

RESEARCH ARTICLE

# Plasmids conferring resistance to extended-spectrum betalactamases including a rare IncN+IncR multireplicon carrying $bla_{\text{CTX-M-1}}$ in *Escherichia coli* recovered from migrating barnacle geese (*Branta leucopsis*) [version 1; peer review: 2 approved]

Paula Kurittu<sup>1</sup>, Banafsheh Khakipoor<sup>1</sup>, Michael S.M. Brouwer<sup>2</sup>, Annamari Heikinheimo<sup>1,3</sup>

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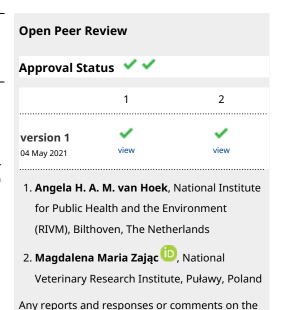
#### **Abstract**

**Background:** Increasing antimicrobial resistance (AMR) is a global threat and wild migratory birds may act as mediators of resistant bacteria across country borders. Our objective was to study extended-spectrum beta-lactamase (ESBL) and plasmid-encoded AmpC (pAmpC) producing *Escherichia coli* in barnacle geese using whole genome sequencing (WGS) and to identify plasmids harboring *bla* genes. **Methods:** Barnacle geese feces (n=200) were collected during fall 2017 and spring 2018 from an urban area in Helsinki, Finland.

ESBL/AmpC-producing *E. coli* were recovered from nine samples (4.5%) and isolates were subjected to WGS on both short- and long-read sequencers, enabling hybrid assembly and determination of the genomic location of *bla* genes.

**Results:** A rare multireplicon IncN+IncR was recovered from one isolate carrying  $bla_{\text{CTX-M-1}}$  in addition to aadA2b, lnu(F), and qnrS1. Moreover, rarely detected IncY plasmids in two isolates were found to harbor multiple resistance genes in addition to the human-associated  $bla_{\text{CTX-M-15}}$ . Poultry-associated  $bla_{\text{CMY-2}}$  was identified from the widely distributed IncI1 and IncK plasmids from four different isolates. One isolate harbored an IncI1 plasmid with  $bla_{\text{CTX-M-1}}$  and flor. A chromosomal point mutation in the AmpC promoter was identified in one of the isolates. WGS analysis showed isolates carried multiple resistance and virulence genes and harbored multiple different plasmid replicons in addition to bla-carrying plasmids.

**Conclusions:** Our findings suggest that wild migratory birds serve as



article can be found at the end of the article.

<sup>&</sup>lt;sup>1</sup>Department of Food Hygiene and Environmental Health, Faculty of Veterinary Medicine, University of Helsinki, Helsinki, Finland

<sup>&</sup>lt;sup>2</sup>Wageningen Bioveterinary Research, Lelystad, The Netherlands

<sup>&</sup>lt;sup>3</sup>Laboratory and Research Division, Microbiology Unit, Finnish Food Authority, Seinäjoki, Finland

a limited source of ESBL/AmpC-producing *E. coli* and may act as disseminators of the epidemic plasmid types IncI1 and IncK but also rarely detected plasmid types carrying multidrug resistance. Human and livestock-associated ESBL enzyme types were recovered from samples, suggesting a potential for interspecies transmission. WGS offers a thorough method for studying AMR from different sources and should be implemented more widely in the future for AMR surveillance and detection. Understanding plasmid epidemiology is vital for efforts to mitigate global AMR spread.

#### **Keywords**

Antimicrobial resistance, whole genome sequencing, extendedspectrum beta-lactamases, multidrug resistance, migratory birds, hybrid sequencing, One Health



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Corresponding author: Paula Kurittu (paula.kurittu@helsinki.fi)

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#### Introduction

As antimicrobial resistance (AMR) continues to increase unevenly worldwide (CDC, 2019; EFSA & ECDC, 2020; WHO, 2020), it is becoming increasingly urgent to study the transmission routes of resistant bacteria and mobile genetic elements harboring resistance genes. The drivers behind increasing AMR levels in different niches, including humans, animals, and the environment, have been studied (Holmes *et al.*, 2016), but occurrence and transmission routes of resistant bacteria need to be continuously monitored to enable early mitigation efforts.

Extended-spectrum beta-lactamase (ESBL) and plasmidencoded AmpC (pAmpC) producing bacteria, especially *Escherichia coli*, have been successfully spreading in both humans and animals (Ewers *et al.*, 2012). Moreover, increasing numbers of carbapenemase-producing *E. coli* are worrisome (Nordmann & Poirel, 2019). The successful spread of AMR is partly attributed to epidemic plasmids harboring resistance genes (Carattoli, 2011).

Regarding AMR in humans, studies have indicated human-to-human contact as the main route of transmission of ESBL-producing bacteria (Day *et al.*, 2019; Mughini-Gras *et al.*, 2019). However, resistant bacteria may also spread via animals, food, and the environment (EFSA, 2011; Mughini-Gras *et al.*, 2019) and their role should be continuously monitored as part of a One Health approach.

When studying bacteria resistant to antibiotics in different environments, it is important to assess the true threat any finding possesses to human or animal health. Whole genome sequencing (WGS) offers an efficient method for comparative epidemiological analysis. Different species and locations may have their unique set of bacterial sequence types (STs) and typical resistance genes but overlap between human- or animalassociated resistant bacteria and plasmids has been shown (Carattoli, 2009; Carattoli, 2011; Rozwandowicz et al., 2018). Combining short-read and long-read sequence technology provides a more accurate assembly of sequence data, which is especially important when determining the presence and structure of AMR-encoding plasmids. Plasmids occurring in multiple different regions, bacterial species, and sources can be considered epidemic (Carattoli, 2009). Certain traits, such as pilus formation and effective conjugation machinery in IncI1 and IncN type plasmids or plasmid addiction systems, may aid in the successful spread of plasmids (Carattoli, 2009; Carattoli, 2013).

One of the drivers of AMR has been identified to be international travel (Holmes *et al.*, 2016; Lääveri *et al.*, 2018; van der Bij & Pitout, 2012; Woerther *et al.*, 2017). As people and goods travel across country borders, so do wild animals. Migratory birds are able to travel across continents, potentially carrying resistant bacteria with them which they have picked up from anthropogenic waste sources (Bonnedahl & Järhult, 2014; Dolejska & Papagiannitsis, 2018). Wild birds have been shown to carry and spread ESBL-producing *E. coli* effectively between individuals in a flock (Sandegren *et al.*, 2018),

emphasizing the potential rapid spread of AMR. Waterfowl, especially barnacle geese (*Branta leucopsis*), have become a regular sight in many countries. Thousands of geese feed and defecate in densely human-populated areas, such as near housing and recreational parks (Elmberg *et al.*, 2017). Hundreds of thousands of barnacle geese migrate over Finland each year and feed on crop fields (SYKE, 2016). In addition to direct contact with bird feces, food-producing animals may come into contact with crops contaminated with fecal matter, as barnacle geese increasingly forage on maintained grasslands and pastures (Jensen *et al.*, 2018).

Barnacle geese are a protected species under the European Union Birds Directive (2009/147/EC) and listed under Annex I, which has enabled them to grow exponentially in number during recent decades, reaching a total population size of 1,390,000 in the 2010s (Fox & Leafloor, 2018). The populations are divided into three groups: east Greenland / Scotland & Ireland, Svalbard / southwest Scotland, and Russia / Germany & the Netherlands (Jensen et al., 2018). From the 1980s to 2010s, the Russia / Germany & the Netherlands population size increased by 30 times, with birds breeding now also in the Baltic and North Sea area (Jensen et al., 2018). Originally, barnacle geese only migrated through the Baltic Sea and bred in the Arctic zone, but since 1985 increasing numbers are also breeding in coastal areas of Finland and partly inland, especially the Turku archipelago region and the capital area (SYKE, 2019; Yrjölä et al., 2017). The geese included in this study were expected to belong to this latter group.

The increasing number of birds breeding in and migrating through new areas has resulted in birds being in closer contact with humans. Increased numbers of birds have also resulted in worries of contaminated recreational and drinking water, crop damage, and conflict in cities between humans and birds. Regarding AMR in birds, gulls in particular have been studied previously, and they have been found to harbor ESBLproducing bacteria frequently, although with varying prevalence between different countries (Stedt et al., 2015). The less studied geese and other waterfowl have also been shown to carry multiple pathogens (Elmberg et al., 2017), such as Campylobacter jejuni in barnacle geese in Finland (Llarena et al., 2015) and Yersinia spp. in barnacle geese in Sweden (Niskanen et al., 2003), but studies on the role of barnacle geese in AMR dissemination and potential transmission to humans or food-producing animals is limited. Barnacle geese breeding in Svalbard and wintering in the UK were found to carry intestinal E. coli with a high level of resistance against colistin (100%) but only a low level against ceftazidime (2%) (Hatha et al., 2013). Waterfowl in the Netherlands and Poland have been found to have different levels of resistance against amoxicillin, enrofloxacin, and tetracycline in one study (Kuczkowski et al., 2016), and the difference was speculated to be partly attributed to differences in the environment and the proximity to humans. The resistance patterns in wild birds can mirror those found in humans in different geographic locations (Stedt et al., 2015). As technologies have advanced, hybrid sequencing now offers a powerful tool for in-depth analysis of resistance genes along with plasmid replicons, virulence genes, and phylogeny between samples extracted from different sources. The goal of the present study was to study ESBL/AmpC-producing *E. coli* in barnacle geese residing in close proximity to humans, to identify resistance and virulence genes, STs, and plasmid replicons, and to conduct comparative phylogenetic analysis to identify potential epidemic plasmids.

