# A curated collection of *Klebsiella* metabolic models reveals variable substrate usage and gene essentiality

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The *Klebsiella pneumoniae* species complex (KpSC) is a set of seven *Klebsiella* taxa that are found in a variety of niches and are an important cause of opportunistic health care–associated infections in humans. Because of increasing rates of multi-drug resistance within the KpSC, there is a growing interest in better understanding the biology and metabolism of these organisms to inform novel control strategies. We collated 37 sequenced KpSC isolates isolated from a variety of niches, representing all seven taxa. We generated strain-specific genome-scale metabolic models (GEMs) for all 37 isolates and simulated growth phenotypes on 5II distinct carbon, nitrogen, sulfur, and phosphorus substrates. Models were curated and their accuracy was assessed using matched phenotypic growth data for 94 substrates (median accuracy of 96%). We explored species-specific growth capabilities and examined the impact of all possible single gene deletions using growth simulations in 145 core carbon substrates. These analyses revealed multiple strain-specific differences, within and between species, and highlight the importance of selecting a diverse range of strains when exploring KpSC metabolism. This diverse set of highly accurate GEMs could be used to inform novel drug design, enhance genomic analyses, and identify novel virulence and resistance determinants. We envisage that these 37 curated strain-specific GEMs, covering all seven taxa of the KpSC, provide a valuable resource to the *Klebsiella* research community.

[Supplemental material is available for this article.]

*Klebsiella pneumoniae* is a ubiquitous bacterium that inhabits a variety of host- and non-host-associated environments and is a major cause of human disease. It is an opportunistic pathogen and a significant contributor to the spread of antimicrobial resistance globally (Pendleton et al. 2013; Navon-Venezia et al. 2017; Thorpe et al. 2021). Multi-drug-resistant *K. pneumoniae* with resistance to the carbapenems (the "drugs of last resort") cause infections that are extremely difficult to treat and are considered an urgent public health threat (Pendleton et al. 2013). Understanding the biology and ecological behavior of these organisms is essential to inform novel control strategies.

The last 6–7 yr have seen an explosion of *K. pneumoniae* comparative genomics studies, revealing numerous insights into its epidemiology, evolution, pathogenicity, and drug resistance, and informing a genomic framework that facilitates surveillance and knowledge generation (for a recent summary, see Wyres et al. 2020). It is now clear that isolates identified as *K. pneumoniae* through standard microbiological identification techniques actually comprise seven distinct closely related taxa known as the *K. pneumoniae* species complex (KpSC): *K. pneumoniae sensu stricto*, *Klebsiella variicola* subsp. *variicola*, *K. variicola* subsp. *tropica*, *Klebsiella quasipneumoniae* subsp. *quasipneumoniae*, *K. quasipneumoniae* subsp. *similipneumoniae*, *Klebsiella quasivariicola*, and

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Article published online before print. Article, supplemental material, and publication date are at https://www.genome.org/cgi/doi/10.1101/gr.276289.121. Freely available online through the *Genome Research* Open Access option. *Klebsiella africana* (Gorrie et al. 2017; Long et al. 2017; Rodrigues et al. 2019; Wyres et al. 2020). *K. pneumoniae sensu stricto* accounts for the majority of human infections and is therefore the most well-studied of these organisms.

Each individual K. pneumoniae genome encodes between 5000 and 5500 genes; approximately 2000 are conserved among all members of the species (core genes), and the remainder vary between individuals (accessory genes) (Holt et al. 2015). The total sum of all core and accessory genes is estimated to exceed 100,000 protein-coding sequences that can be assigned to various functional categories, many of which are not well-characterized. For example, the diversity, mechanism, and phenotypic impact of antimicrobial resistance genes, accounting for 1% of the total gene pool, is well understood. In contrast the functional implications of metabolic genes, which account for the largest single fraction of the gene pool (37%) (Holt et al. 2015), are relatively poorly understood. The sheer number of genes in this category suggests that substantial metabolic variability exists within the KpSC, a hypothesis supported by two studies that have generated growth phenotypes for multiple isolates (Brisse et al. 2009; Blin et al. 2017). However, these data are limited by the number and variety of substrates tested and it is difficult to consolidate the genotype data in the context of these phenotypes. Moreover, these phenotyping methods are slow, expensive, and nonscalable across large numbers of isolates.

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Genome-scale metabolic modeling represents a powerful approach to bridge the gap between genotypes and phenotypes. Drawing on the accumulated biochemical knowledge base, it is possible to infer the metabolic network of an individual organism from its genome sequence and subsequently apply in silico modeling approaches to predict its metabolic capabilities (growth phenotypes) (Thiele and Palsson 2010; O'Brien et al. 2015). Such models allow exploration of metabolic diversity (Monk et al. 2013; Bosi et al. 2016; Seif et al. 2018), prediction of the impact of gene deletions or the response to drug exposure (Tong et al. 2020), identification of novel virulence factors or drug targets (Bartell et al. 2017; Ramos et al. 2018; Zhu et al. 2018), and optimization for the production of industrially relevant compounds (Jung et al. 2015; Li et al. 2016).

To date, two curated and validated single-strain genome-scale metabolic models (GEMs) have been reported for K. pneumoniae. The first was generated for the MGH78578 laboratory strain and published in 2011 (model ID iYL1228) (Liao et al. 2011). It comprised 1228 genes, 1188 enzymes, and 1970 reactions, and was validated by comparison of in silico growth predictions to true phenotypes generated for 171 substrates using a Biolog phenotyping array. The estimated accuracy of iYL1228 was 84% when compared to Biolog growth phenotypes. A second K. pneumoniae GEM, for laboratory strain KPPR1, was published in 2017 (model ID iKp1289) (Henry et al. 2017). This model contained 1289 genes and 2145 reactions. The KPPR1 model was found to be 79% accurate when compared to Biolog phenotype data in terms of predicting substrate-growth phenotypes. More recently, Norsigian and colleagues (Norsigian et al. 2019a) reported nonvalidated draft GEMs for 22 antimicrobial-resistant K. pneumoniae clinical isolates built from the iYL1228 model through a subtractive approach. Subsequent in silico growth predictions indicated variability between isolates in terms of carbon, nitrogen, and sulfur, but not phosphorus utilization. There was evidence that nitrogen substrate usage could be used to classify strains associated with distinct drug resistance phenotypes. However, none of these models were experimentally validated.

