Food Safety and Food Access: A Pilot Study

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Abstract (WC 238)

Objectives: This study was designed to evaluate the feasibility of testing the hypothesis that differences in neighborhood level food access may be associated with consumer exposure to food borne microbial contamination.

Methods: This study was carried out in Baltimore MD in 2011 among selected neighborhoods defined as high or low food access. In each category, packages of chicken thighs and ground beef were purchased from small stores and supermarkets. We evaluated presence of E. coli and Staphylococcus aureus and also tested isolates for antimicrobial resistance.

Results: Microbial contamination of both chicken and beef products was highly prevalent (S. aureus-13/32 for chicken and 14/32 for beef; E. coli 21/32 for chicken and 12/32 for beef). Small stores were more likely to sell food carrying these microbes as well as MDR strains of both E. coli and S. aureus, and chicken was more likely to carry E. coli as compared to ground beef.

Conclusions: This is the first study of this hypothesis. While it is limited in size and in focus on one US city, the results indicate that further research is appropriate to examine neighborhood level risk factors for differential exposures to food borne microbes.

Abbreviations: S. aureus (Staphylococcus aureus); E. coli (Escherichia coli), MRSA (methicillin resistant Staphylococcus aureus), MDR (multi-drug resistant), LFA (low food access), HFA (high food access). FDA (US Food and Drug Administration), USDA (US Department of Agriculture), CLSI (Clinical and Laboratory Standards Institute).

Keywords: food safety, food access, food borne illness, Staphylococcus aureus, Escherichia coli, MRSA

(WC = 3656)

1. Introduction

Food access is increasingly recognized as a public health issue (NAS, 2011). It is a community/neighborhood level concept that refers to the availability, affordability and ease of getting to sources of food (Ver Ploeg & Breneman, 2009) including the number and types of different food stores and other sources that are accessible within a given community, as well as the types and costs of foods available at these stores (Drewnowski, 2004; Powell et al., 2007b; Casagrande et al., 2011). Differences in food access have been found to contribute to both food insecurity and to health disparities among socioeconomic and ethnic groups in terms of both poorer diets and increased risks of adverse health outcomes including obesity, diabetes, and cardiovascular disease (Drewnowski, 2009; Coleman-Jensen et al., 2011; NAS, 2011).

Many studies have demonstrated that food access differs in many countries, including Canada and the US by ethnicity and socioeconomic status (SES) with the lowest access reported in low-income, urban minority neighborhoods and American Indian communities (Lewis et al. 2005; Powell et al., 2007a; Larson et al., 2009; Cerin et al., 2011; O’Connell et al., 2011; Fuller et al., 2013; Vahabi & Damba, 2013) and among food stamp recipients (Oberholser & Tuttle 2004; D’Angelo et al. 2011). These factors may differ in terms of importance in different cities and countries (Boone-Heinonen et al., 2011).
Very few studies have examined the possible relationship at the neighborhood level between food access and microbial food safety despite shared sociodemographic risk factors related to both access and food borne infections (Wallace et al., 2000; Hardnett et al., 2004). Koro et al. (2010) and Signs et al. (2011) reported that ready to eat foods and fresh produce purchased in stores in low SES areas of Philadelphia had higher plate counts of yeast, mold, and bacteria (total aerobes, anaerobes, and coliforms) as compared to the same items purchased in higher SES areas. A national level analysis of listeriosis in the UK reported that incidence of diagnosed listeriosis increased with census-based indicators of neighborhood deprivation and that listeriosis cases frequently reported use of local food stores as compared to supermarkets (Gillespie et al., 2010).

No studies have directly examined differences in the prevalence of specific food borne bacteria, including pathogens, or of antimicrobial resistance in bacteria isolated from consumer products available in neighborhoods that differ in terms of food access. Studies of food borne bacteria and pathogens in food purchased at retail stores rarely identify the neighborhood source of items analyzed for microbial carriage. Exposures to antimicrobial resistant bacteria on meat and poultry products is an important aspect of food safety et al. (2011b). At the national level, this is driven by producer decisions to utilize subtherapeutic levels of antimicrobials in animal feeds (Angulod et al., 2004; Silbergeld et al., 2008). At the local level differences in frequency of health inspections at food stores and food service outlets may also be important (Darcey & Quinlan 2011). Practices at the store or outlet level can also affect food safety, such as inadequate refrigeration, stocking items past sell by dates, and practices in meat grinding or cutting (FDA, 2009). At the household level, consumer behaviors can enhance risks through lack of information and food safety practices (Fein et al., 2011) as well as cultural traditions of preparation that result in incomplete cooking (Anderson Steeves et al., 2012).

