Population genomics of hypervirulent *Klebsiella pneumoniae* clonal-group 23 reveals early emergence and rapid global dissemination

Margaret M. C. Lam¹, Kelly L. Wyres¹, Sebastian Duchêne¹, Ryan R. Wick¹, Louise M. Judd¹, Yunn-Hwen Gan², Chu-Han Hoh², Sophia Archuleta³, James S. Molton³, Shirin Kalimuddin⁴, Tse Hsien Koh⁵, Virginie Passet⁶, Sylvain Brisse⁶ & Kathryn E. Holt¹,7

Severe liver abscess infections caused by hypervirulent clonal-group CG23 *Klebsiella pneumoniae* have been increasingly reported since the mid-1980s. Strains typically possess several virulence factors including an integrative, conjugative element ICE*Kp* encoding the siderophore yersiniabactin and genotoxin colibactin. Here we investigate CG23’s evolutionary history, showing several deep-branching sublineages associated with distinct ICE*Kp* acquisitions. Over 80% of liver abscess isolates belong to sublineage CG23-I, which emerged in ~1928 following acquisition of ICE*Kp10* (encoding yersiniabactin and colibactin), and then disseminated globally within the human population. CG23-I’s distinguishing feature is the colibactin synthesis locus, which reportedly promotes gut colonisation and metastatic infection in murine models. These data show circulation of CG23 *K. pneumoniae* decades before the liver abscess epidemic was first recognised, and provide a framework for future epidemiological and experimental studies of hypervirulent *K. pneumoniae*. To support such studies we present an open access, completely sequenced CG23-I human liver abscess isolate, SGH10.
Klebsiella pneumoniae (Kp) is a ubiquitous bacterium and important cause of multidrug-resistant healthcare-associated infections. The past three decades have also seen the emergence of severe community-acquired hypervirulent Kp disease, usually manifesting as pyogenic liver abscess with accompanying bacteraemia, but also meningitis, brain abscess or ophthalmitis.\(^1,2\) Earliest reports of this syndrome emerged across parts of Asia including Taiwan, China, Hong-Kong, Singapore and South Korea and more recently from Europe, the United States, South America, the Middle East, and Australia.\(^3,4\)

The majority of Kp liver abscess isolates belong to a small number of clonal groups (CGs) \(^5,6\) defined by multi-locus sequence typing (MLST).\(^6\) In particular, CG23 was shown to account for 37–64% isolates in Taiwan,\(^6\) Singapore\(^7\) and mainland China,\(^8,9\) and includes sequence types ST23, ST26, ST57 and ST163.\(^7\) CG23 Kp are associated with the highly serum-resistant K1 capsule and a number of virulence factors; the genotoxin colibactin, microcin E492, and the iron-scavenging siderophores aerobactin, yersiniabactin and salmochelin.\(^1,9,10\) The CG23 sublineage I, CG23-I has become globally distributed and includes sequence types ST23, ST26, ST57 and ST163.\(^3,7\) CG23-I was separated from the rest of CG23 by 49 SNPs, including one nonsense mutation (Supplementary Data 2). The nonsense mutation is predicted to truncate the outer membrane usher domain of KpcC, likely preventing expression of the Kpc 

Two CG23 comparative genomic analyses have been published, incorporating up to 27 isolates.\(^15,16\) Both studies reported limited nucleotide variation in core chromosomal genes and high conservation of the acquired virulence genes. These include the K1 capsule synthesis locus KKL, the chromosomally encoded yersiniabactin locus (ybt), plus the iro (salmochelin), iuc (aerobactin) and rmpA/rmpA2 genes (upregulators of capsule expression) located on the pK2044 virulence plasmid.\(^17\) The ybt locus is mobilised by the ICEKp integrative conjugative element,\(^18\) for which we have recently described over a dozen variants in the wider Kp population.\(^14\) Variants harbouring ybt, plus iro (ICEKpI) or the colibactin locus (icl, ICEKp10) have been reported in CG23.\(^14,18,19\) ICEKpI was the first ICE to be described in Kp,\(^18\) originating from the sequence type (ST) 23 liver abscess strain NTUH-K2044, which was also the first CG23 strain to be completely sequenced.\(^17\) However ICEKp10 is more common and was observed in all other CG23 genomes investigated in Struve et al.\(^16\)

CG23 has not generally been associated with acquired antimicrobial resistance (AMR), but the last few years have seen increasing reports of resistant strains, including those resistant to third generation cephalosporins and carbapenems.\(^15,20–22\)

