Identification and antibiotic resistance profiling of bacterial isolates from septicaemic soft-shelled turtles (*Pelodiscus sinensis*)

T.H. $Chung^{1\dagger}$, S.W. $Yi^{2\dagger}$, B.S. Kim^2 , W.I. Kim^2 , G.W. $Shin^{2*}$

ABSTRACT: The present study sought to identify pathogens associated with septicaemia in the Chinese soft-shelled turtle (Pelodiscus sinensis) and to characterise antibiotic resistance in these pathogens. Twenty-three iso-lates recovered from the livers of diseased soft-shelled turtles were genetically identified as Aeromonas hydrophila (n = 8), A. veronii (n = 3), Citrobacter freundii (n = 4), Morganella morganii (n = 3), Edwardsiella tarda (n = 2), Wohlfahrtiimonas chitiniclastica (n = 1), Chryseobacterium sp. (n = 1), and Comamonas sp. (n = 1). Most isolates (n = 21) were resistant to ampicillin whereas a low percentage of isolates was susceptible to aminoglycosides (amikacin, gentamicin, and tobramycin). PCR assays and sequence analysis revealed the presence of the qnrS2 and bla_{TEM} antibiotic resistance genes in all isolates. The bla_{DHA-1} , $bla_{CTX-M-14}$ and bla_{CMY-2} genes were harboured by 17.4% (n = 4), 13.5% (n = 3) and 8.7% (n = 2) of the strains, respectively. One or more tetracycline resistance genes were detected in 60.9% (n = 14) of the isolates. Four isolates (17.4%) harboured single or multiple class 1 integron cassettes. Collectively, a variety of bacterial pathogens were involved in the occurrence of septicaemia in Chinese soft-shelled turtles and most of the isolates had multi-antibiotic resistant phenotypes. To our knowledge, the present report is the first to identify W. chitiniclastica and Comamonas sp. as causes of septicaemia in soft-shelled turtles and the first to identify Aeromonas spp. with $bla_{CTX-M-14}$ and bla_{DHA-1} resistance genes.

Keywords: pet; ulcer disease; liver; lesions; septicaemia; antibiotic resistance

The Chinese soft-shelled turtle (*Pelodiscus sinensis*) is a reptile that lives in fresh water. In addition to its popularity as an indoor pet, the species is a traditional nutrient-rich food in Asian countries (Feng et al. 1996; Yin et al. 2005), which has led to commercial aquaculture of these animals in land-based tanks or ponds in various countries including China and Japan. In Korea, they are farmed to a lesser extent than other aquatic animals. Over the past two decades, the soft-shelled turtle has been one of the most intensively cultured fresh-water animals. However, infectious diseases that cause problems

such as skin ulcers, shell necrosis, and septicaemia continue to threaten soft-shelled turtle farms.

In turtles, septicaemia manifests as various clinical symptoms including anorexia, lethargy, shell necrosis, and skin ulcers. Another significant postmortem observation is liver necrosis (Kohler 2006). Citrobacter freundii is a major pathogen responsible for septicaemic cutaneous ulcerative disease in turtles (Kohler 2006). In China, Aeromonas spp., C. freundii, and Edwardsiella tarda have frequently been reported as aetiological agents of septicaemia in the Chinese soft-shelled turtle (Hu

Supported by the Fishery Commercialization Technology Development Program (Project No. 20160055), Ministry of Oceans and Fisheries and the Cooperative Research Program for Agriculture Science and Technology Development (Project No. PJ0119802), Rural Development Administration, Republic of Korea.

¹Department of Animal Science, Joongbu University, Chungnam, Republic of Korea

²Bio-safety Research Institute and College of Veterinary Medicine, Chonbuk National University, Jeonju, Republic of Korea

^{*}Corresponding author: shingw@chonbuk.ac.kr

[†]These authors contributed equally to this work, share co-first authorship

et al. 2010; Chen et al. 2013a; Chen et al. 2013b). Chryseobacterium spp. and Morganella morganii can also be isolated from turtles with shell disease, if only rarely (Hernandez-Divers et al. 2009). In the absence of vaccines against specific bacteria, antibiotic treatment is the best way to control these pathogens in aquaculture. Most previous studies on the antimicrobial susceptibilities of turtle isolates have focused on the bacterial flora of wild and captive turtles as bio-indicators for polluted effluents (Al-Bahry et al. 2009; Foti et al. 2009; Al-Bahry et al. 2012; Wheeler et al. 2012). In addition, Salmonella spp. in pet turtles have been demonstrated to be a risk factor for human salmonellosis (Seepersadsingh and Adesiyun 2003; Diaz et al. 2006; Chen et al. 2010; Guerra et al. 2010). To date, the prevalence of antibiotic resistance genes has been examined in only a few studies in pathogenic bacteria isolated from farmed soft-shelled turtles.

