Diversity and prevalence of colibactin- and yersiniabactin encoding mobile genetic elements in enterobacterial populations: insights into evolution and co-existence of two bacterial secondary metabolite determinants

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1 **1 Abstract**

2 The bacterial genotoxin colibactin interferes with the eukaryotic cell cycle by causing double-3 stranded DNA breaks. It has been linked to bacterially induced colorectal cancer in humans. Colibactin is encoded by a 54-kb genomic region in Enterobacteriaceae. The colibactin genes 4 commonly co-occur with the yersiniabactin biosynthetic determinant. Investigating the 5 6 prevalence and sequence diversity of the colibactin determinant and its linkage to the versiniabactin operon in prokaryotic genomes, we discovered mainly species-specific lineages 7 8 of the colibactin determinant and classified three main structural settings of the colibactin-9 yersiniabactin genomic region in Enterobacteriaceae. The colibactin gene cluster has a similar 10 but not identical evolutionary track to that of the yersiniabactin operon. Both determinants could have been acquired on several occasions and/or exchanged independently between 11 12 enterobacteria by horizontal gene transfer. Integrative and conjugative elements play(ed) a central role in the evolution and structural diversity of the colibactin-yersiniabactin genomic 13 14 region. Addition of an activating and regulating module (clbAR) to the biosynthesis and transport module (*clbB-S*) represents the most recent step in the evolution of the colibactin 15 16 determinant. In a first attempt to correlate colibactin expression with individual lineages of colibactin determinants and different bacterial genetic backgrounds, we compared colibactin 17 18 expression of selected enterobacterial isolates in vitro. Colibactin production in the tested Klebsiella spp. and Citrobacter koseri strains was more homogeneous and generally higher 19 20 than that in most of the *E. coli* isolates studied. Our results improve the understanding of the 21 diversity of colibactin determinants and its expression level, and may contribute to risk 22 assessment of colibactin-producing enterobacteria.

23 2 Introduction

The non-ribosomal peptide/polyketide hybrid colibactin is a secondary metabolite found in a variety of bacterial species of the family *Enterobacteriaceae*. The colibactin biosynthetic machinery is encoded by a 54-kb large polyketide synthase (*pks*) or *clb* genomic island[1], which includes 19 genes. The largest part of the island consists of a section of overlapping or closely spaced genes: *clbB* to *clbL* and *clbN* to *clbQ*, which are aligned on the same strand and code for components of the biosynthesis complex. The colibactin assembly line is supplemented with a dedicated transporter, encoded by *clbM*, and a resistance-conferring 31 protein encoded by *clbs* [2, 3]. Two additional genes required for colibactin production are 32 located ca. 400 bp upstream of the first biosynthesis gene *clbB* in the opposing reading direction: the *clbR* gene coding for an auto-activating, *pks* island-specific transcription factor 33 and the phosphopantetheinyl transferase-encoding gene *clbA*, which is crucial for activation 34 35 of polyketide biosynthesis complexes (Fig. 1) [4-6]. Between these two divergent transcription units, there is a "variable number of tandem repeat" (VNTR) region, which comprises a varying 36 number of a repeating octanucleotide sequence (5'-ACAGATAC-3') depending on the isolate 37 38 [2].

Recently, the structure of the colibactin molecule has been proposed [3, 7-9]. Yet, the biological role of colibactin is still under discussion. Colibactin can interfere with the progression of the eukaryotic cell cycle, presumably by cross-linking DNA resulting in doublestrand DNA breaks in genomic instability in eukaryotes [1, 10, 11]. The ability to produce colibactin has been described to increase the pathogenic potential of the producing bacteria and to promote colorectal cancer development [12-17], but has also been related to beneficial effects to the host [3, 18-20].

46 Initially, the *pks* island has been described in extraintestinal pathogenic *E. coli* to be chromosomally inserted into the *asnW* tRNA locus in close proximity to another tRNA(*Asn*) 47 gene-associated pathogenicity island, the so-called "high pathogenicity island" (HPI). The HPI 48 49 harbors an additional polyketide determinant coding for the metallophore yersiniabactin biosynthetic machinery [1, 21]. As members of the Enterobacteriaceae are generally not 50 known as archetypal secondary metabolite producers the origin of the *pks* island remains to 51 52 be further investigated. Interestingly, the colibactin determinant has also been detected as 53 part of an "integrative and conjugative element" (ICE) in different enterobacteria. This ICE also integrates near a tRNA(Asn) locus into the bacterial chromosome and commonly carries the 54 versiniabactin gene cluster [2, 22]. It has been suggested that the close linkage observed 55 between the colibactin and yersiniabactin gene clusters results from the functional 56 57 interconnection between the colibactin and yersiniabactin biosynthetic pathways via the 58 phosphopantetheinyl transferase ClbA, which can also contribute to the biosynthesis of versiniabactin [5, 23]. The highly conserved colibactin determinant has so far been detected in 59 a spectrum of strains belonging to the *Enterobacteriaceae* family: most commonly among *E*. 60 61 coli strains of the phylogroup B2, followed by Klebsiella pneumoniae isolates, but also in 62 Citrobacter koseri and Klebsiella aerogenes [2, 18, 24]. Less conserved variants or homologs of 63 the colibactin gene cluster have been described phenotypically or based on nucleotide 64 sequence data in an *Erwinia oleae* strain, the honey bee symbiont *Frischella perrara*, and the 65 marine α -proteobacterium *Pseudovibrio* [24, 25].

66 Based on the low sequence similarity between the enterobacterial colibactin genes and the 67 two homologous polyketide determinants in *F. perrara* and *Pseudovibrio* as well as on their association with mobile genetic elements (MGEs) or at least mobility-associated genes, one 68 69 can hypothesize that the colibactin gene cluster is spread by horizontal gene transfer, maybe 70 via an ICE-like element [2, 24]. While in most *Enterobacteriaceae* the colibactin determinant is 71 typically associated with an ICE, the characteristic mobility and transfer features of an ICE are absent in the sequence context of the *pks* island in *E. coli* phylogroup B2 strains. Nevertheless, 72 the pks island in E. coli remains mobilizable and transferable through external factors, 73 74 supporting the hypothesis that former MGEs can undergo a stabilization (homing) process upon their chromosomal integration [26-28]. 75

76 Studies addressing the prevalence of the colibactin genes were so far mainly focused on 77 Klebsiella spp. or Escherichia coli backgrounds. The scarcity of data in other prokaryotic 78 species regarding its distribution and the structure of the associated MGE makes it challenging 79 to reliably further characterize the transmission and evolution of this polyketide determinant [18]. Previous data show that the prevalence varied from 5.3% to 25.6% in *Klebsiella* and from 80 81 9.5% to 58% in Escherichia, highlighting an enrichment of pks island in specific ecological 82 niches, whereas studies with a broader screening approach resulted in a prevalence of 14% in 83 Klebsiella and 9.5% in Escherichia isolates, respectively [2, 29-34]. Notably, in health-related 84 studies, a higher association of the *clb* genes was observed amongst strains with an increased 85 virulence potential, with a prevalence of as much as 78.8% for Klebsiella subgroups and 72.7% 86 for colorectal cancer-associated E. coli isolates [29, 32, 35-37]. The colibactin genes are frequently found in hyper-virulent and multidrug-resistant *K. pneumoniae* isolates [38, 39]. 87

The obvious prevalence of the colibactin gene cluster in specific enterobacterial species combined with the description of more distantly related homologous determinants has sparked our interest in a better understanding of the spread and evolution of the colibactin determinant and its genetic context in bacteria. Therefore, our study aimed to investigate the prevalence and diversity of the colibactin determinant also in isolates outside of the *Enterobacteriaceae* family. Furthermore, we compared colibactin expression levels among enterobacterial isolates carrying different lineages of colibactin determinants as a first

95 attempt to assess the functional context of the bacterial genetic background, pathogenicity,

96 and colibactin expression.

