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Molecular Characterization of Multidrug-Resistant *Salmonella enterica* subsp. *enterica* Serovar Typhimurium Isolates from Swine

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As part of a longitudinal study of antimicrobial resistance among salmonellae isolated from swine, we studied 484 *Salmonella enterica* subsp. *enterica* serovar Typhimurium (including serovar Typhimurium var. Copenhagen) isolates. We found two common pentaresistant phenotypes. The first was resistance to ampicillin, chloramphenicol, streptomycin, sulfamethoxazole, and tetracycline (the AmCmStSuTe phenotype; 36.2% of all isolates), mainly of the definitive type 104 (DT104) phage type (180 of 187 isolates). The second was resistance to ampicillin, kanamycin, streptomycin, sulfamethoxazole, and tetracycline (the AmKmStSuTe phenotype; 44.6% of all isolates), most commonly of the DT193 phage type (77 of 165 isolates), which represents an unusual resistance pattern for DT193 isolates. We analyzed 64 representative isolates by amplified fragment length polymorphism (AFLP) analysis, which revealed DNA fingerprint similarities that correlated with both resistance patterns and phage types. To investigate the genetic basis for resistance among DT193 isolates, we characterized three AmKmStSuTe pentaresistant strains and one hexaresistant strain, which also expressed resistance to gentamicin (Gm phenotype), all of which had similar DNA fingerprints and all of which were collected from the same sampling. We found that the genes encoding the pentaresistance pattern were different from those from isolates of the DT104 phage type. We also found that all strains encoded all of their resistance genes on plasmids, unlike the chromosomally encoded genes of DT104 isolates, which could be transferred to *Escherichia coli* via conjugation, but that the plasmid compositions varied among the isolates. Two strains (strains UT08 and UT12) had a single, identical plasmid carrying *bla* _TEM_ (which encodes ampicillin resistance), _aphA1-Iab_ (which encodes kanamycin resistance), _strA_ and _strB_ (which encode streptomycin resistance), class B _tetA_ (which encodes tetracycline resistance), and an unidentified sulfamethoxazole resistance allele. The third pentaresistant strain (strain UT20) was capable of transferring by conjugation two distinct resistance patterns, AmKmStSuTe and KmStSuTe, but the genes were carried on plasmids with slightly different restriction patterns (differing by a single band of 15 kb). The hexaresistant strain (strain UT30) had the same plasmid as strains UT08 and UT12, but it also carried a second plasmid that conferred the AmKmStSuGm phenotype. The second plasmid harbored the gentamicin resistance methylase (*grm*), which has not previously been reported in food-borne pathogenic bacteria. It also carried the _sulI_ gene for sulfamethoxazole resistance and a 1-kb class I integron bearing _aadA_ for streptomycin resistance. We also characterized isolates of the DT104 phage type. We found a number of isolates that expressed resistance only to streptomycin and sulfamethoxazole (the StSu phenotype; 8.3% of serovar Typhimurium var. Copenhagen strains) but that had AFLP DNA fingerprints similar or identical to those of strains with genes encoding the typical AmCmStSuTe pentaresistance phenotype of DT104. These atypical StSu DT104 isolates were predominately cultured from environmental samples and were found to carry only one class I integron of 1.0 kb, in contrast to the typical two integrons (InC and InD) of 1.0 and 1.2 kb, respectively, of the pentaresistant DT104 isolates. Our findings show the widespread existence of multidrug-resistant *Salmonella* strains and the diversity of multidrug resistance among epidemiologically related strains. The presence of resistance genes on conjugative plasmids and duplicate genes on multiple plasmids could have implications for the spread of resistance factors and for the stability of multidrug resistance among *Salmonella* serovar Typhimurium isolates.

The frequency of resistance among food-borne pathogens has increased dramatically, presumably due to the extensive use of antimicrobial agents in human and veterinary medicine (8, 26). Furthermore, resistance to combinations of several classes of antimicrobials has led to the emergence of multidrug-resistant (MDR) strains that may pass from food animals to humans (30, 48). One important pathogen known to harbor multiple resistance factors is *Salmonella*, one of the leading causes of food-borne bacterial diseases. It is estimated that the annual economic costs due to food-borne *Salmonella* infections in the United States are $2.4 billion (http://www.ers.usda.gov).

MDR *Salmonella* isolates have been reported since the 1960s (1), with the resistance patterns of *Salmonella* serovars of public health importance often associated with specific phage types. One notable MDR strain is *Salmonella enterica* subsp. *enterica* serovar Typhimurium definitive type 104 (DT104). It was first recognized in the United Kingdom (39) and since has been reported in many parts of the world (6, 16, 18, 19, 27) and from various host species including food animals and pets (12, 22, 43, 46) as well as from processed ready-to-eat meat.
products (47). DT104 strains are commonly known to be pentaresistant, exhibiting resistance to ampicillin, chlorampheni-
locin, streptomycin, sulfamethoxazole, and tetracycline (the Am-
CmStSuTe resistance phenotype). The drug resistance spectra
of MDR strains of Salmonella serovars have also been expand-
ing in recent years (13, 23, 28, 40, 49).

