

RESEARCH ARTICLE

Identifying virulence determinants of multidrug-resistant *Klebsiella pneumoniae* in *Galleria mellonella*

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One sentence summary: As *Klebsiella pneumoniae* is often difficult to study in mice, the authors used the wax moth *Galleria mellonella* to describe the global fitness landscape of this important human pathogen.

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ABSTRACT

Infections caused by *Klebsiella pneumoniae* are a major public health threat. Extensively drug-resistant and even pan-resistant strains have been reported. Understanding *K. pneumoniae* pathogenesis is hampered by the fact that murine models of infection offer limited resolution for non-hypervirulent strains which cause the majority of infections. The insect *Galleria mellonella* larva is a widely used alternative model organism for bacterial pathogens. We have performed genome-scale fitness profiling of a multidrug-resistant *K. pneumoniae* ST258 strain during infection of *G. mellonella*, to determine if this model is suitable for large-scale virulence factor discovery in this pathogen. Our results demonstrated a dominant role for surface polysaccharides in infection, with contributions from siderophores, cell envelope proteins, purine biosynthesis genes and additional genes of unknown function. Comparison with a hypervirulent strain, ATCC 43816, revealed substantial overlap in important infection-related genes, as well as additional putative virulence factors specific to ST258, reflecting strain-dependent fitness effects. Our analysis also identified a role for the metalloregulatory protein NfeR (YqjI) in virulence. Overall, this study offers new insight into the infection fitness landscape of *K. pneumoniae*, and provides a framework for using the highly flexible and easily scalable *G. mellonella* infection model to dissect molecular virulence mechanisms of bacterial pathogens.

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Keywords: *Klebsiella pneumoniae*; *Galleria mellonella*; TraDIS; Tn-seq; ST258

INTRODUCTION

Klebsiella pneumoniae is a Gram-negative, capsulated bacterial pathogen responsible for a high proportion of hospital-acquired infections (Podschun and Ullmann 1998; Pendleton, Gorman and Gilmore 2013). *Klebsiella pneumoniae* is part of the so called “ESKAPE” group of human pathogens which are the leading cause of healthcare-associated infections worldwide (Rice 2008; Boucher et al. 2009). Of special concern are carbapenem-resistant *K. pneumoniae*, classified by the WHO as a critical priority for new drug development (Tacconelli et al. 2018). Infections caused by multidrug resistant isolates are associated with significantly higher costs for the healthcare system (Zhen et al. 2019) and in Europe, multidrug resistant *K. pneumoniae* strains cause more than 90 000 infections per year, almost 16 000 of those are caused by carbapenem resistant isolates, leading to more than 7000 deaths per year (Cassini et al. 2019). Classical *K. pneumoniae* (cKp) causes a range of opportunistic infections (e.g. pneumonia, skin/soft tissue and catheter-associated urinary tract infections) in the elderly and immunocompromised, while hypervirulent *K. pneumoniae* (hvKp) causes community-acquired invasive disease. Classical and hypervirulent *K. pneumoniae* can be distinguished by the presence of a specific virulence markers, or by their lethality in mice. Though hypervirulent strains are a serious public health threat, the majority of *Klebsiella* disease burden is currently associated with classical strains (Wyres, Lam and Holt 2020). *Klebsiella pneumoniae* virulence factors include its protective polysaccharide capsule, O antigen, adhesive pili, capsule overproduction regulators and several different siderophores. Of these, the capsule overproduction genes and the siderophores aerobactin and salmochelin are hypervirulence markers that are absent from the majority of cKp isolates.

HvKp strains are mouse-virulent with a lethal dose of less than 10^6 colony forming units (cfu), while cKp strains generally are not (Yu et al. 2007; Russo and Marr 2019; Russo and MacDonald 2020). The use of mouse models to study cKp infections is typically limited to enumeration of surviving bacteria following challenge with very high inocula (for example, (Diago-Navarro et al. 2018; Palacios et al. 2018)). Such models capture intermediate points during self-resolving infections and may miss subtle virulence phenotypes. Furthermore, these models cannot easily be used in conjunction with the functional genomics approaches often used to identify infection-related genes *en masse* (e.g. TnSeq, RNA-seq), as these methods require large numbers of bacteria to provide enough material for sequencing and avoid population bottlenecks (Cain et al. 2020). *In vitro* approaches like cell culture (Shames et al. 2017) or *in vitro* organ culture (Brockmeier et al. 2018) are more amenable to the high bacterial cell numbers needed for functional genomics, but these single cell-type or single organ approaches can be highly technically challenging and lack the complexity of an infection in a eukaryotic host. Due to these difficulties the majority of *K. pneumoniae* pathogenesis studies use infection of mice by hvKp strains as the primary measure of virulence. Studying putative cKp-relevant virulence genes in hvKp strains may fail to identify relevant virulence activities, because mechanisms of pathogenesis vary between *K. pneumoniae* isolates (Xiong et al. 2015) and because even shared *K. pneumoniae* phenotypes can be underpinned by strain-specific gene sets (Dorman et al. 2018; Short et al. 2020).

Alternative infection models for *K. pneumoniae* infections include *Dictyostelium discoideum*, *Drosophila melanogaster*, *Galleria mellonella*, *Caenorhabditis elegans*, zebrafish (reviewed in (Bengoechea and Sa Pessoa 2019)), and the *ex vivo* porcine lung (Dumigan et al. 2019). Of these models, the greater wax moth *Galleria mellonella* larval model is by far the most established. *Galleria mellonella* larvae are susceptible to infection by both cKp and hvKp strains (Li et al. 2020; Russo and MacDonald 2020), and recapitulate many relevant features of mammalian infections (Insua et al. 2013). *Galleria mellonella* larvae used in research are not standardised – larvae are usually obtained from pet food or fishing bait suppliers – so published lethal doses in this model vary widely (e.g. MGH 78578 in (Insua et al. 2013; Wand et al. 2015), reviewed in (Pereira et al. 2020)). Despite this lack of standardisation, the virulence of different *K. pneumoniae* strains in *Galleria* broadly agrees with virulence in mice; though this relationship is not strong enough to reliably differentiate between hvKp and cKp strains in the absence of other information ((Li et al. 2020; Russo and MacDonald 2020), note the latter study used bait-grade larvae). The flexibility of the *Galleria* model, and its susceptibility to cKp infection, makes this a valuable animal model for high-throughput functional genomics studies of cKp.

We have performed transposon directed insertion sequencing (TraDIS) to identify genes in *K. pneumoniae* RH201207 – a multidrug-resistant cKp strain of the global, outbreak-associated clonal group ST258 – that contribute to *G. mellonella* infection, and compared the *in vivo* fitness requirements of this cKp strain to that of the hvKp strain ATCC 43816. Our results identify known virulence genes along with newly identified putative virulence factors, and we have validated the TraDIS screen with defined single-gene mutants. One gene of interest was the Nickel-dependent transcriptional repressor NfeR, which has not previously been linked to virulence in any species. An NfeR mutant showed reduced virulence and increased expression of a neighbouring ferric reductase gene – effects that were reversed by complementation – but, unexpectedly, could not be linked to any *in vitro* phenotypes. Overall, our results show that the *Galleria mellonella* model is well-suited to high-throughput functional genomics studies of cKp strains, and suggest that even very subtle disruptions to metal homeostasis may be important during cKp infections.

MATERIAL AND METHODS

Bacterial strains and culture conditions

Bacterial strains, TraDIS libraries, plasmids and oligonucleotides used in this work are listed in Table 1. *Klebsiella pneumoniae* RH201207 (ST258) and ATCC 43816 (ST493) were grown in LB medium at 37°C with shaking for routine culture. Where necessary, antibiotics were added in the following concentrations: tetracycline 15 µg mL⁻¹, chloramphenicol 25 µg mL⁻¹. Viable counts of bacterial cultures were determined by serial dilution in PBS followed by spot-plating of the entire dilution series with technical duplicates.

Galleria mellonella infection experiments

Research-grade *G. mellonella* larvae at their final instar stage (TruLarv; Biosystems Technology Ltd, Exeter, UK) were kept at room temperature (approx. 20°C) in the dark for a maximum of

Table 1. Bacterial strains, TraDIS libraries, plasmids and primers used in this study.