Phylogenetic analyses indicated that CG23 has a number of sublineages separated by deep-branches, one of which (labelled CG23 sublineage I, CG23-I) has become globally distributed (Fig. 1, Supplementary Fig. 1). The CG23-I clade was strongly supported by two independent analysis methods (>99% posterior support in Bayesian tree, 100% bootstrap support in ML tree). This clade comprised 81 isolates (83.5% of all CG23) collected from Asia, Australia, North America, Europe and Africa (Fig. 1), including 82% of all liver abscess strains (Supplementary Data 1). Neither of the two oldest strains in our analysis, M109 (1932, Murray Collection)\(^23\) and NCTC9494 (1954, human sputum), were part of CG23-I (Fig. 1); nor was NTUH-K2044, the first sequenced ST23 strain that has served as a reference for much of the reported experimental and genomic work on ST23.\(^16–18\)

CG23-I was separated from the rest of CG23 by 49 SNPs, including one nonsense mutation (Supplementary Data 2). The nonsense mutation is predicted to truncate the outer membrane usher domain of KpcC, likely preventing expression of the Kpc efflux pumps, the AmpC beta-lactamase, the PmrB sensor kinase, four membrane transport secretion proteins, an acid shock protein and the cardiolipin synthase 2 protein (Supplementary Data 2).

Kp is considered an important cause of sexually transmitted disease in horses\(^25\) but little is known about the molecular epidemiology of this group or sources of infection. The horse isolates included in this study (including genital tract, sperm, foetus and metritis specimens isolated between 1980 and 2004) were originally selected for sequencing only on the basis of K1 serotype, but clustered together within a monophyletic subclade nested within CG23-I (Fig. 1), separated from the rest of CG23-I by 83 SNPs (Supplementary Data 3). Of note, four non-synonymous mutations arose within genes encoding putative oxidoreductases and two intergenic mutations were within close proximity to predicted oxidoreductase genes.

The nested positioning within CG23-I suggests that CG23 Kp may have entered the horse population on a single occasion where it now circulates via sexual contact, and there is no evidence of transmission between humans and horses.

We detected a strong temporal signal in the CG23 genome alignment (see Supplementary Fig. 2a–c, Supplementary Methods), sufficient to estimate evolutionary rates and dates for the most recent common ancestors (MRCA) of key CG23 lineages with BEAST (best-fitting Bayesian model was the UCLD constant model, see Supplementary Methods). The mean evolutionary rate for CG23 was estimated to be \(3.40 \times 10^{-7}\) substitutions site\(^{-1}\) year\(^{-1}\) (95% HPD; \(2.43 \times 10^{-7}-4.38 \times 10^{-7}\)). The MRCA for the entire CG23 population, CG23-I and equine sublineage nested within CG23-I were estimated to be 1878 (95% HPD; 1961–1915), 1928 (95% HPD; 1908–1953) and 1972 (95% HPD, 1961–1975), respectively. We hypothesised that the successful spread of CG23-I in the human population may have been associated with a population expansion; i.e. an increase in diversification rate and effective population size, compared to the rest of the CG23 population. We used an epidemiological model (birth-death) to compare these dynamics between the different lineages (see Supplementary Methods), which demonstrated population decline in the horse lineage and expansion of the entire human-associated CG23 population (Supplementary Fig. 3a, c), particularly among CG23-I, which was associated with five times the rate of expansion than the rest of the population (Supplementary Fig. 3b, d). Importantly, there was a population decline in the horse lineage and expansion of the entire human-associated CG23 population (Supplementary Fig. 3a, c), particularly among CG23-I, which was associated with five times the rate of expansion than the rest of the population (Supplementary Fig. 3b, d). Importantly, there was a
significant overlap in the Bayesian parameter estimate distributions from four alternative models, and with estimates derived from an alternative method known as least-squares LSD analysis (Supplementary Fig. 2d-f, see Supplementary Methods). This provides confidence that CG23 emerged some time in the late 19th century and the globally distributed CG23-I emerged in the 1920s.

Isolate CAS686 is a hybrid of ST23 and ST281-like Kp. The genome of isolate CAS686 (ST260, a double locus variant of ST23) was previously shown to contain a large recombination import spanning approximately half of the genome (2.4Mbp)\(^\text{16}\). Here, we identified the likely hybridisation partner to be a ST281-related Kp. We used the RedDog mapping and variant calling pipeline to compare the genome of CAS686 to those of CG23 and other genomes in our curated collection\(^\text{14,26}\). The distribution of SNPs differentiating CAS686 from other CG23 genomes confirms the previous observation that approximately half of the CAS686 genome is closely related to ST23 (Supplementary Fig. 4a). However, the region of similarity appears to be split into two, separated by regions of similarity to the genomes of QMP M1-975 (isolated from asymptomatic human gut carriage\(^\text{9}\)) and INF211 (isolated from a case of pneumonia in an Australian hospital\(^\text{27}\)), both of which are double locus variants of ST281 (Supplementary Fig. 4b).

