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# Clostridium difficile in Retail Meat Products, USA, 2007

J. Glenn Songer, Hien T. Trinh, George E. Killgore, Angela D. Thompson, L. Clifford McDonald, and Brandi M. Limbago

To determine the presence of *Clostridium difficile*, we sampled cooked and uncooked meat products sold in Tucson, Arizona. Forty-two percent contained toxigenic *C. difficile* strains (either ribotype 078/toxinotype V [73%] or 027/toxinotype III [NAP1 or NAP1-related; 27%]). These findings indicate that food products may play a role in interspecies *C. difficile* transmission.

The incidence and severity of *Clostridium difficile* infections (CDIs) are increasing in North America (*I*), probably because of emergence of an epidemic strain (NAP1/BI/027, toxinotype [TT] III) (2,3). *C. difficile* transmission occurs primarily in healthcare facilities, but community-associated CDI (CA-CDI) appears to be increasing and may now account for 20%–45% of positive diagnostic assay results (4,5). Up to 35% of patients with CA-CDI report no antimicrobial agent use within 3 months before disease onset (4,5), although nonantimicrobial drugs (e.g., proton pump inhibitors, nonsteroidal antiinflammatory agents) are also implicated as risk factors (4). Sources of *C. difficile* acquisition in community settings are unknown.

CDI is increasingly important in food animals (6). Infection rates of >95% have been documented among neonatal pigs in farrowing facilities, resulting in diarrhea and typhlocolitis (6). Toxigenic *C. difficile* is also implicated as a cause of diarrhea in calves (7). *C. difficile* was identified in raw meat intended for pet consumption (8) and in  $\approx$ 20% of retail ground beef in Canada (9). We report the isolation of *C. difficile* from uncooked and ready-to-eat meats in retail markets in a US metropolitan area.

### The Study

Packaged meats were purchased from 3 national-chain grocery stores in the Tucson, Arizona, area on 3 occasions at 1-month intervals from January to April 2007. Prod-

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ucts sampled were both uncooked (ground beef, ground pork, ground turkey, pork sausage, and pork chorizo) and ready to eat (beef summer sausage, pork braunschweiger) (Table). Pork chorizo was produced and distributed locally; all other samples were national brands. Products with different sell-by dates (a surrogate for production date) were sampled for each meat type. Samples were not representative of all meat products in each grocery store.

For each sample, 1 g of meat was added to two 10-mL tubes of prereduced brain heart infusion (BD, Franklin Lakes, NJ, USA), which had been supplemented with 0.5% yeast extract (BD), 0.05% DL-cysteine (Sigma-Aldrich, St. Louis, MO, USA), and 0.1% taurocholate (MP Biomedicals, Solon, OH, USA). One tube was heat shocked (80°C, 10 min), and both were then incubated anaerobically at 37°C for 72 h. Aliquots were subcultured onto taurocholate cycloserine cefoxitin fructose agar (TCCFA) (10) and incubated anaerobically for 24–72 h at 37°C. Colonies were subcultured onto anaerobic blood agar, TCCFA (with or without antimicrobial agents), and confirmed as *C. difficile* by *p*-cresol odor, yellow-green fluorescence under UV illumination, a positive L-proline aminopeptidase reaction, and negative indole reaction.

Isolates were characterized by PCR ribotyping (11), toxinotyping (3), and pulsed-field gel electrophoresis (PFGE) (12). Presence of tcdA, tcdB, cdtB (binary toxin), and deletions in tcdC was determined by PCR (2).

MICs were determined by Etest (AB Biodisk, Solna, Sweden) on *Brucella* blood agar with vitamin K and hemin (Remel, Lenexa, KS, USA) that was incubated anaerobically at 35°C. Reference interpretive criteria for *C. difficile* susceptibility to clindamycin and moxifloxacin were used; MICs for levofloxacin and gatifloxacin were interpreted by using criteria for moxifloxacin (*13*). *Bacteroides fragilis* ATCC 25285, *B. thetaiotaomicron* ATCC 29741, *C. difficile* ATCC 700057, and *Enterococcus faecalis* ATCC 29212 were included as controls.

Proportions were compared by  $\chi^2$  or Fisher exact test. Thirty-seven (42.0%) of 88 retail meats yielded *C. difficile*, including 42.4% of beef, 41.3% of pork, and 44.4% of turkey products (Table). Ready-to-eat products were more commonly culture positive (11/23; 47.8%) than were uncooked meats (26/65; 40.0%), although the difference was not significant (p = 0.34). The highest percentages of C. difficile isolates were recovered from pork braunschweiger (62.5%) and ground beef (50.0%). Culture-positive results came from both heat-shocked and non-heat-shocked cultures, whereas culture-negative specimens were negative in both types of culture, and no specimen was positive by both methods (not shown). No association was found with the meat processor, the sell-by date, the store, or the month sampled (not shown). Multiple independent cultures from 2 braunschweiger samples vielded indistinguishable isolates

Table. Source and characteristics of Clostridium difficile isolates obtained from retail meats sold in Tuscon, Arizona, USA, 2007\*

