Raw Comminuted Chicken Sampling Data –
Data Documentation

Overview

The data covered by this documentation are the sampling results of FSIS’ routine microbiological sampling of raw comminuted chicken products. Additional information can be found on the FSIS Laboratory Sampling Data web page.

Data postings are part of the Agency’s efforts to prevent pathogens from entering the food supply throughout the farm-to-fork continuum. Posting these datasets may help industry identify repetitive subtypes and implement control measures. It may allow researchers to identify trends to address basic research questions or to develop new diagnostics or therapies such as vaccines.

These datasets are posted for informational purposes only and are not sufficient to determine if there is an association between multiple samples. Therefore, this data should not be used to identify foodborne illness outbreaks, associate samples with foodborne illness outbreaks or determine whether two or more samples are causally related.

The FSIS number is a unique identifier for retrieving whole genome sequence data from the National Center for Biotechnology and Information (NCBI) Pathogen Detection Isolates Browser. The allele codes included in this dataset provide a convenient naming method for reporting Whole Genome Sequencing (WGS) data. Because allele codes can change over time as more WGS data becomes available, a date stamp allows the data to be used in reports.

FSIS, the Centers for Disease Control and Prevention (CDC) and other public health partners monitor WGS information gathered from samples in real time and use sophisticated epidemiological tools to identify whether the cumulative findings might signal a foodborne illness outbreak. Outbreaks identified through this process are announced to the public through CDC’s website and FSIS’ Outbreak Response Page. The FSIS investigative process is described in FSIS Directive 8080.3.

Two datasets will be provided: archived and current. The archived dataset will provide data starting from October 1, 2013, up to the final day of the previous fiscal year (FY) of the report execution date (e.g., the data posted in April 2022 was through the end of FY21). The archived dataset will be updated annually. The current dataset will provide data starting from the first day of the FY following the final date available in the archived dataset through the end of the previous fiscal quarter of the report execution date (e.g., data posted in April 2022 was through the end of FY22 Quarter 1). The current dataset will be updated quarterly.
Each row in these datasets represents one sample collected and sent to an FSIS laboratory for analysis. Each sample is analyzed for *Salmonella* species and *Campylobacter*. In this dataset, samples analyzed before August 27, 2018, were tested for *Campylobacter* using the direct plate method. Samples analyzed on August 27, 2018, and after were tested for *Campylobacter* using the enrichment method. FSIS announced this change in the [*August 27, 2018, Constituent Update*](https://www.fsis.usda.gov/emergencyidency/constituent-update).

Isolate characterization data will not be publicly posted in the datasets until the full characterization profile is completed.

Data contained in this dataset on tested product from establishments are not sufficient to determine an association with human illnesses. Further epidemiologic information is needed to determine if there is an association among the non-clinical isolates and human illnesses.

**Data Dictionary**

- **EstablishmentID**
  - A unique numeric identifier that is used to identify an establishment across data tables in the FSIS databases.
- **EstablishmentNumber**
  - A letter/number combination uniquely identifying each establishment.
- **EstablishmentName**
  - The name of an establishment on its FSIS grant of inspection.
- **State**
  - The state where the establishment is located.
- **ProjectCode**
  - A short name given to easily identify an FSIS sampling project.
  - Project code used in this dataset:
    - HC_CH_COM01 - Sampling for ground and other comminuted chicken (not mechanically separated)
- **ProjectName**
  - The name of the FSIS sampling project.
- **FormID**
  - The form number used to uniquely identify a specific sample.
- **CollectionDate**
  - The date the FSIS inspector collected the sample at the FSIS-regulated establishment.
- **SampleSource**
  - The type of product collected in the sample.
- **SalmonellaSpAnalysis**
  - The result of the analysis for *Salmonella* species in the sample.
    - Negative = *Salmonella* was not found in the sample.
    - Positive = *Salmonella* was found in the sample.
- **SalmonellaSerotype**
- The name of the distinct variation of the tested species of bacteria. A list of the serotypes that are more commonly associated with human illness can be found on the Centers for Disease Control and Prevention (CDC) web site on their National Salmonella Surveillance web page.

