Molecular characterization of blaESBL-harboring conjugative plasmids identified in multi-drug resistant Escherichia coli isolated from food-producing animals and healthy humans

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INTRODUCTION

Global dissemination of multiple antibiotic resistance and virulence traits by plasmids poses an increasing threat to the successful treatment of bacterial infectious diseases in animals and humans alike (Bush, 2010). Extended-spectrum β-lactamase (ESBL)-producing Escherichia coli has become one of the most important causes of nosocomial and community acquired infections (Coque et al., 2008; Foxman, 2013) and most of the ESBL enzymes recognized are expressed from genes mapped to plasmids (Paterson and Bonomo, 2005; Garcellan-Barcia et al., 2011). It is acknowledged that the horizontal transfer of plasmids carrying these ESBL genes is an important contributory factor in the epidemiology of this bacterial ecosystem (EFSA Panel on Biological Hazards [BIOHAZ], 2011). The predominant ESBL families of clinical importance include TEM, SHV, and CTX-M (Bush and Jacoby, 2010); and the genes conferring this resistance phenotype in animals include blaTEM-1 (the most frequently identified ESBL), blaCTX-M-1, blaCTX-M-14, blaTEM-52, and blaSHV-12 (EFSA Panel on Biological Hazards [BIOHAZ], 2011). E. coli is the most common bacterial species identified with these genes.

Plasmids evolve as an integral part of the bacterial genome, consisting of several extra-chromosomal traits, one of which is their resistance genes, which can be exchanged among bacteria of different origins by conjugation (Carattoli, 2011). Those expressing an ESBL phenotype frequently carry genes encoding resistance to other commonly used antimicrobial drug classes (such as aminoglycosides, chloramphenicol, fluoroquinolones, or tetracycline; Haralaj et al., 2010; Rogers et al., 2011). Plasmid-mediated transfer of ESBL-encoding genes frequently map to plasmids, yet few of these structures have been characterized at the molecular level to date.

Methods: Eighty-seven ESBL-producing Escherichia coli were isolated from fecal samples of food-producing animals and healthy humans in Switzerland from 2009 to 2011. Plasmid DNA of all isolates was purified. Broth mating assays were carried out individually for 32 isolates to determine if the ESBL marker could be transferred by conjugation. The plasmid sizes were determined by S1-nuclease pulsed-field gel electrophoresis (PFGE) and the plasmids were typed by PCR-based replicon typing. Susceptibility tests by disk diffusion followed with a re-analysis S1-nuclease PFGE and PCRs were performed to confirm plasmid transfer. Microarray was performed to detect additional antibiotic resistance markers and multi-locus sequence typing was also performed in selected donor strains. The phylotypes were identified by triple PCR.

Results: About half (n = 46) of the 87 isolates carried small (<20-kb) plasmids. All selected 32 isolates contained large plasmids (ranging in sizes from 20- to 600-kb). Eleven plasmid replicon types were detected. Of these, IncFIA (n = 5), IncFIB (n = 9), and IncK/B (n = 4) were common. Nine isolates demonstrated the ability to transfer their cefotaxime resistance marker at high transfer rates. Plasmid profile re-analysis of these transconjugants identified 16 plasmids. IncFIB and IncI1 were the most prevalent replicon types. Phylogenetic grouping showed that five of the nine donor strains belonged to phylogroup B1. Nine different sequence types were identified in nine tested donor strains.

Conclusion: Characterization of these ESBL-encoding conjugative plasmids extends our understanding on these resistance markers in multi-drug resistant E. coli cultured from healthy human and animal sources.

Keywords: ESBLs, E. coli, horizontal gene transfer, replicon typing, conjugation, S1-nuclease PFGE, plasmid profiling, plasmids
of drug-resistance genes among various bacterial species is considered to be one of the most important mechanisms driving the spread of multi-drug resistance (MDR) markers. The use of antimicrobial compounds in human and veterinary medicine also constitutes a risk factor for the selection and dissemination of resistant clones, and also plasmids containing the corresponding resistance genes. Consequently, these features limit the treatment options available when ESBL-producing organisms are encountered. Therefore, as a first step to address this problem, it is important to monitor the dissemination of ESBL-producing bacteria, characterizing these microbes when they arise. Moreover, studies describing the genetic basis of plasmids derived from bacteria recovered from humans, animals, and the environment are essential.

