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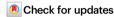
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Combined comparative genomics and clinical modeling reveals plasmid-encoded genes are independently associated with *Klebsiella* infection

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Members of the *Klebsiella pneumoniae* species complex frequently colonize the gut and colonization is associated with subsequent infection. To identify genes associated with progression from colonization to infection, we undertook a case-control comparative genomics study. Concordant cases (N = 85), where colonizing and invasive isolates were identical strain types, were matched to asymptomatically colonizing controls (N = 160). Thirty-seven genes are associated with infection, 27 of which remain significant following adjustment for patient variables and bacterial phylogeny. Infection-associated genes are not previously characterized virulence factors, but instead a diverse group of stress resistance, regulatory and antibiotic resistance genes, despite careful adjustment for antibiotic exposure. Many genes are plasmid borne, and for some, the relationship with infection is mediated by gut dominance. Five genes were validated in a geographically-independent cohort of colonized patients. This study identifies several genes reproducibly associated with progression to infection in patients colonized by diverse *Klebsiella*.

The *Klebsiella pneumoniae* species complex (hereby referred to as *Klebsiella*) is comprised of several species capable of causing severe infections, including bacteremia, pneumonia, and urinary tract infection (UTI). The member species are *K. pneumoniae*, *K. variicola*, *K. quasipneumoniae*, *K. quasivariicola* sp. nov.¹, and *K. africana*², which are genetically distinct but often clinically indistinguishable (reviewed in³). *Klebsiella* infections are a serious public health concern because they are a leading cause of healthcare-associated infections⁴, can

harbor multiple antimicrobial resistance (AMR) determinants, and can be hypervirulent. Antimicrobial resistant *Klebsiella* complicate disease treatment, leading to high mortality and healthcare costs whereas hypervirulent *Klebsiella* cause unique clinical manifestations such as pyogenic liver abscess and meningitis^{3,5,6}. *Klebsiella* colonization is the primary risk factor for disease^{7,8}, where the gut acts as a reservoir for disease-causing *Klebsiella* strains^{7,9}. Patient variables, including comorbidities and baseline laboratory values, partially explain which

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patients will progress to infection¹⁰. Furthermore, the high relative abundance of *Klebsiella* colonizing the gut is associated with an increased risk of infection in colonized patients¹¹. For hypervirulent *Klebsiella*, the genes associated with their unique presentations have been well-defined by molecular epidemiology and experimental studies (reviewed in¹²); however, little is known about the *Klebsiella* genes contained by non-hypervirulent strains, including multi-drug resistant (MDR) strains, that increase the risk of disease in colonized patients more broadly. Given that most infections are caused by non-hypervirulent strains¹³, more data concerning the genetic determinants of infection are necessary for predicting, diagnosing, and experimentally evaluating these strains.

In general, the ability of bacteria to progress from colonization to infection is dependent on the immune status of the host, environmental and iatrogenic exposures such as antimicrobials, and the virulence potential of the strain. Known Klebsiella virulence determinants are found in both the core genome, which are ~1700 genes present in >95% of all sequenced strains, and the accessory genome, which are genes that vary among sequenced strains¹⁴. The overall Klebsiella accessory genome is currently estimated to include >100,000 unique genes³. With an average of >5000 genes per genome, thousands of genes vary between strains. The size of the Klebsiella accessory genome provides a unique opportunity to perform comparative genomics studies to understand pathogenesis. Comparative genomics approaches are used to understand hospital outbreaks^{15,16}, identify in-host adaptation¹⁷, and infer conserved virulence determinants or pathways. The latter approach has been used to interrogate the pathogenicity of numerous bacterial pathogens, such as *Pseudomonas aeruginosa*¹⁸, *Streptococcus* pneumoniae¹⁹, Escherichia coli²⁰, and invasive non-typhoidal Salmo*nella enterica* spp.²¹. Previously, we used a comparative genomics approach to identify Klebsiella factors associated with infection in a small, hospital-wide cohort of *Klebsiella*-colonized patients²². Two infection-associated factors identified in this study, the psicose utilization locus and ter operon, were experimentally validated and characterized^{22,23}, confirming the value of this approach for understanding Klebsiella pathogenesis.

The strong association between gut colonization and *Klebsiella* disease indicates a potential point of intervention to prevent infection

in the most at-risk patients, such as those in intensive care and hematology/oncology units. If patients with risk factors for infection are colonized by high-risk Klebsiella isolates, early detection and intervention may prevent disease. The development of these approaches is likely to be critical as therapeutic options become limited due to rising antimicrobial resistance. We previously performed a large cohort study of Klebsiella-colonized patients in the intensive care and hematology/oncology units and identified patient variables associated with subsequent infection¹⁰. Additionally, we identified an association between gut dominance by Klebsiella and infection, even after adjustment for the patient variables¹¹. Here, we aimed to rigorously identify Klebsiella genes that are associated with disease in colonized patients through whole-genome sequencing (WGS) in a nested casecontrol study from a cohort of over 1900 colonized patients. This approach differed from other comparative genomics studies in that it focused on concordant infections (e.g., same colonizing and infecting isolate) after infection, directly compared cases of infection to wellmatched controls from the same patient population and incorporated careful adjustment with patient variables associated with infection. Through this approach, we identified 27 genes associated with infection in colonized patients. We identified which genes may act through gut dominance, and determined that many genes were found on large, conjugative plasmids. Finally, we demonstrate that five of these genes are associated with infection in a geographically-independent cohort of intensive care patients colonized with Klebsiella.

Results

Selection of *Klebsiella* for WGS and description of the casecontrol study

A nested case-control study was performed using a cohort of 1978 *Klebsiella* colonized, intensive care, and hematology/oncology patients at a single academic medical center in Michigan, USA¹⁰. Over the course of this study patient rectal swabs were screened for the presence of *Klebsiella*. If a patient was colonized with *Klebsiella*, up to three rectal isolates were banked, and the patient was enrolled in the study. Subsequent rectal swab isolates and clinical culture *Klebsiella* isolates (blood, respiratory, urinary, and other) were also banked (Fig. 1a). Cases of infection were considered for inclusion based on chart review using strict clinical definitions¹⁰. To facilitate a rigorous

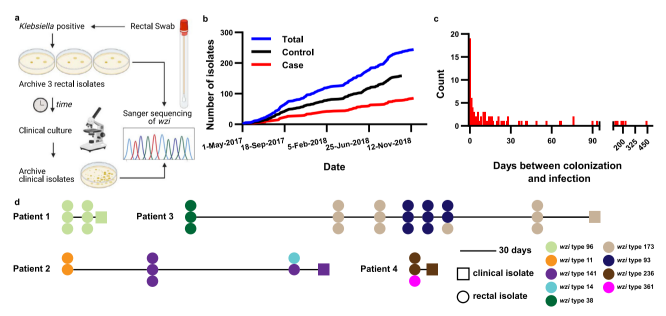


Fig. 1 | **Description of** *Klebsiella* **isolate collection.** a Schematic representation of *Klebsiella* isolate collection and *wzi*-based concordance testing for WGS selection. **b** Rectal isolate selected for WGS collection dates stratified by case status.

c Histogram of the number of days between identification of a rectal *Klebsiella* isolate concordant with the clinical isolate from the same patient. **d** Trace diagrams of *wzi* typing in *Klebsiella* colonizing and infecting isolates in infected patients.

Table 1 | Patient demographics and select baseline characteristics

Patient variable		Case (N = 85)	Control (N = 160)	Odds ratio (OR)*	95% CI	Fisher P-value
Sex	Female	40 (47.1%)	76 (47.5%)			
	Male	45 (52.9%)	81 (50.6%)		,	
	Missing	0 (0.0%)	3 (1.88%)			
Median age (range)		60 (83–25)	59 (85–27)			
Infection site	Blood	39 (45.9%)				
	Respiratory	19 (22.4%)	,			
	Urine	23 (27.1%)	1			
	Other	4 (4.71%)	,			
Mean weighted Elixhauser score (±SD)		22.2 ± 11.6	19.7 ± 11.9	1.02	0.994 to 1.04	0.127
Depression	Yes	30 (35.3%)	44 (27.5%)	1.31	0.761 to 2.26	0.327
	No	55 (64.7%)	113 (70.6%)			
	Missing	0 (0.0%)	3 (1.88%)			
Prior diuretics	Yes	30 (35.3%)	38 (23.8%)	1.69	0.905 to 3.05	0.101
	No	55 (64.7%)	119 (74.4%)			
	Missing	0 (0.0%)	3 (1.88%)			
Prior Vitamin D	Yes	18 (21.2%)	19 (11.8%)	1.66	0.972 to 4.44	0.0591
	No	67 (78.8%)	138 (86.3%)			
	Missing	0 (0.0%)	3 (1.88%)			
Prior pressors/inotrope	Yes	20 (23.5%)	14 (8.75%)	3.06	1.40 to 6.68	0.00502
	No	65 (76.6%)	143 (89.4%)			
	Missing	0 (0.0%)	3 (1.88%)			
High-risk antibiotics	Yes	30 (35.3%)	32 (20.0%)	2.21	1.18 to 4.14	0.013
	No	55 (64.7%)	125 (78.1%)			
	Missing	0 (0.0%)	3 (1.88%)			
Albumin	<2.5 g/dL	35 (41.2%)	34 (21.3%)	2.86	1.49 to 5.48	0.00157
	≥2.5 g/dL	47 (55.3%)	116 (72.5%)			
	Missing	3 (3.53%)	10 (6.25%)			
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*Two-sided test. No adjustments were made for multiple comparisons. OR for sex was incalculable due to use of sex as a matching variable. Bold text indicates significant results (P-value ≤ 0.01 after adjustment for clinical variables).

comparative genomics approach, Sanger sequencing of the wzi gene²⁴ was used to screen for concordance between the clinical culture isolate and a preceding colonizing rectal isolate from the same patient (Fig. 1a). Cases and controls accumulated at a similar rate throughout the enrollment period (Fig. 1b). The median time between detection of Klebsiella colonization and subsequent infection was 10 days (range = 0-443 days, Fig. 1c). In some instances, the progression from colonization to infection was simple, wherein a single wzi type was detected in all colonizing Klebsiella isolates and the subsequent infecting isolate (Fig. 1d, "Patient 1"). In other instances, the infection trajectory was more complicated, wherein some patients exhibited single or multiple strain replacements in the gut prior to infection (Fig. 1d, "Patient 2" and "Patient 3"), or mixed colonization where only one colonizing isolate caused the subsequent infection (Fig. 1d, "Patient 2" and "Patient 4"). Collectively, these data indicate concordant infections are detectable in colonized patients in the context of highly dynamic Klebsiella populations.

