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ESBL-Producing *Escherichia coli* from Cows Suffering Mastitis in China Contain Clinical Class 1 Integrons with CTX-M Linked to ISCR1

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The prevalence of pathogenic multi-drug resistant (MDR) extended-spectrum β-lactamase (ESBL)-producing Escherichia coli is rapidly increasing, becoming a global concern. In a veterinary context, ESBL-producing E. coli are mostly reported in poultry and pigs. Here, we report on the prevalence and characterize ESBL-producing E. coli isolated from diverse dairy farms in China. Overall, 36 (23.53%) out of 153 E. coli isolates from mastitic milk samples (n = 1252) were confirmed as ESBL-producers by double-disc synergy testing and PCR. Nucleotide analysis of PCR amplicons revealed that blacTX-M was the predominant ESBL gene detected in 28 (77.78%) isolates, with bla_{CTX-M-15} being the major (78.57%) allele encoding for ESBLs. Also, 20 (55.56%) and 6 (16.67%) of the ESBL isolates were carrying *bla*_{TEM} and *bla*_{SHV} genes, respectively, in singlet or in combination. The majority of these isolates belonged to phylo-group A (69.44%) and D (16.67%). Strikingly, all these isolates were found to be MDR showing high resistance to cephalosporins including the fourth generation cefepime and common non β -lactams. Additionally, class 1 integrons (*intl* 1) were found in 30 (83.33%) isolates. Analysis of the class 1 integrons variable regions indicated that they were carrying up to five different gene cassettes conferring resistance to various drugs with a predominant combination of dfrA17-aadA5 genes in tandem, conferring resistance to aminoglycosides and trimethoprim. However, no ESBL encoding genes were found in the cassettes. Interestingly, 22 (66.11%) of the ESBL isolates were also carrying insertion sequence common region 1 (ISCR1) which was found to be associated with most of the CTX-M genes. Altogether, the current study reports on the high prevalence of ESBL-positive E. coli, particularly CTX-M-15, carrying clinical class 1 integrons and ISCR1 elements are likely indicative of their rapid and wider dissemination, posing threats to veterinary and public health. To the best of our knowledge, this is the first comprehensive study to report on the alarming high occurrence of ESBL-producing E. coli from mastitic cows in China.

Keywords: E. coli, ESBLs, CTX-M-15, integrons, gene cassettes, bovine mastitis

INTRODUCTION

Bovine mastitis, inflammation of the mammary gland, is the most prevalent and economically important disease of dairy animals (Halasa et al., 2007). Mastitis can be caused by a variety of bacterial pathogens, but Escherichia coli is one of the leading causes (Dahmen et al., 2013). Antimicrobial agents are used for therapeutic as well as preventive measures against bacterial infections including bovine mastitis in farm animals. Beta-lactams, such as ampicillin and amoxicillin, remain the first-line treatment in veterinary medicine but an increase in drug-resistance to these antibiotics has been observed. Therefore, extended-spectrum cephalosporins (ESC) such as ceftiofur have been approved in China for the treatment of animal diseases (MAO, 2010). Unfortunately, several recent studies have reported the increasing occurrence of highly resistant extendedspectrum β-lactamase (ESBL)-producing *Enterobacteriaceae*, mainly E. coli, isolated from food-producing animals from various countries including China (Rao et al., 2014; Xu et al., 2015; Seni et al., 2016).

Bacterial resistance to β -lactams, popular antibiotics due to their proven safety and efficiency, is increasing at an alarming rate. This resistance is mainly achieved through β-lactamases that can hydrolyse most β -lactam antibiotics including the third and fourth generation ESCs and monobactams (Bush and Jacoby, 2010). ESBLs are predominantly produced in gram negative bacteria, particularly in E. coli, and are considered a key mechanism conferring resistance to cephalosporins (Perez et al., 2007). Multi-drug resistance (MDR) has been commonly observed in most ESBL-producers and more alarmingly, co-resistance to other commonly used antibiotics like aminoglycosides, fluoroquinolones, tetracycline has been often reported (Chen et al., 2010; Timofte et al., 2014; Xu et al., 2015). This renders these organisms resistant to a wide range of antibiotics with limited therapeutic options. ESBL encoding genes have been categorized into three main types: bla_{CTX-M}, *bla*_{SHV}, and *bla*_{TEM}. The *bla*_{CTX-M} has been further categorized into five sub-groups (*bla*_{CTX-M-1}, *bla*_{CTX-M-2}, *bla*_{CTX-M-8}, bla_{CTX-M-9}, bla_{CTX-M-25}) and more than 150 variants have been documented (http://www.lahey.org/studies). In the past few years, CTX-M, especially CTX-M-15, has emerged as the most dominant type of ESBLs globally (D'Andrea et al., 2013). Recently, CTX-M-15 producing E. coli have been frequently documented from various sources including humans and food producing animals (Timofte et al., 2014; Liu et al., 2015; Xu et al., 2015), showing the broad spectrum of reservoirs carrying and spreading these genes. Food-animals are well established reservoirs of ESBL-producing E. coli, which can be transmitted from animals to humans by various direct and indirect means (Dahmen et al., 2013; Geser et al., 2015). This is also verified by Madec et al. (2012), they reported that the plasmids carrying CTX-M-15 genes in E. coli isolated from cattle were highly similar to those found in ESBL-producing E. coli isolates from human beings.