Bacterial strains		
Strain	Description/genotype (Genome accession no)	Source
<i>K. pneumoniae</i> ATCC 43816	Hypervirulent, ST493, K-type 2, commonly used in mouse studies (CP009208.1)	Isolate: American Type Culture Collection; Genome: (Broberg et al. 2014)
<i>K. pneumoniae</i> RH201207	Colistin-resistant UK gut isolate, ST258, K-type 106	Isolate: (Jana et al. 2017); Genome: This study
<i>Escherichia coli</i> β 2163	F ⁻ RP4-2-Tc::Mu DdapA::(<i>erm</i> -pir)	(Demarre et al. 2005)
<i>K. pneumoniae</i> RH201207	Tn5- <i>rfaH</i>	(Short et al. 2020)
<i>K. pneumoniae</i> RH201207	Tn5- <i>wza</i>	(Short et al. 2020)
<i>K. pneumoniae</i> RH201207	Tn5- <i>wzxE</i>	This study
<i>K. pneumoniae</i> RH201207	Δ <i>nfeR</i> (KPNRH.00645)	This study
<i>K. pneumoniae</i> RH201207	Δ <i>nfeR</i> (KPNRH.00645) complemented on chromosome	This study
<i>K. pneumoniae</i> RH201207	Δ <i>phoQ</i> (KPNRH.03335)	This study
Transposon mutant libraries		
Parent strain	Transposon (primers)	Source
<i>K. pneumoniae</i> RH201207	Tn5, TetR (Tn5tetR-5Seq, Tn5tetR-5PCR)	(Jana et al. 2017)
<i>K. pneumoniae</i> ATCC 43816	Tn5-derived from pDS1028 vector, CmR (FS107, FS108)	(Dorman et al. 2018)
Plasmids		
Name	Description	Source
pKNG101-Tc	Allelic exchange vector, Tc ^R	(Poulter et al. 2011)
pFLS27	RH201207 <i>nfeR</i> knockout vector, pKNG101-Tc-derived, constructed with FS273-276	This study
pFLS28	RH201207 <i>phoQ</i> knockout vector, pKNG101-Tc-derived, constructed with FS277-280	This study
pFLS36	RH201207 <i>nfeR</i> complementation vector, pKNG101-Tc-derived, constructed with FS273-274	This study
Primers		
Name	Sequence 5' – 3'	Description
FS107	GAGCTCGAATTCATCGATGATGGTTGAGATGTGTA	pDS1028 TraDIS 5' Seq
FS108	AATGATACGGCGACCACCGAGATCTACACCAGGAACACTTAACGGCTGACATGG	pDS1028 TraDIS 5' PCR
Tn5tetR-5PCR	AATGATACGGCGACCACCGAGATCTACACTGTGATAAACCACCGATTAAAGCTTATCG	Tn5-Tet TraDIS 5' PCR
Tn5tetR-5Seq	CGATGATAAGCTGTCAAACATTGATGGTTGAGATGTGTA	Tn5-Tet TraDIS 5' Seq
FS273	gctACTAGTTGCCCTACATTGAAGATGC	Mutant construction, SpeI site
FS274	gctACTAGTCGTAAGCCAGCTCGTGA	Mutant construction, SpeI site
FS275	GGTGGTGAAAAAAGAGGCGGTTACCTCCTGCTGTTTT	Mutant construction, overlap
FS276	ACAGCAGGAGGTAACCGCCTCTTTTTTCACCACCGGC	Mutant construction, overlap
FS277	gctACTAGTCGGGCAGTGCTGTTTCA	Mutant construction, SpeI site
FS278	gctACTAGTGCGAACATCTCCCGGAT	Mutant construction, SpeI site
FS279	GTGAAAAATCTCAACGACAGCGCAATTCGAACAGAT	Mutant construction, overlap
FS280	TCGAATTGCGCTGTCGTTGAGATTTTTTCACGGCG	Mutant construction, overlap
FS133	TTAAACAGGCCGAATTCACG	qRT-PCR <i>recA</i>
FS134	CCGCTTTCTCAATCAGCTTC	qRT-PCR <i>recA</i>
FS135	TCGCCGATGCGTCGTATAAA	qRT-PCR <i>yqjH</i>
FS136	TGCTGATCGATGCTCTCGTC	qRT-PCR <i>yqjH</i>

seven days before use. Larvae were reared under standardised conditions, age and weight defined and decontaminated prior to shipping (Champion, Titball and Bates 2018). Injections and haemolymph extractions were performed as described (Harding et al. 2013). For survival analyses, *K. pneumoniae* strains were grown to late exponential phase ($OD_{600} = 1$, approx. cfu 8×10^8 RH201207; 5×10^8 ATCC 43816), harvested by centrifugation, and resuspended in sterile PBS. 10 μ L doses of diluted bacterial suspensions were injected into the right hind proleg of the larvae using 30-gauge hypodermic needles (BD Diagnostics, Wokingham, UK), and the infected larvae were incubated in the dark without food at 37°C for up to 72 hours. Larvae were examined every 24 hours and were scored as dead when they were unresponsive to touch.

TraDIS infection experiments were performed in biological triplicate using previously reported high-density mutant libraries (Jana et al. 2017; Dorman et al. 2018). Aliquots of frozen

pooled transposon mutant libraries (minimum of 10^8 cells) were grown overnight, subcultured and grown to late exponential phase, then resuspended and diluted in PBS to an approximate density of 10^7 cfu mL⁻¹. Groups of larvae were injected with 10 μ L prepared TraDIS library per larva (approx. dose 10^5 cfu) and incubated at 37°C in the dark. Bacteria were extracted from infected *Galleria* as follows: haemolymph was recovered from all of the larvae in each group and pooled in four volumes of ice-cold eukaryotic cell lysis buffer (1 \times PBS + 1 % Triton X-100) and the mixture was held on ice for ten minutes. Treated haemolymph was centrifuged at $250 \times g$ for 5 minutes to pellet eukaryotic cell debris while leaving the majority of bacterial cells in the supernatant. The supernatant was centrifuged at $8000\text{--}10000 \times g$ for 2 minutes to pellet bacterial cells, and bacteria were resuspended in 5 mL LB and outgrown at 37°C to approx. OD_{600} of 1 in order to generate enough material for gDNA extraction and sequencing. Bacteria were grown for a limited number of

generations to late exponential phase to increase cell numbers and therefore genomic material without altering relative mutant frequencies. The final post-infection time points were chosen as the time where there was visible melanisation of the majority of larvae in each group, but no reduction in the volume of recoverable haemolymph. This corresponded to an endpoint per-larva bacterial load of $\sim 2 \times 10^7$ cfu (Fig 1B).

Specific parameters for *G. mellonella* TraDIS infection experiments were as follows. RH201207: 20 larvae per group, 1.5×10^5 cfu inoculum, time points at 2 hours post-infection (hpi) and 6 hpi, outgrowth periods 4 hours (2 hpi samples) and 1.5 hours (6 hpi samples); ATCC 43816: 12 larvae per group, 1.1×10^5 cfu inoculum, time point at 4 hpi, outgrowth period 2 hours. See also Fig 1B for per-larva bacterial loads over the course of the experiment.

Genome re-sequencing and annotation of RH201207

Genomic DNA of RH201207 was extracted using the MasterPure Complete DNA and RNA Purification Kit (Lucigen, Middleton, WI, USA) with DNA resuspended in 50 μ L nuclease free water by carefully flicking the tube. Purity was checked on a NanoDrop spectrophotometer (Thermo Fisher, Waltham, MA, USA) with values of 260/280 nm of 2.01, and 230/260 nm of 2.12; quality and quantity was assessed on a TapeStation (Agilent, Santa Clara, CA, USA) with DIN of 9.7 and concentration of 36.9 ng μ L⁻¹.

