Prevalence of Campylobacter in a turkey production facility

Aaron Shawn Kiess
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Prevalence of *Campylobacter* in a Turkey Production Facility

Aaron S. Kiess

Thesis submitted to the
Davis College of Agriculture, Forestry, and Consumer Sciences
At West Virginia University
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Master of Science
in
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ABSTRACT

Prevalence of *Campylobacter* in a Turkey Production Facility

Aaron Shawn Kiess

Frequency of detection was monitored in three flocks of turkeys from May, 2000 to March 2001. The effect of time was considered for hens in flocks 1 and 2, and the effect of time, gender, and litter (fresh or used) was determined for flock 3. Poult, poult-box liners, waterers, and fecal droppings were monitored throughout production for the presence of *Campylobacter* using Campy-Cefex agar incubated at 42 °C under microaerophilic conditions (85% N₂, 10% CO₂, and 5% O₂). Peak colonization occurred near 3 weeks of production. Frequency of *Campylobacter* isolation from bird sources paralleled isolation from waterers. Frequency of detection from birds placed originally on used litter was significantly lower than detection from birds placed originally on fresh litter (2 v. 58%). Gender did not affect rate of detection. Controls to minimize peak colonization at 3 weeks and appropriate litter management are opportunities to reduce the level of this organism in turkeys.
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Introduction:

*Campylobacter*, a pathogenic organism, is a food borne health concern. In the past, *Campylobacter*, formerly known, as *Vibrio* was only acknowledged as an organism responsible for causing stillborn births in cattle and sheep. This limited understanding is due to the fact that the organism is very fragile and requires specific environmental conditions to exist. It was not until the 1970’s that scientists began understanding *Campylobacter’s* contribution to food borne illness. *Campylobacter* is now a leading cause of food borne illness in the United States (U.S. Food and Drug Administration, 1999). Each year it is estimated that over 2 million individuals are infected by this pathogen, resulting in approximately 500 deaths (Center for Disease Control and Prevention, 2000). This rate of infection has made *Campylobacter* even more prevalent than some more recognized pathogens like *Salmonella*, *Shigella* and *E.coli* O157:H7 (U.S. Food and Drug Administration, 1999). *Campylobacter* infects the gastrointestinal tracts of household pets (cats and dogs), domestic livestock (cattle, sheep, swine and poultry), as well as humans. Exposure to less than 500 organisms has the potential to cause the disease, campylobacteriosis. Infection occurs within 2 to 10 days after exposure to the organism. Symptoms include fever, headaches, muscle pain, nausea and bloody diarrhea (Food Safety and Inspection Service, 1999). The severity of these infections, in most cases, is not serious, and symptoms are only experienced for a week. In a few incidences, the infection can spread to other parts of the body like the vascular or nervous system. Campylobacteriosis can also cause arthritis and Guillan-Barre syndrome (GBS), a disease that affects the nervous system causing paralysis (Patterson, M.F., 1995)
Campylobacter is transmitted to humans by improper processing, handling and consumption of raw or undercooked food products. The cross contamination of food items like fruits and vegetables used in salads with raw poultry is thought to be a major source of infection. By thoroughly cooking and properly handling the product, infection by this organism can be reduced or stopped (Center for Disease Control and Prevention, 2000). The objectives of this study were to 1). evaluate the frequency of Campylobacter in turkeys and 2). to determine if vertical transmission of the bacteria from the parent to the egg may be a source for contamination of the poult. Since most research in this area of food microbiology has been performed on broilers more information is needed for turkey producers.
LITERATURE REVIEW

Characteristics of the Organism

Organisms in the genus *Campylobacter* are defined as slender, vibroid, gram-negative cells, 0.2-0.5 µm wide and 0.5-5 µm long (Hensyl, 1994). *Campylobacter* can also take a rod or spiral shape, where they can measure up to 8 µm. It is non-spore forming and may take on a coccoid or spherical form in older cultures. When two or more organisms come together, they may appear as S-shaped or gull winged. It is motile and moves in a characteristic corkscrew motion. The motion is possibly due to a single polar flagellum that is attached at either one or both ends of the cell. The flagella can be up to three times as long as the main body of the organism. *Campylobacter* is microaerophilic requiring less than 5% O₂ to live. In a few strains of *Campylobacter*, up to 20% O₂ can be tolerated. It is a chemoorganotroph that relies on a respiratory type of metabolism in which amino acids and tricarbocyclic acid cycle intermediates are needed for energy. Carbohydrates are neither fermented nor oxidized by this organism. Some species of *Campylobacter* can grow under anaerobic conditions. For this to happen, fumarate, formate and fumarate, or H₂ and fumarate must be present in the growth media or environment. Serum or blood is not a requirement for growth or when testing for *Campylobacter* conformation. They are able to reduce nitrates, gelatin is not hydrolyzed and methyl red and Voges-Proskauer tests are negative. No lipase activity can be seen. Oxidase is positive and urease is negative, except for a few strains of *C. lari*. Some species are pathogenic to humans and animals. They are found in the intestinal tracts, reproductive organs and oral cavities of humans and animals (Hensyl, 1994).
Turkey Production System

History

Over 8000 years ago the first species of fowl (red jungle fowl) was captured and domesticated. The selection process was based on fighting behavior. This selection process was motivated by entertainment value. It was then discovered by Romans, that the fowl offered other characteristics, that were more beneficial to people. The fowl, once domesticated, could be used as a source of meat and eggs that added to the nutrition of the people (Appleby, et al., 1992). The Aztec Indians of Mexico first achieved domestication of the wild turkey. Although the wild turkey was also native to North America, the Native Indians only hunted turkey, and this provided food for consumption and feathers, for clothes and headdresses. The Spanish also domesticated the wild turkey, it was then sent back to their homeland of Spain. In Spain domestication was continued, and turkey was eventually accepted by most of the European countries. By 1573 many other countries like France, Italy and England also adopted the turkey. The first settlers reintroduced the domesticated turkey of the European countries to the Americas. From that time, turkey production has grown into an industry (Moreng, et al., 1985).

The turkey production system is a highly regulated operation. When raising turkeys, many steps are considered so that production will be successful. To achieve the level of production demanded by the industry, special attention is paid to breeding, hatching, housing, brooding and rearing, nutrition, disease and flock management.

Breeding

In today’s turkey production systems, the most prominent breeds used are the large white and large bronze turkeys (Appleby, et al., 1992). To establish a healthy and productive line of turkeys, genetics is the starting point. Economically important traits are governed by a combination of genes. The industry has become indebted to the
Due to this field of genetics, meat birds have been selected for growth rate, meat yield, ratio of white to dark muscle and rapid feathering. When a turkey is selected to carry one or two of these genes, it becomes a part of the grandparent flock. Then by crossing the grandparents, a parent is made. This turkey then carries genes to produce the meat bird. Scientists have looked for genes that will enhance feed efficiency. Producers want a turkey that has high output to low food intake. This is desirable, since the highest cost to the production system is feed. Further research in the field of genetic engineering, will improve efficiency of production and turkey welfare (Moreng, et al., 1985).

**Hatching**

Incubating and hatching turkey eggs are complicated processes, with critical limits that must be monitored thoroughly for maximum hatch to occur. Incubation can occur in two ways. The single-stage method of incubation allows for all the eggs, which are all at the same stage of development, to be set in the incubator at one time. The single stage incubator allows for fine-tuning, of the temperature, humidity and ventilation. Another reason for using this method of incubation is that the hatching cabinet can be thoroughly cleaned after the hatch is finished. On the other hand, multistage incubation allows eggs to be set in the incubator at different times, creating an incubator with eggs at different stages of development resulting in a continuous hatch. In most cases when using this method of incubation, eggs are set on a weekly basis, where one quarter of the incubator is set at a time. Turkey eggs usually incubate for 25 days and then the eggs are set into a hatching box for an additional 3 days so the poults can pip out. During this period of time, much care must be taken to ensure that the correct temperature, humidity and ventilation are maintained. The temperature at which turkey eggs should be incubated is
around 99.5°F. A deviation of just 2 to 3 degrees can be problematic. When temperature rises above the optimum temperature the poult may be forced into using its yolk early, therefore dehydration or emaciated poults may hatch. If the temperature is too low, poults may become lazy, fat and piped or unhatched embryos may result. To prevent these problems from occurring, temperature should be monitored on a consistent basis. If adjustments are needed, they should be done in increments of 0.2°F. Another area that must be constantly monitored is humidity. If proper humidity is not maintained inside the entire incubator, the hatching embryos can experience a number of defects. When the humidity is too high, the eggs may not lose enough water. This results in poor air cell formation. If the humidity is too low, eggs may lose too much water. If this happens, the embryos may become dehydrated causing poor hatchability. For these problems not to occur, a relative humidity of 83% is recommended. Proper ventilation is also very important. As the embryos grow within the egg, the need for oxygen increases. When oxygen requirements increase, carbon dioxide levels will also increase. This is why ventilation is so important. With ventilation, the right amount of O₂ can be brought in and CO₂ removed. There are two systems that can be used to ventilate incubators. The first method is regulated with computer controls linked to the heating and cooling system. The second method is manual. For either method, the need to bring in O₂ and exhaust CO₂ is the main function. Attention to this critical step in incubation must be taken. With too much ventilation, a decrease in temperature and humidity may occur, whereas without enough ventilation, CO₂ can build up resulting in a poor hatch. Following hatching, vaccination, debeaking and declawing may be carried out.
**Housing**

Once poults have hatched, the next need in the production system is housing. Housing today is not as simple as it was a hundred years ago. The design of today’s turkey house is based on production objective and location of the house and it is focused on maximizing growth (Moreng, et al., 1985). In the past minimal housing was needed. This was due to the fact that turkeys were mostly free ranged, meaning they were allowed out of the building into an enclosed area. The house in this system provided protection to the birds from predators at night (Appleby, et al., 1992). Today, turkeys are raised in environmentally controlled buildings. Birds are raised from poults until they are finished. The house provides protection, efficient space, proper lighting, ventilation and heat. Turkeys, used on their genetic predisposition to growth, are raised for specific periods of time in the house. Female turkeys are usually finished by 17 weeks of age and male turkeys are usually finished by 24 weeks (Austic, et al., 1990). All the items mentioned above are all regulated by good management practices. If management is not considered when the house is being designed, it will not be effective. Engineers, when designing a building take great care in deciding how the building should be ventilated, heated, insulated, sanitized, lighted, and plumbed (Austic, et al., 1990). Laborers comfort is also considered. If laborers are not comfortable, birds will suffer. Laborers want to get in and out of the house as quickly as possible, which may result in certain indicators of birds well being overlooked (Appleby, et al., 1992). All of these factors depend on location, climate and space needed by the birds, ensuring the building will produce a profitable end product.
**Brooding and Rearing**

Brooding poults is similar to a nursery period; when poults arrive at the house, strict care must be taken to maintain their health. The first step taken is to ensure that the house has been properly cleaned and sanitized. Once the house has been inspected, the producer will then heat the house. This occurs approximately 12h before poults arrive (Moreng, et al., 1985). Turkey poults are raised on the floor of the house; fresh shavings are laid down for the comfort of the poults. Brooding rings are set up to keep the poults in and around the brooder so they will not get too cold. The brooder is a heating device that maintains temperature around 99.5°F. Waters are adjusted to the correct height or depth to allow for easy access (Austic, et al., 1990). When poults arrive at the house, behavior is observed. Observations will indicate whether poults are getting too hot or cold. The producer will note if the birds are consuming water, turkeys may be hard to start on feed and water. If this occurs, it is not uncommon for the laborers to dip the poults head in the water to familiarize them with it. Feed is not put into the brooding ring for about 4h after poults arrive. This practice allows birds to consume some water first in case they are dehydrated from the trip to the house (Moreng, et al., 1985). During brooding, laborers and producers carefully monitor heat, airflow, moisture and space allowed for each poult. Once poults have been monitored for a week within the brooding ring and all poults are eating and drinking, the ring can be taken down and the brooder turned off and raised. More room will be needed for growing poults and this need should be taken care of by building design (Appleby, et al., 1992). Feeders and waters should be placed throughout the house to allow for access. As poults grow, all behavior and areas
previously mentioned should be continuously monitored. If not watched, bird performance will decrease and so will profit.