To our knowledge, no studies have focused on the aetiological agents responsible for the occurrence of infectious bacterial diseases in commercial Chinese soft-shelled turtles farmed in Korea. Furthermore, little information has been obtained with regard to antibiotic resistance and its genetic determinants in this reptile. Data on the antibiotic resistance of pathogens are of crucial importance for efforts directed towards the prevention and control of infectious diseases in soft-shelled-turtle farms. Therefore, the aims of the present study were to identify the bacterial pathogens associated with septicaemia in the soft-shelled turtle and to characterise the antibiotic resistance of these pathogens. Furthermore, we investigated the prevalence of the following antibiotic resistance determinants: β-lactamase encoding genes, tet, plasmid-mediated qnr and the class 1 integron cassette.

MATERIAL AND METHODS

Isolation. Bacteria were isolated from the liver lesions of 23 farm-raised Chinese soft-shelled turtles (*P. sinensis*) with various clinical symptoms (lethargy, slow growth, and skin and shell ulceration) in Korea. The bacterial isolations were performed by streaking liver samples onto tryptic soy agar (TSA) agar plates followed by incubation overnight at 27 °C. The isolated colonies were then subcultured on TSA plates to obtain pure colonies. A total of 23 strains were obtained and stored at -70 °C

in CryoCare Bacteria Preservers (Key Scientific Products, Stampford, USA) until required for further laboratory procedures.

Identification. The stored bacteria were incubated in tryptic soy broth overnight at 27 °C. The bacterial suspensions were processed using an AccuPrep® genomic DNA extraction kit (Bioneer, Daejeon, Korea) for the purification of bacterial DNA. The concentration of purified DNA was determined with the aid of an Epoch spectrophotometer system (Biotek, USA). Bacterial identifications were performed using standard primer sets (27F: 5'-AGAGTTTGATCMTGGCTCAG-3', 1492R: 5'-TACGGYTACCTTGTTACGACTT-3') for 16S rDNA sequencing (Macrogen Service Center, Daejeon, Korea). Partial gyrB sequencing was conducted to verify 11 Aeromonas strains to the species level, according to our previous study (Yi et al. 2013) using the same primer sets listed in Table 1.

Disc-diffusion assays. The antimicrobial susceptibility profile for each of the strains was investigated using the disc-diffusion method with 16 different antimicrobial agents: amikacin (AN, 30 mg), gentamicin (GM, 10 mg), tobramycin (NN, 10 mg), enrofloxacin (ENR, 5 mg), norfloxacin (NOR, 10 mg), oxolinic acid (OA, 2 mg), sulfamethoxazole/trimethoprim (Sxt, 300 mg + 5 mg), tetracycline (25 mg), amoxicillin/clavulanic acid (AMC, 30 mg), ampicillin (AM, 10 mg), piperacillin (PIP, 100 IU), cefotaxime (CTX, 30 mg), cephalothin (CF, 30 mg), cefaclor (CEC, 5 mg), imipenem (IMP, 10 mg), and chloramphenicol (C, 30 mg). The strains were recovered from the freezer with growth on TSA and then tested on Muller-Hinton agar plates. Various antimicrobial discs were then applied to the cultures. The inhibition zones were measured after incubation for 18 h at 27 °C. The resistance of the strains to the antimicrobials was determined according to the manufacturer's instructions and the M100-S17 document of the Clinical and Laboratory Standards Institute.

Detection of antibiotic resistance genes. All of the strains were tested by PCR assays to detect the genetic determinants associated with resistance to tetracycline, quinolones, and β-lactams, in addition to the class 1 integron gene cassette encoding resistance to various antimicrobials. The primer sets and PCR conditions are summarised in Table 1. The assays were carried out in 20 ml AccuPower[®] PCR premix (Bioneer) containing 1 ml of each forward and reverse primer (10mM) and 1 ml of bacterial genomic

Table 1. PCR primers used in this study

PCR	Target	Primer pair	Sequence (5'-3')	Annealing temperature (°C)	Amplicon size (bp)	Reference	
S	gyrB	gyrB-3F gyrB-14R	TCCGGCGGTCTGCACGGCGT TTGTCCGGGTTGTACTCGTC	1100	Martinez-Murcia et al. (2011)		
S	class 1 integron	5'-CS 3'-CS	GGCATCCAAGCAGCAAG AAGCAGACTTGACCTGA	64	VR	Lee et al. (2008)	
M-1	bla _{CTX-M-1} group	CTXGp1-F CTXGp1-R	TTAGGAARTGTGCCGCTGYA CGATATCGTTGGTGGTRCCAT	68	688		
	bla _{CTX-M-2} group	CTXGp2-F CTXGp2-R			404	Dallenne et al. (2010)	
	bla _{CTX-M-9} group	CTXGp9-F CTXGp9-R	TCAAGCCTGCCGATCTGGT TGATTCTCGCCGCTGAAG	68	561		
M-2	bla_{TEM}	TEM-F TEM-R	CATTTCCGTGTCGCCCTTATTC CGTTCATCCATAGTTGCCTGAC	60	800		
	$bla_{{SHV}}$	SHV-F SHV-R	AGCCGCTTGAGCAAATTAAAC ATCCCGCAGATAAATCACCAC	60	713	Dallenne et al. (2010)	
	$bla_{_{OXA ext{-}A ext{-}like}}$ variants	OXAA-F OXAA-R	GGCACCAGATTCAACTTTCAAG GACCCCAAGTTTCCTGTAAGTG	60	564		
M-3	qnrA	qnrA-F qnrA-R	AGAGGATTTCTCACGCCAGG TGCCAGGCACAGATCTTGAC	60	580		
	qnrB	qnrB-F qnrB-R	GGAATCGAAATTCGCCACTG TTTGCCGTCCGCCAGTCGAA	60	264	Cattoir et al. (2007)	
	qnrS	qnrS-F qnrS-R	GCAAGTTCATTGAACAGGGT TCTAAACCGTCGAGTTCGGCG	60	428		
M-4	tetA	tetA-F tetA-R	GTAATTCTGAGCACTGTCGC CTGCCTGGACAACATTGCTT	62	1000		
	tetE	tetE-F tetE-R	GTGATGATGGCACTGGTCAT CTCTGCTGTACATCGCTCTT	62	1200		
M-5	tetB	tetB-F tetB-R	CTCAGTATTCCAAGCCTTTG CTAAGCACTTGTCTCCTGTT	57	400	Akinbowale et al.	
	tetD	tetD-F tetD-R	ATTACACTGCTGGACGCGAT CTGATCAGCAGACAGATTGC	57	1100	(2007)	
S	tetC	tetC-F tetC-R	TCTAACAATGCGCTCATCGT GGTTGAAGGCTCTCAAGGGC	62	588		
S	tetM	tetM-F tetM-R	GTTAAATAGTGTTCTTGGAG CTAAGATATGGCTCTAACAA	55	600		

