

Genomic characterisation of the population structure and antibiotic resistance of *Salmonella enterica* serovar Infantis in Chile, 2009–2022



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Summary

Background Multidrug-resistant (MDR) *Salmonella* Infantis has disseminated worldwide, mainly linked to the consumption of poultry products. Evidence shows dissemination of this pathogen in Chile; however, studies are primarily limited to phenotypic data or involve few isolates. As human cases of *Salmonella* Infantis infections have substantially increased in recent years, this study aimed to characterise the genomic epidemiology and antimicrobial-resistance profiles of isolates obtained from different sources, aiming to inform effective surveillance and control measures.

Methods We sequenced 396 *Salmonella* Infantis genomes and analysed them with all publicly available genomes of this pathogen from Chile (440 genomes in total), representing isolates from environmental, food, animal, and human sources obtained from 2009 to 2022. Based on bioinformatic and phenotypic methods, we assessed the population structure, dissemination among different niches, and antimicrobial resistance (AMR) profiles of *Salmonella* Infantis in the country.

Findings The genomic and phylogenetic analyses showed that *Salmonella* Infantis from Chile comprised several clusters of highly related isolates dominated by sequence type 32. The HC20_343 cluster grouped an important proportion of all isolates. This was the only cluster associated with pESI-like megaplasmids, and up to 12 acquired AMR genes/mutations predicted to result in an MDR phenotype. Accordingly, antimicrobial-susceptibility testing revealed a strong concordance between the AMR genetic determinants and their matching phenotypic expression, indicating that a significant proportion of HC20_343 isolates produce extended-spectrum β -lactamases and have intermediate fluoroquinolone resistance. HC20_343 *Salmonella* Infantis were spread among environmental, animal, food, and human niches, showing a close relationship between isolates from different years and sources, and a low intra-source genomic diversity.

Interpretation Our findings show a widespread dissemination of MDR *Salmonella* Infantis from the HC20_343 cluster in Chile. The high proportion of isolates with resistance to first-line antibiotics and the evidence of active transmission between the environment, animals, food, and humans highlight the urgency of improved surveillance and control measures in the country. As HC20_343 isolates predominate in the Americas, our results suggest a high prevalence of ESBL-producing *Salmonella* Infantis with intermediate fluoroquinolone resistance in the continent.

Funding Partially supported by the Food and Drug Administration (FDA) of the U.S. Department of Health and Human Services as part of an award, FDU001818, with 30% percent funded by FDA/HHS; and by Agencia de

The Lancet Regional Health - Americas 2024;32: 100711

Published Online xxx <https://doi.org/10.1016/j.lana.2024.100711>

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Investigación y Desarrollo de Chile (ANID) through FONDECYT de Postdoctorado Folio 3230796 and Folio 3210317, FONDECYT Regular Folio 1231082, and ANID—Millennium Science Initiative Program—ICN2021_044.

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Keywords: *Salmonella* Infantis; Americas; Antibiotic resistance; pESI; Megaplasmid; CTX-M-65; Fluoroquinolones; Chile

Research in context

Evidence before this study

In the last decade, emergent multidrug-resistant *Salmonella* Infantis has spread worldwide, primarily linked to poultry product consumption. However, in most countries from the Americas Region, such as Chile, the extent of the dissemination of emergent *Salmonella* Infantis and its molecular epidemiology remains unknown. In May and September 2023, an online search was conducted using the Google engine and the PMC database with the terms “*Salmonella*,” “Infantis,” and “Chile,” with no language restrictions. We assessed the results to select those presenting antimicrobial resistance, epidemiologic, or genomic data directly associated with isolates from Chile (13 studies). The selected studies showed that the prevalence of *Salmonella* Infantis in poultry-meat production systems, its resistance to different antibiotics, and the number of human cases of infection caused by this serovar have increased since 2014–2016. However, these reports were limited to phenotypic data or involved the genomic analysis of a few isolates (<50) obtained from the same source. No study has assessed the genomic epidemiology of the *Salmonella* Infantis population at the country level.

Added value of this study

Here, we present the first large-scale genomic epidemiology analysis of *Salmonella* Infantis in Chile, including isolates from environmental, food, animal, and human sources obtained from 2009 to 2022. We found that *Salmonella* Infantis in Chile is divided into several clusters of highly related isolates and that only a single cluster, the HC20_343, was associated with multiple antimicrobial-resistance determinants and pESI-like

megaplasms. We also report that isolates from this cluster are widespread among most sources, including irrigation water, poultry, food, and human cases. Detection of AMR determinants coupled with antimicrobial-susceptibility testing indicated that most HC20_343 isolates are ESBL-producers and have intermediate resistance to ciprofloxacin. Population structure analysis of this foodborne pathogen evidenced an active transmission of MDR *Salmonella* Infantis between different niches. This study reveals the widespread dissemination of MDR *Salmonella* Infantis in Chile.

Implications of all the available evidence

The evidence indicates that emerging *Salmonella* Infantis from the HC20_343 cluster is spreading among various niches, including irrigation water, poultry, and food, causing human infections in Chile. Its resistance to first-line antibiotics used for treating salmonellosis in individuals with a higher risk of severe or invasive infections is concerning. Currently, most surveillance and control efforts to reduce salmonellosis in Chile are focused on the poultry industry, and the study of outbreaks does not include whole-genome sequence analyses. Our findings highlight the urgent necessity to improve the surveillance and control measures to include agricultural waters to prevent contamination of produce and the further dissemination of resistance genes in the environment. As the HC20_343 cluster is highly prevalent in the Americas, further research involving large-scale genomic population analyses would shed light on the extent of the dissemination and transmission routes of emergent *Salmonella* Infantis in the continent and may contribute to informing surveillance and control policies.

Introduction

Non-typhoidal *Salmonella* (NTS) are one of the leading causes of foodborne disease globally, mainly affecting children under five and the elderly. In 2019, NTS caused 215,000 deaths and the loss of approximately 15 million years of healthy life worldwide, according to the estimates of the Global Burden of Disease Study.¹ The burden posed by NTS is in part fuelled by the emergence and spread of antimicrobial resistance, especially to critically-relevant antimicrobials such as third-generation cephalosporins (3 GC) and fluoroquinolones, which are the first-line treatment option for severe NTS infections.^{2,3}

In the past decade, multidrug-resistant (MDR) and extended-spectrum β -lactamase (ESBL)-producing *Salmonella enterica* serovar Infantis have emerged in different continents as a zoonotic pathogen causing outbreaks of foodborne illness associated with poultry products consumption.^{4–6} In some countries, this pathogen has displaced the historically most prevalent *Salmonella* serovars, such as Typhimurium and Enteritidis.^{7–10} The success of emerging MDR *Salmonella* Infantis is linked to the acquisition of a \approx 300 Kbp pESI-like megaplasmid, which encodes virulence, fitness-enhancing, and antibiotic-resistance factors that favour its capacity of biofilm production, adhesion, invasion, and resistance to 3 GC.^{4–6}

Moreover, isolates of this emerging pathogen are also associated with the chromosomal *gyrA* (D87Y) mutation, involved in fluoroquinolone resistance.^{5,11} The enhanced virulence and antimicrobial resistance traits of emergent *Salmonella* Infantis make this pathogen a global threat to public health.