Here, we present an updated version of the MGH78578 GEM in addition to novel GEMs for 36 KpSC strains, including representatives of all seven taxa in the species complex. We curate and validate the models using a combination of Biolog growth assays and additional targeted growth phenotype data, resulting in a median accuracy of 96%. We define the core reactomes of *K. pneumoniae* and the broader species complex, and we identify species-specific metabolic capabilities. We then explore these models to identify strain-specific gene essentiality and metabolic pathway redundancy across growth on 145 core carbon substrates.

# Results

#### Completed KpSC genomes

We collated 37 previously described isolates from the KpSC complex, including at least one representative per taxon (Blin et al. 2017; Rodrigues et al. 2019). The collection spanned a variety of sequence types (STs) within species with more than one strain and represented a wide range of isolation sources (including humanhost-associated, water, and the environment). The strains were geographically and temporally diverse, sampled from five continents, and with isolation dates spanning from 1935 to 2010 (Supplemental Table 1).

Eight strains had previously published complete genome sequences available, and we generated complete genome sequences for the remaining 29 strains using a combination of short- and long-read sequencing (Methods). The median genome size was 5.5 Mbp (range 5.1–6.0 Mbp) with a median of 5145 genes (range 4798–5704 genes). The majority of strains carried at least one plasmid (n=29, 78%), with seven strains carrying five or more plasmids.

## Model generation, curation, and validation

Using these completed genomes we created strain-specific GEMs, initially using the curated MGH78578 GEM (iYL1228) as a reference to identify conserved genes and reactions, followed by manual curation (Methods). The latter was enabled by the availability of matched phenotype data (Blin et al. 2017) indicating the ability of each strain to grow in minimal media supplemented with each of 94 distinct sole carbon substrates for which we were able to predict growth in silico using the GEMs (Supplemental Table 2). Our phenotypic data included 12 carbon substrates for which growth was shown for at least one strain and for which the corresponding metabolite transport and/or processing reactions were not present in the original iYL1228 model. Literature searches were undertaken to identify the putatively responsible candidate genes and reactions for GEM inclusion. For example, all strains were able to use palatinose as a carbon substrate; the reaction required to catabolize this compound was added based on the presence of core genes with  $\geq 99\%$  nucleotide homology with *aglAB* (that encode AglAB), which has been shown to catabolize palatinose in K. pneumoniae (Supplemental Table 3; Thompson et al. 2001). When the model-based predictions and our phenotypic growth data disagreed, we attempted to correct the models by identifying alternative pathways from the literature or homologous genes in other Klebsiella or Enterobacteriaceae species with sufficient evidence to allow inclusion in our models (Methods; Supplemental Table 3). Overall, we added 49 genes and 56 reactions across all models.

The final curated, validated models were highly accurate for the prediction of growth phenotypes measured through Biolog (median accuracy 95.7%, range 88.3%-96.8%) (Supplemental Table 1). The majority (87%) of the discrepancies were false positives, that is, the model predicted growth on a carbon substrate, but we did not observe any phenotypic growth. False positives usually occur because of gene regulation, in which strains carry the genes encoding the enzymes required to import and metabolize a substrate; however, these genes are not expressed during the phenotypic growth experiments. False positives can also be related to technical issues with measuring metabolic phenotypes, for example, the limit of detection, sensitivity of growth detection, and use of correct standards for measurements (Ibarra et al. 2002). Every model had at least one false positive (median 4, range 1-11) (Supplemental Table 1) across 31 different carbon substrates. The most common false positive calls were predicted growth in 2-oxoglutarate (n = 35 strains), ethanolamine (n = 29), L-ascorbate (n = 10028), and 3-hydroxycinnamic acid (n = 20); false positive calls for the remaining 27 carbon substrates were associated with six or fewer strains each (Supplemental Table 4).

Five carbon substrates had at least one strain with a false negative call, in which the model did not predict growth, but we observed a growth phenotype: L-tartaric acid (n = 12 strains), Llyxose (n = 5), L-sorbose (n = 2), propionic acid (n = 2), and L-galactonic acid-gamma-lactone (n = 1) (Supplemental Table 4). In such cases it is assumed that the models are missing information required to optimize for growth on these substrates (Orth and Palsson 2012). Despite thorough literature and database searches,

Table 1.	Summary of	genomes a	nd the core	elements of	of the GEMs
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Species	Genomes	STs	Model genes (core)	Reactions (core)	Phenotypes (core)
K. pneumoniae	20	18	1202–1243 (1183)	2288–2305 (2276)	277–282 (277)
K. quasipneumoniae subsp. quasipneumoniae	5	5	1197–1209 (1190)	2283-2289 (2283)	270–274 (268)
K. quasipneumoniae subsp. similipneumoniae	5	5	1200–1220 (1194)	2283–2299 (2287)	273–280 (273)
K. variicola subsp. variicola	4	4	1212–1227 (1214)	2294–2301 (2299)	279–282 (279)
K. africana	1	1	1216	2289	279
K. quasivariicola	1	1	1228	2299	279
K. variicola subsp. tropica	1	1	1237	2310	281

we were unable to identify alternate biological pathways that could plausibly fill these gaps in the models. This was particularly notable among the five *K. quasipneumoniae* subsp. *quasipneumoniae* strains, which all had false negative predictions for L-lyxose utilization. These genomes were each missing *sgaU* (KPN\_04590), which was present in all other KpSC genomes and encodes an enzyme that converts L-ribulose-5-phosphate to L-xylulose-5-phosphate. We were unable to detect any other proteins belonging to this enzyme class or carrying similar domains. As the phenotypic results indicated that all *K. quasipneumoniae* subsp. *quasipneumoniae* can use L-lyxose, we hypothesize that they must contain unknown functional ortholog/s to *sgaU*, which can perform isomerase activity on L-ribulose-5-phosphate.