**Nutrition**

The leading cost to turkey production is feed, thus nutrition is a primary concern to the industry. Energy requirement for turkeys cannot be designed with amino acids, vitamins and minerals alone (Austic, et al., 1990). The major goal of the nutritionist is to design a feeding program that includes all essential ingredients and maximizes muscle growth as economically as possible (Moreng, et al., 1985). Nutritionists have designed feeding programs that maximize growth efficiency of turkeys. As turkeys age, metabolic energy requirements increase. For young poults, requirement for metabolic energy is around 1250 to 1350 kcal; whereas older turkeys require 1400 to 1500 kcal of metabolic energy (Austic, et al., 1990). Concurrently, protein requirements decrease with age. Most diets use carbohydrate sources to substitute for protein such as corn or soy. Essential vitamins and minerals increase or decrease with the age of the turkey (BUTA). A well-developed management plan will recognize the turkey requirement and, if a ration is needed for a longer or shorter period of time, adjustments can be made (Appleby, et al., 1992).

**Disease**

Diseases are another major concern to the turkey industry. If a disease does become a problem, it could affect the entire flock possibly causing death to all birds. For this reason, all attempts to ensure that diseases do not occur are taken. A disease may be defined as any deviation from the normal state of well being. This deviation could appear as a slight ailment in a single bird or one that endangers the life of the entire flock (Moreng, et al., 1985). Disease agents can be specific pathogens, nutritional deficiency,
toxic agents or even environmental changes. Diseases that are most common and managed by sanitation practices and vaccination are viruses, bacteria, fungi, protozoan, poisons, and internal and external parasites (Austic, et al., 1990). Methods of disease prevention include vaccination, sanitation, farm security, and the practice of all-in/all-out management. When proper hygiene is practiced and observation of flock health is recorded accurately, disease outbreaks are minimized (Appleby, et al., 1992).

**Flock Management**

Management is essential to every area of the system, and if not properly practiced by all personnel, it could cause failure of the entire operation. Moreng (1985) stated “The design of a management program for a poultry flock should be based upon the efficiency of operation necessary to meet the production objectives of the unit. Efficient use of space, equipment, time, and employee’s knowledge should combine with many other factors which must be applied toward the production goals of a specific program”. Breeding, hatching, housing, rearing and marketing should be considered in development of a management program. When poult comfort, employee welfare, breeding, hatching, disease prevention, and rearing are managed properly, benefits will be realized. If problems are not solved promptly and production is halted, management has failed. The goal of management is efficient and profitable production.

**Prevalence of *Campylobacter* in turkeys and their products**

Over the years, interest in *Campylobacter* has increased because of increasing links between it and gastroenteritis (Quinones-Ramirez et al., 2000). Poultry, sheep, pigs and cattle, at various levels, carry it. In poultry, a major reservoir for *Campylobacter*, studies have indicated levels of 8.1 to 100% (Quinones-Ramirez et al. 2000). Doyle (1984) associated *Campylobacter* with all parts of the poultry system from live birds to retail
products. In the live bird, many studies had been carried out to determine if they were a source of contamination. Initially, experiments were designed to see if poultry actually harbored the organism (*Campylobacter*). Smith and Muldoon (1974) were a few of the first to document the presence of *Campylobacter jejuni* associated with poultry carcasses. Acuff and coworkers (1982) looked for *Campylobacter* in turkey eggs, poultts and brooding houses. They reported that poultts, 15 to 19 days of age, had a frequency of contamination of 16 to 76% respectively. They also sampled litter, grit, feed and water from within the house. Water and litter were positive for *Campylobacter*, but grit and feed were not. When fertile eggs and newly hatched poultts were sampled, neither harbored *Campylobacter*. Doyle (1984) investigated the excretion patterns of *C. jejuni* in individually caged laying hens. He found that 8.1% of the laying hens were chronic carries of the bacteria. He also found that an increase of 25% in the shedding of *Campylobacter* occurred during the months of October and late April through early May. Doyle (1984) concluded that *Campylobacter* could not penetrate the egg, but when eggs had fecal material on them, it was possible to isolate *Campylobacter*. Poultry provide an optimum environment for *Campylobacter*, but the bacterium does not cause disease in the host.

Stern and coworkers (1988) investigated this commensal relationship between *Campylobacter* and poultry. They evaluated colonization dosage, age of the host and strain variability relative to cecal colonization influence of competitive exclusion (CE). All chicks challenged with $10^5$ cfu/chick were consistently colonized. When six different strains of *C. jejuni* were used as a challenge, four showed permanent attachment properties. One other strain did show consistent colonization, but it was by oral-fecal
passage among the flock. When birds were given competitive exclusion cultures from mature adult microflora, no difference was seen in the colonization rate of *Campylobacter* in the birds. For birds treated with CE cultures, 81 of 84 were colonized and 45 of 46 of the control birds were also colonized.

Kwiateck and coworkers (1990) looked at the presence of *Campylobacter* on poultry and slaughter animals in Poland. They sampled chickens, ducks, geese, turkeys, pigs and cattle. Out of 839 samples, 80.3% of chicken, 48% of duck, 38% of geese, 3% of turkey, 2.9% of porcine and 0.9% of bovine samples taken were positive for *Campylobacter*. Jones (1991) evaluated the rate of *C. jejuni* contamination at various points in broiler production and processing systems. He indicated that feed mills, hatcheries, insects, and mice were all negative for *C. jejuni*. Insects were externally cleaned with ethanol, so the negative counts were all based on internal colonization of insects, nonetheless external transmission may be possible. In this experiment the actual source for contamination could not be determined. Broilers entering the slaughter facility had cloacal swabs taken; of those samples, 20% were positive for *C. jejuni*. It was also found that 52% of carcasses, post chilling were positive. Whole broiler carcasses, once processed, were also contaminated with *C. jejuni* at a frequency of 31.6%. Grados and coworkers (1988) studied free-range chickens. *C. jejuni* was present in birds that had access to a backyard lot. Adekeye and coworkers (1989) observed that *Campylobacter* acts as if it was part of the normal flora. They found that in intensive management and free-range management systems *Campylobacter* colonization was almost identical. Pearson and coworkers (1993) investigated an outbreak of *C. jejuni* infections in Bournemouth, United Kingdom. The outbreak was associated with a catering college that was supplied with chicken from
a single wholesaler. Pearson and colleagues traced the \textit{C. jejuni} back to a farm that had a contaminated water supply.

Stern and coworkers (1994) examined the possibility that when chicks are challenged with viable but non-culturable (VBNC) suspensions of \textit{Campylobacter}, colonization may occur. They indicated that some chicks challenged by oral gavages of VBNC suspensions became colonized. Confirmation was done by two laboratories that used heat-stable and heat-labile serotyping schemes. Stern and coworkers (1995) studied the role of transport and holding on \textit{Campylobacter} colonization rates. They found that 9 of 10 farms sampled had broilers contaminated with \textit{Campylobacter}. When birds were examined before and after transport and holding, a higher level of contamination occurred after transport and holding. To reconfirm this finding, five additional farms, and 200, six-week old chickens were studied. Before transport, 12% of birds harbored an average of 2.71 log$_{10}$ cfu/carcasses. After transport, this level of contamination increased to 5.15 log$_{10}$ cfu/carcass in 56% of the birds sampled. On two Dutch broiler farms, Jacobs-Reitsma and coworkers (1994) found that 100% of the samples collected were positive for \textit{Campylobacter}. Neill and coworkers (1984) found that 10 of 12 flocks sampled did not produce \textit{Campylobacter} positive samples when the chicks were at 2-weeks of age. As the birds aged, the flock became colonized, but the source of colonization could not be found. Aho and Hirn (1988) reported that \textit{C. jejuni} colonization was very low, around 1.7%, at 4-5 weeks of age. At slaughter (6-7 weeks), the frequency of \textit{C. jejuni} colonization had increased significantly to around 24%. With levels as high or higher than seen by Aho and Hirn (1988) the possibility for contamination during processing is high. Grant and coworkers (1980) examined broiler
chickens to see if they presented a potential source of infection with *Campylobacter*. In 46 broiler chickens, 38 (83%) harbored *Campylobacter* at an average level of $4.4 \times 10^6$ cfu/g of feces. The organism survived in feces for 96 h at 4°C. This study showed that the potential for contamination in the processing plant is high.

*Campylobacter* in processing plants has been studied extensively. Leuchtefeld and Wang (1981) studied the patterns of *Campylobacter* contamination in a turkey processing plant. In this study, 600 cecal and fecal droppings were collected over a 1-year period, and many samples were positive for *Campylobacter*. Thirty-three, fresh-dressed carcasses were examined before chilling in chlorinated ice water. From those samples, 94% were positive, and after overnight soaking, 34% of 83 carcasses were still contaminated. These investigators sampled gutters, chutes, conveyor belts and water treatment lagoons, and positive samples were collected from all sources while the plant was in operation. Four days after shutting the plant down during the winter all samples taken were negative.

Svedhem and coworkers (1981) looked at the occurrence of *C. jejuni* in fresh foods as well as the survival of the bacteria under different conditions. They found that *C. jejuni* survived on food items at 4°C for a week and at -20°C for up to 3 months. They indicated that by cooking food for 15 min at 60°C was enough to destroy *C. jejuni*. Park and coworkers (1981) examined the incidence of *C. jejuni* in fresh, eviscerated whole chickens. *C. jejuni* was recovered at a frequency rate of 62 and 42% from Ontario and Ohio samples respectively. In Sydney, Australia, Shanker and coworkers (1982) found *C. jejuni* from 18 of 40 processed carcasses and 134 of 327 cloacal swabs. Eighty two
percent of chicken and 98% of human isolates from the same area were identical biotypes.

Kramer and coworkers (2000) applied an epidemiological typing strategy to analysis of fresh meat and poultry fecal samples, and to human isolates from campylobacteriosis cases. All samples and isolates were from the same geographical area and they were collected within the same time frame. *Campylobacter spp.* were isolated from 73.2% of 489 samples. Frequency of isolation was highest for chickens at a rate of 83.3%, followed by lamb at 72.9%, pig at 71.7% and beef livers at 54.2%. For human cases, 89.3% were *C. jejuni* and 10.7% *C. coli*. Only 30% of the positive isolates collected from each species were multiple strains, reinforcing the fact that more than one isolate should be selected from each sample.

Barot and coworkers (1983) looked for the location of *C. jejuni* on chicken livers. They found that of 117 livers sampled from New York retail outlets, 56 were positive for *C. jejuni*. Thirty-six of the livers showed surface contamination, and two had *C. jejuni* contamination in the tissue. This contamination was most likely due to unhygienic handling of the offal. Oosterom (1983) evaluated the occurrence of *C. jejuni* contamination during poultry processing in the Netherlands. Birds, equipment, workers hands and air were determined to be sources for *Campylobacter* contamination. Intestinal contamination in the birds was found as high as 7 log$_{10}$ cfu/gram. Scalding reduced *C. jejuni* levels, but levels increased during defeathering and evisceration. Carcasses and livers were contaminated at a frequency of 50 to 75%, respectively.

Wempe and coworkers (1983) studied the prevalence of *C. jejuni* at different stages of slaughter in two California chicken processing plants. They found that *C. jejuni* was
isolated from 68% of ready-to-eat products. Sixty to 100% of cecal and fecal samples had levels of \textit{C. jejuni} as high as \(10^6\) cfu/g when entering the slaughterhouse. This study showed that fecal and cecal contamination of carcasses is possible during the slaughtering process. Harris and coworkers (1986) investigated prevalence of \textit{C. jejuni/coli} (\textit{C. j/c}) in fresh meats available to the consumers in King County, Washington. A total of 297 samples were collected from a poultry processing plant. From those samples, 56.6% were positive for \textit{C. jejuni/coli}. When 862 retail chickens were sampled, 23.1% were positive and 17.2% of 29 retail, game hens were also positive. In turkey, pork and beef sampled, \textit{C. jejuni} and \textit{C. coli} were not frequently isolated.

Berrang and coworkers (2000) examined levels of \textit{Campylobacter} associated with broiler chickens entering the processing plant. \textit{Campylobacter} populations (log\textsubscript{10} CFU/g) were 5.4 on feathers, 3.8 on skin, 4.7 in the crop, 7.3 in the ceca, and 7.2 in the colon. Juven and Rogol (1986), looked at the incidence of \textit{Campylobacter} in chickens at the pre-salt immersion stage in a kosher processing plant in Israel. For carcasses sampled before immersion, 85% were contaminated with \textit{Campylobacter}. After immersion, the frequency of detection (85%) did not change. These data suggest that salt-water is not an effective way to remove \textit{Campylobacter} from carcasses. In 1983, Kinde and coworkers estimated the prevalence of \textit{C. jejuni} in chicken wings sold at supermarkets in California. \textit{C. jejuni} was found in 82.9% of 94 wing packages. However, after a few days at the supermarket, only 15.5% of 45 packages were positive. Another study in 1994 by Flynn and coworkers looked at the prevalence of \textit{Campylobacter spp.} in retail chicken wings in Northern Ireland. They sampled 153 chicken wings that were purchased from retail outlets over a 10-wk period. For these samples, 64.7% were contaminated with
*Campylobacter spp.* Confirmation was performed with the API-Campy Identification system for *Campylobacter*.