Chile is a South American country organized into 16 administrative Regions, that concentrates its population and most agricultural activities in the central area.¹² Ranking third after Brazil and the United States, Chile is one of the leading exporters of poultry meat in the region.¹³ Recent research, primarily phenotypic or involving few isolates, has shown the spread of MDR *Salmonella* Infantis in the country, mainly linked to poultry. Analysis of the poultry and pig production systems indicated increased *Salmonella* Infantis prevalence and resistance to multiple antibiotics such as β -lactams, aminoglycosides, and tetracyclines.^{14,15} This was concurrent with a high proportion of reported ESBL-producing isolates in poultry products sold at Santiago de Chile's supermarkets.^{16–18} In 2019, the Instituto de Salud Pública de Chile (ISP; Public Health Institute of Chile) reported a 431% increase in intestinal and invasive human infections caused by *Salmonella* Infantis in 2018, compared to 2014,¹⁹ clearly documenting the farm-to-fork transmission of this pathogen. Furthermore, MDR strains of this pathogen were also isolated from irrigation water and a Magellanic Horned-Owl,^{20,21} suggesting its dissemination in the environment outside poultry sources. These data document the emergence and spread of this foodborne pathogen in Chile in recent years; however, its magnitude and molecular epidemiology remain unknown.

In this study, we conducted the first large-scale genomic analysis of the *Salmonella* Infantis population in the country, aiming to assess its population structure, antimicrobial resistance, and dissemination among different sources, in order to produce valuable information for the surveillance and control of this emergent pathogen and the spread of antibiotic resistance genes in food and the environment.

Methods

Study design, sample collection and isolation of *Salmonella* Infantis

A collaborative effort between different institutions studying *Salmonella* (see author list and affiliations) was assembled to rapidly gather isolates of *Salmonella* Infantis from different sources and collection years, and to obtain and access whole-genome sequencing (WGS) data. 396 isolates (from 2014 to 2022) were obtained from environmental (surface water: 264; poultry-farm environment: 41), poultry (28), food (chicken meat: 37), animal feed (10), animal (dog feces: 5), and clinical (human: 11) sources from three regions of central Chile:

Región de Valparaíso, Región Metropolitana, and Región del Maule (Fig. 1A and Supplementary Table S1).

Surface water samples (10 L) were collected at various points from five watersheds in central Chile (the Maipo, Mapocho, Claro, Lontué, and Mataquito rivers) using modified Moore swabs.²² Poultry-associated samples (boot swabs, chicken crops, and cecal content) were collected from poultry farms and chicken-meat production systems located in Región Metropolitana. These samples were processed according to a modified FDA-BAM protocol to isolate *Salmonella* as previously described.¹² Isolation of *Salmonella* was confirmed by PCR-amplification of the *invA* gene with primers *invAF* (5'-GAATCCTCAGTTTTCACGTTTC-3') and *invAR* (5'-TAGCCGTAA-CAACCAATACAAATG-3').²³ Raw meat-based dog diets and faecal samples (rectal swabs; Copan® Transystem™ 132C) from raw-fed dogs were processed according to the FDA-BAM protocol with modifications; raw food (25 g) was enriched in 225 mL lactose broth (BD Difco), and faecal samples were enriched in 10 mL buffered peptone water. Isolation of *Salmonella* was confirmed by PCR as described for the water isolates. Dog-samples collection was approved by the Institutional Committee for the Care and Use of Animals from the University of Chile under the code 20411-VET-UCH.

Raw and ready-to-eat poultry products were collected from supermarkets, restaurants, and meat-producer facilities by the Subsecretaría de Salud Pública from Valparaíso and transported at 0–4 °C to the Laboratorio de Salud Pública, Ambiental y Laboral (SEREMI Salud—Valparaíso). *Salmonella* isolation from these samples was performed according to ISO 6579-2017, and serotyping was performed at the ISP.

Human clinical samples (e.g., stool, blood, urine) representing human salmonellosis cases from different Región Metropolitana areas were received at the Laboratory of Microbiology of the UC-Christus Health network and processed for *Salmonella* isolation. Samples were inoculated in Hektoen Enteric agar and incubated at 35 ± 2 °C for 24–48 h in aerobiosis; then, *Salmonella* confirmation was performed using MALDI-TOF mass spectrometry, and serotyping was performed at the ISP. Permission to use the human clinical isolates was granted by the Comité Ético Científico de Ciencias de la Salud UC (Protocol ID 220614005).

Salmonella Infantis available from the above-described sources were all selected. All isolates were stored in 20% glycerol stocks and maintained at –80 °C.

Genome sequencing and construction of the genome dataset

Isolates from surface water were sequenced at the United States Food and Drug Administration Center for Food Safety and Applied Nutrition. Isolates from other sources were sequenced at the GenomeTrakr New York State Department of Health laboratory. Whole genome sequencing was performed on 385 isolates, and the