In total, 85 cases of infection with an identical *wzi* type to a colonizing rectal isolate (cases) were selected for WGS, including 39 bloodstream infections, 23 urinary tract infections, 19 pneumonias, and 4 other sites of infection (including two cases of cholecystitis, one case of empyema, and one case of ecthyma). Based on antibiotic susceptibility data from medical records, 44 of 85 (51.8%) of the clinical isolates from cases were pan-sensitive apart from intrinsic ampicillin resistance. The most common AMR phenotypes were resistance to first- and second-generation cephalosporins, the combinatorial β -lactam/ β -lactamase-inhibitors ampicillin-sulbactam and piperacillin-

tazobactam, and trimethoprim-sulfamethoxazole (Supplementary Fig. S1). Cases were matched to two colonized controls (who remained asymptomatic) based on sex, age, and date of rectal swab collection. In addition, controls had the same type of clinical culture collected as their corresponding case, but with no cultivatable bacteria. Two strata included only a single control because of the inability to find an appropriate match, and an additional 8 controls were removed from the study following WGS based on species classification outside of *Klebsiella*. In the final case-control design, 85 cases, including the clinical and rectal isolates, and 160 control rectal isolates were evaluated (Table 1).

Description of sequenced Klebsiella isolates

Following the determination of isolates for WGS, genomic DNA was extracted from all *Klebsiella* isolates and subjected to Illumina sequencing (Supplementary Data 1). Assembled genomes were generally of high quality, wherein the median number of contigs was 110.5 (range = 43–876 contigs) and the median N50 was 459,591 bp (range = 147,972–3,285,432 bp). All genomes were of expected length, wherein the median genome size was 5.50 Kb (range = 5.06–6.26 Kb). Sequencing quality was similar between rectal case and control isolates, which were used for most subsequent analyses (Supplementary Fig. S2). As expected, species assignment based on WGS included the three most common members of the *Klebsiella pneumoniae* species complex (*K. pneumoniae*, *K. variicola*, *K. quasipneumoniae*). *K. pneumoniae*, *K. variicola*, and *K. quasipneumoniae* were evenly distributed amongst the four sites of infection and control rectal swabs

(Supplementary Fig. S3a). Case and control rectal isolates within a stratum may be the same or different species, as reflected in either high (100-80%) or low (60-30%) overall nucleotide identity (Supplementary Fig. S3b). For the rectal and clinical isolate of a given case of infection, matching wzi types were highly predictive of overall nucleotide identity (median % nucleotide identity = 99.65%, range = 95.04-99.88%, Supplementary Fig. S3c). The time between detection of Klebsiella colonization and subsequent infection was negatively correlated with sequence similarity; however, the effect was minimal ($r^2 = 0.06$, Supplementary Fig. S3d). Interestingly, instances of lower nucleotide identity similarity (<98%) between the rectal and clinical isolates from concordant cases of infection appeared to be due to large genomic losses or additions (Supplementary Fig. S3e), which may be driven by plasmids or other mobile genetic elements (MGEs). Indeed, these case clinical and rectal isolate pairs displayed differences in the number of annotations that corresponded to differences in genome size (Supplementary Fig. S3f-g).

To confirm concordance between case clinical and rectal isolates, wzi strain typing was performed from the assembled genomes. All wzi types matched results from Sanger sequencing, and construction of a wzi approximately-maximum-likelihood tree revealed a total of 147 unique wzi types identified (Supplementary Fig. S4). Then, multi-locus sequence typing was performed across all isolates. One-hundred thirty-one unique sequence types (STs) were identified, and all case rectal and case clinical STs were identical. Rarefaction of STs indicated that cases and controls were equally rich (Supplementary Fig. S5). ST253 was the most abundant ST but still low frequency (10/245 rectal isolates, 4.08%), and 44 rectal isolates were non-typable (44/245, 17.96%). Overall, hypervirulent clonal groups (CGs)²⁵ were rare, comprising 2.35% of cases and 1.35% of controls. Only two isolates of hypervirulent ST23 were identified, and only one caused an infection. MDR CGs²⁵, frequently containing extendedspectrum β-lactamases, carbapenemases, or both, were more common, representing 18.8% of cases and 9.38% of controls. These included 1 ST11, 4 ST14, 3 ST15, 1 ST16, 3 ST17, 5 ST37, 2 ST147, 1 ST152. 1 ST258, and 3 ST307. 85.7% (210/245) of the isolates in our collection were from non-hypervirulent, non-MDR CGs. Combined, MDR CGs were significantly associated with case status (odds ratio = 2.24, 95% C.I. 1.03 to 4.93, Fisher P = 0.043). Construction of a core genome phylogeny demonstrated the presence of three Klebsiella species within our cohort and multiple lineages across colonizing and invasive isolates of each species (Supplementary Data 1; Fig. 2a). Together, these data indicate that sequenced Klebsiella isolates were highly diverse, without a dominant hypervirulent or MDR CG within the population.

To enable the identification of infection-associated genes in rectal samples, we assembled a pangenome (Supplementary Data 2). The rectal isolate pangenome consisted of 26,089 total annotations. The core genome (genes present in \geq 95% genomes) was comprised of 3921 genes, and the accessory genome (genes present in <95% genomes) consisted of 22,168 genes (Supplementary Fig. S6a). To assess the similarity of the accessory genome between *Klebsiella* isolates in our collection, pairwise Jaccard distances were calculated after excluding the core genome and summarized as a mean for each isolate (244 pairwise comparisons per isolate). Interestingly, the mean pairwise Jaccard distance for all rectal isolates was 0.73 \pm 0.04, indicating that the accessory genome of each isolate was highly dissimilar from other isolates. No differences were observed between mean pairwise Jaccard distances for cases and controls (Supplementary Fig. S6b). Overall, the cohort of *Klebsiella* isolates exhibited high pangenome diversity.

Genes associated with Klebsiella infection

To identify genes in the accessory genome associated with infection we used Scoary, collapsing linked genes that displayed identical presence/absence patterns across all sequenced isolates (e.g., organized

in operons, linked to MGEs, etc.) into units and comparing unit frequencies between cases and controls (Supplementary Data 3) 26 . Thirty-seven genes across 34 units were significantly associated with infection (Q-value \leq 0.2, 2166 tests). Two units contained genes with identical presence/absence patterns ($group_5875$ -tehA_2- $group_10494$ and $group_5455$ -sul2); the remaining units contained a single gene. Overall, 36 genes were associated with cases, one was associated with controls (Table 2).

To control for patient variables in our comparative genomics analysis, we leveraged a previously constructed multivariate model for infection risk in the larger cohort from which this nested case-control study was derived. This model was comprised of mean weighted Elixhauser comorbidity score, depression, prior diuretic use, vitamin D use, pressors/inotrope use, high-risk antibiotic use, and low albumin. Prior medication use was defined as >48 h but <90 days prior to the colonizing swab and albumin was measured within 48 h of the colonizing swab¹⁰. High-risk antibiotics, based on their impact on indigenous gut microbiota, were defined as β-lactam/β-lactamase inhibitor combinations, carbapenems, third- and fourth-generation cephalosporins, fluoroquinolones, clindamycin, and oral vancomycin²⁷. Consistent with the explanatory value of these variables in the overall cohort, the variables used for adjustment trended towards or were significantly associated with infection in the nested case-control study (Table 1). We then used inverse probability of treatment weighting to adjust for these variables in measuring the association of each gene with the outcome of infection.

Following adjustment for clinical variables including comorbidities and antibiotic exposure, 28 of 37 genes remained significantly associated with infection (P-value ≤ 0.01 ; Table 2, Fig. 2b). These 28 infection-associated genes fell into four categories. First, many infection-associated genes encode AMR proteins even after correction for exposure to high-risk antibiotics. These included aac(6')-lb-cr5, encoding an aminoglycoside acetyltransferase that also has activity against fluoroquinolones²⁸, and *bla_{OXA-1}*, *bla_{CTX-M-15}*, bla_{TEM-1} encoding broad-spectrum β -lactamases. Second, several infection-associated genes have predicted protein functions that may alter the physiology of the bacteria directly. These include the predicted aquaporin gene aqpZ_2, tellurite-resistance gene tehA_2, arsenate resistance gene arsD 1, and diguanylate cyclase gene dgcE. Third, certain MGEs and plasmid maintenance genes were associated with infection (e.g., IS1380 family transposase ISEcp1, Transposon Tn3 resolvase, IS6 family transposase IS6100, umuDC_2). The remainder of infection-associated genes encode hypothetical proteins. Interestingly, certain genes were highly associated with one another, suggesting the presence of linked genes that may be acquired simultaneously through horizontal gene transfer (HGT) events (Supplementary Fig. S7).

None of the Klebsiella genes associated with infection encoded known virulence factors. To directly assess the association of known virulence genes with infection, we used the Kleborate genotyping tool, which is an in silico platform for rapid genotyping of Klebsiella genomes (Supplementary Data 4)25. Consistent with the paucity of hypervirulent STs, most well-characterized Klebsiella virulence factors were rare in our dataset, including salmochelin (iro), aerobactin (iuc), and colibactin (clb) synthesis loci, and hypermucoviscosity regulators RmpA/A2 (Table 3). The yersiniabactin (ybt) synthesis locus was more frequent overall, and in cases versus controls, but was not associated with infection (odds ratio = 1.7, P = 0.144). Correspondingly, the virulence score, which is a composite metric of the virulence loci listed above, was not significantly different between cases and controls (Case virulence score = 0.294 ± 0.784 , Control virulence score = 0.163 ± 0.513 , P = 0.115, Student's *t*-test). Even when assessed directly and after controlling for clinical variables, canonical virulence factors are not associated with infection in this group of colonized, intensive care patients.