Another important phage type of serovar Typhimurium of-
ten exhibiting multidrug resistance is DT193, an MDR strain
responsible for outbreaks in humans in the late 1980s and early
1990s, mainly in Europe. Two of the major outbreaks due to
DT193 (in Italy and the United Kingdom) were traced to
contaminated pork products (25, 32). Preliminary analysis of
the resistance determinants has shown that the isolates col-
mcted from the outbreak in the United Kingdom carried their
resistance factors on conjugative plasmids (17). This phage
type has been among the most common MDR strains that we
identified among swine isolates (15). Since at least two DT193
outbreaks affecting humans have been traced to contaminated
pork products within the last decade (25, 32) and since serovar
Typhimurium isolates with similar resistance patterns have been
reported to be increasingly prevalent in recent years (1998 report
of the National Antimicrobial Resistance Moni-
toring System [http://www.cdc.gov/ncidod/dbmd/narms/an-
nual/1998_anu.htm]), we thought that it would be important
to characterize the genetic basis of multidrug resistance in these
strains commonly isolated from swine in our study.

The genetic characterization of antimicrobial resistance
genes as well as their location and diversity is important in
identifying factors involved in resistance, understanding the
diversity of MDR strains, identifying genetic linkages among
markers, understanding potential transfer mechanisms, and
developing efficient detection methods. In this study, we iden-
tified two predominant pentaresistant MDR phenotype
among serovar Typhimurium (and serovar Typhimurium var. 
Copenhagen) isolates, determined the genetic diversities of
these isolates using amplified fragment length polymorphism
(AFLP) DNA fingerprinting techniques, identified the resis-
tance genes involved, and determined the locations and diver-
sities of these resistance genes.

### MATERIALS AND METHODS

#### Collection of Salmonella isolates

We collected fecal samples from two swine production companies using all-in and all-out management systems. Samples from a total of 6 nurseries and 18 finishing farms were included in the study. At each visit, 96 fecal samples and 30 feed samples were collected. In addition, environmental samples were collected from each farm. Ten dram swab samples were collected from cleaned and apparently disinfected empty barns at the finishing stage before pigs were moved in from nursery barns. Sample collection was performed for three replicates (cohorts of pigs) between 1997 and 2000. A total of 7,452 samples (6,912 fecal samples and 540 dram swab environmental samples) were collected for this study. With an average prevalence of 21% per group, a total of 1,565 Salmonella isolates of 30 different serovars were detected. The most common serovar, serovar Typhimurium (including serovar Typhi-
murium var. Copenhagen), constitutes more than 75% of the total. In this study, a subset of the total, 484 Salmonella serovar Typhimurium and Salmonella
serovar Typhimurium var. Copenhagen isolates, was characterized.

#### Isolation, identification, and antimicrobial resistance

We isolated salmonel-
lae using conventional methods, as described previously (2). Antimicrobial sus-
citability was tested with the Vitek Jr. (VITEK system manual, 1996; Bio-
merieux Vitek, Hazelwood, Mo.) semiautomated system for 11 antimicrobials. The antimicrobials (and the respective breakpoints for resistance, given in pa-
netheses) were amikacin (32 g/ml), amoxicillin-clavulanic acid (16 g/ml),
ampicillin (16 g/ml), cefotaxime (16 g/ml), cephalothin (16 g/ml), chloram-
phenicol (16 g/ml), ciprofloxacin (2 g/ml), gentamicin (8 g/ml), piperacillin
(32 to 64 g/ml), tetracycline (8 g/ml), and trimethoprim-sulfamethoxazole (40
to 80 g/ml). The breakpoints of the Vitek Jr. system indicated above were
matched with the NCCLS standard breakpoints for gram-negative enteric
organisms, and the following quality control strains were routinely used: Esche-
richia coli ATCC 25922, Pseudomonas aeruginosa ATCC 27853, Sphingobacter-
ium aureus ATCC 29213, and Enterococcus faecalis ATCC 29212 (29). The Kirby-
Bauer disk diffusion method for three more antimicrobials, kanamycin, strepto-
mycin, and sulfamethoxazole, was performed as described previously (15). Se-
rotyping and phage typing were performed at the National Veterinary Services
Laboratories, Ames, Iowa.

#### Identification of resistance gene alleles

We identified resistance genes either by PCR or, when necessary, by cloning of PCR products and sequencing of the
cloned genes. The primers used for PCR are listed in Table 1. We used the following PCR conditions: denaturation at 95°C for 5 min and then 40 cycles of
denaturation at 95°C for 1 min, annealing at 53°C for 30 s, and extension at 72°C
for 30 s.

In order to identify the resistance gene cassette(s) carried on integrons among non-DT104 strains with pentaresistant phenotypes, we cloned and sequenced the integron amplicon. Briely, class 1 integron primers with ends compatible with XbaI were designed, and DNA fragments were amplified with primers common
to class 1 integrons by using the PCR conditions listed above. The DNA product
was purified and digested with XbaI and cloned into a plasmid vector (pPCR-
Script Cam; Stratagene, La Jolla, Calif.). E. coli DH5a was transformed by
electroporation, and candidate β-galactosidase-negative white colonies were se-

