Institute of Infectious Diseases, Tokyo, Japan

⁵ Zhuhai Entry-Exit Inspection and Quarantine Bureau, Guangdong province, Zhuhai, PR China

^{*} Author for correspondence: Dr C.-W. Ke, Center for Diseases Control and Prevention of Guangdong Province, Guangzhou 510300, PR China.

a report on the DF epidemic in Guangzhou in 2005 suggested that DENV had recently become endemic in some areas in China [14]. Phylogenetic analysis of the entire sequence of the E gene of DENV-1 is a powerful molecular epidemiological method for the confirmation of endemic and imported cases. Studies of DENV molecular evolution and sequence data were performed to determine the phylogenetic relationships among viruses within each serotype. A threshold of 6% divergence is currently used to separate different genotypes within a dengue serotype [15–17]. Several phylogenetic studies analysing a large number of DENV-1 virus genomic sequences have been published [12, 15]. These studies analysed either a 240-nt sequence in the E/NS1 junction region of the genome or a 180-nt sequence in the E gene and suggested that there were potentially five or three DENV-1 virus genotypes, respectively. A recent study, based on nucleotide sequences of the entire E gene, showed that DENV-1 viruses can be classified into five genotypes [18, 19]. One study on full-length sequences of DENV-1 showed that DENV-1 viruses can be classified into four genotypes [20]. In this study, the entire E gene sequence was used to trace the origin of the DENV-1 virus/strain known to have caused the DF outbreak in Guangdong province in 2006.

In the second half of 2006, DF outbreaks occurred in Guangzhou, Yangjiang, Nanhai and Shantou City in Guangdong province, with the most severe epidemic occurring in Guangzhou. The cause of the outbreak was confirmed as DENV-1. A total of 1010 DF cases were reported in Guangdong province (CDC of Guandong Province, unpublished data). As the origin of these outbreaks was not determined by epidemiological investigation, the question arose again as to whether the DF/DHF outbreaks had become endemic in Guangdong province. To answer this question, we isolated DENV from serum samples obtained from patients in these cities that were found to be positive by TaqMan real-time PCR [21]. The E gene of the Guangdong isolates were sequenced and compared to those of DENV strains isolated in recent years in other parts of China and Southeast Asia.

METHODS

Serum samples, cell lines and viruses

After obtaining informed consent, a total of 103 serum samples were collected from febrile patients during

the DF outbreak in Guangdong province in 2006. The mosquito cell line, C6/36, was passaged in Dulbecco's modified Eagle's medium supplemented with 10% fetal calf serum and used for virus isolation. Twenty-three serum samples in which DENV RNAs were found to be positive by real-time PCR [15], were inoculated to C6/36 mosquito cells, and 16 isolates were identified as DENV-1. The DENV strains isolated from patients in recent years and which were identified as DENV-1 were also inoculated to C6/36 cells for passaging. The viruses analysed in the present study are listed in Table 1 together with their GenBank accession number and geographic origin.

RNA extraction, primer design and reverse transcription-polymerase chain reaction (RT-PCR)

Viral RNAs were extracted from the supernatant of cell culture, to which serum from DF patients or previous DENV isolates had been inoculated and shown a CPE (cell pathogenic effect), using a QIAamp Viral RNA mini kit (Qiagen, Hilden, Germany) according to the manufacturer's protocol. To amplify the entire sequence of the DENV E gene, we designed three pairs of primers for RT-PCR with Primer Express version 3.0 (Applied Biosystems, Foster City, CA, USA), as shown in Table 2. RNA samples (5 µl) were used for one-step RT-PCR (TaKaRa, Shiga, Japan). Briefly, the RT reaction was performed at 50 °C for 30 min and the RT enzyme was denatured at 95 °C for 15 min. Next, the PCR reaction was performed for three cycles at 94 °C for 1 min, 60 °C for 40 s, and 72 °C for 1 min. Subsequently, 30 cycles at 94 °C for 30 s, 60 °C for 30 s, and 72 °C for 40 s and a final 72 °C for 10 min were performed. PCR products were purified with a QIAquick PCR Purification kit (Qiagen) according to the manufacturer's protocol.