A subsequent Bayesian phylogenetic analysis showed that the ST23-like regions of CAS686 cluster within CG23-I (models and parameters as described for the analysis presented above, all CG23 genomes were included and the recombination regions in CAS686 were masked). However, it is not possible to know whether CAS686 represents a ST23 strain that has acquired two large regions from an ST281-related strain, or vice versa. Note though that CAS686, but neither QMP M1-975 nor INF211, harbours the virulence plasmid encoding iro, iuc and rmpA/rmpA2 loci that is characteristic of CG23, indicating that import into CG23 is most parsimonious.

Virulence gene content variation in CG23. To investigate the ongoing evolution of virulence in CG23, we performed detailed analyses of variation in virulence gene content within the
population. All CG23 genomes (including the hybrid strain CAS686) carried the K1 locus associated with biosynthesis of the K1 capsule serotype. The K-locus of the 1954 isolate NCTC9494 harboured a 1065 bp insertion between the wzi and wza genes, which contained an IS102-like transposase; insertions in K-loci are common amongst historical isolates\(^{(26,27)}\) and likely arise during long-term storage.

The ybt locus was present in 90 CG23 genomes (93%). Comparison of ybt lineages, ICEKp structures and chromosomal integration sites indicated at least six independent ybt + ICEKp acquisitions (triangles in Fig. 1). CG23-I was characterised by the presence of ICEKp10 encoding ybt lineage 1 (ybt-1) and clb lineage 2 A sequence variants, integrated at tRNA-Asn site 1, consistent with a single integration event in the ancestor of the sublineage. One isolate from this group lacked the ybt genes (Fig. 1) but harboured the left and right ends of ICEKp10 (Supplementary Fig. 5), consistent with this genome sharing the historical acquisition of ICEKp10 followed by subsequent loss of ybt and the ICEKp mobilisation machinery.

At least five unrelated ICEKp integration events were identified outside of CG23-I (Fig. 1), none of which included clb. Three separate introductions of ICEKp3 were detected: ED23 (from Taiwan) carried ybt-9 integrated at tRNA-Asn site 3, three isolates from Australia carried ybt-9 variant integrated at tRNA-Asn site 4, and two related Singapore isolates carried ybt-8 integrated at tRNA-Asn site 1. ICEKp1 encoding ybt-2 was identified in two strains from Taiwan (including the NTUH-K2044 reference strain) and one strain from France. These strains were monophyletic in the tree (grey shaded clade in Fig. 1) and share a MRCA close to the root, however the chromosomal locations of ICEKp1 suggest they may have been acquired via distinct integration events in each strain: at site 1 in NTUH-K2044 and site 4 in SB3926 (the integration site could not be resolved for the French isolate SB4446).

The virulence plasmid-associated loci iuc, iro, rmpA and rmpA2 were present in the vast majority of CG23 genomes (see Fig. 1). Comparison of genome sequences to the reference sequence of virulence plasmid pK2044 (strain NTUH-K2044) confirmed that the plasmid backbone was present in 94 genomes, including all those carrying iuc, iro, rmpA and/or rmpA2, although some plasmids harboured deletions that affected these virulence loci (see Fig. 1, Supplementary Fig. 6a; note it is unclear whether the three plasmid-negative isolates were actually lacking the plasmid in vivo or had lost it during laboratory culture). Other virulence loci detected in CG23 included microcin E492 (only three deletion variants detected, see Supplementary Fig. 7) and allantoinase, both of which were found in the chromosome.

The microcin E492 ICE was present in 89 of the CG23 genomes (92%, Fig. 1), integrated at tRNA-Asn site 2 in all cases (except four genomes in which the site could not be resolved from the available short-read data, see Supplementary Data 1). In the genomes lacking microcin, the tRNA-Asn 2 region was intact and showed no other integration at this site. Sequence homology and structure of the microcin locus was generally conserved, with no more than four SNPs between pairs of microcin ICE sequences. However three strains carried putative deletion variants (Supplementary Fig. 7), consistent with three independent deletion events (grey in Fig. 1 heatmap). The chromosomal allantoinase locus was present in all CG23 genomes, consistent with previous reports\(^{(15,16)}\).

**Antimicrobial resistance determinants in CG23.** All CG23 genomes carried the intrinsic beta-lactamase gene bla\(_{ESBL}\) (which confers ampicillin resistance) and the oxacillinase genes (which confer reduced susceptibility to quinolones). However no fluoroquinolone resistance-associated mutations were identified in gyrA or parC, and acquired AMR determinants were rare. The equine clade and three other unrelated CG23-I isolated from humans (CAS813, BG130 and BG141; * in Fig. 1) carried multiple AMR genes.