---

### TABLE 1. PCR primers used in identification of resistance genes

<table>
<thead>
<tr>
<th>Gene</th>
<th>Oligonucleotide sequencesa</th>
<th>Product size (bp)</th>
<th>Reference or source</th>
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</thead>
<tbody>
<tr>
<td>blqPSE1</td>
<td>F, TTT GGT TCC GCG TTA TCT G; R, TAC TCC GAC CAA ATC CG</td>
<td>150</td>
<td>9</td>
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<tr>
<td>blaTEM</td>
<td>F, GCA GCA GTT GGT TAC ATC GA; R, GGT CCT CCG ATC GTT GTC AG</td>
<td>310</td>
<td>9</td>
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<tr>
<td>aphA1-lab</td>
<td>F, AAA CTT GCT CGA GGC; R, CAA ACC GTT ATT CAT TCG TGA</td>
<td>500</td>
<td>14</td>
</tr>
<tr>
<td>aadA</td>
<td>F, GTG GAT GGC GGC CTT ACG AA; R, AAT GGC CAG TCG GCA GCG</td>
<td>528</td>
<td>24</td>
</tr>
<tr>
<td>strA</td>
<td>F, CTT GGT GAT AAC GGC AAT TC; R, CCA ATC GCA GAT AGA AGG C</td>
<td>548</td>
<td>AP273682</td>
</tr>
<tr>
<td>strB</td>
<td>F, ATC GTC AAG GGA TGG AAA CC; R, GGA TCG TAU AAC ATA TGT TGC</td>
<td>509</td>
<td>AP246402</td>
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<tr>
<td>sulA</td>
<td>F, CAC TCG CAC AAG CCG TAA; R, GTC GGC CTC AGC AAT ATC</td>
<td>363</td>
<td>AP171555</td>
</tr>
<tr>
<td>tetA</td>
<td>F, GCT ACA TGC TGC TGT CCT TC; R, CAT AGA TCG CCG TGA AGA GG</td>
<td>210</td>
<td>X61367</td>
</tr>
<tr>
<td>tetB</td>
<td>F, TGT GTG AGG GGC AAT TTG TG; R, CTA TGG GGC CAA TAA CAC CG</td>
<td>659</td>
<td>19</td>
</tr>
<tr>
<td>tetG</td>
<td>F, CAG TGT CGT GAT TCT TAC GG; R, GAT TGG TGA GCG TCG TTA GC</td>
<td>844</td>
<td>19</td>
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<tr>
<td>gm</td>
<td>F, AAG CCG ACG AAG CCG GGC CTG; R, AAG GCG GCC CTC AAG GAG GTC</td>
<td>414</td>
<td>14</td>
</tr>
<tr>
<td>aadB</td>
<td>F, GAG GCA AAT CTT GCG CTC TGC TG; R, CTG TTA CAA CCG ACT GCC GC</td>
<td>310</td>
<td>14</td>
</tr>
<tr>
<td>aac(6)-I</td>
<td>F, TGA GCA TGA CCT TGC GAT; R, GAA CAG CAA GGC AAC CAG</td>
<td>337</td>
<td>AP282595</td>
</tr>
<tr>
<td>int</td>
<td>F, GCC ATC CAA GCA GCA AG; R, AAG CAG ACT CCT GAT GA</td>
<td>Variable</td>
<td>19</td>
</tr>
</tbody>
</table>

a F, forward; R, reverse.
ected on 5-bromo-4-chloro-3-indolyl-p-d-galactopyranoside. We verified the presence of intact isolation of plasmid DNA, and inserts were sequenced with an Applied Biosystems automated sequencer.

**DNA fingerprinting.** We used a previously described AFLP fingerprinting method (41), with modifications. This method has been shown to have superior discriminatory power and reproducibility for the subtyping of *Salmonella* serovars (21, 31), and, in addition, we found this method to be both economical and efficient. Briefly, cells were grown overnight on Luria-Bertani broth (Difco, Atlanta, Ga.), and the genomic DNA was purified with a Qiagen DNAeasy tissue kit (Qiagen, Valencia, Calif.), as described in the DNAeasy Tissue Kit Handbook (p. 16-18; Qiagen GmbH, Hilden, Germany, 1999) and adjusted to a concentration of 50 ng/μl in a volume of 10 μl of water. The DNA was digested with *EcoRI* and *MseI* at 37°C for 1 h. Adapters were then ligated to each end of the restriction fragments with T4 DNA ligase (New England Biolabs) at 16°C overnight. The fragments were then amplified with primers specific for the *EcoRI* adapter (forward primer, 5′-GACTCGGTCTTGATGTA) and the *MseI* adapter (reverse primer, 5′-GATGAGTCCTGAGTAA), ensuring that only those fragments with one *EcoRI* end and one *MseI* end were amplified. The conditions for amplification were 94°C for 15 s, 60°C for 30 s with increases of 1 s per cycle for 28 cycles, and then incubation at 72°C for 2 min. The amplified fragments then underwent a second round of amplification, this time with an infrared-labeled *EcoRI* primer (Liorc, Lincoln, Neb.). An additional adenine was added to this primer at its 3′ end (5′-GACTCGGTCTTGATGTA) in order to obtain the fewer bands optimum for band scoring and fingerprint analysis. This final selective amplification consisted of 13 cycles at 94°C for 10 s, 65°C for 30 s, and 72°C for 1 min and then 25 cycles of 94°C for 10 s, 58°C for 30 s, and 72°C for 1 min, with ramping at 1 s/cycle for the final step. The reaction mixture was finally incubated for 2 min at 72°C. The fragments were then separated on a Licor 4200 DNA sequencer. A detailed AFLP protocol can be found at http://www4.ncsu.edu/~wagebrey/AFLP-Salm.pdf. The bands were scored and analyzed with Quantity One software (version 4.1.1; Bio-Rad, Richmond, Calif.). The fingerprints were analyzed by using the Dice coefficient algorithm, and dendrograms were constructed by the unweighted pair group method with arithmetic averages clustering method.