We performed an independent validation of the models by comparing growth phenotypes from the VITEK GN card with simulated phenotypes (n = 13 substrates; Methods). The models were highly accurate in this setting (median accuracy 100%, range 92.3%–100%) (Supplemental Table 5). All discrepancies were false positives (n = 4): two for growth in succinate, one in tagatose, and one in 5-keto-D-gluconate (Supplemental Table 5).

#### Novel GEMs reveal species- and strain-specific metabolic diversity

Our strain collection provided us with a novel opportunity to compare predicted metabolic functionality between all seven taxa within the KpSC. Overall there were median 1219 genes and 2294 reactions in each curated strain-specific GEM (ranges 1190-1243 and 2283-2305, respectively), representing median 23.6% of all coding sequences in each genome (Supplemental Table 1). Each species had approximately 1200 core model genes and about 2200 core reactions (Table 1), with a slight decreasing trend with increasing sample size. Conversely, the total number of distinct reactions detected among the best represented species, K. pneumo*niae* (2312, n=20 genomes), was higher than those detected among each of the species represented by fewer genomes (2299 in K. quasipneumoniae subsp. quasipneumoniae; 2307 in both K. quasipneumoniae subsp. similipneumoniae, and K. variicola subsp. variicola). In terms of the reactions themselves, the vast majority were core across all species (Fig. 1); however, there was variability in reactions associated with carbohydrate metabolism, for which 16% (n = 37/234) were not conserved across all models (Fig. 1). Among these variable reactions we identified three involved in the N-acetylneuraminate pathway (ACNAMt2pp, ACNML, and AMANK) that were species-specific and were found to be core in all five K. quasipneumoniae subsp. similipneumoniae in our study, but absent from all other genomes. A BLASTN screen of all 307 K. quasipneumoniae subsp. similipneumoniae genomes from Lam et al. (2021) revealed that these three genes were present in all 307 genomes, indicating that this pathway is likely to be core across all members of the species.

We simulated growth on 511 substrates as the sole sources of either carbon (n = 272), nitrogen (n = 155), phosphorus (n = 59), or sulfur (n = 25) (Methods; Supplemental Table 2). A total of 224 (44%) were unable to support growth for any strain (carbon = 107, nitrogen = 87, phosphorus = 15, sulfur = 15). Overall, the number of core growth-supporting phenotypes was very similar across taxa, with a median of 279 (range 268–281) (Table 1). Of the 287 that were predicted to support growth for at least one strain, 262 were conserved across all 37 strains (carbon = 145, nitrogen = 64, phosphorus = 43, sulfur = 10), with only 25 (5%) substrates variable between strains. Substrates that could be used as a carbon source had the most variation, with 7% of carbon substrates displaying variable predicted growth phenotypes by strain (Fig. 2). This was in stark contrast to substrates used as a source of sulfur, in which no variation was observed (Fig. 2).

Among the 20 variable carbon substrates, there was some species-specific variation. Six of these reflect core growth capabilities in all but one of the seven species (3-hydroxycinnamic acid, 3-(3-hydroxy-phenyl)propionate, D-arabitol, L-ascorbate, L-lyxose, tricarballylate) (Fig. 3). In the case of tricarballylate, we identified a new pathway that was absent from the original K. pneumoniae MGH78578 model: all KpSC species except for K. pneumoniae carried the *tcuABC* operon, which encodes the enzymes responsible for oxidizing tricarballylate to cis-aconitate (Lewis et al. 2009) through the TCBO reaction (Fig. 3). In contrast, all KpSC were able to use L-ascorbate with the exception of K. quasipneumoniae subsp. quasipneumoniae, in which all five genomes were lacking the ulaABC operon encoding the transport reaction ASCBptspp (Fig. 3). This reaction converts L-ascorbate into L-ascorbate-6-phosphate as it is transported into the cytosol (Zhang et al. 2003). We screened all 149 K. quasipneumoniae subsp. quasipneumoniae genomes from Lam et al. (2021) for ulaABC with BLASTN and found that this operon was missing from all members of the species, suggesting that this is a conserved deletion in K. quasipneumoniae subsp. quasipneumoniae.

The remaining 14 variable carbon substrates were specific to five or fewer strains. For example, sn-glycero-3-phosphocholine could be used by all strains as a carbon and phosphorus substrate, except for the single *K. africana* and *K. quasivariicola* representatives, which share a common ancestor in the core gene phylogenetic tree (Fig. 3). Both these genomes lacked *glpQ*, encoding the enzyme required to convert sn-glycero-3-phosphocholine into sn-glycero-3-phosphate and ethanolamine (Brzoska and Boos 1988). We confirmed that *glpQ* was absent in all 13 *K. quasivariicola* genomes listed in Lam et al. (2021) by screening for the gene using BLASTN. To check the result if the *glpQ* deletion is present in other *K. africana* (because we have only a single genome), we screened six *K. africana* genomes (all ST4838) for *glpQ* from Vezina et al. (2021) and found that this gene was present in all strains. There was only a single carbon substrate, N-acetylneuraminate, which supported growth for all



**Figure 1.** Number of model reactions by category. Bars are colored to indicate core reactions (black, conserved in all strains) and accessory reactions (gray, variably present).

*K. quasipneumoniae* subsp. *similipneumoniae*, because of the presence of the *nan* operon (Vimr and Troy 1985), encoding the proteins required to catalyze the ACNAMt2pp, ACNML, and AMANK reactions, which were absent in all the other species (Fig. 3).