Meat product	No. samples cultured	Total no. (%) positive	Ribotype	Toxinotype	∆tcdC, bp†	PFGE type	No. (%) pos
Ground beef (uncooked)	26	13 (50)	027	Ш	18	NAP1	1 (3.8)
						NAP1-related	2 (7.7)
			078	V	39	NAP7	8 (30.8)
						NAP8	2 (7.7)
Summer sausage (ready to eat)	7	1 (14.3)	027	Ш	18	NAP1	1 (14.3)
Ground pork (uncooked)	7	3 (42.9)	027	III	18	NAP1-related	1 (14.3)
			078	V	39	NAP7	2 (28.6)
Braunschweiger (ready to eat)	16	10 (62.5)	027	Ш	18	NAP1	2 (12.5)
						NAP1-related	1 (6.2)
			078	V	39	NAP7	7 (43.8)
Chorizo (uncooked)	10	3 (30.0)	027	Ш	18	NAP1-related	1 (10.0)
			078	V	39	NAP7	2 (20.0)
Pork sausage (uncooked)	13	3 (23.1)	027	III	18	NAP1-related	1 (7.7)
			078	V	39	NAP7	2 (15.4)
Ground turkey (uncooked)	9	4 (44.4)	078	V	39	NAP7	4 (44.4)
Totals	88	37 (42.0)	027	III	18	NAP1	4 (4.4)
						NAP1-related	6 (6.7)
			078	V	39	NAP7	25 (27.8
						NAP8	2 (2.2)

<sup>\*</sup>All samples were positive for *cdtB*, which encodes the binding component of binary toxin. PFGE, pulsed-field gel electrophoresis. †Deletions in *tcdC* regulatory gene.

in the same meat sample (10/10 from 1 package and 12/12 from another; not shown), which suggests that a single strain may predominate when *C. difficile* is present. Our percentage of recovery of *C. difficile* from retail meat products is higher than that reported (20%) in a similar study of Canadian ground beef (9), possibly because of differences in culture methods, the meats sampled, or national or geographic variation.

Isolates were grouped into ribotype 078/TT V (27/37, 73.0%) and ribotype 027/TT III (10/37, 27.0%). Strain types were not specific to meat type, store, or sampling month (Table). All isolates were PCR positive for binary toxin (*cdtB*), *tcdA*, and *tcdB*. Characteristic 18-bp and 39-bp deletions in *tcdC* were present in 027/TT III and 078/TT V isolates, respectively (2,12). PFGE divided 027/TT III isolates into NAP1 (≥80% related to human NAP1) and NAP1-related (78% related to human NAP1) groups and 078/TT V isolates into NAP7 and NAP8 groups (Figure).

Ribotype 027 isolates are described almost exclusively in context of the current human epidemic strain, NAP1/027/TT III (2). In this study, we also found 027/TT III isolates that were only 78% similar to NAP1 (i.e., NAP1-related). Ribotype 078 strains were previously uncommon causes of healthcare-associated CDI in humans (12), but now they are emerging in pigs and calves with diarrhea (7; J.S. Weese, pers. comm.) and in persons with CDI (12). Two epidemiologically unrelated 078/TT V isolates from human CDI patients are indistinguishable by PFGE from pig isolates (12).

The 078/TT V isolates were uniformly susceptible to levofloxacin, moxifloxacin, and gatifloxacin. Like human

TT V isolates (12), most 078/TT V meat isolates were nonsusceptible to clindamycin (56% resistant, 41% intermediate). This may not be surprising given the widespread use of tylosin, erythromycin, virginiamycin, and lincomycin in food animals and the potential for selection of macrolidelincosamide-streptogramin resistance (14).

NAP1 isolates have demonstrated high-level resistance to levofloxacin, moxifloxacin, gatifloxacin (>32  $\mu$ g/mL), and clindamycin (>256  $\mu$ g/mL), consistent with current human strains (2). NAP1-related isolates were susceptible to levofloxacin, moxifloxacin, and gatifloxacin but resistant to clindamycin, similar to the pattern of historic NAP1 strains (2).

### **Conclusions**

Fluoroquinolones are widely used in human therapy, and the current epidemic strain may have emerged because of its resistance to these agents. Fluoroquinolone use is limited in food animal production (14), with the exception of enrofloxacin for treatment of bovine respiratory disease (now approved for use in swine).

The source of *C. difficile* in retail meats may involve antemortem deposition of spores in the animal's muscle or other tissues, fecal or environmental contamination of carcasses, or contamination during processing. Spores could persist in packing plants, resulting in contamination of carcasses or food products during processing. Contamination may also occur in retail meat markets.

Direct or indirect human-to-human transmission is responsible for most healthcare-related CDIs (15) and most likely contributes to CA-CDI. Therefore, stopping such

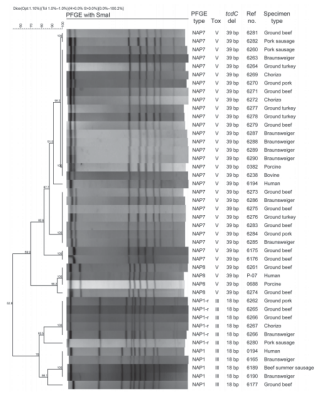


Figure. Origin, NAP types, and relatedness of strains from foods and humans, Arizona, USA, 2007. All strains were positive by PCR for binary toxin. Scale bar indicates genetic relatedness. Tox, toxinotype; Ref, reference; NAP1-r, NAP1-related.

transmission remains the critical control point for preventing most human CDIs. Nonetheless, our findings highlight the potential both for selection of virulent or resistant strains in animals and interspecies transmission through the food supply. Our data do not prove transmission of *C. difficile* from foods to humans but highlight the need for studies to characterize risks posed by this organism in the human food supply.

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Dr Songer is professor of Veterinary Science and Microbiology at the University of Arizona. His research interests focus on bacterial diseases of food animals, mainly those affecting the gastrointestinal tract.

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