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Meta-Taxonomic Analysis of Poultry and Slaughterhouse Microbiota: A Comprehensive **Examination of Resident Microbial Communities**

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Meta-Taxonomic Analysis of Poultry and Slaughterhouse Microbiota: A Comprehensive Examination of Resident Microbial Communities

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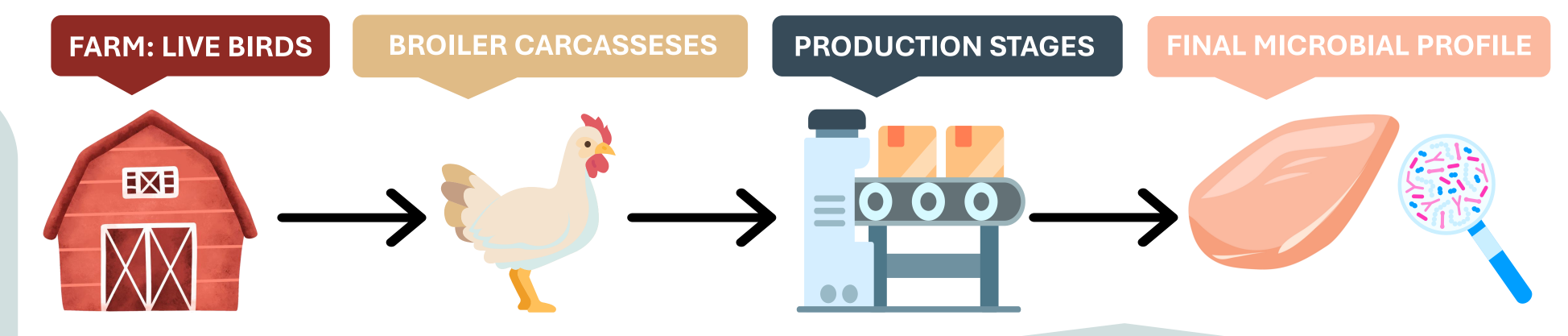
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Poultry contaminations occur during the processing chain and are found from farm to table.

The final microbial profile of poultry meat is influenced by each step of the process.

The consumption of contaminated poultry poses a significant threat to public health. It is associated to infections caused by pathogens such as: Campylobacter, Salmonella, Listeria monocytogenes and Arcobacter.



The **objective** is to assess the contribution of broiler skin and gut to the indigenous slaughterhouse microbiota. In addition, the study aims to clarify how the **persistence** and environmental distribution of Arcobacter spp. are influenced by the crosscontamination patterns associated with these specific sources.

4. BIOINFORMATIC ANALYSIS:

MATERIALS & METHODS

1. BROILERS AND ENVIRONMENTAL SAMPLING

154 samples were collected from a local poultry abattoir

taxonomy was assigned with a 99% of sequence similarity by 49 samples of broiler matching ASVs to the Silva neck skin (BNS) prokaryotic reference database 49 samples of broiler caecum (BC) **V4 V3** 56 from the samples 465 bp **WHERE** slaughterhouse equipment (SE) **North-Italy**