Integrons are genetic elements that play a vital role in the development and dissemination of MDR in clinical isolates due to their ability to capture, integrate and express gene cassettes (Vinue et al., 2008; Chen et al., 2010). Three main classes of integrons (1-3), carrying the gene cassettes encoding for antimicrobial resistance genes, are generally found to be associated with antibiotic resistance genes in pathogenic E. coli, Class 1 integrons are the most common in clinical E. coli, followed by less frequent class 2 integrons (Vinue et al., 2008; Xu et al., 2015). Class 1 integrons contain a 5' conserved segment (CS) and 3'CS, followed by a variable region that contains one or more gene cassettes. The 5'CS consists of an integrase gene (intI1), a recombination-site (attI1), and the Pc promoter(s), and the 3'CS includes $qacE\Delta 1$ and sull genes which encode for quaternary ammonium compound and sulphonamide resistance (Hall and Stokes, 1993). Moreover, insertion sequences like ISCR1 (insertion sequence common region 1) as part of the complex class 1 integrons are found to be associated with ESBL and other resistance encoding genes and are probably involved in their mobilization and transposition. ISCR1 may mobilize the truncated 3'CS and nearby sequences from one integron to the 3'CS of another integron utilizing rolling circle transposition, thus facilitating dissemination of resistance elements (Eckert et al., 2006; Toleman et al., 2006).

Limited studies, particularly from China, have characterized ESBL-producing *E. coli* isolated from diseased food producing animals, mainly from mastitic cows (Lu et al., 2010; Timofte et al., 2014). Thus, we designed the current study to investigate the prevalence of pathogenic ESBL-producing *E. coli* and to characterize the ESBL genes and genetic elements which are likely to be responsible for their mobility and dissemination. To the best of our knowledge, this is the first comprehensive study into the molecular characterization of ESBL genes in *E. coli* isolated from dairy cows in China.

MATERIALS AND METHODS

Statement of Ethics

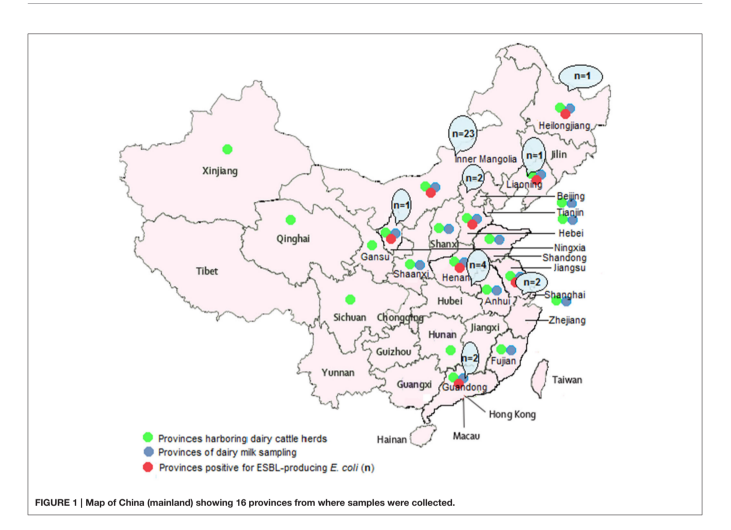
The present study was conducted in accordance with the ethical guide lines of China Agricultural University (CAU), Beijing. Proper ethical approval was granted by the departmental committee of College of Veterinary Medicine, CAU. Sampling was carried according to the standard protocols and with prior consent of the dairy herd's authority.

Sample Collection and Location

Milk samples of mastitic cows (n = 1252) were collected from 61 large commercial dairy herds (2000–40,000 cows/herd) located in 16 provinces of China during January 2015 to May 2016 (**Figure 1** and **Table 1**). Sampling was carried out when the cows were suffering from mastitis and not according to a fixed schedule. The guidelines of the National Mastitis Council (NMC, 1999) were followed for the collection of milk samples from cows. Samples were taken in 50 mL sterile tubes and transported on ice to the laboratory for further processing.

Isolation and Identification of *E. coli*

Milk samples, shortly after arrival, were streaked (10 μ L) onto MacConkey Agar (DifcoTM, Becton Dickinson, Sparks, MD USA)



and incubated at 37°C for 18–24 h. Presumptive *E. coli* colonies with the dark pink to red colors, were further confirmed with the API-20E kit (bioMérieux, Marcy l'Etoile, France) as per instruction of the manufacturer. Biochemically confirmed *E. coli* isolates were further verified by PCR as described previously (Tantawiwat et al., 2005). Confirmed *E. coli* isolates were stored in brain heart infusion broth (BHI; Sigma-Aldrich) containing 30% glycerol at -80° C.

Phenotypic Screening of ESBL-Producers

E. coli isolates were first screened for the phenotypic identification of ESBLs-producers on MacConkey agar containing cefotaxime (1 mg/L). These presumptive ESBL-producing *E. coli* were further confirmed by double-disc synergy testing in accordance with recommendations of the Clinical and Laboratory Standards Institute (CLSI, 2014), using antimicrobial discs (Becton Dickinson, Sparks, MD USA) of cefotaxime (30 μ g), cefotaxime plus clavulanic acid (30/10 μ g), ceftazidime (30 μ g), and ceftazidime plus clavulanic acid (30/10 μ g). The test was recorded positive when the zone of inhibition of cefotaxime plus clavulanic acid or ceftazidime plus clavulanic acid was \geq 5 mm larger than their respective single discs (CLSI, 2014).