**Conjugation and restriction patterns of plasmids.** We performed conjugation experiments to determine whether antimicrobial resistance markers were located on conjugative plasmids. Candidate MDR donor strains were mated with a spontaneous rifampin-resistant derivative of *E. coli* K-12 strain MG1655. Single colonies of each of the donor and the recipient were mixed on Luria-Bertani agar, and the mixture was incubated for 6 h at 37°C. The mixture was then transferred to a selective plate containing rifampin and one of the antimicrobials to be tested and incubated at 37°C overnight. Transconjugants were further purified and confirmed to be *E. coli* rather than spontaneous rifampin-resistant *salmonellae* by growth on MacConkey agar. Selective plating and PCR amplification were used to test for the presence of other resistance markers. The mixture was then ampliﬁed with primer speciﬁc for the EcoRI adapter (forward primer, 5′-GACTCGGTCTTGATGTA) and the MseI adapter (reverse primer, 5′-GATGAGTCCTGAGTAA), ensuring that only those fragments with one EcoRI end and one MseI end were ampliﬁed. The conditions for ampliﬁcation were 94°C for 15 s, 60°C for 30 s with increases of 1 s per cycle for 28 cycles, and then incubation at 72°C for 2 min. The amplified fragments then underwent a second round of amplification, this time with an infrared-labeled EcoRI primer (Liorc, Lincoln, Neb.). An additional adenine was added to this primer at its 3′ end (5′-GACTCGGTCTTGATGTA) in order to obtain the fewer bands optimum for band scoring and fingerprint analysis. This final selective amplification consisted of 13 cycles at 94°C for 10 s, 65°C for 30 s, and 72°C for 1 min and then 25 cycles of 94°C for 10 s, 58°C for 30 s, and 72°C for 1 min, with ramping at 1 s/cycle for the final step. The reaction mixture was finally incubated for 2 min at 72°C. The fragments were then separated on a Licor 4200 DNA sequencer. A detailed AFLP protocol can be found at http://www4.ncsu.edu/~wagebrey/AFLP-Salm.pdf. The bands were scored and analyzed with Quantity One software (version 4.1.1; Bio-Rad, Richmond, Calif.). The fingerprints were analyzed by using the Dice coefficient algorithm, and dendrograms were constructed by the unweighted pair group method with arithmetic averages clustering method.

**RESULTS**

Phenotypic characterization of MDR serovar Typhimurium and serovar Typhimurium var. *Copenhagen.* In a longitudinal study of swine salmonellosis in commercial swine production systems at multiple sites in North Carolina, we analyzed a total of 484 isolates (156 *Salmonella* serovar Typhimurium isolates and 328 *Salmonella* serovar Typhimurium var. *Copenhagen* isolates) collected in three replicate samples of 18 groups of pigs. We found two common pentaresistant MDR phenotypes: resistance to ampicillin, chloramphenicol, streptomycin, sulfamethoxazole, and tetracycline (the AmCmStSuTe phenotype; 36.2%) or resistance to ampicillin, kanamycin, streptomycin, sulfamethoxazole, and tetracycline (the AmStSuTe phenotype; 44.6%). Phage typing resulted in the classification of the serovar Typhimurium isolates (including serovar Typhimurium var. *Copenhagen* isolates) into eight different phage types: DT104, DT193, DT21, DT208, DT12, U302, DT169, and DT120. Analysis of isolates with the aforementioned two pentaresistance patterns showed that DT104 predominated among isolates with the AmCmStSuTe resistance pattern (106 of 121 isolates for which the phage type was obtained), while phage type U302 was less commonly represented among isolates with this pentaresistant phenotype (11 of 121 isolates). The isolates with the AmKmStSuTe pentaresistance pattern were predominantly of the DT193 phage type (53 of 59 isolates) among serovar Typhimurium var. *Copenhagen* isolates and the DT21 phage type among the serovar Typhimurium isolates (82 of 106 isolates).

**Genetic diversity of MDR strains.** In order to determine the extent of genetic diversity among these MDR strains of *Salmonella* serovar Typhimurium (including *Salmonella* serovar Typhimurium var. *Copenhagen*), we performed DNA fingerprinting using AFLP analysis of 64 isolates originating from eight farms. The dendrogram in Fig. 1 shows the similarities of these strains on the basis of the fragment sizes of the amplified bands. Isolates with the same letter code (e.g., UT) were collected from the same farm on the same visit. The results show that two major clusters of Typhimurium and serovar Typhimurium var. *Copenhagen* were identified. The first cluster (Fig. 1, cluster 1, A to C) included 19 of the 64 isolates fingerprinted. The predominant resistance pattern was pentaresistance of the AmKmStSuTe phenotype (15 of 19 isolates), but isolates with the AmKmStSuTeSgm (1 of 19 isolates), StTe (1 of 19 isolates), and AmCmStSuTe (2 of 19 isolates) phenotypes were also seen. In this cluster, only one isolate was found from an environmental sample (a drag swab of the barn floor), and this isolate exhibited the AmCmStSuTe resistance phenotype. The predominant phage type in this cluster was DT193 (11 of 19 isolates), followed by DT12 (4 of 19 isolates), DT104 (1 of 19 isolates), and U302 (1 of 19 isolates), along with two untypeable isolates. Isolates in this cluster were derived from only four of the eight farms.