The primary objective of this study was to carry out a pilot study to test the hypothesis that risks of exposure to food borne bacteria and pathogens, and to drug resistant strains, are increased for consumers in neighborhoods with low food access as compared to those with high food access. Because this issue has not been previously studied, a pilot study is an appropriate first step for determining feasibility and likely sample size required for a more rigorous test of this hypothesis. In other studies we have examined household level factors such as food safety knowledge and practices in food selection, handling, preparation and storage (Anderson Steeves et al., 2012).

2. Methods

2.1 Study Site

The study was carried out in the Baltimore MD (USA) metropolitan area among neighborhoods stratified as high food access (HFA) or low food access (LFA) on the basis of measured characteristics such as SES, housing status, presence or absence of large retail food stores and healthy food availability at these stores. This analysis was separately carried out by Casagrande et al. (2011) prior to our study and the results are mapped in Figure 1a. Standard metrics, including census data on socioeconomic variables as well as assessment of stores in terms of size, numbers of employees, and availability of fresh and healthy foods were utilized in this assessment (Casagrande et al., 2011; O'Connell et al., 2011). From these neighborhoods, we selected those that contained both supermarkets and small grocery stores. Stores classified as “convenience stores” were not included in this study since a preliminary survey by us indicated that none of these outlets carried raw poultry or ground beef products, which were the focus of this study. To reduce other potential sources of variability, all selected neighborhoods were defined by the US Census (2000) as mixed in terms of race, that is, neighborhoods in which no one race or ethnicity accounted for more than 59% of the population. A map of store locations is shown in Figure 1b. Most stores were located within the city (Figure 1b), and those stores outside the city limits were considered to be within neighborhoods that extended into the city based on resident surveys (Casagrande et al., 2011; O'Connell et al., 2011).
Figure 1a. Food map of Baltimore City (from Johns Hopkins Center for a Livable Future (www.mdfoodsystemmap.org)
Initially, we randomly selected four stores in each neighborhood, two of which represented smaller stores (selling meat and poultry products) and two of which represented supermarkets (O’Connell et al., 2011). This design was modified by expediency, as many of the LFA areas originally selected did not contain both store types at the time of our survey initiation in 2011. The resulting sample was still drawn from our randomized design. From each outlet, we collected 2 packages each of poultry (thighs) and ground beef over a period of three months, from February to May 2011. For those stores sampled more than once, sampling from the same store was carried out within 7 to 12 weeks of the first visit. We analyzed poultry and ground beef for the following reasons: these are major items in the consumer food basket; they are major components of exposure to food borne pathogens in the US (Zhao et al., 2012); and they represent different producer and retail risk factors relevant to pathogen carriage. That is, in many cases, poultry products are packaged at the processing plant, while ground beef is often prepared at the store or another post-primary processing plant level.

2.2 Food Sampling

All food items were unfrozen at the time of purchase. Immediately after purchase the food items were placed in plastic grocery store bags, and transported in coolers without ice to the laboratory within 1 h. Since no quantitation of microbial contamination was planned, changes in bacterial numbers did not affect our study. At the time of purchase, we noted the integrity of packaging, whether the product was wrapped in heavy heat sealed plastic (indicative of packaging at the processing plant) or in thinner material (indicative of post processing rewrap), the presence of information on expiration date, safe food handling and cooking, and the presence of tracer code information. Because no food items labeled as “organic” or “antibiotic free” were available in stores sampled from LFA areas, we only sampled conventionally produced items at all stores. At the lab, each package was labeled with the store of purchase, and then immediately placed in a refrigerator (4°C) in their original
packaging until processing. At the time of processing, each package was coded and then photographed to record package information.

2.3 Microbiological Analyses

All microbiological analyses were conducted on coded samples to mask identification of source. Analyses were initiated between 10 min and 3 h after receipt of samples (time differences related to numbers of samples collected on a given day). We evaluated the presence of two bacterial species, *E. coli* and *Staphylococcus aureus*. In other studies, we have examined carriage of *Campylobacter jejuni* in poultry (Price et al., 2005, 2007). We selected these species since our goal in this study was to utilize the same microbial indicators for both beef and poultry. These two species were selected for different reasons: *E. coli* are a nearly omnipresent reservoir strain for resistance. We did not define the strain of *E. coli* (with respect to specific pathogenic strains). This method is used by FDA in its surveys of bacterial contamination of retail meats (Zhao et al., 2012). *S. aureus* (including MRSA) is an increasingly documented food borne disease risk reported to be carried by both beef and poultry products (Hanson et al., 2011; Kelman et al., 2011; Waters et al., 2011).