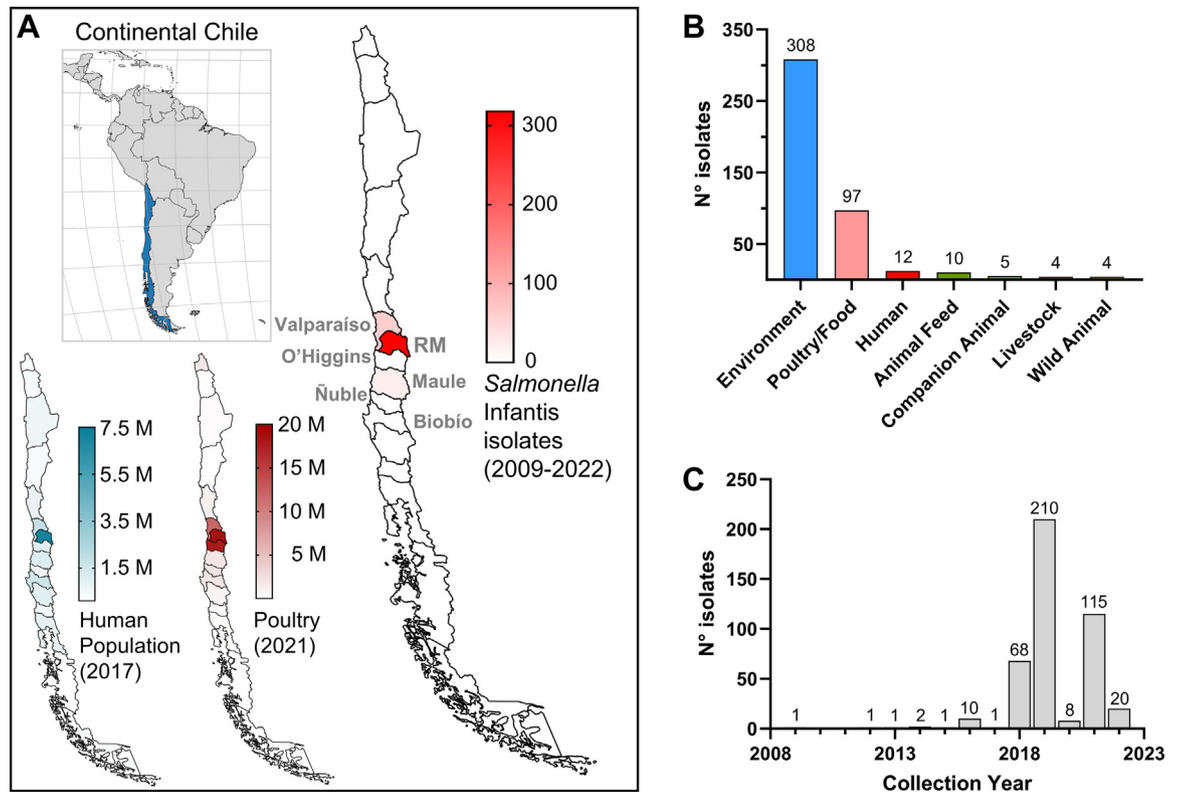


Fig. 1: Region of origin, isolation source, and collection year of the *Salmonella* *Infantis* isolates analysed in this study. A) Location of continental Chile in South America and distribution of its human population, poultry production, and collected *Salmonella* *Infantis* isolates among the 16 Regions (first-level administrative divisions). Note the concentration of these three variables in central Chile, especially in the Metropolitan Region. Population and poultry data were obtained from the corresponding last population and agricultural censuses carried out in 2017 and 2021, respectively (available at <http://resultados.censo2017.cl/> and <https://www.ine.gob.cl/censoagropecuario/resultados-finales/graficas-regionales>). The administrative Regions from which the *Salmonella* *Infantis* isolates were collected are indicated. RM: Región Metropolitana. B) *Salmonella* *Infantis* isolates distribution per isolation source and per C) collection year.

reads were deposited in the Sequence Read Archive, NCBI. Additionally, the *Salmonella* *Infantis* strains from human cases were sequenced at SeqCenter, Pittsburgh, PA, and the reads were directly uploaded to Enterobase.²⁴ All sequencing was conducted using Illumina platforms.

On April 20th, 2023, the Enterobase database for *Salmonella* was queried for genomes from Chile and the serovar *Infantis* as predicted by SISTR1²⁵ or SeqSero2.²⁶ A total of 440 *Salmonella* *Infantis* records were retrieved, including the 396 sequenced by us, along with their associated metadata (Table 1; Supplementary Table S1). Only two genomes lacked the information regarding the collection year and were not included for the analysis of the isolate distribution per collection year. To avoid/reduce methodological discrepancies in genome assemblies, all genomes were downloaded from Enterobase, where they were assembled from raw reads and annotated, with the use of a standardized bioinformatic pipeline.²⁴

Population structure and phylogenetic analyses

The 7-gene MLST, core genome MLST (cgMLST), hierarchical clustering based on cgMLST profiles, core SNP-based phylogeny, and minimum spanning trees were all carried out directly in Enterobase.²⁴ Annotated genomes were used for allele calling to classify the *Salmonella* *Infantis* isolates in sequence types (STs) by MLST (based on genes *aroC*, *dnaN*, *hemD*, *hisD*, *purE*, *sucA* and *thrA*) or core genome STs (cgSTs) by cgMLST (based on a 3002 allele scheme, cgMLST V2; https://pubmlst.org/bigsub?db=pubmlst_salmonella_seqdef&page=schemeInfo&scheme_id=4). HierCC V1²⁷ was used to cluster isolates based on the cgMLST profiles. Clusters grouping isolates with links no more than 20 alleles apart (HC20) were used to describe the *Salmonella* *Infantis* population. Based on cgMLST profiles, a minimum spanning tree (MST) was constructed with MSTree V2. A maximum likelihood phylogenetic tree based on core-SNPs was built with the *Salmonella* *Infantis* N55391 genome (Enterobase Barcode SAL_EA1888AA; GenBank

Feature	# genomes (Total = 440)
7-gene MLST	
ST32 (<i>thrA19</i>)	435 (98.9%)
ST9835 (<i>thrA1541</i>)	3 (0.7%)
ST9853 (<i>thrA1552</i>)	2 (0.5%)
cgMLST + HierCC	
HC20_343	322 (73.2%)
HC20_775	83 (18.9%)
HC20_2398	8 (1.8%)
Other HC20 clusters	27 (6.1%)
Isolation source	
Environment ^a	308 (70.0%)
Food/Poultry	97 (22.0%)
Human	12 (2.7%)
Other	23 (5.2%)
Isolation year	
2022	20 (4.5%)
2021	115 (26.1%)
2020	8 (1.8%)
2019	210 (47.7%)
2018	68 (15.5%)
2009–2017	17 (3.9%)
Unknown	2 (0.4%)

^aEnvironmental isolates were obtained from surface water (n = 265) and soil/dust (n = 43).

Table 1: Features of the publicly available *Salmonella* Infantis genomes from Chile (April 20th, 2023).

accession NZ_CP016410.1) as a reference. The phylogenetic tree was constructed with RAxML V8 based on 2008 variant sites called in $\geq 95\%$ of the genomes.

Identification of antibiotic-resistance genes/ mutations and presence of pESI-like megaplasmids

All 440 genome assemblies were downloaded from Enterobase and stored locally. Antibiotic resistance genes and point mutations involved in antimicrobial resistance in *Salmonella* were identified using AMRFinderPlus v3.11.4.²⁸ Since the *mdsA* and *mdsB* genes were found in all isolates and their presence did not result or explain any phenotypic resistance, these genes were not included in the analyses. The presence of pESI-like megaplasmids was assessed with ABRicate v1.0.1 (<https://github.com/tseemann/abricate>) and a custom database that included all 315 genes from the pESI-like megaplasmid pN55391 (GenBank accession NZ_CP016411.1).

Intra-source genomic diversity analysis

The cgMLST allelic profiles for each *Salmonella* Infantis isolate were downloaded from Enterobase. Pairwise allelic differences (PAD) were calculated for any pair of isolates within each source to assess the genomic diversity within each source. Only isolates representing unique cgSTs within a given source were included in the analysis. When more than one isolate per cgST was

detected, only one was randomly selected and included in the analysis.

Antibiotic-susceptibility testing

A sub-sample of 23 *Salmonella* Infantis isolates representing the diversity of the HC20_343 cluster was selected for antimicrobial susceptibility testing. These isolates were chosen because they represented different isolation sources and collection years and had the highest intra-source genomic diversity based on their PADs. Susceptibility testing was carried out in cation-adjusted Mueller-Hinton agar by the agar-dilution method following the recommendations of the Clinical and Laboratory Standards Institute.²⁹ The minimum inhibitory concentration (MIC) was interpreted according to the CLSI breakpoints available in the M100Ed33 document.³⁰ The following antibiotics, or antibiotic-inhibitor, were tested: amikacin (AMK), ampicillin (AMP), ampicillin-sulbactam (SAM), cefazoline (CFZ), cefepime (FEP), cefotaxime (CTX), cefotaxime/clavulanate (CTX/CLA), ceftazidime (CAZ), ceftazidime/clavulanate (CAZ/CLA), ciprofloxacin (CIP), fosfomycin (FOS), gentamycin (GEN), imipenem (IPM), meropenem (MEM), piperacillin-tazobactam (TZP), and trimethoprim/sulfamethoxazole (SXT). ESBL production was detected when the MIC of CTX and CAZ showed ≥ 3 2-fold reduction in the presence of clavulanate, an ESBL-inhibitor. A multidrug-resistant phenotype was assigned to isolates that displayed resistance to one or more antibiotics from at least three different classes.