Extended-spectrum-β-lactamase transmission is mainly driven by insertion sequences, transposons, integrons, and plasmids, some of which are homologous in isolates from both food-producing animals and humans (van Hoek et al., 2011; Stalder et al., 2012). Plasmid-encoded genes can arise from multiple sources (Boyd et al., 1996) being subsequently disseminated by horizontal gene transfer (HGT). HGT is responsible for the dissemination of many traits associated with bacteria, including antibiotic resistance and virulence. In addition, broad-host-range plasmids play an important role in bacterial adaptation to new environments. Taken together these characteristics provide much of the motivation of extending our understanding of the structural relationships that exist between plasmids from a variety of bacterial sources.

Following the dissemination routes of antibiotic resistance genes on ESBL–harboring plasmids is important to broaden our knowledge of how these genes are shared horizontally across species boundaries as well as their evolution (Zhou et al., 2012). Moreover, characterization based on plasmid profiling and the definition of corresponding incompatibility (Inc) groups is an integral part of plasmid epidemiological surveillance enhancing discrimination between E. coli strains (Carattoli et al., 2005).

In this paper, we report on the preliminary characterization of ESBL-producing E. coli isolates that were originally described following culture and phenotypic studies in healthy humans and animals in Switzerland. The aims of this study were (i) to identify the plasmids relating to ESBL markers, (ii) to characterize the donor E. coli isolates by phylogenetic grouping and multi-locus sequence typing (MLST), and (iii) to determine the replicon types and S1 nuclease-based plasmid profiles, following pulsed-field gel electrophoresis (PFGE). The epidemiological data presented extends our understanding of this resistance type in humans and animals.

MATERIALS AND METHODS

BACTERIAL STUDY STRAINS COLLECTIONS

Extended-spectrum-β-lactamase-producing E. coli (n = 87) were cultured from fecal samples of healthy humans (n = 34) and healthy food-producing animals (n = 52) in Switzerland, as well as one from a mastitis milk sample. To prevent sample clustering, at most two animal samples per farm were taken throughout Switzerland. All human samples were collected in urban areas, and each individual was tested only once. Prior to testing, all isolates were streaked for purity on MacConkey agar (Oxoid, Basingstoke, England). The protocols for collecting and isolating the strains were described previously (Geser et al., 2012a,b).

PLASMID PROFILING

Plasmid DNAs from all isolates were purified using the WizardR Plus SV DNA Purification System (Promega, Madison, WI, USA) according to the manufacturer’s instructions, followed by separation in a 0.8% (w/v) agarose gel (SeaKemR LE Agarose, Lonza Wokingham Ltd, UK) staining with GelRedTM (Biotium, Hayward, CA, USA). S1-nuclease (Promega, Madison, WI, USA) digestion as well as PFGE analysis were performed for selected S2 isolates based on their small plasmid subgroups and corresponding ESBL genotypes reported previously (the complete isolate information is listed in Figure A1 in Appendix). Briefly, the procedure included a lysis step of the bacterial cells embedded in agarose plugs followed by digestion with 8 U S1 nuclease at 37°C for 45 min. Finally, each plasmid sample was resolved by PFGE in a ChefMapper® XA System (Bio-Rad, USA) at 14°C, with a switch time between 1 and 12 s, at 6 V/cm on a 120° angle in 0.5 × TBE buffer for 18 h. Each DNA band was considered a unit length of linear plasmid (Barton et al., 1995). The approximate molecular mass of plasmids was determined by comparing with E. coli 39R 861, containing four reference plasmids, of known molecular weights 6.9-, 36-, 63-, and 147-kb (Macrina et al., 1978).