All packages were opened under sterile conditions and food items were placed in stomacher bags with appropriate broth. Our laboratory methods for *E. coli* are similar to those of Zhao et al. (2012) and as described in Price (Price et al., 2007); methods for *Staphylococcus sp* are described in Waters et al. (2011). Briefly, we utilized standard culture methods in which one loop of broth was streaked on appropriate plates and one well-defined colony was selected for further culture, and resistance testing as well as confirmation with PCR (*S. aureus*) or biochemical testing by indole (*E. coli*). We tested for antimicrobial resistance using standard disk diffusion assays (CLSI, 2009, M100-S19 http://www.clsi.org), for *E. coli* isolates: Amikacin (AMK), Ampicillin (AMP), Amoxicillin/Clavulanic acid (AmC), Cefoxitin (CFX) (recommended for methicillin), Ceftriaxone (AXO), Chloramphenicol (CHL), Ciprofloxacin (CIP), Fosfomycin (FOS), Gentamicin (GEN), Imipenem (IMP), Kanamycin (KAN), Nalidixic acid (NAL), Streptomycin (STR), Sulfamethoxazole/Trimethoprim (SXT), and Tetracycline (TET); and for *S. aureus* isolates: Ampicillin (AMP), Ceftriaxone (AXO), Chloramphenicol (CHL), Ciprofloxacin (CIP), Clindamycin (CLI), Gatifloxacin (GAT), Gentamicin (GEN), Kanamycin (KAN), Levofloxacin (LEVO), Linezolid (LZD), Nitrofurantoin (NIT), Oxacinil (OXA), Penicillin (PEN), Quinupristin/Dalfopristin (SYN), Rifampin (RIF), Sulfamethoxazole/Trimethoprim (SXT), Tetracycline (TET), and Vancomycin (VAN). This list includes antimicrobials from drug classes utilized in poultry and beef production (Zhao et al., 2006; Silbergeld et al., 2008) Reference strains were included at all steps to ensure validity of results: *Escherichia coli* ATCC 25922; *Staphylococcus aureus* subsp aureus ATCC 25923; and for MRSA *Staphylococcus aureus* subsp aureus ATCC 43300.

2.4 Definition of Resistance Phenotype

In reporting the results of resistance testing, we utilized both standard CLSI nomenclature of “susceptible”, “intermediate”, and “fully resistant”, and we also utilized the dichotomous definition of “susceptible” or “nonsusceptible” (to include both intermediate and full resistance phenotypes) as recommended by Magiorakos et al. (2011) This a consensus report proposes this nomenclature for classifying the results of *in vitro* testing for purposes of epidemiological and other research, as distinct from testing for clinical purposes, because of different definitions in the literature. This paper also recommends use of the term “MDR” to denote nonsusceptibility to three or more classes of antimicrobial agents.

3. Results

3.1 Summary

In total, we collected 32 samples of chicken and 32 samples of ground beef from 6 stores in HFA neighborhoods and 3 stores in LFA neighborhoods. In HFA neighborhoods we sampled 2 small stores (both twice) and 4 large stores (2 twice, 2 once); in LFA neighborhoods we were able to sample only one large store (twice) and 2 small stores (both twice) due to the lack of retail outlets selling chicken or ground beef in LFA neighborhoods, contrary to the database collected between 1-2 years prior to our study. Repeated sampling was not undertaken to produce replicable results, since shipments of food products occurred throughout the intervening period between sampling; thus each visit was considered as an independent event. In terms of packaging, ground beef samples were more likely to be wrapped loosely (22/32) as compared to chicken (6/32). Most of the heat wrapped ground beef packages came from stores in HFA neighborhoods (one store in a LFA neighborhood from HFA stores sold heat wrapped ground beef) and most of the loosely wrapped chicken packages came from small stores in LFA neighborhoods. All packages contained information on expiration date, safe food handling, preparation, and storage. No package had information relevant to tracking.
3.2 Pathogen Prevalence

The microbiological findings are reported in Table 1.