The equine clade was associated with addA1 and aph3’-1a (aminoglycosides), sul1 (sulphonamides) and tetAR (tetracycline), carried in an IncFI plasmid backbone (see Supplementary Fig. 8). We could not resolve complete resistance regions from all the short-read assemblies, but we were able to extract a putative plasmid sequence from the assembly graph of isolate SB4816, which was subsequently resolved into the complete plasmid sequence pBSB4816 using long read sequencing (deposited under GenBank accession MF363048, see Methods). The pBSB4816 sequence comprised an IncFII type plasmid backbone harbouring a Tn21-like element that carried a class I integron with addA1 in the gene cassette and succ downstream of the pacEdelta1 fragment (similar to GenBank accession: AP071413\(^{(28)}\), see Supplementary Fig. 8a). The Tn21-like element was interrupted by IS26 sequence inserted in the tnpR and tniA genes. The plasmid backbone showed close similarity (99% nucleotide identity) to plasmid pEC_L8 from a clinical E. coli isolate\(^{(29)}\). Mapping of reads to the plasmid sequence from SB4816 confirmed that a variant of the plasmid was present in all of the AMR equine clades genomes and absent from the rest of the CG23 population (Supplementary Fig. 6). Where aph3’-1a or tetAR were present they were also located in sequences flanked by IS26 (see Supplementary Fig. 8b), hence we propose that variation in AMR gene content within the equine clade is due to IS26-mediated mobilisation or deletion of these AMR genes.

The multidrug-resistant strains isolated from humans each harboured unique sets of AMR genes conferring resistance to multiple classes of drugs, as previously reported\(^{(15,16)}\) (see Supplementary Data 1). While precise locations of the AMR genes were not resolvable in the public draft genome assemblies, all three were found to harbour large conjugative plasmids that are frequently associated with multidrug resistance in Enterobacteriaceae: IncA/C\(_2\) plasmid sequence type (PST) 3 in CAS813 (Denmark, 2008) and BG130 (Vietnam, 2008); IncN PST6 in BG141 (Madagascar, 2007).

**Other gene content variation in CG23.** We compared the seven completely assembled CG23 chromosomes, comprising SGH10 plus five clinical Kp isolates from Taiwan and China collected between 2002 and 2015 (NTUH-K2044; \(^{(17)}\) 1084; \(^{(30)}\) RFJ999; ED2; ED23) and NCTC9494 from the Public Health England National Culture Type Collection (PHE NCTC, collected in 1954 and recently sequenced as part of the NCTC3000 genomes project, see http://www.sanger.ac.uk/resources/downloads/bacteria/nctc/). The analysis revealed remarkable conservation of gene content and synteny (Fig. 2). Relative to the majority consensus, chromosomal inversions of \(~1.45\) Mbp and \(~472\) kbp regions were observed in strains ED2 and RFJ999, respectively (Fig. 2). The boundaries of the RFJ999 inversion were flanked by copies of the transposase ISKpn1. Phage sequences flanked the inverted region in ED2. Variations in gene content were mainly due to integrations of prophage, ICEKp variants and the microcin ICE.

Pan-genome analysis of the full set of 97 genomes identified 4170 core genes (present in \(\geq95\%\) of genomes) and 5493 accessory genes (see Supplementary Fig. 9). Nineteen coding sequences were uniquely present in CG23-I and conserved within this lineage (\(\geq95\%\) CG23-I genomes). Eighteen of these were part of the colibactin synthesis locus clb (detailed below) and the other was an ISKpn1 transposase inserted within an ethanamine transporter gene, eat (KPI1165 in NTUH-K2044).
To estimate the extent to which phage account for differences in gene content, we submitted all 97 assemblies for PHAST\textsuperscript{31} analysis. The number of complete phage sequences detected (listed in Supplementary Data 1) ranged from 0 to 3 per genome (mean = 1.4), and the sizes of these phage varied from 20 kbp to >100 kbp. These phage sequences account for 1238 (12.8\%) of the coding sequences amongst CG23 genomes. The median number of spacers per genome was 22 (range 11–55 amongst strains with intact cas loci), with a median of seven spacers shared with NTUH-K2044, suggesting the CRISPR/Cas system is also active in the majority of CG23 Kp, which may explain the relatively low level of horizontal gene transfer (including plasmid burden\textsuperscript{34} and recombination\textsuperscript{35}) detected in this clone compared to other clonal groups such as CG258\textsuperscript{36}.