#### Methods

#### Sampling

Fecal samples from barnacle geese (n=200) were collected during one day on two occasions, September 2017 (samples H1–H100) and May 2018 (samples H101–H200), in Helsinki, Finland. The sample location, a recreational park area, was the same on both occasions. During both sampling days, a flock of approximately 500 barnacle geese were present in the area. Samples were selected on the basis of collecting fresh droppings from the area, each sample being located approximately 0.5 - 2 meters from the previous sample, totaling at covering a sampling area of approximately 100 x 100 meters. Samples were aseptically collected into individual 1 1 plastic bags and transported to the laboratory for further analysis within 1 h.

#### Isolation and confirmation of Escherichia coli

From each sample,  $1.0 \pm 0.1$  g of feces was enriched in 9 ml sterile buffered peptone water (Oxoid, Basingstoke, UK) by incubating at  $37^{\circ}\text{C}$  overnight. Subsequently,  $10 \, \mu \text{l}$  of the pre-enrichment was streaked onto selective MacConkey agar plates (Oxoid, Basingstoke, UK) supplemented with 1 mg/l cefotaxime and incubated at  $44^{\circ}\text{C}$  for  $18-22 \, \text{h}$ . One colony from each plate with bacterial growth was re-streaked onto MacConkey agar plates with 1 mg/l cefotaxime supplement and incubated at  $37^{\circ}\text{C}$  for  $18-22 \, \text{h}$ . If a plate had morphologically different colonies, a representative colony from each different growth was streaked onto an agar plate.

After achieving a pure bacterial culture, the isolate was streaked onto a bovine blood agar plate and incubated at 37°C overnight for bacterial species determination with a matrix-assisted laser desorption ionization—time of flight mass spectrometry (MALDI-TOF MS) based Bruker Biotyper (Bruker Daltonics). A score value of 2.0–3.0 was considered high-confidence and was set as the criteria. All isolates identified as *E. coli* were stored at -70°C for further characterization.

#### Antimicrobial susceptibility testing

To confirm ESBL, AmpC, and/or carbapenemase production, antimicrobial susceptibility testing was performed on  $E.\ coli$  isolates with the disk diffusion method (EUCAST, 2017). Susceptibility to third-generation cephalosporins was tested with ceftazidime (10 µg) (Neo-Sensitabs, Rosco Diagnostica, Taastrup, Denmark) and cefotaxime (5 µg) (Oxoid, Basingstoke, UK), to fourth-generation cephalosporin with cefepime (30 µg), to cephamycin with cefoxitin (30 µg), and to carbapenem with meropenem (10 µg) (Neo-Sensitabs, Rosco

Diagnostica, Taastrup, Denmark). Epidemiological cutoff values were used as a reference (EUCAST, 2019). Synergism between third-generation cephalosporins and clavulanic acid was tested with a combination disk diffusion test with cefotaxime+clavulanic acid (30  $\mu g + 10~\mu g)$  and ceftazidime+clavulanic acid (30  $\mu g + 10~\mu g)$  (Neo-Sensitabs, Rosco Diagnostica, Taastrup, Denmark). E. coli ATCC 25922 was included as a quality control. In addition to resistance to third-generation cephalosporins, resistance to cephamycin and < 5~mm difference in inhibition zones in the combination disk diffusion test were used as criteria for AmpC production, whereas ESBL production was evidenced by resistance to third-generation cephalosporins and  $\geq 5~mm$  difference in the combination disk diffusion test.

#### DNA extraction and whole genome sequencing

**Short-read sequencing.** All ESBL/AmpC-producing *E. coli* isolates (n=9) were subjected to WGS with Illumina to study the presence of AMR and virulence genes and plasmid replicons, as well as to assess the multilocus ST.

Bacterial DNA was extracted and purified with the Pure-Link Genomic DNA Mini Kit (Cat# K182002, Invitrogen, Thermo Fischer Scientific, Carlsbad, CA, USA) according to the manufacturer's instructions. The assessment of DNA quality was carried out using a NanoDrop ND-1000 spectrophotometer (Thermo Fischer Scientific, Wilmington, DE, USA) and DNA quantity was measured using a Qubit 2.0 fluorometer (Invitrogen, Life Technologies, Carlsbad, CA, USA). Optical density OD260/280 of 1.8–2.0 and concentration of  $\geq$  50 ng/µl were set as thresholds. Library preparation was performed with Illumina Nextera XT and sequencing with Illumina Novaseq 6000 (Center for Genomics and Transcriptomics, Tübingen, Germany) with paired-end reads. Samples were sequenced with 100 × coverage, 2×100 bp read length, and a minimal phred quality score of 30.

Long-read sequencing. To study the complete sequences and to identify plasmid replicons carrying bla genes, all ESBL/AmpCproducing E. coli isolates (n=9) were additionally long-read sequenced. DNA extraction and purification were performed as described above. DNA extracts from all isolates were multiplexed in a random order with either SQK-LSK 108 or SQK-LSK 109 ligation sequence kit (Oxford Nanopore Technologies, Oxford, UK), depending on the availability of the respective flow cells, as described in more detail in the following sentences. DNA extracts from four geese isolates (H11, H21, H29, and H163) were multiplexed using the SQK-LSK108 ligation sequence kit (Oxford Nanopore Technologies, Oxford, UK) according to the manufacturer's protocol. Libraries were loaded onto FLO-MIN106D R9.4.1 MinION flow cells (Oxford Nanopore Technologies, Oxford, UK) used with the MinION Mk1B sequencing device and sequenced with MinKNOW software v19.06.8 for 48 h. For five isolates (H5, H58, H68B, H98, and H193) DNA extracts from three or two isolates at a time were multiplexed using the SQK-LSK109 ligation sequence kit (Oxford Nanopore Technologies, Oxford, UK)

according to the manufacturer's protocol. Libraries were loaded onto FLO-FLG001 R9.4.1 Flongle flow cells (Oxford Nanopore Technologies, Oxford, UK) used with the MinION Mk1B sequencing device and sequenced with MinKNOW software v19.06.8 for 20–24 h.

#### Bioinformatic analyses

Nanopore FAST5 read files were basecalled using Guppy v3.4.1 (Oxford Nanopore Technologies, Oxford, UK) with FASTQ output and demultiplexed with Qcat v1.1.0 (Oxford Nanopore Technologies, Oxford, UK). Quality trimming was performed with BBDuk (BBTools v38.71, Joint Genome Institute, USA) using a OTRIM value of seven. Hybrid assembly of Illumina and Nanopore sequences was performed with Unicycler v0.4.9b (Wick et al., 2017) set at default values. Bioinformatic analyses of bacterial DNA sequences were run on a web-based service (Center for Genomic Epidemiology, DTU, Denmark). Hybrid-assembled FASTA files were uploaded to ResFinder 4.1 (Zankari et al., 2012) to determine acquired beta-lactamase resistance genes with a 90% identity threshold and minimum length of 60%. In addition, chromosomal point mutations were searched for isolate H68B with the tool. Bacterial species identification was confirmed with KmerFinder 3.1 (Clausen et al., 2018; Hasman et al., 2014; Larsen et al., 2014). Bacterial multilocus sequence typing (MLST) was determined with MLST 2.0 (Larsen et al., 2012) using E. coli scheme 1 (Wirth et al., 2006). Virulence genes for E. coli isolates were determined with VirulenceFinder 2.0 (Joensen et al., 2014), using an identity threshold of 90% and a minimum length of 60%. PlasmidFinder 2.1 (Carattoli et al., 2014) was used to determine plasmid replicons located in the same contigs as beta-lactamase genes using an identity threshold of 95% and a minimum length of 60%. Plasmid STs were determined for beta-lactamase harboring plasmid replicons with pMLST 2.0 (Carattoli et al., 2014).

The plasmid sequences were annotated with Prokka v1.13 (Seemann, 2014) and manually curated with BLASTn/BLASTp. Plasmid structures were compared with previously published reference plasmids for each different replicon type identified and visualized in BRIG v0.95 (Alikhan et al., 2011). The structure of plasmid pZPK-H11 was visualized with SnapGene software (from Insightful Science, available at https:// www.snapgene.com). Visualization of the plasmid structure can also be achieved with BRIG v0.95 (Alikhan et al., 2011), for example. Studied plasmids were compared using Bac-Compare (Liu et al., 2019) with previously published plasmids found with a BLASTn search against the National Center for Biotechnology Information (NCBI) database and the 20 best matches with available metadata for each studied incompatibility type (for IncY plasmids only six similar, previously published plasmids with available metadata were found) were used to build a core genome MLST (cgMLST) based tree with 95% occurrence for discriminatory loci. The minimum spanning tree was visualized using GrapeTree v1.5 (Zhou et al., 2018). Information on included previously published plasmids with available metadata from NCBI GenBank are provided in Table 1.

#### **Ethics statement**

The study did not include any handling or disruption of animals, and therefore ethical approval of the research was not needed.

#### **Results**

#### Phenotypic identification of ESBL/AmpC-producing Escherichia coli

Out of 200 samples, 98 (49%) yielded bacterial growth on MacConkey agar plates supplemented with cefotaxime (1 mg/l). Of these, 55 (56%) samples were collected in fall 2017 and 43 (44%) in spring 2018. One colony was tested from each sample by using MALDI-TOF, revealing *E. coli* from nine (4.5%) samples. Seven (78%) of these originated from fecal samples collected on the first sampling period and two (22%) from the second sampling period. All *E. coli* samples originated from fresh, wet droppings.

Antimicrobial susceptibility testing. All nine isolates subjected to antimicrobial susceptibility testing were resistant to third-generation cephalosporins (cefotaxime and ceftazidime) (Table 3). According to the combination disk diffusion test, four isolates were phenotypically ESBL producers, four isolates AmpC producers, and one isolate both an ESBL and AmpC producer. Antimicrobial susceptibility testing results are shown in Table 3.