97

98 3 Impact Statement

99 Colibactin can act as a bacterial genotoxin and thus promote colorectal cancer development. Little is known about the origin, diversity and prevalence of the colibactin genes (*clb*) within 100 101 prokaryotes. The *clb* genes are closely associated with pathogenicity islands or integrative and 102 conjugative elements (ICEs). We screened roughly 375,000 prokaryotic genomes to analyze the diversity and evolution of such mobile genetic elements among bacterial populations. 103 Interestingly, *clb* genes are only present in a subgroup of enterobacteria, mainly *E. coli*, 104 105 Klebsiella spp. and Citrobacter koseri. The clb determinant, together with the versiniabactin (vbt) gene cluster, belong to an ICE in most of the *clb*-positive enterobacteria, especially in 106 Klebsiella. We show that both determinants, though in principle freely transferable within 107 108 bacteria, have a mainly species-specific phylogeny, and that colibactin expression levels were 109 species-independent. Recombination promoted the structural diversification of the ICE in different species, including its successive degeneration that led to the establishment of the 110 111 colibactin and versiniabactin islands in *E. coli* phylogroup B2 strains. Our results not only 112 illustrate differing evolutionary tracks of the *clb* and *ybt* determinants in different 113 enterobacterial species, but also highlight the important of ICEs for genomic variability in enterobacteria and the evolution of archetypal pathogenicity islands. 114

116 4 Methods

117 4.1 Bacterial strains and media

- 118 For cultivation, bacteria were grown as batch cultures in lysogeny broth (LB) (10g/l tryptone,
- 119 5g/l yeast extract, 5g/l sodium chloride) at 37°C. Strains used in this study are listed in the
- 120 Supplemental Material (Table S1).

121 4.2 DNA extraction and sequencing

DNA extraction of the enterobacterial strains was performed using the MagAttract[®] HMW DNA Kit (Qiagen, Hilden, Germany) according to the manufacturer's recommendations. To prepare paired-end libraries we used the Nextera XT DNA Library Preparation kit (Illumina, San Diego, CA, USA). Libraries were sequenced on the Illumina MiSeq sequencing platform using v2 sequencing chemistry (500 cycles) or on the Illumina NextSeq500 system using v2.5 chemistry (300 cycles). Accession numbers of in-house sequences submitted to the NCBI GenBank database are included in Supplemental Material (Table S1).

129 4.3 Genome selection and phylogenetic analysis

All genome sequences not generated in this study were obtained from publically available 130 prokaryotic genomes (NCBI GenBank). The quality of in-house sequenced genomes was 131 132 checked with FastQC v0.11.5 (https://github.com/chgibb/FastQC0.11.5/blob/master/fastqc), and low-quality reads were trimmed using Sickle v1.33 (https://github.com/najoshi/sickle). The 133 134 processed reads were de novo assembled with SPAdes v3.13.1 [40] and annotated with prokka v1.12 [41]. The genomes were screened for the presence of >45 kb of the complete 135 pks genomic island using standalone BLAST+ v2.8.1 [42] and antiSMASH v5.0.0 [43]. The pks 136 island found in the genome of the E. coli strain M1/5 (accession no. CP053296) was used as a 137 138 reference sequence. The VNTR and the sequence stretch of the *pks* island that spans between *clbJ* and *clbK* were excluded from analysis as these regions are prone to misassembly. 139

The contigs that align to the colibactin genes were ordered using ABACAS v1.3.1 [44] and multiple sequence alignment was generated using Kalign v3.1.1 [45]. Recombinant regions were detected and removed using Gubbins v2.4.1 [46]. The recombination filtered polymorphisms were then used to generate a maximum likelihood phylogeny of the colibactin determinant using RAxML v8.2.11 [47] under the GTR-GAMMAX model from 9974 polymorphic sites. The branch support of the maximum likelihood tree was estimated by

bootstrap analysis of 200 replicate trees. The homologous gene cluster found in *F. perrara*was used as an outgroup. The phylogeny of the corresponding *ybt* islands was generated with
a similar approach. The generated trees were visualized using itoL (https://itol.embl.de).

149 4.4 Phylo-grouping of *pks*-positive strains

The *E. coli* and *Klebsiella* strains that harbored the colibactin gene cluster were allocated to their corresponding sequence types using mlst v2.16.1 (<u>https://github.com/tseemann/mlst</u>), which detects sequence types using the PubMLST typing schemes. The *Escherichia* strains were further classified into their phylogenetic lineages using the standalone tool, EzClermont v0.4.5 (<u>https://github.com/nickp60/EzClermont</u>).

The analysis of the diversity of the colibactin and versiniabactin gene clusters involved 155 virulence gene multi-locus sequence typing (MLST) for both polyketide determinants as 156 157 previously described [38]. Briefly, the allele sequences of sixteen genes of the colibactin gene 158 cluster (*clbACDEFGHILMNOPQR*) as well as of eleven genes of the versiniabactin determinant (fyuA, ybtE, ybtT, ybtU, irp1, irp2, ybtA, ybtP, ybtQ, ybtX, ybtS) were extracted from the 159 individual genomes and analyzed for allelic variations. Each observed combination of alleles 160 161 was assigned a unique colibactin sequence type (CbST, listed in Table S5) or versiniabactin 162 sequence type (YbST, listed in Table S6).

163 4.5 Variable number tandem repeat (VNTR) detection

164 The VNTR copy number present within the colibactin determinant (upstream of *clbR*) was 165 detected using the standalone version of tandem repeats finder v4.09 [48]. The VNTR copy 166 number distribution was visualized using R v3.4.3 (https://www.r-project.org/index.html).

167 4.6 Detection of *E. coli* virulence markers for pathotyping

For pathotyping, the *clb*-positive *E. coli* strains were *in silico* screened for the presence of
different *E. coli* pathotype marker genes using BLAST+ v2.8.1 (Supplemental Material, Tab. S).
These genes were used as markers for *E. coli* pathotypes: enteroaggregative *E. coli* (EAEC),
enterohemorrhagic *E. coli* (EHEC), enteropathogenic *E. coli* (EPEC), enterotoxigenic *E. coli*(ETEC), diffusely adhering *E. coli* (DAEC), uropathogenic *E. coli* (UPEC), and newborn
meningitis-causing *E. coli* (NMEC).

174 4.7 Quantification of colibactin expression through N-myristoyl-D-asparagine

Following an approach described by Bian and colleagues [49], a collection of colibactin-175 producing strains of the main species harboring the colibactin determinant was characterized 176 for their ability to produce colibactin under in vitro growth conditions. For this purpose, we 177 178 quantified N-myristoyl-D-asparagine (N-Myr-D-asparagine) a byproduct during colibactin 179 maturation. The amount of this intermediate extrapolates the resulting colibactin amount produced. After growing the bacteria for 24 h at 37°C in glass tubes in 5 ml LB supplemented 180 with 200 μ l of a water/XAD-16-resin slurry, bacterial cells were harvested by centrifugation. 181 182 The pelleted bacteria-slurry mixes were sedimented, filtered, and dissolved three times in 183 acetone with increasing volume (12ml, 100 ml, and finally 200ml). Afterward, the solvent was exchanged by rotary evaporation and replaced by 1.6 ml methanol. The sample was further 184 concentrated by centrifugation (10 min, 15000 rpm at 4°C), followed by drying 1,5 ml of the 185 186 solution in a vacuum centrifuge and subsequent resuspension in 50 μ l methanol. 30 μ l of these processed samples were measured by Ultra Performance Liquid Chromatography 187 coupled to High-Resolution Mass Spectrometry (UPLC-HRMS) conducted on a Thermo 188 Scientific Ultimate 3000 RS with a Waters Acquity BEH 100*2.1mm 1.7µm 130A column 189 190 (eluent A: 0.1% formic acid in ddH2O, eluent B: 0.1% formic acid in acetonitrile), where a flow 191 rate of 0.6ml/min followed by a Bruker Maxis II-4G, 150-2500 m/z and a scan rate of 2Hz was applied. To enable quantification of N-Myr-D-asparagine, we used 250 mM cinnarizine as 192 193 internal standard and normalized peak areas based on the internal standard and the optical density (OD₆₀₀) of the bacterial culture. 194