**SGH10 as a novel reference strain for hypervirulent CG23.** Our data on CG23 population structure and virulence loci show that none of the currently available finished genome sequences represent a ‘typical’ genome (Figs. 1 and 2b). NTUH-K2044 (liver abscess), ED23 (blood) and NCTC 9494 (sputum) are not part of the predominant CG23-I sublineage and do not harbour the colibactin locus (labelled A–C in Fig. 1; note that NTUH-K2044 also carries additional chromosomal copies of iro and mmpA within ICEKp1 that are absent from the majority of CG23). Strains 1084 (liver abscess) and ED2 (blood) belong to CG23-I but lack the virulence plasmid, and RJF999 is a blood isolate of undetermined virulence. We therefore propose strain SGH10 as a reference strain for experimental and genomic studies of hypervirulent CG23 *K. pneumoniae* associated with human liver abscess because it belongs to the predominant CG23-I sublineage and is responsible for the majority of CG23-associated liver abscess infections, and it has all of the common virulence loci intact with no atypical accessory genes (see Fig. 1 and Supplementary Fig. 9). SGH10 was isolated from a 35-year-old liver abscess patient with...
no diabetes and no detectable underlying disease, in Singapore in 2014. We have previously demonstrated SGH10 to be hyper-mucoid and highly resistant to human serum7. Here we confirmed its virulence potential in the murine model via oral infection of C57BL/6 mice, which resulted in translocation to the liver, lungs and spleen 48 h post-inoculation (see Methods and Fig. 3a). We performed additional long and short-read sequencing of the SGH10 genome (see Methods), yielding complete circular sequences for the chromosome (5485,114 bp, 57.43% GC content) (Fig. 3b) and virulence plasmid (pSGH10; 231,583 bp, 50.15% GC) (Fig. 3c). The annotated genome sequence was deposited in GenBank under accessions CP025080 (chromosome) and CP025081 (virulence plasmid), and the strain submitted to the National Collection of Type Cultures in the United Kingdom (NCTC number 14052).

Discussion
The earlier comparative genomic studies of up to 27 CG2315,16 included only four genomes outside of CG23-I, which somewhat obscured the diversity and population structure of the clonal group, and provided insufficient temporal signal to reconstruct evolutionary dynamics. Here, the expanded genome collection revealed that the majority of human clinical isolates of CG23 belong to a clonally expanded sublineage, CG23-I, which emerged in the early 20th century (Fig. 1). This predates the identification of serotype K1 Kp as a significant cause of liver abscess by over 50 years7,38, suggesting it has been circulating undetected for many decades. We also found evidence that the entire population of CG23 associated with human infections has been expanding since its emergence, but that the CG23-I sublineage has undergone particularly accelerated population growth associated with rapid dissemination to five of the six inhabited continents (Supplementary Fig. 3).

Our genomic analyses suggest some potential mechanisms for the success of the CG23-I sublineage in the human population. The colibactin synthesis locus clb (also referred to as pks), encoded downstream of the ybt locus in ICEKp10, is its most notable feature. Colibactin is a hybrid nonribosomal peptide-polypeptide that has a genotoxic effect on host cells by cross-linking DNA and inducing double-strand DNA breaks19,39,40. It was first discovered in E. coli40 but has since been reported in 3.5–4% of K. pneumoniae14,41, in which it has also been shown to induce DNA double-strand breaks in HeLa cells41. Genotoxicity of colibactin in ST23 Kp has been demonstrated for strain 1084 (which belongs to CG23-I, but lacks the virulence plasmid-encoded iro, iuc and rmpA/rmpA2 loci, see Figs. 1 and 2b), both in vitro in mouse liver cells and in vivo in liver parenchymal cells of orally infected BALB/c mice19. In these experiments, the genotoxic effect of the ST23 Kp strain was clearly attributed to colibactin production, using isogenic ΔclbA and complemented mutants19. Subsequent infection experiments using the same strains revealed that loss of colibactin caused a reduction in dissemination to the blood, liver, spleen and brain. Crucially, this work also indicated that the ΔclbA mutant was attenuated in its ability to colonise the intestinal mucosa, which is considered a critical prerequisite for invasive disease42. Similarly, colibactin has been shown to promote gut colonisation and to be essential for disseminated infection by E. coli40. Therefore, we speculate that acquisition of the clb locus promotes gut colonisation and/or mucosal invasion of CG23-I Kp, leading to enhanced transmissibility and increased virulence relative to ICEKp + clb- strains, which may explain the dissemination of the lineage and its dominance amongst hypervirulent infections globally. Infection experiments using the ICEKp + clb- strain NTUH-K2044 show that it is highly virulent despite the lack of colibactin13, but virulence in this strain is likely also influenced by additional features encoded downstream of the

Fig. 3 Virulence and genomic properties of proposed CG23-I strain SGH10. a Bacterial burden (CFU/g) in organs and stool of n = 8 mice 48 h following oral infection, bars indicate mean values. b SGH10 chromosome and plasmid. Tracks shown are (from inner to outer): GC skew (G – C/G + C), G+C content, key capsule and virulence features as labelled, coding sequences on reverse strand and coding sequences on forward strand.