- **SalmonellaPFGEPattern**
  - The specific pattern identified from Pulsed-Field Gel Electrophoresis (PFGE), which is the laboratory technique used to produce a deoxyribonucleic acid (DNA) fingerprint for a group of the same type of bacteria. FSIS discontinued PFGE characterization in March 2019 and replaced the technique with the more modern whole genome sequencing (WGS) characterization.

- **SalmonellaAlleleCode**
  - Definition: A code assigned by CDC-PulseNet based on the number of differences in pre-defined genes in the WGS data. The allele code also includes the date when FSIS retrieved the allele code from PulseNet. It is possible for PulseNet to adjust the allele code after it was retrieved. The data format is: allele code[space][pipe][space]mm/dd/yyyy (e.g., SALM1.0 – 1.2.3.4.5.6 | 01/01/2021). When PulseNet is unable to assign an allele code, the entry will be: Allele Code Ineligible[space][pipe][space]mm/dd/yyyy (e.g., Allele Code Ineligible | 01/01/2021).

- **SalmonellaFSISNumber**
  - A unique identifier for retrieving Whole Genome Sequencing (WGS) data for a Salmonella isolate from the National Center for Biotechnology and Information (NCBI) Pathogen Detection Isolates Browser. NCBI developed the Browser to help users learn about the sequences they contribute. NCBI has provided a video introduction to this browser, and this document contains a table outlining information available in NCBI’S Pathogen Detection Isolates Brower for additional reference.

- **SalmonellaAMRResistanceProfile**
  - The antimicrobial resistance profile of the antimicrobial drugs phenotypically tested to which isolates are found to be resistant using the National Antimicrobial Resistance Monitoring System (NARMS) panel 5. The Food and Drug Administration (FDA) in its Guidance 152 classified antimicrobial drugs based on importance of the drug to human medicine. Isolates displaying resistance to multiple antimicrobial drugs tested by the NARMS panel are classified according to the antimicrobial drug(s) with the highest classification of risk. A resistance profile that is “pan-susceptible” means that the isolate is not resistant to any of the antimicrobial drugs tested. See the FDA Antimicrobial drug classification table in this document.

- **CampylobacterAnalysis1ml**
  - The result of the analysis for Campylobacter species in the sample that was tested using the direct plate method.
    - Negative = Campylobacter was not found in the sample.
    - Positive = Campylobacter was found in the sample.

- **CampylobacterAnalysis30ml**
The result of the analysis for *Campylobacter* species in the sample that was tested using the enrichment method.

- **Negative** = *Campylobacter* was not found in the sample.
- **Positive** = *Campylobacter* was found in the sample.

- **CampylobacterSpecies**
  - Species designation for isolates belonging to the genus *Campylobacter*. Isolates are positive if they are species jejuni, coli, or lari.

- **CampylobacterPFGEPattern**
  - The specific pattern identified from PFGE, the laboratory technique used to produce a DNA fingerprint for a group of the same type of bacteria. PFGE characterization was discontinued in November 2018.

- **CampylobacterAlleleCode**
  - A code assigned by CDC-PulseNet based on the number of differences in pre-defined genes in the WGS data. The allele code also includes a date stamp, the date when FSIS retrieved the allele code from PulseNet. It is possible for PulseNet to adjust the allele code after it was retrieved.
  - **NOTE:** *Campylobacter* allele codes are not expected to be available until FY2023.

- **CampylobacterFSISNumber**
  - A unique identifier for retrieving Whole Genome Sequencing (WGS) data for a *Campylobacter* isolate from the National Center for Biotechnology and Information (NCBI) Pathogen Detection Isolates Browser. NCBI developed the Browser to help users learn about the sequences they contribute. NCBI has provided a video introduction to this browser, and this document contains a table outlining information available in NCBI’S Pathogen Detection Isolates Brower for additional reference.