Nanopore 1D sequencing library was prepared using the genomic DNA by ligation sequencing kit SQK-LSK109 (Oxford Nanopore Technologies (ONT), Littlemore, UK), barcoded using the barcoding extension kit EXP-NPB104 and sequenced on a GridION X5 using a R9.4.1 flow cell (ONT) together with five bacterial genomes from a different study. Bases were called with Albacore v2.0 (ONT) and adapter sequences were trimmed and sequencing reads de-multiplexed with Porechop v0.2.3 (<https://github.com/rwick/Porechop>). Genome assembly was performed in combination with the previously described paired-end Illumina reads of RH201207 (Jana et al. 2017) accessible at the ENA (study PRJEB1730). The hybrid read set was assembled with Unicycler v0.4.7 (Wick et al. 2017) using the normal mode and assembly graphs were visualised with Bandage (Wick et al. 2015). The final assembly was annotated using Prokka v1.14.5 (Seemann 2014) with additional functional gene annotation by KEGG (Kanehisa and Goto 2000) and UniProt (The UniProt Consortium 2019). Plasmid replicons were identified with PlasmidFinder (Carattoli et al. 2014). Typing of the *Klebsiella* K- and O-loci was performed with Kaptive Web (Wick et al. 2018). Iron uptake genes were annotated with SideroScanner (<https://github.com/tomdstanton/sideroscanner>).

TraDIS sequencing and analysis.

Genomic DNA was extracted by phenol-chloroform extraction. At least 1 μ g DNA per sample was prepared for TraDIS as described in (Barquist et al. 2016). Sequencing statistics and accession numbers are given in Table S1 (Supporting Information). Sequencing reads were mapped to the RH201207 or ATCC 43816 genomes using the Bio::TraDIS pipeline as described previously (Langridge et al. 2009; Barquist et al. 2016), with a 96 % mapping threshold, multiply-mapped reads discarded and one transposon tag mismatch allowed (script parameters: “-v -smalt.y 0.96 -smalt.r -1 -t TAAGAGACAG -mm -1”). Insertion sites and reads were assigned to genomic features with reads mapping to the 3' 10% of the gene ignored, and between-condition comparisons were performed without read

count filtering. Gene essentiality was determined by running tradis.essentiality.R (Barquist et al. 2016) on the combined gene-wise insertion count data of all three biological replicates. Essential or ambiguous-essential genes of the input samples (the initial transposon library) as defined by the Bio::TraDIS pipeline were excluded from further analysis. Gene functional categories and Cluster of Orthologous Groups (COG) (Tatusov et al. 2000) were assigned to genes using EggNOG Mapper (Huerta-Cepas et al. 2017) and enrichment of clusters of orthologous groups (COG) was determined by two-tailed Fisher's exact test in R version 3.6.2 (R-function fisher.test) (R Core Team 2019).

Mutagenesis and complementation

Details of all plasmids, strains and oligonucleotides used are given in Table 1. Markerless single-gene deletion mutants of *K. pneumoniae* RH201207 *phoQ* (RH201207_003326) and *nfeR* (RH201207_00640) were generated by allelic exchange mutagenesis using vectors derived from pKNG101-Tc as described (Dorman et al. 2018). Mutagenesis vectors containing the up- and downstream sequences for each target gene (>500 bp) were constructed by overlap extension PCR and delivered by conjugation with *Escherichia coli* β 2163. Complementation of *nfeR* was performed by restoring the gene in its native location by allelic exchange mutagenesis of *K. pneumoniae* RH201207 Δ *nfeR* with pFLS36 (Table 1). Transposon-insertion mutants of *K. pneumoniae* RH201207 were generated by subjecting the TraDIS mutant library to two rounds of density-gradient selection then identifying mutants by random-primed PCR as described (Short et al. 2020).

Klebsiella pneumoniae phenotypic tests

All quantitative phenotypic tests reported are from three biological replicates. Serum susceptibility was determined by incubation of late log-phase cells with 66 % normal human serum (Sigma-Aldrich, St. Louis, MO, USA) as described (Short et al. 2020). Siderophore production was visualised by the chrome azurol S assay (Schwyn and Neilands 1987; Loudon, Haarmann and Lynne 2011) with the following measures to remove trace iron: glassware was soaked in 6 M HCL for two hours and rinsed three times with ultrapure water, and Casamino acid solution was treated with 27 mL 3 % 8-hydroxyquinoline in chloroform for 20 minutes, then the supernatant was removed, extracted once with a 1:1 volume chloroform, and filter-sterilised. Overnight cultures of *K. pneumoniae* strains were normalised to an OD₆₀₀ of 0.5 prior to spotting on CAS agar, and plates were incubated at 37°C for 48 hours. For qRT-PCR, total RNA was extracted from bacteria grown in LB medium to OD = 1 using an RNeasy kit (Qiagen, Hilden, Germany), and 2 μ g RNA was treated with TURBO DNase (Thermo Fisher). TURBO DNase was removed from treated RNA using the supplied inactivation reagent according to the manufacturer's instructions. Transcripts of *nfeF* and the *recA* housekeeping gene were quantified using a KAPA SYBR FAST one-step qRT-PCR master mix (Sigma-Aldrich) according to the manufacturer's instructions. Three biological replicates and two technical replicates were performed, with 2.5 ng total RNA per reaction. Sensitivity to hydrogen peroxide was determined by diluting stationary phase cultures 1:100 in Mueller-Hinton II (Oxoid, Basingstoke, UK) medium, then adding hydrogen peroxide (30 % v/v, Sigma-Aldrich) to a final concentration of 4–8 mM. Samples were incubated at 37°C for 120 minutes before serial dilution and plating to enumerate surviving bacteria. Nickel toxicity and dipyriddy sensitivity tests

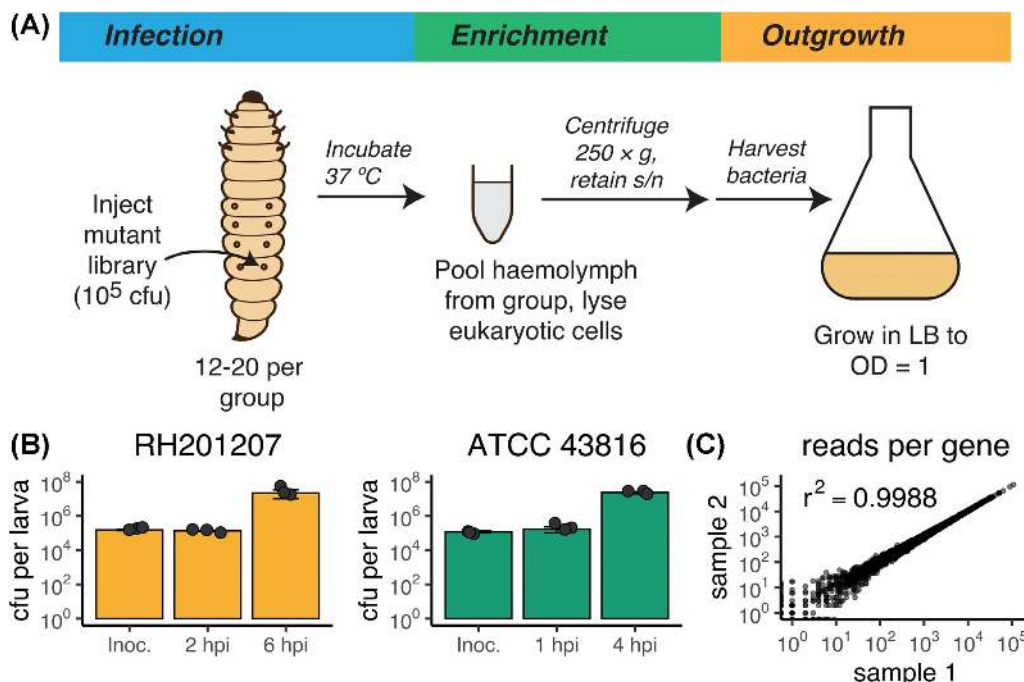


Figure 1. Overview of transposon insertion screening in *Galleria mellonella* larvae **A:** Schematic of experimental procedure. Groups of larvae were injected with a pooled library of *Klebsiella pneumoniae* transposon mutants and incubated at 37°C for up to 6 hours. Haemolymph was extracted from infected larvae, pooled and treated to remove eukaryotic cells. Surviving bacteria were then grown in LB to generate sufficient material for sequencing. **B:** Per-larva bacterial counts over the course of *G. mellonella* infection. Bacteria were recovered when the load per larva reached approximately 10^7 cfu. The experiment was done in triplicates, closed circles represent one measurement, error bars denote standard deviation. Abbreviation: Inoc., inoculum. **C:** Reads per gene of two biological replicates of the unchallenged RH201207 input transposon library and their Pearson correlation coefficient.

were performed in a 96-well plate format with 100 μ l volume per well and an initial cell density (seeded from overnight culture) of $OD_{600} = 0.05$. Nickel toxicity tests were performed in LB medium supplemented with nickel(II) sulfate hexahydrate (Sigma-Aldrich). Dipyrindyl sensitivity tests were performed in M9 minimal medium supplemented with 0.2 % glucose, with the inoculum washed three times in M9 salts. Plates were sealed with air-permeable film and incubated with shaking for 18 hours at 37°C prior to measurement of OD_{600} .