196 **5 Results**

197 5.1 Prevalence of colibactin determinant

Of the 374,754 publicly accessible prokaryotic genomes (as of 30.06.2019) that were screened 198 199 for the presence of the colibactin gene cluster, 1,969 genomes carried this polyketideencoding operon. An additional 200 *clb*-positive enterobacterial genomes determined in-200 201 house were added to the analysis (Tab. 1, Supplemental Material, Tab. S1). The *clb* gene 202 cluster was detected in several enterobacterial species, most frequently in E. coli and K. 203 pneumoniae isolates, but also to a lesser extent in K. aerogenes, C. koseri, E. cloacae, E. hormaechei, K. michiganensis, S. marcescens, and E. oleae. The colibactin determinant was, 204 205 however, not detectable in 112,546 Salmonella enterica and 41 S. bongori genomes 206 (Supplemental Material, Tab. S4), but in one out of eight genomes of unspecified Salmonella isolates. We did not detect the *clb* genes in 2,634 *Shigella* spp., 861 *Yersinia* spp., 677 *Serratia* 207 spp., 186 Proteus spp. and 69 Morganella spp. genomes (Supplemental Material, Tab. S4). A 208 less well-conserved homolog of the colibactin determinant was detected in three F. perrara 209 210 genomes. It should be noted that the number of genomes of Klebsiella spp. and E. coli 211 analyzed in this study are markedly higher than those of the other species and lineages due to 212 the sequencing bias towards *Klebsiella* spp. and *E. coli* strains. Accordingly, a reliable 213 statement on the prevalence of the colibactin determinant in the different species cannot be 214 made.

215 **5.2** Diversity of the colibactin determinant

To find out whether the prevalence of the colibactin gene cluster is restricted to specific 216 217 phylogenetic lineages of *E. coli* and *Klebsiella* spp., the sequence types of the corresponding *E*. coli and Klebsiella spp. isolates were further analyzed. As shown in Fig. S1, the clb gene cluster 218 was enriched in a small subset of E. coli STs (twelve out of 11,537 STs, as of 30.10.2020), K. 219 220 aerogenes STs (two out of 214 STs, as of 30.10.2020), and K. pneumoniae STs (six out of 5,237 221 STs, as of 30.10.2020), respectively. In these twelve E. coli STs, between 58% and 94% of the allocated isolates carry the *clb* determinant. A high percentage (ca. 96%) of the *K. aero*genes 222 223 ST4 and ST93 included in our study harbored the colibactin genes. In the tested K. pneumoniae strains, all ST3 isolates were *clb*-positive, more than 75% of the analyzed ST23 224 225 and ST234 isolates carried the colibactin gene cluster, whereas this was only the case for a 226 significantly lower percentage of the K. pneumoniae isolates allocated to ST11, ST258, and

227 ST48. Table S2 (Supplemental Material) contains a complete list of STs to which colibactin-228 positive *E. coli* and *Klebsiella* isolates have been assigned.

229 The nucleotide sequences of the *clb* gene cluster extracted from the 2,169 strains 230 (Supplemental Material, Tab. S1) were used to generate a recombination-free phylogeny of 231 the colibactin determinant as shown in Fig. 2A (Fig. S2 and S3 for branch support values and strain labels/assembly IDs, Fig. S4 indicates predicted recombination events in the *clb* gene 232 cluster). Serratia marcescens strain MSU97 isolated from a plant source, Erwinia oleae strain 233 234 DAPP-PG531 isolated from olive tree knot, Klebsiella michiganensis strain NCTC10261 of an 235 unknown source, and the *E. coli* phylogroup E strain 14696-7 isolated from the pericardial sac of a white-tailed deer (Odocoileus virginignus) harbor the most genetically distant variants of 236 the colibactin determinant (Fig. 2A). Within the Enterobacteriaceae, a large group of clb gene 237 238 clusters can be defined, which is dominated by two highly conserved clades present in E. coli 239 phylogenetic lineage B2 and different Klebsiella spp. isolates, respectively. The colibactin determinants detected in E. cloacae and E. hormaechei belong to the Klebsiella clades of clb 240 241 loci, whereas the clb gene clusters found in C. koseri and in an unspecified Salmonella isolate represent an independent clade, i.e. *clb*6 (Fig. 3). In a few other *E. coli* and *Klebsiella* spp. 242 243 genomes, the *clb* determinant can be distinguished from the two major conserved clades of 244 colibactin determinants observed in Klebsiella or E. coli. An even more divergent group 245 comprises the *clb* gene clusters of mainly *E. coli* phylogroup A, B1, and a few B2 isolates, but 246 also of some *K. pneumoniae* strains (Fig. 3, belonging to *clb* clades *clb*1 and *clb*2).

247 Although the *clb* gene clusters found in the large clade of *E. coli* B2 strains are highly conserved, individual ST-specific lineages, such as *clb*10 (from *E. coli* ST141 and ST2015 248 249 strains) and *clb*12 (from *E. coli* ST121 strains) can still be described within this clade. Beyond that, we also observed multiple lineages per sequence type, such as *clb*2, *clb*11 and *clb*16 250 251 found in E. coli strains of ST73 (Fig. 3). Three lineages of the clb locus were predominantly detectable in Klebsiella pneumoniae strains. They belong to the most distant Klebsiella 252 253 ST3/ST380 clade (*clb*1), the remaining large and diverse ST258/ST11 clade (*clb*8), and finally the hypervirulent ST23 clade (clb9) (Fig. 3). A phylogeny of the colibactin gene cluster inferred 254 255 from concatenated amino acid sequences of the 17 *clb* genes (Fig. S8) was very similar to the 256 aforementioned recombination-free nucleotide-based phylogeny (Fig. 3).

257 5.3 Diversity of the yersiniabactin determinant in colibactin-positive bacteria

The majority (>98%) of the *clb*-positive enterobacterial strains also harbored the 258 versiniabactin genes (*ybt*) (Fig. 3, 3rd circle). The *E. coli* strains of phylogroup A, B1, and E as 259 260 well as the tested K. michiganensis and E. oleae strains carrying the most genetically distant 261 lineages of the colibactin gene cluster, together with the F. perrara strains used as an outgroup, are ybt-negative (Fig. 2B). There were no strains that carried multiple copies of the 262 263 colibactin gene cluster; yet two well-separated copies of the ybt determinant were, however, found in *C. koseri* strains ATCC BAA-895 and 0123A 53 520. It should be noted that the latter 264 265 strain is derived from a metagenome (Fig. S6). The phylogenetic analysis indicated that all ybt 266 operons from *E. coli* clustered together (Fig. 2B). Alike the *clb* gene cluster, also the *ybt* locus of the *E. coli* phylogroup B2 strains was highly conserved. In contrast, the sequence 267 comparison of the *ybt* determinants of *Klebsiella* spp. resulted in different lineages, which 268 269 correlate with lineages ybt1, 12, and 17 (ICEKp10), ybt9 (ICEKp3), and ybt4 (originating from a 270 plasmid) previously described by Lam and colleagues [38].