#### Single gene knockout simulations reveal variable gene essentiality

Strain-specific GEMs provide an unparalleled opportunity to simulate the impact of single gene knockout mutations for diverse strains. Because carbon substrates were associated with the greatest amount of variation, we focused on the impact of single gene knockouts in this group. For each strain we simulated the impact of deletion of each unique gene in its

GEM on growth in each of the core carbon substrates (those predicted to support growth of all strains, n = 145), resulting in 6,544,865 unique simulations (Supplemental Table 6). Among these simulations, 639,365 (9.8%) were predicted to result in a loss-of-growth phenotype.

To compare the diversity of knockout phenotypes between strains, we focused on simulations representing core gene-substrate combinations; 1133 genes that were present in all GEMs × 145 substrates) and excluded those representing noncore gene-substrate combinations (n = 19,140 combinations), because the former can be directly compared for all strains, whereas the latter cannot (by definition not all strains harbor all of the genes). A total of 146,385 core gene-substrate combinations (89.1%) resulted in no loss-ofgrowth phenotype in any strain, whereas 7170 (10.5%) combinations resulted in a loss-of-growth phenotype in all strains. At the gene level, 807 genes (71.2%) were not predicted to be essential for growth for any substrate in any strain, and just 57 genes (5.0%) were predicted to be essential for all substrates in all strains. The latter were associated with 194 distinct reactions (1-32 reactions each, median = 1, Supplemental Table 7), encompassing eight subsystem categories: cell membrane metabolism (n= 76 reactions), lipid metabolism (n = 42), amino acid metabolism (n=33), transport, inner- (n=29) or outer-transport (n=6), nucleotide metabolism (n=5), carbohydrate metabolism (n = 2), and cofactor and prosthetic group biosynthesis (n = 1).

Gene essentiality varied by strain, with reasonable consistency within species. The number of core gene-substrate combinations predicted to result in a loss-of-growth phenotype ranged from 0 to 519 (median = 143) (Fig. 4), and the

number of core genes resulting in a phenotype on at least one growth substrate ranged from 0 to 15 (median = 3). The vast majority of these genes (31 of 36 unique genes, 86.1%) were associated with loss-of-growth phenotypes for six or fewer substrates, with minimal variation in the total number of substrates among those strains that were impacted. In contrast, a small number of genes were associated with loss of growth for all or almost all substrates for some strains (four genes, 11.1%, each affecting 143 or more substrates per strain) (Fig. 4).

We further investigated the core gene deletions predicted to result in loss-of-growth phenotypes for 143 or more substrates in







**Figure 3.** Variable growth phenotypes across all seven taxa in KpSC. (*Left*) Core gene phylogeny for all 37 strains, with tips colored by species per legend. (*Middle*) Heatmap of variable substrates for which both phenotypic growth results and model predicted results were available. White indicates no growth, and color indicates growth. False positive calls are shown in yellow, and false negative calls are in gray (per legend). (*Right*) Heatmap of variable substrates for which indicates no growth, color indicates growth, with substrate type indicated per legend.

only a subset of strains, beginning with an apparent K. quasipneumoniae subsp. quasipneumoniae species-specific phenotype. The associated gene, KPN\_03428, encodes the enzyme for catalysis of two reactions in the models: CYSDS (cysteine desulfhydrase) and CYSTL (cystathionine b-lyase), the latter of which may also be encoded by KPN\_01511 (malY). malY was present in all other models but absent from all K. quasipneumoniae subsp. quasipneumoniae (closest bidirectional BLASTP hit had 30.07% identity, well below the threshold required for inclusion as a homolog and considerably lower than the expected divergence between KpSC species; 3%-4% nucleotide divergence) (Holt et al. 2015), and no alternate genes encoding putative cystathionine b-lyases could be identified by search of the KEGG database, indicating a lack of genetic redundancy for these reactions. Direct comparison of the K. quasipneumoniae subsp. quasipneumoniae 01A030T chromosome to K. pneumoniae MGH78578 revealed that the former harbored a ~5 kbp deletion relative to the latter, spanning the zntB, malY, and malX genes as well as part of malI. The lack of malY (KPN\_ 01511) in combination with the KPN\_03428 deletion resulted in predicted loss of ability to produce three key metabolites (L-homocysteine, ammonium, and pyruvate) and ultimately the predicted loss of biomass production. This deletion was replicated in all five K. quasipneumoniae subsp. quasipneumoniae strains. Inspection of an additional 149 publicly available K. quasipneumoniae subsp. quasipneumoniae genome assemblies (Methods) found this region to be present in only 37 genomes (24%), suggesting that the most recent common ancestor of this species is lacking this region, with occasional reacquisition in some lineages.

Unlike the KPN\_03428 deletion, deletion of KPN\_04246 resulted in predicted loss-of-growth phenotypes for all 145 substrates for the single K. africana strain plus 13 of 20 K. pneumoniae strains (comprising multiple distantly related lineages including representatives of the well-known globally distributed ST14, ST23, ST86, and ST258). KPN\_04246 encodes a protein that catalyzes two reactions-ACODA, acetylornithine deacetylase, and NACODA, N-acetylornithine deacetylase-both of which may also be encoded by the product of KPN\_01464 (homologs of this gene were identified in only those genomes that were not associated with loss-of-growth phenotype). Comparison of the K. pneumoniae strain CG43 (ST86) chromosome lacking KPN\_01464 to K. pneumoniae MGH78578 harboring KPN\_01464 showed that CG43 contained a ~10 kbp deletion resulting in the loss of KPN\_01464. This deletion was replicated in the K. africana 200023T genome and the remaining 12 K. pneumoniae genomes that lacked KPN\_01464 (≤33.24% identity for the best bidirectional BLASTP hit, no alternate genes encoding putative acetylornithine deacetylases/N-acetylornithine deacetylases were identified in KEGG).