Availability of sequencing data

The TraDIS sequencing data is available in the European Nucleotide Archive (<https://www.ebi.ac.uk/ena/browser/home>) under the Study Accession No. PRJEB20200. Individual sample accession numbers are available in Table S1 (Supporting Information). Oxford Nanopore reads of RH201207 are available in the ENA repository under study accession number PRJEB40551.

RESULTS AND DISCUSSION

TraDIS analysis of *G. mellonella* infection determinants in *K. pneumoniae* RH201207

Klebsiella pneumoniae RH201207 is a multidrug-resistant isolate of clonal group CG258 used in previous transposon insertion sequencing studies (Jana et al. 2017; Short et al. 2020). We first measured *K. pneumoniae* RH201207 infection parameters in *G. mellonella*. In order to minimise variability between larvae and improve the sensitivity of the TraDIS analysis, research-grade *G. mellonella* larvae were used; these are age- and weight-standardised, genetically identical, and reared without antibiotics or hormones. An inoculum of $\sim 10^5$ cfu was sufficient to

kill the majority of infected larvae, with melanisation evident at 5 hours post infection. For TraDIS analysis, three groups of *G. mellonella* larvae were infected with replicate cultures of the *K. pneumoniae* RH201207 mutant pool, and we recovered and sequenced surviving bacteria from the larval haemolymph at 2 hpi and 6 hpi. The volume of haemolymph that could be recovered declined at later time points due to melanisation of the larvae; while reducing the inoculum to increase the infection time would be possible, this could compromise the experiment by introducing a population bottleneck. Haemolymph was treated with detergent to lyse eukaryotic cells, then bacteria were recovered and grown in rich medium (Fig. 1A). This method allowed high recovery of infecting bacteria without antibiotic selection, while avoiding co-isolation of DNA from either the host or its gut or skin microbiota (Allonsius et al. 2019). Viable counts were measured on infection and at each sampled time point, and showed approximately seven generations of bacterial replication at 6 hpi (Fig. 1B). Our infection screening parameters therefore allow identification of mutations that impair replication in this host, as well as mutations that cause sensitivity to killing by *G. mellonella* immune system components.

Re-sequencing of RH201207

During the initial analysis of the TraDIS experiments, we identified a possible mis-assembly of the original RH201207 genome, which could not be improved by optimising the assembly parameters. To improve the assembly and generate a circularised chromosome sequence, we sequenced RH201207 by long-read Oxford Nanopore Technologies (ONT) sequencing. Nanopore GridION sequencing yielded 2.3 G bases with an average read length of 10.5 kb and maximum read length of 165.9 kb, corresponding to

~400-fold theoretical coverage. Hybrid assembly of these reads together with existing MiSeq reads from Jana *et al.* (1.24 million reads, ~35-fold genome coverage) yielded a circularised chromosome of 5475789 bp and three circularised potential plasmids (Table S2, Supporting Information) of 113640 bp (contig RH201207_2, IncFIB(pQil) and IncFII(K) replicons), 43380 bp (contig RH201207_5, IncX3 replicon) and 13841 bp length (contig RH201207_8, ColRNAI replicon). The remaining contigs of 202245 bp length could not unambiguously be closed and circularised due to the presence of potential duplicated sequences. Nonetheless, typical plasmid replication proteins and the two replicons IncFIB(K) and IncFII(K) were present on these contigs, indicating that they are likely to be derived from plasmids (Table S2 and Fig. S1, Supporting Information). The re-annotated RH201207 genome has 5787 genes in total (5546 protein-coding) and encodes a capsule of type KL106 and an O-antigen of type O2v2 as determined by Kaptive Web (Wick *et al.* 2018).

Identification of infection-related genes

TraDIS sequencing, read mapping and quantification of each transposon insertion site was performed using the Bio::TraDIS pipeline (Barquist *et al.* 2016). Each sample yielded from 12.9 to 15.1 million transposon-tagged reads, > 89 % of which unambiguously mapped to the RH201207 chromosome (excluding unscaffolded contigs and plasmids) (Table S1, Supporting Information). Analysis of the unchallenged RH201207 TraDIS library showed a total of more than 500 100 unique transposon insertion sites distributed across the chromosome, which corresponds to one insertion in every 11 nucleotides (Table S3, Supporting Information). About 638 genes (or 11.84 % of the 5390 chromosomal genes) were either essential or ambiguous-essential as defined by the Bio::TraDIS pipeline (Barquist *et al.* 2016), which is in good concordance with the first description of this library (Jana *et al.* 2017). A challenge in applying highly saturated transposon insertion libraries to infection models is the occurrence of bottlenecks, that is a stochastic drastic reduction in population size, which results in reduced genetic diversity of the population (Cain *et al.* 2020). We found a slight reduction in the diversity of the mutant library post-infection, with recovered unique insertion sites of nearly 380 000 and 360 000 total insertions (of 500 100) at 2 and 6 hpi, respectively, however transposon insertion density remained very high with an insertion approximately every 15 bp (Table S3, Supporting Information). To test if the loss of mutant library diversity had compromised the resolution of our experiment, we compared individual replicates using linear correlation analyses. Pearson correlation coefficients between *in vivo* replicates were very high with r^2 values greater than 0.98 when analysing reads per gene and insertion indices per gene, and 0.69 to 0.77 when analysing the reads per unique insertion site (Fig. 1; Fig. S2, S3 and Table S4, Supporting Information). Therefore, although our TraDIS experiments show a mild bottleneck, this is highly unlikely to affect any downstream analyses that use gene-level metrics. To identify *in vivo* fitness genes, the numbers of transposon insertion reads within each gene were compared between the *Galleria* infection and the inoculum pools, with essential and ambiguous-essential genes excluded to reduce false positives. Our analysis identified mutants of 133 (of 4752) nonessential genes to be significantly less abundant at 6 hpi and two features, *rseA* and KPNRH.05271 to be slightly, but significantly enriched (Fig. 2 and Table S5, Supporting Information). About 35 genes were so severely depleted after infection that they can be considered conditionally essential in *G. mellonella* (Table S5, Supporting Information).

Surface polysaccharides, cell envelope and iron acquisition genes are critical to cKp *G. mellonella* infection

Capsule, lipopolysaccharide and enterobacterial common antigen

Capsule and lipopolysaccharide (LPS) are essential to the virulence of *K. pneumoniae* (Podschun and Ullmann 1998). The capsule allows *K. pneumoniae* to circumvent host detection and prevent an early immune response; acapsular mutants are less virulent in mouse models and are unable to spread systemically (Paczosa and Mecsas 2016). The lipopolysaccharide, consisting of lipid A, core and O antigen is able to bind and sequester parts of the complement system (Paczosa and Mecsas 2016). Both surface polysaccharides also play a role in modulating innate immunity (Bengoechea and Sa Pessoa 2019). Following *G. mellonella* infection, the eight genes with the most dramatic reduction in read counts (log2 fold-change < -12) all belonged to the capsule or O-antigen loci (Fig. 2B & C), and almost all genes in these clusters were significantly depleted (Fig. 2D–F).