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copies of the iro and rpmA virulence loci that are not found in the majority of CG23 Kp (Figs. 1 and 2, also see below for discussion of other virulence determinants). Concerningly, colibactin also increases the likelihood of serious complications, such as metastatic spread to the brain42 and potentially tumorigenesis43,44. Colibactin-positive Kp is particularly common in Taiwan, where it has been reported in 17–25% of non-abscess infections and significantly associated with K1 strains (which are most likely ST23)43,45. In this setting pyogenic liver abscess, particularly caused by K. pneumoniae, has been shown to be a significant risk factor for colorectal cancer43,46 and liver cancer47. K1 and/or ST23 Kp gut colonisation rates up to 10% have been reported across East Asia48,49. The prevalence of colibactin among these strains has not yet been investigated in most of these countries, however Kp liver abscess has been associated with colorectal cancer across East Asia50, and based on our genome data we predict the majority of the strains in question will belong to the predominant clb + CG23-I sublineage.

The only other gene content changes characteristic of CG23-I were loss-of-function mutations in the fimbral protein KpcE (via nonsense mutation) and the etanolamine transporter Eut (via IS integration). Whether these differences in surface structures or metabolic pathways reflect adaptive selection, or simply loss-of-functions that are not advantageous to the clone, remains a topic for future research. Kp has two operons involved in etanolamine metabolism: eatRKLCAHGEJMTDQF and eat-eutBC. The former is typically associated with facultative anaerobes that live in the gut and/or mouth, and the latter with obligate aerobes that may be found in the environment;51 hence the loss of Eat may reflect adaptation to the host-associated, anaerobic niche. A recent metabolic study of Kp growth under aerobic conditions included two ST23 strains NTUH-K2044 (intact eat) and SB4385 (CG23-I, disrupted eat). Both grew on etanolamine with no apparent differences in efficiency52, indicating no detrimental effect of the loss of Eat in CG23-I strains during aerobic growth, hence the available evidence indicates this change is unlikely to be of particular functional significance to the lineage.

The K1 capsule synthesis locus KLI1, microcin ICE and virulence plasmid encoding aerobactin, salmochelin and rpmA (Fig. 1, Supplementary Fig. 6) were present in most strains, suggesting they were in the ancestor of CG23 and have been selectively maintained. In contrast, distinct variants of ICEKp carrying ybt have been acquired on multiple separate occasions after divergence from the CG23 ancestor. Notably only one such acquisition—that of clb-positive ICEKp10 in CG23-I—was associated with detectable clonal expansion, suggesting positive selection for colibactin production (discussed above). However, given that current genome collections are biased towards clinical strains, we must acknowledge the possibility that CG23 representing other sublineages with or without yersiniabactin and/or colibactin are circulating more frequently among asymptomatic carriers. Indeed, both asymptomatic carriage isolates in our study lacked ICEKp. Asymptomatic gastrointestinal carriage of K1 CG23 has also been reported previously, with up to 10% prevalence in Asian populations48,49 but whether these strains carry ICEKp10 or belong to CG23-I is unknown. On the other hand, three liver abscess strains in our collection also lacked ICEKp, and it was recently reported that the 1932 Murray Collection isolate lacking ICEKp is highly virulent in the Galleria mellonella infection model23. Hence although ybt has been implicated as a key determinant of virulence in mouse models41,12 and has been positively associated with invasive infections in population studies9,14, possession of ybt does not appear to be a strict requirement for invasion. In addition, CG23-I strain 1084, which lacks the virulence plasmid but carries ybt and clb in ICEKp10 (Figs. 1 and 2b), was isolated from human liver abscess and is reportedly highly invasive in murine models of pneumonia and liver abscess19,53. Taken together these data support the notion that no particular virulence factor of CG23 is necessary or sufficient for invasive disease. Future testing using the G. mellonella and/or murine models may shed further light on the interactions between these virulence determinants.