#### Whole genome sequencing and sequence analysis

WGS revealed seven different *E. coli* STs in the sequenced isolates (Table 4). Each sequenced isolate harbored from one to six different plasmid replicons. Three isolates carried an IncI1 plasmid replicon. Multidrug resistance was found in all of the sequenced isolates. Four of the isolates harbored genes conferring resistance to seven different antibiotic classes. In addition to beta-lactams, resistance genes were detected against aminoglycoside, fluoroquinolone, macrolide, lincosamide, streptogramin B, phenicol, sulfonamide, tetracycline, and trimethoprim. No fosfomycin or rifampicin resistance genes were detected.

Human-associated resistance gene  $bla_{\text{CTX-M-15}}$  was found in two isolates sharing the same sequence type ST453 and plasmid replicon IncY. Additionally, four and two isolates harbored  $bla_{\text{CMY-2}}$  and  $bla_{\text{CTX-M-1}}$ , respectively, which are commonly found from poultry (EFSA, 2011; Rozwandowicz et~al., 2018). Interestingly, no beta-lactamase genes were recovered from isolate H68B, although this isolate was phenotypically an AmpC producer. The phenotype correlated with the genotype in all but two isolates—the aforementioned isolate H68B and isolate H193—which was phenotypically both an AmpC and an ESBL producer, but only  $bla_{\text{CMY-2}}$  was recovered from the WGS analysis. In addition, isolate H21 was phenotypically an AmpC producer but harbored both  $bla_{\text{CMY-2}}$  and  $bla_{\text{TEM-32}}$ .

The isolates harbored a wide variety of different virulence genes with 38 different genes found altogether. The increased serum survival gene *iss*, tellurium ion resistance gene *terC*,

Table 1. Plasmids from GenBank database included for comparison with plasmids obtained in the current study.

Year of isolation	2010–2012	2013	2016	2013	2013	2008	2010-2012	2012-2016	2014	2014	2002-2005	2012	N/A	2002	2016	2009	2006	2010-2012	2000	2015	2011	2013	2011	2014
Source	Healthy broiler caecal sample	Healthy human	Poultry feces	Rat	Human blood sample (pyelonephritis)	Pig digestive tract	Healthy broiler caecal sample	Dog	Chicken meat	River	Human urinary tract sample	Australian silver gull	Human clinical sample	Dog urinary tract infection	Poultry feces	Pig	Pig digestive tract	Broiler (colibasillosis)	Human fecal sample	Cattle/pork meat	Human urinary tract infection	Poultry meat	Broiler fecal sample	Human urinary tract infection
Country	France	Switzerland	Norway	Guadeloupe / France	Guadeloupe / France	France	France	Switzerland	Denmark	France	France	Australia	Australia	Scotland	Norway	Z	France	France	Vietnam	Denmark	Switzerland	Switzerland	Japan	Switzerland
Accession number	MG648932.1	KJ484629.1	MN419437.1	CP053679.1	CP053677.1	MH847511.1	MG648914.1	MG948334.1	MK181561.1	MT230257.1	MG844436.1	MT468651.1	HG970648.1	CP023385.1	MN419430.1	KF787110.1	MH847502.1	MG649043.1	MG569891.1	MK181558.1	KR905389.1	KR905384.1	LC501565.1	KR905385.1
Bacterial species / ST	E. coli	E. coli / ST1638	E. coli / ST1638	E. coli / ST196	E. coli / ST349	E. coli	E. coli	E. coli	E. coli / ST602	E. coli	E. coli	E. coli / ST457	E. coli	E. coli	E. coli / ST57	E. coli	E. coli	E. coli	Shigella sonnei	E. coli / ST156	E. coli / ST1431	E. coli / ST1564	E. coli / ST648	E. coli / ST131
Inc group / pMLST	Ind1/ST3	IncI1/ST3	IncI1-Iy/ST3	Ind1/ST3	Ind1/ST3	Ind1/ST3	IncI1/ST3	Ind1/ST3	IncI1/ST3	Incl1/ST3	Incl1	IncI1/ST23	IncI1/ST2	IncI1/ST23	IncI1-Iy/ST3	Incl1/ST108	IncI1/ST3	Incl1/ST3	Incl1	IncI1/ST3	IncK2	IncK2	IncB/O/K/Z	IncK2
<i>bla</i> gene	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	CMY-136	CMY-2	CMY-2	CMY-2	CTX-M-1	CTX-M-1	CTX-M-1	CTX-M-1	TEM-1	CTX-M-1	CMY-2	CMY-2	CMY-2	CMY-2
Identity (%)	99.79	06.66	99.83	99.87	06.66	99.90	06.66	98.66	99.91	99.91	99.95	96.66	98.75	66.66	98.44	98.95	60.66	99.10	90.66	98.51	99.85	99.81	99.84	99.88
Query coverage (%)	96	96	95	95	92	94	94	94	92	92	100	100	100	97	93	92	92	92	06	92	94	94	94	92
Plasmid name (GenBank)	pCOV15	pH2291-112	p22638	pEC7	pEC38	p08-1118	pCOV12	p2305	p14011252	pESBL26	pCMY-136	pCE1628_I1	pJIE512b	p87	p17437	pIFM3804	p07-024	pCOV30	pDE105	p15076331	p4809.66	pDV45	p23C57-3	p5312.29
<i>bla</i> gene	CTX-M-1										CMY-2										CMY-2, TEM-32			
Inc group / pMLST	IncI1/ST38 CC-3										IncI1/ST23 CC-2										IncK			
Plasmid name (this study)	pZPK-H5										pZPK-H163, pZPK-H193										pZPK-H21			

Year of isolation	2014	2010–2012	2012	2013	2014	2012	2011	2005	2010	2012	2012	2004	2010	2006	2007	2012	2018	2016–2017
Source	Broiler meat (imported from Germany)	Healthy broiler caecal sample	Broiler meat	Poultry meat	Broiler meat (imported from Germany)	Poultry cloacae	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Broiler fecal sample	Poultry cloacae	Human clinical sample	Human blood culture
Country	Switzerland	France	Norway	Switzerland	Switzerland	Switzerland	Japan	Japan	Japan	Japan	Japan	Japan	Japan	Japan	Japan	Switzerland	Pakistan	Pakistan
Accession number	KY565558.1	MG648907.1	KU312044.1	KR905390.1	KY689635.1	KR905388.1	LC501559.1	LC501529.1	LC501547.1	LC501577.1	LC501570.1	LC501526.1	LC501544.1	LC501531.1	LC501535.1	KR905387.1	CP044008.1	LT906492.1
Bacterial species / ST	E. coli / ST38	Escherichia coli	Escherichia coli/ ST38	Escherichia coli/ ST38	Escherichia coli/ ST226	Escherichia coli/ ST420	Escherichia colil ST23	Escherichia coli/ ST373	<i>Escherichia colil</i> ST362	Escherichia colil ST155	Escherichia coli/ ST57	Escherichia coli! ST155	Escherichia coli/ ST10	Escherichia colil ST90	<i>Escherichia colil</i> ST354	Escherichia coli/ ST420	Salmonella Typhi	Salmonella Typhi
Inc group / pMLST	IncK2	IncB/O/K/Z	IncK2	IncK2	IncK2	IncK2	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncB/O/K/Z	IncK2	IncY	IncY
<i>bla</i> gene	TEM-1B (also mcr-1)	CMY-2	CMY-2	CMY-4	TEM-1B (also mcr-1)	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CMY-2	CTX-M-15, TEM-1B	CTX-M-15
Identity (%)	94.81	99.87	98.86	66.66	97.82	98.86	86.98	99.95	99.78	99.94	76.96	99.95	99.95	99.95	99.95	96.96	99.92	99.98
Query coverage (%)	06	∞ ∞	∞ ∞	∞ ∞	88	98	66	66	98	98	97	97	97	97	97	97	87	87
Plasmid name (GenBank)	pMbl488	6/00/d	pNVI1292	pDV10	pMbl536	pTMSA970	p23C16-2	p17C9-3	p22C48-3	p24C117-3	p24C25-2	p16C96-3	p22C25-2	p18C3-2	p19C79-2	pTMSA992	p2018K-0756	90009d
<i>bla</i> gene							CMY-2										CTX- M-15, TEM-1B	
Inc group / pMLST							IncK										IncY	
Plasmid name (this study)							pZPK-H58										pZPK-H29, pZPK-H98	