 $S = single \ PCR$, $M = multiplex \ PCR$, VR = variable

DNA (30–40 ng). The amplicons for each gene were resolved by electrophoresis on 1.5% agarose/TBE gels including RedSafe (iNtRON Biotechnology) and visualized under UV light. The class 1 integron amplicons were purified using the AccuPower® gel purification kit (Bioneer) and sequenced directly using 5'-CS and 3'-CS primers (Macrogen Service Center, Korea). The class 1 integron gene cassette homology searches were performed using the Basic Local

Alignment Search Tool (BLAST) from the National Center for Biotechnology Information (NCBI) website (www.ncbi.nlm.nih.gov/BLAST).

RESULTS

Twenty-three isolates were independently recovered from the livers of diseased Chinese soft-

shelled turtles with skin/shell ulcers. Based on 16S rDNA sequences, the 23 isolates were divided into eight different species belonging to six families: $A.\ hydrophila\ (n=8),\ A.\ veronii\ (n=3),\ M.\ morganii\ (n=3),\ C.\ freundii\ (n=4),\ Wohlfahrtiimonas\ chitiniclastica\ (n=1),\ Chryseobacterium\ spp.\ (n=1),\ Comamonas\ spp.\ (n=1),\ and\ E.\ tarda\ (n=2)\ (Table\ 2).$ There are many reports of strong similarities between the 16S rDNA sequences of $Aeromonas\ species\ (Yanez\ et\ al.\ 2003);$ therefore, phylogenetic analysis using partial gyrB sequences was applied to verify the species-level identification of the $11\ Aeromonas\ isolates$. The results corresponded to those obtained using the $16S\ rDNA\ method$ for those same isolates.

The antibiotic resistance of the isolates was determined using the disc diffusion test (Table 2).

Resistance phenotypes of all the isolates to 16 different antibiotics and determinants are summarised in Table 3. A high percentage of the isolates were resistant to ampicillin (95.7%), cefaclor (78.3%) and cephalothin (73.9%), whereas a low percentage was resistant to aminoglycosides (13.1%). The trend of resistance according to major bacterial family was found as follows: AM (100%) > CF = CEC = TE (72.7%) > IPM = PIP (63.6%) > OA = Sxt (27.3%) >AMC = C = CTX (18.2%) for the Aeromonasceae and AM (88.9%) > CEC (77.8%) > CF = TE = AMC (66.7%) > PIP = C = CTX (55.6%) > ENR = NOR =OA = Sxt (44.4%) > IPM (22.2%) > GM = NN (11.1%)for the Enterobacteriaceae. Twenty-one of the 23 isolates (91.3%) exhibited greater than intermediate resistance to three or more antimicrobial agents (the exceptions were two *E. tarda* isolates).