Acuff and coworkers (1986) evaluated the effects of roasting, braising, stewing, and microwaving on contamination of turkey thighs. Effects of various utensil and hand-washing procedures on the survival of *C. jejuni* were also evaluated. Roasting, braising and stewing were all effective in decontaminating turkey thighs of *C. jejuni*. Microwaving was most successful when a thermometer was used to evaluate internal temperature. Manually washing utensils with detergent and water was sufficient to remove *Campylobacter jejuni*, except on wooden cutting boards where dishwashers were needed for proper sterilization. Quinones-Ramirez and coworkers (2000) examined the contamination of *C. jejuni* on poultry that was available to consumers on street stands, the most common for eating in Mexico City. Out of 100 samples from 3 locations, 600 isolates were grown. Of the 600 isolates, 121 were positive. Fifty-one samples were *C. jejuni*, 21 were *C. coli* and 49 were other species. Twenty-seven positive isolates were taken from one stand. Previously cooked poultry had been placed under raw chicken during roasting, allowing cross-contamination to occur.

In 1983, Hopkins and Scott investigated an outbreak of *Campylobacter* in Colorado. This outbreak was suspected to be due to the mishandling of raw chicken. In ten cases, interviews were conducted on infected persons as well as the family members who were not infected. Nine of 10 infected individuals had handled raw chicken before the onset of symptoms. They concluded that handling raw chicken was more of a risk than the consumption of cooked chicken. In another study, Lammereding and coworkers (1988) developed a national monitoring program in Canada that provided information on the
status of thermophilic *Campylobacter* in food animals at the slaughterhouse level.

Thermophilic *Campylobacter* was isolated from 16.9% of pork, 22.6% of beef, and 43.1% of veal samples. *Campylobacter* was isolated from 73.7 and 38.2% of turkey and chicken samples, respectively.

**Preharvest (Production) factors affecting the frequency of *Campylobacter***

Poultry is a major reservoir of *Campylobacter* and the high frequency of *Campylobacter* occurrence in poultry has also been implicated as a predisposing factor for gastroenteritis in humans. Due to its prevalence, elimination of the pathogen has been the focus of recent research. To eliminate *Campylobacter* from poultry, a logical step is to identify factors that influence the frequency of *Campylobacter* in the live bird. Doyle and Roman (1981) looked at the effect that temperature and pH would have on *Campylobacter* growth. They indicated that an optimum temperature for *Campylobacter* was in the range of 42 to 45°C and that *Campylobacter* could grow in a pH range of 5.5 to 8.0. Jones and coworkers (1993) studied the effect of temperature, microaerophilic conditions and air on *Campylobacter* growth. Optimum growth occurred around 42°C, but growth was seen at 37°C and as low as 4°C. After 2-3 days in air, sub cultures could change to aerobic metabolism and grow without microaerophilic conditions. Changes in morphology and outer-membrane proteins were seen, but serotyping reactions were not changed when identifying the organism.

Season affects carriage rate of *Campylobacter* in poultry. Jacobs-Reitsma and colleagues (1994) investigated risk factors for *Campylobacter* colonization in Dutch broiler flocks. Of 187 broiler flocks 82% (153) were contaminated with *Campylobacter*. Colonization rate varied with season; June through September had the highest rates of
colonization and March had the lowest colonization rate. In this seasonal study, slaughterhouse and husbandry practices correlated to *Campylobacter* contamination, but broiler line and age and geographical location did not affect colonization rate. Stern and coworkers (1995) examined seasonal influence on colonization of broilers with *Campylobacter*. The lowest levels of *Campylobacter* were detected in the spring. In the summer and fall, levels were at their highest level. Willis and Murray (1997) determined that the highest number of *Campylobacter* on carcasses was from May to October. The lowest levels were detected in December and January. These findings support the fact that seasonality influences the detectability of *Campylobacter* in market broilers.

Other risk factors are associated with the frequency of *Campylobacter* colonization of poultry. Humphrey and coworkers (1993) conducted a longitudinal study that lasted approximately 12 months and examined the *Campylobacter* status of broiler flocks. Ceca of up to 100 birds per flock were examined at slaughter. This sampling rate allowed for a study for a variety of environmental and production factors. They determined that bird colonization were not associated with water or the floor structures used in the house; also, no seasonal variation was seen in carriage rate. They indicated that dipping boots prior to entering a house helped to reduce or even eliminate *C. jejuni* in samples collected from the house.

Smith and Fratamico (1995) reviewed factors that were involved in the emergence/recognition and persistence of several bacterial, parasitic, viral and viral-like agents that are associated with food-borne outbreaks. Relative to *Campylobacter*, they stated that increased emergence or recognition is related to 1) increased awareness of *Campylobacter* as a food-borne pathogen and 2) the development of better selective
media for stool and food samples. Relative to factors contributing to the persistence of the pathogen, they implicated raw milk, untreated water, poultry consumption, contact with farm animals and pets, contamination of food by food handlers, and concentration of chickens, pigs and cattle in production systems.

Jacobs-Reitsma and coworkers (1995) conducted a longitudinal study of *Campylobacter* in broiler flocks and associated environmental sources at two Dutch poultry farms. They determined that the hatchery, water, feed, and fresh litter were not sources of contamination. Darkling beetles carried the same serotype of *Campylobacter* as the broilers thus acting as a vector for horizontal transmission. In 1996, Cawthraw and coworkers examined the mechanism by which *Campylobacter* spread through large broiler flocks. They indicated that a dose of 40 cfu/gram was large enough to maximize colonization in poultry. Van de Giessen and colleagues (1996) identified risk factors as well as risk-reducing measures for *Campylobacter* infection in Dutch broiler flocks. Risk factors were boots, hand washing, foot baths, improper cleaning with detergents and failure to clean the yard between flocks. Animals and ground water used for cleaning were sources of flock infections.

Transportation contributes to *Campylobacter* contamination of poultry. Stern and colleagues (1995) investigated the role of transport from the farm to the processing facility in *Campylobacter* colonization of chickens and contamination of carcasses. Stern indicated that the level of *Campylobacter* before transport was around 5.44 log<sub>10</sub> cfu/g of cecal matter. After transport and a holding for 16 to 18h, counts increased to 6.15 log<sub>10</sub> cfu/g of cecal matter. Achen and coworkers (1998) studied time of onset and duration of *C. jejuni* shedding and observed that peak excretion of *Campylobacter* occurred 13-19
days post inoculation. By market age, 37.5% of the birds were shedding *C. jejuni* and 12.5% were chronic carriers. Lindblom and coworkers (1986) examined the natural colonization of chickens by *C. jejuni* during commercial breeding and rearing. No *C. jejuni* was found in newly hatched chicken feces. By 5-9 weeks of age, many samples became positive for *C. jejuni*. Once a bird became colonized, it spread rapidly, but the for the rate of transmission was not known. Feed and water were negative, and environmentally protected birds were colonized over time. These authors suggested that transmission was due to flies and other insects within the house.

The pathway whereby poultry become colonized is still an enigma. Many studies have looked at vertical or horizontal transmission as a pathway by which colonization occurs. Shanker and colleagues (1986) evaluated the role of vertical transmission in *C. jejuni* colonization of a broiler flock. They stated that, in 6 breeder flocks supplying eggs, 74% of the breeders were positive for *C. jejuni*. Out of 187 eggs sampled, only 2 were identified as *Campylobacter* carriers. When eggs were incubated and/or challenged with *Campylobacter* isolates, only two were positive. Thus, vertical transmission was not considered a possible route of infection in broilers. Jacobs-Reitsma (1995) investigated the role of breeder flocks in the epidemiology of *Campylobacter* in poultry production. *Campylobacter* was isolated from 67% of the flocks. *Campylobacter* colonization of breeder flocks suggested a potential role for vertical transmission, but serotype data did not support this link.

Pearson and coworkers (1996) found that out of 12,233 samples collected, 27% were positive for *Campylobacter*. When 251 broiler houses were sampled, 35.5% were *Campylobacter* positive. Of those positive houses, only 9.2% had consecutive samples in
which *Campylobacter* could be isolated. In sampling two hatcheries, this group found that the first hatchery had 17.6% *Campylobacter* positive samples. In the second hatchery, 42.9% of the samples were positive. These data support the idea that vertical transmission is a way whereby *Campylobacter* colonizes chicks.

In 1992, van de Giessen and colleagues used the Penner serotyping and DNA-typing systems to assess roles of vertical and horizontal transmission in *C. jejuni* infection of poultry flocks. Samples were collected from two broiler houses and the first house had strains of *C. jejuni* that could not be isolated from subsequent flocks. The second house had identical strains of *C. jejuni* isolated from subsequent flocks. In this study, horizontal transmission was suspected to be the major contributor to colonization of poultry flocks. Shanker and colleagues (1990) provided information on horizontal transmission. *C. jejuni* was investigated in *Campylobacter*-free broiler chickens. When chicks were orally challenged with the organism, 64% were positive for *Campylobacter* within 3 days and 89% were positive by day 7. When chicks were exposed to contaminated water or seeder chicks, colonization occurred within 2-7 days. When all chicks were removed from the house and environmental samples collected, the house was negative by day 3. Colonization of 1-day old chicks was not affected when adult cecal microbiota was introduced.

Due to the many opportunities for *Campylobacter* to colonize poultry, scientists are looking for ways to reduce the frequency of positive birds entering the processing plant. Stern and coworkers (1994) looked at alternative sources of flora antagonistic to *C. jejuni*. Mucosal competitive exclusion flora (MCE) was given to chicks. Forty-eight hours later, chicks were challenged with *C. jejuni* to examine colonization characteristics.
Exposure to MCE reduced the level of *Campylobacter* that colonized the chick, but after the MCE was stopped, its effectiveness was reduced. In 1997, Morishita and colleagues evaluated whether the probiotic effect of *Lactobacillus acidophilus* and *Streptococcus facium* would reduce colonization and frequency of fecal shedding of *C. jejuni* in broilers. Birds given probiotics for 1 to 3 days had a 70% reduction in the frequency of *C. jejuni* shedding and a 27% reduction in jejunal colonization at slaughter compared to control groups. Line and coworkers (1997) evaluated the ability of *Saccharomyces boulardii* to reduce populations of *Salmonella* and *Campylobacter* in broiler chickens subjected to feed withdrawal and transport stress. Yeast did not reduce the frequency of *Campylobacter* isolation from the ceca; however, *Campylobacter* populations in the ceca were significantly reduced when the culture was given to a chick that was contaminated with both *Campylobacter* and *Salmonella*. Line and colleagues (1998) also studied the potential for using a viable, dried preparation of *S. boulardii* in the feed as a defined culture for controlling colonization of broiler chicks with *Salmonella typhimurium* and *C. jejuni*. *Salmonella* colonization was significantly reduced due to the yeast treatment. *Campylobacter* colonization on the other hand was not affected by yeast treatment.

Bailey and coworkers (1993) evaluated technology that would prevent communal intestinal colonization of chickens by human bacterial enteropathogens. Bailey concluded by indicating that the only method to reduce or eliminate *C. jejuni* at the processing plant is to achieve a bird that is *C. jejuni* free.

**Isolation and Detection of *Campylobacter***

*Campylobacter* was not recognized as a food-borne pathogen until the 1970’s. This
was due to its fragile state outside its preferred environment. Over the years much research has been conducted on media that can support growth of *Campylobacter spp.* Many selective media use antibiotics; enrichment media to recover injured cells have been developed. With the advent of DNA-based technology, new products are being developed that can identify *Campylobacter spp.* using specific deoxyribonucleic acid (DNA) segments through techniques like polymerase chain reactions (PCR) and pulsed field gel electrophoresis (PFGE). Patton and colleagues (1981) compared (Skirrow’s, Butzler’s and a modified Butzler’s, containing a higher concentration of colistin) for there efficiency in the primary isolation of *C. fetus* subsp. *jejuni*. Skirrow’s and Butzler’s and modified Butzler’s media were comparable in their isolation of *C. fetus* subsp. *jejuni*. Modified Butzler’s gave the highest level of isolation, and when combined with Skirrow media, 98% of the isolates obtained could be detected. Butzler and coworkers (1983) compared new selective medium that consisted of Butzler’s medium with the addition of cefoperazone, rifampicin, colistin and amphotericin. They found that out of 3,404 stool samples, *C. jejuni* was isolated with the same frequency (7-9%) for both media. However, competing fecal flora were strongly suppressed by the new media. Bolton and colleagues (1983) also compared Skirrow’s, Butzler’s, Campy-BAP and Preston media for *Campylobacter spp.* isolation from human, animal and environmental specimens. They indicated that the Butzler medium gave the lowest isolation rate; whereas Preston medium, the most selective medium, gave the highest isolation rate. Again in 1984, Bolton and coworkers compared charcoal, cefazolin, sodium deoxycholate (CCD), a blood-free selective agar, to Preston medium for isolation of *C. jejuni* from human feces.
Both media resulted in similar isolation rates. CCD was less selective for Campylobacter than Preston media.