Four genes (*wzxE*, *wecD*, *rffH*, *wecA*) of the Enterobacterial common antigen (ECA) synthesis locus (encoded by genes *rho*/KPNRH.05213 to *yifK*/KPNRH.05200) were also significantly depleted at 6 hpi (Fig. 2B and C). The ECA is a conserved carbohydrate antigen common to most Enterobacterales and plays an important role in bacterial physiology and its interaction with the environment (Rai and Mitchell 2020). *Klebsiella pneumoniae* ECA synthesis mutants were attenuated in *in vitro* murine lung and spleen tissues infections, but not *in vivo* in a murine intranasal infection model (Lawlor *et al.* 2005). Therefore, our analysis identified mutants of all three major polysaccharide antigens (O, K and ECA) present on the cell surface (Rai and Mitchell 2020) to be attenuated in *G. mellonella*.

Cell envelope proteins and regulators

Genes that encode cell envelope proteins appeared to be important during *G. mellonella* infection. The major outer membrane lipoprotein Lpp (also known as Braun's lipoprotein) encoded by *lpp*/KPNRH.02211, and the genes of the Tol-Pal system (*pal*/KPNRH.03757, *tolB*/KPNRH.03758, *tolA*/KPNRH.03759 and *tolQ*/KPNRH.03761) (located at 3.8 Mbp) were significantly depleted at 6 hpi. The Tol-Pal system is involved in maintaining outer membrane integrity and consists of five proteins: TolA, TolQ, and TolR in the inner membrane, TolB in the periplasm, and the peptidoglycan-associated lipoprotein Pal anchored to the outer membrane (Lloubès *et al.* 2001). Lpp is a crucial protein in the outer membrane covalently linking it with the peptidoglycan layer (Asmar and Collet 2018) and it is important for complement resistance in multiple, phylogenetically distinct *K. pneumoniae* strains (Short *et al.* 2020). Mutations in both Lpp and Tol-Pal have been previously linked to attenuated virulence in diverse bacteria (Sha *et al.* 2008; Godlewska *et al.* 2009; Asmar and Collet 2018). One of the most abundant proteins in the outer membrane is the porin OmpK36 (Hernández-Allés *et al.* 1999) and mutants thereof were attenuated in a *G. mellonella* model (Insua *et al.* 2013) and a pneumonia mouse model (March *et al.* 2013). Likewise, our experiments identified a significant underrepresentation of transposon insertion mutants in *ompK36* (KPNRH.01613) at 6 hours post-infection.

Also required for infection were regulators of cell envelope composition and integrity. Mutants of *phoPQ* were significantly less abundant following infection. PhoP-PhoQ is a two-component system, comprising the inner membrane sensor PhoQ and the cytoplasmic regulator PhoP. This system regulates

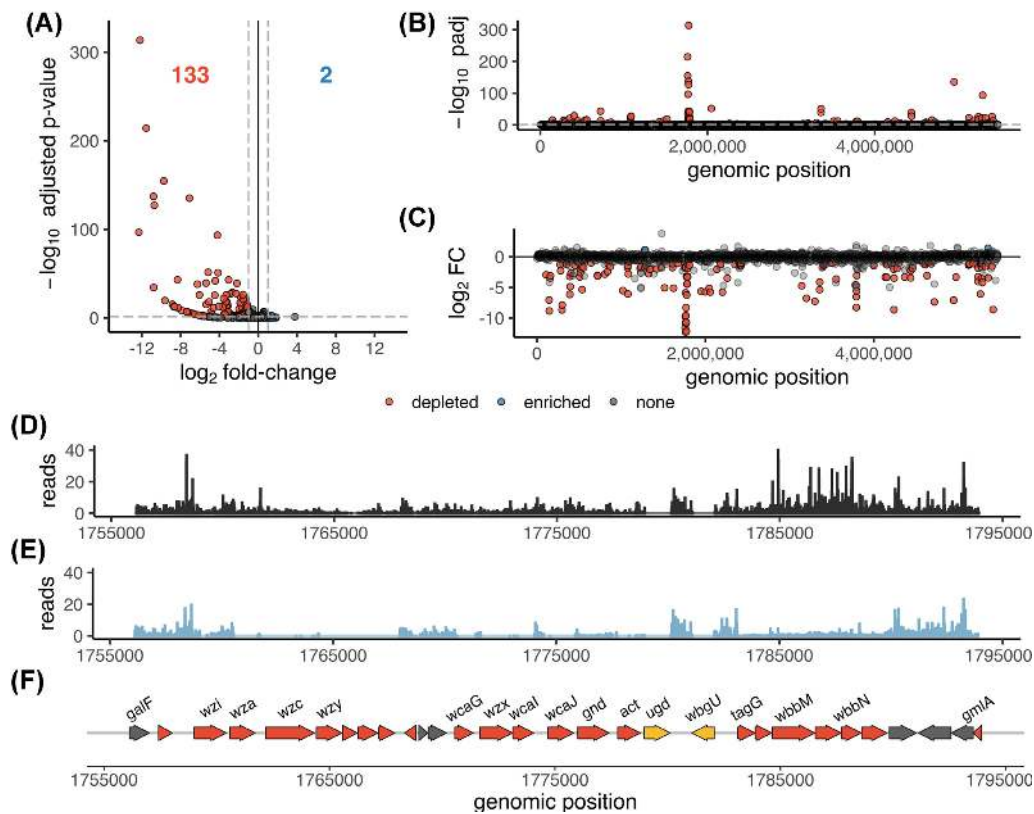


Figure 2. TraDIS analysis identifies multiple potential *K. pneumoniae* fitness factors important during *G. mellonella* infections. (A), Volcano plot showing significantly less abundant genes 6 hours after infecting *G. mellonella* larvae in red and significantly higher abundant genes in blue. Non-significant genes are shown in grey. Genes were considered significant if they possessed a Benjamini-Hochberg corrected P-value below a threshold of 0.05 and an absolute log₂ fold-change greater than 1. Essential and ambiguous-essential genes are removed from the analysis. (B and C), Manhattan plots with Benjamini-Hochberg corrected P-value and log₂ fold-change on the y-axis, respectively. Abbreviations: padj, Benjamini-Hochberg corrected p-value; FC, fold-change. (D and E), Reads per nucleotide of the inoculum and at 6 hpi, respectively of the chromosomal region containing the K- and O-locus. For this graphical representation, triplicate TraDIS samples were combined and reads were normalised to the total number of reads in all three replicates. (F), Gene annotations of the *Klebsiella* K-locus (*galF* to *wbgU*) and O-locus (*tagG* to *gmlA*), encoding the polysaccharide capsule and lipopolysaccharide O antigen, respectively are coloured according to the TraDIS results at 6 hours post-infection; significantly depleted genes are shown in red, essential genes which were not included in the analysis in yellow and genes without significant changes in grey.

lipid A remodelling in *K. pneumoniae* *in vivo* and *in vitro* (Llobet et al. 2015) and other virulence-associated genes in many enteric pathogens (Groisman 2001; Bijlsma and Groisman 2005; Alteri et al. 2011; Lin et al. 2018). *Salmonella* Typhimurium knockout strains of *phoP* and *phoQ* are highly attenuated for virulence in macrophages and a mouse infection (Miller, Kukral and Mekalanos 1989) and this two-component system also makes a small contribution to *K. pneumoniae* virulence during *G. mellonella* infections (Insua et al. 2013). We also noted significant changes in mutant abundance of genes related to the alternative sigma-factor RpoE (σ^{24} or σ^E), one of the major regulators of cell envelope stress response systems (Treviño-Quintanilla, Freyre-González and Martínez-Flores 2013, Flores-Kim and Darwin 2015; Roncarati and Scarlato 2017). RpoE activity is tightly controlled by a proteolytic cascade after dissociation of RseB from RseA (the anti-sigma factor which binds RpoE), RseA is partially cleaved by the proteases DegS and RseP, then fully degraded by other cellular proteases such as ClpP/X-A, Lon and HslUV, releasing RpoE in the cytoplasm (Roncarati and Scarlato 2017). While *rpoE* itself and *rseP* are essential genes and therefore not included in our analysis, transposon insertion mutants of *degS*, *clpP*, *clpX* and *lon* were significantly less abundant, highlighting the role of RpoE gene regulation during infection. Transposon mutants of the RpoE inhibitor RseA and insertions in the promoter of the RseA-degrading protease HslUV (KPNRH.05271)