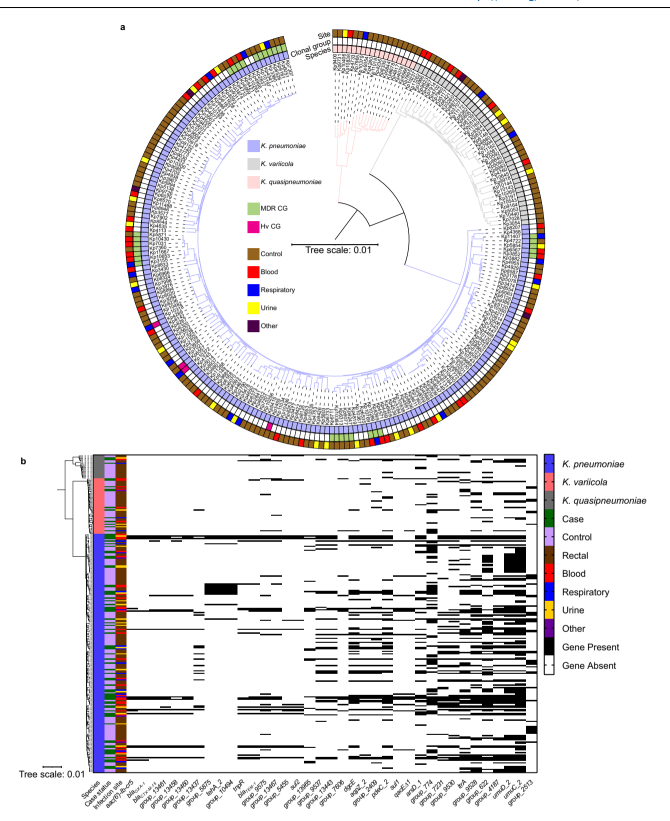


Fig. 2 | *Klebsiella* rectal isolates are highly diverse and infection-associated genes span that diversity. a Core genome phylogeny tree of sequenced *Klebsiella* rectal isolates. The inner ring indicates the species, the middle ring indicates a clonal group of concern (hypervirulent [Hv] CG: clonal groups 23, 25, 380; MDR CG: 14/15, 17/20, 37, 147, 258, 307), and the outer ring indicates site of infection for

concordant clinical isolate. **b** Heatmap of infection-associated gene presence/ absence organized vertically by core genome phylogeny and horizontally by the strength of association (Odds ratio) between gene presence and infection following adjustment for patient variables.

Table 2 | Genes associated with patient case status

Gene	Annotation	Case fre- quency	Control frequency	Unadjusted		Adjusted for clin- ical variables		Mediation				
				Odds ratio	P-value	Q-value**	Odds ratio	P-value**	Direct β	Direct P-value**	Indirect β	Indirect P-value**
aac(6')-lb-cr5	fluoroquinolone-acetylating aminoglycoside 6'-N-acetyltransferase AAC(6')-lb-cr5	0.176	0.006	30.68	0.0011	0.00131	52.41	0.0003	3.0601	0.0045	0.086	0.01
bla _{OXA-1}	oxacillin-hydrolyzing class D β- lactamase OXA-1	0.153	0.006	25.54	0.002	0.0053	49.29	0.0003	2.9698	0.0064	0.077	0.03
bla _{CTX-M-15}	class A extended-spectrum β -lactamase CTX-M-15	0.165	0.013	13.94	0.0007	0.0053	19.17	0.0004	2.3386	0.004	0.047	0.1567
group_13461	IS1380 family transposase ISEcp1	0.165	0.013	13.94	0.0007	0.0053	18.69	0.0003	2.482	0.0022	0.019	0.5833
group_13458	hypothetical protein	0.141	0.013	11.46	0.0018	0.01959	17.57	0.0006	2.0832	0.0112	0.034	0.3483
group_13460	hypothetical protein	0.153	0.013	12.69	0.0011	0.01065	16.73	0.0005	2.3118	0.0047	0.024	0.5217
group_13437	hypothetical protein	0.176	0.031	5.97	0.001	0.05505	7.28	0.0014	1.6137	0.0054	0.005	0.875
group_5875- tehA_2- group_10494*	HTH-type transcriptional regulatory protein GabR - tellurite-resistance protein TehA - hypothetical protein	0.129	0.025	5.65	0.004	0.1908	7.09	0.0029	1.8251	0.0053	0.007	0.8383
tnpR	transposon Tn3 resolvase	0.176	0.031	5.97	0.001	0.05505	6.24	0.0017	1.6379	0.0054	0.023	0.45
bla _{TEM-1}	class A broad-spectrum β-lactamase TEM-1	0.212	0.038	6.28	0.0002	0.01236	5.46	0.0009	1.6623	0.0021	0.035	0.1783
group_9575	tyrosine recombinase XerC	0.259	0.056	5.39	0.0001	0.0053	4.72	0.0019	1.3993	0.0044	0.048	0.05
group_13467	IS6 family transposase IS6100	0.224	0.044	5.75	0.0002	0.01093	4.58	0.0026	1.5338	0.003	0.023	0.4017
group_5455- sul2*	IS91 family transposase - sulfonamide- resistant dihydropteroate synthase Sul2	0.129	0.025	5.07	0.0077	0.1908	4.52	0.0258	2.0214	0.0053	-0.033	0.3617
group_13965	hypothetical protein	0.153	0.038	4.11	0.0066	0.16685	3.59	0.0264	1.0196	0.0806	0.074	0.01
group_9537	hypothetical protein	0.235	0.081	3.71	0.0009	0.13482	3.3	0.0035	1.0141	0.0216	0.049	0.025
group_13443	hypothetical protein	0.224	0.075	3.81	0.001	0.16685	3.23	0.005	1.0762	0.0178	0.06	0.0083
group_7606	hypothetical protein	0.141	0.025	6.26	0.0021	0.10991	3.07	0.0916	1.106	0.0975	0.097	0.0001
dgcE	hypothetical protein	0.341	0.138	3.37	0.0003	0.07663	3.06	0.0016	0.9197	0.0133	0.048	0.0083
aqpZ_2	aquaporin Z	0.247	0.094	3.96	0.0005	0.16685	2.97	0.0085	0.923	0.0385	0.074	0.0001
group_2409	ISL3 family transposase ISEc38	0.259	0.106	3.07	0.002	0.1908	2.67	0.0083	0.803	0.0487	0.05	0.01
pdeC_2	putative cyclic di-GMP phosphodiesterase PdeC	0.365	0.169	2.88	0.0008	0.10991	2.66	0.0026	0.7709	0.0296	0.041	0.015
sul1	sulfonamide-resistant dihydropteroate synthase Sul1	0.141	0.031	4.97	0.0037	0.17478	2.64	0.0939	1.102	0.0994	0.12	0.0001
qacE∆1	quaternary ammonium compound efflux SMR transporter QacE Δ 1	0.153	0.031	5.46	0.0019	0.12853	2.63	0.1011	1.1732	0.0745	0.107	0.0001
arsD_1	arsenical resistance operon trans-acting repressor ArsD	0.318	0.144	3.19	0.0006	0.17478	2.62	0.0043	0.7263	0.0563	0.068	0.0001
group_774	hypothetical protein	0.459	0.25	2.55	0.0013	0.1391	2.55	0.0016	0.8601	0.0088	0.042	0.0033
group_7231	hypothetical protein	0.306	0.138	2.69	0.0032	0.16685	2.54	0.0071	0.7807	0.0373	0.026	0.1733
group_9530	hypothetical protein	0.459	0.238	2.64	0.0009	0.08496	2.38	0.0037	0.8449	0.0094	0.015	0.3533
ltrA	hypothetical protein	0.424	0.225	2.57	0.0013	0.16683	2.33	0.0051	0.7495	0.0229	0.037	0.015
group_9528	hypothetical protein	0.353	0.163	2.87	0.001	0.12853	2.31	0.0134	0.7418	0.0384	0.032	0.0917
group_622	hypothetical protein	0.482	0.269	2.71	0.0006	0.12566	2.24	0.0063	0.9086	0.0049	0.015	0.3683
group_4187	Antirestriction protein KlcA	0.424	0.219	2.54	0.0017	0.12566	2.24	0.0092	0.7747	0.0191	0.021	0.1983
umuD_2	Protein UmuD	0.6	0.369	2.42	0.0016	0.10991	2.1	0.0114	0.8272	0.0098	0.006	0.7
umuC_2	Protein UmuC	0.682	0.456	2.52	0.0012	0.10991	2.09	0.0141	0.8239	0.0107	0.006	0.71
group_2513	hypothetical protein	0.071	0.225	0.26	0.0038	0.16685	0.25	0.0031	-1.2577	0.0119	-0.02	0.3617

^{*}Loci have identical presence/absence patterns across all sequenced isolates and are clustered for analysis.

Next, we aimed to adjust for the effects of phylogeny on the association between each group of genes and infection. We used treeWAS to test for the confounding potential of clonality and recombination in the population structure by simulating a null genetic dataset across the phylogenetic tree²⁹. The output of treeWAS is three complimentary scores: The Terminal score counts the four possible combinations of genotype and phenotype without regard to phylogeny, whereas the Simultaneous and Subsequent scores account for population structure²⁹ in measuring the association between genotype and phenotype. Of the 28 infection-associated genes after adjustment (Table 2), 23 genes were also significant ($P \le 0.01$) by the Terminal score; group_774, group_9530, ltrA, group_622, group_4187, and group_2513 were not (Supplementary Fig. S8, Supplementary Table 1). Other than group 2513, 27 genes were significantly associated with

infection by either the Simultaneous or Subsequent score, or both (Supplementary Fig. S8). For subsequent analyses, we focused on these 27 genes associated with infection both after adjustment for clinical variables and after at least one test of independence from bacterial phylogeny.

AMR genes are associated with infection independent of prior exposure to antibiotics

The identification of multiple AMR genes associated with infection after adjustment for exposure to high-risk antibiotics was unexpected. To investigate the association between AMR genes and infection further, we used two independent approaches. First, we identified classes of AMR genes as defined by Kleborate²⁵ in our strain set. Overall, the prevalence of many of these AMR determinants was low (<25%).

^{**}Bold text indicates significant results (unadjusted Q-value ≤ 0.2, P-value ≤ 0.01) after adjustment for clinical variables, and direct and/or indirect effect P-value ≤ 0.01).