The second cluster (Fig. 1, cluster 2, A to E) included 45 of the 64 isolates fingerprinted. The predominant resistance phenotype in this cluster were AmCmStSuTe (31 of 45 isolates) and StSu (9 of 45 isolates). Rarely, isolates with the hexaresistance patterns were also found: UF19 with the AmCmSt-SuTeCe phenotype (2 of 45 isolates) and UH03 with the AmCmStSuTeGm phenotype (1 of 45 isolates). The remaining two isolates, U002 and UM06, showed a pentaresistance pattern (the AmKmStSuTe phenotype), which was the predominant resistance phenotype of cluster 1. The predominant phage type in the second cluster was DT104 (41 of 45 isolates), with isolates of this phage type commonly exhibiting one of two resistance phenotypes: AmCmStSuTe or StSu. In addition to DT104, one isolate of each of phage types DT193, DT21, DT169, and U302 was found within this cluster. MDR strains in this cluster were isolated from all eight farms: 15 (34%) from environmental samples (drag swabs) and 30 (66%) from fecal samples.

Characterization of non-DT104 pentaresistant serovar Typhimurium and serovar Typhimurium var. *Copenhagen* isolates. We have identified a group of MDR strains with the AmKmStSuTe resistance pattern. Strains of phage types
DT193, DT12, DT21, and DT208 exhibited this resistance phenotype, the most common one being DT193. This phage type has been implicated in disease outbreaks, but the resistance profile that we have most often seen (AmKmStSuTe) is different from those that have commonly been observed in previous studies: either AmStSuTe (3, 11, 17) or Te alone (17). We chose four isolates for genetic characterization: isolates UT08, UT12, and UT20 (all with the AmKmStSuTe resistance phe-
found that UT08 and UT12 had identical isolates obtained from the same environment. Second, we intended to study the similarity (or diversity) of phenotypically similar isolates obtained from the same environment. Thus, characterization of these isolates enabled us to study the similarity (or diversity) of phenotypically similar isolates obtained from the same environment.

First, all four isolates were collected from the same farm at the same time. Thus, characterization of these isolates enabled us to study the similarity (or diversity) of phenotypically similar isolates obtained from the same environment. Second, we found that UT08 and UT12 had identical fingerprints, consistent with their identical resistance phenotypes, but that UT20 had a different fingerprint, although it shared the same penta-resistance phenotype. In addition, we noticed that the fingerprint of the hexaresistant UT30 strain was similar to those of UT08 and UT12, even though its resistance pattern included resistance to gentamicin (the Gm phenotype). Thus, we were interested in studying further this disparity between fingerprint similarity and resistance phenotype.

Conjugal transfer of resistance genes and plasmid characterization. To determine whether any of the resistance genes of these MDR strains were located on conjugative plasmids, we conjugated each of the four donors (isolates UT08, UT12, UT20, and UT30) with a rifampin-resistant derivative of E. coli K-12 strain MG1655. We found that all of the resistance markers in all four isolates could be transferred by conjugation and thus were encoded on conjugative plasmids. However, the numbers and restriction patterns of the conjugative plasmids carrying resistance markers varied within and between strains. For UT08 and UT12, conjugation with selection for resistance to ampicillin, kanamycin, or tetracycline always yielded transconjugants resistant to all five antimicrobials to which the donor strains were resistant. Restriction enzyme analysis revealed that UT08 and UT12 each transfer approximately 140 kb of plasmid DNA harboring all five of the resistance genes (Fig. 2, lanes 1 and 2). These plasmids also had identical restriction fragment patterns when they were cut with HindIII (Fig. 2) and other enzymes (data not shown). Thus, it is likely that these two strains, which have identical resistance patterns and fingerprints and which carry the same plasmid, are in fact clonal.

UT30, however, was able to transfer by conjugation plasmids with two distinct resistance patterns and restriction maps. Despite the similarity of its fingerprint to those of UT08 and UT12, it could transfer by conjugation a plasmid encoding resistance to amikacin, kanamycin, streptomycin, sulfamethoxazole, and tetracycline as well as one producing resistance to amikacin, kanamycin, streptomycin, sulfamethoxazole, and gentamicin. The first of these, which encoded the AmKmStSuTe resistance phenotype, was identical to the plasmids isolated from UT08 and UT12; it was 140 kb and had the same HindIII restriction pattern as the plasmids isolated from UT08 and UT12 (Fig. 2, lane 5). The second plasmid, which encoded the AmKmStSuGm resistance phenotype, was approximately 145 kb and had a different restriction pattern from those of the other plasmids (Fig. 2, lane 6). Thus, it is possible that UT30 is a derivative of UT08 and/or UT12 that has acquired a second conjugative plasmid that encodes resistance to gentamicin and duplicates the resistance to the other four antimicrobials already present in these strains. UT20 also carried two conjugative plasmids with restriction patterns different from those of the plasmids found in the other three strains: an 85-kb plasmid (pUT20A) carried genes encoding resistance to all five antimicrobials found in UT20 (Fig. 2, lane 3), and a second plasmid (pUT20B) of about 70 kb harbored genes encoding resistance to kanamycin, streptomycin, sulfamethoxazole, and tetracycline (Fig. 2, lane 4). Apart from the presence of a 15-kb band in pUT20A, the two plasmids from UT20 exhibited similar restriction patterns. The similarities of the restriction patterns of these two plasmids suggest that they may in fact be related plasmids.