Finally, we investigated the two gene deletions (KPN\_02238 and KPN\_00456) resulting in predicted loss of growth on all



**Figure 4.** Variable loss-of-growth phenotypes. (*Left*) Core gene phylogeny per Figure 3, with tips colored by species as indicated in legend: (Ka) *K. africana*; (Kp) *K. pneumoniae*; (Kqq) *K. quasipneumoniae* subsp. *quasipneumoniae*; (Kqs) *K. quasipneumoniae* subsp. *similipneumoniae*; (Kqv) *K. quasivariicola*; (Kvt) *K. variicola* subsp. *tropica*; (Kvv) *K. variicola* subsp. *variicola*. (*Middle*) Heatmap showing core genes for which variable loss-of-growth phenotypes were predicted (columns). Shading indicates the number of substrates where loss of growth was predicted for each strain (rows) per the scale legend. (*Right*) Bars show the total number of loss-of-growth phenotypes predicted for each strain.

substrates in only *K. pneumoniae* NJST258-1. KPN\_02238 encodes the protein responsible for catalyzing phosphoribosylpyrophosphate synthetase (PRPPS), for which no redundant genes were included in any of our KpSC models. This reaction converts alpha-Dribose 5-phosphate to 5-phospho-alpha-D-ribose 1-diphosphate, a key substrate used as input for 14 downstream reactions. Although the *K. pneumoniae* MGH78578 reference model contains a redundant pathway to support this conversion, one of the required reactions (R15BPK, catalyzed by a ribose-1,5-bisphosphokinase) was missing from the NJST258-1 model because the associated genome lacked a homolog of KPN\_04492 (best bidirection BLASTP hit 26.19% identity), whereas all other genomes contained a homolog of this gene. Further investigation showed that the NJST258-1 chromosome was missing a ~17 kbp region compared to MGH78578. In the NJST258-1 chromosome, this region, which included KPN\_04492, was replaced by the insertion sequence IS*1294* (99% nucleotide identity). We were not able to identify a similar deficiency to explain the strain-specific loss-of-growth phenotype associated with KPN\_00456, which encodes a protein implicated in 14 distinct reactions.

# Discussion

Here, we present an updated GEM for *K. pneumoniae* MGH78578 plus novel GEMs for 36 members of the KpSC, capturing all seven taxa and representing the first reported GEMs for the *K. variicola* (subsp. *variicola* and *tropica*), *K. quasipneumoniae* (subsp. *quasipneumoniae* and *similipneumoniae*), *K. quasivariicola*, and *K. africana* 

species. All models were validated and curated by comparison of predicted and true growth phenotypes, and they had a median accuracy of 95.7% (range 88.3%–96.8%), higher than estimated for the previously published *K. pneumoniae* MGH78578 (84%) and KPPR1 (79%) models.

Our in silico growth phenotype predictions for a diverse set of substrates highlighted variability among strains within the K. pneumoniae species, as has been indicated by previous smaller scale GEM comparisons and phenotypic comparisons (Brisse et al. 2009; Blin et al. 2017; Henry et al. 2017; Norsigian et al. 2019a). Similar variability was also indicated within and between the other species in the KpSC (Fig. 3). Carbon substrates were associated with the greatest diversity; a total of 145 substrates (53%) predicted to support growth of all 37 strains and 20 (7%) predicted to support growth of 1-36 strains each (Fig. 2). These predictions were consistent with the observed reaction variability, in which the highest proportion of accessory reactions was identified among those associated with carbohydrate metabolism (16%) (Fig. 1). This is consistent with a previous pan-genome analysis of 328 K. pneumoniae that indicated that ~50% of the total gene pool predicted to encode proteins with metabolic functions were specifically associated with carbohydrate metabolism (Holt et al. 2015). This trend is also consistent with previous studies of the closely related species, Escherichia coli, which showed carbohydrate metabolism as the most diverse category for this organism (Monk et al. 2013; Fang et al. 2018).

The extent of diversity reported for E. coli and Salmonella spp. (Seif et al. 2018) was much higher than reported here for KpSC. We propose two likely explanations for these differences. First, the current analysis for KpSC comprises just 37 strains, compared to 55 and 110 strains included in the E. coli studies (Monk et al. 2013; Fang et al. 2018) and 410 in the Salmonella study (Seif et al. 2018). With greater sample size we expect to capture greater gene content diversity (Tettelin et al. 2008), including genes associated with metabolic functions that drive metabolic diversity (as was shown to be the case for Salmonella spp.) (Seif et al. 2018). Second, our draft KpSC strain-specific models were generated using the reference-based protocol (Norsigian et al. 2019b), in which homology search is used to identify genes in the reference model that are absent from the strain of interest and are therefore removed from the strain-specific model. We added novel genes/reactions to the models based on comparison of predicted versus observed growth phenotypes and manual sequence/literature search, but we did not conduct an automated screen to identify additional genes that are present in the novel strain collection. The latter approach is expected to reveal further diversity, but it requires significant manual curation and validation to ensure the high-quality status of the models is maintained; therefore, it should be addressed in future studies.

In addition to growth capabilities, our analyses revealed considerable variation in terms of predicted gene essentiality as has been implicated for other bacterial species (Breton et al. 2015; Poulsen et al. 2019; Tong et al. 2020; Rousset et al. 2021). Specifically, our data indicate that (1) deletion of a single core gene in a given strain may result in loss of growth on all, none, or only a subset of growth substrates; and (2) the impact of such deletions may vary between strains (Fig. 4). Among genes for which deletion was predicted to have variable impact, most were associated with the loss of growth for only a small number of substrates in the impacted strains. However, four genes were associated with predicted loss of growth on 143 or more of 145 substrates for between one and 14 strains each. In two cases (genes KPN 03428 and KPN 04246), the impacted strains were missing redundant genes that were present in the MGH78578 reference model-that is, those encoding proteins with the same functional annotation as the deleted gene. Comparisons of the chromosomes of these strains suggested that the genes were lost through largescale chromosomal deletions (5-10 kbp). One of these deletions was uniquely conserved among strains belonging to K. quasipneumoniae subsp. quasipneumoniae, suggesting that it may have occurred in the most recent common ancestor of this subspecies and has been inherited through vertical descent, with evidence from additional public genome data pointing toward recent reacquisition of this region in some lineages. The other chromosomal deletion was found among a distantly related subset of K. pneumoniae as well as the single K. africana isolate; therefore, its distribution cannot be explained by simple vertical ancestry. Rather, we speculate that this deletion has been disseminated horizontally by chromosomal recombination, as is known to occur frequently among K. pneumoniae (Bowers et al. 2015; Wyres et al. 2019) and has been reported between KpSC species (Holt et al. 2015).