were enriched after infection, indicating that increased RpoE signalling can enhance fitness in *G. mellonella*. Both RseA and HslUV are, via RpoE, involved in stress response and the regulation of virulence genes in multiple Enterobacteriaceae species (Flores-Kim and Darwin 2015). Interestingly, none of the RpoE regulated cell envelope stress response systems CpxAR, BaeRS, Rcs and Psp (Flores-Kim and Darwin 2015) were identified in our screen, indicating either a redundancy in these systems or the involvement of other factors. Such factors could be the periplasmic chaperones Skp and SurA; both are members of the RpoE regulon in *E. coli* (Dartigalongue, Missiakas and Raina 2001) and have previously been shown to be involved in pathogenicity in *E. coli*, *Salmonella* Typhimurium, *Shigella flexneri* and *Pseudomonas aeruginosa* (Sydenham et al. 2000; Redford and Welch 2006; Purdy, Fisher and Payne 2007; Klein et al. 2019). Mutants of *skp* and *surA* were less abundant in the *Galleria* TraDIS output.

Iron acquisition systems

Iron acquisition is essential during *K. pneumoniae* infections (Paczosa and Mecsas 2016), and RH201207 produces the siderophores enterobactin and yersiniabactin to chelate host iron. Because siderophores are 'common goods' that can be utilised by any bacterium possessing a suitable uptake system, loss of siderophore biosynthesis genes is not expected to influence

fitness in a mutant population where the majority of bacteria still produce siderophores. Siderophore biosynthesis genes were not identified by the *Galleria* infection TraDIS. Utilisation of enterobactin was, however, important: mutations in three of the four ferric iron-enterobactin uptake complex components *tonB* (KPNRH.02155), *exbB* (KPNRH.00722), and *exbD* (KPNRH.00723) reduced fitness in *G. mellonella*. TonB is essential for pathogenicity in multiple bacteria, including hypervirulent *K. pneumoniae* (Hsieh et al. 2008). The fourth component of this complex, *fepB*, was present in multiple copies in the RH201207 genome, so presumably mutation of just one gene does not cause loss of function. Mutation of the yersiniabactin receptor gene *fyuA* did not impair fitness, suggesting that enterobactin is sufficient to allow bacterial replication in the *G. mellonella* haemolymph. This is consistent with the roles of *K. pneumoniae* siderophores in murine infections, where enterobactin alone allows replication in serum by sequestering iron from transferrin (Bachman et al. 2012). Transferrin is also present in the haemolymph of *G. mellonella* (Vogel et al. 2011).

Further genes involved in *Galleria mellonella* fitness

Additional metabolic and hypothetical genes also contributed to fitness during *Galleria* infection. These included the majority of genes in the *aro* operon for synthesis of chorismite (a precursor to aromatic amino acids), *cys* genes for sulphate assimilation and cysteine biosynthesis, purine biosynthesis genes, several components of the electron transport complex, and *sspAB* stringent response proteins. Only 11 genes of unknown function were identified as infection-related (note some K- and O-locus genes were initially annotated as hypothetical). Several transcription regulators were also identified that compromised fitness when mutated: these were the arginine/lysine synthesis repressor *argP*, the fatty acid regulator *fabR*, the zinc-dependent repressor *nrdR* and the nickel-dependent repressor *yjiJ*.

Temporal analysis of the *K. pneumoniae* RH201207 *G. mellonella* infection

We also analysed transposon mutant fitness at 2 hpi, in order to gain insight into the events of early infection, and distinguish mutations influencing survival in the presence of *G. mellonella* immune system components from those influencing replication in this host. Mutants in 54 genes were significantly less abundant at 2 hpi, and only insertions in the cell division protein *FtsB* were enriched (Fig. 3; Fig. S4-S6 and Table S5, Supporting Information). *FtsB* is a conserved, essential transmembrane protein involved in bacterial cell division and part of the so-called “divisome” (den Blaauwen, Hamoen and Levin 2017). Manual examination of the insertion plot files showed that the *ftsB* transposon insertions were all located close to the 3' end of the gene, just outside of our inclusion cut-off of 90% of gene length, therefore it is likely that the enrichment of *ftsB* is based on insertions that do not inactivate the gene. Comparison of both timepoints showed 18 genes depleted only at 2 hpi, 36 genes less abundant at both timepoints and 97 genes for which mutant abundance was reduced only at 6 hours post-infection (Fig. 3A). The genes that were depleted at both time points consists of two subsets: 25 genes that are depleted within the first 2 hours and whose abundance does not change further, and 11 genes that are depleted at 2 hpi and then further decrease in abundance over time. Ten out of these 11 genes are part of the K-locus, the other one is *tagG*, the first gene of the neighboring O antigen locus.

Of the 18 genes less abundant only at the beginning of the infection, most were barely within our threshold for fitness-related genes. But six genes had log₂ fold-changes of -3 up to -7, among them two tRNAs, one hypothetical protein and the three genes *oxyR* (KPNRH.05248), *mhA* (KPNRH.04380) and *tadA* (KPNRH.01302). It is very likely that the tRNAs are false positives due to their very short length and therefore low number of insertions. *OxyR* is a conserved LysR-type transcription factor which plays a key role in the regulation of defence mechanisms against oxidative stress (Christman, Storz and Ames 1989), and has previously been linked to *K. pneumoniae* pathogenesis (Hennequin and Forestier 2009). *RnhA* is the ribonuclease HI that cleaves RNA of RNA-DNA hybrids and is involved in DNA replication, DNA repair and RNA transcription (Kochiwa, Tomita and Kanai 2007) and *tadA* encodes a tRNA-specific adenosine deaminase. This protein is responsible for adenosine to inosine RNA editing of tRNAs and mRNAs, and is involved in the regulation of a toxin-antitoxin system in *E. coli* (Bar-Yaacov et al. 2017).

Of all significantly less abundant insertion mutants, by far the largest set, consisting of 97 genes, is depleted only at the later timepoint 6 hours after infection. This group of genes contains for example the two-component system *phoPQ*, the Tol-Pal system and the majority of genes of the O antigen cluster. Our data indicates that these genes might only have a minor impact on fitness during very early stages of infection.

Shared *in vivo* fitness determinants in cKp RH201207 and hvKp ATCC 43816

We then sought to compare the *G. mellonella* fitness landscape of the cKp ST258 strain RH201207 with a representative hvKp strain. *G. mellonella* infection TraDIS was performed with the well-studied mouse-virulent strain ATCC 43816, which is an ST493 strain possessing type O1v1 O antigen and K2 capsule type. Experiments were performed in the same way as for RH201207 except that bacteria were recovered for sequencing at 4 hpi, due to the faster progression of infection when using this strain (Fig. 1B). Each sample of ATCC 43816 yielded from 1.85 million to 2.08 million reads, more than 96 % of which were reliably mapped to the chromosome of ATCC 43816 KPPR1 (CP009208.1) (Table S1, Supporting Information). The insertion density of the unchallenged ATCC 43816 mutant library was 415,000 or one insertion per 13 nucleotides, which is similar to the RH201207 library (Tables S3), and 502 genes were classified as essential or ambiguous-essential (9.62 % of 5217 genes). Reproducibility between experimental replicates was very high in this experiment, with Pearson correlation coefficient > 0.97 for reads or insertion indices per gene, and > 0.91 when comparing reads per unique insertion site (Fig. S2, S3 and Table S4, Supporting Information).