Lai-King and colleagues (1985) compared growth of C. coli and C. jejuni in the presence of antibiotics used in selective growth media. C.coli was more susceptible to antibiotics that C. jejuni; 1 out of 9 antibiotics did not inhibit C. coli. Inhibition of C. coli on media developed specifically for Campylobacter spp. confirms that C. coli may be underestimated when antibiotics are used in the media. In 1988, Lai-King and colleagues evaluated the ability of C. coli to grow on a range of media in use for selective culture of Campylobacter spp. C. coli isolates were inhibited more than C. jejuni on selective media developed by Hutchinson and Bolton. Merino and coworkers (1986) compared seven selective media for isolating C. jejuni; these media were Butzler, Blaser, Skirrow, Preston, Preston Blood free agar, Butzler Virion and modified Preston with amphotericin.

B. All media isolated a similar number of C. jejuni. The Preston, Campylobacter blood free medium with cefoperazone yielded the highest number of C. jejuni isolates; all others allowed abundant growth of other fecal flora. The presence of this fecal flora made detection of suspect colonies difficult and increased the time spent in reading each plate. Gun-Munro (1987) evaluated six selective isolation media for their ability to support the growth of C. jejuni. Gun-Munro also found that Preston medium, charcoal, cefoperazone, sodium deoxycholate agar (m-CCDA), charcoal-based selective medium (CSM) and charcoal, cefazolin, sodium deoxycholate agar (CCDA) produced the highest recovery rate with the greatest suppression of other fecal flora. Yang Chih-Shih and coworkers (2000) evaluated the API-Campy Identification kit (Biomerieux, Marcy-l’Etoile, France) for its applicability and compared the efficacy of three selective media
(charcoal, cefoperazone, sodium deoxycholate agar (m-CCDA), Campy-Cefex agar (CCA) and charcoal-based selective medium (CSM)); they observed no differences. The API-Campy kit efficiently detected 87 *Campylobacter spp.* isolates from chicken samples examined with 100% agreement at the genus level and up to 94% at the species level when compared to conventional methods.

Hodge and Terro (1984) compared isolation of *C. jejuni* from human fecal specimens by direct inoculation on selective Columbia agar and liquid enrichment medium. They found that the liquid enrichment medium produced a 30% higher isolation rate for *C. jejuni*. The overall isolation rate achieved by both methods was 8.2% for 1,249 specimens. Agulla and coworkers (1987) evaluated the growth of *Campylobacter* on alkaline peptone water (APW), Bruce-Zochowsky medium broth (BZ), *Campylobacter* enrichment broth (CEB) and Campy-thio broth (CT). *C. jejuni* was isolated from 43 of 359 specimens with CT, 45 with APW, 46 with BZ and 46 with CEB. No significant differences were found for the number of isolates obtained with and without enrichment procedures. In 1983, Fricker and Girdwood compared enrichment of fecal samples for *Campylobacter* to direct plating on Preston and Skirrow media. Enrichment culture had little effect on the frequency of *Campylobacter* from most patients with acute diarrhea, provided that good selective medium is used and that the delay in culturing specimens is minimal. In 1983, Hutchinson and Bolton examined the role of enrichment culture in isolation of *C. jejuni* from feces. They reported that enrichment culture was only necessary for specimens when the number of organisms is likely to be low. Martin and coworkers (1983) tested a new selective enrichment broth for the isolation of *C. jejuni* from fecal specimens of human, poultry and bovine origin. Compared to direct plating,
the new enrichment increased isolation rate of \textit{C. jejuni} by 46.3\%. In 1984, Francis and colleagues demonstrated the advantage of using enrichment-culture techniques to isolate \textit{C. jejuni} from stools. Data indicated that enrichment broth supplemented with antibiotics markedly increased \textit{Campylobacter} isolation rate. In 1985, Garcia and coworkers examined, by direct plating and enrichment techniques, the prevalence and distribution of \textit{C. jejuni} and \textit{C. coli} at various sites in the digestive tract of cattle. Isolates were found 40.2\% more frequently when enrichment techniques were used. Humphrey (1989) appraised the efficacy of pre-enrichment for isolation of \textit{Campylobacter jejuni} from food and water. With the broth culture, isolation could be increased by pre-enrichment in basal or selective media at 37°C for 4 hours. Jeffrey and colleagues (2000) studied production of an economical, easy to prepare field-suitable enrichment medium for detection of \textit{C. jejuni} in small numbers. The medium was able to detect, with 75\% accuracy, \textit{Campylobacter} at $10^0$ and $10^1$ dilution rates. When challenged by inclusion of \textit{E. coli}, the recovery rate was 50 to 100\% when the medium was inoculated with one to 1 million cfu/ml.

Due to advances in technology, molecular techniques have been developed to identify \textit{Campylobacter}. In 1997, Linton and coworkers evaluated newly designed PCR techniques for detection and identification to the species level and for typing of \textit{Campylobacter} directly from human fecal specimens. Out of 20 clinical samples from which \textit{Campylobacters} had been cultured, \textit{C. jejuni} was in 17, \textit{C. coli} in 2 and co-infection of \textit{C. jejuni} and \textit{C. hypointestinalis} in 1 sample. Results agreed with culture and phenotypic identification to the species level. Lawson and coworkers (1997) developed a rapid and simple PCR assay for detecting and differentiating \textit{C. upsaliensis} and \textit{C.
helveticus in fecal samples. The PCR assay was compared with culture detection by a membrane filter technique and a selective agar containing cefoperazone. The PCR assay and membrane filter technique were similar in isolation rates. The selective agar could detect *Campylobacter* at lower levels than the PCR assay, but the PCR assay only took 8 h for results; the selective agar and membrane filter technique required 48 to 96 h in a microaerophilic environment to culture *Campylobacter*. Thunberg and coworkers (2000) compared detection efficacy of a PCR technique with a standard plating method for detection of *C. jejuni* in a variety of foods. Charcoal and iron used in the enrichment broth interferes with the PCR assay. Once the problem was corrected, there was virtually no difference in detection of *C. jejuni* among enriched samples analyzed by PCR and the selective agar isolation (SAI) method. Using 48 h enriched cultures in combination with PCR analysis could possibly save one day in the time required for presumptive identification of *C. jejuni* in suspected foods. Nielsen and coworkers (2000) compared 6 methods for subtyping *C. jejuni* isolates from animal, human and water outbreaks. All isolates were typeable by each of the 6 methods.

Stern and Robach (1995) evaluated non-destructive sampling methods (i.e., fecal droppings, cecal droppings and cloacal swabs) to monitor the presence of *Campylobacter spp.* in broiler chickens. During an entire growout period, 45% of 964 fecal droppings, 58% of 284 cecal droppings and 41% of 786 cloacal swabs presented positive isolates of *Campylobacter spp.* Sampling of cecal droppings was the most sensitive, non-destructive sampling method. In 1990, Arimi and coworkers investigated the haemolytic activity, on blood agar plates, of some Campylobacters to determine if haemolysis might be a useful aid in strain differentiation. Distinct haemolysis occurred for 92.3% of *C. jejuni* and
21.7% of *C. coli* strains on sheep blood heart infusion agar incubated for 4 d microaerophilically at 42°C. Haemolysis was also detected when horse blood heart infusion agar was used. No other strains of *Campylobacter* tested were haemolytic. The plate haemolysis test may aid differentiation within the thermophilic *Campylobacters*. On and Holmes (1991) studied data on reproducibility using three different basal media for tolerance tests. Reproducibility of the tests with each medium exceeded 89%. The proportion of strains able to grow in a reproducible manner on the basal media varied from 100% for blood agar, to 50% for nutrient agar, to 5% for brucella agar. In 1992, Doyle and Roman provided information on the response of *Campylobacter* and nalidixic acid-resistant, thermophilic *Campylobacter* to sodium chloride at 4, 25 and 42°C. Growth occurred at 42°C in 1.5% NaCl, but not at 2.0%. At the same temperature nalidixic acid-resistant, thermophilic *Campylobacter* could grow in 2.0% NaCl and was tolerant up to 4.5%. At 4°C, all strains were sensitive to 1.0% NaCl and higher, but cell death was slower than what was observed at 25°C at a constant level of sodium chloride. Saha and colleagues (1991) studied the ability to resuscitate freeze-thaw injured *C. jejuni* strains to a fully virulent form. After thaw, direct plating did not demonstrate growth. When 16 freeze-thaw injured *C. jejuni* strains were passed through rat guts for 18 h, seven strains were resuscitated. After consecutive samplings, *Campylobacter* strains regained full virulence capacity. In 1999, Doan and colleagues determined the relative recoveries of important periodontal bacteria in Coy anaerobic chambers, Gas Paks and AnaeroPack culture systems. The Coy anaerobic chamber yielded the highest proportional recoveries of *Campylobacter*, which is considered one of the important periodontal bacteria.
Chapter 1

Determining possible effects of *Campylobacter* on a turkey production facility
Introduction:

Colonization of birds by *Campylobacter* predisposes the carcass to contamination during processing and increases the potential to cause disease in humans. Initially, *Campylobacter* was known as *Vibrio* and thought to only pose risk to cattle and sheep where stillborn deaths were experienced. *Campylobacter*, 0.5\(\mu\)m wide and 5.0\(\mu\)m long, requires a microaerophilic environment (85% N\(_2\), 10% CO\(_2\) and 5% O\(_2\)) (Hensyl, 1994). It relies on a form of metabolism that uses amino acids and tricarbocyclic acid cycle intermediates for energy. These requirements make the intestinal tracts of most mammalian and avian species ideal for *Campylobacter* colonization. As a result, poultry share a commensal relationship with *Campylobacter*. This type of a relationship causes no harm to the host (the host being the bird), but still provides the nutrients needed for *Campylobacter* to survive. The kind of relationship poultry has with *Campylobacter* makes it a major reservoir for this pathogen. Stern and coworkers (1992) stated, “The first report on the presence of *Campylobacter jejuni* associated with chicken carcasses was discovered by Smith and Muldoon”.

Recent studies have reported levels of *Campylobacter* from 8.1 to 100% in poultry (Quinines-Ramirez, et al., 2000).

*Campylobacter* has been isolated at all phases of poultry production, from the live bird throughout the production cycle to the retail products sold in supermarkets (Doyle, 1994). Neill and colleagues (1984) evaluated broiler chickens for *C. jejuni* colonization. They found that 10 of 12 flocks sampled did not produce positive samples when chicks were at 2 weeks of age. As the birds aged, the flocks became colonized, but the source of colonization could not be found. Stern and coworkers (1995) studied the role of transport and holding on *Campylobacter* colonization rates. In their results, they found that 9 of 10
farms sampled had broilers contaminated with *Campylobacter*. When birds were examined before and after transport and holding, a higher level of contamination occurred in birds that had been transported and held. Pearson and colleagues (1993) investigated an outbreak of *C. jejuni* in Bournemouth, United Kingdom. The outbreak was associated with a catering college that was supplied chicken from a single wholesaler. Pearson and colleagues traced the *C. jejuni* back to a farm that had a contaminated water supply.