Identification of antimicrobial resistance genes. To characterize the genetic basis of resistance in the four plasmids isolated and described above, we identified resistance genes using PCR and cloning. Since strains UT08, UT12, and UT20 appear to carry the same plasmid, we used the plasmid originally obtained from UT08 (pUT08) to represent the other two in these studies. Unlike DT104, which encodes β-lactam resistance using the blαPS₁₁ gene, all three plasmids have a phenotype of resistance to β-lactams (pUT08, pUT20A, and pUT30) encoded ampicillin resistance by means of the blαPS₁₁ gene (Fig. 3A, B, D, and E). Further PCR testing of an additional 59 MDR isolates with the AmKmStSuGm resistance pattern revealed that they all carried the blαPS₁₁ gene (data not shown). We found that all plasmids encoding kanamycin resistance did so by means of the aminoglycoside phosphotransferase gene aphA1-1ab and that all plasmids encoding tetracycline resistance had tetB (Fig. 3A to E). However, we identified two different streptomycin resistance alleles: streptomycin-6-phosphotransferase (strA and strB) was the sole streptomycin resistance determinant in three of the plasmids (pUT08, pUT20A, and pUT20B). On plasmid pUT30B, however, we also found the aminoglycoside adenyltransferase gene (aadA), in addition to strAB (Fig. 3D and E). Using a primer pair homologous to
regions of class I integrons that flank the antibiotic resistance cassettes (Table 1), we amplified by PCR an approximately 1-kb product from plasmid pUT30B. Cloning and sequencing of this product revealed that aadA is encoded on this class I integron. We found sulfamethoxazole resistance encoded by the dihydropteroate synthase gene (sul1) in pUT30, but we were not able to detect this gene on the other three plasmids (Table 2). The gene responsible for sulfonamide resistance on these plasmids remains to be identified. To identify the gene responsible for gentamicin resistance, we designed primers specific for three distinct bacterial genes known to encode resistance to gentamicin [aac(6)-I, aadB, and grm] and attempted to amplify each from plasmid pUT30. We found that we could detect only the gentamicin resistance methylase (grm) of Micromonospora purpurae, an allele not previously identified in pathogenic bacteria (Fig. 3E). The presence of both a class I integron and the gentamicin resistance gene on the same plasmid derived from UT30 could have important implications.

### Table 2. Summary of antimicrobial resistance genes and class I integrons identified among Salmonella isolates of phage types DT193, DT104, and U302

<table>
<thead>
<tr>
<th>Phage type</th>
<th>Plasmid or pattern</th>
<th>Gene(s) for resistance to antimicrobials</th>
<th>Class I integron size/location</th>
</tr>
</thead>
<tbody>
<tr>
<td>DT193</td>
<td>pUT08 (AmKmStSuTe)</td>
<td>blaTEM, aphA1-Iab, strAB, ND*</td>
<td>tetB</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT12</td>
<td>blaTEM, aphA1-Iab, strAB, ND*</td>
<td>tetB</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT20A</td>
<td>blaTEM, aphA1-Iab, strAB, ND</td>
<td>tetB</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT20B</td>
<td>blaTEM, aphA1-Iab, strAB</td>
<td>ND</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT30A</td>
<td>blaTEM, aphA1-Iab, strAB</td>
<td>ND</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT30B</td>
<td>blaTEM, aphA1-Iab, strAB, sul1</td>
<td>grm, 1.0 kb/plasmid</td>
</tr>
<tr>
<td>DT193</td>
<td>pUT30B</td>
<td>blaTEM, aphA1-Iab, strAB, sul1</td>
<td>grm, 1.0 kb/plasmid</td>
</tr>
<tr>
<td>DT104</td>
<td>AmCmStSuTe</td>
<td>blaTEM, aadA, sul1</td>
<td>tetG, 1.0 and 1.2 kb/chromosome</td>
</tr>
<tr>
<td>DT104</td>
<td>StSu</td>
<td>aadA, sul1</td>
<td>tetG, 1.0 and 1.2 kb/chromosome</td>
</tr>
<tr>
<td>U302</td>
<td>AmKmStSuTe</td>
<td>blaTEM, aphA1-Iab, strAB, sul1</td>
<td>tetA, 1.0 kb/ND</td>
</tr>
<tr>
<td>U302</td>
<td>AmKmStSuTe</td>
<td>blaTEM, aphA1-Iab, strAB, sul1</td>
<td>tetG, 1.0 and 1.2 kb/ND</td>
</tr>
</tbody>
</table>

*ND, not determined.*
in the expansion of the multidrug resistance spectrum, with the possibility of additional resistance genes being captured by the integron and the potential for the horizontal transfer of resistance genes within or between bacterial species.