Sequencing and genetic analysis

The PCR product was sequenced in both directions using a BigDye Terminator version 3.1 cycle sequencing kit (Applied Biosystems). Sequencing reaction products were purified with Autoseq G-50 (GE Healthcare–Amersham, Piscataway, NJ, USA), and the nucleotide sequences were determined by ABI Prism[®] 3100 genetic analyser. The sequences of D1E1, D1E2 and D1E3 were edited using Sequencher[™] 4.7 (Gene Codes Corporation, Ann Arbor, MI, USA). The sequences of the DENV-1

Table 1. Dengue fever virus strains used in this study

Geographic	Year	Ctura in	C . 1	Accession
origin	isolated	Strain	Code	no.
Thailand	1963	2543-63	Thai63	AF425629
Thailand	1964	16007	Thai64	AF180817
Malaysia	1972	P72-1244	Mala72	AF425622
Philippines	1974	PRS 228682	Phil74	AF425627
Nauru Island	1974	West Pac 74	Naur74	U88535
Guangzhou, China	1980	GZ/80	GZ80	AF350498
Thailand	1980	PUO 359	Thai80	AF425630
Thailand	1982	ThD1_0041_82	Thai82	AY732378
Mexico	1983	1378	Mexi83	AF425624
Taiwan, China	1987	765101	Taiw87	AF425628
Indonesia	1988	A88	Indo88	AB074761
Thailand	1991	ThD1_0119_91	Thai91	AY732413
Guangzhou, China	1993	GZ01/03	GZ93	DQ211348
Guangdong, China	1995	GD23/95	GD95	AY373427
Guangzhou, China	1995	GZ01/95	GZ95	DQ855297
Thailand	1996	ThD1_0214_96	Thai96	AY732422
Brazil	1997	BeH 584526	Braz97	AF425614
Guangdong, China	1997	GD14/97	GD97	AY376737
Thailand	1997	ThD1_0277_97	Thai97	AY732418
Thailand	1998	ThD1 0388 98	Thai98	AY732434
Nanhai, China	1998	D98039	D98039	EF508198
Ivory Coast	1999	D1/H/IMTSSA-	IvCo99	AF298807
Troff Coust	1000	ABID/99/1056	1,0000	711 250007
Thailand	1999	ThD1 K0051 99	Thai99	AY732458
Zhongshan, China	1999	D99020	D99020	EF508199
Argentina	2000	301arg00	Arg00	AF514876
Paraguay	2000	280par00	Par00	AF514878
Thailand	2000	ThD1_0141_00	Thai00	AY732408
Cambodia	2001	01-61-1HuNIID	Cam01	AB111071
Micronesia	2004	—	Mic04	AB178040
Brazil	2001	BR/01-MR	Braz01	AF513110
Myanmar	2001	My01D1m193	My01	AY620953
French Polynesia	2001	FP/01/192206	FP01	AY630407
Thailand	2001	ThD1_0049_01	Thai01	AY732482
Yangjiang, China	2001	D01048	D01048	EF508200
Thailand	2002	02-33-1HuNIID	Thai02	AB111077
Indonesia	2002	D1/Hu/Indonesia/	Indo02	AB232666
muonesia	2002	NIID09/2002	1110002	AB232000
Guangzhou, China	2002	D02031	D02031	EF508201
Guangzhou, China	2002	GZ01/02	GZ02	DQ855296
Guangzhou, China	2002	A04137	A04137	EF508202
Zhejiang, China	2004	Zhejiang/07/04	ZJ04	AY871812
Fujian, China	2004	Fj231/04	ZJ04 FJ04	DQ193572
Guangzhou, China	2004	GZ01/04	GZ04	EF032589
Yangjiang, China	2004	D06045	D06045	EF508204
Nanhai, China	2006	D06068	D06043	EF508204
Shantou, China	2006	D06098	D06098	EF508205 EF508206
Chaozhou, China	2006 2006		D06098 D060117	
		D060117		EF508207
Guangzhou, China	2006	GZ16/2006 GZ061707	GZ06 CZ061707	EF113152
Guangzhou, China	2006	GZ001/0/	GZ061707	EF508203