271 5.4 Congruent phylogeny of the colibactin and yersiniabactin determinants

The strong coexistence of the colibactin and versiniabactin determinants on the one hand and 272 the description of different evolutionary lines of *clb* and *ybt* determinants in different 273 enterobacterial species on the other hand led us to analyze whether both gene clusters can 274 275 predominantly be transferred individually or together. Our results indicate that the clades of 276 the evolutionary lineages of the *clb* and *ybt* loci are chiefly species/genus-specific. The 277 phylogeny of *clb* and *ybt* determinants is largely congruent, with the *ybt* gene cluster being, 278 even more, species/genus-specific than that of the *clb* gene cluster. However, in some strains 279 we observed evidence of interspecies transfer of these genes: The *clb* and corresponding *ybt* 280 determinants of the C. koseri isolates NCTC10769, ATCC BAA-895 and E. coli strains 239A, 926, 281 GN02370, MOD1-EC5674/5 were allocated to the large Klebsiella-dominated lineage clb8 (Fig. 3). Additionally, the *clb* gene cluster of *K. pneumoniae* strain k2265 was found within lineage 282 283 clb6, which predominantly represents C. koseri isolates. However, the ybt determinant of K. pneumoniae strain k2265 belonged to the ybt12 lineage represented by K. oxytoca isolates. 284 285 Similarly, the vbt determinant of the aforementioned strains E. coli strains GN02370 and C. koseri ATCC BAA-895 belonged to lineage ybt4 (plasmid originating ybt loci) instead of ybt17. 286

Regardless of the aforementioned exceptions, clades of the *clb* gene cluster usually correlated with the corresponding clade of *vbt* genes (Fig. 3, 3^{rd} and 4^{th} circle).

289 5.5 Genetic structure of the MGEs harboring the colibactin determinant

To further investigate whether the colibactin and yersiniabactin determinants are jointly 290 291 distributed by horizontal gene transfer and to obtain clues to the underlying mechanism, we 292 compared the chromosomal context of the two polyketide biosynthesis gene clusters and the genetic structure of associated mobile genetic elements (MGEs). We observed species-specific 293 294 structural differences of the chromosomal regions harboring the *clb*, and *ybt* gene clusters 295 (Fig. 4). In *E. coli* phylogroup B2 strains, the *clb*, and *ybt* determinants are present as part of 296 two individual pathogenicity islands (PAIs) with their cognate integrase and different tRNA 297 genes (class I of *clb*-harboring MGE). Both PAIs are located neighboring each other in the 298 chromosome. Within class Ia, the two PAIs are separated by a type 4 secretion system (T4SS)-299 encoding operon (virB) and a region that includes two conserved gene sets (Set 1 and Set 2), 300 different tRNA(Asn) loci and an integrase gene (see Table S3 for genes present in the different 301 conserved gene sets). This region is shown to have been diminished in class Ib structural 302 variants, where only gene virB1 of the T4SS determinant is left alongside the conserved 303 region. In the predominant E. coli structural variant, class Ic, the complete T4SS operon 304 (including virB1) has been lost together with gene set 1, the integrase gene was exchanged 305 and a DNA stretch comprising the genes *yeeO*, tRNA(Asn), cbl and gltC was inverted (shown in 306 red, Fig. 4). The region separating the pks island and the HPI was reduced from a 40-kb (in class Ia) to a 15-kb stretch in class Ic. In contrast, within all *Klebsiella* strains, the *clb* and *ybt* 307 gene clusters are part of an ICE, and are separated by a T4SS-encoding operon (virB) and the 308 309 *hha* gene coding for the hemolysin expression-modulating protein. Downstream of the *ybt* gene cluster an integrase gene is located followed by a set of genes involved in Fe/Mn/Zn 310 311 metabolism (structural class II of *clb*- and *ybt*-harboring chromosomal regions) (Fig. 4). Interestingly, one type of such ICEs, is located next to genes necessary for microcin E492 312 313 biosynthesis (class IIc, Fig. 4). Enterobacter hormaechei strains harbor a structurally similar ICE to that of Klebsiella strains. In most C. koseri strains, however, the two polyketide 314 determinants are separated by a large 250-kb chromosomal region. The T4SS-related genes 315 are closely positioned to the *clb* genes while the gene set involved in Fe/Mn/Zn metabolism is 316 317 located downstream of the ybt determinant. Only a minor fraction of enterobacterial isolates 318 analyzed displayed some variation regarding gene content and synteny of these three main

319 classes of colibactin and versiniabactin-encoding chromosomal regions. The structure of *clb* 320 and ybt regions that do not conform to these major classes are as shown in Figure S5. Instead 321 of class I, several E. coli strains carried class II like chromosomal regions where the T4SS and 322 the Fe/Mn/Zn metabolism genes were present. In E. coli strain HVH128 none of the three 323 main classes colibactin and versiniabactin-encoding regions could be identified. Although both polyketide determinants are co-localized with one integrase gene each, they are widely 324 325 separated on this strain's chromosome. K. pneumoniae strains TUM14001, TUM14002, 326 TUM14009, TUM14126, and WCHKP13F2 harbored two T4SS-encoding gene clusters in close 327 proximity of the *clb* and *ybt* gene clusters and lacked the Fe/Mn/Zn genes, whereas K. 328 pneumoniae strain UCI110 was also missing the Fe/Mn/Zn metabolism-related genes. In contrast to the other C. koseri isolates, we detected a class II- instead of a class III-type clb-ybt 329 330 region in *C. koseri* isolate BAA-895.

331 5.6 Organization of the colibactin gene cluster

The *clb* gene cluster is composed of 19 genes, which are required for the regulation, 332 biosynthesis, and transport of colibactin. The origin of this gene cluster in unclear. We, 333 therefore, compared the structure of the gene clusters representing the homologous *clb* locus 334 335 found in F. perrara and the phylogenetically most distant and potentially older clb 336 determinants relative to the E. coli B2 type of clb determinant (Fig. 2A), which are present in S. 337 marcescens, E. oleae, K. michiganensis, and E. coli phylogroup E strain 14696-7(Fig. 5). The 338 various *clb* determinants correspond in terms of the structure of the genes coding for the biosynthesis machinery, transport, and resistance to colibactin (*clbB-clbS*) and resemble the 339 structure of the well-described *clb* locus in *K. pneumoniae/K. aero*genes/*E. coli* B2 strains. The 340 341 individual *clb* determinants differ more clearly in the presence and localization of the genes involved in the regulation and activation of colibactin biosynthesis (clbR and clbA). These two 342 343 genes are absent in the homologous gene cluster found in F. perrara and in the clb determinant in S. marcescens. However, in F. perrara, a phosphopantetheinyl transferase 344 345 coding for a homolog of ClbA (43% amino acid similarity) and a radical S'-adenosylmethionine (SAM) enzyme-encoding gene are found directly downstream of *clbS*. In *S. marcescens* a helix-346 turn-helix (HTH)-type regulatory protein homologous to *clbR* is encoded by a gene located 347 upstream of clbB (78% amino acid similarity). In E. oleae and K. michiganensis a SAM enzyme-348 349 encoding gene is present directly downstream of *clbA*. Although the colibactin gene clusters in 350 E. oleae and K. michiganensis have a high nucleotide sequence similarity of ca. 99.77%, the

predicted coding regions of *clbC/D* and *clbH/I/J/K/L/M/N* are noticeably different due to multiple frameshift deletions in *K. michiganensis*. The structure of the *clb* locus found in phylogroup E *E. coli* strain 14696-7 already corresponds to the structure of *E. coli* strains of phylogroup B2 (Fig. 1 and Fig. 5), yet this gene cluster shows the least sequence similarity of all tested *clb* gene clusters in non-B2 *E. coli* isolates (Fig. 2A) to the determinant occurring in *E. coli* strains of phylogroup B2.