**Class I integron polymorphism among DT104 isolates.** Among *Salmonella* serovar Typhimurium isolates, strains of phage type DT104 are known to be some of the MDR strains most important to public health. Strains of this phage type also most commonly have a pentaresistant AmCmStSuTe resistance phenotype. It has recently been shown that pentaresistant DT104 strains harbor two class I integrons, encoding resistance to ampicillin, streptomycin, and sulfamethoxazole, and an intervening region encoding resistance to chloramphenicol and tetracycline, all within a span of 13 kb (7). However, in this study, we have identified DT104 isolates with the StSu resistance pattern (8.3% of serovar Typhimurium var. Copenhagen isolates). These isolates had epidemiological significance in that they were more commonly isolated from an environmental source, i.e., drag swabs of cleaned barns (seven of the nine isolates). To characterize these StSu isolates, we first performed DNA fingerprinting to determine the genetic similarity between DT104 isolates with pentaresistance and those with the StSu resistance phenotype. As shown in the dendrogram in Fig. 1, in most cases StSu isolates exhibited a fingerprint identical to that of at least one pentaresistant DT104 isolate. Based upon this observation, we examined the genetic basis of resistance of the StSu DT104 isolates from our study. Although pentaresistant DT104 is known to harbor two integrons of 1.0 and 1.2 kb (InC and InD, respectively), the StSu strains evaluated in this study carried only one class I integron of 1.0 kb (Fig. 4), implying the absence of part of the pentaresistant architecture of DT104. Pentaresistant DT104 harbors two class I integrons on a 13-kb chromosomal region, and these integrons encode the complete pentaresistance phenotype. In strains of the StSu resistance phenotype, the integron encoding the β-lactam resistance and the intervening segment encoding chloramphenicol and tetracycline resistance may have been excised. In addition to the lack of one integron, we verified the absence of ampicillin, chloramphenicol, and tetracycline resistance genes by PCR testing of the known resistance genes (Table 2).

**DISCUSSION**

An increased incidence of MDR *Salmonella* serovar Typhimurium has been widely reported in the last decade, presumably due to the extensive use of antimicrobial agents in human and veterinary medicine. Here we have described the common resistance patterns and genetic characterization of MDR *Salmonella* serovar Typhimurium strains cultured from commercial swine operations. We found two common pentaresistance patterns: AmCmStSuTe and AmKmStSuTe. The former resistance pattern was most commonly found in strains of the DT104 phage type, not an unexpected finding, since numerous studies have shown the association of pentaresistance of the AmCmStSuTe type with DT104. We further found, however, that the majority of pentaresistant DT104 isolates (92 of 93 tested) were of the Copenhagen variant of serovar Typhimurium. It is difficult to assess whether DT104 is universally associated with the Copenhagen variant since serotyping schemes in past studies have often not differentiated between serovar Typhimurium and serovar Typhimurium var. Copenhagen. It is clear, however, that there is an association between host species and some *Salmonella* serovars and that serovar Typhimurium var. Copenhagen is adapted to the swine host (36). It is also known that swine-associated phage types have been reported to be common causes of human salmonellosis (4, 35, 45). Thus, our observations that serovar Typhimurium var. Copenhagen was common among pigs and was associated with phage type DT104 suggest that contaminated pork products present a substantial risk for the acquisition of strains with this type of multiresistance by humans. In addition to DT104, 11 of 14 phage type U302 isolates also exhibited pentaresistance of the AmCmStSuTe type. Genetic characterization also revealed that these U302 isolates carried the same resistance genes that DT104 carries (Table 2), a finding consistent with recent reports (33, 42). Although this phage type is infrequently isolated in most present studies, its close relationship to DT104 may signify its potential public health importance. Its genetic architecture also presents a challenge for the identification of DT104 by using amplification of resistance alleles (9), since the two phage types encode identical resistance genes.

In addition to the large number of DT104 isolates cultured in this study, we also identified phage types with pentaresistance of the AmKmStSuTe type. This pattern was found predominantly in two phage types, DT193 (77 of 165 isolates) and DT21 (82 of 165 isolates). As was the case for DT104, there was a clear association of phage type with serovar; the great majority of DT193 isolates with pentaresistance of the AmKmStSuTe type were serovar Typhimurium var. Copenhagen, while most DT21 isolates were serovar Typhimurium. The spread of MDR DT193 was reported as early as 1978 in studies by Threlfall et al. (38) in bovine and human isolates, which showed that phage type conversion of MDR DT204 by the acquisition of resistance factors of specific compatibility group I2 was important in the emergence and expansion of phage type DT193. Further characterization of DT193 in several studies showed that this phage type usually exhibits the AmStSuTe or Te resistance pattern (3, 11, 17, 18). However,
we consistently found DT193 isolates with the AmKmStSuTe pentaresistance pattern, which was uncommon in previous studies. Rarely, isolates from our study also carried resistance to gentamicin. These findings thus imply that DT193 is extending its spectrum of multiresistance.