PCR-BASED REPLICON TYPING

Plasmids were assigned to incompatibility groups on the basis of the presence of specific replicon sequences identified by PCR using the primers previously designed and the corresponding amplification protocols described (Carattoli et al., 2005).

CONJUGATION-BASED MATING EXPERIMENTS AND VERIFICATION

Thirty-two selected isolates were analyzed individually, for their ability to transfer cefotaxime resistance to a rifampicin-resistant, plasmid-free E. coli recipient (26R 793). Conjugation experiments were carried out using a broth mating protocol. Transconjugants were selected on Luria-Bertani (LB) agar plates containing 30 μg/ml cefotaxime (Thermo Fisher Scientific Inc., USA) and 100 μg/ml rifampicin (Sigma, Dublin, Ireland). Transfer frequencies were calculated per donor cell. Susceptibility tests were performed to confirm the plasmid transfer as described below, followed by S1-nuclease PFGE and PCR-based replicon typing (PBRT) to test which plasmids and resistance markers were transferred.

ANTIMICROBIAL SUSCEPTIBILITY TESTING

Donor strains and the corresponding transconjugants were tested for their susceptibility to a panel of antimicrobial compounds by disk diffusion, following recommendations of the Clinical and Laboratory Standards Institute (CLSI). Susceptibility/resistance was interpreted according to the CLSI document M100-S21 (CLSI, 2011). The panel of antibiotic-containing disks (Becton, Dickinson, USA) along with their abbreviated names consisted of ampicillin (AM), amoxicillin–clavulanic acid (AMC), chloramphenicol (C), ciprofloxacin (CIP), cefpodoxime (CPD), cefotaxime (CTM), ceftazidime (CFT), and vancomycin (VAN).
cefotaxin (FOX), cephalothin (KF), tetracycline (TE), streptomycin (S), imipenem (IPM), nalidixic acid (NA), trimethoprim–sulfamethoxazole (SXT), and trimethoprim (W). E. coli ATCC™ 25922 was included as a quality control strain. The strains were classified as susceptible or resistant to each antimicrobial agent. Strains giving intermediate values were considered susceptible. In addition, minimal inhibitory concentrations (MICs) of cefotaxime for donors and transconjugants combinations were determined using E-test strips (AB Biodisk, Solna, Sweden).

DETECTION OF RESISTANCE GENES BY MICROARRAY (GENE CHIP)

A DNA microarray analysis (AMR-ve Genotyping Kit, Clontech Laboratories, Inc., Mountain View, CA, USA) was performed with the donor strains to detect the genes encoding resistance to aminoglycosides, β-lactams, chloramphenicol, erythromycin, quinolone, sulphonamides, tetracycline, trimethoprim, and an integrase-encoding gene (int).

PHYLLOGENETIC CLASSIFICATION

Donor isolates that transferred cefotaxime resistance to E. coli 26R 793 were classified into the four main phylogenetic groups (A, B1, B2, and D) using a previously described triplex PCR-based protocol (Clermont et al., 2000). Purified DNA served as a template. Thereafter, three specific primer sets (custom-synthesized by MWG-Biotech AG, Ebersberg, Germany) were used to identify the following markers, chuc (279 bp), yjcd (211 bp), and tcpB/F (152 bp) (Clermont et al., 2000).

MULTI-LOCUS SEQUENCE TYPING

Internal amplicons of seven housekeeping genes (adhE, fumC, gyrB, icdF, mdi, pucA, recA) from the donor strains were sequenced (Wirth et al., 2006); alleles as well as sequence types (ST) were verified using the E. coli MLST website.

RESULTS

PLASMID PROFILING

Eighty-seven E. coli isolates were included in this study and low molecular weight (<20-kb) plasmids could be identified by purification and conventional agarose gel electrophoresis. About half of the isolates (n = 46; 53%) carried several plasmids ranging in size from approximately 2- to 20-kb, and 41 isolates (47%) were devoid of small plasmids.