Looking at the G+C plot of the colibactin gene clusters, it is obvious that all investigated 357 358 enterobacterial *clb* gene clusters show a very similar G+C plot, which has a significantly higher 359 average G+C content and differs significantly from that of the *clb* homologous gene cluster in F. perrara (Fig. 5). The G+C content profile of these gene clusters indicates that there are two 360 regions of low G+C content in the enterobacterial *clb* determinants: the region including *clbA* 361 362 and *clbR* (at position ca. 1,500 bp - 3,000 bp of the colibactin gene cluster) and the region 363 spanning *clbD* and *clbE* (at position ca. 15,000 bp – 16,500 bp of the *clb* gene cluster). The G+C content drop in the region including *clbA* and *clbR* (at position ca. 1,500 bp - 3,000 bp of the 364 365 colibactin gene cluster) is associated with a predicted recombination site, which is located 366 upstream of or interrupting *clbB* (Fig. S4).

The comparison of structural features of the *clb* gene cluster also included the VNTR region located upstream of *clbR* in the *clbR-clbB* intergenic region. The size of the VNTR region has been described to range from 2 to 20 (Putze *et al.*, 2009). The VNTR copy number distribution in ca. 1,300 *clb*-positive genomes demonstrated that there is a preference for VNTR regions ranging from 7-10 copy numbers. Copy numbers from 18-34 were present in only a few strains (Figure S6). Species and/or ST-specific copy number variation was not observed.

373 Comparative genomic analysis of multiple colibactin-encoding determinants based on (draft) genome sequences led to the observation that the homologous genes *clbJ* and *clbK* are prone 374 375 to fusion/deletion (Lam et al., 2018). We also observed that in several assemblies of the clb gene cluster 625 bp from the 3' end of *clbJ* and 3,540 bp from the 5' end of *clbK* including the 376 11 bp intergenic region are missing, for a total of 4,174 bp (Fig. S5). The assembly of our 377 378 internally generated genome sequences produced by short read (Illumina) sequencing 379 showed this *clbJ*/K fusion/deletion. However, since assemblies of sequence data of the same 380 strains generated by a long-read sequencing technology (PacBio), where the long reads 381 covered both genes, had both *clbJ* and *clbK* completely present, we assume that the *clbJ/K*

fusions described are artificial and result from erroneous assemblies of short-read sequencingdata.

384 5.7 Quantification of colibactin synthesis in selected strains

To investigate a possible correlation between the genetic structure of the *clb* determinant or 385 386 the genetic background of the corresponding host strain with colibactin expression, we 387 quantified N-myristoyl-D-asparagine levels produced in vitro by selected clb-positive E. coli, Klebsiella spp. and C. koseri strains covering the diversity of clb determinants in these species 388 (Fig. 6). Based on the detected relative amount of N-myristoyl-D-asparagine produced, the 389 390 investigated isolates can be roughly divided into two groups: One group included most of the 391 measured E. coli strains that produced only very low relative amounts of N-myristoyl-Dasparagine. In contrast, the tested C. koseri, K. aerogenes, and K. pneumoniae isolates and the 392 E. coli isolates CFT073 and N1 showed a 3 to 70-fold higher N-myristoyl-D-asparagine 393 394 production. Also within the C. koseri, K. aerogenes, and K. pneumoniae isolates, we found 395 differences in the relative N-myristoyl-D-asparagine levels. However, these differences were 396 not as strong as among the eight E. coli isolates studied. The observed relative N-myristoyl-Dasparagine levels do not indicate phylogroup, ST, or species-specific differences in colibactin 397 398 production. For example, the E. coli strain 1873, although the clb gene cluster present in this 399 strain is phylogenetically more closely related to that of *E. coli* strain N1, shows a significantly 400 weaker N-myristoyl-D-asparagine production than E. coli N1. Similarly, it should be noted that 401 C. koseri MFP3 produces less N-myristoyl-D-asparagine than other closely related C. koseri.

402

403 6 Discussion

404 *Prevalence and mobility of the colibactin gene cluster in* Enterobacteriaceae

405 The ability of MGEs to be exchanged between and within species plays a major role in the 406 extent and speed of microbial evolution. Because MGEs are known to emerge and evolve separately from the host, it is important to explain the development and diversity of MGEs 407 408 independently. In our study, we investigated on the one hand the nucleotide sequence variability of the colibactin gene cluster with the associated versiniabactin determinants, and 409 410 on the other hand the structural diversity of the MGEs hosting the colibactin and yersiniabactin determinants responsible for their horizontal distribution. Both polyketide 411 412 determinants together could be detected in certain members of the Enterobateriaceae,

413 mainly in E. coli, K. pneumoniae, K. aerogenes, and C. koseri, but also in Enterobacter cloacae, Enterobacter hormachei, and possibly in uncharacterized Salmonella sp. isolates. The 414 415 colibactin, but not the versiniabactin determinants could also be detected in *K. michiganensis*, 416 E. oleae, and S. marcescens and a few phylogroup A and B1 E. coli isolates (Fig. 2A). 417 Interestingly, the colibactin genes have not yet been described in not only archaeal genomes but also S. enterica and S. bongori genomes, although we have screened more than 112,000 418 419 Salmonella spp. genomes. Overall, it is remarkable that the *clb* gene cluster was only found in 420 extraintestinal clinical or in fecal isolates of healthy hosts, but not in enterobacterial 421 diarrhoeal pathogens such as Yersinia spp., Salmonella spp., Shigella spp. as well as the various intestinal pathogenic *E. coli* pathotypes. This observation is consistent with previously 422 423 published data [2, 50]. In this context, it would be interesting to study in the future whether 424 and in which way colibactin expression supports extraintestinal pathogenicity or intestinal 425 persistence and colonization, but is detrimental to the pathogenesis of diarrhoeal pathogens. Possibly, the genetic background also plays an important role in the horizontal distribution 426 427 and establishment of MGEs carrying both polyketide determinants. The fact that the 428 colibactin determinant has so far been preferentially distributed in only some, often highly 429 virulent STs in E. coli and K. pneumoniae or K. aerogenes and not more broadly within the 430 respective species (Fig. 2) [38, 39], could also indicate that the transmission, uptake or 431 chromosomal integration of these MGEs is restricted. It is interesting to note that the *clb* gene 432 cluster as a whole is highly conserved and usually characteristic of the respective species or 433 genus (Fig. 3). Nevertheless, ST-specific variants have been found within a species, e.g. in K. pneumoniae ST3 and ST23. Several groups of sequence variants of the *clb* gene cluster can 434 435 also be found within one ST, such as in the E. coli ST73 and ST95 (Fig. 3). These results show that, on the one hand, intraspecies transfer of the colibactin determinant can happen, but on 436 the other hand, certain adaptations of the *clb* genes to a specific genetic background can also 437 occur at the nucleotide level. In addition, examples of interspecies transfer of the *clb* and *ybt* 438 439 genes can be seen between Klebsiella spp. and E. hormachei strains. Similarly and in contrast 440 to the majority of *E. coli* isolates, we found *clb* and *ybt* gene clusters in some *E. coli* isolates, which are assigned to the large clade of *Klebsiella/Enterobacter/Citrobacter-specific clb* and 441 442 *ybt* variants (Fig. 2A and 3). Apart from the interspecies transfer of the entire *clb*- and *ybt*containing MGE, we also identified an example that shows that the two polyketide gene 443 clusters can also be exchanged independently, as in the case of the *clb* gene cluster of K. 444

pneumoniae strain k2265, which belongs to colibactin clade *clb*6 (predominantly *C. koseri* lineage of *clb* loci), whereas the *ybt* determinant in this strain is assigned to the yersiniabactin clade *ybt*12 instead of *ybt*3, which is usually associated with *C. koseri* strains carrying clade *clb*6 (Fig. 3).