Table 1. Overall prevalence of bacterial carriage by chicken and ground beef (note that susceptibility testing was only conducted on those items carrying either SA or EC, and multi drug resistance was only tested in those isolates that were nonsusceptible)

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Chicken (n = 32)</th>
<th>Ground Beef (n = 32)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>S. aureus</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-susceptible</td>
<td>9/13 (69%)</td>
<td>11/14 (79%)</td>
</tr>
<tr>
<td>Multi-drug resistant</td>
<td>2/9 (22%)</td>
<td>4/11 (36%)</td>
</tr>
<tr>
<td><strong>E. coli</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-susceptible</td>
<td>17/21 (81%)</td>
<td>7/12 (58%)</td>
</tr>
<tr>
<td>Multi-drug resistant</td>
<td>8/17 (47%)</td>
<td>0/7 (0%)</td>
</tr>
</tbody>
</table>

This table presents data on SA prevalence in all samples, and then on drug resistance in those samples testing positive for SA; thus, the denominators change with the category since not all samples tested positive for SA. In the overall sample set, contamination of both chicken and beef products by *S. aureus* was prevalent (13/32 or 40% for chicken and 14/32 or 44% for beef), while prevalence of *E. coli* was higher in chicken (21/32, 66%) as compared to beef (12/32, 38%). Using the definitions of Magiorakos et al. (2011), there was no difference between chicken and beef in carriage of nonsusceptible or multidrug resistant *S. aureus* (that is, including both intermediate and full resistance in the definition of non-susceptible and for MDRSA). Chicken was more likely to carry nonsusceptible and multidrug resistant strains of *E. coli* as compared to beef. Overall there was a highly significant association between carriage of *S. aureus* and *E. coli* (both positive and negative) (Pearson chi square 0.004 odds ratio).

Table 2. Prevalence of bacterial carriage by neighborhood and store size. Figures in parentheses are percentages of the preceding row

<table>
<thead>
<tr>
<th>Neighboring</th>
<th>LFA</th>
<th>HFA</th>
<th>LFA</th>
<th>HFA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Store Size</td>
<td>Small</td>
<td>Large</td>
<td>Small</td>
<td>Large</td>
</tr>
<tr>
<td><strong>S. Aureus</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-susceptible</td>
<td>1 (50)</td>
<td>3 (75)</td>
<td>5 (71)</td>
<td></td>
</tr>
<tr>
<td>Multi Drug Resistant</td>
<td>0 (0)</td>
<td>1 (33)</td>
<td>1 (20)</td>
<td></td>
</tr>
<tr>
<td><strong>E. Coli</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Non-susceptible</td>
<td>3 (75)</td>
<td>4 (66)</td>
<td>5 (83)</td>
<td>5 (100)</td>
</tr>
<tr>
<td>Multi Drug Resistant</td>
<td>2 (66)</td>
<td>1 (25)</td>
<td>5 (100)</td>
<td>0 (0)</td>
</tr>
</tbody>
</table>

Table 2 presents data on microbial contamination by neighborhood (HFA or LFA) and by store size (large or small). Given the challenges in locating equal numbers of both store types in both neighborhood types, we did not attain a sufficient sample size for statistical analysis. The results suggest that store size was more likely to be associated with microbial contamination by nonsusceptible strains of *E. coli* for chicken, with small stores more
likely to carry MDR strains, while for ground beef, there was a suggestion of more prevalent carriage of nonsusceptible strains of \textit{E. coli} in samples from small stores.

3.3 Antimicrobial Resistance

All SA isolates were tested for antimicrobial resistance and, in the case of MRSA, we used PCR to confirm phenotypic results by detection of the \textit{mecA} gene. MDR was defined as a phenotype that was nonsusceptible to three or more drugs (Magoriakis et al., 2011). Results are presented in Figure 2 by food product, neighborhood, and store size, for both \textit{E. coli} and \textit{S. aureus} in three categories: susceptible, intermediate, and resistant.
We analyzed the data in two ways: as two categories of susceptible/nonsusceptible (including intermediate and resistant phenotypes), and as three categories of susceptible/intermediate/fully resistant. As shown in Table 2, there were very few isolates resistant to oxacillin (n = 4 equally distributed by store type and neighborhood) and none of these tested positive for the mecA gene by PCR. As can be seen from these analyses, chicken and ground beef products from small stores in both HFA and LFA neighborhoods were more likely to carry MDR S. aureus. Similarly, small stores in both LFA and HFA neighborhoods were more likely to carry MDR E. coli and small stores in HFA neighborhoods were more likely to carry MDR E. coli as compared to large stores in the same neighborhoods. All isolates were susceptible to vancomycin. Since no isolates were found to be nonsusceptible by disk diffusion assays, we did not carry out further analyses to confirm resistance.