Table 2. Antibiotic resistance of the isolates

T1-4-	AM	AMC	CEC	CF	CTX	PIP	IPM	AN	GM	NN	ENR	NOR	OA	С	Sxt	TE
Isolate	DI															
SST001	0 R	18 S	10 R	0 R	29 S	21 S	9 R	24 S	20 S	20 S	30 S	26 S	26 S	30 S	26 S	26 S
SST002	0 R	21 S	17 I	18 S	29 S	25 S	19 S	27 S	20 S	18 S	25 S	20 S	0 R	19 S	0 R	21 S
SST003	0 R	12 R	0 R	30 S	12 R	0 R	0 R	22 S	18 S	17 S	0 R	12 R	0 R	0 R	0 R	0 R
SST004	0 R	19 S	20 S	21 S	33 S	20 I	0 R	22 S	21 S	20 S	24 S	21 S	0 R	27 S	26 S	0 R
SST005	0 R	21 S	0 R	0 R	22 I	17 R	14 I	24 S	21 S	22 S	29 S	26 S	23 S	26 S	26 S	11 R
SST006	0 R	19 S	12 R	12 R	27 S	20 I	12 R	22 S	22 S	23 S	26 S	26 S	24 S	27 S	25 S	10 R
SST007	0 R	15 I	0 R	0 R	27 S	25 S	20 S	21 S	16 S	20 S	23 S	25 S	20 S	0 R	25 S	20 S
SST008	0 R	20 S	12 R	11 R	29 S	21 S	16 I	23 S	21 S	23 S	23 S	24 S	22 S	30 S	26 S	12 R
SST009	0 R	12 R	0 R	0 R	22 I	18 I	19 S	17 S	17 S	16 S	0 R	12 R	0 R	19 S	0 R	0 R
SST010	0 R	13 R	0 R	0 R	22 I	19 I	19 S	20 S	20 S	20 S	29 S	26 S	24 S	17 I	25 S	0 R
SST011	0 R	13 R	0 R	0 R	17 I	17 R	24 S	21 S	15 S	18 S	0 R	10 R	0 R	0 R	0 R	0 R
SST012	0 R	16 I	0 R	0 R	0 R	13 R	15 I	16 I	15 S	18 S	22 S	25 S	19 S	28 S	22 S	25 S
SST013	0 R	19 S	10 R	0 R	26 S	19 I	11 R	22 S	21 S	21 S	24 S	24 S	21 S	24 S	27 S	10 R
SST014	0 R	20 S	18 S	14 R	30 S	19 I	18 S	20 S	22 S	19 S	24 S	25 S	25 S	26 S	22 S	25 S
SST015	0 R	19 S	10 R	0 R	26 S	21 S	15 I	24 S	20 S	18 S	30 S	27 S	24 S	25 S	0 R	15 R
SST016	0 R	18 S	20 S	22 S	28 S	25 S	12 R	22 S	19 S	20 S	31 S	29 S	31 S	27 S	32 S	11 R
SST017	0 R	12 R	0 R	11 R	23 S	10 R	22 S	20 S	15 S	16 S	0 R	12 R	0 R	0 R	0 R	10 R
SST018	0 R	16 I	0 R	0 R	0 R	14 R	17 S	11 R	0 S	0 S	33 S	22 S	28 S	10 R	26 S	18 I
SST019	0 R	18 S	13 R	0 R	0 R	16 R	16 I	11 R	0 S	0 S	24 S	12 R	22 S	12 R	24 S	11 R
SST020	0 R	23 S	0 R	0 R	19 I	21 S	25 S	20 S	16 S	16 S	29 S	23 S	20 S	28 S	29 S	26 S
SST021	21 S	25 S	24 S	22 S	28 S	22 S	25 S	23 S	16 S	15 I	31 S	28 S	29 S	24 S	27 S	25 S
SST022	16 I	22 S	25 S	20 S	31 S	18 S	22 S	18 S	12 R	16 S	28 S	27 S	27 S	27 S	27 S	21 S
SST023	0 R	11 R	0 R	0 R	21 I	13 R	24 S	23 S	16 S	20 S	0 R	12 R	0 R	9 R	0 R	10 R

AM = ampicillin, AMC = amoxicillin/clavulanic acid, AN = amikacin, C = chloramphenicol, CEC = cefaclor, CF = cephalothin, CTX = cefotaxime, D = diameter (mm), ENR = enrofloxacin, GM = gentamicin, I = intermediate, IMP = imipenem, NN = tobramycin, NOR = norfloxacin, OA = oxolinic acid, PIP = piperacillin, R = resistance, S = susceptible, Sxt = sulfamethoxazole/trimethoprim, TE = tetracycline

Table 3. Characterisation of the 23 strains isolated from Chinese soft-shelled turtles according to resistance phenotypes and genetic determinants, including class 1 integron gene cassettes