Thirty-two isolates from the original set of 87 were selected for further study, based on their diverse plasmid profiles and the corresponding ESBL genotypes, reported previously (Geser et al., 2012a,b). All were assessed for the presence of large plasmids (ranging in size from 30- to 600-kb) by S1 nuclease digestion followed by PFGE (Barton et al., 1995). S1 nuclease plasmid analysis revealed that all 32 contained detectable large plasmids; most possessed two plasmids (n = 13, 40.6%) and some (n = 9, 28.1%) had three plasmids. Heterogeneity among the profiles was a common feature noted, although most of the plasmid profiles were related among strains isolated from particular sources. Majority of the human isolates (n = 12, 86%) carried two or more large (>20-kb) plasmids and a similar situation was noted for isolates cultured from the animal sources (n = 16, 89%; Figure 1).

PBRT OF ISOLATES EXPRESSING AN ESBL-PHENOTYPE

Eighteen plasmid replicons were detected by qualitative PBRT (Carattoli et al., 2005) among those 32 isolates carrying large (>20-kb) plasmids. Replicon typing was identified 11 of 18 replicons. Interestingly, IncFIA, IncFrep, IncHI2, IncI1M, IncW, IncT, or IncX types could not be detected, by PCR analysis in our collection. Nine Inc-types were identified among the isolates cultured from healthy humans. Of these, IncFIB (n = 13), IncFIA (n = 5), and IncK/B (n = 4) replicon types were the predominant types. Isolates recovered from poultry group had the highest number of IncI1 (n = 8) types. Three other replicon types were also detected in E. coli isolates cultured from poultry and these included IncFIB (n = 7), IncK/B (n = 4), and IncT (n = 3). When compared with the Inc-types identified in isolates cultured from humans, replicon types from the animal strains were less diverse. Several isolates were positive for more than one replicon type. The reason could be that they possessed multiple plasmids, or a single plasmid-encoded replication, or partitioning genes from more than one replicon family. A summary of these features along with the corresponding antimicrobial resistance profiles for all 32 isolates is shown as a heat-map in Figure 1.

CONJUGATION EXPERIMENTS WITH ESBL-PRODUCING ISOLATES

Each isolate of the 32 selected E. coli, elaborating an ESBL-phenotype, were tested for their ability to transfer the ESBL-resistant phenotype, by conjugation under laboratory conditions. Nine isolates transferred the cefotaxime resistance marker to a susceptible E. coli recipient with transfer rates ranging from 3.1 × 10^{-2} (in the case of L-2) to 5.7 × 10^{-3} (for C-49) transconjugants per donor cell. Transconjugants recovered were characterized as described below.

HORIZONTAL TRANSFER OF ANTIMICROBIAL RESISTANCE AND ASSOCIATED DETERMINANTS

With the exception of E. coli H-1519 cultured from a human and which contained two small plasmids (of 2- and 7-kb; data not shown), none of the other smaller (>30-kb) plasmids could be transferred to the E. coli rifampicin-resistant recipient under laboratory conditions. Four isolates (H-2291, H-2332, C-25, and C-60) transferred a single large plasmid only (Figure 2).

Conjugation studies and plasmid profile analysis of all transconjugants showed that the nine isolates possessed 16 plasmids. Of these, 14 were large (>30-kb) plasmids and the replicon types were also identified, by PCR. Six plasmids belonged to different donors and possessed IncI1 replicons. Three large plasmids contained IncFIB replicons and two contained IncN types. Single plasmids were typed as being members of the Incp or IncBO replicon families (Figure 2).