Table 3 | Klebsiella genotype association with infection

Genotype	Case frequency	Control frequency	Unadjusted		Adjusted for clinical variables	
			Odds ratio	P-value	Odds ratio	<i>P</i> -value
Virulence determinants						
ybt	0.207	0.133	1.7	0.144	1.78	0.155
col	0.024	0.007	3.72	0.286	0.629	0.709
iuc	0.024	0.007	3.72	0.286	0.436	0.507
iro	0.012	0.007	1.84	0.668	0.476	0.602
rmpA	0.024	0.007	3.72	0.286	0.436	0.507
rmpA2	0.012	0.007	1.84	0.668	0.937	0.964
AMR determinants*						
AGly acquired (aminoglycosides)	0.268	0.067	5.133	7.01E-05	3.77	0.00427
Flq (fluoroquinolones)	0.220	0.040	6.75	1.14E-04	6.69	0.000507
MLS (macrolides)	0.098	0.013	8.0	0.010	6.41	0.0649
Phe (phenicols)	0.171	0.027	7.51	0.001	6.35	0.00332
Rif (rifampin)	0.012	0.000	N/A	N/A	N/A	N/A
Sul (sulfonamides)	0.268	0.047	7.49	1.22E-05	5.19	0.00226
Tet (tetracyclines)	0.183	0.100	2.01	0.076	1.88	0.128
Tmt (trimethoprim)	0.244	0.053	5.73	8.86E-05	4.28	0.00285
Omp mutation	0.037	0.020	1.86	0.453	1.55	0.676
Bla (β-lactamase that is "other")	0.573	0.520	1.24	0.438	1.27	0.379
Bla Carb (carbapenemase)	0.061	0.007	9.68	0.040	36.2	0.00151
Bla ESBL (extended-spectrum β-lactamases)	0.207	0.087	2.76	0.011	2.94	0.0116
Bla broad (broad-spectrum β-lactamases)	0.354	0.347	1.03	0.915	0.902	0.734
Bla broad inhR (broad-spectrum $\beta\text{-lactamases}$ with resistance to $\beta\text{-lactamase}$ inhibitors)	0.110	0.107	1.03	0.942	1.05	0.922

*Kleborate²⁵ predicted AMR genotypes absent in our dataset are not shown. Bold text indicates significant results (*P*-value ≤ 0.01).

Consistent with results from Scoary, AMR determinants for aminoglycosides, fluoroquinolones, phenicols, sulfonamides, and trimethoprim were significantly associated with infection after adjustment. In addition, carbapenamase genes were also significantly associated with infection, and extended-spectrum β-lactamases (ESBL) approached significance (P = 0.011; Table 3). Notably, one case rectal isolate (Kp10372, Supplementary Data 4) displayed a hypervirulent (ST23, ybt⁺, col⁺, iuc⁺, rmpA/A2⁺), MDR (KPC-3) genotype. The Kleborate Resistance Score²⁵ was not significantly different between cases and controls (Case resistance score = 0.2 ± 0.483 , Control resistance score = 0.163 ± 0.418 , P = 0.528, Student's *t*-test); however, this score is heavily weighted by the presence of carbapenemases in the context of colistin resistance, the latter of which was not detected in our dataset²⁵. Next, we assessed phenotypic resistance across four antibiotic classes, using an agar dilution method with antibiotic concentrations at the Clinical & Laboratory Standards Institute breakpoint for resistance³⁰. Screen results were confirmed for 12 randomly selected Klebsiella isolates using the broth microdilution method. Overall, phenotypic AMR rates were low (<25%). Phenotypic resistance to chloramphenicol and gentamicin did not correlate well to genotypic resistance and was not associated with infection (Supplementary Table 2). Conversely, phenotypic resistance to ciprofloxacin, cefazolin, cefuroxime, and cefepime was associated with infection following adjustment for clinical variables (Supplementary Table 2).

Given the consistent association of AMR genes and phenotypes despite adjustment for exposure to high-risk antibiotics as a group, we evaluated whether their association with infection is due to exposure to specific cognate antibiotics. For example, *aac(6)-lb-cr5* may be associated with exposure to aminoglycosides, a class that was not included in adjustment. Interestingly, prior antibiotic exposure by specific classes was not associated with the presence of most infection-

associated genes in unadjusted analysis (Supplementary Fig. S7). Next, we explicitly adjusted for each class of antibiotic, in addition to clinical variables by IPTW, and assessed the association of each gene with infection. Even after individual adjustment for 18 classes of antibiotics (no patients had a prior exposure to daptomycin or polymyxin), all AMR genes remained significantly associated with infection (Supplementary Fig. S9). These data suggest an alternative explanation exists for the association between infection and AMR genes.

Gut relative abundance mediates association between certain genes and *Klebsiella* infection

We previously identified that high Klebsiella relative abundance in the gut is associated with infection¹¹, and that genes associated with infection can enhance fitness in the gut²³. Therefore, we hypothesized that the Klebsiella genes associated with infection may act by enabling gut dominance that in turn increases the risk of infection. Klebsiella relative abundance measurements from the rectal swabs most proximal to infection for cases and matched controls were available for 233 of 245 sequenced rectal isolates¹¹. Reanalysis of these data was consistent with that of the original study, wherein cases had significantly higher Klebsiella relative abundance in the gut compared to controls (Fig. 3a). Mean relative abundance was not significantly different between species (Supplementary Fig. S10a). Abundance was higher in patients exposed to high-risk antibiotics, and when stratified by antibiotics exposure the abundance in cases was significantly higher than controls (Supplementary Fig. S10b, c). Analysis of *Klebsiella* gut relative abundance based on gene presence/absence (rather than case status) identified 11 of the 27 infection-associated genes associated with increased relative abundance (Fig. 3b), whereas the other 16 infection-associated genes were not associated with increased relative abundance (Supplementary Fig. S10d).

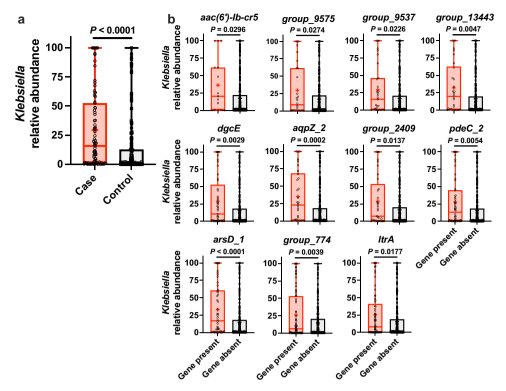


Fig. 3 | Specific infection-associated genes are associated with increased *Klebsiella* gut relative abundance. a Comparison of *Klebsiella* gut relative abundance in patient rectal swabs between cases (n = 83) and controls (n = 149, mean displayed, ****P-value ≤ 0.00005 , two-sided Student's t-test) and **b** in patient rectal swabs stratified by gene presence/absence (25% percentile and 75% percentile bound the box, median displayed as line, mean displayed as plus sign, and range displayed as whiskers, *P-value ≤ 0.005 , ***P-value ≤ 0.0005 , two-sided Student's t-test). Each data point represents a

single patient rectal swab. Source data are provided as a Source Data file. Sample sizes for aac(6)-lb-cr5 present n=15, aac(6)-lb-cr5 absent n=217, $group_9575$ present n=28, $group_9575$ absent n=204, $group_9537$ present n=31, $group_9537$ absent n=201, $group_13443$ present n=30, $group_13443$ absent n=202, dgcE present n=49, dgcE absent n=183, $aqpZ_2$ present n=34, $aqpZ_2$ absent n=198, $group_2409$ present n=37, $group_2409$ absent n=195, $pdeC_2$ present n=54, $pdeC_2$ absent n=178, $pdeC_2$ absent n=178, $pdeC_2$ absent p=178, $pdeC_2$ absent p=184, $pdeC_2$ absent $pdeC_$

To test the hypothesis that infection-associated genes may act through gut dominance we undertook a mediation analysis, a statistical approach that estimates the strength of both the direct associations between each gene and the outcome of infection and indirect association through gut relative abundance. For seven gene units, the association with infection was mediated, at least in part, by gut abundance (Table 2; indirect P-value, indicated in bold). The AMR gene aac(6)-lb-cr5 had both a strong direct association with infection and an indirect mediation by abundance (*P*-value \leq 0.01). The hypothetical protein group_774 also had both significant direct and indirect effects. The remaining five genes (hypothetical protein group_13443, dgcE, aqpZ_2, transposase group_2409, and arsD_1) had only significant indirect effects, indicating that their association with infection is mediated by gut relative abundance. Collectively, these findings begin to suggest biological roles for Klebsiella genes associated with infection, in which some genes may act through high gut relative abundance, but others likely have separate functions.

Most infection-associated genes are carried on plasmids

Given the diversity of functions and frequencies of the infectionassociated genes, we sought to determine the genetic context of these genes as part of the accessory genome. Based on nucleotide homology, many of these genes have been previously detected on plasmids^{31–33}. Consistent with plasmid carriage, certain infectionassociated genes, including both AMR and non-AMR genes, are highly associated with one another (Supplementary Fig. S7) and appear to be linked (Fig. 2b). Furthermore, we detected the loss or gain of groups of infection-associated genes between some case clinical and rectal pairs, rather than individual genes (strata 211, 214, 701, 713, 716, Supplementary Fig. S3f). To determine the association between plasmids and infection-associated genes, we used the PlasmidFinder tool to detect replicons in the rectal isolates in our dataset³⁴. We detected a replicon in 163 of 245 isolates (66.5%, 72.94% of case rectal isolates, 63.13% of control isolates), with a median of 2 replicons per isolate (Supplementary Data 5, range = 0–10 replicons). There was no significant difference between the number of replicons per isolate between cases and controls (Supplementary Fig. S11a). The three most common replicons were Col(pHAD28), IncFIB(K), and IncFII(K), and these replicons were distributed across the core genome phylogeny (Supplementary Fig. S11b); however, no replicon was significantly associated with infection (Supplementary Table 3).