These antibiotics were selected as they are routinely tested by the United States National Antimicrobial Resistance Monitoring System for Enteric Bacteria-NARMS³¹ (GEN, STX, MEM, AMP, CIP), allows the detection of ESBL production (CTX, CTX/CLA, CAZ, CAZ/CLA) or represent different members of included antibiotic classes (SAM, CFZ, FEP, IPM, TZP, AMK). FOS was tested as it is a first-line antibiotic used for uncomplicated urinary tract infections (UTIs) and there are reports of UTI caused by MDR *Salmonella* Infantis.³²

Statistical analyses and graphs

Independence between the presence of AMR genes/ mutations and the results of antibiotic-susceptibility was tested with the Fisher's exact test in GraphPad Prism v10.1.0, with $\alpha = 0.05$. Multiple correspondence analysis was carried out in DisplayR (<https://www.displayr.com/>) on metadata for all 440 genomes regarding 35 categories within 5 variables: Isolation source (8 categories), collection year (13 categories), 7-gene MLST (3 categories), HC20 cluster (9 categories), and megaplasmid presence (two categories).

Bar charts and violin plots were created in GraphPad Prism. MSTs were made with GrapeTree v1.5.1.³³ The iTOL v6 online tool was used to display and annotate the phylogenetic trees (<https://itol.embl.de/>). The multiple correspondence analysis plot was made in DisplayR.

Reporting guidelines

This work was written following the STrengthening the REporting of Genetic Association Studies (STREGA) reporting guidelines.³⁴

Role of the funding source

The funders of the study had no role in study design, data collection, data analysis, data interpretation, or writing of the report.

Results

Region of origin, isolation source, and collection year of the isolates

To study the structure of the *Salmonella* *Infantis* population from Chile, we sequenced 396 isolates of this pathogen. We analysed their genomes with all public *Salmonella* *Infantis* genomes from Chile available in Enterobase on April 20th, 2023. Our dataset was mainly composed of genomes from isolates coming from central Chile, specifically from Región de Valparaíso (n = 60), Región Metropolitana (n = 318), Región del Libertador General Bernardo O'Higgins (n = 1), Región del Maule (n = 21), Región Ñuble (n = 1), and Región del Biobío (n = 2). Most isolates (85.9%; 378/440) came from Región Metropolitana and Región de Valparaíso (Fig. 1A), and the data about the Region of origin for 37 isolates was not available. Genomes from environmental (river/creek/lagoon/irrigation water; boot swabs of soil/dust) and poultry/food (chicken carcass/crop/caecal content/faeces; chicken meat) origin made 92.0% (405/440) of total genomes, followed by human clinical cases which accounted for 2.7% (12/440 genomes) (Fig. 1B). While the collection year ranged from 2009 to 2022, most isolates (95.7%; 421/440) were distributed from 2018 to 2022 (Fig. 1C).

The *Salmonella* *Infantis* population in Chile belongs to ST32 and comprises different cgMLST clusters of highly related isolates

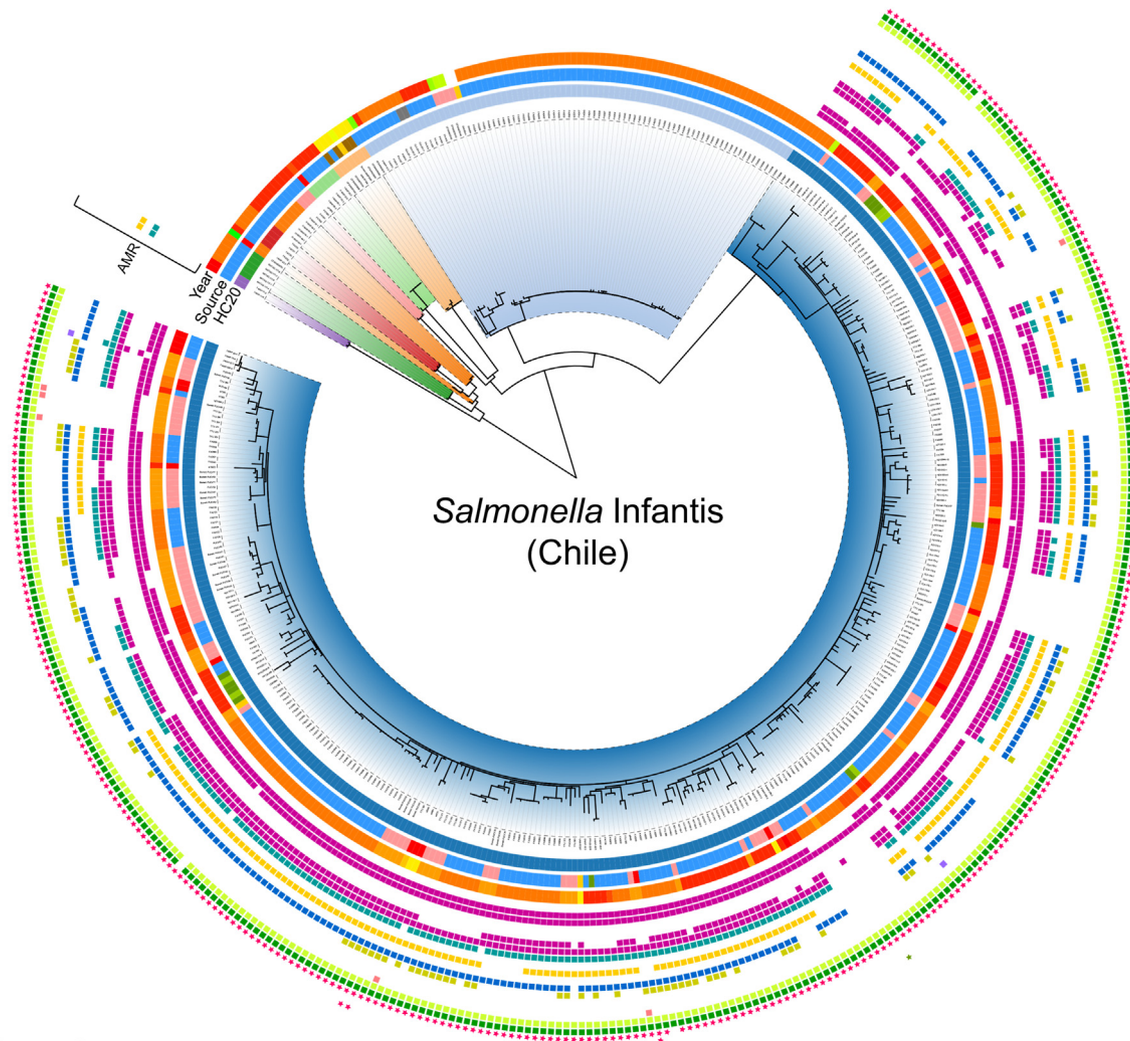
Among all isolates, 435/440 (98.9%) were ST32 (7-gene MLST), while the other five (three ST9835 and two ST9853) were single locus variants of ST32 that differed in the *thrA* allele (Table 1). The core SNP-based maximum likelihood phylogeny showed the presence of several clades comprising highly related genomes (Fig. 2). The hierarchical clustering of genomes linked by no more than 20 cgMLST-allele differences (HC20) revealed that the clades shown by the phylogeny corresponded with different HC20 groups, dominated by clusters HC20_343 and HC20_775 (405/440; 92.0% of genomes) (Table 1, Fig. 2). Isolates belonging to cluster HC20_343 came from the widest variety of isolation sources and some clusters (e.g., 2398, 422, 215390, 327110) seemed to group isolates only from environmental sources (Fig. 2). However, this is most likely an effect of a sample-size bias, with the most populated