- **CampYAMRResistanceProfile**
  - The antimicrobial resistance profile of the antimicrobial drugs phenotypically tested to which isolates are found to be resistant using the National Antimicrobial Resistance Monitoring System (NARMS) panel 5. The Food and Drug Administration (FDA) in its Guidance 152 classified antimicrobial drugs based on importance of the drug to human medicine. Isolates displaying resistance to multiple antimicrobial drugs tested by the NARMS panel are classified according to the antimicrobial drug(s) with the highest classification of risk. A resistance profile that is “pan-susceptible” means that the isolate is not resistant to any of the antimicrobial drugs tested. See the FDA Antimicrobial drug classification table in this document.
# FDA’s Antimicrobial Drug Classification According to Their Importance to Human Medicine

<table>
<thead>
<tr>
<th>Antimicrobial Class</th>
<th>Antimicrobial Drug</th>
<th>Abbreviation</th>
<th>FDA Classification</th>
</tr>
</thead>
<tbody>
<tr>
<td>1st Generation Cephalosporins (Cephems)</td>
<td>Cephalothin (Cefazolin)</td>
<td>CEP</td>
<td>Important</td>
</tr>
<tr>
<td>3rd Generation Cephalosporins (Cephems)</td>
<td>Ceftiofur</td>
<td>TIO</td>
<td>Critically Important</td>
</tr>
<tr>
<td></td>
<td>Ceftriaxone</td>
<td>AXO</td>
<td>Critically Important</td>
</tr>
<tr>
<td>Aminoglycosides</td>
<td>Amikacin</td>
<td>AMI</td>
<td>Highly Important</td>
</tr>
<tr>
<td></td>
<td>Apramycin</td>
<td>APR</td>
<td>Highly Important¹</td>
</tr>
<tr>
<td></td>
<td>Gentamicin</td>
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<tr>
<td></td>
<td>Kanamycin</td>
<td>KAN</td>
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</tr>
<tr>
<td></td>
<td>Streptomycin</td>
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<tr>
<td>B-Lactam/B-Lactamase Inhibitor Combinations</td>
<td>Amoxicillin-Clavulanic Acid (Amoxicillin)</td>
<td>AUG</td>
<td>Highly Important</td>
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<tr>
<td>Carbapenems</td>
<td>Imipenem</td>
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<tr>
<td>Carboxypenicillins</td>
<td>Ticarcillin</td>
<td>TIC</td>
<td>Highly Important</td>
</tr>
<tr>
<td>Cephamycins (Cephems)</td>
<td>Cefoxitin</td>
<td>FOX</td>
<td>Important</td>
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<tr>
<td>Fluoroquinolones</td>
<td>Ciprofloxacin</td>
<td>CIP</td>
<td>Critically Important</td>
</tr>
<tr>
<td>Folate Pathway Inhibitors</td>
<td>Sulfamethoxazole (1998-2003)</td>
<td>SMX</td>
<td>Not Classified</td>
</tr>
<tr>
<td></td>
<td>Sulfisoxazole (2004-2009)</td>
<td>FIS</td>
<td>Not Classified</td>
</tr>
<tr>
<td></td>
<td>Trimethoprim-Sulfamethoxazole</td>
<td>COT</td>
<td>Critically Important</td>
</tr>
<tr>
<td>Macrolides</td>
<td>Azithromycin</td>
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</tr>
<tr>
<td></td>
<td>Erythromycin</td>
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<td>Phenicols</td>
<td>Chloramphenicol</td>
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<tr>
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<td>Florfenicol</td>
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<tr>
<td>Quinolones</td>
<td>Nalidixic Acid</td>
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<td>Ketolides</td>
<td>Telithromycin</td>
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<tr>
<td>Lincosamidies</td>
<td>Clindamycin</td>
<td>CLI</td>
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</tr>
<tr>
<td>Penicillins</td>
<td>Ampicillin</td>
<td>AMP</td>
<td>Highly Important</td>
</tr>
<tr>
<td>Tetracyclines</td>
<td>Tetracyclines</td>
<td>TET</td>
<td>Highly Important</td>
</tr>
</tbody>
</table>

**Critically Important (C):** Antimicrobial drugs which meet BOTH criteria 1 and 2 in Appendix A of the FDA Guidance for Industry #152 are considered critically important to human medical therapy.