In terms of full resistance, we observed a suggested pattern of greater likelihood of full resistance to the antimicrobials tested for both S. aureus and E. coli isolates from chicken and ground beef in small as compared to large stores and in small stores in LFA as compared to HFA neighborhoods.

The most commonly observed nonsusceptibility phenotypes (observed in 3 or more isolates) for S. aureus in chicken were ampicillin and penicillin, and in beef ampicillin, gatifloxacin and penicillin. For E. coli the most common nonsusceptibility phenotypes were ampicillin and amoxicillin/clavulanic acid, gentamycin, streptomycin and tetracycline in chicken, and streptomycin and tetracycline in beef.

4. Discussion

This is the first study on prevalence of microbial contamination of poultry and ground beef products in stores located in neighborhoods differing by food access, including specific pathogen identification and antimicrobial resistance profiles. It complements research by Signs et al. (2011) who assessed nonspecific microbial contamination of ready-to-eat foods as distinct from our study of uncooked consumer meat and poultry products.
While the sample size was small, some trends are noted. The data from this pilot study suggest that size of store was a factor in the likelihood of microbial contamination and that small stores in LFA neighborhoods appeared to be the most likely to carry fully resistant isolates of both \textit{S. aureus} and \textit{E. coli}.

The prevalence of microbial contamination in this sample of urban stores from the Baltimore metropolitan region was relatively high although not dissimilar from some recent studies. Our results cannot be compared to these studies since we did not attempt quantitative analyses of microbial contamination, since this would be expected to vary with these conditions.

Interestingly, we observed marked decreases in the prevalence of full resistance to ciprofloxacin in isolates from chicken as compared to our studies in 2004 and 2006 conducted before and after the FDA banned the use of ciprofloxacin in poultry feed (Price et al., 2007). In 2006, we found average prevalence of 26% for full resistance to CIP in \textit{Campylobacter jejuni} isolates from conventionally raised chicken products in this study, we found no \textit{S. aureus} or \textit{E. coli} isolates that were fully resistant to CIP and only four \textit{S. aureus} isolates were intermediate resistant to CIP. This is similar to a recent report on \textit{Campylobacter} from Canada (Deckert, Valdivieso-Garcia et al., 2010) but lower than that reported in a recent study on \textit{S. aureus} carriage in the US (Waters et al., 2011). Other observations from this study indicate concerns requiring further investigation. For the first time, we investigated the associations between carriage of \textit{E. coli} and \textit{S. aureus}. There was a highly significant correlation between both positivity and negativity of carriage. This information is relevant to assessing overall health risks as well as designing effective interventions to reduce consumer exposure. The prevalence of loose wrapping of ground beef may be a preventable risk for food contamination. Finally, the absence of label information on producer origin may impede trace backs through the production chain from consumers and retail level to processors and producers.

Overall, this study must be considered as exploratory as it is the first study designed to test associations between food access and food safety. The study is limited by its final sample size and also by lack of information on specific strain level data for \textit{E. coli}. Our decision not to quantify bacterial presence was related to the lack of information available on factors in the overall food production system, including conditions at the retail outlets from which samples were purchased. Our decision to study \textit{E. coli} without specification of pathogenic strains was driven by a similar strategy utilized in the most recent large survey of consumer food products reported by the FDA (Zhao et al., 2012). This, along with data on actual food purchases and consumption within these neighborhoods, limit these results in terms of inferences for health risks. We also decided not to quantify bacterial contamination in this study since we did not have information on the stages of production from farm through primary and secondary processing owing to lack of label information. Moreover, we were not able to obtain data on store conditions, such as temperature and length of holding, which are key factors in microbial food safety. For these reasons, we did not attempt quantitative analyses of microbial contamination, since this would be expected to vary with these conditions.

The study demonstrated challenges to such research as well as the necessity to carry out a larger project to investigate food safety related to differing conditions of food access. Our decision to design a neighborhood based, rather than city-based, study of this hypothesis resulted in selection of neighborhoods with fewer food stores than had been reported in an earlier survey. Conclusions are also limited because of restriction to one urban area. The results support the need for further systematic investigation of this topic in order to define the risks of food safety related to food access, store size, and other factors. The results suggest that monitoring programs for food borne microbes, conducted by USDA and FDA among others, should include sampling from stores in a range of neighborhoods and that analyses of pathogen prevalence should include information on store type and neighborhood.
References