cluster being more likely to harbour genomes from different sources. Notably, 10 out of 12 *Salmonella* *Infantis* isolates from human infections belonged to the HC20_343 cluster while the remaining two were found in clusters HC20_215390 and HC20_2398. These HC20_343 isolates were closely related to, and shared temporal proximity with, isolates from different sources, forming subclades in the phylogeny that included poultry (chicken), food (chicken meat), and environmental (surface water and soil/dust from boot swabs) isolates (Fig. 2).

Salmonella *Infantis* isolates from the HC20_343 cluster include MDR ESBL-producing strains and carry pESI-like megaplasמידs

The presence of AMR genes/mutations among the 440 genomes from Chile was assessed (Fig. 2; Supplementary Tables S2 and S3). Interestingly, the HC20_343 cluster (322 genomes) was the only one associated with up to 12 acquired antimicrobial resistance genes or mutations predicted to result in resistance to aminoglycosides, cepheids, folate pathway antagonists, chloramphenicol, fosfomycin, lincosamides, fluoroquinolones and tetracycline (Fig. 3A). The most frequent resistance genes among HC20_343 isolates were *tet(A)* (99.4%; 320/322), *sul1* (99.4%; 320/322), and *aadA1* (98.8%; 318/322), encoding predicted resistance to tetracycline, sulfonamide, and aminoglycosides, respectively. Genes *aph(3')-Ia*, *aph(4)-Ia*, *aac(3)-IVa* (aminoglycoside resistance), *bla_{CTX-M-65}* (3 GC resistance), *dfrA14* (trimethoprim resistance), and *floR* (phenicol resistance) were found in 60.6%–81.7% (195/322–263/322) of the isolates. Other identified resistance genes [*aadA12*, *aadA22*, *bla_{TEM-1}*, *lnu(A)*, *lnu(G)*, and *qnrB19*] were present in 6/322 or less isolates only, except by *fosA3* (fosfomycin resistance) that was carried by 28.9% (93/322) of isolates. Noteworthy, all but one HC20_343 isolate (99.7%; 321/322) carried the *gyrA* (D87Y) mutation involved in fluoroquinolone resistance.

Within the HC20_343 cluster, 210/322 genomes (65.2%) harboured nine or more AMR genes/mutations, and 321/322 genomes (99.7%) carried genes/mutations predicted to encode resistance to at least one antibiotic from three or more antibiotic classes, potentially resulting in an MDR phenotype (Fig. 3B). We found association (Fisher's exact test) between the content of genetic AMR determinants and the phenotypic resistance in all the isolates for which the MIC was assessed (Supplementary Table S4). Accordingly, all *bla_{CTX-M-65}*-positive isolates were resistant to the cephalosporins CFZ and CTX and displayed an ESBL-phenotype ($p < 0.0001$). Isolates harbouring *aadA1*, *aac(3)-Iva*, and *aph(4)-Ia* were resistant to GEN ($p < 0.0001$). Some isolates (9/23) were susceptible dose-dependent (SDD) to the fourth-generation cephalosporin FEP. However, this phenotype did not correlated with any of the identified AMR genes. The presence of *aadA1* alone was not



Legends (inside to outside rings)

HC20 (cgMLST)	Isolate source	Collection Year	AMR genes/mutations
HC20_343	Environment	2009	<i>aac(3)-IVa</i> ; <i>aadA1</i> ; <i>aadA12</i> ; <i>aadA22</i> ;
HC20_775	Poultry/food	2010	<i>aph(3')-Ia</i> ; <i>aph(4)-Ia</i>
HC20_10642	Human	2011	<i>bla_{CTX-M-65}</i> ; <i>bla_{TEM-1}</i>
HC20_326950	Animal feed	2012	<i>dfrA14</i> ; <i>dfrA8</i>
HC20_422	Companion animal	2014	<i>floR</i>
HC20_2398	Livestock	2015	<i>fosA3</i>
HC20_238389	Wild animal	2016	<i>Inu(A)</i> ; <i>Inu(G)</i>
HC20_215390		2018	<i>qnrB19</i>
HC20_327110		2019	<i>sul1</i>
		2020	<i>tet(A)</i>
		2021/2022	★ <i>gyrA(D87Y)</i> ; <i>gyrA(S83F)</i> ; <i>gyrB(S464Y)</i>
			☆ <i>ramR(T18P)</i>

Fig. 2: Phylogenetic analysis of Chilean *Salmonella Infantis* genomes. A core SNP-based maximum likelihood phylogeny (2008 variant sites; 95% presence) was constructed with 440 Chilean *Salmonella Infantis* genomes using *Salmonella Infantis* N55391 (Enterobase barcode SAL_E-A1888AA), a USA strain isolated from poultry in 2014, as the reference. Additionally, metadata regarding the HC20 clusters (clade colours and first ring), isolation source (second ring), isolation year (third ring), and presence of antibiotic-resistance genes/mutations as determined by AMRFinderPlus (coloured squares/stars) were incorporated into the phylogenetic tree.