To determine whether infection-associated genes are located on plasmids or the chromosome, we performed long-read sequencing and hybrid assembly with existing short read data on 10 isolates from cases of infection (Figs. 4a-c, S12). All of these isolates contained plasmids with infection-associated genes, and no two plasmids containing infection-associated genes were identical in terms of size, organization and gene content. At one extreme, the rectal case isolate Kp11407 contained a 208,703 bp hybrid IncFIB(K)/IncFII(K) plasmid encoding all 18 infection-associated loci present in this isolate (Fig. 4a). This plasmid closely aligns to the conjugative plasmid pUUH239.2 (CP002474.1), which was linked to a nosocomial K. pneumoniae outbreak³⁵. Notably, this plasmid also contains several heavy metal resistance operons, including ars, pco, and sil³⁵, which may enhance resistance to innate immune responses³⁶. A second case rectal isolate, Kp5854, encoded 23 of 27 infection-associated loci on a 247,093 bp hybrid IncFIB(K)/IncFII(K) plasmid (Fig. 4b). This plasmid closely aligns to pF16KP0070-1 (CP052586.1), which also encodes ars, pco, and sil.

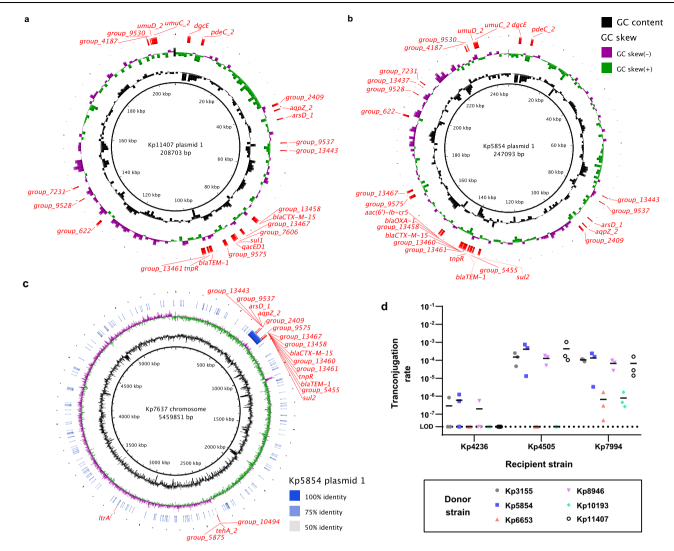


Fig. 4 | Infection-associated genes are present on large, conjugative plasmids. a Plasmid map of Kp11407 plasmid 1. b Plasmid map of Kp5854 plasmid 1. c Alignment of Kp7637 chromosome and Kp5854 plasmid 1. d Transconjugation rates using Kp3155, Kp5854, Kp6653, Kp8946, Kp10193, and Kp11407 as donor

strains to Kp4236, Kp4505, and Kp7994, which are *K. quasipneumoniae*, *K. pneumoniae*, and *K. variicola*, respectively, as recipient strains (mean displayed, N.D. = none detected). For **d**, each data point represents a single biological replicate. Source data are provided as a Source Data file.

Interestingly, in the corresponding clinical isolate, this plasmid appears to have integrated into the chromosome with the loss of some infection-associated genes (Fig. 4c; stratum 713, Supplementary Fig. S3g). These isolates also contained four additional infection-associated genes that mapped to the chromosome. The number of plasmid-encoded, infection-associated genes varied widely between plasmids (Supplementary Fig. S12). In contrast to Kp11407 and Kp5854, Kp11162 contained one plasmid with two infection-associated genes and a second plasmid with four (Supplementary Fig. S12g, h). Across these isolates 23 of all 27 infection-associated genes were found on at least one plasmid (Supplementary Fig. S12). Only *group_774* and the *tehA_2* gene cluster were not found on at least one plasmid. This suggests that these large plasmids facilitate acquisition of infection-associated loci and are likely drivers of the association between many genes and infection.

Infection-associated genes are transmissible by plasmids

To determine if infection-associated genes could be horizontally transmitted, we performed transconjugation assays between rectal isolates. Kp3155, Kp5854, Kp6653, Kp8946, Kp10193, and Kp11407 were selected as donor strains, as these *K. pneumoniae* case rectal isolates contained various infection-associated genes, including

 $bla_{\text{CTX-M-15}}$ and dgcE, as well as the IncFIB(K) replicon. Three cefuroxime-sensitive control isolates lacking these genes, *K. pneumoniae* (Kp4505), *K. variicola* (Kp7994), and *K. quasipneumoniae* (Kp4236), were selected as recipient strains. Kp7994 and Kp4236 lack plasmid replicons, and Kp4505 contains an IncFIB(K) replicon. Conjugation to Kp4236 was limited, with inconsistent or lack of transfer across donor strains. In contrast, conjugation was readily detected from donors Kp3155, Kp5854, Kp8946, Kp11407 to recipient Kp4505, and from all donors to Kp7994 (Fig. 4d). All transconjugants were cefuroxime resistant, indicating the transfer of $bla_{\text{CTX-M-15}}$, and were PCR-positive for dgcE. Therefore, we concluded that these large plasmids are conjugative, resulting in the transfer of infection-associated genes.

Validation of infection-associated genes in a geographically distinct patient cohort

Finally, we aimed to validate our findings across multiple geographical locations. To this end, we screened a cohort of *Klebsiella* colonized patients from a single academic medical center in Maryland, USA for six infection-associated genes: aac(6)-lb-cr5, $bla_{CTX-M-15}$, $group_13461$, $group_13467$, dgcE, and $aqpZ_2$. These genes were selected because they were representative of the many facets of our findings,

Table 4 | Gene association with positive clinical cultures in a geographically independent cohort of *Klebsiella* colonized patients

Gene	Sample	Frequency	Odds ratio	95% CI	<i>P</i> -value
aac(6')-lb-cr5	Case clinical	0.167	4.88	1.96 to 11.1	0.0009
	Case rectal	0.115	3.18	1.22 to 8.41	0.0276
	Control rectal	0.039			
dgcE	Case clinical	0.481	2.28	1.29 to 3.99	0.0074
	Case rectal	0.5	2.46	1.37 to 4.41	0.0039
	Control rectal	0.289			
group_13461	Case clinical	0.204	4.55	2.06 to 9.6	0.0005
	Case rectal	0.192	4.23	1.93 to 9.24	0.0042
	Control rectal	0.053			
group_13467	Case clinical	0.296	3.81	1.94 to 7.29	0.0002
	Case rectal	0.25	3.02	1.48 to 6.1	0.0043
	Control rectal	0.1			
bla _{CTX-M-15}	Case clinical	0.185	4.94	2.19 to 11.4	0.0005
	Case rectal	0.173	4.55	1.83 to 10.9	0.0013
	Control rectal	0.044			
aqpZ_2	Case clinical	0.296	1.58	0.83 to 2.93	0.164
	Case rectal	0.327	1.82	0.98 to 3.38	0.076
	Control rectal	0.213			

Bold text indicates significant results (P-value ≤ 0.05).

encapsulating known AMR genes (aac(6')-lb-cr5, bla_{CTX-M-I5}), MGEs (group_13461, group_13467), and those where the relationship between gene presence and infection was (aac(6')-lb-cr5, dgcE, aqpZ_2) and was not (bla_{CTX-M-15}, group 13461, group 13467) significantly mediated by Klebsiella gut relative abundance. Moreover, these genes are found on conjugative plasmids (Fig. 4). This second cohort consisted of 55 cases of Klebsiella infection and 432 asymptomatically colonized control patients. Since wzi concordance data was not available, we screened both case isolates and a rectal isolate from each patient for each gene. Rectal isolates were not available for three infected patients and a clinical isolate was not available for one patient. In total, 538 Klebsiella isolates (54 clinical isolates, 52 case rectal isolates, 432 control rectal isolates) were screened for the presence or absence of target genes using a multiplex RT-PCR assay. As an internal control for Klebsiella detection, we included a previously validated assay for the fiu gene¹¹. Based on either clinical or rectal isolates from cases, we observed a significant association with infection for all genes except aqpZ_2 (Table 4). Overall, 82.4% (42/51 of available pairs) of clinical-rectal pairs returned concordant RT-PCR results, which was highly consistent with our previous study using wzi sequencing to determine concordance¹⁰. As in the primary cohort (Supplementary Fig. S7), the presence of these genes in case rectal isolates were associated with one another (Supplementary Fig. S13), suggesting that these genes in the Maryland isolates may also be linked through conjugative plasmids. The strongest associations were between dgcE and apqZ 2 and between aac(6)-Ib-cr5, bla_{CTX-M-15}, group_13461 and group_13467. Overall, five of six genes were reproducibly associated with case status in colonized patients in two geographically-distinct cohorts.

Discussion

In this study, we have identified novel *Klebsiella* genes associated with infection in colonized patients. The findings from this diverse collection of *Klebsiella* isolates, carefully adjusted for patient risk factors and phylogeny, and validated in a second geographically distinct cohort, provides important insights into the broader dynamics of *Klebsiella* pathogenesis. Most of these genes were found across a diverse set of plasmids, suggesting that these genes are either direct mediators of fitness and virulence or indirect markers of large, conjugative plasmids

that increase infection risk as a unit. These genes can be separated into two categories: those whose association with infection is mediated through increased abundance in the gut, and those that have a direct association with infection. Genes whose effect on infection risk is not mediated through intestinal dominance may act at a subsequent step of pathogenesis. These genes associated with infection are readily mobilized within and across members of the *Klebsiella pneumoniae* species complex, and changes in infection-associated gene content and genetic context were detectable within colonized patients who progressed to infection. Collectively, this study provides a set of genetic signatures of infection that have potential for predicting risk of infection in patients colonized by diverse *Klebsiella*.

The majority of infection-associated loci in our dataset are present on plasmids, which can vary in size, gene content, copy number, and conjugative ability (Fig. 4a-c, Supplementary Fig. S12). This plasmid carriage creates genetic linkage between genes that are associated with infection, raising the possibility that some are direct effectors of fitness or virulence whereas others are robust markers for these effectors. In some instances, clear relationships exist between linked genes. For example, nucleotide cyclases are often present in tandem with cyclic nucleotide phosphodiesterases, thus the linkage of dgcE and pdeC 2 is expected. dgcE and pdeC 2 may regulate expression of type 1 and 3 pili through modulation of cyclic di-GMP levels. Expression of the type 1 pilus reduces virulence in the lung and spleen but enhances gut colonization in murine models^{37,38}. Therefore, dgcE may enhance fitness in the gut while pdeC 2 may enhance fitness in extraintestinal sites. Likewise, the linkage of $bla_{CTX-M-15}$ and $group_13461$ is expected since ISEcp1 transposases are known to mobilize and promote the expression of CTX-M-type β -lactamases^{39,40}. Conversely, relationships between other linked genes, such as arsD_1 and the various AMR genes, are more difficult to understand. It may be the case that these infection-associated genes act at different stages of infection or one or both are robust genetic markers for fitness genes elsewhere on the plasmid.