Following conjugation the antimicrobial susceptibility of all nine transconjugants was determined by disk diffusion. In addition to cefotaxime resistance, resistance to several non-β-lactam-based antimicrobial compounds such as chloramphenicol, tetracycline, trimethoprim, and trimethoprim–sulfamethoxazole were also transferred to the recipient, suggesting that these markers are genetically linked. The MIC values recorded for all nine
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FIGURE 1 | A heat-map summary of the sources, a schematic showing the IS1 nucleic plasmid profiles, ESBL present/absent profiles, the resistance profile and the corresponding Inc plasmid type(s) for all 32 ESBL-positive E. coli previously reported from healthy food-producing animals and humans in Switzerland. Black squares shown indicate a feature present in that isolate denoting its original source the ESBL marker(s) detected, its corresponding antimicrobial resistance profile and the Inc types detected. White squares denote features that are lacking in the corresponding bacterial isolate. Isolate numbers denotes by the grayed boxes on the left of the heat-map indicate those bacteria that contained self-transmissible ESBL-harboring plasmids. (they are investigated further, as described in Figure 2). Antimicrobial compounds (it is known that many ESBL producers may appear susceptible or intermediate to certain oxyimino cephalosporins in vitro, if CLSI criteria are applied strictly, but do not respond to the respective therapies. Consequently, for clinical reporting these results have to be corrected to “resistant.”) used are abbreviated as follows: AM, ampicillin; AMC, amoxicillin–clavulanic acid; CAZ, ceftazidime; CF, cephalothin; CPD, cefpodoxime; CIP, ciprofloxacin; CTX, cefotaxime; CXM, cefuroxime; FEP, cefepime; GM, gentamicin; S, streptomycin; SXT, trimethoprim-sulfamethoxazole; TE, tetracycline.

Donors ranged from 96 to >256 μg/ml by cefotaxime E-test. When these MIC values were measured for the transconjugants, all were recorded at 256 μg/ml and in three cases there was between a 2- to 2.7-fold increase, in this value the MICs of transconjugants correspondingly were higher (>256 μg/ml; Figure 2).

Twenty-eight antibiotic resistance-encoding genes were tested on commercial microarray in all nine donor strains. Only nine of these markers, including aadA, cat, cti, dfr, tem, tet, strA, strB, and sul were detected. A CTX-encoding gene was detected in eight isolates, with an additional isolate positive for the TEM gene. These data were subsequently confirmed by PCR and the genes identified as blaCTX-M-15 and blaTEM-52, respectively. Five strains were positive for the blatem-1 gene. Six isolates were positive for the integra-encoding gene (intI1) associated with class 1 integrons. The genotypes of all nine donors and their corresponding transconjugants are summarized in Figure 2 and Table 1.

DISCUSSION
By definition, plasmids do not carry genes essential for the growth of host cells under non-stressed conditions but most plasmids confer positively selectable phenotypes including antimicrobial resistance genes among others (Carattoli, 2011). Plasmids
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Table 1 | Summary of genotypic and phenotypic characteristics of nine ESBL-producing E. coli isolates cultured from healthy food-producing animals and humans in Switzerland.

| Isolate no. | Phylogenetic groups | Serotype | MLST | Plasmid(s) (kb) isolated from transconjugants | Additional resistance determinants identified using a commercial microarray a |
|-------------|---------------------|----------|------|-----------------------------------------------|---------------------------------------------------------------|
| H-1038      | B1                  | O8:H21   | Unknown ST | 140 - 3 - 7 - 80 - 100 - | aadA-strA-strB-cat-dfr-sul-tet-intl |
| H-1519      | A                   | O2:H48   | ST10 complex | 110 - 140 - | aadA-dfr-sul-tet-intl |
| H-2291      | A                   | D53.H18  | ST1638 | 100 - 150 - | aadA-dfr-sul-tet-intl |
| H-2332      | D                   | D24.H26  | ST57/ST350 complex | 100 - 150 - | aadA-dfr-sul-tet-intl |
| C-23        | B1                  | D3.Hnt   | ST389 | 100 - | - |
| C-49        | B1                  | Dnt.H21  | ST446 complex | 110 - | aadA-dfr-sul-tet-intl |
| C-59        | B1                  | D86.H51  | ST155 complex | 110 - 150 - | aadA-cat-dfr-sul-tet-intl |
| C-60        | B1                  | D86.H21  | ST374 complex | 110 - | aadA-dfr-sul-tet-intl |
| L-2         | B1                  | Dnt.H11  | ST295 | 50 - 100 - | strB-cal-sul-tet |

aMethod used: Identi Bac Chip (ALIMVE Genotyping Kit, Version 05m); the symbol “–” signifies the fact that the bacterial isolate is negative for those genes included on the commercial microarray; “Ont” denotes an isolate whose serotype was non-typable.