Genotypic Screening of ESBL-Producing *E. coli* Isolates

Bacterial DNA from ESBL-positive *E. coli* was isolated by the TIANamp Bacteria DNA Kit (TIANGEN, Beijing, China) according to the manufacturer's instructions. PCR assays were used for the detection of bla_{CTX-M} , bla_{SHV} , bla_{TEM} genes as described previously (Chen et al., 2010). Details of the primers used in this study are shown in **Table 2**. All ESBL genes relevant PCR amplicons were purified by the TIANquick Midi Purification Kit (TIANGEN, Beijing, China), bi-directionally sequenced and aligned with sequences available in GenBank (Chen et al., 2010). *Klebsiella pneumoniae* ATCC 700603 (ESBLspositive strain) and ddH₂O, instead of template DNA, was used as positive and negative controls, respectively, in all PCR assays.

Phylogenetic Grouping

ESBL-positive *E. coli* isolates were placed in one of the four phylogenetic groups: phylo-group A, group B1, group B2 or group D. For this purpose, a triplex PCR assay targeting the *chuA* and *yjaA* genes and *TspE4* was used as described previously by Clermont et al. (2000). The primers sequences and the annealing temperatures are listed in **Table 2**.

Provinces of sampling	No. of dairy herds ^a	No. of milk samples ^b	<i>E. coli</i> isolates ^c	ESBL <i>E. coli</i> from each herd ^d
Anhui	А	63	4	0
Beijing	B/B1/B2	26 (6/9/11)	5 (2/2/1)	0/0/0
Fujian	F	12	0	0
Guangdong	G/G1	98 (23/75)	3 (1/2)	2 (0/2)
Hebei	Hb/ Hb1/ Hb2/ Hb3	220 (11/10/16/16	36 (2/0/0/5	2 (0/0/0/1
	/Hb4/Hb5/Hb6/Hb7 /Hb8/Hb9/Hb10	/6/11/12/18/23 /24/12/38/14/9)	/1/0/0/4/5 /3/3/6/6/1)	/0/0/0/0/1 /0/0/0/0/0)
	/Hb11/Hb12/Hb13			
Heilongjiang	H/H1/H2/H3	73 (10/13/10/40)	7 (4/2/0/1)	1 (1/0/0/0)
Henan	Hn/Hn1/Hn2/Hn3/Hn4	43 (12/6/7/12/6)	5 (2/0/0/2/1)	4 (2/0/0/2/0)
nner-Mongolia	1/11/12/13/14/15	425 (17/14/12/18	45 (1/2/0/1	23 (0/0/0/0
	/16/17/18/19/110	/42/37/49/22/24	/6/8/7/5/3	/5/7/6/3/0
	/111/12/113	/61/33/53/22/21)	/6/0/3/2/1)	/2/0/0/0/0)
Jiangsu	J	9	4	2
Liaoning	L	20	4	1
Ningxia	N/N1/N2/N3	97 (15/32/20/30)	19 (2/7/5/5)	1 (0/1/0/0)
Shaanxi	Sx-Bj	13	2	0
Shandong	S/S1/S2/S3	83 (14/14/15/40)	8 (0/1/5/2)	0/0/0/0
Shanxi	Sx-Cz	6	0	0
Shanghai	S/S1/S2/S3	59 (16/16/10/17)	10 (1/0/6/3)	0/0/0/0
Tianjin	Т	5	1	0
Total	61	1252	153 (12.22%)	36 (23.53%)

TABLE 1 | Occurrence of ESBL-producing E. coli isolated from dairy herds located in 16 provinces of China.

^a The letters in the column represent the farm number.

^bThe numbers in parenthesis indicate no. of milk samples corresponding to the respective farm in second column.

^cThe numbers in parenthesis show E. coli isolates from the respective farm.

^d The numbers in parenthesis indicate ESBL-producing E. coli isolated from the respective farm.

Antibiotic Susceptibility Testing

Antibiotic susceptibility of ESBL isolates was carried out on Mueller-Hinton agar (DifcoTM) against 16 different antibiotics discs (Becton Dickinson, Sparks, MD, USA), using the standard Kirby-Bauer disk diffusion method according to recommendations of the CLSI (2014). The panel of antimicrobial agents consisted of both β -lactam and non- β -lactam antibiotics as listed in **Table 3**. *E. coli* ATCC 25922 (ESBL-negative strain) and *K. pneumoniae* ATCC 700603 (ESBL-positive strain) were used as quality control strains (CLSI, 2014). The isolates were declared as multi-drug resistant (MDR) when found resistant to three or more categories of antimicrobial drugs.

Detection of Integrons, Gene Cassettes and ISCR1

A PCR assay was used to detect Class 1, 2, and 3 integrons in all ESBL-producing *E. coli* using integron-integrase gene specific primers, *int11*, *int12*, and *int13*, respectively (Dillon et al., 2005). Subsequently, the *int11* positive genotypes (n = 24) were determined by sequencing amplicons derived from PCR for the class 1 integron variable regions as described previously (White et al., 2000). The ISCR1 region was PCR amplified from ESBL-producing isolates using specific primers (**Table 2**). The sequenced amplicons of the ISCR1 elements were confirmed by BLAST analysis (see below). A combination of primers specific to the IS*CR1* elements (Kiiru et al., 2013) elements and consensus primers of ESBL genes were used to verify their association.