E gene amplified in the study were BLASTed against GenBank sequences. The sequences were analysed with the edited sequences of the E gene using

Clustal X v.1.83 software. The sequence homologies of nucleic acids and amino acids were calculated and a phylogenetic tree was constructed by the

Primer name	Sequence (from 5' to 3')	Polarity	Position*	Amplicon name
D1E1854	TTAGCACACGCCATAGGAACATC	+	854–876	D1E1
D1E11440	GGAGGTTGAGGTGTTATGGTTGC	_	1418-1440	
D1E21292	AAGTGTGTGACAAAACTGGAAGG	+	1292-1314	D1E2
D1E21972	GGTCACTCCCTTCTCATCTTGG	_	951-1972	
D1E31857	TAGAGAAGGAAGTGGCTGAGACC	+	1857-1879	D1E3
D1E32518	TTTGTATTGCTCTGTCCAAGTGTG	_	2495–2518	

^{*} Based on the numbering of 01-61-1HuNIID strain (accession number AB111071).

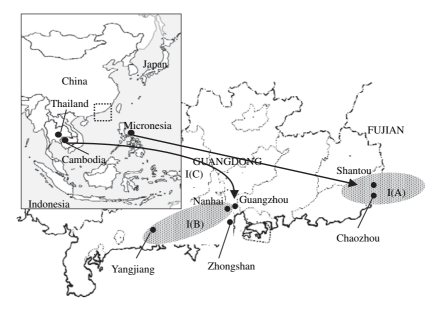


Fig. 1. Geographic representation of locations from which the dengue viruses were isolated. The inset map on the left shows the Southeast Asian region from which the dengue viruses, whose sequences were used in this study, were isolated. The main map shows an enlargement of Guangdong province and the marked areas indicate the locations from which DENV strains were collected. Our hypothesis regarding the origins of the dengue outbreaks is shown schematically. The isolates D06045 and D06068 in cluster B might have originated from a virus imported from Thailand in 2001 before circulating within Guangdong province, while the isolates D06098 and D060117 in cluster A might be derived from a virus imported from Micronesia before circulating within Guangdong province. The two isolates, GZ06 and GZ061707, identified in Guangzhou in 2006, might have two different origins. The former might have come from Thailand and the latter might have originated from Cambodia in 2001. These viruses re-emerged in Guangdong province as endemic in 2006.

neighbour-joining method using MEGA 3.1 software. One thousand bootstrap repetitions were used for confirmation of the statistical significance of the phylogenetic analysis. The geographic regions from which serum samples were obtained are shown in Figure 1.

RESULTS

Five of 16 DENV-1 isolates obtained from patient serum samples collected during the DF outbreak in Guangdong province in 2006 were sequenced. The sequence of the E gene was determined and analysed phylogenetically along with 43 previously reported isolates (Fig. 2). DENV-1 is divided into five genotypes. The DENV-1 endemic in Southeast Asia consists of genotypes I and IV, and the isolates from Guangdong in 2006 belong to genotype I. For genotype I, seven independent sub-genotypes, A–G, were identified by sequence analysis of the E gene and the homologies of the nucleic acids and amino acids within each sub-genotype were >99%. Sub-genotype A consisted of two isolates from the Guangdong DF outbreak in 2006, Shantou (D06098) and Chaozhou

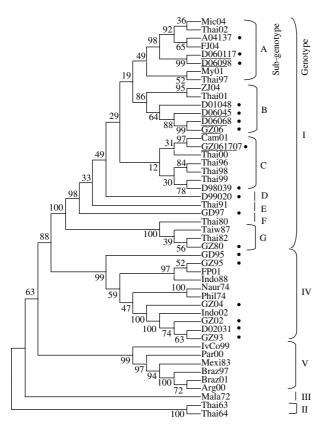


Fig. 2. Phylogenetic analysis of DENV strains isolated during outbreaks, including that in Guangdong province in 2006. The phylogenetic tree is based on the *E* gene sequences of five isolated DENV strains we sequenced, and 43 sequences from Genbank. The strains isolated in Guangdong province are indicated by black dots (●) to the right of the strain name. The Guangdong isolates are distributed across two genotypes, genotype I and genotype IV, while those in the main sub-genotypes A−G are clustered in genotype I.