represent an important pool of adaptive and transferable genetic information and large plasmids (defined as being >30-kb in size) are commonly being capable of transferring their genetic information in bacteria (Norman et al., 2009; Smillie et al., 2010; Williams et al., 2013). Previously E. coli isolates were cultured from a number of healthy food-producing animals and humans in Switzerland and all expressed an ESBL-phenotype. In the collection of 32 isolates which were further studied, 82 megaplasmids were identified. Four plasmids had molecular sizes ranging from 200- to about 350-kb and a further 78 (95%) were large plasmids of sizes between 50- and 200-kb were observed. The approximate molecular sizes of plasmids were determined by comparing with the reference
plasmids (Macrina et al., 1978). At the outset, these data suggested that the ESBL markers may be mapped to larger sized plasmids (50–200 kb).

Currently, 27 Inc groups are recognized among the Enterobacteriaceae family (Carattoli, 2011). Particular plasmid incompatibility groups are more frequently encountered among E. coli and these play a major role in the dissemination of specific resistance genes. For instance, in Netherlands, genetically related IncI1 plasmids carrying the bla_{CTX-M-52} gene were identified in E. coli isolates from poultry meat samples and in infected individuals suggesting that transfer of these plasmids may have occurred among strains from food-producing animals and humans (Leverstein-van Hall et al., 2011).CTX-M-1 was also identified on plasmids belonging to the IncF family in Europe among porcine (Moodley and Guardabassi, 2009), canine (Schink et al., 2011), and human E. coli isolates (Novais et al., 2007). In contrast, IncFI, IncAC, IncM1, IncN, and IncI1 plasmids carrying ESBL determinants are currently considered to be epidemic resistance plasmids, being detected in Enterobacteriaceae of different origin and sources (Carattoli, 2011). In the study reported here, PFGE typing identified 11 of 18 replicons (including IncA/C, IncFIA, IncFIB, IncFIC, IncHI1, IncK/B, IncI1, IncN, and IncY). IncFIB, IncK/B, and IncI1 were the more commonly identified replicon families in this study with 67% (n = 6) being IncI1 plasmids and which were isolated from the poultry samples. Moreover, conjugative transfer of plasmids belonging to incompatibility groups IncI1, IncFIB, and, less frequently, IncN, IncVO, and IncF could be demonstrated under laboratory conditions, suggesting that the potential dissemination of these markers may exist in the wildlife (Figure 2). This observation is interesting given that plasmids of IncI (both IncI1 and IncI2) and IncF groups are commonly recovered from E. coli and Salmonella species cultured from both animals and humans, and they are also linked to a number of genes coding for ESBL-enzymetypes (Johnson et al., 2007; Carattoli, 2011; Doumith et al., 2012).