We performed DNA fingerprinting using AFLP analysis in order to discern genetic similarities among a representative sample of the serovar Typhimurium var. Copenhagen isolates obtained in this study. This analysis divided the isolates into two groups (clusters 1 and 2; Fig. 1) composed primarily of phage types DT193 and DT104 with the AmKmStSuTe and AmCmStSuTe resistance patterns, respectively, with 90 to 100% similarity within each group. The similarity between and within groups was expected, since most isolates of serovar Typhimurium have been shown to be genetically very similar (5). Our finding is also consistent with those of earlier studies that showed that diversity among DT193 strains was based upon differences in plasmid compositions (17). DT104 has also been proposed to have originated as a single clone which spread epidemiologically (10, 20), predicting similar fingerprints among isolates. The heterogeneity that we found in each cluster is presumably due to recent genetic events such as deletions, insertions, or mutations of one or a few loci or by the acquisition of plasmids. The identical fingerprints between pentaresistant and StSu DT104 isolates suggest that these isolates are genotypically closely related and that the loss of resistance genes is likely to be a recent genetic event (discussed further below). We did find, however, that a few isolates were genetically more similar to members of the group dominated by the opposite serovar and the opposite resistance pattern (for example, strains UU01 and FV06 have the AmCmStSuTe pattern and are phage types DT104 and U302, respectively, but are similar to strains with the AmKmStSuTe pattern, while U002 and UF02 are phage types DT193 and DT21, respectively, but map to the cluster with the AmCmStSuTe pattern). There is evidence that subtle changes in the bacterial genome insufficiency to change the fingerprint pattern may cause a change in phage type (44), thus suggesting the need for phenotypic methods complementary to genotypic fingerprinting for the ultimate discrimination of strains.

We characterized the molecular basis of resistance of the DT193 isolates in this study both because this phage type represents a significant cause of food-borne disease and because we had identified the AmKmStSuTe resistance pattern in many of these isolates, a pattern similar to that for pentaresistant DT104 isolates and previously not commonly reported among DT193 isolates. We chose for study four isolates that were both phenotypically and genotypically similar, as well as epidemiologically related, in order to investigate the means by which differences in antibiotic resistance can be encoded in similar strains. We found that all resistance genes were encoded on conjugative plasmids that could be efficiently transferred to E. coli. It has long been recognized that antimicrobial resistance genes among Salmonella serovars can be carried on conjugative plasmids; however, we isolated plasmids of different sizes and numbers and with different resistance gene contents from strains collected from the same location and time, strengthening the evidence of genetic heterogeneity of epidemiologically related Salmonella serovar Typhimurium isolates. We found, for example, that the addition of gentamicin resistance in one strain was due to the acquisition of a second conjugative plasmid that also encoded duplicative resistance to ampicillin, kanamycin, streptomycin, and sulfamethoxazole. Gentamicin resistance was encoded by a homologue of grm of M. purpurea, a gene that, to our knowledge, has not previously been found in pathogenic bacteria. We also found this plasmid to have a class I integron harboring aadA, which encodes streptomycin resistance, and sul1, which encodes sulfamethoxazole resistance and which is commonly found on class I integrons. It is unclear whether the duplication of resistance in this strain provided any selective advantage (for example, aadA also encodes resistance to spectinomycin and so might broaden the spectrum of resistance). The presence of an integron on this plasmid is of particular concern. Integrons are capable of trapping one or more resistance gene cassettes and furnishing a promoter for the efficient expression of antimicrobial resistance genes (37), leading to the rapid evolution of multiresistance. Salmonella strains carrying this plasmid might thus acquire new resistance factors and might easily transfer those resistance genes within and between species by means of conjugation.

In addition to the many AmCmStSuTe pentaresistant DT104 isolates cultured in this study, we also found atypical DT104 strains with only the StSu resistance pattern, a pattern that was observed in 8.3% of DT104 isolates. Similar atypical DT104 strains have also been reported previously (11, 34); however, none of those reports examined the epidemiological significance of these strains or performed detailed genetic characterization. We detected these StSu isolates predominantly from environmental samples consisting of drag swabs collected from cleaned and apparently disinfected barns before they were filled with new groups of pigs. As expected from their phenotype, PCR analysis of isolates with the StSu resistance phenotype showed that they did not carry genes encoding resistance to ampicillin, chloramphenicol, and tetracycline. We also found that, in contrast to the 1.0- and 1.2-kb integrons of pentaresistant DT104, isolates with the StSu phenotype had only one integron of 1.0 kb. We hypothesized that pentaresistant DT104 isolates excise a chromosomal fragment encoding ampicillin, chloramphenicol, and tetracycline resistance, including the 1.2-kb integron, by homologous recombination in the presence or absence of some unidentified selective pressure. Since most of the StSu DT104 isolates were isolated from cleaned barns, it is possible that the loss of this chromosomal piece provides a selective advantage for survival during the cleaning and disinfecting procedures. Alternatively, expression of all five resistance factors may reduce the competitive fitness of Salmonella strains when growing outside of the animal host, in the absence of antimicrobial agents, thus allowing the selective proliferation of strains that have reduced their levels of expression of resistance genes by deleting them.

The results of this study have important implications for public health. We found MDR Salmonella serovar Typhimurium isolates to be common on commercial swine farms. A frequent isolate was phage type DT193, which had the AmKmStSuTe resistance phenotype and which is classified as one of the phage types of public health importance. We have shown here that DT193 isolates encode the genes for resistance to all five antimicrobials on plasmids that can easily be transferred by conjugation. This poses a significant threat not
only through the food-borne salmonellosis attributed to pork consumption but also through the horizontal transfer of resistance determinants to commensal organisms of the human digestive tract or other pathogens of importance to human health. Studies on the epidemiological significance of this panterturessant drug resistance phenotype on human health are under way. The high prevalence of phage type DT104, which has been known to cause outbreaks globally for the last decade, poses an additional threat through the food-borne acquisition of this panterturessant strain.

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