PCR-RFLP Genotyping of Class 1 Integron Variable Region Amplicons

A PCR-based restriction fragment length polymorphism (PCR-RFLP) assay was adopted to identify genetic variation in the amplified products using restriction enzyme *Hinf1* (Takara, Shiga Japan) as published previously (Gu et al., 2008). PCR-RFLP products with similar band profiles were regarded as the same genotypes carrying identical gene cassette(s).

Nucleotide Sequencing and Data Analysis

Regardless of the similar PCR-RFLP genotypes, all amplicons of gene cassettes, IS*CR1* elements, and ESBL genes were bidirectionally sequenced using ABI 3730 sequencer (Applied Biosystems, Foster City, CA, USA). PCR amplicons of >1.8 kb were further sequenced using primer walking based on the sequenced amplicons. The obtained sequences were subjected to BLAST homology searches in the INTEGRALL database (http:// integrall.bio.ua.pt). Other sequence analyses were compared with BLASTN software (http://www.ncbi.nlm.nih.gov/BLAST/). Clone Manager 7 (Sci-Ed software, Denver, USA) and ClustalW2 (http://www.ebi.ac.uk/Tools/msa/clustalw2/) were also used for detailed analysis such as alignments and open reading frames.

TABLE 2 | Details of primers used in this study.

Primers	Sequence (5' to 3')	Target gene	Annealing temperature	Amplicons size	References
β-lactamases					
CTX-MA CTX-MB	CGC TTT GCG ATG TGC AG ACC GCG ATA TCG TTG GT	bla _{CTX-M}	54°C	550-bp	Villegas et al., 2004
SHV-F SHV-R	GGG TTA TTC TTA TTT GTC GC TTA GCG TTG CCA GTG CTC	bla _{SHV}	58°C	567-bp	Chang et al., 2001
TEM-F TEM-R	ATA AAA TTC TTG AAG ACG AAA GAC AGT TAC CAA TGC TTA ATC	bla _{TEM}	56°C	1086-bp	Yao et al., 2007
INTEGRONS					
intl1-F intl1-R	CCT CCC GCA CGA TGA TC TCC ACG CAT CGT CAG GC	intl1	54°C	280-bp	Dillon et al., 2005
intl2-F intl2-R	AAA TCT TTA ACC CGC AAA CGC ATG TCT AAC AGT CCA TTT TTA AAT TCT A	intl2	54°C	439-bp	Dillon et al., 2005
intl3-F intl3-R	AGT GGG TGG CGA ATG AGT G TGT TCT TGT ATC GGC AGG TG	intl3	54°C	599-bp	Dillon et al., 2005
intl1-VR-F ntl1-VR-R	TCA TGG CTT GTT ATG ACT GT GTA GGG CTT ATT ATG CAC GC	intl1variable region	56°C	variable	White et al., 2000
E. coli-SPECIE					
UAL	TGG TAA TTA CCG ACG AAA ACG GC ACG CGT GGT TAC AGT CTT GCG	uidA	62°C	147-bp	Tantawiwat et al., 200
PHYLO-GROU	JPS				
ChuA-F ChuA-R	GAC GAA CCA ACG GTC AGG AT TGC CGC CAG TAC CAA AGA CA	ChuA	55°C	279-bp	Clermont et al., 2000
YjaA-F YjaA-R	TGA AGT GTC AGG AGA CGC TG ATG GAG AAT GCG TTC CTC AAC	YjaA	55°C	211-bp	Clermont et al., 2000
TspE4C2-F	GAG TAA TGT CGG GGC ATT CA	TspE4C2	55°C	152-bp	Clermont et al., 2000
TspE4C2-R ISCR1	CGC GCC AAC AAA GTA TTA CG				
ISCR1-F ISCR1-R	CGC CCA CTC AAA CAA ACG GAG GCT TTG GTG TAA CCG	ISCR1	55°C	469-bp	Kiiru et al., 2013

F, forward; R, reverse.

TABLE 3 | Antibiotic susceptibility profiles of ESBL-producing *E. coli* isolates (*n* = 36) from milk of mastitic cows.