(D060117), and one isolate from Guangzhou (A04137) and one from Fujian (DQ193572) in 2004. An isolate from Micronesia in 2004 (AB178040) also belonged to this sub-genotype. Sub-genotype B contained three DENV-1 isolates from the 2006 DF outbreak in Yangjiang (D06045), Nanhai (D06068) and Guangzhou (GZ06) cities, Guangdong province, and one DENV-1 isolate from 2001 in Yangjiang (D001048) (Fig. 2). Yangjiang and Nanhai cities are located to the west of Guangzhou city and are relatively close to each other (Fig. 1). Interestingly, subgenotype A also contained a virus isolate AY732482 from Thailand in 2001. Sub-genotype C contained four DENV-1 isolates from Thailand obtained during 1996 or 2000 (Thai96, Thai98 Thai99 Thai00) and one from Nanhai in 1998 (D98039), one from Cambodia in 2001 (Cam01) and one from Guangzhou in 2006 (GZ061707). In sub-genotypes D-G, the isolates were

obtained from China and other Southeast Asian countries prior to 1990.

DISCUSSION

In previous DF/DHF outbreaks in Guangdong, China, the circulating viruses could be traced back to other countries through the analysis of DENV gene sequences [12]. Thus, it is believed that the DF/ DHF outbreaks in Guangdong were caused mainly by cases imported from Southeast Asian countries around the time of the epidemics in Guangdong [3-10, 14]. For example, the DF outbreak in Guangzhou, 2002, may have been caused by a case entering the province from Indonesia (Fig. 2, genotype IV), as a DF outbreak was reported at the same time in Indonesia, and the E gene sequences of the viruses isolated from Guangzhou and Indonesia were very similar. Therefore, we hypothesize that the DF epidemics in Yangjiang in 2001 (Fig. 2, sub-genotype B), Nanhai in 1998 (Fig. 2, sub-genotype C), Zhongshan in 1999 (Fig. 2, sub-genotype D), and Guangzhou in 2004 (Fig. 2, sub-genotype A) may have been caused by cases originating in Thailand (for sub-genotype A, and B and C isolates) or Micronesia (Fig. 1).

The scenario for the DF outbreak in Guangdong province in 2006 was different from those of previous outbreaks in Guangdong as the sequence comparison of the E gene of the viruses suggested that the 2006 outbreak was a case of endemic infection of dengue circulating locally in the province in 2006. The E gene sequences of the viruses isolated in Shantou (D06098) and Chaozhou (D060117) were very similar to those isolated in Guangzhou (A04137) and Fujian in 2004 (FJ04) (Fig. 2, sub-genotype A). Although the possibility that the isolated virus may have been imported from another province in 2006 still remains, our hypothesis is that the viruses circulating in Micronesia in 2004 were imported into Guangdong and Fujian provinces in 2004, and re-emerged endemically in Shantou and Chaozhou cities, Guangdong province, in 2006. The E gene sequences of the viruses isolated in Yangjiang and Nanhai cities (D06045 and D06068, respectively) were similar to that of the virus isolated in Yangjiang (D01048) in 2001 (Fig. 2, sub-genotype B). The virus isolated in Yangjiang (D01048) may have been imported from Thailand during a DF epidemic that occurred there in 2001. One of the isolates from Guangzhou in 2006 (GZ061707), together with isolate AB111071 obtained in Cambodia in 2001, belonged to sub-genotype C. No viruses isolated in Guangdong province have been reported with an *E* gene sequence similar to that isolated in Cambodia in 2001. However, the epidemic might have been caused by an imported virus that had been circulating in Cambodia in 2001 (Fig. 1).

According to our experimental data, we demonstrated the existence of endemic DF cases in the 2006 outbreak in Guangdong province, and that virus strains imported from Southeast Asian countries were also in circulation at the same time.

ACKNOWLEDGEMENTS

This work was supported by Grant-in-aids for Scientific Research and the Programme of Excellence for Zoonosis Control, the 21st Century COE Program, Japanese Ministry of Education, Culture, Sports, Science and Technology (18580301 and 17255009) and Japanese Ministry of Health, Labour and Welfare (H15-Shinkou-17 and H17-Shinkou-ippan-018). We thank Dr H.-M. Wang for his careful revision of the manuscript.