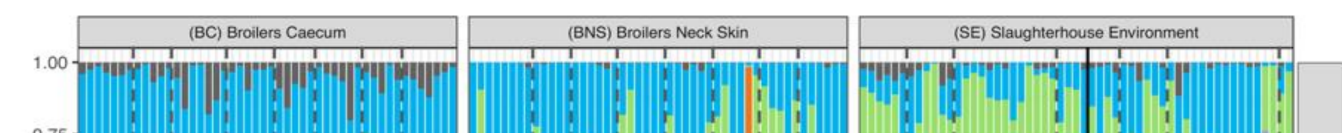
2. DNA extraction

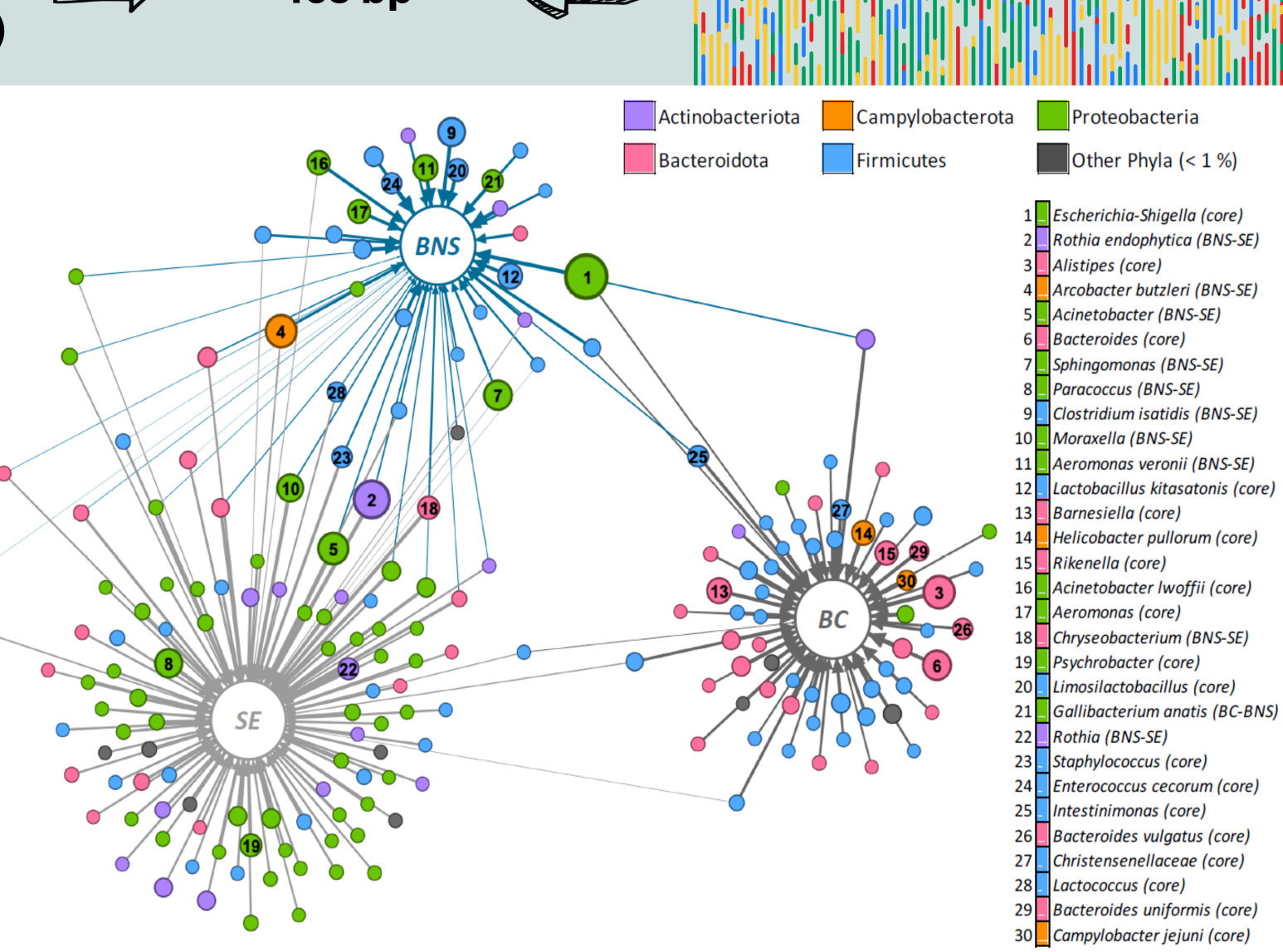
3. V3-V4 region library constructed

from the 16S rRNA gene region

RESULTS & DISCUSSIONS

The BC microbial community is significantly distinct from the BNS and SE community (Fig. 1, 2).