Year of isolation	d culture 2019	sample 2011	sample 2015																
	Human blood culture	a Human fecal sample	Human bile sample (cholangitis)		Pig fecal sample	Pig fecal sampl Human urine													pu pu pu
Accession Country number	430.1 Taiwan	153.1 Nigeria	423.1 France	70E 1									_						
/ ST number	yphi CP046430.1	oli KM023153.1	oli MF510423.1	VXXAX2095 1				Jilo Oli	Jilo Jilo		ος,	co.	co co	0,000	<u> </u>	20 00 14 14 14 14 14 14 14 14 14 14 14 14 14	33 45	23 25 26 27 27 27 27 27 27 27 27 27 27 27 27 27	20 00 7
/st	Salmonella Typhi	Escherichia coli	Escherichia coli	0 mm   mm   mm   mm   mm   mm   mm   mm	Shigella flexneri	Shigella flexneri Escherichia coli/ ST167	Snigella flexneri Escherichia coli, ST167 Escherichia coli	Snigella flexne Escherichia co ST167 Escherichia co Klebsiella pneumoniae/ ST2748	Sngella flexneri Escherichia colli ST167 Escherichia colli Mebsiella pneumoniae/ ST2748 Escherichia colli	Escherichia coli/ ST167 Escherichia coli/ Klebsiella pneumoniae/ ST2748 Escherichia coli/ Escherichia coli/ ST295	Snigella flexneri Escherichia colif ST167 Escherichia coli Klebsiella pneumoniael ST2748 Escherichia colif Escherichia colif ST295 Salmonella enterica serovar London	Shigella flexneri Escherichia coli/ ST167 Escherichia coli/ Klebsiella pneumoniae/ ST2748 Escherichia coli/ ST295 Salmonella enterica serovar Wirchow	Escherichia co ST167 Escherichia co ST2748 Escherichia co Escherichia co Escherichia co ST295 Salmonella en serovar Londa Salmonella en serovar Wirch Uncultured bacteria	Snigelia flexneri Escherichia coli ST167 Escherichia coli ST2748 Escherichia coli Fscherichia coli Salmonella enteri Scovar London Salmonella enteri serovar Wirchow Serovar Wirchow Serovar Wirchow Serovar Serovar S	Singelia flexineri Escherichia colii ST167 Escherichia colii Klebsiella pneumoniae/ ST2748 Escherichia colii ST295 Salmonella ente, serovar London Salmonella ente, serovar Wirchou Uncultured bacteria Escherichia coliii ST635 Escherichia coliii ST635	Singelia flexineri Escherichia coli ST167 Escherichia coli S12748 Escherichia coli S1295 Salmonella enterica serovar London Salmonella enterica serovar Wirchow Uncultured bacteria Escherichia colii ST635 Escherichia colii ST635 Escherichia colii ST635	Singelia flexineri Escherichia colli ST167 Escherichia colli ST2748 Escherichia colli ST295 Salmonella entei serovar London Salmonella entei serovar Wirchov Uncultured bacteria Escherichia colli ST635 Escherichia colli ST131 Klebsiella pneumoniae/ST8	Escherichia coli ST167 Escherichia coli ST2748 Escherichia coli ST2748 Escherichia coli ST295 Salmonella ente serovar Londor Salmonella ente serovar Wircho Uncultured bacteria Escherichia coli ST131 Klebsiella pneumoniae/ST	Escherichia colii ST167 Escherichia colii ST167 Escherichia colii Escherichia colii Escherichia colii ST295 Salmonella ente serovar Wircho Salmonella ente serovar Wircho Salmonella colii ST635 Escherichia colii ST635 Escherichia colii ST131 Klebsiella pneumoniae/ST ST131 Escherichia colii ST131 Escherichia colii ST131 Escherichia colii Escherichia colii
pMLST	i, IncY	i, IncY	i, IncY		IncY	IncY IncF-IncN													
and general	CTX-M-15, TEM-1B	CTX-M-15, TEM-1B	CTX-M-15, TEM-1B	TFM-1B	1	NDM-5	NDM-5 CTX-M-1, TEM-1	NDM-5 CTX-M-1, TEM-1 CTX-M-1	CTX-M-1, CTX-M-1	NDM-5 CTX-M-1, TEM-1 CTX-M-1 CTX-M-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 (also qnrS1)	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 (also qnr51) TEM-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 TEM-1 TEM-1 (also qnrS1) TEM-1	CTX-M-1, TEM-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 EM-1B TEM-1B TEM-1B CTX-M-65, TEM-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 TEM-1B TEM-1B CTX-M-65, TEM-1 TEM-1 TEM-1 TEM-1 KPC-4, TEM-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 TEM-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 EM-1 EM-1 EM-1 EM-1 EM-1 EM-1 EM-1	CTX-M-1, TEM-1 CTX-M-1 CTX-M-1 CTX-M-1 CTX-M-1 TEM-1 TEM-1 TEM-1 TEM-1 CTX-M-65 TEM-1 (also qnr51) CTX-M-65 TEM-1 (also qnr51) (arm 4)
Query Identity coverage (%) (%)	99.92	86.98	06.66	99.88	0	76.99	79.99 79.99 79.99	99.97	99.96	99.97	99.96	99.97	99.97	99.97	99.97	99.97	99.97	99.97	99.97
	98	71	70	61	9 - 2		59												
name (GenBank)	pR19.2839_ 83k	pPGR46	pEco-CTX- M-15	pRC960-1	p100_NDM5_	TICIN	pH1038-142	pH1038-142	pH1038-142 pC5_41608	pH1038-142 pC5_41608 pHHA45	pH1038-142 pC5_41608 pHHA45 pL2-43	pH1038-142 pC5_41608 pHA455 pL2-43 pYUHAP5-2	pH1038-142 pC5_41608 pL2-43 pYUHAP5-2 pVQS1	pH1038-142 pH4445 pHA45 pL2-43 pVQS1 pVQS1 pABWA45_3	pH1038-142 pC5_41608 pL2-43 pVQS1 pVQS1 pRSB203 pRSB203 pKC396	pH1038-142 pC5_41608 pL2-43 pVQ51 pVQ51 pRSB203 pABWA45_3 pABWA45_3	pH1038-142 pH1445 pH445 pL2-43 pVQS1 pVQS1 pABWA45_3 pKC396 pKC396	pH1038-142 pH1445 pHH445 pL2-43 pVQ51 pVQ51 pKC396 pKC396 pKC394 pKT58A	pH1038-142 pH1038-142 pC5_41608 pL2-43 pVQS1 pVQS1 pABWA45_3 pKC396 pKC396 pKC394 pKC384
gene gene					/ CTX-M-1														
Inc group /					IncN+IncR/ ST1(IncN)														
Plasmid name (this study)					pZPK-H11														

asmid ame (this udy)	Plasmid Inc group / bla name (this pMLST gene study)	<i>bla</i> gene	Plasmid name (GenBank)	Query coverage (%)		Identity bla gene (%)	Inc group / pMLST	Bacterial species / ST	Accession number	Country	Source	Year of isolation
			pRSB206	48	66.66	TEM-1	IncN	Uncultured bacteria	JN102344.1	Germany	Wastewater treatment N/A plant effluent	A/N
			pRSB205	47	66.66	TEM-1	IncN	Uncultured bacteria	JN102343.1	Germany	Wastewater treatment N/A plant effluent	A/N
			plasmid IncN 48	48	99.78	TEM-1A	IncN	Klebsiella pneumoniae/ST258	CP027050.1	Greece	Human clinical sample 2012–2014 (stool)	2012–2014
			pRSB201	50	99.70	TEM-1	IncN	Uncultured bacteria	JN102341.1	Germany	Wastewater treatment plant effluent	N/A
			pQNR2078	53	99.61	(qnrB19)	IncN	Escherichia coli	HE613857.1	Germany	Horse clinical sample (genital tract infection)	2005

Inc group = incompatibility group; pMLST = plasmid multilocus sequence type; ST = sequence type; E coli = Escherichia coli.

**Isolate** Sample secondary Illumina run **Oxford Nanopore** Plasmid sequence Sample primary name accession accession accession run accession accession SAMEA7856498 ERR5188293 ERR5190298 H5 ERS5602973 ERZ1738234 H11 ERS5602974 SAMEA7856499 ERR5188294 ERR5190299 ERZ1738235 H21 ERS5602975 SAMEA7856500 ERR5188295 ERR5190300 ERZ1738236 H29 ERS5602976 SAMEA7856501 ERR5188296 ERR5190301 ERZ1738237 H58 ERS5602977 SAMEA7856502 ERR5188297 ERR5190302 ERZ1738238 H68B ERS5602978 SAMEA7856503 ERR5188298 ERR5208198 No ESBL-plasmid

ERR5188299

ERR5188300

ERR5188301

ERR5190303

ERR5190304

ERR5190305

SAMEA7856504

SAMEA7856505

SAMEA7856506

Table 2. Accession numbers for isolates deposited to European Nucleotide Archive project number PRIEB42655.

and glutamate decarboxylase gene *gad* were found in all isolates, and the long polar fimbriae gene *lpfA* in seven isolates. Enterobactin siderophore receptor gene *iroN* and temperature sensitive hemagglutinin gene *tsh* were both found in three isolates. Two of the isolates harbored the adhesin intimin coding *eae* gene, which is associated with enteropathogenic *E. coli* (Frankel *et al.*, 1998; Müller *et al.*, 2016), but no Shiga toxin coding genes were found.

ERS5602979

ERS5602980

ERS5602981

H98

H163

H193

**Plasmid analysis.** Eight out of the nine hybrid sequenced *E. coli* isolates were found to harbor a plasmid replicon with a *bla* gene. All hybrid-assembled *bla*-carrying plasmids were found to be in one circular contig.

One isolate, H68B, did not carry any plasmids with bla genes, but instead a chromosomal point mutation in the AmpC promoter was identified. This isolate did, however, harbor multiple IncF type replicons and resistance genes mdf(A) and tet(A) (Table 4).

#### IncI1 type plasmids

In three isolates, an IncI1 type replicon harboring either  $bla_{\rm CMY-2}$  (H163 and H193) or  $bla_{\rm CTX-M-1}$  (H5) was identified (plasmids pZPK-H5, pZPK-H163, and pZPK-H193). Plasmid pZPK-H5 from isolate H5 was 98.7 kb in size with a G+C content of 50.2% and 112 predicted coding sequences (CDSs). Plasmid multilocus sequence typing (pMLST) analysis indicated that the plasmid belonged to ST38 CC-3. In addition to  $bla_{\rm CTX-M-1}$ , pZPK-H5 harbored the florfenicol resistance gene floR. The only virulence gene identified on this plasmid was the channel-forming colicin cia.

pZPK-H163 from isolate H163 and pZPK-H193 from isolate H193 were found to be identical with pairwise alignment with BLASTn. Both plasmids were 89.6 kb in size with a G+C content of 50.31% and predicted 101 CDSs. pMLST analysis

identified the plasmids as ST23 CC-2. In addition to  $bla_{\text{CMY-2}}$ , the plasmids carried the macrolide-associated resistance gene mdf(A). Similar to pZPK-H5, the only virulence gene identified on these plasmids was cia.

ERZ1738239

ERZ1738240

ERZ1738241

Pairwise alignment with BLASTn against IncI1 type reference plasmid R64 (GenBank accession: AP005147) demonstrated high similarity between R64 and pZPK-H163 and pZPK-H193, with 89% coverage and 97.57% identity. BLASTn alignment between R64 and pZPK-H5 indicated less similarity, with 79% coverage and 98.52% identity. Alignments of the studied plasmids with the reference R64 are visualized in Figure 1A for pZPK-H5 and Figure 1B for PZPK-H163 and pZPK-H193.