Identification Isolates		Resistance phenotype	Resistance genes	Class 1 integron gene cassette	
A. hydrophila	SST001	AM, IPM, CF, CEC	qnrS2, bla _{TEM-171}		
	SST002	OA, Sxt, AM, CEC(I)	qnrS2, bla _{TEM-171} , tetC	dfrA12-orfF-aadA catB3-aadA1	
	SST005	TE, AM, IPM, PIP, CTX(I), CF,CEC	qnrS2, $bla_{TEM-171}$, $tetA$, $bla_{CTX-M-14}$		
	SST006	TE, AM, IPM, PIP(I), CF, CEC	qnrS2, bla _{TEM-171} , tetA		
	SST013	TE, AM, IPM, PIP(I), CF, CEC	qnrS2, bla _{TEM-171} , tetA		
	SST014	AM, PIP(I), CF	qnrS2, bla _{TEM-171}		
	SST015	Sxt, TE, AM, IPM(I), CF, CEC	qnrS2, bla _{TEM-171} , tetA		
	SST018	AN, TE(I), AmC(I), AM, PIP, C, CTX, CF, CEC	qnrS2, bla _{TEM-171}		
A. veronii	SST004	OA, TE, AM, IPM, PIP(I)	qnrS2, bla _{TEM-171} , tetA		
	SST016	TE, AM, IPM	qnrS2, $bla_{TEM-171}$		
	SST017	ENR, NOR, OA, Sxt, TE, AmC, AM, PIP, C, CF, CEC	qnrS2, bla_{TEM-2} , bla_{DHA-1} , $tetA$, $tetB$		
C. freundii	SST007	AmC(I), AM, C, CF, CEC	qnrS2, bla $_{\scriptscriptstyle TEM-171}$		
	SST008	TE, AM, IPM(I), CF, CEC	qnrS2, bla _{TEM-2} , tetD, bla _{CMY-2}		
	SST009	ENR, NOR, OA, Sxt, TE, AmC, AM, PIP(I), CTX(I), CF, CEC	qnrS2, qnrB14, bla $_{TEM-171}$, bla $_{CMY-2}$, tet A	dfrA1-aadA1	
	SST010	TE, AmC, AM, PIP(I), C(I), CTX(I), CF, CEC	qnrS2, qnrB1, $bla_{TEM-171}$, $bla_{CTX-M-14}$, $tetB$		
M. morganii	SST003	ENR, NOR, OA, Sxt, TE, AmC, AM, IPM, PIP, C, CTX, CEC	qnrS2, bla $_{TEM-171}$, tetB, bla $_{DHA-1}$	bla _{PSE-1} -aadA2, aadB-catB3	
	SST011	ENR, NOR, OA, Sxt, TE, AmC, AM, PIP, C, CTX(I), CF, CEC	qnrS2, bla _{TEM-171} , bla _{DHA-1} tetB	bla _{PSE-1} -aadA2, aadB-catB3	
	SST023	ENR, NOR, OA, Sxt, TE, AmC, AM, PIP, C, CTX, CF, CEC	qnrS2, bla _{TEM-171} , bla _{DHA-1} , tetB		
E. tarda	SST021	NN	qnrS2, bla _{TEM-171}		
	SST022	GM, AM(I)	qnrS2, bla _{TEM-171} , tetA		
Chryseobacterium sp.	SST019	AN, NOR, TE, AM, IPM, PIP, C, CTX, CF, CEC	qnrS2, bla _{TEM-171}		
Comamonas sp.	SST020	AM, CTX(I), CF, CEC	qnrS2, bla _{TEM-171}		
W. chitiniclastica	SST012	AN(I), AmC(I), AM, IPM(I), PIP, CTX, CF, CEC	qnrS2, bla _{TEM-171} , bla _{CTX-M-14}		

AmC = amoxicillin/clavulanic acid, AN = amikacin, AN = ampicillin, C = chloramphenicol, CEC = cefaclor, CF = cephalothin, CTX = cefotaxime, ENR = enrofloxacin, GM = gentamicin, I = interpretation according to CLSI M100-S17, IPM = imipenem, NN = tobramycin, NOR = norfloxacin, OA = oxolinic acid, PIP = piperacillin, Sxt = sulfamethoxazole/trimethoprim, TE = tetracycline

Multiple resistance to more than 10 antibiotics was observed in the *A. veronii* SST017, *C. freundii* SST009, *M. morganii* SST011, SST023, and SST003 and *Chryseobacterium* sp. SST019 isolates.

All isolates were subjected to PCR amplification and sequence analysis to identify the antimicrobial

resistance determinants β -lactamase (bla), plasmid-mediated quinolone resistance (qnr), tetracycline resistance (tet), and class 1 integron (intl1) genes. The $bla_{TEM-171}$ and bla_{TEM-2} genes were detected in 21 and two isolates, respectively. The 14 isolates carrying only the $bla_{TEM-171}$ gene varied in resistance

patterns to β-lactams. In addition, *E. tarda* isolates exhibited below intermediate resistance to ampicillin. On the other hand, nine of the isolates that harboured bla_{TEM} also carried $bla_{CTX-M-14}$ (A. hydrophila SST005, C. freundii SST010 and W. chitiniclastica SST012), bla_{DHA-1} (A. veronii SST017, M. morganii SST003, SST011 and SST023), or bla_{CMY-2} (C. freundii SST008 and SST009). The qnrS2 gene was detected in all isolates investigated in the present study, of which C. freundii SST009 and SST010 simultaneously carried qnrB14 and qnrB1, respectively. The tet genes were detected in 14 isolates and the genes detected included tetA (8/14), tetB (5/14), tetC (1/14), and tetD (1/14). In Aeromonas isolates, tetA (6/11) was the most common tet determinant whereas tetC was detected in only A. hydrophila SST002. In addition, the simultaneous detection of tetA and tetB was observed in only A. veronii SST017. In the case of enteric bacteria, all the isolates possessed one of three tet determinants, tetA, tetB or tetD. The class 1 integron gene cassette was detected in the following four isolates: dfrA12-orfF-aadA2 and catB3-aadA1 in A. hydrophila SST002, dfrA1-aadA1 in C. freundii SST009 and blaPSE-1-aadA2 and aadB-catB3 in M. morganii SST003 and SST011.