Antimicrobial agents	Abbreviations	Conc.* (µg)	Susceptible (%)	Intermediate (%)	Resistance (%)
Ampicillin	AM	10	11.11 (04/36)	02.78 (01/36)	86.11 (31/36)
Amoxicillin/clavulanic acid	AMX/CA	20/10	25.00 (9/36)	11.11 (04/36)	63.89 (23/36)
Cefalexin	CX	30	00.00 (00/36)	00.00 (00/36)	100 (36/36)
Cefaclor	CEC	30	05.56 (02/36)	00.00 (00/36)	94.44 (34/36)
Cefoxatin	FOX	30	83.34 (30/36)	08.33 (03/36)	8.33 (3/36)
Cefotaxime	CTX	30	00.00 (00/36)	00.00 (00/36)	100.0 (36/36)
Ceftazidime	CAZ	30	33.33 (12/36)	00.00 (00/36)	66.67 (24/36)
Cefepime	FEP	30	41.67 (15/36)	11.11 (04/36)	47.22 (17/36)
Aztreonam	AZT	30	13.89 (05/36)	00.00 (00/36)	86.11 (31/36)
Meropenem	MPN	10	100.0 (36/36)	00.00 (00/36)	00.00 (00/36)
Tetracycline	TE	30	16.67 (06/36)	11.11 (04/36)	72.22 (26/36)
Gentamicin	G	10	27.78 (10/36)	11.11 (04/36)	61.11 (22/36)
Ciprofloxacin	CIP	05	55.56 (20/36)	08.33 (03/36)	36.11 (13/36)
Chloramphenicol	С	30	47.22 (17/36)	11.11 (4/36)	41.67 (15/36)
Nalidixic acid	NAL	30	19.44 (07/36)	02.78 (01/36)	77.78 (28/36)
Trimethoprim/sulphamethoxazole	STX	1.25/23.75	25.00 (09/36)	02.78 (01/36)	72.22 (26/36)

*Conc: concentrations.

RESULTS

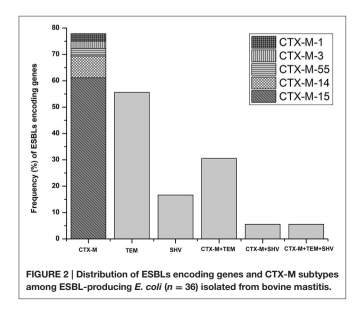
Prevalence and Characterization of ESBL-Producing *E. coli*

Overall, 153 *E. coli* isolates were recovered from 1252 milk samples of mastitic dairy cows from 16 different provinces of China. Thirty six (23.53%) isolates were detected as ESBL-producing *E. coli* by phenotypic confirmatory tests and this was also verified by ESBL genotype specific PCR assay. The distribution of these isolates among different cattle herds is shown in **Table 1**. The highest occurrence of ESBL producers was observed in the Inner Mongolia province (23 isolates), followed by the Henan region (four isolates).

Figure 2 shows the frequency (%) of various ESBL encoding genes among 36 ESBL-producing E. coli isolated from mastitic milk. Overall, bla_{CTX-M} was the most prevalent ESBL gene (77.78%; 28/36), while blaTEM and blaSHV genes were present in 55.56% (20/36) and 16.67% (6/36) of ESBL-positive isolates, respectively. The *bla*_{TEM} and *bla*_{SHV} genes were most frequently observed together with *bla*_{CTX-M}, rather than alone (Figure 2). Notably, two of the isolates from Inner Mongolia carried three β -lactamase genes ($bla_{CTX-M-15}+bla_{TEM-1}+bla_{SHV-12}$) in combination. Sequence analysis revealed that *bla*_{CTX-M-15} was the dominant (78.57%; 22/28) subtype. The other *bla_{CTX-M}* subtypes were: *bla_{CTX-M-14}* (10.71%; 3/28), *bla_{CTX-M-1}* (3.57%; 1/28), *bla*_{CTX-M-3}(3.57%; 1/28), and *bla*_{CTX-M-55} (3.57%; 1/28). The phylo-group A was the most prevalent (69.44%; 25/36) among 36-ESBL-positive E. coli followed by group D (16.67%; 6/36), B1 (8.33%; 3/36), and B2 (5.56%; 2/36) as depicted in Table 4.

Antibiotic Susceptibility Profiles

All 36 ESBLs-producing *E. coli* isolates were found to be multiple-drug resistant (MDR). However, different isolates exhibited slight variation in their antibiotic susceptibility profiles against the 16 tested antibiotics (**Table 3**). The majority of the isolates were resistant to first (cephalexin, 100%), second



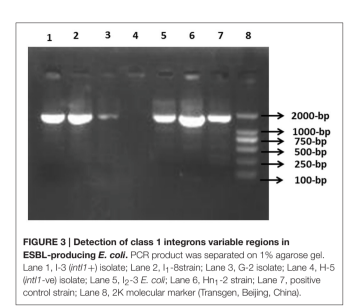
(cefaclor, 94.4%), third (cefotaxime and ceftazidime, 100% and 66.67%, respectively), and fourth (cefepime, 58.33%) generation cephalosporins. However, a high rate of susceptibility was observed toward cefamycin (cefoxatin, 83.34%) and carbapenem (meropenem, 100%), but susceptibility to monobactams (aztreonam, 13.89%) was low. The isolates were also resistant to other β -lactam and non- β -lactam antibiotics including ampicillin (88.89%), amoxicillin/clavulanic acid (75.00%), chloramphenicol (52.78%), ciprofloxacin (44.44%), gentamicin (72.22%), nalidixic acid (80.56%), tetracycline (83.33%) and trimethoprim/sulphamethoxazole (75%).