The studied IncI1 plasmids demonstrated typical IncI plasmid backbones with maintenance and stability-related regions with parAB, impCAB, ssb, psiAB, ardA, and pndAC genes, and transfer-associated, shufflon, and pilus formation regions (Figure 1A–B). In pZPK-H5, flor and cia genes were located in the accessory module, whereas  $bla_{\text{CTX-M-1}}$  was integrated into the shufflon region. As previously described,  $bla_{\text{CTX-M-1}}$  was integrated into shufflon segment B by a copy of ISEcp1, although in pZPK-H5 this IS element contains an integration of ISKpn26 (Brouwer et al., 2015). In pZPK-H163 and pZPK-H193,  $bla_{\text{CMY-2}}$  was located in the accessory module after the replication region.

Comparison of the studied IncI1 plasmids with the most closely related previously published plasmids from GenBank using BLASTn demonstrated that pZPK-H163 and pZPK-H193 were similar with a  $bla_{\text{CMY-136}}$ -harboring IncI1 plasmid obtained from a human urinary tract sample in France with 100% coverage and 99.95% identity (GenBank accession: MG844436.1), a  $bla_{\text{CMY-2}}$ -harboring IncI1 ST23 plasmid from an Australian silver gull with 100% coverage and 99.96% identity (GenBank

Table 3. Antimicrobial susceptibility testing<sup>a</sup> for presumptive extended-spectrum beta-lactamase (ESBL)/AmpC-producing *Escherichia coli* from barnacle geese.

Phenotype <sup>b</sup>	ESBL	ESBL	AmpC	ESBL	AmpC	AmpC	ESBL	AmpC	ESBL + AmpC
Meropenem Cefoxitin Cefepime Cefotaxime+clavulanic acid (10 μg) (30 μg) (30 μg) (30 μg + 10 μg), difference in zone diameter (mm)	7	7	4	12	3	2	12	4	5
Cefotaxime+clavulanic acid (30 µg + 10 µg), difference in zone diameter (mm)	18	20		21	3	_	20	8	0
Cefepime (30 µg)	~	~	S	~	S	S	~	~	~
Cefoxitin (30 µg)	S	S	~	S	~	~	S	~	~
Meropenem (10 µg)	S	S	S	S	S	S	S	S	S
	~	~	2	~	~	~	~	~	22
Isolate Cefotaxime Ceftazidime (5 μg) <sup>a</sup> (10 μg)	Rc	~	~	~	~	~	~	~	22
Isolate	H5	H11	H21	H29	H58	H68B	H98	H163	H193

<sup>a</sup> Antimicrobial susceptibility testing with disk diffusion method according to EUCAST (2017); EUCAST (2019).

b Criteria for ESBL production: resistance to third-generation cephalosporins and ≥ 5 mm difference in combination disk diffusion test; criteria for AmpC production: resistance to third-generation cephalosporins and cephamycin and < 5 mm difference in inhibition zones in combination disk diffusion test.

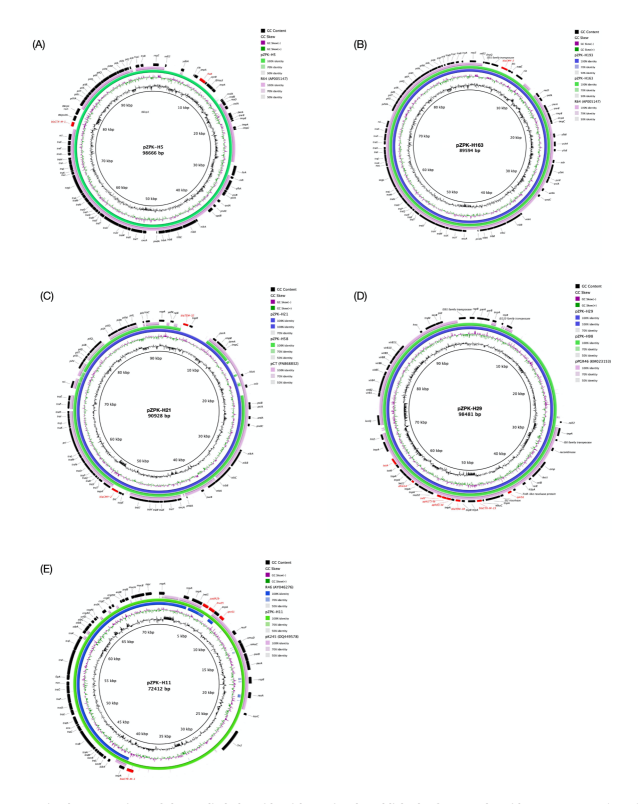
 $^{\circ}$  R = phenotypically resistant; S = phenotypically susceptible.

Table 4. Genomic characteristics of hybrid sequenced Escherichia coli isolates and plasmid replicons from barnacle geese.

Other virulence genes in isolate	astA, cea, etsC, gad, hlyF, iroN, iss, iucC, iutA, lpfA, ompT, sitA, terC, traT, tsh	cea, cia, cvaC, etsC, fyuA, gad, hlyF, iroN, irp2, iss, iucC, iutA, lpfA, mchF, ompT, sitA, terC, traT, tsh	air, celb, chuA, cia, eilA, etsC, gad, hlyF, hra, iha, iss, iucC, iutA, kpsE, kpsMIL K5, mcbA, ompT, papC, sitA, terC, traT	fyuA, gad, irp2, iss, kpsE, kpsMII <u>I</u> K10, lpfA, papC, sitA, terC	astA, celb, gad, gad, hra, iha, iss, iucC, iutA, kpsE, kpsMII, ompT, sitA, terC	cba, cma, etsC, gad, hlyF, iroN, iss, iucC, iutA, lpfA, ompT, sitA, terC, traT, tsh	fyuA, gad, irp2, iss, kpsE, kpsMII <u>I K</u> 10, lpfA, papC, sitA, terC	cif, eae, espA, espF, espJ, gad, iss, lpfA, nleB, sepA, terC, tir	cif, eae, espA, espF, espJ, gad, iss, lpfA, nleB, sepA, terC, tir
Other resistance genes in isolate	mdf(A), aph(6)- Id, aph(3″)-Ib, sul2, dfrA	mdf(A), aph(6)- Id, aph(3″)-Ib, dfrA5, sul2, tet(A)	mdf(A)	dfrA1, mdf(A), aadA1, tet(A)	mdf(A)	mdf(A), tet(A)	dfrA1, mdf(A), aadA1, tet(A)	mdf(A)	mdf(A)
Other plasmid replicons in isolate	IncFIB, IncFIC(FII), IncFII(29)	IncFIB, IncFII, IncFII(pCoo), IncQ1	Col156, Col8282, IncFIB, IncFII, IncFII	1	Col8282, ColpVC	IncFIA, IncFIB, IncFIC(FII), IncFII(pHN7A8)	1	ColpVC, IncFIB, IncFII	ColpVC, IncFIB, IncFII
Phenotype <sup>a</sup>	ESBL	ESBL	AmpC	ESBL	АтрС	AmpC	ESBL	AmpC	AmpC+ESBL
Virulence genes on ESBL- plasmid	cia	I	traT	1	traT	1	1	cia	cia
Other resistance genes on ESBL- plasmid	floR	qnrS1, aadA2b, Inu(F)	1	qnrS1, aph(6)- Id, aph(3")-Ib, sul2, dfrA14, tet(A)	I	1	qnrS1, aph(6)- Id, aph(3")-Ib, sul2, dfrA14, tet(A)	I	1
<i>bla</i> gene(s) on ESBL- plasmid	bla <sub>CTX-M-1</sub>	Ыа <sub>стх:м-1</sub>	Ыа <sub>сму-2</sub> , Ыа <sub>тем-</sub> <sup>32</sup>	<i>bla</i> <sub>СТХ-М-15</sub> ′ <i>bla</i> <sub>ТЕМ-18</sub>	bla <sub>CMY-2</sub>	Chromosomal point mutation in AmpC promoter	<i>bla</i> <sub>СТХ-М-15</sub> ′ <i>bla</i> <sub>ТЕМ-18</sub>	bla <sub>CMY-2</sub>	bla <sub>CMY-2</sub>
ESBL- plasmid (replicon; pMLST)	pZPK-H5 (IncI1; ST38, CC-3)	pZPK-H11 (IncN+IncR; IncN: ST1)	pZPK-H21 (IncK)	pZPK-H29 (IncY)	pZPK-H58 (IncK)	I	pZPK-H98 (IncY)	pZPK-H163 (IncI1; ST23, CC-2)	pZPK-H193 (IncI1; ST23, CC-2)
Number of total contigs	9	<del>-</del>	<b>o</b>	m	20	m	м	6	4
Sampling period	Fall	Fall	Fall	Fall	Fall	Fall	Fall	Spring	Spring
Isolate ( <i>E. coli</i> MLST)	H5 (ST359)	H11 (ST58)	H21 (ST115)	H29 (ST453)	H58 (ST1594)	H68B (ST3580)	H98 (ST453)	H163 (ST2178)	H193 (ST2178)

CC = donal complex, ESBL = extended-spectrum beta-lactamase; MLST = multilocus sequence type; pMLST = plasmid multilocus sequence type.

<sup>a</sup> Based on phenotypic tests.