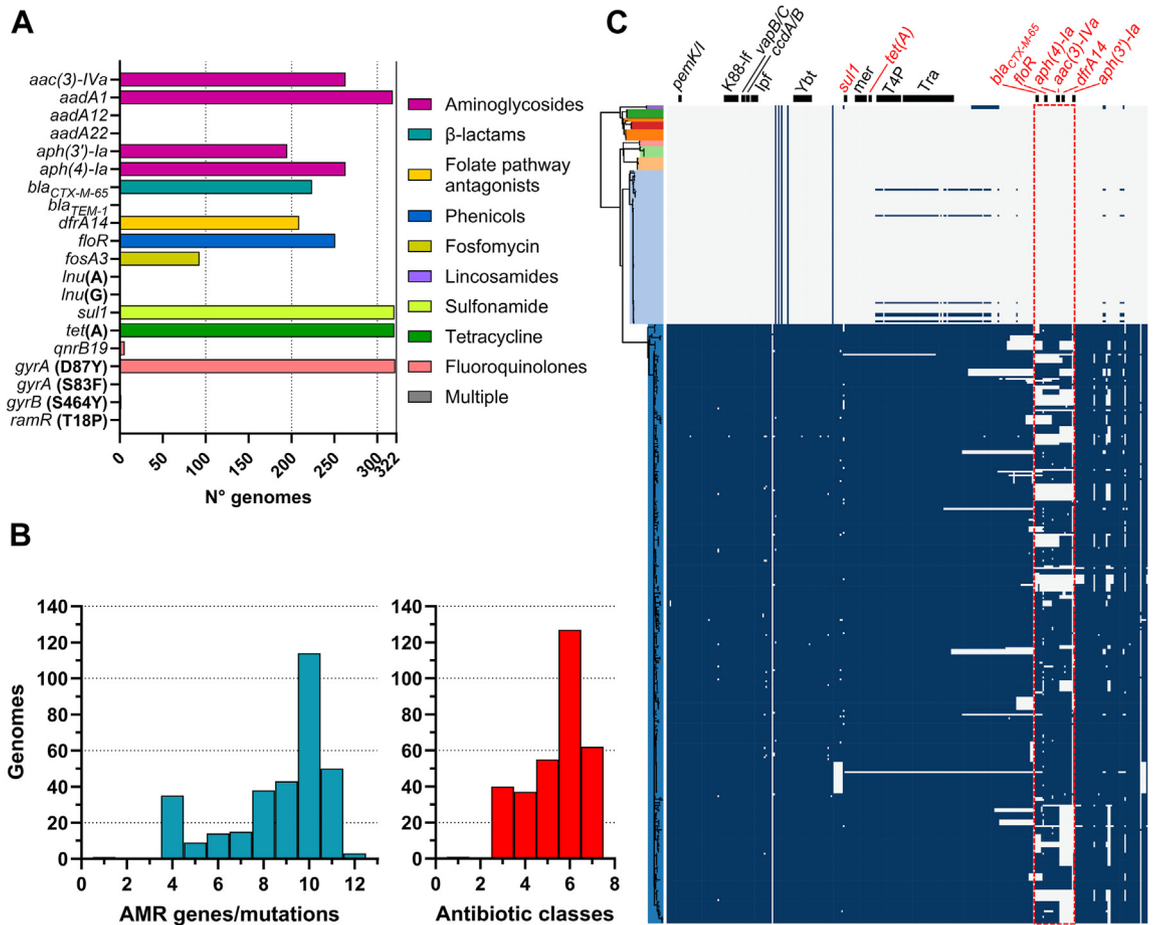


Fig. 3: Antibiotic-resistance determinants and pESI-like megaplasmid presence among *Salmonella Infantis* from the HC20_343 cluster. A) Frequency of individual antibiotic-resistant genes/mutations in the HC20_343 coloured by antibiotic class. B) Frequency of overall antibiotic-resistance genes/mutations per genome (blue bars) and antibiotic classes targeted per genome (red bars). C) Presence of pESI-like megaplasmids among the Chilean *Salmonella Infantis* genomes. A vertical representation of the same phylogenetic tree of Fig. 2 is coloured according to the HC20 clusters. The presence of each of the 315 genes from the pESI-like megaplasmid pN55391 was assessed by ABRicate using a custom database (blue squares). Black horizontal bars above the presence/absence matrix indicate the backbone regions of the megaplasmid (genes in black font) or the antibiotic-resistance regions (genes in red font). A red dashed line rectangle delimits the most variable region among the Chilean pESI-like megaplasmids relative to pN55391. K88-I_f (K88-like fimbria), Ip_f (Infantis plasmid-encoded fimbria, Ybt (Yersiniabactin synthesis cluster), mer (mercury-resistance cluster), T4P (type-IV pili encoding cluster), Tra (transfer region).

sufficient to confer GEN resistance, and *aph(3')-Ia* carriage did not show agreement with the resistance profile ($p_{AMK} > 0.9999$; $p_{GEN} = 0.115$). All tested isolates were susceptible to AMK. SXT-resistance was found in isolates carrying *sul1* plus *dfrA14* ($p < 0.0001$), and lack of *dfrA14* resulted in SXT susceptibility. All *gyrA* (D87Y)-positive isolates displayed intermediate resistance to CIP ($p < 0.048$). Notably, the presence of *qnrB19* plus *gyrA* (D87Y) resulted in CIP-resistance ($p < 0.046$). Conversely, all isolates lacking at least one of the genes/mutations mentioned above were susceptible to the corresponding antibiotics.

Since many of the identified antibiotic-resistance genes have been reported to be carried by the pESI-like

megaplasmid associated with emerging MDR *Salmonella Infantis*, we screened the entire genome dataset for the presence of the pN55391 pESI-like megaplasmid genes (Fig. 3C; Supplementary Table S5). We found that only the HC20_343 cluster harboured most of the pESI-like megaplasmid genes (from 164 to 310 out of 315 genes), including those encoding the three toxin-antitoxin systems (*pemK/I*, *vapB/C* and *ccdA/B*), the K88-like and Ip_f fimbria, the yersiniabactin synthesis cluster, the mercury resistance cluster, and the conjugative transfer region (*tra* and type-IV pili-encoding genes), which are part of the pESI-like megaplasmids backbone.³⁵ Most differences between the megaplasmids harboured by the Chilean strains and the pN55391

megaplasmid were located in the antibiotic resistance region, previously reported as a variable region.⁵ Nevertheless, the resistance genes contained in this region were present in most of the megaplasmid-harboured genomes (*bla*_{CTX-M-65}: 69.6%, 224/322; *floR*: 78.0%, 251/322; *aph(4)-Ia*: 81.7%, 263/322; *aac(3)-IVa*: 81.7%, 263/322; *dfrA14*: 64.9%, 209/322; and *aph(3')-Ia*: 60.6%, 195/322; [Supplementary Table S5](#)). Our analyses revealed that the HC20_343 *Salmonella* Infantis isolates acquired multiple antimicrobial-resistance genes/mutations, partly associated with the presence of pESI-like megaplasms.

HC20_343 *Salmonella* Infantis isolates are disseminated among different niches and show a low intra-source genomic diversity

An MST was constructed to visualize the genomic structure of the *Salmonella* Infantis population based on the cgMLST profiles ([Fig. 4A](#)). In agreement with the core SNP-phylogeny, the MST shows the bacterial population grouped in nine HC20 clusters of highly related isolates linked by no more than 20 allele-differences. All isolates from food, animal feed, and companion animals (dogs) belonged to cluster HC20_343. Accordingly, multiple correspondence analysis (MCA), carried out with metadata of all 440 genomes (variables: isolation source, collection year, 7-gene MLST, megaplasmid presence, and HC20 cluster), also evidenced the occurrence of HC20_343 isolates with majority of sources in recent years ([Supplementary File, Fig S1](#)). MCA clustered the HC20_343 category together with megaplasmid presence, environmental, companion animal, food, and human sources, and years from 2017 to 2022. Conversely, the remaining HC20 clusters grouped with megaplasmid absence, and years 2009–2015. Together, these findings highlight the foodborne, zoonotic, and human-pathogenic potential of HC20_343 *Salmonella* Infantis.