The strong association between multiple AMR genes and infection following adjustment for high-risk antibiotic exposure indicates that the relationship between antibiotic resistance and infection is complex in *Klebsiella*. Antibiotic resistance genes could be associated with

infection by enabling intestinal outgrowth and dominance after exposure to a cognate antibiotic. However, the ESBL determinants bla_{CTX-M-15}, bla_{OXA-1}, and bla_{TEM-1} were associated with infection independent of antibiotic exposure and their association was not significantly mediated by gut relative abundance. Notably, antibiotic exposure rates are low (<10% for most classes) in our original study population¹⁰. Therefore, detection of these antibiotic resistance genes is more likely a marker of other infection-associated genes and/or plasmids, rather than conferring resistance to a cognate antibiotic. It may be the case that these strains acquired these genes earlier in their evolutionary history; however, prospective comparative genomics studies are necessary to pinpoint such events. Potential fitness benefits of AMR genes in the absence of antibiotics are intriguing, yet highly AMR *Klebsiella* STs are generally less virulent in experimental models than their hypervirulent counterparts, and specific mutations that confer AMR can have deleterious effects on Klebsiella fitness⁴¹⁻⁴⁴. On the other hand, plasmid acquisition of AMR is less costly than chromosomal mutations⁴⁵. Therefore, AMR genes may be linked to plasmidencoded genes that enhance infection risk, and then complicate treatment of infections after they are initiated.

The identification of non-AMR genes associated with infection as mediated by gut dominance adds to our understanding of bacterial genotypes and the risk for infection in Klebsiella colonized patients. We previously demonstrated that the ter operon, annotated as a tellurite-resistance operon, is a microbiome-dependent gut fitness factor²³. Similarly, the arsD₁ gene identified in this study is part of an arsenic resistance (ars) operon. Tellurite and arsenic are both toxic metalloids, implying that the ars operon may be playing an analogous role to ter. Consistent with this, gut relative abundance mediates the association between arsD_1 and infection (Table 2). Conversely, we did not find evidence that gut relative abundance mediates the association between the tellurite-resistance gene tehA 2 gene and infection, although it is predicted to encode a different molecular function than ter. These data highlight the complex interplay between bacterial genotype and gut fitness, and the need to understand the role of colonization fitness to both assess infection risk and prophylactic decolonization therapies. The risk of Klebsiella disease in colonized patients may not warrant prophylactic antibiotic administration, especially in the context of rising rates of AMR; however, nonantibiotic options are currently being explored (reviewed in⁴⁶). A better understanding of how bacterial pathogens, such as Klebsiella, successfully colonize and dominate the gut is needed to ensure the efficacy of any decolonization approach.

Although this study advances our understanding of infection in patients colonized by diverse Klebsiella, it is not without its limitations. First, this study is retrospective and relies on the availability of clinical data in the electronic medical record. Prospective studies with longer timeframes aimed at evaluating the association between genes identified in this study and infection would be useful for validating our findings. Second, many infection-associated genes are associated with one another and these genes are plasmid-borne. This finding is important: however, it complicates the evaluation of the causal relationship between individual genes and infection. Experimental studies are necessary to determine the causal relationship between each gene and disease. Additionally, we did not comprehensively assess the genomic architecture and HGT potential of our strains. It may be the case that only a subset of genes is frequently transferred between strains and/or only a subset of strains is able to easily donate and receive these genes. Notably, all plasmid donor strains that we tested were K. pneumoniae that encoded an IncFIB(K) replicon (Supplementary Data 5), therefore limiting our ability to assess the universality of the HGT phenotype. Further assessment of the *Klebsiella* plasmidome will be critical to fully interpret the role of the infection-associated genes identified in this study, as well as the characterization of cryptic plasmids that could be missed with replicon-based analysis such as PlasmidFinder. Moreover, more complicated approaches such as shotgun metagenomics are necessary to contextualize plasmid borne genes in polymicrobial reservoirs of bacteria with pathogenic potential, such as the gut. In our combined approach across three species, the majority of our genomes were from K. pneumoniae. There may be K. variicola or K. auasipneumoniae-specific infection-associated genes that we were unable to detect. Third, the Klebsiella genes associated with infection likely differ by patient population. We found different genes associated with infection (ter, psicose utilization locus) in a hospital-wide comparative genomics study and experimentally validated them as encoding fitness factors^{22,23}. That patient population may have different risk factors and stronger barriers to infection. Among intensive care and hematology/oncology patients, the genes associated with infection were highly reproducible in a geographically independent cohort. Finally, this study focused on gene presence in the accessory genome and does not consider the role of mutations within the core or accessory genomes.

This study provides insights into infection risk from diverse lineages of the K. pneumoniae species complex. The well-characterized virulence genes of hypervirulent clonal groups, many of which are plasmid-borne²⁵, were rare and not informative in assessing infection risk in this population. MDR isolates were more frequent representing six clonal groups, but overall, 85.7% (210/245) of the isolates in our collection were from non-hypervirulent, non-MDR CGs. Given the predominance of non-hypervirulent, or "classical" strains worldwide²⁵, efforts to understand the pathogenesis and enact preventative therapies is likely to have a substantial payoff. The varied combinations of infection-associated genes found across case isolates suggests that a diverse accessory genome acquired through HGT drives infection risk in Klebsiella. Certain combinations have become fixed; some hypervirulent Klebsiella clones have carried their plasmids for decades. However, new combinations can arise within a colonized patient (Fig. 4; Supplementary Fig. S12). Even in seemingly fixed plasmid-strain combinations, exchange has occurred leading to multiple instances of the convergence of virulence and MDR determinants²⁵. In total, this study adds to the growing understanding of the Klebsiella accessory genome as a critical determinant of infection risk.

The insights from this study can be used to predict infection risk in colonized, hospitalized patients and trigger infection prevention interventions. We have previously identified certain patient risk factors associated with infection, and that gut dominance is associated with infection independently of these clinical factors^{10,11}. Here we identified infection-associated genes independent of patient risk factors and that part of the association is mediated by gut dominance. We have also developed the tools to measure these risks, including a robust qPCR assay to measure intestinal dominance and the genotyping PCR assays implemented in this study. The fact that AMR genes and phenotypes are strongly associated with infection could also enable the use of standard culture approaches to detect these genes, and therefore, high-risk colonizing strains. It is possible that these approaches can be combined with gueries of the electronic medical record for clinical variables to generate an integrated risk score for a colonized patient. In the near term, this score could be used to identify patients for trials of decolonization therapies. Eventually, integrated risk models could identify at-risk patients who could most benefit from effective prevention interventions.

Methods

Ethics statement

Patient enrollment and sample collection at the University of Michigan was approved by and performed in accordance with the Institutional Review Boards (IRB) of the University of Michigan Medical School (Study number HUM00123033). Patient enrollment and sample collection at Johns Hopkins University was approved and performed in accordance with the IRB of the Johns Hopkins

University (Study number IRB00129775). Patients were retrospectively enrolled after routine samples were taken in the course of clinical care. This study was performed with a waiver of informed consent since the research involves no more than minimal risk to the subjects, could not practicably be carried out without the waiver, and uses discarded samples. Patient enrollment was identical at both sites.

Sample collection and strain selection

At Michigan Medicine, Klebsiella isolates were selected from a large cohort of Klebsiella colonized patients. Cohort identification, enrollment, clinical data extraction, chart review, and case definitions are described in detail elsewhere¹⁰. During the study period, patient rectal swabs were screened for the presence of Klebsiella by plating on MacConkey agar followed by taxonomic identification using MALDI TOF mass spectrometry. Up to three Klebsiella isolates per rectal swab, and any subsequent Klebsiella positive clinical cultures, were archived. Minimum inhibitory concentration data from case clinical isolates was extracted from the Michigan Medicine clinical microbiology lab. To identify concordant infections. Klebsiella rectal swab and case clinical isolates were subjected to wzi PCR and Sanger sequencing, and the wzi type was assigned by uploading the consensus wzi sequence to the BigsDB database (https://bigsdb.pasteur.fr). Each case was matched to two controls based on sex, age (±10 years), and date of rectal swab collection (±90 days). Several cases could only be matched to a control swab within 120 days (n=6). Case matching was conducted in R, version 4.0.3.

At the Johns Hopkins University, adult patients admitted to medical or surgical ICUs are routinely screened for VRE rectal colonization on admission and weekly thereafter until ICU discharge. Residual Amies broth from rectal ESwabs (COPAN Diagnostics, Inc., Murrieta, California) were inoculated on a MacConkey agar plate using the WASP DT (COPAN) automated specimen processer, and isolates were archived as above.

Whole-genome sequencing, assembly, and annotation

Archived Klebsiella isolates were cultured in Luria-Bertani (LB, Becton, Dickinson and Company, Franklin Lakes, NJ) broth at 37 °C with shaking for genomic DNA extraction. Genomic DNA was extracted using the DNeasy UltraClean Microbial Kit (Qiagen, Hilden, Germany) per manufacturer's instructions. Extracted genomic DNA was then checked for quality and purity and submitted to the University of Michigan Advanced Genomics Core, where it was sequenced on an Illumina NovaSeq using a 300 cycle S4 flow cell (Illumina, San Diego, USA) to generate approximately 6,000,000 150 bp paired-end reads. Resulting reads were trimmed using Trimmomatic v0.39⁴⁷ assembled into scaffolds using SPAdes v3.13.0⁴⁸. Genome assemblies within each stratum were compared using QUAST v5.0.249, wherein each genome assembly was compared to the case rectal isolate genome assembly. Genome annotation was performed using Prokka v1.14.6⁵⁰, and a pangenome was assembled using Roary v3.13.0⁵¹, with settings "-i 90" and "-s." Finally, assembly genotyping was performed using Kleborate v1.0.0²⁵ and PlasmidFinder v2.0.1³⁴. For long-read sequencing, genomic DNA was checked for quality and purity and submitted to the University of Michigan Advanced Genomics Core, where it was sequenced on a GridION X5 platform using a MinION flow cell (Oxford Nanopore Technology, Oxford, UK). Low-quality read ends were trimmed with NanoFilt v2.5.052 and assembled using Unicycler v0.4.8 using the "bold" setting⁵³. The core gene alignment was created using Illumina assemblies with cognac using the default parameters⁵⁴. The *wzi* gene sequences from each Illumina assemblies were identified by BLAST v2.9.0⁵⁵. The BLAST output was parsed to extract the aligned sequence from each query, and the gene sequences were aligned with MAFFT v7.310. Approximate maximum likelihood trees were generated from the wzi alignment and cognac alignment with fastTree v2.1.10⁵⁶. Phylogenetic trees were visualized with the APE R package v5.3⁵⁷. Rarefaction was performed using iNEXT v2.0.20⁵⁸, with setting "q = 0." All analyses were performed in R v4.0.5. Plasmid maps and corresponding alignments were constructed and visualized using BRIG v0.95⁵⁹. All code developed for this manuscript is available at https://github.com/rdcrawford/Bachman_CU8.