Deletion of two genes (KPN\_02238 and KPN\_00456) resulted in the loss of growth on all substrates for only a single strain (K. pneumoniae NJST258-1). This strain is of particular interest because it was associated with the highest number of deletion phenotypes (Fig. 4), and it belongs to ST258, a globally distributed cause of carbapenem-resistant K. pneumoniae infections (Bowers et al. 2015; Wyres et al. 2020). We were unable to identify the cause of this rare knockout phenotype (lacking adenylate kinase, encoded by KPN\_00456), which converts D-ribose 1,5-bisphosphate to 5phospho-alpha-D-ribose 1-diphosphate at the cost of 1 ATP. Comparison of the metabolic networks of NJST258-1 and MGH78578 indicated that NJST258-1 was lacking an additional reaction pathway (phosphoribosylpyrophosphate synthetase) present in MGH78578, allowing an alternative means of 5-phospho-alpha-D-ribose 1-diphosphate production in the absence of ribose-1,5bisphosphokinase. Further investigation showed that the NJST258-1 chromosome was missing a ~17 kbp region containing one of the genes required to express this redundant pathway, which had been replaced by an insertion sequence (IS). ISs are frequently identified among Klebsiella and other Enterobacteriaceae where they are particularly associated with large plasmids and the dissemination of antimicrobial resistance (Adams et al. 2016; Che et al. 2021). The carbapenem-resistant K. pneumoniae lineage, ST258, has been associated with particularly high IS burden (Adams et al. 2016), and we hypothesize that such insertions contribute to the increased number of gene deletion phenotypes predicted for NJST258-1 compared to other K. pneumoniae strains. We screened an additional 1021 nonredundant ST258 genomes from Lam et al. (2021) for the presence of KPN\_02388 (the gene which encodes for phosphoribosylpyrophosphate synthetase) and found that this gene was present in all 1021 ST258 genomes, suggesting that the deletion of this pathway is unique to NJST258-1. This highlights the importance of assessing multiple strains when attempting to draw conclusions regarding observed phenotypes.

These findings indicate that KpSC can differ substantially in terms of metabolic redundancy. We cannot exclude the possibility that the predicted knockout phenotypes might be rescued by products of nonorthologous genes that are not currently captured in our models, but at least for the aforementioned examples, a search of the KEGG database did not indicate any additional known redundant metabolic pathways. Additionally, our findings are consistent with a recent experimental exploration of gene essentiality in *E. coli* (Rousset et al. 2021), which showed that 7%– 9% of roughly 3400 conserved genes were variably essential among 18 *E. coli* strains grown in three different conditions. Genomic comparisons of these *E. coli* implicated a key role for horizontal gene transfer in driving strain-specific essentiality patterns and redundancies through the mobilization of homologous or analogous genes and/or those driving epistatic interactions (Rousset et al. 2021).

Taken together our findings highlight the importance of strain-specific genomic variation in determining strain-specific metabolic traits and redundancy. More broadly, these analyses show the value of an organism investing in redundant systems, either through (1) encoding multiple genes capable of performing the same reaction, or through (2) encoding multiple, alternative pathways for producing key metabolites from different substrates. Given what is known about the extent of genomic diversity among K. pneumoniae and the broader KpSC (Holt et al. 2015; Wyres et al. 2019; Thorpe et al. 2021), it is clear that studies seeking to understand the metabolism of these species-for example, for novel drug design, or to identify novel virulence and drug resistance determinants-should include a diverse set of strains. In this regard, we anticipate that the GEMs, growth predictions, and single gene deletion predictions presented here will provide a valuable resource to the Klebsiella research community that can be used to understand the fundamental biology of these organisms and to derive clinically relevant insights-for example, to understand how substrate usage patterns influence pathogenicity and virulence, or to identify universal or clone-specific metabolic choke points wherein the associated essential genes/proteins could be targeted by novel therapeutics. As exemplified for the E. coli K-12 reference strain, such resources can be continually improved and expanded to maximize their utility and facilitate biological discovery for years to come (Schilling et al. 1999; Monk et al. 2017).

# **Methods**

#### Genome collection

The 37 strains used in this study were sourced from two previous studies (Blin et al. 2017; Rodrigues et al. 2019). Eight strains had completed genome sequences already publicly available, generated using various sequencing and assembly methods (for details, see Supplemental Table 1). For the remaining 29 strains, shortand long-read sequencing was conducted as follows. Genomic DNA was extracted from overnight cultures, using GenFind v3 reagents (Beckman Coulter). The same DNA extraction was used for both Illumina and MinION libraries. Illumina sequencing libraries were made with Illumina DNA Prep reagents (catalog no. 20018705) and the Illumina Nextera DNA UD Indexes (catalog no. 20027217) per manufacturer's instructions with one major deviation from described protocol: Reactions were scaled down to 25% of recommended usage. Illumina libraries were sequenced on the NovaSeq platform using the 6000 SP Reagent Kit (300 cycles; catalog no. 20027465), generating 250-bp paired-end reads. A total of 21 strains were sequenced across multiple long-read sequencing libraries, prepared using the ligation library kit (LSK-109, Oxford Nanopore Technologies [ONT]) with native barcoding expansion pack (EXP-NBD104 and NBD114, ONT). The libraries were run on a R9.4.1 MinION flow cell and base called with Guppy v3.3.3 using the dna\_r9.4.1\_450 bps\_hac (high-accuracy) base calling model. The remaining seven strains had their DNA extracted using Qiagen Genomic DNA kits (Qiagen Genomic-tip 100/G) and sequenced using Pacific Biosciences (PacBio) RS II.