IncI1 plasmids are often associated with resistance to multiple antimicrobial compounds, particularly extended-spectrum cephalosporinases of both the CTX-M and CMY types (Hopkins et al., 2006; Marcade et al., 2009). These have also been linked to TEM-52 enzymes. IncI1 plasmids producing TEM-52 have been identified in E. coli cultured from chicken and turkey meat in Denmark and the UK (Jensen et al., 2006; Randell et al., 2011). Interestingly an IncI1 plasmid carrying bla_{CTX-M-52} and bla_{TEM-52} was associated with the recent 2011 outbreak of the E. coli/O104:H4 strain in Germany (Jensen et al., 2006). Three conjugative IncI1 plasmids were identified among the poultry samples. Moreover, characterization of the resistance genes together with an investigation of their potential for dissemination among bacteria of animal origin provides valuable insights into this important resistance gene pool (Norman et al., 2009). Use of extended-spectrum cephalosporins, in animal and human medicine can select for ESBL-enzymetypes. Dating back to the late 1990s, CTX-M enzymes are recognized as the predominant ESBL-types of animal origin as well as in human isolates in Europe (Livermore et al., 2007). TEM-1 is the most common plasmid-mediated ß-lactamase identified in enteric Gram-negative bacilli, high rates of ampicillin-resistant, ß-lactamase-positive isolates are to be expected (Paterson and Bonomo, 2005). In the current study, the bla_{CTX-M-1} gene was identified in eight transconjugants (including four from healthy humans, three from chickens, and one from a lamb; Figure 2). The ß-lactamase gene was also identified in five transconjugants (four from healthy humans and one from a chicken sample; Figure 2). To our knowledge, neither bla_{CTX-M-1} nor bla_{TEM-1} associated with conjugative plasmids has been reported in healthy humans or food-producing animals in Switzerland.

In the laboratory, cefotaxime is used for optimum detection of ß-lactamase encoding isolates aiding in the identification of CTX-M carrying bacteria (Rodriguez et al., 2006). In this study, cefotaxime resistance was observed to occur concomitantly with resistance to AMP/KF/CPD in eight transconjugants and this feature could be linked directly to the presence of bla_{CTX-M-1} (in eight of nine transconjugants), to bla_{TEM-1} (in five isolates), and to bla_{TEM-52} (in one isolate). However, only one isolate showed resistance to amoxicillin–clavulanic acid (H-1519). As no evidence for mecanistic resistance, such as the expression of inhibitor-resistant TEM or SHV enzymes of the Bush–Jacoby groups 2be or 2ber (Bush and Jacoby, 2010), was found, this phenotype was most likely due to the hyper-expression of co-linear TEM-1 markers (Ortega et al., 2012; Zurfluh et al., 2013).

Interestingly, no IncFI1 type was identified. These data are in accordance with the previous reports (Johnson et al., 2007; Marcade et al., 2009). IncF replicons are widely distributed among E. coli and these plasmid types seem to be well adapted to this species. Three large potential conjugative plasmids (100–150 kb) of IncFIB replicon type were identified. This finding supports the view that IncF family plasmids play a major role in the dissemination of antibiotic resistance in Enterobacteriaceae, and some have been associated with specific genes conferring resistance to aminoglycosides, ß-lactams, and quinolones (Hopkins et al., 2006; Marcade et al., 2009). Plasmids of the IncF group are recognized as a significant antibiotic resistance marker reservoir due to their self-transmissibility and broad host range (Schlueter et al., 2007). Only a single possible conjugative IncP plasmid was identified among the transmissible plasmids studied. A similar situation was noted for the IncBO type. This feature was interesting, given the fact that in previous studies IncP and IncBO were the dominant Inc-types identified among transmissible plasmids characterized (Karczmarczyk et al., 2011). ß-Lactamase-encoding (bla) genes have benefited from the various mechanisms available to promote HGT between bacteria, thereby ensuring the spread of these markers to new hosts. It is often included as component parts of multi-resistance plasmids commonly found in clinical isolates (Bush and Jacoby, 2010). Identification of the resistance genes together with an investigation of their potential for dissemination among bacteria of animal origin provides valuable insights into this important resistance gene pool (Norman et al., 2009). Use of extended-spectrum cephalosporins, in animal and human medicine can select for ESBL-enzymetypes. Dating back to the late 1990s, CTX-M enzymes are recognized as the predominant ESBL-types of animal origin as well as in human isolates in Europe (Livermore et al., 2007). TEM-1 is the most common plasmid-mediated ß-lactamase identified in enteric Gram-negative bacilli, high rates of ampicillin-resistant, ß-lactamase-positive isolates are to be expected (Paterson and Bonomo, 2005). In the current study, the bla_{CTX-M-1} gene was identified in eight transconjugants (including four from healthy humans, three from chickens, and one from a lamb; Figure 2). The ß-lactamase gene was also identified in five transconjugants (four from healthy humans and one from a chicken sample; Figure 2). To our knowledge, neither bla_{CTX-M-1} nor bla_{TEM-1} associated with conjugative plasmids has been reported in healthy humans or food-producing animals in Switzerland.