DECLARATION OF INTEREST

None.

REFERENCES

- 1. **Heinz F,** *et al.* Family Flaviviridae. In: van Regenmortel MHV, *et al.* eds. *Virus Taxonomy*. *Seventh Report of the International Committee on Taxonomy of Viruses*. San Diego: Academic Press, 2000, pp. 859–878.
- Thomas SJ, Strickman D, Vaughn DW. Dengue epidemiology: virus epidemiology, ecology, and emergence. In: Chambers TJ, Monath TP, eds. Advances in Virus Research, Vol. 61, The Flaviviruses: Detection, Diagnosis and Vaccine Development. San Diego: Elsevier, Academic Press, 2003, pp. 235–289.
- Qiu FX, et al. Dengue in China a clinical review. Bulletin of the World Health Organization 1993; 71: 349–359.
- 4. Wen J, Liang F, Du F. An etiological and serological study on Dengue epidemic in Guangdong Province. *Chinese Journal of Experimental and Clinical Virology* 1998; **12**: 165–168.

- Qiu FX, et al. The first epidemic of dengue hemorrhagic fever in the People's Republic of China. American Journal of Tropical Medicine and Hygiene 1991; 44: 364–370.
- Yang PY, et al. Antigenic signature analysis of dengue-2 viruses strains in Hainan China. Acta Microbiologica Sinica 1989; 29: 299–302.
- Kho LK, et al. Dengue hemorrhagic fever accompanied by encephalopathy in Jakarta. Southeast Asian Journal of Tropical Medicine and Public Health 1981; 12: 83–86.
- 8. van Peenen PF, et al. Dengue with hemorrhage and shock in Jakarta, Indonesia. Southeast Asian Journal of Tropical Medicine and Public Health 1978; 9: 25–32.
- 9. **Gubler DJ.** Dengue. In: Monath TP, ed. *The Arboviruses: Epidemiology and Ecology*, vol. 2. Florida: CRC Press, 1988, pp. 223–260.
- 10. **Anon.** Dengue fever/dengue hemorrhagic fever in the WHO western pacific region. *WPRO Communicable Diseases Bulletin* 2000; December, 1–6.
- Luo H, et al. Analysis on the epidemiologic features of Dengue fever in Guangdong province, 1990–2000. Chinese Journal of Epidemiology 2002; 23: 427–430.
- Zheng K, et al. Nucleotide sequencing of E/NS1 gene segment of dengue type 1 viruses isolated in Guangdong province. Chinese Journal of Experimental and Clinical Virology 2002; 16: 382–384.
- Ren RW, et al. Isolation, identification and sequence analyses of dengue virus type 2 strain GD19/2001. Chinese Journal of Epidemiology 2003; 24: 288–290.
- Zhang FC, et al. Analysis on clinical and epidemiological characterization of 1032 patients with Dengue fever in Guangzhou. Chinese Journal of Epidemiology 2005; 26: 421–423.
- Rico-Hesse R, et al. Molecular evolution and distribution of dengue viruses type 1 and 2 in nature. Virology 1990; 174: 337–343.
- 16. Lewis JA, et al. Phylogenetic relationships of dengue-2 viruses. *Virology* 1993; 197: 216–224.
- Chungue E, et al. Molecular epidemiology of dengue-1 and dengue-4 viruses. *Journal of General Virology* 1995; 76: 1877–1884.
- Goncalvez AP, et al. Diversity and evolution of the envelope gene of dengue virus type 1. Virology 2002; 303: 110–119
- Laille M, et al. Comparison of dengue-1 virus envelope glycoprotein gene sequence from French Polynesia. American Journal of Tropical Medicine and Hygiene 2004; 71: 478–484.
- Nukui Y, et al. Novel dengue virus type 1 from travelers to Yap State, Micronesia. Emerging Infectious Diseases 2006; 12: 343–346.
- 21. **Ke C**, *et al.* Detection of Dengue virus by real-time polymerase chain reaction with TaqMan MGB probe. *Chinese Journal of Zoonoses* 2005; **21**: 716–720.