**Highly Important (H):** Antimicrobial drugs which meet EITHER criteria 1 or 2 in Appendix A of the FDA Guidance for Industry #152 are considered highly important to human medical therapy.

**Important (I):** Antimicrobial drugs which meet EITHER criterion 3 and/or 4 and/or 5 in Appendix A of the FDA Guidance for Industry #152 are considered important to human medical therapy.
**Not Classified (NC):** Antimicrobial drugs which are not given a classification in FDA’s Guidance for Industry #152 (dated October 23, 2003).

1Where noted, FSIS has classified drugs approved for animal use only using the same classification that FDA has designated for drugs in the same antimicrobial class that are approved for human use.

**Relationship to Other Data**

This data can be combined with other FSIS datasets using the EstablishmentID variable.

**Notes and Limitations**

Information about FSIS sampling laboratories and procedures can be found on the FSIS website on the Laboratories & Procedures web page and the Microbiology Laboratory Guidebook (MLG) web page.

NULL values indicate that the specific variable is not available for that record.

Samples analyzed for *Salmonella* and *Campylobacter* may lack results for one organism. Lack of analysis for one organism is due to either insufficient volume to perform both analyses or not being analyzed due to not meeting all receipt temperature criteria ([MLG Chapter 1.01](https://www.fsis.usda.gov/wps/portal/fsis/topics/food-safety-health-consumer/publications/microbiology-laboratory-guidebook)). Delays in sample analysis can occur when a sample is non-viable or when there is a need to repeat the WGS analysis.

When a sample screens positive for a pathogen, there normally is only one isolate (e.g., subtype determined using serology [serotype], PFGE, antibiotic resistance, allele code, or whole genome sequence) derived from laboratory confirmation procedures. During such procedures, the enrichment broth is streaked on agar plates, and those plates are subsequently examined for typical pathogen colonies. The laboratory staff ordinarily picks no more than one typical isolated colony from any one plate. On very rare occasions, more than one typical colony may be picked for confirmation. In such circumstances, the multiple isolate data (e.g., *Salmonella* serotype) are separated by a semicolon.

WGS data must be interpreted within the context of how it will be used, e.g., to detect outbreaks or contamination events. Additional corroborating information, including case-patient food exposure and product distribution records, may be necessary to properly interpret the WGS data.

**Prior Analysis**

Prior analysis using this data can be found on the FSIS website, specifically “Progress Report on *Salmonella* and *Campylobacter* Testing of Raw Meat and Poultry Products, 1998-2014” and the Quarterly Sampling Results on *Salmonella* web page.
### Information Available in NCBI’s Pathogen Detection Isolates Browser

<table>
<thead>
<tr>
<th>Category</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Single Nucleotide Polymorphism (SNP)</td>
<td>A SNP cluster is a group of isolates whose genomes are closely related. In the Pathogen Detection Browser this element will contain a link that opens a new page with information about closely related sequences in the database.</td>
</tr>
<tr>
<td>Min-same or Min-diff</td>
<td>Minimum SNP distance from the query isolate to one of the same or a different isolation type. Isolation types are clinical (including human or animal) or environmental (including food).</td>
</tr>
<tr>
<td>BioSample</td>
<td>Further information (metadata) pertaining to the sample from which the sequence was isolated.</td>
</tr>
<tr>
<td>Assembly</td>
<td>Technical information pertaining to the sequence.</td>
</tr>
<tr>
<td>AMR Genotype</td>
<td>Information pertaining to antimicrobial resistant (AMR) genes found in the isolate sequence. Additional information about each AMR gene in this field is provided by a Reference Gene Catalog. Note: Empty cells do not necessarily indicate a lack of AMR genes.</td>
</tr>
</tbody>
</table>