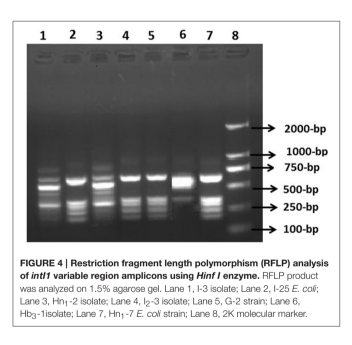
Detection of Integrons, Gene Cassettes and ISCR1

Thirty (83.33%) of the ESBL-producing *E. coli* carried clinical class 1 integrons but class 2 and class 3 integrons were not detected in any of the isolates. Among the *intI1*+ESBL-producing *E. coli*, 24 (80.00%) isolates tested positive for the presence of variable regions, while six of the isolates could not be amplified (**Table 4**). Furthermore, these 24 isolates were also positive for *qacE* Δ *1/sul1* indicating a complete clinical class 1 integron. Integrons lacking 3'CS were not PCR amplified for *qacE* Δ *1/sul1* (results not shown).

The PCR-amplicon sizes of the inserted gene cassettes ranged between ~ 1.0 and ~ 2.2 kb with the most predominant being \sim 1.7 kb amplicons (Figure 3). Most of the PCR amplicons of the variable regions of gene cassette arrays were a single band. However, two of the isolates produced a double band (of ~ 1.7 and \sim 0.2 kb). Subsequent sequence analysis of the gel extracted amplicons indicated that the smaller band was nonspecific amplification. Different band profiles of PCR-RFLP products indicated five distinct genotypic configurations (Figure 4). The most predominant PCR-RFLP genotype produced a profile of \sim 0.6, \sim 0.4, \sim 0.45, \sim 0.2, and \sim 0.22 kb restriction fragments consistent with digestion of 1.7 kb PCR amplicon of the variable regions. Amplicons sequence analysis of the variable regions revealed five gene cassettes carrying single or two genes in tandem. The predominant combination was dfrA17-aadA5 in tandem that conferred resistance to aminoglycosides and trimethoprim. Interestingly, all the CTX-15-positive isolates, except two, carried dfrA17-aadA5 genes in combination. This was consistent with the antibiotic susceptibility profile of these isolates reflecting resistance to the relevant drugs (Table 4). Surprisingly, no ESBL genes were found encoded in the variable region of the gene cassette array of these isolates. Therefore, ISCR1 elements were investigated by targeted-PCR. The PCR amplicons of ISCR1 elements were sequenced and confirmed by homology. Results indicated that ISCR1 was found in 22 (66.11%) ESBL positive isolates (Table 4). Moreover, ISCR1 (Accession number KY095113) was found associated with blaCTX-M, blaTEM and bla_{SHV} in 16, 3, and 4 isolates, respectively. Interestingly, all bla_{CTX-M-15} positive isolates, except one (Hn1-6), have been always found associated with ISCR1 elements. However, bla_{TEM}, when found alone or in combination with others, except *bla*_{CTX-M-15}, was mainly negative for ISCR1 elements. The amplicon size resulting from PCR using primer combinations