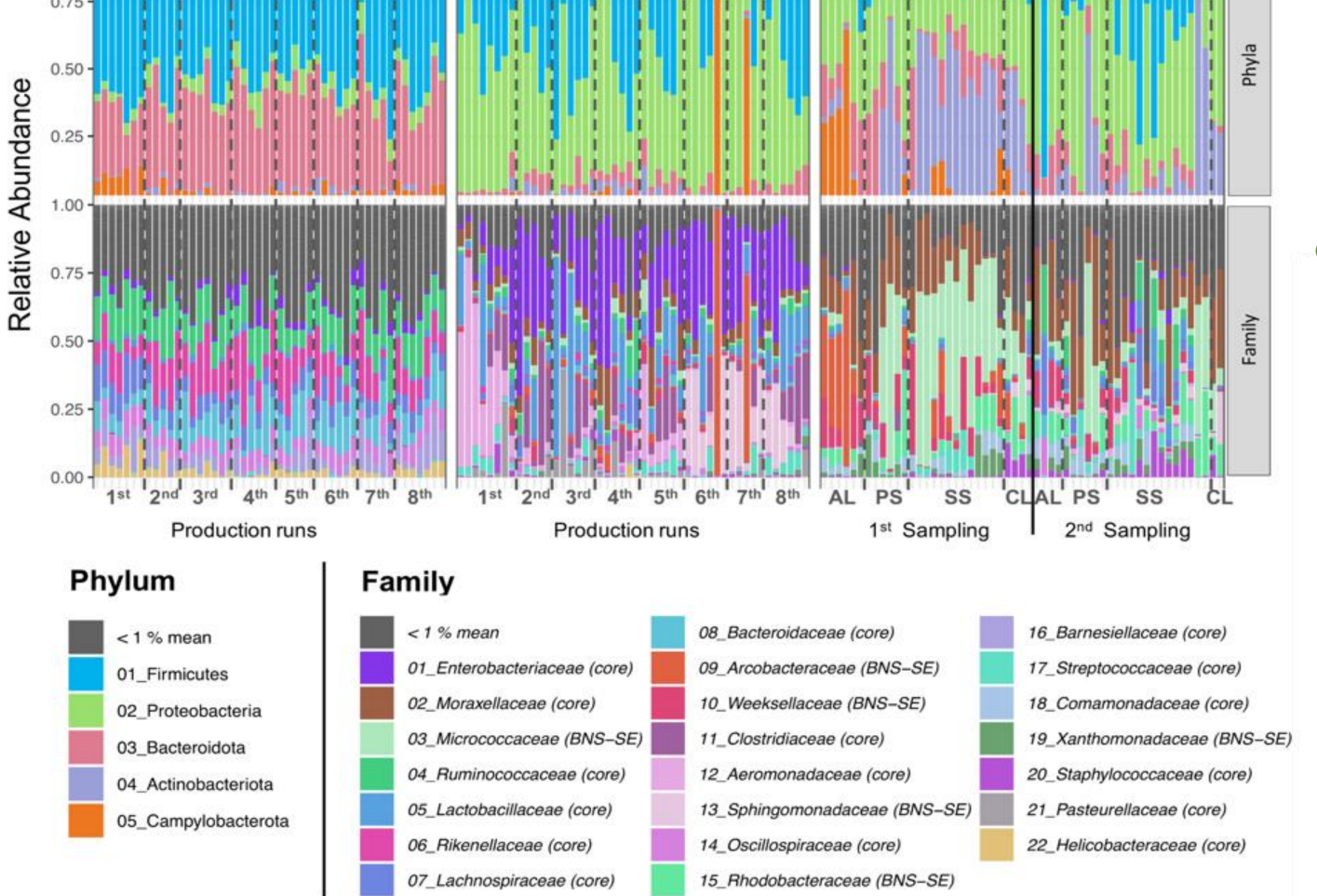


Figure 1. Graphic rapresentation of the network revealing the taxa associated to the three sampling sources. Taxa (colored nodes) are unidirectionally connected with arrows to the sampling sources if significant associations were detected (Indicator Species Analysis: *multipatt* statistics; R > 0.4and *P* value < 0.001).

Arcobacter butzleri emerged as one of the most abundant species and was detected throughout the slaughterhouse, showing a higher prevalence compared to other Campylobacterota (Fig. 1, 2). It was uniquely and significantly associated with BNS and SE, while *Helicobacter pullorum* and Campylobacter jejuni were indicators of BC (Fig. 1). A. butzleri was constantly

Figure 2. Microbiota composition and distribution in phylum and family taxa. The samples are grouped according to the temporal order of the sampling and according to the order of the slaughter of the flock and the processing stage, BC-

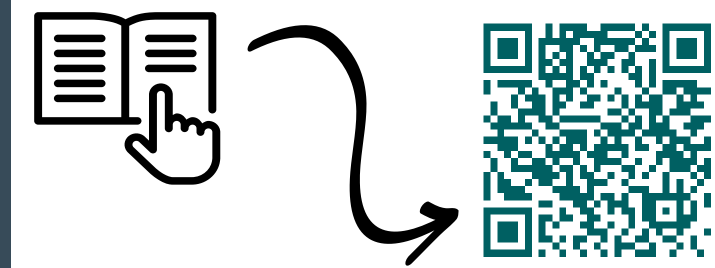
BNS and SE. Slaughterhouse sectors and transport lines are indicated: FL = first row; PS = plucking sector; SS = slaughter sector; SL = second row. Taxa are ordered in the legend from most to least abundant (mean >1%).

present on BNS along production runs, but significantly more abundant the first sampling day in SE (Fig.2).

CONCLUSIONS & IMPORTANCE

Our findings have emphasised the persistence of A. butzleri in a modern poultry abattoir and its establishment as part of the resident microbiota in specific environmental niches. The analysis conducted underlines the significance of early monitoring of food pathogens in the production chain, supported by meta-taxonomic analysis. Using these detection approaches, the presence of these pathogens could soon be considered an indicator of food safety and quality in slaughtered poultry.







Environmental Microbiology | Full-Length Text

Microbial contamination pathways in a poultry abattoir provided clues on the distribution and persistence of Arcobacter spp.

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