The emergence of AMR CG23 is a significant potential health threat. Treatment of pyogenic liver abscess relies on drainage of the abscess and effective antimicrobial therapy1. AMR has been occasionally reported in human isolates of CG2320–22, and our genomic analysis confirmed that acquisition of AMR plasmids was rare, with distinct plasmids identified in just three sporadic strains from human infections40,46, and no evidence of long-term plasmid maintenance or transmission amongst human isolates (Fig. 1). This is in contrast to the frequency, diversity and apparent stability of AMR plasmids found in other Kp clones such as CG258 or CG1554,34–36, which also show evidence of frequent ICEKp integration14. Overall, our population genomics data suggest there may be barriers to plasmid acquisition and maintenance in CG23 that have so far protected us against the emergence of widespread AMR in this hypervirulent clone. These may include the hyper-expression of K1 capsule, which may provide a physical barrier against transformation and conjugation and also CRISPR/Cas systems, which defend against foreign DNA that does manage to penetrate the cell. The significant association between upregulated capsule production and reduced uptake of DNA has previously been documented in Streptococcus pneumoniae57. Notably, these mechanisms could also explain the lack of homologous recombination in CG23, which is common in other clonal groups35. However, the occasional acquisition of plasmids, including AMR plasmids, in CG23 shows that these barriers are not complete and the maintenance of an IncFII AMR plasmid in the equine clade over 20 years (Supplementary Figs. 6 and 8) shows that long-term stability of AMR plasmids is possible in CG23. Hence our data indicate we must anticipate and carefully monitor for the emergence of stable AMR in CG23.

Taken together our findings have important implications for future study of hypervirulent Kp. Firstly, we suggest that epidemiological studies of Kp liver abscess infections should investigate and report not only chromosomal MLST and virulence plasmid marker genes such as rpmA, but also the ybt and clb loci, which can be used as markers for the CG23-I sublineage. These loci can be rapidly identified in genome data using our Klebsiella genotyping tool, Kleborate (https://github.com/katholt/Kleborate) or can be identified by conventional PCR (primers in Lee et al.)7. Secondly, our observations regarding the population structure and virulence loci show that the two strains that have been used as reference genomes and/or experimental models for hypervirulent ST23 associated with liver abscess are rather atypical (Fig. 1): NTUH-K204417,19 does not belong to the dominant sublineage CG23-I and lacks colibactin, while 108419,30 lacks the virulence plasmid. We therefore suggest that future studies aiming to explore the virulence, transmissibility or evolution of CG23 Kp should consider genome-based profiling prior to the selection of clinical isolates for experimental work. In addition, we present the human liver abscess isolate SGH10 as an open access reference strain for future studies into hypervirulent CG23 Kp associated with liver abscess (strain available under NCTC 14052; complete genome available under GenBank accessions ‘CP025080’, ‘CP025081’). SGH10 exemplifies the most clinically relevant CG23 genotype i.e. that of the globally expanded CG23-I sublineage, including intact copies of all virulence loci. SGH10 is hypermutoid and serum-resistant, is capable of causing disseminated infection in a murine model and in the otherwise healthy human patient from which it was isolated. Finally, we
advocate that researchers, clinicians and public health authorities co-ordinate efforts to improve the surveillance of hypervirulent Kp and facilitate rapid identification of emerging threats, such as convergence of hypervirulence and AMR, that may otherwise remain undetected for decades.

**Methods**

**Genome collection.** All CG23 genomes that were available at the time of study were included in this work. We identified 83 CG23 genome sequences from our curated collection (six finished and 77 draft genomes, see Supplementary Data 1) and 14,26. These included 43 genomes from four of our own previous studies and 40 publicly available genomes from seven published studies, Genbank and the NCTC3000 genomes project (as described previously14,26 and including the 27 genomes analysed by Struve et al.16). We also sequenced 15 novel genomes that were identified as ST23 by MLST (02A029, 12A041 and 16A151) or identified as K1 by classical serotyping and then confirmed as ST23 by genome sequencing (12 strains isolated between 1980 and 1987 from samples from horses; see Supplementary Data 1). DNA was extracted by the phenol–chloroform method, libraries prepared using Nextera technology and paired end reads of either 100 bp (Illumina HiSeq 2000) or 300 bp (Illumina MiSeq) were generated. In total 83 of the 98 genomes represented isolates from human infections (38 liver abscess; 16 bacteremia with/without liver abscess; 5 pneumonia; 15 miscellaneous; 7 unknown) or asymptomatic carriage (n = 2), collected from Africa, Asia, Australia, Europe and South America between 1932 and 2015 (see Supplementary Data 1). One genome was from an unknown source. The remaining genomes represented Kp isolated from horses in France between 1980 and 2004 that were originally selected for whole-genome sequencing solely on the basis of K1 capsule type and were subsequently found to belong to CG23. Accession numbers for all 98 genomes included in this work are listed in Supplementary Data 1 (note though that the hybrid strain CA6868 was excluded from most analyses, as explained below).