Many health risks are possible if food is not prepared properly since as few as 500 *Campylobacter* cells can cause infection. Symptoms of an infection usually consist of headaches, muscle pain, nausea, fever and bloody diarrhea. The Food and Drug Administration estimates that 2 million individuals become infected with *Campylobacter* annually, resulting in approximately 500 deaths. The United States Department of Agriculture estimates the annual cost due to infections by *Campylobacter* at 0.7 to 4.3 billion dollars. In order to reduce *Campylobacter* in turkey products, the frequency of *Campylobacter* colonization in live turkeys must be reduced. The objectives of this study were to 1) determine the level of *Campylobacter* throughout turkey production, 2) to assess strain and gender effects, and 3) to determine when the flock becomes colonized. Our goal was to determine when and where preharvest controls could be implemented to reduce the frequency of *Campylobacter*. In conclusion by possibly chlorinating the water in the house while birds are in pens, dipping boots before and after entering the house at all times and using methods like competitive exclusion when poults are first hatched may reduce the frequency of *Campylobacter* or even eliminated it. It was best said by Bailey
(1993), “The only method to reduce or eliminate *C. jejuni* at the processing plant is to achieve a bird that is *C. jejuni* free”.
Material & Methods

Placement

Three flocks were examined from May 2000 to March 2001. The first 2 flocks were housed for a period of 6 weeks and for each flock, poults were placed in the house at 2-3 days of age and removed 6 weeks later. Flock 1, occupied 12 pens within the facility (Fig. 1) and was placed on fresh wood shavings. Flock 1 (F1) was sampled at weeks 0, 3 and 6 of production. Flock 2, also occupied 12 pens within the facility (Fig. 1) and were placed on used wooden shavings after removal of F1 birds. Litter was aerated and disinfected with TEMPO®, an insecticide, before the placement of F2. Samples from F2 were collected on weeks 0, 1, 2, 3, and 6 of production. Poults in flock 3 (F3) were housed from 2-3 days of age to 20 weeks, and they were placed in two stages, designated as placement 1 and 2 (Fig. 2). Placement 1 was located at the south end of the facility, and placement 2 poults were housed on the north end of the facility 4 weeks later. First placement poults were sampled at weeks 0, 4, 7, 12, 18 and 20, and second-placement poults were sampled at weeks 0, 3, 5, 8, 14 and 20. At week 0 for each flock, gastrointestinal tracts and box liners were sampled, and for all other periods, fecal droppings and water from drinkers were sampled. Due to the separation between placement of the F3 poults, data were collected from both ends of the building and compared (Fig. 2). The F3 sampling schedule was performed in the manner previously described to allow for both placements to be sampled on one day when overlap began in the production of the birds. Fresh wood shavings were provided as litter before poults arrived for F3 placements.
Sample Collection and Transport

Samples consisted of gastrointestinal tracts (GI), box liners, drinkers and fecal droppings. Throughout the study, all flocks were sampled in the same manner. On week 0, entire gastrointestinal tracts and 5 cm\(^2\) sections of box liner were sampled.

For GI tracts, poult's were euthanized by cervical disarticulation. A thin layer of skin was removed from the tip of the keel to the base of the neck, and cranially, to the cloaca, caudally, exposing the breast and abdomen. The abdominal cavity was opened with sterile scissors to expose the GI tract. The GI tract was aseptically removed with a pair of sterile tweezers, it was cut into pieces and placed in a stomacher bag (Fisher Scientific, Pittsburgh, PA). Nutrient broth (NB) \# 2 (Appendix III), was added (100-mL) to each bag containing GI tracts and sealed with an ANPRO bag sealer. Bags with GI tracts were placed on ice for transportation.

A 5-cm\(^2\) section was removed from each box liner with sterile scissors. Once removed, it was placed inside a stomacher bag with 25 ml of NB #2. The bag was sealed by the bag sealer and placed on ice for transport.

Fecal droppings and drinker samples were collected throughout the remainder of the production period for each flock. During this collection process, 5 fecal droppings and one drinker sample was collected from each pen. Warm and moist fecal samples were collected in sterile Whirl-pak\textsuperscript{®} bags (Fisher Scientific, Pittsburgh, PA), using sterile gloves and samples were placed on ice for transport. Sampling drinkers consisted of collecting 100-mL of water from each drinker with a 50-mL pipette. These samples were stored in a 250-ml sterile plastic bottle and placed on ice for transport. Samples were
transported from the Reymann Memorial farm in Wardensville, WV to the West Virginia University Poultry lab in Morgantown, WV, and transport did not exceed 3.5 h.

**Isolation and Confirmation**

GI tracts were placed in a stomacher blender (Model 400, Tekmar, Fisher Scientific, Pittsburgh, PA) and mixed for 30 s. Following mixing, 0.1 mL was plated onto a Campy-Cefex agar (Appendix II) plate. Box liner samples were placed in a sterile stomacher bag and blended for 30 s. Following thorough mixing, 0.1 mL was directly plated onto a Campy-Cefex agar plate. For fecal samples, 1g was combined with 10 mL of NB #2 (Appendix III) in a Whirl-pak® bag (Fisher Scientific, Pittsburgh, PA). The sample was hand massaged for 30s, and a 0.1 mL sample was directly plated on a Campy-Cefex agar plate. For drinker samples, 50 mL of the 100 mL water sample was poured into a Gelman 300 mL, magnetic filter funnel (Gelman, Ann Arbor, MI). The sample was filtered through a 47 mm diameter, 0.45 um grid GN-6 metrical sterile filter (Gelman, Ann Arbor, MI). Once filtered, the filter was aseptically transferred, face down, to a Campy-Cefex agar plate. All agar plates were placed in a 3.79L zip-lock® bag. Each bag, containing 10 plates, was flushed with microaerophilic air (85% N₂, 10% CO₂ and 5% O₂), sealed, and placed into an Imperial II incubator 422 (Labline Instruments, Inc., IL) for 36h at 42°C. After 36h, plates were examined for small, white, translucent colonies. Plates showing no characteristic colonies or no growth were discarded. Plates showing characteristic colonies were saved and a colony was picked and streaked to isolation onto a fresh Campy-Cefex agar plate. Once all plates were streaked to isolation, they were again placed into a 3.79L zip-lock® bag, flushed with microaerophilic air and returned to the Imperial II incubator 422 (Labline Instruments, Inc., IL) for an additional
24-h at 42°C. After 24 h, plates were again examined for characteristic small white translucent colonies. If plates had no growth, plates from the previous incubation period were used to streak for isolation again to confirm that the first transfer missed no bacteria. All plates showing characteristic colonies were confirmed as *Campylobacter* by selecting a single colony, in some cases 2-3 colonies, and using a campy latex agglutination test (Appendix IV) which is specific for *C. jejuni*, *C. laridis* and *C. coli*.

**Long-term Storage**

A single colony, from the same area of the plate that the colony for confirmation was selected, was transferred to a tube containing Protect beads (Appendix V). The tube was sealed and shaken 6 to 7 times and glycerol was aspirated off. All samples were stored at -80°C until needed.

**Statistical Analysis**

A straightforward-randomized design was used to evaluate the data from both 6-week trials to test the effect of flock on frequency of detection. Orthogonal comparisons were used to test linear and quadratic relationships between frequency of colonization and week of production. For the 20-week flock, a randomized complete block design with a split plot arrangement of treatments was employed. The whole plot factor was 2 line X 2 gender treatment combinations and the subplot factor was sampling period. The survey was repeated twice; once on the northern end and once on the southern end of the facility (Fig. 2). For each placement, 3 replications of the 2 X 2 combination was used with pen as the experimental unit. Analysis of variance was performed using Proc GLM of SAS and linear, quadratic, and cubic effects were tested for effect of time on frequency of detection (SAS version 8, SAS Institute Inc., Cary, NC, USA. 1999).
Results

Six-week Production Study

In the first phase, frequency of *Campylobacter* isolation was affected by time for both six-week flocks (Table 1). At week 0, all samples collected from both flocks were *Campylobacter* negative (Table 1). Sixty-five percent of the F1 fecal droppings were positive by week 3 of production. *Campylobacter* peaked at week 3 for flock 1. Frequency of *Campylobacter* detection then began a gradual decline over time. By week 6 frequency of *Campylobacter* detected had dropped to 55%. In F2, a similar pattern of *Campylobacter* detection was observed. However, F2 had lower numbers of positive samples than F1. By week 3, 8.3% of the F2 fecal droppings collected were confirmed positive. Frequency of detection decreased to 0% by week 6. This decrease can not be explained by the data collected in this study. Lower levels of detection in F2 may have been due to housing. Flock 2 was housed on previously used litter. Although the litter had been treated with TEMPO®, an insecticide, and aerated, it may have been possible for poult to obtain existing microflora from the litter. This mature microflora may have out competed *Campylobacter* in the GI tract, resulting in reduced colonization. *Campylobacter* was not isolated from drinker samples (Table 1) in F1 until week 6 of the production cycle. At week 6, the frequency of *Campylobacter* isolation was 41.7%. In F2, drinker samples were contaminated with *Campylobacter* by week 3 at a frequency of 25%. Detection declined over time to 16.7% by the end of the production cycle. The peak and subsequent decline in the frequency of *Campylobacter* in drinker samples of F2 coincided with fecal dropping data.
Twenty-week Production Study

In the second phase of the study, a 20-wk production cycle was evaluated. Data sets were collected and compared on 1\textsuperscript{st} and 2\textsuperscript{nd} placement poult in the southern and northern ends of the facility, respectively. For both placements, week-0 samples were negative (Table. 2). Frequency of \textit{Campylobacter} detection in fecal droppings was 100\% by week 4 of production (Table. 2). Frequency of \textit{Campylobacter} declined gradually, and by the end of the production cycle, week-20, positive samples had fallen to 55\% (Table. 2). In the second placement birds, a slower rate of colonization was observed. The majority of the birds were colonized at week 5, and frequency of \textit{Campylobacter} detection increased to 93.3\% by week 8. Frequency of \textit{Campylobacter} detection declined to 50\% by week 20 (Table. 2). A similar trend was observed among drinker samples for both placements. Positive samples peaked between weeks 4 and 5 and then gradually declined throughout the remainder of the production cycle. Frequency of positive drinkers did not fall below 50\% except at week-7 of production, where the frequency of \textit{Campylobacter} dropped to 41.7\% (Table. 2). Perhaps this was due to farm personnel cleaning the drinkers prior to week 7 sampling. The effect of gender was evaluated and the data indicated that tom turkeys had a slightly higher frequency of \textit{Campylobacter} than hen turkeys, 57.5\% compared to 55.3\% respectively. These data do not agree with data collected in a preliminary study carried out earlier in 1999 (Appendix X). In the preliminary study, tom turkeys had a frequency of \textit{Campylobacter} at 40\% and hens had a frequency of 22.5\%. The studied indicated that tom turkeys carry a higher frequency of \textit{Campylobacter} compared to hens, which was not seen in this study. An explanation for the increased frequency of \textit{Campylobacter} seen in tom turkeys of the preliminary study may have been
due to the fact that genders were separated by putting toms on one side of the house and hens on the other. In this study, toms and hens were only separated by pens, allowing for toms and hens to be influenced by the presence of one another. This influence may have allowed for cross contamination to occur or stress to build in the birds resulting in a weaker defense against the pathogen. Two distinct lines of turkeys were also compared in this study. T1 and Big 6 turkeys were housed in alternating pens in the facility. Pen assignments are shown in (Appendix III). Frequency of Campylobacter detection was not affected by gender, but the Big 6 line showed a slightly higher level than the T1 line, (57.2% versus 55.6%).
Discussion

Our data show that *Campylobacter* is a food safety challenge to the industry. It is currently being isolated from turkeys as it has been in chickens for the past several years. This study indicated that flocks became colonized with *Campylobacter* by 3 to 4 weeks of production. Our findings are in agreement with many other related studies (Acuff, et al., 1982; and Jacobs-Reitsma, et al., 1995). These investigators found that, in both broilers and turkeys, the majority of colonization occurs between 3 to 4 weeks of production. By weeks 3 to 4, the majority of the water samples became colonized, and the frequency of *Campylobacter* declined gradually in later weeks. Pearson and coworkers (1993) reported that water is a major source by which whole flocks of poultry become colonized with *Campylobacter*.

Data from this study showed that gender and line had no effect on the frequency of *Campylobacter* detected. In a preliminary study (Appendix X), an increase in the frequency of *Campylobacter* was seen in the toms, when compared to hens. This discrepancy in our findings may have been due to the arrangement of the birds in the house, since in F3 birds, toms and hens were placed adjacent to each other and in the preliminary study the toms were on one side of the house and hens were on the other. Tom turkeys may contribute to the frequency of *Campylobacter* found in the hens due to cross contamination between pens or that when genders are mixed, a higher level of excitement is reached amongst the birds making them more susceptible to this pathogen.