DISCUSSION

In the present study, most of the identified bacterial species have been reported as infectious agents responsible for septicaemia in chelonia (Oros et al. 2005; Hernandez-Divers et al. 2009; Hu et al. 2010). C. freundii is known to be the aetiological agent of septicaemic cutaneous ulcerative disease (SCUD), which is accompanied by liver necrosis (Kohler 2006). Aeromonas spp. have frequently been isolated from the liver or kidney of soft-shelled turtles with various clinical manifestations, such as soft shell and white abdominal shell (Chen et al. 2013a; Chen et al. 2013b). In addition, *M. morganii* and *C.* indologenes have been suggested to be infectious agents associated with shell necrosis in map turtles (Graptemys spp.; Hernandez-Divers et al. 2009). E. tarda and C. freundii have been isolated from clinical samples of soft-shelled turtles with fulminant septicaemia (Hu et al. 2010). In agreement with previous studies (Kohler 2006; Hernandez-Divers et al. 2009; Hu et al. 2010; Chen et al. 2013a; Chen et al. 2013b), we identified aeromonads and enteric bacteria as the major pathogens associated with septicaemia and skin/shell diseases in soft-shelled turtles. However, to the best of our knowledge, there is no information regarding infection with *W. chitiniclastica* and *Comamonas* spp. in chelonians. In addition, infections with both bacterial isolates have been reported only rarely in animals, including humans (Rebaudet et al. 2009; Almuzara et al. 2011; Nseir et al. 2011; Farshad et al. 2012). Moreover, *W. chitiniclastica* has yet to be recorded at all in Korea.

The literature data regarding the antimicrobial resistance patterns according to bacterial species and isolate sources are inconsistent (Foti et al. 2009; Chen et al. 2010; Al-Bahry et al. 2011; Al-Bahry et al. 2012; Aravena-Roman et al. 2012; Jang et al. 2013). In contrast to these previous reports, our major findings include higher resistance to cefotaxime and imipenem among enteric bacteria and aeromonads, respectively. These two antibiotics have not been approved for use in aquatic animals worldwide; therefore, resistance to these antibiotics has rarely been reported among bacterial isolates from aquatic animals, including turtles. The reason why the present isolates are highly resistant to both antibiotics remains to be established. Further studies are therefore warranted to examine whether the aforementioned antibiotic resistance is the result of contamination by polluted effluent.

Many studies have indicated that the aquatic environment is a vehicle for the spread of antibioticresistant bacteria and resistance genes (Jang et al. 2013; Marti et al. 2014). The present data further support this idea, and show a high prevalence and diversity of genetic determinants among the present isolates recovered from freshwater turtle aquaculture. The 14 isolates described here that harboured only the $bla_{\mathit{TEM-171}}$ gene varied in resistance patterns to β -lactams. These results might be due to varying levels of enzyme produced in different bacterial species and/or isolates, and indeed previous findings demonstrated that the degree of resistance to β -lactams depends on the amount of TEM or SHV enzyme produced among isolates (Wu et al. 1994; Livermore 1995). In addition, a high prevalence of the $bla_{\mathit{TEM-171}}$ gene was reported in only one previous study with *E. coli* isolates collected from a river in Korea (Jang et al. 2013). Although determining the prevalence of the bla_{TEM} gene requires further study, this gene might be widely disseminated in natural Korean environments. The genes encoding

β-lactamases that hydrolyse clinically important third-generation cephalosporins such as cefotaxime are widespread in clinical and environmental Enterobacteriaceae isolates (Livermore 1995; Kim et al. 2005; Jacoby 2009; Jang et al. 2013). However, CTX-M-14- and DHA-1-encoding genes have not been reported previously in aeromonads. In addition, there is limited information about antibiotic resistance and its genetic background in *W. chitiniclastica*. To our knowledge, the present report is the first demonstrating that *A. hydrophila* and *W. chitiniclastica* harbour CTX-M-14-encoding genes and that *A. veronii* harbour a DHA-1 type AmpC-encoding gene.

In spite of the presence of a variety of genetic determinants, the present study revealed some discrepancies between antibiotic resistance phenotypes and genetic determinants among the isolates; for example, two isolates with ampC genes were susceptible to cefotaxime. Previous studies showed that induction of AmpC enzymes was dependent on regulatory genes (ampR and ampD) and relative ampC promoter strength (Barnaud et al. 1998; Jacoby 2009). Therefore, cefotaximesusceptible isolates carrying the *ampC* gene might lack regulatory genes and/or might have a weaker ampC promoter sequence. Thus, although the discrepancy could be caused by the presence of a variety of other genetic determinants, it might due to insufficient production of proteins or mutations in the promoter sequences of genes for antibiotic resistance (Davies and Davies 2010; Wellington et al. 2013; Lazar et al. 2014).

This study is the first to profile bacterial pathogens and genetic determinants of antibiotic resistance from septicaemic Chinese soft-shelled turtles with ulcer disease. Many Gram-negative pathogens may collectively be associated with septicaemia in these animals. A multi-antibiotic resistance phenotype was frequently observed among the present isolates and all isolates harboured two or more genetic antibiotic resistance determinants. This finding might be due to the wide dissemination of antibiotic-resistant bacteria and resistance genes in aquatic environments around turtle farms.