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MIDs of cefotaxime of three transconjugants (C-23-(T), C-49-(T) and C-59-(T), Figure 2) were between 2- and 2.7-fold higher than the corresponding donor strains. These increases indicated that an amplification of multiple copies of the ESBL gene or occurrence of a point mutation in one gene copy might increase specific activity of \( \beta \)-lactamase and downregulate its catalytic power to increase resistance (Tenover et al., 2006; Sun et al., 2009).

All nine transconjugants showed phenotypic resistance to more than one class of antimicrobial compound. One isolate (H-1538-(T); Figure 2) was found to be resistant to members of four different drug classes (aminoglycosides, chloramphenicol, sulfamethoxazole/trimethoprim, and tetracycline), and a further three (H-2291-(T), H-2332-(T) and C-59-(T), Figure 2) were resistant to two different antimicrobial compound classes (aminoglycosides, sulfamethoxazole/trimethoprim, and tetracycline or chloramphenicol). To our knowledge, this is the first evidence of this emerging multi-resistant clone with large conjugative plasmids from healthy humans and animals in Switzerland. Furthermore, ESBL-producing \( E. \) coli ST57 and ST371 were cultured from chickens in the UK, Germany, and Canada (Randall et al., 2011). Of the nine donor strains investigated in this study, the clonal lineages detected among healthy humans included phylogroup B1-08:H21-unknown ST type, phylogroup A-O2:H48-ST10 complex, and phylogroup D-01707-08. Among the five animal isolates the lineages included phylogroup B1-035:H1-ST389, phylogroup B1-035:H1-ST3174 complex, phylogroup B1-036:H51-ST315 complex, and phylogroup D-01701-ST295 from a lamb source (Table 1). Our study represents the first report of ESBL-producing \( E. \) coli of nine different ST types from animals and humans in Switzerland. Interestingly, these shared a distribution of plasmid replicon types similar to those reported previously in unselected \( E. \) coli, whereas in contrast the distribution of STs was markedly different (EFSA Panel on Biological Hazards [BIOHAZ], 2011; Randall et al., 2011). These findings may indicate the presence of systems affecting the incorporation and/or maintenance of antibiotic resistance genes not related to plasmid restriction (Bengtsson et al., 2012).

Characterizing plasmids from bacteria of different origins may provide early insights into the epidemiology of important resistance markers, including ESBL-determinants. Few studies have provided clear evidence of a direct transmission of ESBL-producing \( E. \) coli isolates from food-producing animals and/or food to humans. Nonetheless, these data showed that MDR phenotypes, including resistance to newer generations of antimicrobial compounds, were transferable and in some cases were directly linked to large conjugative plasmids carrying multiple resistance genes. This situation could, in the future give rise to the co-selection, promoting the dissemination of ESBL-phenotypes among bacteria of healthy food-producing animal and human origins alike.

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APPENDIX

FIGURE A1 | A heat-map summary of the original 87 ESBL-positive E. coli isolates previously reported from healthy food-producing animals and humans in Switzerland. Black squares shown indicate a feature present in that isolate denoting its original source and the ESBL marker(s) detected; and white squares denote features that are lacking in the corresponding bacterial isolate. The approximate sizes of small plasmids are also showing in separate columns. The symbol “−” signifies no small (<30-kb) plasmid in the corresponding strain. Isolate numbers denoted by the grayed boxes on the left of the heat-map indicate those bacteria that were selected for conjugation experiments and S1-nuclease-based PFGE studies (as described in Figure 1).