The data from this study failed to indicate a route whereby *Campylobacter* is able to colonize turkeys. In a study by Pearson and coworkers (1996) it was indicated that vertical transmission might be a route whereby *Campylobacter* can be transmitted,
because high levels of *Campylobacter* were isolated from fecal excretions. Jacobs-Reitsma and colleagues (1995) suggested that vertical transmission is not likely to occur. Transmission among flocks would be better explained by horizontal transmission from contaminated boots, feed, water, insects or other animals, which may come into contact with the birds. Humphrey and coworkers (1993) found that simply dipping boots into a disinfectant before entering a house could delay or possibly prevent *Campylobacter* from colonizing the birds. Our data indicated that turkeys placed on used litter had a lower frequency of *Campylobacter* than turkeys placed on fresh litter. Used litter may already contain a mature microflora. Due to this microflora, *Campylobacter* is out competed in the gastrointestinal tracts of turkeys, which results in the observed lower frequency. This does not suggest that the microflora out competing *Campylobacter* is not another pathogenic organism that can have the same effects as *Campylobacter*. Another possibility to why used litter has a lower frequency of *Campylobacter* than fresh litter may be due to its appearance. Used litter may not be as eye-catching to the poults as fresh litter, which would result in less pecking and initial colonization of the chicks, which was observed in our F2 poults. The main goal of this study was to establish baseline information on frequency of *Campylobacter* detection in turkeys that could assist establishment of a program to reduce or eliminate *Campylobacter* from turkey production facilities. In summary, the frequency at which *Campylobacter* was isolated supports the implementation of on-farm practices to reduce the levels of this organism in birds prior to entering the processing facility.
Tables
Samples Collected from two 6-wk studies

Table 1. Frequency of Campylobacter isolation from GI tracts, box liners, fecal droppings, and drinkers

<table>
<thead>
<tr>
<th>Period</th>
<th>GI tracts</th>
<th>Box liners</th>
<th>Fecal Droppings</th>
<th>Water</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>no. positive/no. sampled</td>
<td>% positive</td>
<td>no. positive/no. sampled</td>
<td>% positive</td>
</tr>
<tr>
<td><strong>1st Flock</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 wk</td>
<td>0/36 0%</td>
<td>0/15 0%</td>
<td>39/60 65%</td>
<td>0/12 0%</td>
</tr>
<tr>
<td>3 wk</td>
<td></td>
<td></td>
<td>31/60 52%</td>
<td>5/12 41.7%</td>
</tr>
<tr>
<td>6 wk</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total</td>
<td>0/36 0%</td>
<td>0/15 0%</td>
<td>70/120 58%</td>
<td>5/24 21%</td>
</tr>
<tr>
<td><strong>2nd Flock</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 wk</td>
<td>0/36 0%</td>
<td>0/15 0%</td>
<td>0/60 0%</td>
<td>0/12 0%</td>
</tr>
<tr>
<td>1 wk</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>2 wk</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>3 wk</td>
<td></td>
<td></td>
<td>6/60 10%</td>
<td>3/12 25%</td>
</tr>
<tr>
<td>6 wk</td>
<td></td>
<td></td>
<td>0/60 0%</td>
<td>2/12 17%</td>
</tr>
<tr>
<td>Total</td>
<td>0/36 0%</td>
<td>0/15 0%</td>
<td>6/240 2.5%</td>
<td>5/48 10%</td>
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</table>
Samples collected from 20-wk study

Table 2. Frequency of *Campylobacter* isolation from GI tracts, box liners, fecal droppings, and drinkers

<table>
<thead>
<tr>
<th>Period</th>
<th>GI Tracts</th>
<th>Box liners</th>
<th>Fecal Droppings</th>
<th>Water</th>
</tr>
</thead>
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<tr>
<td></td>
<td>no. positive/no. sampled</td>
<td>% positive</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1st Placement</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0 wk</td>
<td>0/36</td>
<td>0%</td>
<td>0/12</td>
<td>0%</td>
</tr>
<tr>
<td>4 wk</td>
<td>60/60</td>
<td>100%</td>
<td>11/12</td>
<td>91.70%</td>
</tr>
<tr>
<td>7 wk</td>
<td>55/60</td>
<td>91.70%</td>
<td>5/12</td>
<td>41.70%</td>
</tr>
<tr>
<td>12 wk</td>
<td>57/60</td>
<td>95%</td>
<td>12/12</td>
<td>100%</td>
</tr>
<tr>
<td>18 wk</td>
<td>38/60</td>
<td>63.30%</td>
<td>9/12</td>
<td>75%</td>
</tr>
<tr>
<td>20 wk</td>
<td>33/60</td>
<td>55%</td>
<td>7/12</td>
<td>58.30%</td>
</tr>
<tr>
<td>Total</td>
<td>0/36</td>
<td>0%</td>
<td>0/12</td>
<td>0%</td>
</tr>
<tr>
<td></td>
<td>243/300</td>
<td>81%</td>
<td>44/60</td>
<td>73.30%</td>
</tr>
<tr>
<td>2nd Placement</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>0 wk</td>
<td>0/36</td>
<td>0%</td>
<td>0/12</td>
<td>0%</td>
</tr>
<tr>
<td>3 wk</td>
<td>24/60</td>
<td>40%</td>
<td>4/12</td>
<td>33.30%</td>
</tr>
<tr>
<td>5 wk</td>
<td>47/60</td>
<td>78.30%</td>
<td>11/12</td>
<td>91.70%</td>
</tr>
<tr>
<td>8 wk</td>
<td>56/60</td>
<td>93.30%</td>
<td>12/12</td>
<td>100%</td>
</tr>
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<td>14 wk</td>
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<td>20 wk</td>
<td>30/60</td>
<td>50%</td>
<td>7/12</td>
<td>58.30%</td>
</tr>
<tr>
<td>Total</td>
<td>0/36</td>
<td>0%</td>
<td>0/12</td>
<td>0%</td>
</tr>
<tr>
<td></td>
<td>200/300</td>
<td>66.70%</td>
<td>46/60</td>
<td>76.70%</td>
</tr>
</tbody>
</table>
Figures
Figure 1

Pen Configuration for 6-wk Trials

North End

South End

2nd half of flock

Entrance

1st half of flock
Figure 2.

Housing Design for 20-wk Trial
Chapter 2

Vertical Transmission as a route of *Campylobacter* colonization in turkeys
**Introduction:**

*Campylobacter* colonizes the intestinal tract of poultry. Colonization by this organism may result in carcass contamination during processing and it may potentially spread and cause disease in humans. Initially, *Campylobacter* was known as *Vibrio* and only posed risk to cattle and sheep where stillborn deaths were experienced. Since the 1970’s, research on this pathogen has been extensive. It is known that poultry shares a commensal relationship with *Campylobacter*, thus making poultry a major reservoir for this pathogen.

The actual pathway whereby poultry become colonized is still an enigma. Many studies have looked at vertical transmission as the pathway by which colonization occurs. Shanker and colleagues (1986) evaluated the role of vertical transmission in *C. jejuni* colonization of a broiler flock. They stated that, in 6 breeder flocks, 74% of the birds were confirmed positive for carrying *C. jejuni*. Out of 187 eggs sampled from those breeders, only 2 were identified as *Campylobacter* carriers. Thus, vertical transmission was not considered a possible route of infection in broilers. Acuff and coworkers (1982) looked for *Campylobacter* in turkey eggs, poults and brooding houses. When fertile eggs and newly hatched poults were sampled, neither harbored *Campylobacter*. Pearson and coworkers (1996) sampled two hatcheries to determine if *Campylobacter* could be isolated. They found that the first hatchery had 17.6% *Campylobacter* positive samples. In the second hatchery, 42.9% of the samples were positive. These data support the idea that vertical transmission is a mechanism whereby *Campylobacter* colonizes chicks.

The objective of this study was to assess the possibility of vertical transmission as a pathway whereby *Campylobacter* colonizes turkeys. An accurate, sensitive, DNA-based
technique known as flaA SVR (short variable region) sequence typing was used. This technique is a molecular tool that concentrates on specific flagellin genes that are highly conserved and have variable regions present. Results from the study will highlight the possibility of transmission from parents to offspring and will emphasize that for thorough control of *Campylobacter* in turkeys interventions must occur that prevent contamination of the egg.
Materials and Methods

Resuscitation of Environmental Samples

Environmental samples from parent and poults were randomly selected for *flaA* SVR (short variable region) sequence typing. Pure cultures isolated from parent and offspring were removed from an -80°C freezer and a single bead was streaked onto a Campy-Cefex agar (Appendix II) plate. Plates were sealed in a 3.79L Zip-lock® bag and flushed with microaerophilic air (5% O₂, 10% CO₂ and 85% N₂). Flushed bags were then placed into an Imperial II incubator 422 (Labline Instruments, Inc., IL) overnight at 42°C to allow for growth. Plates were removed after incubation and a single colony was selected and streaked to isolation on a fresh Campy-Cefex agar plate. Plates were placed in Zip-lock® bags, and bags were flushed with microaerophilic air; flushed bags were incubated for an additional 24 h. After 24 h, plates were removed from the incubator, and a single colony was selected from the plate and transferred to Wang’s transport medium (Appendix VI). This sample was incubated for 24 h, packed into microbiological containers, and shipped to the USDA-ARS lab in Athens, GA for *flaA* SVR sequence typing.

Sample Preparation

*Campylobacter* was streaked to isolation on a Campy-Cefex agar plate. Plates were placed in Zip-lock® bags, flushed with microaerophilic air, and incubated at 42°C overnight. Plates were removed the following day; one colony from the plate was selected and streaked once more to gain confluent growth on a Campy-Cefex agar plate. Plates were incubated as previously described. After 24 h, all growth was removed from the plate with a sterile loop and transferred into a 0.6-mL microcentrifuge tube containing 300 µL of sterile water. Tubes were heated to 100°C for 10 min to allow cell walls to
rupture. Boiled samples were stored at 4°C until used as a template for the polymerase chain reaction (PCR).

**Polymerase Chain Reaction (PCR)**

The thermocycler was preheated and 10 microliters of the boiled sample was placed into a sterile microcentrifuge tube. Ninety microliters of a reaction premix (Appendix VII), using the FLA245FU (5'CTA TGG ATG AGC AAT TWA AAA T3’) and FLA625RU (5’CAA GWC CTG TTC CWA CTG AAG3’) primers (Meinersmann, et al., 1997), were added to the tube, and the mixture was covered with 2 drops of sterile mineral oil in place of a heated lid. The microcentrifuge tubes were capped and placed in a Perkin-Elmer 480 thermocycler (Perkin-Elmer-Applied Biosystems Inc., Foster City, CA.) programmed for a 35-cycle reaction system. The 35-cycle reaction was 1 min for denaturing at 96°C, 1 min for annealing at 52°C, and 1 min for extension at 72°C, yielding an approximate 425bp product. Product was purified to remove excess primers, dNTP’s, and buffers using a QIAquick PCR purification kit (Qiagen Inc., Chatsworth, CA). Purified samples were sent to a core lab facility and sequenced using either the FLA242FU primer or the FLA625RU primer with the Big-Dye Dye-Terminator Cycle Sequencing Kit (ABI-PE, Foster City, CA). Data were assembled with Sequencher 4.1 (GeneCodes Corp., Ann Arbor, MI) and aligned using ClustalX (Thompson, J., et al., 1994). Once samples were sequenced and aligned, they were compared. Dendograms were generated using the unweighted pair-group method with arithmetic means (UPGMA) algorithm with HKY85 distance measurements in phylogenetic analysis using parsimony (PAUP® 4.0) (Swofford, D., 1988).
Results

The analysis of DNA sequences was exhaustive (repeated many times), using a variety of different algorithms. Analyses performed on the sequences were Parsimony analysis, Absolute distance measurements and unweighted pair-grouped method with arithmetic means (UPGMA) cluster analysis. All analyses resulted in identical dendograms. The dendogram (Fig. 1) indicated there were 5 instances among the parent flock samples where more than one isolate had a sequence identical to another isolate being sampled. These relationships are clonal, thus there were 5 multiple clones. Six multiple clones were found among samples collected from the offspring. Most offspring samples were found to have identical flaA SVR, DNA sequences when compared to parent samples. Identical DNA sequences indicated that samples were clonal (closely related) to each other. Offspring samples 075, 077, 084, 101, and 105, which contained identical sequences to each other and samples 081, 092, and 095, which also contained identical sequences to each other, showed no relationship to any of the parental genotypes. This indicated that those offspring samples were distantly related to parent isolates sampled in this study. Parent samples 007 and 033 had genotypes that were distantly related to all offspring samples analyzed in the study. Additionally, parent sample 003, 028, 029, and 034 are closely related, one base-pair difference, to offspring samples 043, 044, 048, 055, 056, 059, 061, 062, 064, 065, 068, 071, 074, 080, 087, 088, 091, 093, 096, 102, and 103. This close relationship between parent and poult samples suggests these isolates may be clonal, this base-pair difference may be due to genetic drift (random mutation).
Discussion

Vertical transmission of *Campylobacter* in turkeys is not defined. Scientists have suggested that vertical transmission is a likely pathway whereby colonization occurs, but data have been limited to support this hypothesis. Jacobs-Reitsma and colleagues (1995) investigated the role of breeder flocks in the epidemiology of *Campylobacter* in poultry production. Campylobacter colonization of breeder flocks suggested a potential role for vertical transmission, but serotype data were not supportive. Shanker and coworkers (1986) evaluated the role of vertical transmission in *C. jejuni* colonization of broiler breeder flocks. Out of 187 eggs sampled from 6 breeder flocks, 2 were identified as *Campylobacter* carriers. Thus, vertical transmission was ruled out as a possible route of contamination.