Phenotypic antibiotic screen

To screen for phenotypic antibiotic resistance, all *Klebsiella* rectal isolates were arrayed into 96-well plates and stored at –80 °C in 20% LB-glycerol. We were unable to locate one isolate in our freezer following WGS, thus it was omitted from phenotypic analysis. Then *Klebsiella* rectal isolates were sub-cultured in LB broth at 37 °C overnight and replica plated on antibiotic containing LB-agar. Antibiotic concentrations (36 mg/mL ampicillin, 32 mg/mL chloramphenicol, 16 mg/mL gentamicin, 1 mg/mL ciprofloxacin, 32 mg/mL cefazolin, 32 mg/mL cefuroxime, 16 mg/mL cefepime) were selected in accordance with CLSI criteria for resistance. All assays were performed in triplicate.

Transconjugation assay

Single colonies of Kp4236, Kp4505, Kp7994 (recipient strains; chloramphenicol resistant), and Kp3155, Kp5854, Kp6653, Kp8946, Kp10193, Kp11407 (donor strains; cefuroxime resistant) were subcultured overnight in 1 mL of LB broth at 37 °C with shaking. The next day, overnight cultures were harvested by centrifugation and resuspended in 1 mL sterile phosphate-buffered saline (PBS). 50 µL of Kp4236, Kp4505, or Kp7994 resuspension was mixed with 50 μL of each donor strain resuspension and spotted on LB-agar. 50 µL of unmixed resuspension of each strain was spotted in parallel. Bacterial spots were grown overnight at 37 °C, then harvested into 1 mL sterile PBS by scraping and serially plated on differential growth media. Transconjugation rates were calculated by quantifying the bacterial density of dual-resistant colonies divided by the total density of the recipient strain. Transconjugates were confirmed by qPCR targeting the dcgE gene using unmixed strains as a reference (assay details below). All assays were performed in triplicate, and 3 transconjugate colonies and 3 unmixed recipient and donor colonies were selected per assay for PCR confirmation.

RT-PCR design and assay

To screen for the presence of aac(6)-lb-cr5, bla_{CTX-M-15}, group_13461, group_13467, dgcE, and aqpZ_2, multiplex real-time PCR assays were designed using PanelPlex (DNA Software, Ann Arbor, USA) and combined with a previously validated Klebsiella-specific fiu assay11. All primer and probe sequences can be found in Supplementary Table 4. Clinical and rectal isolates were arrayed and sub-cultured in 150 µL LB overnight at 37 °C. To extract genomic DNA, overnight cultures were harvested by centrifugation and resuspended in 100 µL of PCR-grade water, boiled at 95 °C for 5 min, and pelleted. Then, 5 µL of the supernatant was used as a PCR template. Primer (Integrated DNA Technologies, Coralville, USA) and probe (Thermo Fisher Scientific, Waltham, USA) concentrations were 400 nM and 200 nM, respectively¹¹. Real-time PCR was performed using a QuantStudio 3 real-time thermocycler (Thermo Fisher Scientific, Waltham, USA) using the following protocol: 50 °C for 2 min, 95 °C for 15 min, then 40 cycles of 94 °C for 1 min and 60 °C for 1 min. All assays were performed in duplicate unless replicates conflicted, wherein a third replicate was performed, and all isolate arrays included 3 positive controls and 3 randomly dispersed negative controls. The target gene was assigned as present in an isolate if a reaction was (1) positive for fiu, (2) amplified within 2 Ct of the range of the positive control Cts, and (3) results were concordant between all replicates.

Statistical analysis

For clinical variable comparisons, odds ratios and associated 95% confidence intervals were calculated using conditional logistic

regression in R (version 3.6.3) and a *P*-value \leq 0.05 was considered statically significant. For infection-associated gene discovery, a *P*-value \leq 0.05 and *Q*-value \leq 0.2 was used as a cutoff for further exploration. For these analyses, Scoary v1.6.16²⁶ was used with the "-collapse" and "-n" settings with our custom core genome phylogeny, limited to genes that were 5–95% frequent in the dataset. For analyses considering population structure, treeWAS v1.1²⁹ was used with the phen.type = "discrete" setting and our custom core genome phylogeny. For gene-gene associations, odds ratios were calculated using Fishers exact test in R Studio v1.2.5001. To account for the inflated type I error rate due to the large number of comparisons, a *P*-value \leq 0.01 was considered statistically significant for subsequent clinical variable adjustment, mediation, population structure, and genotyping analyses.

Next, we considered adjustment of clinical variables. Inverse probability of treatment weighting method was used for this purpose⁶⁰. The analysis is performed separately for each gene. In the propensity score model, a logistic regression was used with the clinical variables as predictors and binary gene presence/absence as the outcome. The predicted probabilities estimated from the regression model were used to compute weights and then included in the outcome model. The outcome model is a logistic regression with Klebsiella infection as the outcome and gene (presence/absence) as the predictor, while incorporating the matched nature of the data. Following IPTW analysis, we assessed if each gene was associated with infection after individually adjusting for exposure to each class of antibiotics. We repeated the described IPTW analysis using the same weights, controlling for each prior exposure as a covariate in the outcome model one at a time. The "survey" package (v4.1-1) in R was used to obtain the robust standard error.

We then conducted a mediation analysis on each gene that was significantly associated with infection to explore if the relationship between gene presence/absence and infection is mediated through abundance. Specifically, we assessed (1) the direct effect defined as the effect of gene presence on infection, adjusting for abundance as a continuous variable and clinical factors. (2) the indirect effect of the gene on infection mediated through abundance after adjusting clinical factors. To account for the matched case-control design, a generalized linear mixed model was used with a random effect for the strata defining matched case-controls. A significant mediation (i.e., indirect) effect indicates evidence that the effect of the gene on infection is mediated through abundance, and direct effects were also reported. The P-values for these effects were assessed to identify genes to investigate further. We conducted the analyses using the 'mediation' (v4.5.0) and 'lme4' packages (v1.1-29)^{61,62} in R v3.6.3.

For in vitro experiments, one-sample t-test, unpaired Student's t-tests, and ANOVA followed by indicated post-hoc test was used to determine significant differences between groups following log transformation of experimental values. For in vitro experiments, a P-value ≤ 0.05 was considered statistically significant. Experimental replicates represent biological replicates. Statistical testing was two-sided in all circumstances. Statistical analysis of in vitro experiments was performed using Prism 8 (GraphPad Software, La Jolla, CA).

Reporting summary

Further information on research design is available in the Nature Research Reporting Summary linked to this article.

Data availability

The sequencing data generated in this study have been deposited in the Sequence Read Archive (SRA) database under accession code PRJNA789565. The human subjects data are available under restricted access for the protection of the rights and welfare of human research subjects. Deidentified human data are available under restricted access

and can be obtained from MAB within 1 year upon request, pending approval from the University of Michigan Institutional Review Board. The remaining data generated in this study are provided in the Supplementary Information/Source Data file. Source data are provided with this paper.

Code availability

All code developed for this manuscript is available at https://github.com/rdcrawford/Bachman CU8.

References

- Long, S. W. et al. Whole-genome sequencing of a human clinical isolate of the novel species *Klebsiella quasivariicola* sp. nov. *Genome Announc*. https://doi.org/10.1128/genomeA.01057-17 (2017).
- Rodrigues, C. et al. Description of Klebsiella africanensis sp. nov., Klebsiella variicola subsp. tropicalensis subsp. nov. and Klebsiella variicola subsp. variicola subsp. nov. Res. Microbiol. 170, 165–170 (2019).
- Wyres, K. L., Lam, M. M. C. & Holt, K. E. Population genomics of Klebsiella pneumoniae. Nat. Rev. Microbiol. 18, 344–359 (2020).
- Magill, S. S. et al. Multistate point-prevalence survey of health care-associated infections. N. Engl. J. Med. 370, 1198–1208 (2014).
- CDC. The Direct Medical Costs of Healthcare-Associated Infections in U.S. Hospitals and the Benefits of Prevention (Centers for Disease Control & Prevention, 2009).
- O'Neill. Tackling Drug-Resistant Infections Globally: Final Report And Recommendations. https://amr-review.org (2016).
- Martin, R. M. et al. Molecular epidemiology of colonizing and infecting isolates of *Klebsiella pneumoniae*. mSphere 1, e00261–16 (2016).
- Collingwood, A. et al. Epidemiological and microbiome associations between Klebsiella pneumoniae and vancomycin-resistant enterococcus colonization in intensive care unit patients. Open forum Infect. Dis. 7, ofaa012 (2020).
- Gorrie, C. L. et al. Gastrointestinal carriage is a major reservoir of Klebsiella pneumoniae Infection in intensive care patients. Clin. Infect. Dis.: Off. Publ. Infect. Dis. Soc. Am. 65, 208–215 (2017).
- Rao, K. et al. Risk Factors for Klebsiella infections among hospitalized patients with preexisting colonization. mSphere https://doi. org/10.1128/mSphere.00132-21 (2021).
- Sun, Y. et al. Measurement of Klebsiella intestinal colonization density to assess infection risk. mSphere https://doi.org/10.1128/ mSphere.00500-21 (2021).
- Russo, T. A. & Marr, C. M. Hypervirulent Klebsiella pneumoniae. Clin. Microbiol. Rev. https://doi.org/10.1128/CMR.00001-19 (2019).
- Wyres, K. L. et al. Genomic surveillance for hypervirulence and multi-drug resistance in invasive Klebsiella pneumoniae from South and Southeast Asia. Genome Med. 12, 11 (2020).
- Martin, R. M. & Bachman, M. A. Colonization, infection, and the accessory genome of Klebsiella pneumoniae. Front. Cell. Infect. Microbiol. 8, 4 (2018).
- Snitkin, E. S. et al. Tracking a hospital outbreak of carbapenemresistant Klebsiella pneumoniae with whole-genome sequencing. Sci. Transl. Med. 4, 148ra116 (2012).
- Long, S. W. et al. Population genomic analysis of 1,777 extendedspectrum beta-lactamase-producing *Klebsiella pneumoniae* isolates, houston, texas: unexpected abundance of clonal group 307. mBio. https://doi.org/10.1128/mBio.00489-17 (2017).
- Marvig, R. L., Sommer, L. M., Molin, S. & Johansen, H. K. Convergent evolution and adaptation of *Pseudomonas aeruginosa* within patients with cystic fibrosis. *Nat. Genet.* 47, 57–64 (2015).
- Allen, J. P. et al. A comparative genomics approach identifies contact-dependent growth inhibition as a virulence determinant. *Proc. Natl Acad. Sci. USA* 117, 6811–6821 (2020).