449 Structural diversity of the colibactin-yersiniabactin region

The structural analysis of the genomic region comprising the *clb* and *ybt* determinants in their 450 chromosomal sequence context is an important aspect to understand the evolution of these 451 452 polyketide determinants and their origin. In principle, three structural constellations (classes I 453 to III) can be described, in which the *clb* and *ybt* gene clusters are present (Fig. 4). Class I 454 depicts the *clb* and *vbt* gene clusters found in the majority of *E. coli* isolates, each associated 455 with a tRNA(Asn) and an integrase gene. In class Ia, the clb genes are chromosomally inserted 456 at the tRNA locus asnV. Our analyses suggest that in class I, the type 4 secretion system gene 457 cluster ("mobilization module") and conserved neighboring genes (set 1) have been lost in a 458 stepwise process, from class Ia to Ic. A further structural modification in this region is 459 represented by the inversion of the yeeO-tRNA(Asn)-cbl-gltC gene set (Fig. 4, red arrow), as a 460 result of which in class Ic, the tRNA gene asnW is located closest to the *clb* genes. Taking into 461 account that the *E. coli* strains with class Ia and Ib structures are found in the potentially 462 earlier phylogenetic clades, we hypothesize that the MGE harboring the *clb* genes was 463 introduced into the *E. coli* chromosome separately from that carrying the *ybt* determinant. 464 Both MGEs were then progressively modified as described above. We do not yet have an explanation why the resulting class Ic colibactin-yersiniabactin region, which has been 465 described as two pathogenicity islands (PAIs) comprising the colibactin and versiniabactin 466 467 determinants, respectively [1, 2], is only found in phylogroup B2 strains and only there has it 468 become so successful.

Class II includes different variants of an ICE, in which the two polyketide determinants are 469 present in association with a type 4 secretion system-encoding "mobilization module" and a 470 471 "module" consisting of genes that contribute to Fe/Mn/Zn metabolism (Fig. 4, Supplementary Table S3). This type of ICE was found in *Klebsiella* spp. and *E. hormachei* and in a few cases in 472 E. coli and C. koseri. Unlike in class I and III, the ICE in class II does not only have different 473 tRNA(Asn) loci serving as chromosomal insertion sites, but also lacks tRNA(Asn) and integrase 474 475 genes in between the *clb* and *ybt* genes. In a population-wide analysis of *Klebsiella* spp. strains, this ICE was designated ICEKp10 and described as being associated with different 476

477 combinations of *ybt* and *clb* gene lineages [38]. In contrast to class I and II colibactin478 yersiniabactin regions, the *clb* and *ybt* gene clusters are located far apart on the chromosome
479 in most of the *C. koseri* genomes studied (class III) (Fig. 4).

480 The existence of an ICE that unites the *clb* and *ybt* gene clusters is the easiest way to explain 481 the co-localization and the joint transfer of the two determinants and thus the high 482 correlation of *clb* and *ybt* phylogenetic clades (Fig. 3, 4th circle) in *Klebsiella* spp. strains. 483 Despite slight differences in the sequence context and different chromosomal insertion sites 484 (Fig. 4), the ICEs of the four class II variants have an overall identical genetic structure (Fig. 4). 485 The uptake of this ICE thus leads to the acquisition of both the *clb* and the *ybt* gene clusters. The presence of the Fe/Mn/Zn metabolic genes neighboring these ICE variants with an 486 487 additional integrase gene indicates recombination processes that can alter the genetic 488 structure of the ICE. The clear separation between the *clb*/T4SS module and the *ybt*-Fe/Mn/Zn 489 metabolism module in the *C. koseri* genomes points towards rearrangement/relocation of the ybt region, after ICE integration into the chromosome. The fact that C. koseri strain ATCC BAA-490 491 895 possesses in addition to the complete ICE (class II) a second ybt gene cluster (99.98% nucleotide similarity to the ybt genes present in the ICE) that is located far away from the ICE 492 493 (Fig. S6), supports the hypothesis that the individual polyketide gene clusters can also be 494 integrated into the genome independently of each other. This state could result, for example, 495 from the initial chromosomal integration of different ICE variants, as described by Lam and 496 colleagues in Klebsiella spp. [38]. As a result of deletion events, through which individual 497 modules are subsequently deleted from one of the two ICEs, the second copy of the ybt gene 498 cluster remains in the genome as a fragment of the degenerated ICE. The presence of two 499 non-identical T4SS modules in the K. pneumoniae strains TUM14001, TUM14002, TUM14009, 500 TUM14126, and WCHKP13F2 (Fig. S6 and Fig. 3, *clb*4) associated with the *clb* or *ybt* module (Fig. S6) could be the result of such degeneration of different ICEs. In this way, our 501 502 observations on the phylogeny (Fig. 2) and structure (Fig. 4) of the two polyketide 503 determinants and their sequence context can be reconciled, which show that despite the 504 predominant species/genus specificity, there are also sequence type-specific lineages of the *clb* genes, which do not necessarily have to match that of the associated *ybt* genes. 505

506 In this context, one could imagine that the arrangement of *clb* and *ybt* determinants in *E. coli* 507 strains (class I) also results from independent integration events of different MGEs, which 508 subsequently degenerated as a result of a stabilization process of these MGEs [26, 28]. This

premise is supported by the absence of *ybt* genes in the *clb*-positive *E. coli* strains, which carry the phylogenetically most distant *clb* determinants compared to the *clb* genes of phylogroup B2 isolates (Fig. 2A), along with the presence of integrase and tRNA(*Asn*) genes in close proximity to both polyketide determinants (Fig. 4, class I).

513 Evolution of the colibactin determinant

Homologs of the *clb* gene cluster were detected in marine alpha-proteobacteria such as 514 515 various Pseudovibrio sp. (isolates AD26, FO-BEG1, POLY-S9) and Pseudovibrio denitrificans (isolates DSM 17465 and JCM12308) [51]. Despite the general conformity of the genetic 516 517 structure of these gene clusters, their nucleotide sequence identity to the colibactin gene 518 cluster is guite low (<26%). Therefore, it was hypothesized that these *Pseudovibrio* isolates 519 have the potential to produce molecules related to colibactin [51]. Another homolog of the 520 colibactin determinant with a higher (62%) amino acid sequence identity is found in F. perrara 521 [24]. While the genes required for biosynthesis and transport of the polyketide are present, 522 the genes corresponding to *clbA* and *clbR* are missing in this gene cluster (Fig. 5). The case is 523 similar with the colibactin gene cluster in S. marcescens. While in F. perrara, a gene coding for 524 a clbA homolog and a gene coding for a SAM enzyme are located immediately downstream of 525 the *clbS* homolog, a *clbR* homolog is located upstream of *clbB* in *S. marcescens*. It is therefore 526 conceivable that the *clbA* homolog in *F. perrara* and the *clbR* homolog in *S. marcescens* are 527 involved in the activation or regulation of colibactin biosynthesis in these bacteria (Fig. 5). It 528 has been described that S-adenosylmethionine (SAM) is used in NRPS modules for colibactin biosynthesis [52]. Looking at the genetic structure of the *clbB-S* homologous genes cluster in 529 F. perrara and the clb gene cluster in S. marcescens, one can assume that genes involved in 530 531 the regulation and activation of the biosynthetic pathway including the SAM enzyme gene as 532 well as *clbA* and *clbR* homologs have been fused upstream to the already existing part of the island (*clbB-clbS*) to improve regulation of polyketide biosynthesis. Without having further 533 534 knowledge about the origin of the colibactin biosynthetic genes themselves, the acquisition of the regulatory/activating genes is obviously among the last evolutionary steps that led to the 535 536 structural organization of what we currently describe as the colibactin gene cluster. This hypothesis is supported by the abrupt decrease in G+C content and the presence of the 537 538 predicted recombination (Fig. S4) directly upstream of *clbB* in many *clb* determinants. 539 Furthermore, a module consisting of a gene for a SAM enzyme and a *clbA* homolog is not only located directly downstream from the *clbS* homolog in *F. perrara*, but also upstream from *clbR* 540

541 in *K. michiganensis* and *E. oleae*, which represent evolutionarily older variants of colibactin-542 positive *Enterobactericeae* (Fig. 5).