**Figure 1. Circular comparison of the studied plasmids with previously published reference plasmids.** GC content and GC skew are depicted in the inner map with distance scale. Predicted coding sequences of the plasmid named within the circle are depicted in the outer ring with antimicrobial resistance genes highlighted in red. (A) pZPK-H5 compared with IncI1 type reference R64 (GenBank accession: AP005147), (B) pZPK-H163 and pZPK-H193 compared with IncI1 type reference R64 (GenBank accession: AP005147), (C) pZPK-H21 and pZPK-H58 compared with IncK type reference pCT (GenBank accession: FN868832), (D) pZPK-H29 and pZPK-98 compared with IncY type reference pPGR46 (GenBank accession: KM023153), (E) pZPK-H11 compared with IncN type R46 (GenBank accession: AY046276) and IncR type pK245 (GenBank accession: DQ449578).

accession: MT468651.1), and a *bla*<sub>CMY-2</sub>-harboring IncI1 ST2 plasmid recovered from a human clinical sample in Australia with 100% coverage and 98.75% identity (GenBank accession: HG970648.1). pZPK-H5 was found to be most similar, with *bla*<sub>CTX-M-1</sub>-harboring IncI1 ST3 plasmids, such as a plasmid obtained from a poultry sample in France (GenBank accession: MG648932.1) with 96% coverage and 99.79% identity, and a plasmid recovered from *E. coli* from a healthy human in Switzerland (GenBank accession: KJ484629.1) with 96% coverage and 99.90% identity. A cgMLST-based minimum spanning tree built with 75 discriminatory loci (95% occurrence) is visualized in Figure 2.

#### IncK type plasmids

Two isolates, H21 and H58, harbored IncK type replicons with  $bla_{\rm CMY-2}$ . Plasmid pZPK-H21 from isolate H21 was 90.9 kb in size with a G+C content of 52.51% and 108 CDSs. In addition to  $bla_{\rm CMY-2}$ , pZPK-H21 carried  $bla_{\rm TEM-32}$ . pZPK-H58 was 79.2 kb in size with a G+C content of 52.16% and 92 predicted CDSs. From both pZPK-H21 and pZPK-H58, only the virulence gene traT was identified. Pairwise alignment with BLASTn showed 88% coverage and 99.99% identity between pZPK-H21 and pZPK-H58.

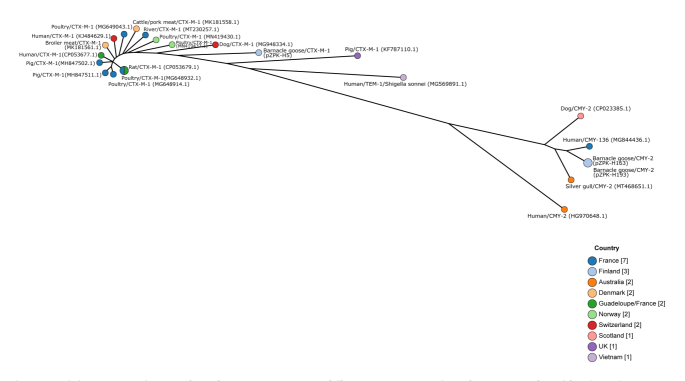
Similar to IncI1 plasmids, pZPK-H21 and pZPK-H58 were found to have a typical I-complex plasmid backbone structure with maintenance and stability, conjugation, shufflon, and pilus formation regions (Figure 1C).  $bla_{\text{TEM-32}}$  in pZPK-H21 was located in the accessory module, whereas  $bla_{\text{CMY-2}}$  in both

pZPK-H21 and pZPK-H58 was located near a transfer-associated region next to *tnpA*. BLASTn pairwise alignment with IncK type reference plasmid pCT (GenBank accession: FN868832) indicated that both pZPK-H21 and pZPK-H58 shared 92.42% identity with pCT, whereas pZPK-H21 shared 81% coverage and pZPK-H58 82% coverage.

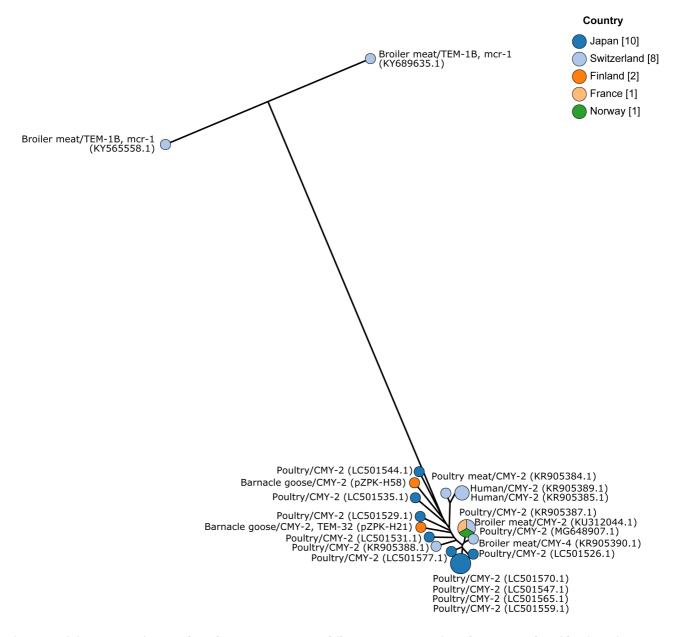
A BLASTn search against the NCBI database indicated that the most similar previously published plasmids consisted of poultry-associated IncK2 or IncB/O/K/Z replicons harboring  $bla_{\text{CMY-2}}$  (Table 1). pZPK-H21 was also found to be similar with two IncK2 plasmids derived from *E. coli* from a human urinary tract infection in Switzerland, with 94% coverage and 99.81% identity with plasmid pDV45 (GenBank accession: KR905384.1) and 92% coverage and 99.88% identity with plasmid p5312.29 (GenBank accession: KR905385.1). A cgMLST-based minimum spanning tree was built with 64 discriminatory loci (95% occurrence) and is visualized in Figure 3.

#### IncY type plasmids

Two isolates, H29 and H98, harbored IncY type replicons. pZPK-H29 from isolate H29 and pZPK-H98 from isolate H98 were highly similar when compared with pairwise alignment with BLASTn, resulting in 100% coverage and 99.94% identity. Both plasmids harbored  $bla_{\text{CTX-M-15}}$  and  $bla_{\text{TEM-1B}}$ . In addition to bla genes, both plasmids carried the resistance genes qnrSI, aph(6)-Id, aph(3")-Ib, sul2, dfrA14, and tet(A). Both plasmids were 98.5 kb in size, had a G+C content of 51.13% and 98



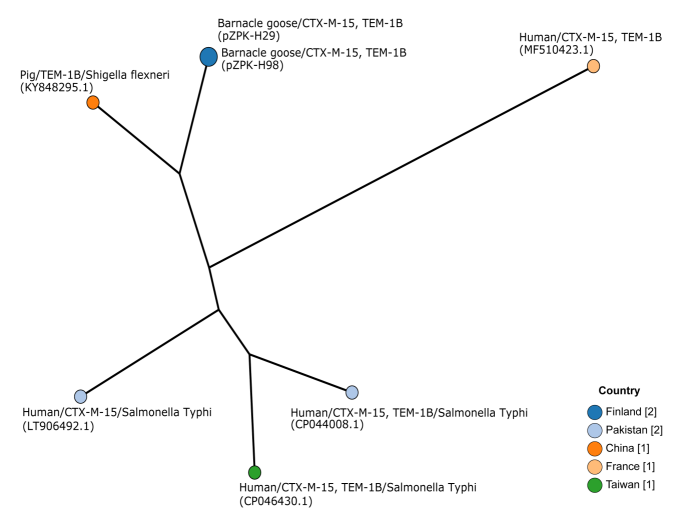
**Figure 2. Minimum spanning tree based on core genome multilocus sequence typing of IncI1 type plasmids.** Plasmids pZPK-H5, pZPK-H163, and pZPK-H193 compared with 20 previously published IncI1 plasmids from GenBank. Nodes are colored by the country of origin and node text includes information on the source, *bla* gene, and bacterial species if other than *Escherichia coli*. GenBank accession numbers are provided in parentheses.



**Figure 3. Minimum spanning tree based on core genome multilocus sequence typing of IncK type plasmids.** Plasmids pZPK-H21 and pZPK-H58 compared with 20 previously published IncK or IncB/O/K/Z plasmids from GenBank. Nodes are colored by the country of origin and node text includes information on the source, *bla* gene, and possible additional *mcr-1*. GenBank accession numbers are provided in parentheses.

predicted CDSs in pZPK-H29, and had a G+C content of 51.12% and 99 predicted CDSs in pZPK-H98. These IncY plasmids were not found to harbor any virulence genes. BLASTn comparison with previously described IncY type plasmid pPGR46 (GenBank accession: KM023153) demonstrated 71% coverage with both pZPK-H29 and pZPK-H98, and 99.98% identity with pZPK-H29 and 99.94% identity with pZPK-H98 (Figure 1D). A BLASTn search against the NCBI database identified only six IncY type previously published plasmids with available metadata to be similar to pZPK-H29 and pZPK-H98 (Table 1). Three of the similar previously

published plasmids were obtained from human clinical samples isolated from  $Salmonella\ Typhi$  (GenBank accessions: CP044008.1, LT906492.1, and CP046430.1) and two from  $E.\ coli$  also of human clinical sample origin (GenBank accessions: KM023153.1 and MF510423.1), all harboring  $bla_{CTX-M-15}$ . One IncY plasmid originated from  $Shigella\ flexneri$  from a pig fecal sample harboring  $bla_{TEM-1B}$  (GenBank accession: KY848295.1). Phylogenetic comparison with these six previously published IncY plasmids with a cgMLST-based minimum spanning tree included 20 discriminatory loci (95% occurrence) and did not indicate clusters (Figure 4).



**Figure 4. Minimum spanning tree based on core genome multilocus sequence typing of IncY type plasmids.** Plasmids pZPK-H29 and pZPK-H98 compared with six previously published IncY plasmids from GenBank. Nodes are colored by the country of origin and node text includes information on the source, *bla* gene, and bacterial species if other than *Escherichia coli*. GenBank accession numbers are provided in parentheses.