**Characterisation of novel CG23-1 reference strain SGH10.** We previously sequenced SGH10 (liver abscess isolate, Singapore 2013) via Illumina MiSeq. Here we generated additional short read (Illumina Miniseq) and long read (Oxford Nanopore) sequences (see Supplementary Methods) and constructed a high quality finished genome sequence using our hybrid assembler Unicycler19. Virulence of SGH10 was assessed using a murine oral infection model as previously described20 with slight modification (see Supplementary Methods). Animal studies were ethically reviewed and approved by the National University of Singapore (NUS) Institutional Animal Care and Use Committee, with the approval number L2015–0135, according to the National Advisory Committee for Laboratory Animal Research (NACLAR) Guidelines.

**Genotyping, yersiniabactin ICEKp and K-locus typing.** Where available (n = 89), short-read sequence data were de novo assembled using Unicycler v0.30.58 with SPAdes v3.8.139 and annotated with Proka v1.1.16. SRST261 was used to (i) determine chromosomal, yersiniabactin and colibactin multi locus sequence types (STs)53–55 and (ii) screen for virulence genes included in the BIGLots database56 and plasmid replicon genes in the PlasmidFinder database63. Nine of the publicly available genomes were available only as pre-assembled sequences. For these we used Kleborate [https://github.com/katholt/Kleborate] to determine MLST, virulence and AMR information, and identified plasmid replicons separately by BLASTn search of the assemblies (coverage ≥ 90%, identity ≥ 90% databases as above). Genomes with novel alleles were submitted to the Kp BIGLots database (http://biglots.pasteur.fr/klebsiella/klebsiella.html) for assignment of novel allele and ST numbers.

Yersiniabactin ICEKp structures were predicted for all genome assemblies using Kleborate17 and confirmed by manual inspection using the Artemis genome browser.54 ICEKp integration sites were determined (from the four possible chromosomal tRNA-Aasn sites) using the Bandage assembly graph viewer65 and BLASTn searches. Micronin E491 ICE sequences were extracted based on their flanking direct repeats and BLASTn searches were used to confirm their integration at tRNA-Aasn site 2. K-loci were typed from the assemblies using Kaptive55.

**Phylogenetic inference and molecular dating analyses.** Sequence reads were mapped to the NTUH-K2044 reference chromosome (accession: ‘AP006726.1’) using RedDog, and each annotated genome was counted as present in a given isolate if the reads covered ≥ 90% of the gene length at read depth ≥ 5. Putative AMR plasmids were investigated using the Bandage assembly graph viewer55 as detailed in Supplementary Methods. To generate a reference sequence for the plasmid in the equine clade, we extracted from the assembly graph of isolate SB4816 a putative novel plasmid sequence comprising three contigs that carried all the acquired AMR genes, was separated from the chromosome assembly graph, and displayed 4-fold read depth compared to the chromosomal sequences. We generated long reads for SB4816 as described for SGH10, yielding sufficient reads to confirm the orientation of the three contigs and resolve a complete circular plasmid sequence, pSB4816. Conservation of this plasmid amongst the other equine isolates was assessed by mapping each read set to the pSB4816 sequence as above. Unfortunately we were not able to extract the complete sequences of the three AMR plasmids from public genome data due to complexities in the assembly graphs, but we were able to identify the putative plasmid replicon types and associated plasmid STs (see Supplementary Methods).

**Data availability.** Accession numbers for all genome data included in this work are summarised in Supplementary Data 1. Illumina sequence reads for the 15 newly sequenced equine isolates have been deposited in the NCBI sequence read archive under accession ‘PRJNA391904’. The MDR plasmid sequence pSB4816 was deposited in GenBank (accession: ‘MF302404’). The finished chromosome sequence for SGH10 (BioSample: ‘SAMN06121188’) was deposited in GenBank (accessions ‘CP025800.1’, ‘CP025801’ and the strain was deposited in the UK National Collection of Type Cultures (NCTC 14052); novel Illumina and Oxford Nanopore sequence reads for this isolate are available as a complete assembly graph read archive under accession ‘SRP263939’, ‘SRP263933’, ‘SRP263932’ and ‘SRP263931’. The Bayesian phylogenetic tree, annotated with strain information, is available for download and interactive visualisation at https://microreact.org/project/ri1_d5C9w-

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**Author contributions**

M.M.C.L. and K.L.W. analysed data and wrote the paper. S.D., R.R.W., S.A. analysed data. L.M.J., V.P., Y.H.G. and C.H.H. performed lab experiments. Y.H.G., S.A., J.S.M., S. K., T.H.K. and S.B. contributed isolates for sequencing. S.B. and K.E.H. conceived the study, analysed and interpreted data and contributed to manuscript writing.

**Additional information**

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