The Illumina and MinION read data were combined to generate completed genomes for n = 28/29 strains with Unicycler v0.4.8 (Wick et al. 2017) using default parameters. SB610 could not be assembled into a completed genome using this approach, so we used Trycycler v0.3.3 (Wick et al. 2021) to combine 12 independent long-read only assemblies into a single consensus assembly. The 12 assemblies were generated from 12 independent subsets of the long reads (randomly selected) at 50× depth, which were assembled with one of three assemblers (n=4 assemblies each): Flye v2.7 (Kolmogorov et al. 2019), Raven v1.1.10 (Vaser and Šikić 2021), and Miniasm v0.3 (Li 2016). The final consensus assembly was then polished with the long reads using Medaka v1.1.3 (https://github.com/nanoporetech/medaka), followed by three rounds of polishing using the Illumina reads with Pilon v1.23 (Walker et al. 2014). The PacBio reads were assembled with HGAP, and overlaps between contigs extremities were manually circularized. All 37 completed genomes were annotated with Prokka v1.13.3 (Seemann 2014), using a trained annotation model (created using 10 genomes with Prodigal v2.6.3) (Hyatt et al. 2010). All genomes were analyzed with Kleborate v2.0.3 (Lam et al. 2021) to obtain ST and other genomic information (Supplemental Table 1).

#### Phenotypic testing

We used the Biolog growth phenotypes for 190 carbon substrates generated previously (Blin et al. 2017; Rodrigues et al. 2019). As determined in Blin et al. (2017), a maximum value in the respiration curve of 150 or greater was used to indicate growth, whereas a value of less than 150 indicated no growth.

We performed additional phenotypic tests on six carbon substrates; two which were not available on Biolog, 3-(3-hydroxy-phenyl)propionate (Sigma-Aldrich PH011597) and 3hydroxycinnamic acid (CAS 14755-02-3); and four Biolog substrates for which we required further evidence, gamma-amino butyric acid (CAS 56-12-2), L-sorbose (CAS 87-79-6), D-galactarate (CAS 526-99-8), and tricarballylate (CAS 99-14-9). Overnight cultures of all 37 isolates were grown in M9 minimal media (2× M9, Minimal Salts [Sigma-Aldrich], 2 mM MgSO<sub>4</sub>, and 0.1 mM CaCl<sub>2</sub>) plus 20 mM D-glucose, at 37°C, shaking at 200 RPM. Each carbon source substrate solution was prepared to a final concentration of 20 mM in M9 minimal media, pH 7.0. Then, 200 µL of each substrate solution was added to separate 96-well cell culture plates (Corning) and 5 µL of overnight cultures added to the wells, diluted to McFarland standard of 0.4-0.55. Negative controls were included on every independent plate and included (1) no substrate solution controls (20 mM M9 minimal media) and (2) no isolate controls but 20 mM substrate solution. For positive controls, each isolate was also grown independently in M9 minimal media containing 20 mM D-glucose. Every growth condition was performed in technical triplicate. Plates were then sealed with Aera-Seal film (Sigma-Aldrich) and then grown aerobically for 18 h at 37°C, shaking at 200 RPM. Plates were then read using the FLUOstar Omega plate reader (BMG Labtech) using Read Control version 5.50 R4, firmware version 1.50, using 595 nm absorbance after 30 sec of shaking at 200 RPM. No isolate controls were used as blanks to generate the optical density (OD) value for each technical replicate. We then averaged all no isolate OD values to obtain a mean M9 media OD value. To determine growth/no growth using the OD method, we calculated the mean OD for growth on a particular substrate for each strain at 24 h and subtracted from this the OD value of M9 media alone. Subsequently, for each carbon substrate, we divided the mean OD value for a strain by the mean OD for that strain in M9 media alone to get an OD fold change. OD fold changes  $\geq 2$  were considered sufficient evidence of growth (Supplemental Fig. 1).

We performed additional growth tests for independent validation of the models using VITEK 2 GN ID cards (bioMérieux). All 37 strains were assayed on the card to evaluate growth on 13 carbon sources (VITEK codes for those 13 sources can be found in Supplemental Table 5). Cards were read on the VITEK 2 Compact (bioMérieux) per the manufacturer's instructions using VITEK 2 software version 8.0.

#### Creating and curating strain-specific GEMs

Using the method outlined by Norsigian et al. (2019b), we extracted and translated all CDS from each genome and used bidirectional BLASTP hits (BBH) to determine orthologous genes compared to the reference *K. pneumoniae* MGH78578 GEM (iYL1228) (Liao et al. 2011). Genes with at least 75% amino acid identity were considered orthologous. Genes and their reactions that did not meet this threshold were removed from their respective models.

During GEM creation, we discovered that the original biomass function (BIOMASS\_) in iYL1228 required the production of both rhamnose, which is a component of the capsule in *K. pneumoniae* MGH 78578, as well as UDP-galacturonate and UDP-galactose, which are components of the variable O antigen. Because both the capsule and O antigens are known to differ greatly between strains (Follador et al. 2016; Wyres et al. 2016), we created a new biomass function (BIOMASS\_Core\_Oct2019) that no longer required the associated metabolites dtdprmn\_c, udpgalur\_c, and udpgal\_c.