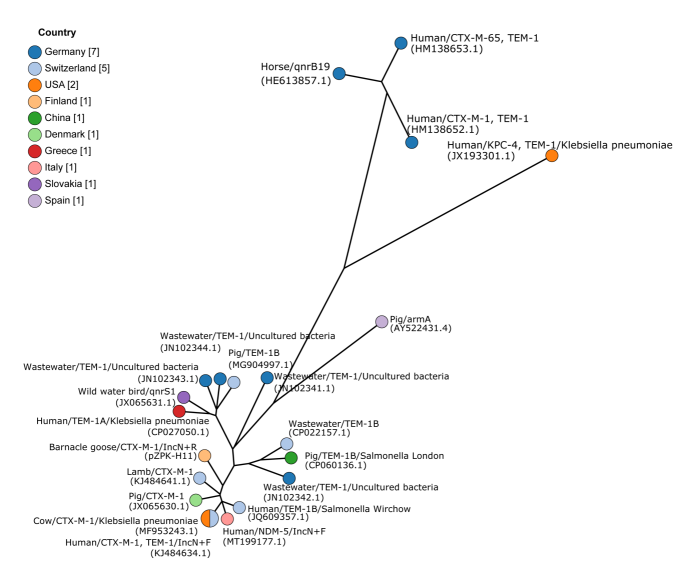
#### IncN+IncR multireplicon

From isolate H11, two plasmid replicons, IncN and IncR, were identified on the same contig. This multireplicon plasmid pZPK-H11 was 72.4 kb in size, with a G+C content of 50.91% and 78 predicted CDSs. Pairwise alignment with BLASTn indicated that the plasmid aligned with IncN reference plasmid R46 (GenBank accession: AY046276) with a coverage of 55% and 99.73% identity, whereas alignment with previously published IncR type plasmid pK245 (GenBank accession: DQ449578) resulted in a coverage of 31% and 98.53% identity (Figure 1E).

The multireplicon plasmid harbored  $bla_{\text{CTX-M-1}}$  and additional resistance genes qnrS1, aadA2b, and lnu(F). No virulence genes were detected on the multireplicon plasmid. The IncN replicon in the plasmid was identified as ST1 with pMLST analysis. Top two hits from a BLASTn search against the NCBI

database were IncF+IncN type multireplicons both isolated from  $E.\ coli$  of human origin, one in Italy with  $bla_{\mathrm{NDM-5}}$  (GenBank accession: MT199177.1; 60% coverage and 99.97% identity) and the other in Switzerland carrying  $bla_{\mathrm{CTX-M-1}}$  (GenBank accession: KJ484634.1; 59% coverage and 99.97% identity) (Table 1). Other top hits with BLASTn included IncN plasmids isolated from various sources, such as uncultured bacteria from wastewater treatment plant effluent in Germany, carrying  $bla_{\mathrm{TEM-1}}$  (GenBank accessions: JN102342.1, JN102344.1, and JN102343.1). A cgMLST-based minimum spanning tree was built with 34 discriminatory loci (95% occurrence) and is visualized in Figure 5.

pPK-H11 shared similar backbone structures with IncN reference R46, including the replicon *repA*, stability-related *stbABC*, *mucA*, and *mucB* involved in mutagenesis enhancement, genes involved in plasmid DNA protection from type I



**Figure 5. Minimum spanning tree based on core genome multilocus sequence typing of IncN+IncR multireplicon.** Plasmid pZPK-H11 compared with 20 previously published plasmids from GenBank. Nodes are colored by the country of origin and node text includes information on the source, *bla* gene, or other resistance gene if the *bla* gene was not present, bacterial species if other than *Escherichia coli*, and replicon type if other than IncN. GenBank accession numbers are provided in parentheses.

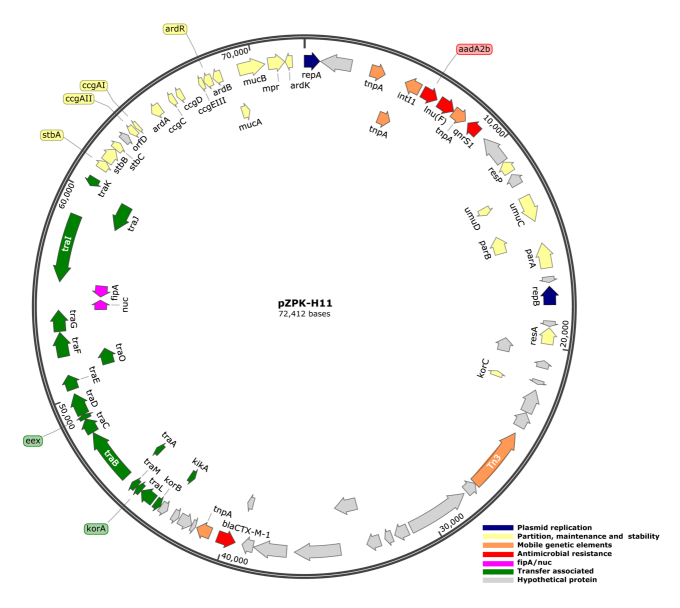
restriction enzymes (ccgC, ccgD, ccgAI, and ccgAII), antirestriction-associated ardA, ardB, and ardR, and two transfer-associated tra loci (traI, traJ, traK; traL, traM, traA, traB, traC, traD, traE, traO, traF, traG) (Carattoli et al., 2010; Delver & Belogurov, 1997; Dolejska et al., 2013). Similar to the R46 structure, fipA and nuc were located between tra loci in pZPK-H11. IncR plasmid backbone structures identified in pZPK-H11 included replicon repB, parAB, and umuCD and multimer resolvase, but toxin–antitoxin vagDC operon involved in plasmid maintenance and retA reverse transcriptase were not identified (Guo et al., 2016). Transcriptional regulator korC was identified in a region not aligning with either IncN or IncR reference plasmids (Ludwiczak et al., 2013). bla<sub>CTX-M-1</sub> was located next to mobile genetic element tnpA outside of a

multidrug resistance cassette containing aadA2b, lnu(F), and qnrS1, which were located downstream of repA. The pZPK-H11 structure is visualized in Figure 6.

### Discussion

The purpose of this study was to identify ESBL/pAmpC-producing *E. coli* in migrating barnacle geese residing near humans and to investigate plasmids harboring *bla* genes. The *bla* genes were encoded on four different plasmid replicons from nine different samples: IncI1 and IncK, the rare IncY, and the multireplicon IncN+IncR.

Interestingly, IncY harboring  $bla_{CTX-M-15}$  was found from two sequenced isolates with the same bacterial ST type (ST453)



**Figure 6. Genetic structure of multireplicon plasmid pZPK-H11.** Predicted coding sequences and their orientation are represented by arrows colored based on the function of the gene product. Size of the plasmid is depicted on the outer circle.

and AMR gene profile, indicating clonal spread of the bacteria between the animals or a common source. Plasmids of the IncY group are considered rarely detected phage-like plasmids with low copy numbers (Meyer et al., 1986; Rozwandowicz et al., 2018). ESBL-producing E. coli harboring IncY have been detected in clinical human isolates (Yasir et al., 2020), a wastewater treatment plant in China (Jiang et al., 2019), and environment and fish samples in Tanzania (Moremi et al., 2016). The aforementioned human-associated finding of IncY raises the question of the origin of IncY in the barnacle geese samples. It can be speculated that the origin of AMR in geese is from anthropogenic sources.

Notable is the finding of the pZPK-H11 multireplicon IncN+IncR harboring  $bla_{\rm CTX-M-1}$  in one of the sequenced isolates. IncN has been commonly found in animal fecal microbiota (Carattoli, 2009) and has been identified as a disseminator of  $bla_{\rm CTX-M-1}$  among animals and humans in Europe (Dolejska et al., 2013).  $bla_{\rm CTX-M-1}$  has often been recovered from IncN ST1 (Rozwandowicz et al., 2018). In addition to  $bla_{\rm CTX-M-1}$ , the IncN+IncR multireplicon pZPK-H11 harbored resistance genes aadA2b, lnu(F), and qnrS1. qnr genes have been previously associated with IncN plasmids from different Salmonella serovars isolated from human and poultry samples in the Netherlands (García-Fernández et al., 2009). It has

been shown previously that IncR plasmids are able to form multireplicons with IncN, IncA/C, IncHI, and IncF type plasmids (Drieux *et al.*, 2013; Jing *et al.*, 2019; Papagiannitsis *et al.*, 2013; Qu *et al.*, 2019). Previous studies indicate that IncR plasmids lack genes involved in conjugation (Bielak *et al.*, 2011; Chen *et al.*, 2006), which indicates that pZPK-H11 is a putatively conjugative plasmid contributing to the transfer-associated genes located on the IncN region of the plasmid.

IncI1 carrying bla<sub>CTX-M-1</sub> was identified in one of the isolates in our study. The finding of two identical IncI1 plasmids harboring bla<sub>CMY-2</sub> from two samples, pZPK-H163 and pZPK-H193, indicates very recent clonal transmission within the flock or the possibility of samples originating from the same goose individual. IncK type plasmids with bla<sub>CMY-2</sub> were recovered from two isolates in our study. IncK type plasmids carrying bla<sub>CMV-2</sub> and  $bla_{\text{CTX-M-14}}$  have been previously isolated mainly from E. coli from animal sources in Europe (Rozwandowicz et al., 2018). The most prevalent resistance genes in the isolates in our study were AmpC type  $bla_{\mathrm{CMY-2}}$  followed by  $bla_{\mathrm{CTX-M-1}}$  and  $bla_{\text{CTX-M-15}}$ . The finding of  $bla_{\text{CMY-2}}$  and  $bla_{\text{CTX-M-1}}$  on IncII plasmids is a common finding among poultry (Accogli et al., 2013; Leverstein-van Hall et al., 2011), and these genes and plasmid replicons have been identified previously also from other wild avian species, such as seagulls and pelicans in Florida (Poirel et al., 2012). Wild birds of different species in Catalonia have also been found to harbor  $bla_{\text{CTX-M-15}}$  and  $bla_{\text{CMY-2}}$ among other beta-lactamases, such as  $\mathit{bla}_{\scriptscriptstyle{\text{OXA-48}}}$  in a barn owl (Darwich et al., 2019). No carbapenemases were recovered in our study. In addition, in a study by Bonnedahl et al. (2009), 9.4% of wild yellow-legged gulls in the south of France were found to harbor ESBL-producing E. coli and 6% more specifically  $bla_{\text{CTX-M-1}}$ . The finding of beta-lactamases, especially in wild birds habituating aquatic environments, is worrisome because of the potential of bacterial transmission between humans and animals via surface water. Wild gulls in Sweden were found to harbor  $bla_{\text{CTX-M-15}}/bla_{\text{CTX-M-14}}$ -producing E. coli similar to human and surface water isolates (Atterby et al., 2017). Wild birds may acquire resistance genes from anthropogenic sources and circulate resistance determinants again back into the human population, although contact with wild birds has been estimated to contribute only 0.3% of community-acquired intestinal carriage of ESBL/pAmpC-producing E. coli in humans in the Netherlands (Mughini-Gras et al., 2019). In addition to human health, wild migratory birds may play a role as disseminators of human-derived AMR to remote areas, potentially endangering the welfare of native animal species (Hernández & González-Acuña, 2016).