REFERENCES

Akinbowale OL, Peng H, Barton MD (2007): Diversity of tetracycline resistance genes in bacteria from aquaculture

sources in Australia. Journal of Applied Microbiology 103, 2016–2025.

Al-Bahry S, Mahmoud I, Elshafie A, Al-Harthy A, Al-Ghafri S, Al-Amri I, Alkindi A (2009): Bacterial flora and antibiotic resistance from eggs of green turtles Chelonia mydas: An indication of polluted effluents. Marine Pollution Bulletin 58, 720–725.

Al-Bahry SN, Mahmoud IY, Al-Zadjali M, Elshafie A, Al-Harthy A, Al-Alawi W (2011): Antibiotic resistant bacteria as bio-indicator of polluted effluent in the green turtles, Chelonia mydas in Oman. Marine Environmental Research 71, 139–144.

Al-Bahry SN, Al-Zadjali MA, Mahmoud IY, Elshafie AE (2012): Biomonitoring marine habitats in reference to antibiotic resistant bacteria and ampicillin resistance determinants from oviductal fluid of the nesting green sea turtle, Chelonia mydas. Chemosphere 87, 1308–1315.

Almuzara MN, Palombarani S, Tuduri A, Figueroa S, Gianecini A, Sabater L, Ramirez MS, Vay CA (2011): First case of fulminant sepsis due to Wohlfahrtiimonas chitiniclastica. Journal of Clinical Microbiology 49, 2333–2335.

Aravena-Roman M, Inglis TJJ, Henderson B, Riley TV, Chang BJ (2012): Antimicrobial susceptibilities of Aeromonas strains isolated from clinical and environmental sources to 26 antimicrobial agents. Antimicrobial Agents Chemotherapy 56, 1110–1112.

Barnaud G, Arlet G, Verdet C, Gaillot O, Lagrange PH, Philippon A (1998): Salmonella enteritidis: AmpC plasmid-mediated inducible beta-lactamase (DHA-1) with an ampR gene from Morganella morganii. Antimicrobial Agents Chemotherapy 42, 2352–2358.

Cattoir V, Poirel L, Rotimi V, Soussy CJ, Nordmann P (2007): Multiplex PCR for detection of plasmid-mediated quinolone resistance qnr genes in ESBL-producing enterobacterial isolates. Journal of Antimicrobial Chemotherapy 60, 394–397.

Chen CY, Chen WC, Chin SC, Lai YH, Tung KC, Chiou CS, Hsu YM, Chang CC (2010): Prevalence and antimicrobial susceptibility of salmonellae isolates from reptiles in Taiwan. Journal of Veterinary Diagnostic Investigation 22, 44–50.

Chen JS, Ding XY, Zhu NY, Kong L, He ZY (2013a): Prevalence and antimicrobial susceptibility of Aeromonas species from diseased Chinese soft-shelled turtles (Trionyx sinens). Aquaculture Research 46, 1527–1536.

Chen JS, Zhu NY, Kong L, Bei YJ, Zheng TL, Ding XY, He ZY (2013b): First case of soft shell disease in Chinese soft-shelled turtle (Trionyx sinens) associated with Aeromonas sobria—A. veronii complex. Aquaculture 406, 62—67.

Dallenne C, Da Costa A, Decre D, Favier C, Arlet G (2010): Development of a set of multiplex PCR assays for the

- detection of genes encoding important β -lactamases in Enterobacteriaceae. Journal of Antimicrobial Chemotherapy 65, 490–495.
- Davies J, Davies D (2010): Origins and evolution of antibiotic resistance. Microbiology and Molecular Biology Reviews 74, 417–433.
- Diaz MA, Cooper RK, Cloeckaert A, Siebeling RJ (2006): Plasmid-mediated high-level gentamicin resistance among enteric bacteria isolated from pet turtles in Louisiana. Applied and Environmental Microbiology 72, 306–312.
- Farshad S, Norouzi F, Aminshahidi M, Heidari B, Alborzi A (2012): Two cases of bacteremia due to an unusual pathogen, Comamonas testosteroni in Iran and a review literature. Journal of Infection in Developing Countries 6, 521–525.
- Feng H, Yamazaki M, Matsuki N, Saito H (1996): Anti-tumor effects of orally administered soft-shelled turtle powder in mice. Biological and Pharmaceutical Bulletin 19, 367–368.
- Foti M, Giacopello C, Bottari T, Fisichella V, Rinaldo D, Mammina C (2009): Antibiotic resistance of Gram negatives isolates from loggerhead sea turtles (Caretta caretta) in the central Mediterranean Sea. Marine Pollution Bulletin 58, 1363–1366.
- Guerra B, Helmuth R, Thomas K, Beutlich J, Jahn S, Schroeter A (2010): Plasmid-mediated quinolone resistance determinants in Salmonella spp. isolates from reptiles in Germany. Journal of Antimicrobial Chemotherapy 65, 2043–2045.
- Hernandez-Divers SJ, Hensel P, Gladden J, Hernandez-Divers SM, Buhlmann KA, Hagen C, Sanchez S, Latimer KS, Ard M, Camus AC (2009): Investigation of shell disease in map turtles (Graptemys Spp.). Journal of Wildlife Diseases 45, 637–652.
- Hu G, Li D, Li T, Su X (2010): Isolation and identification of bacteria from soft-shelled turtle (Trionyx sinensis) associated with fulminant septicaemia. Journal of Fishery Sciences of China 17, 859–868.
- Jacoby GA (2009): AmpC beta-Lactamases. Clinical Microbiology Reviews 22, 161–182.
- Jang J, Suh YS, Di DYW, Unno T, Sadowsky MJ, Hur HG (2013): Pathogenic Escherichia coli strains producing extended-spectrum β -Lactamases in the Yeongsan river basin of South Korea. Environmental Science and Technology 47, 1128–1136.
- Kim J, Lim YM, Jeong YS, Seol SY (2005): Occurrence of CTX-M-3, CTX-M-15, CTX-M-14, and CTX-M-9 extended-spectrum beta-lactamases in Enterobacteriaceae clinical isolates in Korea. Antimicrobial Agents Chemotherapy 49, 1572–1575.
- Kohler G (ed.) (2006): Diseases of Amphibians and Reptiles. Krieger Publishing Co., Malabar. 184 pp.
- Lazar V, Nagy I, Spohn R, Csorgo B, Gyorkei A, Nyerges A, Horvath B, Voros A, Busa-Fekete R, Hrtyan M, Bogos B,