To validate each GEM against its respective phenotypic growth results, we used flux-based analysis (FBA) implemented in the COBRApy framework (Ebrahim et al. 2013) to simulate growth of each GEM in M9 media with all possible sole carbon, nitrogen, phosphorus, or sulfur substrates. The updated BIOMASS function, BIOMASS\_Core\_Oct2019, was used as the objective to be optimized. M9 media was defined by setting the lower bound of the cob(I)alamin exchange reaction to -0.01, and the lower bound of the following exchange reactions to -1000: Ca<sup>2+</sup>, Cl<sup>-</sup>, CO<sub>2</sub>, Co<sup>2+</sup>, Cu<sup>2+</sup>, Fe<sup>2+</sup>, Fe<sup>3+</sup>, H<sup>+</sup>, H<sub>2</sub>O, K<sup>+</sup>, Mg<sup>2+</sup>, Mn<sup>2+</sup>, MoO<sub>4</sub><sup>2-</sup>, Na<sup>+</sup>, Ni<sup>2+</sup>, Zn<sup>2+</sup>. To predict growth on alternate carbon substrates, we set the lower bound of glucose to zero (to prevent the model using this as a carbon source), and then set the lower bound of all potential carbon substrates to -1000 in turn. The carbon substrate was considered growth supporting if the predicted growth rate was  $\geq 0.001$ . The code used to simulate growth on each substrate can be found in Supplemental Code (simulate\_ growth\_single.py).

While identifying carbon substrates, the default nitrogen, phosphorus, and sulfur substrates were ammonium  $(NH_4)$ , inorganic phosphate (HPO<sub>4</sub>), and inorganic sulfate (SO<sub>4</sub>). Prediction of nitrogen, phosphorus, and sulfur supporting substrates was performed in the same way as carbon but setting glucose as the default carbon substrate.

We matched predictions and phenotypic growth data for all strains for 94 distinct carbon substrates. These data were used to (1) curate and update the models and (2) estimate model accuracy. Where we had evidence of phenotypic growth but a lack of simulated growth, we attempted to identify the missing reactions using gene homology searches and literature searches in related bacteria (for a full list of reactions added and the evidence for each, see Supplemental Table 3). During this process it became apparent that the directionality of the following transport reactions in the original iYL1228 GEM were set to export the compound from Strain model accuracy was determined by calculating the percentage of true positive and negative compounds, as well as calculating Matthew's correlation coefficient using the following formula (TP = true positive; TN = true negative; FP = false positive; FN = false negative):

$$\frac{TP \times TN - FP \times FN}{/(TP + FP)(TP + FN)(TN + FP)(TN + FN)}.$$

We assessed accuracy against the VITEK growth data using the same method as described previously. However, these data were used only to estimate model accuracy and were not used to curate or update the models.

#### Gene essentiality for growth on core carbon substrates

To determine which genes were essential for growth in each core carbon substrate (n = 145) for each strain, we used the *single\_gene\_deletion* functions in COBRAPy (Ebrahim et al. 2013). For each GEM, on every core carbon substrate we simulated growth in M9 media with that substrate as the sole carbon source using FBA (as described previously), but with one gene knocked out using the *single\_gene\_deletion* function. Each gene was knocked out in turn, and optimized biomass values  $\geq 0.001$  were considered positive for growth. The code used to perform the knockouts and growth simulations on each substrate can be found in Supplemental Code (single\_gene\_knockouts.py).

Four gene-substrate combinations were selected for further investigation by interrogation of the model gene-protein-reaction rules and search of the KEGG database (Kanehisa et al. 2002) using KofamKOALA (Aramaki et al. 2020) for redundant genes/pathways. When relevant, pairwise chromosomal comparisons were performed using BLASTN (Camacho et al. 2009) and visualized using the Artemis Comparison Tool (Carver et al. 2005). The putative insertion sequence was identified by BLASTN search of the ISFinder database (Siguier et al. 2006).

#### Core genome phylogeny

The core genome for the set of 37 genomes was determined using panaroo v1.1.2 (Tonkin-Hill et al. 2020) in strict mode with a gene homology cutoff of 90% identity, which generated a core gene alignment consisting of 3717 genes with 75,899 variable sites. We generated a phylogeny using this core gene alignment with IQ-Tree v2 (Minh et al. 2020), which selected GTR+F+I+G4 as the best-fit substitution model. The resulting phylogeny was visualized using *ggtree* (Yu et al. 2017) in R (R Core Team 2021).

#### Gene screening in public genomes

To determine whether specific gene deletions or acquisitions are likely to be conserved in all members of a species or clone, we used the curated set of 13,156 *Klebsiella* genome assemblies from Lam et al. (2021). We used BLASTN to screen for (1) the *nan* operon in 307 *K. quasipneumoniae* subsp. *similipneumoniae* genomes; (2) the *ulaABC* operon in 149 *K. quasipneumoniae* subsp. *quasipneumoniae* genomes; (3) *glpQ* in 13 *K. quasivariicola* genomes and six *K. africana* genomes (Vezina et al. 2021); and (4) KPN\_02388 in 1021 nonredundant ST258 genomes. Hits with  $\geq$ 90% coverage and  $\geq$ 90% identity were considered to be present.

#### Data access

All completed genomes and raw sequence data generated in this study have been submitted to the NCBI BioProject database (BioProject; https://www.ncbi.nlm.nih.gov/bioproject/) under accession number PRJNA768294. All strain metabolic models generated in this study have been deposited in JSON format, along with the gene annotations used for the models in figshare (https://doi.org/10.26180/16702840). MEMOTE reports for all models can be found in figshare (https://doi.org/10.26180/19180274.v1).

## Competing interest statement

The authors declare no competing interests.

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Author contributions: J.H., K.E.H., J.M.M., and K.L.W. conceived the study and designed analyses. S.B., C.R., and S.L.-F. provided bacterial isolates and Biolog phenotype data. B.V., L.M.J., T.H., and C.R. generated novel sequence and/or phenotype data. J.H., B.V., and K.L.W. performed data analyses. J.H., J.M.M., S.B., K.E.H., and K.L.W. obtained funding. J.H. and K.L.W. wrote the manuscript. All authors read, commented on, and approved the manuscript.

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