Advances in technology occur every day, such as faster, more reliable and more sensitive techniques to identify bacteria. These techniques facilitate the search for pathways whereby *Campylobacter* colonizes turkeys. Wassenoar and Newell (2000) stated that there are two methods generally accepted for serotyping, the Penner scheme and Loir scheme. Both techniques give high numbers of untypable strains and are time consuming and technically demanding. Recently, molecular subtyping methods have been developed. These genotyping techniques may become universally available making them a major advantage over older techniques.

The discovery of the flagellin genes in *Campylobacter* that have highly conserved and variable regions facilitated development of DNA-based methods for genotyping. In 1997, Stern and colleagues were able to determine potential reservoirs of *Campylobacter* spp. by comparing flagellin A gene of isolates from broiler production facilities.
Camarda and coworkers (2000) also used the flagellin genes to investigate *C. jejuni* isolates from the intestines and oviducts of laying hens. Tsang and colleagues (2001) used the flagellum as a potential marker for *C. jejuni* strains associated with Guillain-Barre’ syndrome.

In this study, the *flaA* SVR sequence typing technique was used to investigate the role of vertical transmission in turkeys. Out of 81 samples evaluated, 5 multiple clones were found within the parent flocks and 6 multiple clones were found among offspring samples. Our data indicate that most offspring samples were identical to the parent samples and clonal (closely related).

The goal of this study was to evaluate the possible role of vertical transmission in turkeys, which could assist establishment of a program to reduce or eliminate *Campylobacter* from turkey production facilities. Identical isolates from parents and offspring support vertical transmission as a pathway whereby turkey flocks become colonized. Improved on-farm practices at the grandparent and parent flock levels to reduce colonization by *Campylobacter* are required to minimize the parent as a source of contamination. Eliminating this source will enhance the effectiveness of preharvest controls for horizontal transmission.
Figure 1.

Unweighted pair-grouped method with arithmetic means analysis
Reference List:


Appendices
Appendix I

Pen assignment for Flock 3
20-wk Trial

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<thead>
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<th>Strain</th>
<th>Gender</th>
<th>Pens</th>
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<td>Hens</td>
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<tr>
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<td>Hens</td>
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Appendix II

Campy-Cefex agar

**Basal medium**

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<td>Brucella agar</td>
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<tr>
<td>Ferrous sulfate (FeSO(_4)7H(_2)O)</td>
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</tr>
<tr>
<td>Sodium bisulfite</td>
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<td>Sodium pyruvate</td>
<td>0.5g</td>
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<td>dH(_2)O</td>
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**Supplements**

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<td>Sodium cycloheximide</td>
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</tr>
<tr>
<td>Lysed horse blood</td>
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</table>

**Replacement for cycloheximide**

- Nystatin
- Rifampicin

Appendix III

Campylobacter Enrichment Broth

Nutrient broth No. 2 (Oxoid) with 0.6% yeast extract

Lab-Lemco powder (Oxoid L29) 10g
Peptone 10g
NaCl 5g
Yeast extract 6g
Distilled water 950 ml

Autoclave 15 min at 121°C in graduated bottles. Use broth within 1 month of preparation (preferably less than 2 weeks). Media will absorb O₂ during storage, which can inhibit recovery of microaerophiles. Keep bottles tightly closed. Before use add 50-ml fresh or frozen-fresh lysed horse blood (5%), 4-ml high concentration FBP, and 4 ml of appropriate antibiotic concentrate (solutions made separately). Store powdered media tightly closed in cool, dry area to reduce oxygen infusion and peroxide formation. Final pH, 7.5 ± 0.2.

Appendix IV

INDX®-Campy (jcl)™

Culture Confirmation Test for
Campylobacter jejuni, C. coli, and C. laridis
Catalog #2200-01-50

Materials:

INDX®-Campy (jcl)™ Latex Detection Reagent (2 X 3.50ml)- consists of rabbit antiserum to common antigens of selected Campylobacter species bound to latex particles suspended in buffer containing a preservative.

INDX®-Campy (jcl)™ Extraction Reagent (2.80ml)- a dilute solution of hydrochloric acid.

INDX®-Campy (jcl)™ Neutralization Reagent (280ml)- Glycine buffer containing a preservative.

INDX®-Campy (jcl)™ Positive Antigen Control Reagent (2.70ml)- consists of a neutralized acid extract of appropriate Campylobacter organisms in buffer containing a preservative.

Test slide

Applicator sticks

High intensity lamp

Slide rotator

Procedure:

1. Remove the reagents from the refrigerator and allow them to warm to room temperature before use.

2. Label one circle on the test slide for each specimen to be tested.

3. Identify one circle for the positive control and another circle for the negative control reactions.

4. Remove the cap and tip protector from the vial of Extraction Reagent. While holding the vial in a vertical position, dispense one free-falling drop of Extraction Reagent
into each specimen circle and the negative control circle. Replace the tip protector and cap.

5. Touch one isolated colony with the end of a wooden applicator stick to remove it from the agar surface. Generally, one colony with a diameter of 2mm (about the diameter of the applicator stick) will provide an adequate inoculum. If colonies are small, yet distinct from the surface of the agar, it may be necessary to pick 2-6 colonies. However, care must be taken as too much inoculum may contribute to poor readability.

6. Make a homogeneous suspension by rotating the inoculum containing stick in the Extraction Reagent within the appropriate specimen circle. **It is very important to dissociate all visible clumps of the inoculum and distribute the suspension over the entire area within the circle.** Repeat steps 5 and 6 for each specimen to be tested. No incubation time is required for this step. Proceed to step 7.

7. Remove the cap and tip protector from the vial of Neutralization Reagent. While holding the vial in a vertical position, dispense one free falling drop of Neutralization Reagent into the fluid spread in each specimen circle and the negative control circle. Replace the tip protector and cap.

8. Remove the cap from the Positive Control Reagent and wipe the tip with a clean lint-free tissue. While holding the vial in a vertical position, dispense one free-falling drop into the positive control circle. Replace the cap.

9. Gently resuspend the Latex Detection Reagent to assure a homogeneous suspension. Do not shake the reagent and avoid the formation of foam or bubbles.

10. Remove the cap from the Latex Detection Reagent and wipe the tip with a clean lint-free tissue. While holding the vial in a vertical position, dispense one free-falling drop of Latex Detection Reagent into each circle, as appropriate, on the slide. Avoid foaming bubbles on the dropper tip as the latex reagent is dispensed. Do not touch the tip of the dropper vial to the material on the slide. Replace the cap.

11. At this point each circle will have received the following:

<table>
<thead>
<tr>
<th>Specimen</th>
<th>Negative Control</th>
<th>Positive Control</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Extraction Reagent</td>
<td>1. Extraction Reagent</td>
<td>1. Positive Control Reagent</td>
</tr>
<tr>
<td>2. Colony(ies)</td>
<td>2. Neutralization Reagent</td>
<td>2. Latex Detection Reagent</td>
</tr>
<tr>
<td>3. Neutralization Reagent</td>
<td></td>
<td>3. Latex Detection Reagent</td>
</tr>
<tr>
<td>4. Latex Detection Reagent</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

12. Use a separate applicator stick to mix the contents of each circle thoroughly.

13. Place the slide on a rotator and rotate at 100-110 rpm for 5 minutes at room temperature.

14. After rotation is completed, immediately observe the reactions for visible agglutination under a high intensity light.
15. A positive test is indicated when the Latex Detection Reagent clearly agglutinates with the test specimen and no agglutination occurs in the negative control circle. The presence of agglutination in the negative control circle renders the test invalid.

16. A negative test is indicated by the absence of agglutination of the Latex Detection Reagent with the test specimen.

INDX-Integrated Diagnostics, Inc. Baltimore, MD 21227 USA.
Appendix V

Protect™ Bacterial Preservers

Composition:

Protect is a sterile vial containing chemically treated porous beads in a cryopreservative fluid of TSB + glycerol with a hypertonic additive. Each bead serves as carrier for the culture during storage.

Procedure for Preparation:

1. Remove the cap being careful not to contaminate the contents. (the GRIPPER helps with this) Inoculate the PROTECT vial with young (18-24 hours) colonial growth of a pure culture of the organism being preserved, to approximate a McFarland 3-4 standard, using a sterile loop. (optional method) Use a sterile pipette to harvest and emulsify the colonies into a PROTECT vial by using a squeezing action. The same pipette may then be used to extract the excess fluid. (Step 3) Liquid cultures can be lightly centrifuged and the deposit used as above.
2. Cap the tube and invert 6 times. Do Not Vortex.
3. Let vial stand for 30 seconds. The organism will now be bound to the beads. Remove the excess cryopreservative fluid leaving the beads as free of liquid as possible. A sterile pipette is best for this procedure. Close the vial finger tight.
4. Record the culture details on the vial and store at minus 70°C.

Procedure For Use – Recovery:

1. Remove the vial from the freezer.
2. Carefully open the vial and remove a single bead with a sterile needle, forceps, or the special PROTECT hook available from KEY Scientific.
3. Rub the bead over solid medium, streak from point of contact, or drop into appropriate growth broth. Some organisms perform better using broth method. Beads should not be returned to the vial after removal.

Medical Laboratory Sciences, 1985, 42:289-290.
Appendix VI

Wang’s Transport Storage Media

Brucella broth 28 g
Brucella agar 4 g
DH2O 950 mL

Bring mixture to a boil and autoclave
Temper solution to 50°C

Lysed horse blood 50 mL
DH2O 50 mL

Add diluted lysed horse blood to the tempered solution

Correspondence:
Stern, N.J., 2000. USDA-ARS Russell Research Center
950 College Station Rd. Athens, GA 30604
Appendix VII

PCR Reaction Premix

Prepare PCR Reaction Premix in a 1.5 mL microcentrifuge tube, for the appropriate number of samples. Include both a positive and a negative control. It is advisable to prepare reaction premix for one additional sample due to possible pipetting errors. Therefore if 10 samples are to be analyzed, prepare a reaction premix for 13 samples (10 samples + 1 positive control + 1 negative control + 1 extra sample).

<table>
<thead>
<tr>
<th>Reagent</th>
<th>1 Sample</th>
<th>5 Samples</th>
<th>10 Samples</th>
<th>20 Samples</th>
</tr>
</thead>
<tbody>
<tr>
<td>10X Buffer</td>
<td>10 uL</td>
<td>50 uL</td>
<td>100 uL</td>
<td>200 uL</td>
</tr>
<tr>
<td>MgCl</td>
<td>6 uL</td>
<td>30 uL</td>
<td>60 uL</td>
<td>120 uL</td>
</tr>
<tr>
<td>dNTP mix</td>
<td>8 uL</td>
<td>40 uL</td>
<td>80 uL</td>
<td>160 uL</td>
</tr>
<tr>
<td>Forward Primer</td>
<td>0.5 uL</td>
<td>2.5 uL</td>
<td>5 uL</td>
<td>10 uL</td>
</tr>
<tr>
<td>Reverse Primer</td>
<td>0.5 uL</td>
<td>2.5 uL</td>
<td>5 uL</td>
<td>10 uL</td>
</tr>
<tr>
<td>Sterile water</td>
<td>64.5 uL</td>
<td>322.5 uL</td>
<td>645 uL</td>
<td>1290 uL</td>
</tr>
<tr>
<td>Taq Polymerase</td>
<td>0.5 uL</td>
<td>2.5 uL</td>
<td>5 uL</td>
<td>10 uL</td>
</tr>
<tr>
<td><strong>Total Volume</strong></td>
<td><strong>90 uL</strong></td>
<td><strong>450 uL</strong></td>
<td><strong>900 uL</strong></td>
<td><strong>1800 uL</strong></td>
</tr>
</tbody>
</table>