- Aggarwal, S. D. et al. Function of BriC peptide in the pneumococcal competence and virulence portfolio. PLoS Pathog. 14, e1007328 (2018).
- Mageiros, L. et al. Genome evolution and the emergence of pathogenicity in avian Escherichia coli. Nat. Commun. 12, 765 (2021).
- Bornstein, K. et al. Genetic changes associated with the temporal shift in invasive non-typhoidal Salmonella serovars in Bamako Mali. PLoS Negl. Trop. Dis. 13, e0007297 (2019).
- Martin, R. M. et al. Identification of pathogenicity-associated Loci in Klebsiella pneumoniae from hospitalized patients. mSystems https://doi.org/10.1128/mSystems.00015-18 (2018).
- Vornhagen, J. et al. A plasmid locus associated with Klebsiella clinical infections encodes a microbiome-dependent gut fitness factor. PLoS Pathog. 17, e1009537 (2021).
- Brisse, S. et al. wzi Gene sequencing, a rapid method for determination of capsular type for Klebsiella strains. J. Clin. Microbiol. 51, 4073–4078 (2013).
- Lam, M. M. C. et al. A genomic surveillance framework and genotyping tool for Klebsiella pneumoniae and its related species complex. Nat. Commun. 12, 4188 (2021).
- Brynildsrud, O., Bohlin, J., Scheffer, L. & Eldholm, V. Rapid scoring of genes in microbial pan-genome-wide association studies with Scoary. Genome Biol. 17, 238 (2016).
- Baggs, J. et al. Risk of subsequent sepsis within 90 Days after a hospital stay by type of antibiotic exposure. Clin. Infect. Dis.: Off. Publ. Infect. Dis. Soc. Am. 66, 1004–1012 (2018).
- Robicsek, A. et al. Fluoroquinolone-modifying enzyme: a new adaptation of a common aminoglycoside acetyltransferase. *Nat. Med.* 12, 83–88 (2006).
- Collins, C. & Didelot, X. A phylogenetic method to perform genome-wide association studies in microbes that accounts for population structure and recombination. PLoS Comput. Biol. 14, e1005958 (2018).
- CLSI. Methods for Dilution Antimicrobial Susceptibility Tests fo Bacteria That Grow Aerobically. 10th edn (Clinical and Laboratory Standards Institute, 2015).
- Alcock, B. P. et al. CARD 2020: antibiotic resistome surveillance with the comprehensive antibiotic resistance database. *Nucleic Acids Res.* 48, D517–D525 (2020).
- Wu, J. & Rosen, B. P. The arsD gene encodes a second trans-acting regulatory protein of the plasmid-encoded arsenical resistance operon. Mol. Microbiol. 8, 615–623 (1993).
- Serfiotis-Mitsa, D. et al. The structure of the KlcA and ArdB proteins reveals a novel fold and antirestriction activity against Type I DNA restriction systems in vivo but not in vitro. *Nucleic acids Res.* 38, 1723–1737 (2010).
- 34. Carattoli, A. & Hasman, H. PlasmidFinder and in Silico pMLST: identification and typing of plasmid replicons in wholegenome sequencing (WGS). *Methods Mol. Biol.* **2075**, 285–294 (2020).
- Sandegren, L., Linkevicius, M., Lytsy, B., Melhus, A. & Andersson, D.

 Transfer of an Escherichia coli ST131 multiresistance cassette has created a Klebsiella pneumoniae-specific plasmid associated with a major nosocomial outbreak. J. Antimicrob. Chemother. 67, 74–83 (2012).
- Djoko, K. Y., Ong, C. L., Walker, M. J. & McEwan, A. G. The role of copper and zinc toxicity in innate immune defense against bacterial pathogens. J. Biol. Chem. 290, 18954–18961 (2015).
- Khater, F. et al. In silico analysis of usher encoding genes in Klebsiella pneumoniae and characterization of their role in adhesion and colonization. PLoS One 10, e0116215 (2015).
- Rosen, D. A., Twentyman, J. & Hunstad, D. A. High levels of cyclic Di-GMP in Klebsiella pneumoniae Attenuate virulence in the lung. Infect. Immun. https://doi.org/10.1128/IAI.00647-17 (2018).

- Bonnet, R. Growing group of extended-spectrum beta-lactamases: the CTX-M enzymes. Antimicrob. Agents Chemother. 48, 1–14 (2004).
- Poirel, L., Decousser, J. W. & Nordmann, P. Insertion sequence ISEcp1B is involved in expression and mobilization of a bla(CTX-M) beta-lactamase gene. Antimicrobial agents Chemother. 47, 2938–2945 (2003).
- 41. Bruchmann, S., Feltwell, T., Parkhill, J. & Short, F. L. Identifying virulence determinants of multidrug-resistant *Klebsiella pneumoniae* in *Galleria mellonella*. *Pathog. Dis.* https://doi.org/10.1093/femspd/ftab009 (2021).
- Xiong, H. et al. Distinct contributions of neutrophils and CCR2+ monocytes to pulmonary clearance of different Klebsiella pneumoniae strains. Infect. Immun. 83, 3418–3427 (2015).
- Olonisakin, T. F. et al. CD36 provides host protection against Klebsiella pneumoniae intrapulmonary infection by enhancing lipopolysaccharide responsiveness and macrophage phagocytosis. J. Infect. Dis. 214, 1865–1875 (2016).
- Wong, J. L. C. et al. OmpK36-mediated Carbapenem resistance attenuates ST258 Klebsiella pneumoniae in vivo. Nat. Commun. 10, 3957 (2019).
- Vogwill, T. & MacLean, R. C. The genetic basis of the fitness costs of antimicrobial resistance: a meta-analysis approach. *Evol. Appl.* 8, 284–295 (2015).
- Gargiullo, L., Del Chierico, F., D'Argenio, P. & Putignani, L. Gut microbiota modulation for multidrug-resistant organism decolonization: present and future perspectives. Front. Microbiol. 10, 1704 (2019).
- Bolger, A. M., Lohse, M. & Usadel, B. Trimmomatic: a flexible trimmer for Illumina sequence data. *Bioinformatics* 30, 2114–2120 (2014).
- Bankevich, A. et al. SPAdes: a new genome assembly algorithm and its applications to single-cell sequencing. J. Comput. Biol. 19, 455–477 (2012).
- Gurevich, A., Saveliev, V., Vyahhi, N. & Tesler, G. QUAST: quality assessment tool for genome assemblies. *Bioinformatics* 29, 1072–1075 (2013).
- 50. Seemann, T. Prokka: rapid prokaryotic genome annotation. *Bioinformatics* **30**, 2068–2069 (2014).
- Page, A. J. et al. Roary: rapid large-scale prokaryote pan genome analysis. *Bioinformatics* 31, 3691–3693 (2015).
- De Coster, W., D'Hert, S., Schultz, D. T., Cruts, M. & Van Broeckhoven, C. NanoPack: visualizing and processing long-read sequencing data. *Bioinformatics* 34, 2666–2669 (2018).
- Wick, R. R., Judd, L. M., Gorrie, C. L. & Holt, K. E. Unicycler: Resolving bacterial genome assemblies from short and long sequencing reads. *PLoS Comput. Biol.* 13, e1005595 (2017).
- 54. Crawford, R. D. & Snitkin, E. S. cognac: rapid generation of concatenated gene alignments for phylogenetic inference from large, bacterial whole genome sequencing datasets. *BMC Bioinforma*. **22**, 70 (2021).
- 55. Altschul, S. F., Gish, W., Miller, W., Myers, E. W. & Lipman, D. J. Basic local alignment search tool. *J. Mol. Biol.* **215**, 403–410 (1990).
- Price, M. N., Dehal, P. S. & Arkin, A. P. FastTree: computing large minimum evolution trees with profiles instead of a distance matrix. *Mol. Biol. Evol.* 26, 1641–1650 (2009).
- Paradis, E., Claude, J. & Strimmer, K. APE: Analyses of phylogenetics and evolution in R language. *Bioinformatics* 20, 289–290 (2004).
- Hsieh, T. C., Ma, K. H. & Chao, A. iNEXT: an R package for rarefaction and extrapolation of species diversity (Hill numbers). Methods Ecol. Evol. 7, 1451–1456 (2016).
- Alikhan, N. F., Petty, N. K., Ben Zakour, N. L. & Beatson, S. A. BLAST Ring Image Generator (BRIG): simple prokaryote genome comparisons. *BMC Genom.* 12, 402 (2011).

- Haukoos, J. S. & Lewis, R. J. The propensity score. *Jama* 314, 1637–1638 (2015).
- Tingley, D., Yamamoto, T., Hirose, K., Keele, L. & Imai, K. mediation: R package for causal mediation analysis. J. Stat. Softw. https://doi. org/10.18637/jss.v059.i05 (2014).
- 62. Bates, D., Mächler, M., Bolker, B. & Walker, S. Fitting linear mixed-effects models using lme4. *J. Stat. Softw.* **67**, 48 (2015).

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Competing interests

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Additional information

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