We carried out a more detailed analysis of cluster HC20_343 isolates. The MST evidenced putative events of transmission between different sources, as exemplified by subclusters 1 to 4 in [Fig. 4B](#). Subclusters 1, 3, and 4 included isolates from environmental, food, and human sources, while subcluster 2 included isolates from environmental, animal feed, and companion animals. Moreover, subclusters 1, 2, and 3 also included *Salmonella* Infantis from poultry. Importantly, isolates from all sources within these clusters were linked to isolates obtained from food (poultry products), and, ultimately, all sources within cluster HC20_343 had links with poultry. We assessed the intra-source genomic diversity of the HC20_343 isolates regarding the PAD between any pair of isolates representing unique cgSTs ([Fig. 4C](#); [Supplementary Table S6](#)). The environmental isolates displayed the highest diversity, with PAD values ranging from 0 to 51. All other sources harboured isolates with PADs ≤ 34 . The median PAD per source ranged from 13

in food isolates to 25 in animal feed. Environmental, poultry, food, and human isolates had the lowest median PADs (from 13 to 17), while the isolates from animal feed and companion animals had the highest (25 and 23, respectively). Overall, a low genomic diversity was found within the different isolation sources.

Discussion

The expansion of MDR *Salmonella* Infantis has been reported worldwide, with the highest proportion of isolates coming from the Americas region, followed by Europe.³⁶ The dissemination of this foodborne pathogen is mainly linked to poultry and poultry products, and different countries have reported a rise in human infections.^{8,15,36–38} Nevertheless, little is known about the extent of the *Salmonella* Infantis dissemination in different niches and its molecular epidemiology.

Here, we report the first large-scale genomic analysis of this foodborne pathogen population in Chile, finding evidence of the transmission of *Salmonella* Infantis carrying pESI-like megaplasms and multiple AMR determinants (cluster HC20_343) between environmental, poultry, food, animals, and human niches. Highly related isolates from different years were found in diverse sources, indicating a constant inter-source transmission. Currently, Chile is working to enhance the surveillance and control of *Salmonella* (including antimicrobial resistance) (SAG, Exempt Resolution 3687; <https://bcn.cl/2pap1>).³⁶ However, these efforts mainly focus on poultry and its derived products as they are known sources of *Salmonella*. Importantly, we report that irrigation waters are a source of *Salmonella* Infantis with MDR phenotypes. The presence of this pathogen in surface watersheds that supply the country's main agricultural region¹² underscores the urgent necessity to improve the current monitoring of irrigation waters and establish effective control measures to prevent the contamination of produce and the dissemination of antibiotic-resistance genes in the environment.

Human infection with non-typhoidal *Salmonella* usually results in self-limited acute gastroenteritis. However, children under five, adults over 65, and immunocompromised people are at higher risk of developing a severe life-threatening infection.³ Antibiotics, such as 3 GC and fluoroquinolones, are recommended to prevent or treat severe diseases.³ In 2017, the World Health Organization presented a list of 12 antibiotic-resistant bacterial pathogens, categorised into critical, high, and medium priority tiers, urgently requiring research and development of novel antibiotics since available treatments are becoming limited.³⁹ This list placed 3 GC-resistant *Enterobacteriaceae* and fluoroquinolone-resistant *Salmonella* spp. as critical and high-priority pathogens. A significant proportion of *Salmonella* Infantis from the HC20_343 cluster (69.6%, 224/322) harboured the ESBL-encoding gene *bla*_{CTX-M-65}

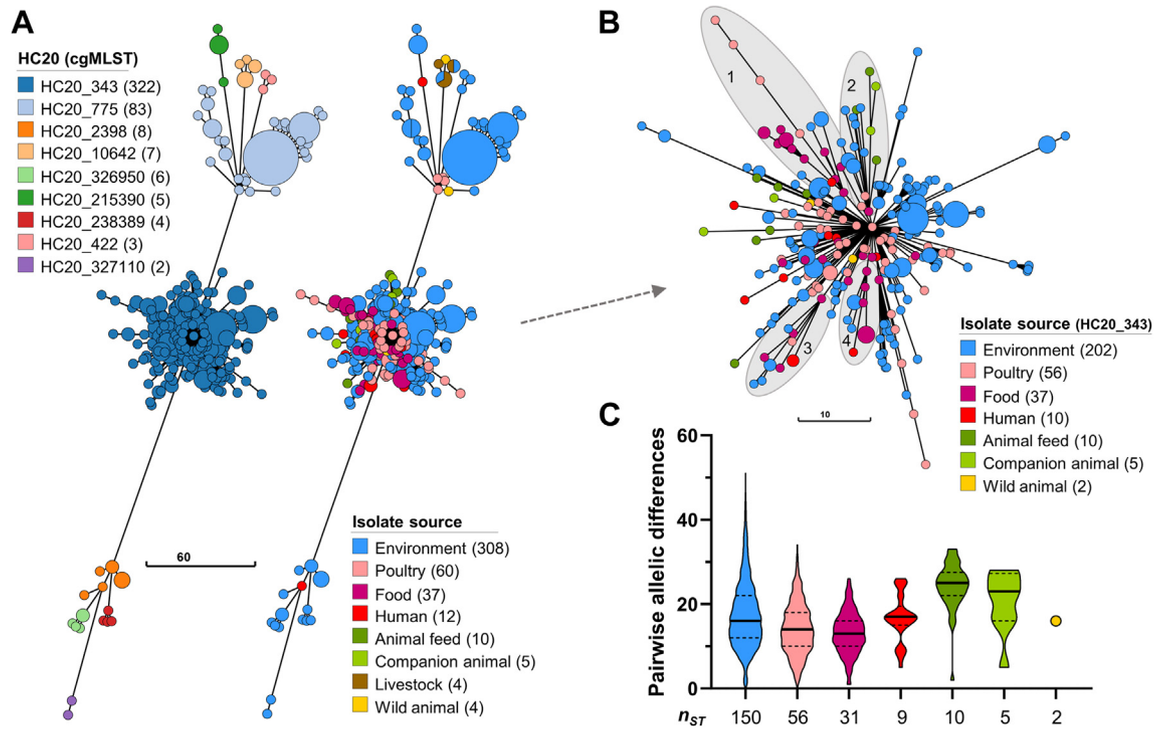


Fig. 4: Population structure of Chilean *Salmonella* *Infantis* and intra-source genomic diversity within cluster HC20_343. A) Minimum spanning tree depicting the population structure of *Salmonella* *Infantis* in Chile based on the cgMLST profiles of 3002 alleles. Nodes are coloured according to the HC20 clusters determined by HierCC or the isolation source, and their size is proportional to the number of isolates included in each node. The legends indicate the different HC20 clusters present, the isolation sources, and the number of isolates. B) Zoom-in view of the HC20_343 cluster structure with nodes coloured according to the isolation source. The legend also indicates the number of isolates per source within HC20_343. The shaded area and numbers indicate subclusters evidencing transmission of *Salmonella* *Infantis* between different sources. C) PAD between unique cgSTs from cluster HC20_343 per isolation source. Each violin plot, truncated at the highest and smallest values, represents the frequency distribution of PADs. The black unbroken and dashed lines represent the median PAD, and the 25th and 75th percentiles, respectively. The n_{ST} value indicates the number of isolates with unique cgSTs within each isolation source. Only isolates representing unique cgSTs were included in the analysis. Note: In panel 4B, the links and node sizes are the same as in panel 4A. However, panel 4B is presented at a different scale, and some branches were adjusted to prevent overlapping and improve subcluster visualization.