We identified 92 nonessential genes of *K. pneumoniae* ATCC 43816 that had significantly lower mutant abundance after the *Galleria* infection and two genes (VK055_RS06420 and VK055_RS19345) that were enriched (Fig. S4-S6 and Table S6, Supporting Information). Mutants in 10 genes of the 18-gene K-locus of ATCC 43816 were significantly depleted after *G. mellonella* infection, thereby mirroring the results of RH201207. In contrast, genes of the O-locus were not implicated in *in vivo* fitness in *G. mellonella*. This was unexpected because our previous study showed that O antigen genes were required for ATCC 43816 serum resistance (Short et al. 2020) and we predicted that resistance to *G. mellonella* humoral immunity may have similar requirements. However other studies of ATCC 43816 O antigen

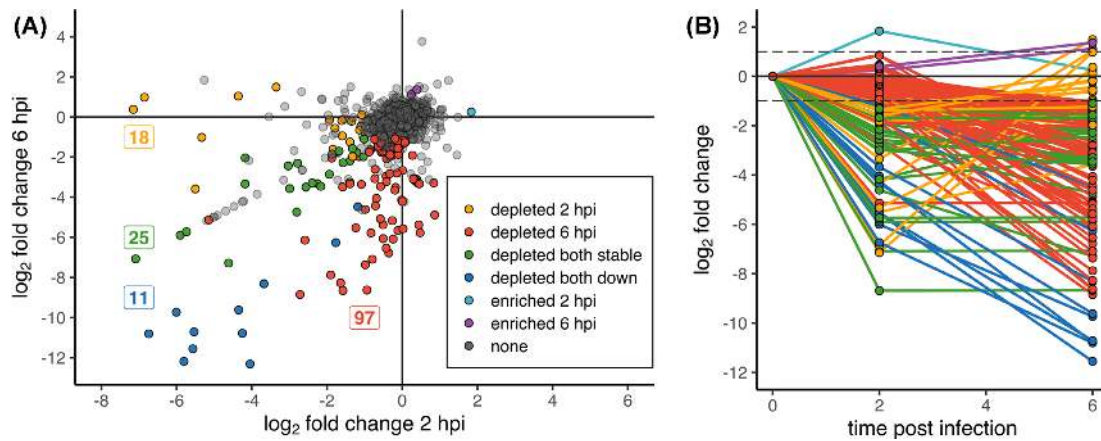


Figure 3. Few genes are important for the onset but not the late stages of a *G. mellonella* infection. (A), The plot shows log₂ fold-changes at 2 hpi and 6 hpi on the x- and y-axis, respectively. Transposon mutants significantly less abundant at 2 and 6 hpi are shown in yellow and red, respectively. Mutants which are depleted at both timepoints are shown in either green or blue (if the log₂ fold-change at 2 hpi roughly equals that at 6 hpi in green and if their abundance was much lower at 6 hpi in blue). Genes significantly higher abundant at 2 hpi and 6 hpi are shown in turquoise and purple, respectively. Non-significant genes are shown in grey. Numbers indicate the number of genes of each group. (B), Graph highlighting the direction of the log₂ fold-change change over time. Genes are coloured according to A; non-significant genes are not shown.

have shown variable effects on virulence and related *in vitro* phenotypes (Shankar-Sinha et al. 2004; Yeh et al. 2016). We speculate that the ATCC 43816 capsule has a dominant role in protection from *G. mellonella* immunity and masks the activity of O antigen.

To compare the global TraDIS results from RH21207 with those of ATCC 43816, we identified their shared genes by bidirectional best hits BLAST search using a sequence similarity cut-off of 80 %. This analysis identified 4,391 shared genes, 3974 of which were nonessential in both strains and therefore included in the comparison. We identified 149 shared genes with a role in *G. mellonella* infection in either RH21207 or ATCC 43816, and 44 genes were implicated in both strains (Fig. 4A). This means that 37.3 % of the hits in RH21207 were also identified in ATCC 43816 and 58.7 % of all hits in ATCC 43816 were identified in either of the two RH21207 datasets, the largest overlap was with the later timepoint of 6 hpi. Amongst those genes were, for example, the membrane-associated genes *ompK36*, *tolABQ*, and *lpp*, the outer membrane protein assembly factor *bamB/yfgL*, the periplasmic chaperone *surA*, and the ECA synthesis genes *rffG* and *wecA*.

We performed a clusters of orthologous groups (COG) enrichment analysis to define and compare the broad pathways and molecular mechanisms that may contribute to fitness during *G. mellonella* infections in both strains. The COG classifications of the genes in the RH21207 and the ATCC 43816 genome was annotated with EggNOG Mapper (Huerta-Cepas et al. 2017) and the COGs of all genes in which mutants were significantly underrepresented after infection were compared to the total gene set. In RH21207, genes of the replication, recombination and repair (L), cell wall/membrane/envelope metabolism (M) and nucleotide transport and metabolism (F) clusters were overrepresented among the infection-related genes at 2 hpi, whereas amino acid transport and metabolism (E) and genes of unknown function (S) were underrepresented (Fig. 4B). At 6 h post-infection, cell wall/membrane/envelope metabolism (M), intracellular trafficking, secretion, and vesicular transport (U) and nucleotide transport and metabolism (F) were overrepresented, whereas only genes of unknown function (S) were underrepresented (Fig. 4C). In the ATCC 43816 infection determinants, genes assigned to carbohydrate transport and metabolism (G), cell wall/membrane/envelope biogenesis (M) and nucleotide transport and metabolism (F) were overrepresented. The cell

wall/membrane/envelope metabolism (M) and nucleotide transport and metabolism (F) clusters were the only ones in which infection-related genes were overrepresented for both cKp and hvKp, and at early and late time points. This finding further stresses the importance of cell membrane integrity and surface polysaccharides in infections, but also demonstrates the essentiality of nucleotide metabolism during the course of an infection. The latter COG included the genes *purA*, *purC*, *purE*, *purH*, and *rdgB*; all purine biosynthesis genes that showed significantly reduced fitness in both strains. The ability to *de novo* synthesize purines has been associated with the intracellular survival of bacterial pathogens such as *Burkholderia pseudomallei*, *Shigella flexneri*, and uropathogenic *Escherichia coli* (Ray et al. 2009; Shaffer et al. 2017).

Comparison with genetic fitness requirements in murine hosts and *in vitro* virulence screens

We then sought to compare our findings in *G. mellonella* to published results of other high-throughput fitness screens of *K. pneumoniae*, with the caveat that such screens by their nature provide only a snapshot of the events of an infection, and are unlikely to be comprehensive.

The *G. mellonella* fitness genes identified in both strains showed considerable similarity to those required for survival in human serum, which we examined previously using the same mutant libraries (Short et al. 2020). This was largely due to the importance of capsule, O antigen and cell envelope proteins such as Lpp for survival under both selections; the many metabolic genes identified as infection-relevant, for example those of purine biosynthesis (Fig. 4), generally did not contribute to serum survival. For RH21207, over half of the genes identified as required for full serum fitness also contributed to fitness in *G. mellonella*.

Fitness factors required for intestinal colonisation of mice have also been examined in an ST258 background (Benoit et al. 2019), although this screen was not comprehensive due to various experimental factors. Several genes required in *G. mellonella* are also required for intestinal colonisation, such as *bamB* (an outer membrane assembly protein), *ompC/ompK36*, *cyaA*, (adenylate cyclase) and *typA* (a GTP-binding protein). Finally, there are