543 *Expression of colibactin in different hosts*

Furthermore, we investigated the question of how differently colibactin is expressed within 544 545 enterobacterial genera or even within different lineages of the same species. Interestingly, we observed an often lower production level of N-myristoyl-D-asparagine in E. coli isolates 546 compared to K. aerogenes, K. pneumoniae, and C. koseri (Fig. 6), which may be expected since 547 548 E. coli is described as a non-optimal producer of complex secondary metabolites [53]. 549 However, it is of interest that the amount of N-myristoyl-D-asparagine produced in E. coli strains CFT073 and N1 is comparable to that of other enterobacterial genera (Fig. 6). A 550 species- or lineage-specific ability to produce N-myristoyl-D-asparagine could not be 551 552 determined so far. Future studies will have to investigate which bacterial factors are 553 important for colibactin production and how the strain-specific differences in the expression of this polyketide come about. The systematic comparison of phenotypic colibactin 554 555 production with information on the genomic context, regulatory and metabolic properties of host strains, and their classification in a phylogenetic context should help us to identify 556 557 bacterial factors that affect colibactin synthesis.

558 7 Conclusion

559 The colibactin and versiniabactin gene clusters are highly conserved polyketide determinants within some Enterobacteriaceae. They usually coexist together in the genome and are also 560 561 linked to each other at the biosynthetic level. With the exception of E. coli, the two gene clusters are part of an ICE, which allows the horizontal transfer of both secondary metabolite 562 563 determinants usually within one species/genus. Bacteria of the genus Klebsiella played an important role in the evolution and distribution of both gene clusters. A large number of 564 different ICEs has been described in Klebsiella spp., which besides several other groups of 565 566 genes include the versiniabactin determinant [38]. Recombination and rearrangements events 567 between different ICE types may have contributed to the evolution of the ICE variants so far identified in Klebsiella spp. and other enterobacteria as well as to the further degeneration of 568 569 such MGEs leading to the colibactin and versiniabactin-encoding PAIs present in phylogroup 570 B2 E. coli strains (Fig. 7).

The phylogeny of the *clb* determinants does not determine the level of phenotypic colibactin production. The underlying bacterial factors responsible for the colibactin production efficiency of individual strains need to be identified in future work.

574 Our investigations provide deeper insights into the evolution of the colibactin gene cluster in 575 Enterobacteriaceae. Based on our findings, we can extend the current explanation for the coexistence and genetic co-localization of both gene clusters. The combination of a PPTase-576 577 encoding gene (*clbA*) with the *clbB-S* biosynthetic gene cluster during the evolution of the *clb* 578 determinant not only enabled the efficient activation of the colibactin biosynthesis machinery, 579 but also linked the colibactin and versiniabactin determinants, which are functionally connected by the activity of PPTase ClbA. This enables the bacteria to synthesize both 580 581 functionally different secondary metabolites, which leads to a stabilization of the co-existence 582 and co-localization of the two gene clusters in the genome. Our data underpin the importance 583 of mobile genetic elements, especially of ICEs, for genomic diversity and variability in 584 enterobacteria as well as for the evolution of more complex bacterial phenotypes, such as the 585 combined expression of the secondary metabolites colibactin and versiniabactin.

586 8 Author statements

587 8.1 Authors and contributors

588 H.W., A.W. and U.D. conceptualized the project. H.W., A.W., D.S. ran the analyses. R.M., M.S., and E.O.

589 contributed reagents and new tools. H.W., A.W., and U.D. analysed the data. H.W., A.W. and U.D.

590 wrote the manuscript. H.W., A.W., M.S., R.B., E.O., R.M. and U.D. edited and revised the manuscript.

591 All authors read, commented on and approved the final manuscript.

592 8.2 Conflicts of interest

593 The authors declare that there are no conflicts of interest.

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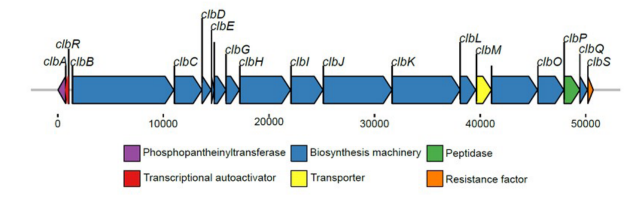
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Figure 1. Schematic representation of the genomic architecture of the *pks* island (ca. 54 kb) present in *E. coli* strain M1/5. The 19 genes within the island are colored with respect to their function. The island codes for a phosphopantetheinyl transferase (*clbA*, in purple), a transcriptional autoactivator (*clbR*, in red), multiple core biosynthetic genes (*clbB-clbL*, *clbN*, *clbO*, and *clbQ* in blue), a transporter (*clbM*, yellow), a peptidase (*clbP*, in green) and a resistance factor (*clbS*, orange).



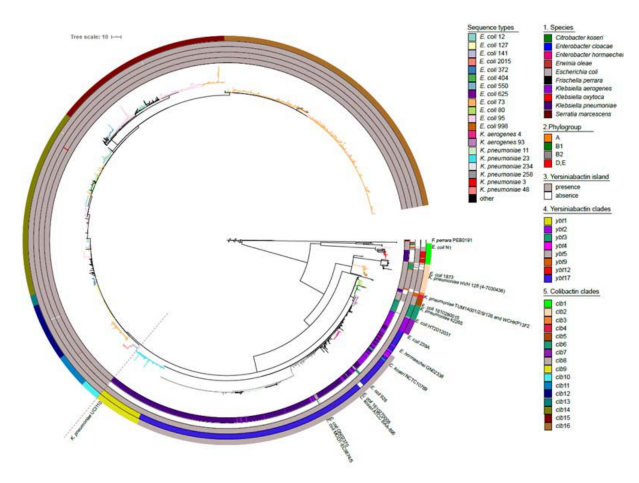
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Figure 2. Maximum-likelihood based phylogenetic analysis of the colibactin and yersiniabactin determinants. (A) Phylogenetic tree of the colibactin gene cluster (collapsed), B) phylogenetic tree of the corresponding *ybt* determinants (collapsed) using the genetically distant *K. michiganensis* strains as outgroup (Lam *et al.*, 2018). Additionally, the yersiniabactin sequence type (YbST) as defined by Lam and colleagues (Lam et al., 2018) associated with individual bacterial clades are indicated. The branch colors in both trees depict the prominent bacterial sequence type of the clade.