isolates	Place of isolation	groups			with <i>bla</i> genes	class 1	amplicons (bp)	5/CS -3/CS	numbers	non β -lactam antibiotics
<u>-</u> ع	Inner Mongolia	۵	TEM-1+SHV-1	+	+	+	2200	dfrA1-aacA4	KY114582	Cip; C; NAL; SXT; TE
-4	Inner Mongolia	A	CTX-M-15	+	+	+	1700	dfrA17-aadA5	KY114583	Cip; C; G; NAL; SXT; TE
-5	Inner Mongolia	Ω	CTX-M-15+TEM-1+SHV-1	I	I	+	2200	dfrA1 - aacA4	KY114584	Cip; G; NAL; SXT; TE
I-6	Inner Mongolia	A	TEM-1	+	+	+	1700	dfrA17-aadA5	KY114585	Cip; C; G; NAL; SXT; TE
I-12	Inner Mongolia	B1	CTX-M-15+TEM-1	+	+	I	I	I	I	Cip; G; NAL; SXT; TE
l-14	Inner Mongolia	A	CTX-M-15+ SHV-1	+	+	+	1700	dfrA17-aadA5	KY114586	Cip; C; G; NAL; SXT; TE
1-17	Inner Mongolia	∢	CTX-M-15	I	I	+	1700	dfrA17-aadA5	KY114587	Cip; G; NAL; SXT; TE
I-18	Inner Mongolia	A	CTX-M-15	+	+	I	I	I	I	Cip; G; NAL; SXT; TE
I-22	Inner Mongolia	Ω	CTX-M-15+SHV-1	+	+	+	I	I	I	Cip; G; NAL; SXT; TE
I-25	Inner Mongolia	A	CTX-M-15+TEM-1+SHV-1	I	I	+	1700	dfrA17-aadA5	KY114588	Cip; G; NAL; SXT; TE
1-1	Inner Mongolia	A	CTX-M-14	+	+	+	1800	dfrA17-aada4	KY114589	I
1-2	Inner Mongolia	A	TEM-1	+	+	+	I	I	I	G; TE
l ₁ -3	Inner Mongolia	A	TEM-1	+	I	Ι	I	I	I	Ι
l ₁ -4	Inner Mongolia	A	CTX-M-15	I	I	+	1700	dfrA17-aadA5	KY114590	Ι
l ₁ -5	Inner Mongolia	A	CTX-M-15	I	I	I	I	I	I	NAL
1 ₁ -7	Inner Mongolia	B1	CTX-M-55+TEM-1	I	I	+	1700	dfrA17-aadA5	KY114591	Cip; C; G; NAL; SXT; TE
l ₁ -8	Inner Mongolia	A	CTX-M-14	I	I	+	1800	dfra17-aada4	KY114592	C; G; NAL; SXT; TE
l ₁ -11	Inner Mongolia	B1	TEM-1	Ι	I	+	1700	dfrA17-aadA5	KY114593	C; G; NAL; SXT; TE
l2-1	Inner Mongolia	A	CTX-M-15+TEM-1	Ι	I	+	1700	dfrA17-aadA5	KY114594	Cip; C; G; NAL; SXT; TE
l ₂ -2	Inner Mongolia	A	TEM-1	I	I	+	1700	dfrA17-aadA5	KY114595	C; G; NAL; SXT; TE
l ₂ -3	Inner Mongolia	A	CTX-M-14+TEM-1	I	I	+	1800	dfra17-aada4	KY114596	C; G; NAL; SXT; TE
l ₃ -1	Inner Mongolia	B2	CTX-M-15	+	+	+	1700	dfrA17-aadA5	KY114597	C; G; NAL; SXT; TE
l ₃ -2	Inner Mongolia	B2	CTX-M-15	+	+	+	1700	dfrA17-aadA5	KY114598	C; G; NAL; SXT; TE
G-1	Guangdong	A	CTX-M-15	+	+	+	1700	dfrA17-aadA5	KY114599	Cip; C; NAL; SXT; TE
G-2	Guangdong	A	CTX-M-15+TEM-1	+	+	+	1700	dfrA17- aadA5	KY114600	C; NAL; SXT; TE
J-4	Jiangsu	D	CTX-M-15+TEM-1	+	+	+	Ι	Ι	I	C; G; NAL; SXT; TE
J-5	Jiangsu	Ω	CTX-M-15+TEM-1	+	+	+	I	I	Ι	C; G; NAL; SXT; TE
Hb ₂ -4	Hebei	Ω	TEM-1	I	I	+	2000	dfrA17-aadA5	KY114601	C; G; NAL; SXT; TE
Hb ₃ -1	Hebei	A	CTX-M-15	+	+	+	1000	aadA1	KY114602	Cip; C; G; NAL; SXT; TE
L-1	Liaoning	Þ	CTX-M-15+TEM-1	+	+	+	1700	dfrA17-aadA5	KY114603	C; G; NAL; SXT; TE
N-1	Ningxia	∢	CTX-M-3+TEM-1	I	Ι	+	1300	aadA5	KY114604	TE
H-5	Heilojinag	A	SHV-12	+	+	I	I	I	I	I
Hn-6	Henan	A	CTX-M-15	I	I	Ι	I	I	I	C; TE
Hn ₁ -2	Henan	A	CTX-M-1	+	+	+	1700	dfrA17-aadA5	KY114605	G; NAL
Hn ₁ -6	Henan	Þ	CTX-M-15+TEM-1	+	I	+	1700	dfrA17-aadA5	KY114606	G; NAL; SXT; TE
Hn ₁ -7	Henan	A	CTX-M-15+TEM-1	+	+	+	1700	dfrA17-aadA5	KY114607	G; NAL; SXT; TE





specific to ISCR1 elements and ESBL genes revealed that the ESBL genes were oriented downstream of ISCR1 elements. Altogether, these results indicated that ISCR1 elements are associated with ESBL genes. The detailed characterizations of all ESBLs-producing *E. coli* are elaborated in **Table 4**.

DISCUSSION

In the past few years, ESBL-producing *E. coli* have been increasingly isolated from food-producing animals raising global concerns for veterinary and public health (Seiffert et al., 2013). The current study reports on the higher occurrence (23.53%) of ESBL-producing *E. coli* (n = 36) among 153 *E. coli* isolates from mastitic cows, as compared to previous reports from China (Yu et al., 2015) and other countries (Dahmen et al., 2013; Geser

et al., 2015; Freitag et al., 2016). Interestingly, the majority of ESBL-producing E. coli (23 isolates) were recovered from herds in Inner Mongolia province possibly linked to dense farming (2.37 million dairy cows) with the largest dairy herds in this province (Dou, 2014). Dense farming is significantly correlated with the incidence of bovine mastitis (Ali et al., 2014), and mastitis is the main reason for frequent and prolonged use of antibiotics that exert selective pressure for emergence and dissemination of resistant isolates (Berge et al., 2005). Our results revealed that CTX-M, mainly the CTX-M-15, was the most prevalent genotype, followed by TEM and SHV. These findings agree with other contemporary studies from China and around the world that also reported CTX-M as the dominant ESBL genotype (Locatelli et al., 2009; Dahmen et al., 2013; Geser et al., 2015; Kar et al., 2015). This goes along with the recent detection of CTX-M-15 producing E. coli from cattle and other foodanimals in east Asia (Ohnishi et al., 2013; Yu et al., 2015), India (Upadhyay et al., 2015), the United Kingdom (Timofte et al., 2014), Germany (Freitag et al., 2016) and Tanzania (Seni et al., 2016). A national resistance surveillance study in China reported that the prevalence of ESBL-producing E. coli in humans has persisted above 50% since 2000 (Xiao et al., 2011), and recently Liu et al. reported even higher prevalence (68.2%) of ESBLs in clinical E. coli isolates, mainly the bla_{CTX-M-15} (Liu et al., 2015). Similarly, in animals the prevalence of ESBL-producing E. coli in China has considerably increased in recent years with CTX-M being the major prevailing gene encoding for ESBLs (Rao et al., 2014). It is known that, generally, ESBL genes are located on plasmids that could spread easily among commensal and pathogenic bacteria in the herd and the environment. Due to limited resources, in the present study, we could not investigate the prevalence of ESBL genes in other bacteria, in other healthy cows, or in cows with subclinical mastitis which is usually 30-40% higher than clinical mastitis (Halasa et al., 2007). We presume that the actual prevalence of ESBL-producers, particularly E. coli, may be much higher than the reported.