- Mehi O, Fekete G, Szappanos B, Kegl B, Papp B, Pal C (2014): Genome-wide analysis captures the determinants of the antibiotic cross-resistance interaction network. Nature Communications 5, 4352.
- Lee MF, Peng CF, Lin YH, Lin SR, Chen YH (2008): Molecular diversity of class 1 integrons in human isolates of Aeromonas spp. from southern Taiwan. Japanese Journal of Infectious Diseases 61, 343–349.
- Livermore DM (1995): β-Lactamases in laboratory and clinical resistance. Clinical Microbiology Reviews 8, 557–584.
- Marti E, Variatza E, Balcazar JL (2014): The role of aquatic ecosystems as reservoirs of antibiotic resistance. Trends in Microbiology 22, 36–41.
- Martinez-Murcia AJ, Monera A, Saavedra MJ, Oncina R, Lopez-Alvarez M, Lara E, Figueras MJ (2011): Multilocus phylogenetic analysis of the genus Aeromonas. Systematic and Applied Microbiology 34, 189–199.
- Nseir W, Khateeb J, Awawdeh M, Ghali M (2011): Catheterrelated bacteremia caused by Comamonas testosteroni in a hemodialysis patient. Hemodialysis International 15, 293–296.
- Oros J, Torrent A, Calabuig P, Deniz S (2005): Diseases and causes of mortality among sea turtles stranded in the Canary Islands, Spain (1998–2001). Diseases of Aquatic Organisms 63, 13–24.
- Rebaudet S, Genot S, Renvoise A, Fournier PE, Stein A (2009): Wohlfahrtiimonas chitiniclastica bacteremia in homeless woman. Emerging Infectious Diseases 15, 985–987.
- Seepersadsingh N, Adesiyun AA (2003): Prevalence and antimicrobial resistance of Salmonella spp. in pet mammals, reptiles, fish aquarium water, and birds in Trinidad. Journal of Veterinary Medicine B 50, 488–493.
- Wellington EM, Boxall AB, Cross P, Feil EJ, Gaze WH, Hawkey PM, Johnson-Rollings AS, Jones DL, Lee NM, Otten W, Thomas CM, Williams AP (2013): The role of the natural environment in the emergence of antibiotic resistance in Gram-negative bacteria. The Lancet Infectious Diseases 13, 155–165.
- Wheeler E, Hong PY, Bedon LC, Mackie RI (2012): Carriage of antibiotic-resistant enteric bacteria varies among sites in Galapagos reptiles. Journal of Wildlife Diseases 48, 56–67.
- Wu PJ, Shannon K, Phillips I (1994): Effect of hyperproduction of tem-1 beta-lactamase on in vitro susceptibility of Escherichia coli to beta-lactam antibiotics. Antimicrobial Agents Chemotherapy 38, 494–498.
- Yanez MA, Catalan V, Apraiz D, Figueras MJ, Martinez-Murcia AJ (2003): Phylogenetic analysis of members of the genus Aeromonas based on gyrB gene sequences. International of Journal of Systematic and Evolutionary Microbiology 53, 875–883.
- Yi SW, You MJ, Cho HS, Lee CS, Kwon JK, Shin GW (2013): Molecular characterization of Aeromonas species iso-

lated from farmed eels (Anguilla japonica). Veterinary Microbiology 164, 195–200.

Yin J, Tezuka Y, Shi L, Ueda JY, Matsushige K, Kadota S (2005): A combination of soft-shell turtle powder and essential oil of a unicellular chlorophyte prevents bone

loss and decreased bone strength in ovariectomized rats. Biological Pharmaceutical Bulletin 28, 275–279.

Received: April 8, 2016 Accepted after corrections: November 5, 2016