950 College Station Rd. Athens, GA 30604
Appendix VIII

SAS Program for Analysis of Six-Week Data

Title 'Aaron Kiess Data - Flock 1&2 Samples From Birds';
options ls=80 ps=52 pageno=1;
proc import datafile="C:\My Documents\kiess files\flocks 1 & 2 birds.xls"
   out=one
   replace;
data two; set one; if week=0 or week=3 or week=6;
proc sort; by flock week pen;
proc means noprint; by flock week pen; var result;
output out=three mean=Mres;

proc glm; classes flock week;
model mres= flock|week;
contrast 'week linear' week -1 0 1;
contrast 'week quadratic' week 1 -2 1;
contrast 'flock' flock -1 1;
contrast 'flock x week linear' flock*week -1 0 1 0 -1;
contrast 'flock x week quadratic' flock*week 1 -2 1 -1 2 -1;
lsmeans flock|week;
means flock|week;
run;
Title 'Aaron Kiess Data - Flock 1&2 Samples From Drinkers';
options ls=80 ps=52 pageno=1;
  proc import datafile="C:\My Documents\kiess files\flocks 1 & 2 pens.xls"
      out=one
      replace;
  data two; set one; if week=3 or week=6;

  proc glm; classes flock week;
  model rslt= flock|week;

  lsmeans flock|week;
  means flock|week;

  run;
Appendix VIII

SAS Program for Analysis of Twenty-Week Data

Title 'Aaron Kiess Data - Flock 3 Samples From Birds 3';
options ls=80 ps=52 pageno=1;
/* samples from pens; tests among genders/lines; 3 sub-blocks within each place;
test linear, quadratic, and cubic effect of week*/
proc import datafile="C:\My Documents\kiess files\flock_3.xls"
  out=one
  replace;
data two; set one; if srce="GI_tract" or srce="Fecal" then src2="Bird"; else src2="Pen";
  if pen=1 or pen=2 or pen=7 or pen=8 then Block=1;
  if pen>=3 and pen<=6 then Block=2;
  if pen>=9 and pen<=12 then Block=3;
  if pen=13 or pen=14 or pen=19 or pen=20 then Block=4;
  if pen>=15 and pen<=18 then Block=5;
  if pen>=21 and pen<=24 then Block=6;
  if src2="Bird";
  proc sort; by plce block gend line pen week;
  proc means noprint; by plce block gend line pen week; var rslt; output out=three
    mean=Mres;
data four; set three; drop _TYPE_ _FREQ_; proc print;
  proc glm; classes plce block gend line week;
  model mres= plce block(plce) gend line line*gend gend*line*block*plce
    week(plce) gend*week(plce) line*week(plce) line*gend*week(plce);
  test h=gend line line*gend e=gend*line*block*plce/etype=1;
  lsmeans plce gend line week(plce);
  means plce gend line week(plce)/deonly;
data five; set three; Wk=week; WkSq=week**2; WkCu=week**3;
  proc glm; classes plce block gend line week;
  model mres= plce block(plce) gend line line*gend gend*line*block(plce)
    week(plce) gend*week(plce) line*week(plce) line*gend*week(plce);
  test h=wk wksq wkcu week(plce)/htype=1 etype=1;
run;
Title 'Aaron Kiess Data - Flock 3 Samples From Drinkers';
options ls=80 ps=52 pageno=1;
/* samples from pens; tests among genders/lines; 3 sub-blocks within each place; test linear, quadratic, and cubic effect of week*/
proc import datafile="C:\My Documents\kiess files\flock_3.xls"
   out=one
   replace;
data two; set one;
if pen=1 or pen=2 or pen=7 or pen=8 then Block=1;
if pen>=3 and pen<=6 then Block=2;
if pen>=9 and pen<=12 then Block=3;
if pen=13 or pen=14 or pen=19 or pen=20 then Block=4;
if pen>=15 and pen<=18 then Block=5;
if pen>=21 and pen<=24 then Block=6;
if srce="Drinker";
proc sort; by plce block gend line pen week;
proc means noprint; by plce block gend line pen week; var rslt; output out=three
   mean=Mres;
data four; set three; drop _TYPE_ _FREQ_; proc print;
proc glm; classes plce block gend line week;
model mres= plce block(plce) gend line line*gend*line*block*plce
   week(plce) gend*week(plce) line*week(plce) line*week(plce); test h=gend line line*gend e=gend*line*block*plce/etype=1;
lsmeans plce gend line week(plce);
means plce gend line week(plce)/deponly;

data five; set three; Wk=week; WkSq=week**2; WkCu=week**3;
proc glm; classes plce block gend line week;
model mres= plce block(plce) gend line line*gend*line*block(plce)
   wk wksq wkcu week(plce);
test h=wk wksq wkcu e=week(plce)/htype=1 etype=1;
run;
Appendix X
Preliminary Study
Effect of gender on frequency of *Campylobacter* isolation from Turkeys
Abstract

*Campylobacter* ingestion can cause campylobacteriosis in humans and most cases have been linked to poultry products. In order to reduce *Campylobacter* in turkey products, frequency of *Campylobacter* colonization in live turkeys must be reduced. From February to March of 2000, one turkey flock at Wardensville, WV was sampled for *Campylobacter*. This flock was the 10th flock produced in the facility. Samples were collected at weeks 18 and 20 of a 21-week production cycle. The facility consisted of 24 pens, divided equally by a service area. Male and female turkeys were segregated in 12 pens on either side of the facility. Fecal droppings and drinkers were sampled to determine frequency of *Campylobacter* contamination. During each sampling period, for each gender, 5 fecal droppings were collected from different locations within each of the 12 pens. Thus, 120 fecal samples were collected at each sampling (24 pens x 5 samples). A 100-mL water sample was collected from each drinker within each pen. Fecal droppings were placed in sterile plastic bags and marked fresh or old. Drinker samples were placed in sterile bottles and all samples were transported on ice to West Virginia University for *Campylobacter* isolation. Water samples were mixed thoroughly; a loopful of the sample was plated directly onto a Campy-Cefex agar plate. Fecal samples were mixed with number 2 nutrient broth (1:10 w/v). The mixture was massaged, and a loopful of the mixture was plated onto a Campy-Cefex agar plate. About 12-14 plates were placed in a 3.79L Zip-lock® bag, and the bag was sealed and incubated at 40 to 42°C for 48h. After 48h, plates were removed and examined for *Campylobacter*. *Campylobacter* positive colonies were round, flat, and translucent. Presumptive positives were confirmed by latex agglutination. *Campylobacter* was present in 26% of all
samples, and it declined from 31% at week 18 to 21% at week 20. *Campylobacter* was detected more frequently in males (40%) than in female turkey (22.5%). *Campylobacter* was not detected in water collected from drinkers. Fresh fecal samples accounted for 32% of all positive samples. Frequency of *Campylobacter* isolation for this flock supports implementation of on-farm practices to reduce levels of *Campylobacter* in birds going to processing facilities.
Introduction

*Campylobacter* is a leading cause of foodborne illness in the United States. It is responsible for infecting over 2 million people each year with an infection known as campylobacteriosis (Center of Disease Control and Prevention. 2000). Humans ingesting fewer than 500 cells may experience symptoms such as fever, headache, stomach pain, bloody diarrhea and in a few cases Guillan-Barre’ Syndrome (GBS) which may cause paralysis (Patterson, 1995). *Campylobacter* is transmitted through foods of animal origin that have been undercooked and/or mishandled. Cross contamination of raw meat with other food items is also responsible for campylobacteriosis.

Raw and processed poultry products have been implicated as the major source of *Campylobacter* transmission to humans (Stern, N.J. and Robach, M.C., 1995). In order to eliminate the infection rate in humans it is necessary to reduce the frequency of *Campylobacter* in live birds and necessitate comprehensive control on the production facilities. Such controls can be implemented by identifying the sources/vectors responsible for *Campylobacter* incidences and establishing Critical Control Points. The objective of this study was to determine the frequency of *Campylobacter* from various preharvest sources within a turkey production facility.
Material and Methods

Placement

One flock was examined from February to March 2000. The flock was housed for a period of 21 weeks and poults were placed in the house at 2-3 days of age and removed 21 weeks later. The flock occupied 24 pens within the facility (Fig. 1) and was placed on fresh wood shavings as litter. Twelve pens on the northern end of the building were occupied by tom turkeys and the southern end of the building was occupied by hen turkeys. Samples were collected at weeks 18 and 21 of production.

Collection and Transport

Samples consisted of drinkers and fecal droppings. Throughout the study, all samples were collected in the same manner. Fecal droppings and drinker samples were collected twice throughout the last three weeks of the production period. During this collection process, 5 fecal droppings and 1 drinker sample was collected from each pen. Warm and moist fecal samples were collected in sterile Whirl-pak® bags (Fisher Scientific, Pittsburgh, PA), using sterile gloves and samples were placed on ice for transport. Sampling drinkers consisted of collecting 100-mL of water from each drinker with a 50-mL pipette. These samples were stored in a 250-ml sterile plastic bottle and placed on ice for transport. Samples were transported from the Reymann Memorial farm in Wardensville, WV to the West Virginia University Poultry lab in Morgantown, WV, and transport did not exceed 3.5 h.

Isolation and Conformation

Fecal samples, 1g was combined with 10 mL of NB #2 (Appendix III) in a Whirl-pak® bag (Fisher Scientific, Pittsburgh, PA). The sample was hand massaged for 30s,
and a 0.1 mL sample was directly plated on a Campy-Cefex agar plate (Appendix II). For drinker samples, the sterile bottle containing the water was shaken vigourously to mix up contents, then a loopful of the water was streaked onto a Campy-Cefex agar plate. All agar plates were placed in a 3.79L zip-lock® bag. Each bag, contained 12 to 14 plates, the bags were flushed with microaerophilic air (85% N₂, 10% CO₂ and 5% O₂), sealed, and placed into an Imperial II incubator 422 (Labline Instruments, Inc., IL) for 36h at 42°C. After 36h, plates were examined for small, white, translucent colonies. Plates showing no characteristic colonies or no growth were discarded. Plates showing characteristic colonies were saved and a colony was picked and streaked to isolation onto a fresh Campy-Cefex agar plate. Once all plates were streaked to isolation, they were again placed into a 3.79L zip-lock® bag, flushed with microaerophilic air and returned to the Imperial II incubator 422 (Labline Instruments, Inc., IL) for an additional 24-h at 42°C. After 24 h, plates were again examined for characteristic small white translucent colonies. If plates had no growth, plates from the previous incubation period were used to streak for isolation again to confirm that the first transfer missed no bacteria. All plates showing characteristic colonies were confirmed as *Campylobacter* by selecting a single colony, in some cases 2-3 colonies, and using a campy latex agglutination test (Appendix IV) which is specific for *C. jejuni*, *C. laridis* and *C. coli*.

**Long-term Storage**

A single colony, from the same area of the plate that the colony for confirmation was selected, was transferred to a tube containing Protect beads (Appendix V). The tube was sealed and shaken 6 to 7 times and glycerol was aspirated off. All samples were stored at -80°C until needed.
Results

The overall frequency of *Campylobacter* was at 26% in this flock (Table 1). It declined from 31% at week 18 to 21% at week 21. *Campylobacter* was detected more frequently in tom turkeys (40%) than in hen turkeys (22.5%). *Campylobacter* was not detected in water collected from drinkers. Fresh fecal samples accounted for 10% of all positive samples isolated. The frequency at which *Campylobacter* was isolated supports the implementation of on-farm practices to reduce levels of this organism in birds going to the processing facilities.
Table
Samples collected from preliminary study

Table 1. Prevalence of *Campylobacter*

<table>
<thead>
<tr>
<th>Sample</th>
<th>Grow-out period</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Wk 18</td>
<td>Wk 21</td>
<td>Total</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Feces</td>
<td>45/120</td>
<td>30/120</td>
<td>75/240</td>
<td>37.50%</td>
<td>25%</td>
</tr>
<tr>
<td>Drinkers</td>
<td>0/24</td>
<td>0/24</td>
<td>0/48</td>
<td>0%</td>
<td>0%</td>
</tr>
<tr>
<td>Total</td>
<td>45/144</td>
<td>30/144</td>
<td>75/288</td>
<td>31.30%</td>
<td>20.80%</td>
</tr>
<tr>
<td>Fresh</td>
<td>16/120</td>
<td>8/120</td>
<td>24/240</td>
<td>13.30%</td>
<td>6.70%</td>
</tr>
<tr>
<td>Male</td>
<td>31/60</td>
<td>17/60</td>
<td>48/120</td>
<td>52%</td>
<td>28.30%</td>
</tr>
<tr>
<td>Female</td>
<td>14/60</td>
<td>13/60</td>
<td>27/120</td>
<td>23.30%</td>
<td>21.70%</td>
</tr>
</tbody>
</table>
CURRICULUM VITAE

Parents
Melvin D. Kiess
Carla E. Kiess

Birthplace
Elkins, WV

Date of Birth
January 27, 1976

Schools Attended
West Virginia University

Degrees Conferred
Master of Science in Animal and Veterinary Science 2001
Bachelor of Science in Animal and Veterinary Science 1999