In ESBL-positive E. coli, phylogenetic group A represented the most prevalent group followed by virulent extra-intestinal group D; however, this was in contrast to our previous study (Liu et al., 2014), which reported that the pathogenic E. coli associated with mastitis mainly belonged to phylogroup B1 (58.6%) rather than group A (35.7%). Nevertheless, similar phylogenetic distributions were observed in some ESBLproducing E. coli isolated from animals (Abraham et al., 2014; Xu et al., 2015). Our results indicated that all the ESBLproducing E. coli isolates were MDR. The majority of these isolates (54-100%) were found resistant to cephalosporins. In addition, low susceptibility was also observed against the common β -lactam and non- β -lactam antibiotics such as ampicillin, aminoglycosides, tetracycline and fluoroquinolones. Recently, many studies have reported MDR ESBL-producing E. coli isolated from cattle (Timofte et al., 2014), poultry (Kar et al., 2015), pigs (Xu et al., 2015), and humans (Gu et al., 2008). Fluoroquinolones, following ciprofloxacin, are the second important antimicrobial drug in veterinary and human medicine (Coque et al., 2008). Quinolone resistance is traditionally caused by chromosomal mutations in gyrase or topoisomerase encoding

genes or efflux pump expression regulating genes (Hopkins et al., 2005); nonetheless, plasmid mediated quinolone resistance is also increasingly reported in ESBL-producing *E. coli* (Xu et al., 2015).

Integrons play an important role in the emergence of MDR bacteria and in the dissemination of resistance genes. Published reports on the characterization of integrons in ESBLs-positive E. coli from dairy cows are scarce, but previous studies have been conducted in other food-animals, humans and the environment (Gu et al., 2008; Chen et al., 2010; Xu et al., 2015). In accordance to these studies, clinical class 1 integrons were found in the majority of ESBL-positive E. coli (83.33%). The gene cassette arrays of the class 1 integron variable regions contained five different gene combinations that likely impart additional resistance features to our isolates (see Table 3). Six of the intI1 positive amplicons were failed to generate gene cassettes which may be related to the absence of 3'CS in these integrons (Lu et al., 2010). The dfrA17-aadA5 was the predominant gene array that corroborates with the previous studies in China (Gu et al., 2008; Xu et al., 2015). Strikingly, we determined that majority of bla_{CTX-M} genes were associated with ISCR1 elements, but no ESBL genes were found in the class 1 integron cassettes. It agrees with other published reports that also did not detect ESBL genes in the cassettes (Kiiru et al., 2013; Kar et al., 2015; Xu et al., 2015). Notably, our findings of the most predominant CTX-M type (bla_{CTXM-15}) and its association with the ISCR1 elements rather than gene cassette arrays indicated that they are more likely mobilized by ISCR1 elements. Conversely, *bla*_{TEM}, when found alone or not associated with *bla*_{CTXM-15}, was not often found linked to ISCR1, and therefore, was comparatively less prevalent. It has been proposed that antibiotic resistance gene elements are added to the 3'-CS of class 1 integrons by co-mobilization with the nearby ISCR1 from the neighbor integron, implying rolling circle transposition and homologous recombination mechanisms, thus facilitating the formation of complex class 1 integrons (Toleman et al., 2006). Taken together, the current high occurrence of multi-resistant ESBL-producing E. coli carrying clinical class 1 integrons and its association with ISCR1 is worrisome. This may suggest these bacteria are armouring against the antibiotics by devising various tools to render antibiotics useless. Fear exists that this co-existence of ESBL genes along with class 1 integrons as

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gene cassettes and *ISCR1* mobile elements would more robustly disseminate resistance elements within bacterial populations. This calls for an efficient control policy with restriction on the consumption of extended spectrum cephalosporins for long term use.

CONCLUSIONS

Here, we report on the high occurrence of ESBL-producing *E. coli* from bovine mastitis. Genotypic characterization indicated a dominance of $bla_{\text{CTX-M-15}}$ genes harboring clinical class 1 integrons associated with IS*CR1* elements, indicative rapid and wider dissemination potential and posing threats to veterinary and public health. To the best of our knowledge, this is the first comprehensive study to report on the alarming high prevalence of $bla_{\text{CTX-M-15}}$ and class 1 integron resistance conferring elements in ESBL-producing *E. coli* from mastitic cows in China.

AUTHOR CONTRIBUTIONS

BH, TA, and SR, conceived and designed the experiment. TA, MS, and SZ, performed the research. JG, GL, and LZ, contributed in reagents/materials/analysis. BH, TA, and SR, wrote the manuscript.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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