in a pESI-like megaplasmid. Moreover, almost all HC20_343 isolates (321/322) harboured the chromosomal *gyrA* (D87Y) mutation involved in fluoroquinolone resistance. Although the resistance profiles of emergent *Salmonella* *Infantis* reported in different countries are variable,^{8,37,40,41} partially as a result of the diversity within the megaplasmid AMR region (see Fig. 3C and ref.⁹), our findings in Chile are in agreement with a recent global survey of reported AMR determinants in *Salmonella* *Infantis*.³⁶ Importantly, we found that the aminoglycoside AMK and the carbapenems IPM and MEM were consistently active against all tested isolates from Chile. Our findings imply that the available options for preventing or treating severe infections in susceptible individuals are limited. The recently reported burden of AMR in the Americas for the year 2019⁴² showed that the resistance to fluoroquinolones and 3 GC accounted for most of the associated deaths to AMR. However, the number of

deaths caused by non-typhoidal *Salmonella* was among the lowest. As the burden report did not include *Salmonella* among the pathogens causing urinary tract infections, and the non-typhoidal *Salmonella*-3GC combination was also not considered, the real burden caused by this foodborne pathogen in the region could be higher, especially in the light of the widespread dissemination of MDR *Salmonella* *Infantis* in the continent (see below).

An unexpected finding was the association of the HC20_343 isolates with the presence of the pESI-like megaplasmids. Alba et al.⁶ found pESI-like megaplasmids carrying the *bla*_{CTX-M-1} gene in European *Salmonella* *Infantis* isolates from a higher diversity of HC20 clusters, and mainly from the HC20_7828 cluster. They also reported that megaplasmid-positive isolates from North America, or associated with traveling to South America, harboured the *bla*_{CTX-M-65} gene on the megaplasmid and belonged to the HC20_343 cluster. Similar

to our study, they found that strains from the HC20_775 cluster lacked the megaplasmid. In addition to the known association of *bla*_{CTX-M-65} with American and *bla*_{CTX-M-1} with European megaplasmid-carrying *Salmonella* Infantis,^{6,8} our findings suggest that the American megaplasmid-positive *Salmonella* Infantis strains might be associated with the HC20_343 cluster. Testing this hypothesis might help to better understand the global dissemination of emerging *Salmonella* Infantis. Importantly, we found that, out of 14706 *Salmonella* Infantis genomes from the Americas, 60.2% (8852/14706) belonged to the HC20_343 cluster (Supplementary File, Fig. S2), suggesting a high prevalence of megaplasmid-positive ESBL-producing isolates with intermediate fluoroquinolone resistance in the continent.

The approximate time for the arrival of the emerging *Salmonella* Infantis into Chilean territory remains unknown. The oldest pESI-like positive genomes in our dataset date from 2016 (4 genomes) and 2017 (1 genome) (Supplementary Table S1). These genomes represent four clinical isolates and one isolate obtained from a Dominican gull (*Larus dominicanus*), indicating that *Salmonella* Infantis carrying *bla*_{CTX-M-65}-positive pESI-like megaplasmids were circulating in Chile before 2016. A minimum spanning tree constructed with the cgMLST profiles from all *Salmonella* Infantis isolates from the Americas (available in Enterobase on July 7th, 2023) revealed that the HC20_343 isolates from the United States and Chile cluster together (Supplementary File, Fig.S2, also seen at NCBI Pathogen Detection). This finding suggests two possible scenarios in which emerging *Salmonella* Infantis from Chile came from the United States, or isolates from both countries share a common origin.⁴³

Our study is not free of limitations. Our genome dataset includes isolates from 6 out of 16 Regions in the country and is primarily concentrated in two Regions. Nevertheless, it is relevant to note that these Regions harbour most of the human population and poultry production in Chile (Fig. 1A). Therefore, the dataset analysed in this study offers a substantial representation of the bacterial population potentially encountered by most individuals and poultry raising/production systems in the country. We acknowledge, however, that a complete picture of the *Salmonella* Infantis population in Chile would require the analysis of isolates spanning the entire territory. Another limitation is that the genome dataset does not evenly represent different isolation sources and years. Isolates from companion animals, livestock, and wildlife, as well as isolates from samples collected from 2009 to 2017 are underrepresented. This limits our capacity to draw strong conclusions about the diversity of *Salmonella* Infantis within those sources. As our results show a widespread dissemination of this bacterium in irrigation waters, it would be of great relevance to obtain genomic data from isolates obtained from

possible sources of water contamination, as well as from produce irrigated with those waters. Importantly, the rise in human cases of MDR *Salmonella* Infantis reported by the ISP suggests the emergence of this pathogen in Chile around 2014–2016.¹⁹ However, the low number of genomes spanning the period before 2018 was a constraint in using the genomic data to answer questions about the time of emergence of the HC20_343 in the country. Future studies addressing questions about the emergence dynamics of MDR *Salmonella* Infantis in Chile should obtain and analyse genomic data from numerous isolates spanning the 2012–2017 period.

We analysed a dataset of 440 genomes representing the population structure of *Salmonella* Infantis in Chile. This population comprises strains lacking genetic determinants of antibiotic resistance and antibiotic-resistant strains that harbor pESI-like megaplasmids, both from the globally spread ST32. The megaplasmid-carrying strains belonged to the HC20_343 cluster, circulating among environmental, food, diverse animals, and human niches. Our results indicate that a significant proportion of the HC20_343 isolates encode ESBLs and display an intermediate resistance to fluoroquinolones, which limits the available treatments for individuals at a higher risk. Our findings and the reported increase in human cases highlight the urgent need to study the dissemination dynamics of this pathogen to devise effective surveillance and control measures.

Contributors

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All authors read and approved the submitted version of the manuscript and had access to the raw data. API and AIMS verified the data and had the final responsibility for the decision to submit the manuscript.

Data sharing statement

All genome metadata, identified AMR genes/mutations, antimicrobial susceptibility testing results, and identified megaplasmid genes are available in the Supplementary Material. Genome assemblies are publicly available in Enterobase (<https://enterobase.warwick.ac.uk/species/index/senterica>) and GenBank (<https://www.ncbi.nlm.nih.gov/genbank/>) using the corresponding accession numbers found in the Supplementary Table S1.

Editor note

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Declaration of interests

All authors declare no competing interests.

Acknowledgements

This study was partially supported by the Food and Drug Administration (FDA) of the U.S. Department of Health and Human Services (HHS) as part of an award, FDU001818, with 30% percent funded by FDA/HHS; and also by Agencia de Investigación y Desarrollo de Chile (ANID) through FONDECYT de Postdoctorado Folio 3230796 and Folio 3210317, FONDECYT Regular Folio 1231082, and ANID—Millennium Science Initiative Program—ICN2021_044.

Appendix A. Supplementary data

Supplementary data related to this article can be found at <https://doi.org/10.1016/j.j.lana.2024.100711>.

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