K. pneumoniae (ST23) - ybt1

K. aerogenes strain TUM15666 E. coli - ybt5

K. pneumoniae, K. aerogenes, E. cloacae, E. hormaechei - ybt4,ybt17



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Figure 3. Maximum-likelihood based phylogeny of the colibactin gene cluster detected in 2169 757 enterobacterial genomes. Every leaf represents a single sequence variant of the *clb* gene 758 cluster, which can be allocated to different lineages and clades. From innermost to outermost, 759 the 1^{st} circle indicates the species harboring the *clb* determinant; the 2^{nd} circle shows the *E*. 760 coli phylogroup, the 3rd circle shows the presence/absence of the *ybt* operon; the 4th circle 761 shows the versiniabactin sequence types (YbST) of the ybt determinant (from Fig. 1B) that 762 correspond to the *pks* island lineage present in the individual genome. The 5th circle shows the 763 different colibactin sequence types (CbST) of the *clb* gene cluster. The branch colors in the 764 center of the tree depict the prominent bacterial sequence types (Fig. 1). The large conserved 765 E. coli phylogroup B2 clade is separated from the large Klebsiella clade with a faint broken 766 767 line.

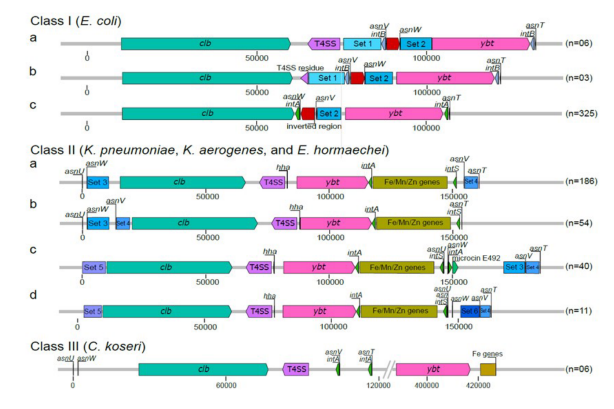
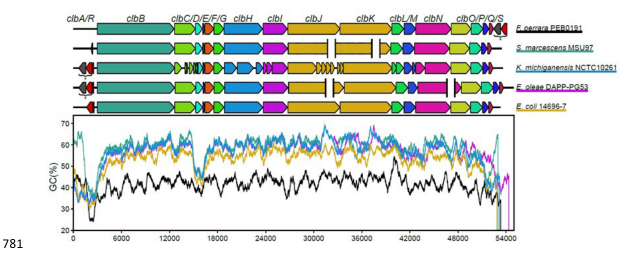


Figure 4. Structural variation of the colibactin and yersiniabactin-encoding chromosomal 770 region in E. coli, E. hormaechei, K. pneumoniae, K. aerogenes and C. koseri. The different 771 genetic structures and chromosomal insertion sites of the colibactin and/or yersiniabactin 772 determinants found within the three main structural classes are shown. The *clb* gene cluster 773 774 (teal green), T4SS module (purple), ybt gene cluster (pink), integrase genes (green), the 775 conserved sets of genes (Table S2) that are present up/downstream the two polyketide determinants, classed into sets (blue boxes) and the Fe/Mn/Zn module (yellow) are shown. 776 The number of genomes included in the tested set of genomes that harbour the different 777 778 structural variants is indicated in brackets. The colibactin-yersiniabactin chromosomal regions 779 that do not conform to these major structures are as shown in Figure S4.

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782 Figure 5. The structural organization and GC profiles of the *clb* determinants in the five most genetically distant bacterial strains (according to Fig. 2A). The genes that make up the 783 784 homologous pks gene cluster found in F. perrara and the most distant clb determinants present in S. marcescens, E. oleae, K. michiganensis and E. coli (phylogroup E) strain 14696-7 785 are depicted. The GC profile of the gene cluster in the different strains is shown alongside 786 with the colors underlining the different species. SAM genes and *clbA* homologues (*) are 787 788 shown downstream of the pks gene cluster in F. perrara and upstream of the clb determinant in *K. michiganensis* and *E. oleae*. The gaps in assembly are shown with white spaces. 789

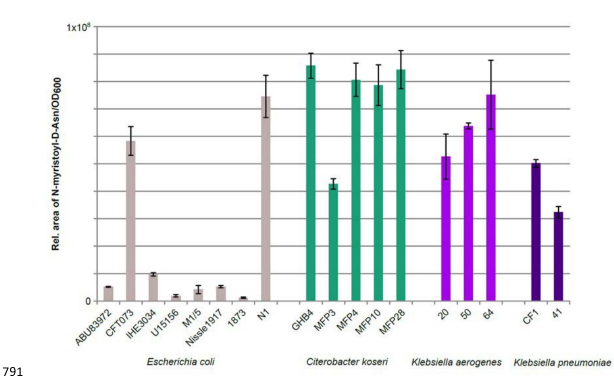


Figure 6. Comparison of colibactin production of different strains accessed by quantification of the precolibactin cleavage product N-Asn-D-myristol. This assay enabled us to compare the ability of different strains and different species to produce colibactin under controlled conditions *in vitro*. Measurements were conducted based on three biological replicates, means with standard deviations are as shown.

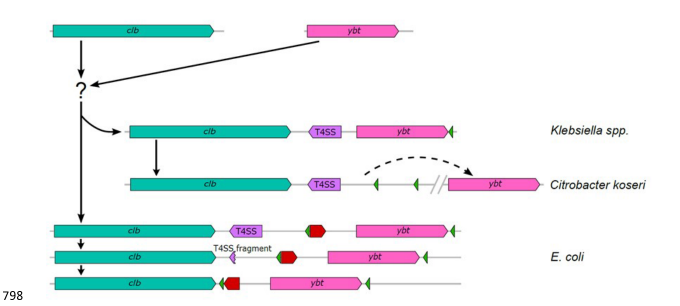


Figure 7. Schematic representation of the predicted evolution of the colibactin-yersiniabactin 799 800 genomic region in Enterobacteriaceae. The different elements of this region, i.e. the clb 801 determinant (teal green), T4SS module (purple), ybt gene cluster (pink), integrase genes 802 (green), and an invertible subset of genes (red arrow) are shown. Based on available genome 803 sequence data, we suggest a development from single MGEs containing the *clb* determinant 804 and the *vbt* gene cluster, respectively, towards the structural arrangement of both polyketide 805 determinants, which is now mainly found in enterobacterial populations. Black arrows (solid 806 or dashed) indicate possible directions of development and DNA rearrangements. After the 807 merge of the *clb* and *ybt* gene clusters into one MGE, represented by ICEKp10, there is 808 evidence that three different structural variants have evolved from it: In *Klebsiella* spp. 809 strains, the ICEKp10 has remained intact, whereas in C. koseri strains, a DNA rearrangement 810 and re-localization of the ybt determinant to a different chromosomal position has taken place. In E. coli, a gradual loss of the T4SS module and the inversion of a gene set between the 811 812 two polyketide determinants led to immobilisation or stabilisation of the ICE thus resulting the 813 two pathogenicity islands known as *pks* island and HPI, respectively.

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816 Supplemental Material

includes Supplementary Tables S1 (*clb*-positive bacterial strains), S2 (*clb*-positive STs in *E. coli*and *Klebsiella*), S3 (conserved gene sets associated with the *clb* and *ybt* determinants), S4 (list

of prokaryotic species included into the *clb* screen), S5 (list of ClbSTs), and S6 (list of YbSTs)

	No. of strains screened	<i>clb</i> -positive strains
Escherichia coli	19,183	1,462
Klebsiella pneumoniae	8,040	572
<i>Klebsiella aero</i> gene <i>s</i>	225	101
Citrobacter koseri	48	27
Enterobacter hormaechei	711	2
Enterobacter cloacae	610	2
Serratia marcescens	515	1
Klebsiella michiganensis	94	1
Salmonella sp.	8	1*
Erwinia oleae	1	1
Frischella perrara	3	3

Table 1. Bacterial species tested positive for the presence of the colibactin determinant

*unverified source organism (excluded from Refseq)