Interestingly, the finding of ESBL/AmpC-producing *E. coli* in 4.5% of the studied barnacle geese fecal samples is slightly less than the prevalence of ESBL/AmpC-producing *E. coli* and *Klebsiella pneumoniae* described in Finnish asymptomatic human carriers (6.3%) (Rintala *et al.*, 2018). AMR has been studied in wild birds previously, but this was the first time it was studied in migratory birds in Finland, and with WGS applied. The prevalence of 4.5% is relatively low compared with other studies conducted with different avian species. In

a study conducted in the Netherlands (Veldman et al., 2013), ESBL/AmpC-producing E. coli were found in 15.7% of 21 different bird species tested, mainly originating from aquaticassociated species. In Spain, 15% of sampled birds were found to be positive for ESBL/pAmpC-producing E. coli (Alcalá et al., 2016), whereas in birds of prey in Germany and Mongolia the prevalence was found to be 13.8% and 10.8%, respectively (Guenther et al., 2012). The lower percentage in the present study could be partly explained by the different bird species examined. Barnacle geese do not feed on landfills or other human-related waste areas like gulls in certain areas (Bonnedahl et al., 2014) or other bird or mammal species like birds of prey but, as a waterfowl, may come into contact with human wastewater. Although the prevalence of ESBL/AmpC-producing E. coli in our study was less than in some other birds studied earlier, barnacle geese that have habituated to densely human-populated areas, such as recreational parks and public beaches (Simões et al., 2010), pose the potential risk of AMR spread to humans. Parks and public beaches can quickly become contaminated with fecal matter while flocks of hundreds of birds feed and defecate in the area. Additionally, household pets and children may come into contact with fecal matter more easily.

It has to be taken into account, however, that the sample size was limited. There have been studies of the effect of ultraviolet light on *E. coli* survival (Vermeulen *et al.*, 2008; Whitman *et al.*, 2004), but to assess the effect of sunlight and other environmental factors on *E. coli* or resistance genes and plasmid survival in bird feces, more studies would need to be conducted. Seven out of nine samples positive for ESBL/AmpC-producing *E. coli* originated from samples collected in fall 2017, and temporal changes have been identified in ESBL-producing Enterobacteriaceae previously in urban Swedish mallards, with higher prevalence during warm summer months (Hessman *et al.*, 2018).

To assess the actual risk of AMR in wildlife, bacteria, resistance genes, and mobile genetic elements have to be typed to a level where comparison between different sources, i.e., humans and animals, is plausible. WGS provides a powerful tool for studying phylogeny between different bacterial and plasmid reservoirs, and also enables the early detection of new resistance genes or successful plasmids. Long-read sequencing of the isolates allows for even more in-depth analysis of the location of resistance genes on plasmid replicons.

As *E. coli* belong to the normal intestinal microbiota of animals and virulence potential varies between strains, the public health relevance of ESBL/AmpC-producing *E. coli* derived from barnacle geese feces should be interpreted with caution. By examining virulence traits, however, it is possible to identify potential pathogens. The increased serum survival gene *iss* was found from all of the sequenced isolates. *Iss* has been identified in extraintestinal pathogenic *E. coli* (ExPEC) as well as in avian pathogenic *E. coli* (APEC) (Johnson *et al.*, 2008; Tivendale *et al.*, 2004). *Iss* and *iroN* have been found to be prevalent in *E. coli*, causing salpingitis in egg-laying

hens (Poulsen et al., 2020), and have also been identified in E. coli in poultry production (Kim et al., 2020; Oikarainen et al., 2019). The glutamate decarboxylase gene gad was also prevalent in the sequenced isolates, and it has been identified frequently in APEC in broilers together with iss and iroN (Azam et al., 2020). These findings highlight the potential of AMR transmission via virulent E. coli strains between bird species, as well as between other animals and potentially humans.

Wildlife may serve as a reservoir for resistant bacteria (Carroll et al., 2015) and more than 60% of emerging infectious diseases since the 1940s have been zoonotic (Jones et al., 2008), which makes the continuous surveillance of wildlife pathogens extremely important. Our study highlights the fact that wild birds living in close contact with humans carry bacteria with plasmid-mediated beta-lactamases and may pose a potential risk of bacterial clonal spread or horizontal gene transfer through the environment.

#### Conclusion

Wild migratory geese were found to carry ESBL/AmpCproducing E. coli with diverse bacterial STs and the assessment of samples with WGS revealed the location of resistance genes in specific plasmid replicons. A novel multireplicon, IncN+IncR, carrying  $bla_{\text{CTX-M-1}}$  was recovered, and humanassociated  $\mathit{bla}_{\text{CTX-M-15}}$  was found from two IncY plasmids. The findings indicate that wildlife carrying plasmids and resistance

genes are potentially worrisome for public health. In addition to successful, more frequently identified plasmid replicons IncI1, IncK, and IncN, the rare IncY plasmid and IncN+IncR multireplicon plasmids were recovered in our study, indicating that wildlife carry epidemic plasmids and also serve as potential disseminators of plasmids currently viewed as uncommon in humans and livestock. Although the prevalence of ESBL/AmpC-producing E. coli was relatively moderate at 4.5%, the transmission potential should not be underestimated, especially in urban areas with dense human and animal populations. Continuous and prudent monitoring of resistance determinants in different environments is vital for understanding evolving resistance patterns and the epidemiology of AMR within and between the different One Health compartments.

#### Data availability

EMBL-EBI European Nucleotide Archive: Raw reads and plasmid sequences. Accession number PRJEB42655; https:// identifiers.org/ena.embl:PRJEB42655.

Accession numbers for each isolate are provided in Table 2.

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Reviewer Report 04 January 2022

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## Magdalena Maria Zając 🗓



Department of Microbiology, National Veterinary Research Institute, Puławy, Poland

The authors of manuscript titled "Plasmids conferring resistance to extended-spectrum betalactamases including a rare IncN+IncR multireplicon carrying blaCTX-M-1 in Escherichia coli recovered from migrating barnacle geese (Branta leucopsis)" have presented interesting and welldescribed research which can have a wide audience. They proved that migratory geese carry ESBL/ampC producing E. coli and identified and described in details plasmids harboring resistance genes by analyses of whole genome sequences.

The manuscript is easy to read and clear. All manuscript sections are high quality writing, detailed description of used methods and tools allows to repeat the research. The sequences are easily accessible at NCBI database. I do not have any remarks.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and does the work have academic merit? Yes

Are sufficient details of methods and analysis provided to allow replication by others?

If applicable, is the statistical analysis and its interpretation appropriate? Not applicable

Are all the source data underlying the results available to ensure full reproducibility? Yes

Are the conclusions drawn adequately supported by the results?

Yes

**Competing Interests:** No competing interests were disclosed.

Reviewer Expertise: AMR in Salmonella and E. coli, whole genome sequencing analysis

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.

Reviewer Report 08 June 2021

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## Angela H. A. M. van Hoek

Centre for Infectious Disease Control (CIb), National Institute for Public Health and the Environment (RIVM), Bilthoven, The Netherlands

The authors of "Plasmids conferring resistance to extended-spectrum beta-lactamases including a rare IncN+IncR multireplicon carrying  $bla_{\text{CTX-M-1}}$  in *Escherichia coli* recovered from migrating barnacle geese (*Branta leucopsis*) have written a very clear and easy to read paper that is of interest to many readers in the current One Health and AMR era. The whole genome sequencing data is publicly available, which allows re-analysis of the results. I have only a few issues with the manuscript.

I find the term core <u>genome</u> MLST (cgMLST) while comparing various IncI1 type (figure 2), IncIK type (figure 3), incY type (figure 4) and IncN+IncR type (figure 5) <u>plasmids</u> confusing. The legends of these figures also miss the number of loci included in the analysis, one has to find this somewhere in the results sections. The number of allelic distances would be much appreciated in the minimum spanning trees of figures 2 - 4.

The finding of EPEC isolates among the spring isolates should be described in more detail. Since so many tools were used from the Center for Genomic Epidemiology website (like MLST, ResFinder, PlasmidFinder) this reader/reviewer would also appreciate the SerotypeFinder results ending up in the manuscript.

Is the work clearly and accurately presented and does it cite the current literature? Yes

Is the study design appropriate and does the work have academic merit?

Are sufficient details of methods and analysis provided to allow replication by others? Yes

If applicable, is the statistical analysis and its interpretation appropriate?

Yes

Are all the source data underlying the results available to ensure full reproducibility?  $\mbox{\em Yes}$ 

Are the conclusions drawn adequately supported by the results?

Yes

Competing Interests: No competing interests were disclosed.

Reviewer Expertise: WGS analysis of AMR (pathogenic) zoonotic bacteria

I confirm that I have read this submission and believe that I have an appropriate level of expertise to confirm that it is of an acceptable scientific standard.