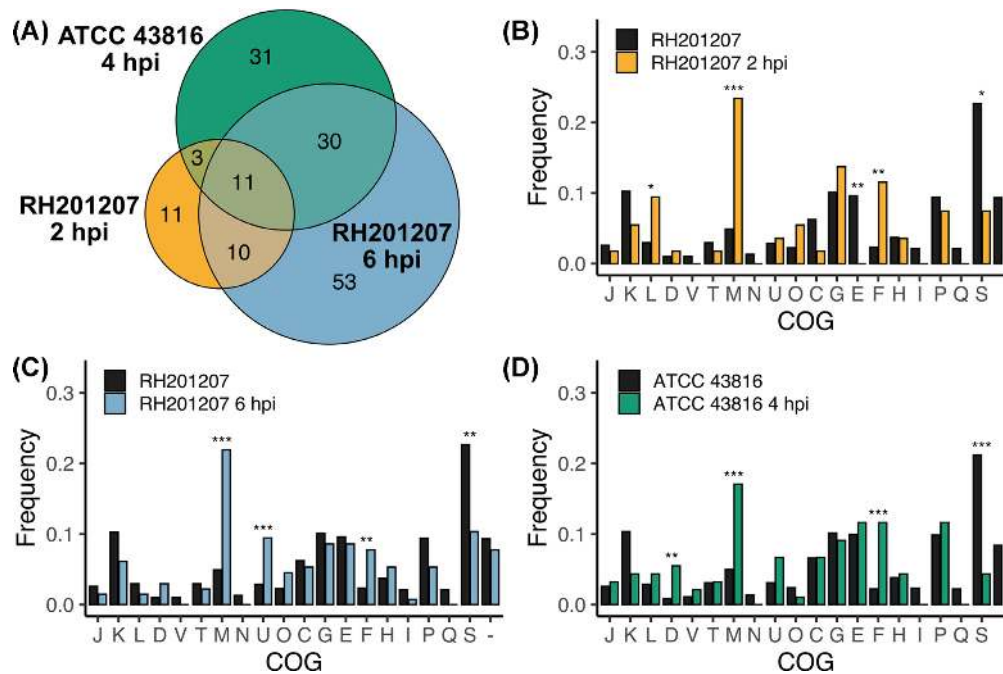


Figure 4. Multiple genes, including the *cps* cluster contribute to the fitness during *Galleria* infections of both classical and hypervirulent *K. pneumoniae*. (A), Venn diagram showing the overlap of significantly less abundant insertion mutants in the classical ST258 strain RH201207 and the hypervirulent strain ATCC 43816. Only non-essential genes shared by both strains as determined by a bi-directional best blast analysis are shown. (B–D), Cluster of orthologous groups (COG) enrichment analysis shows an overrepresentation of outer membrane biogenesis as well as nucleotide transport and metabolism genes in all sets of significantly depleted genes during *G. mellonella* infection. Genes were assigned to Cluster of Orthologous Groups (COG) (Tatusov et al. 2000) with eggNOG. The bars represent the percentage of genes that belong in that category. Black bars denote the frequency of COGs in all non-essential genes of the particular strain, the frequency of all significantly depleted genes after *Galleria* infection is coloured as follows, A: RH201207 2 hpi, B: RH201207 6 hpi, C ATCC 43816 4 hpi. P-values were determined by Fisher's exact test in R and the level of significance is indicated by asterisks (*, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$). COGs without genes assigned to them (A, B, R, W, Y & Z) were removed. Abbreviations of COGs: J, Translation, ribosomal structure and biogenesis; K, Transcription; L, Replication, recombination and repair; D, Cell cycle control, cell division, chromosome partitioning; V, Defence mechanisms; T, Signal transduction mechanisms; M, Cell wall/membrane/envelope biogenesis; N, Cell motility; U, Intracellular trafficking, secretion, and vesicular transport; O, Post-translational modification, protein turnover, chaperones; C, Energy production and conversion; G, Carbohydrate transport and metabolism; E, Amino acid transport and metabolism; F, Nucleotide transport and metabolism; H, Coenzyme transport and metabolism; I, Lipid transport and metabolism; P, Inorganic ion transport and metabolism; Q, Secondary metabolites biosynthesis, transport and catabolism; S, Function unknown; -, no hit.

some important common factors among the requirements for infection of *G. mellonella*, and for an hvKp murine lung infection (Paczosa et al. 2020). Genes contributing to both infection types include some of those encoding ubiquitous virulence factors such as capsule or siderophore importers. Notably, genes involved in aromatic amino acid biosynthesis (e.g. *pabAB* aminodeoxychorismate synthase, *aro* operon genes), and purine biosynthesis (e.g. *purH*) were required both for murine lung infection, and for *Galleria mellonella* infection in both strain backgrounds. The importance of these pathways for two very different types of infection, in representatives of both hvKp and cKp, suggests that they may be general infection requirements for *K. pneumoniae*.

Validation of transposon insertion sequencing results and investigation of NfeR activity

Five single-gene mutants of *K. pneumoniae* RH201207 were tested for lethality in *G. mellonella* to validate the TraDIS screen. The validation set included transposon insertion mutants in the transcription antiterminator *rfaH*, the capsule gene *wzc* and the enterobacterial common antigen gene *wzxE*, as well as clean deletion mutants of *phoQ* and *nfeR*. Research-grade *G. mellonella* larvae were injected with 10^6 cfu of each strain, or a PBS control, and monitored for up to 72 hours. All strains showed a statistically significant virulence defect (Fig. 5A), with the exception of RH201207 Δ *phoQ*; larvae infected with the Δ *phoQ* mutant showed increased survival but the degree did not reach statistical significance. It is possible that the TraDIS screen detected

changes that reflect fitness in a competition environment with other strains, or that are important at early stages of infection but are not relevant in the longer term.

We selected *nfeR* (KPNRH.00645 also called *yqjI*) for further characterisation, as this gene has not previously been linked to virulence in any species. In our TraDIS experiments, this gene was required in *K. pneumoniae* RH201207, but not in ATCC 43816. Complementation of the Δ *nfeR* mutation restored the virulence of the wild-type RH201207 strain (Fig. 5B). NfeR contributes to metal homeostasis in *E. coli* by repressing expression of a neighbouring ferric reductase gene, *nfeF* (Fig. 6A); this repression is relieved under excess nickel conditions (Wang, Wu and Outten 2011; Wang et al. 2014). High nickel levels can disrupt iron homeostasis in *E. coli*, leading to a longer lag phase due to reduced accumulation of iron (Rolfe et al. 2012; Washington-Hughes et al. 2019). We hypothesised that loss of *nfeR* may lead to deregulated metal homeostasis in *K. pneumoniae* RH201207, resulting in reduced virulence in *Galleria mellonella*.

We first tested whether NfeR regulates *nfeF* expression in *K. pneumoniae* RH201207 by qRT-PCR using RNA extracted from late exponential phase bacteria. As shown (Fig. 6B), deletion of *nfeR* caused a dramatic 85-fold increase in *nfeF* expression, and this change was not seen in the complemented strain. We hypothesised that the virulence defect of RH201207 Δ *nfeR* may arise from a reduced defence against humoral immunity, or reduced ability to acquire iron, as *G. mellonella* recapitulates several relevant features of mammalian serum-based immunity, and is iron-limited (Lucidi et al. 2019). However, the mutant did not show any differences relative to its wild type either in its ability to

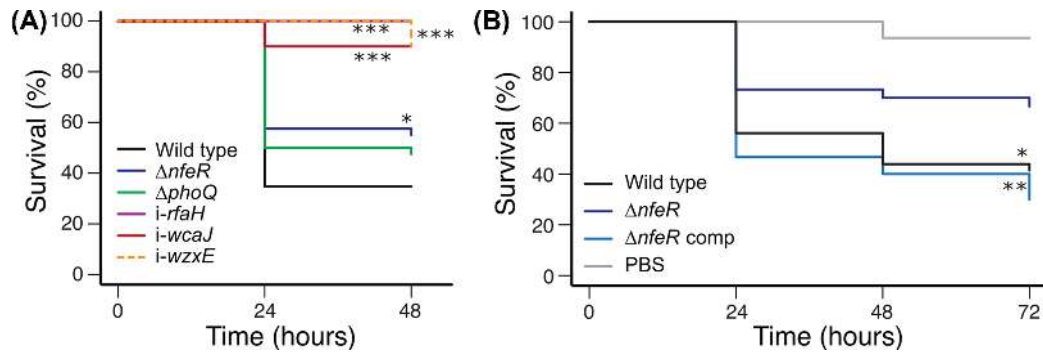


Figure 5. Validation of TraDIS results with single-gene knockouts. (A), Survival curves for *G. mellonella* larvae following infection with *K. pneumoniae* RH201207 and mutants in defined genes. (B), Complementation of the virulence defect of RH201207 $\Delta nfeR$. Mutants where survival is significantly different to wild-type (Kaplan-Meier test) are indicated by asterisks (*, $P < 0.05$; **, $P < 0.01$; ***, $P < 0.001$).

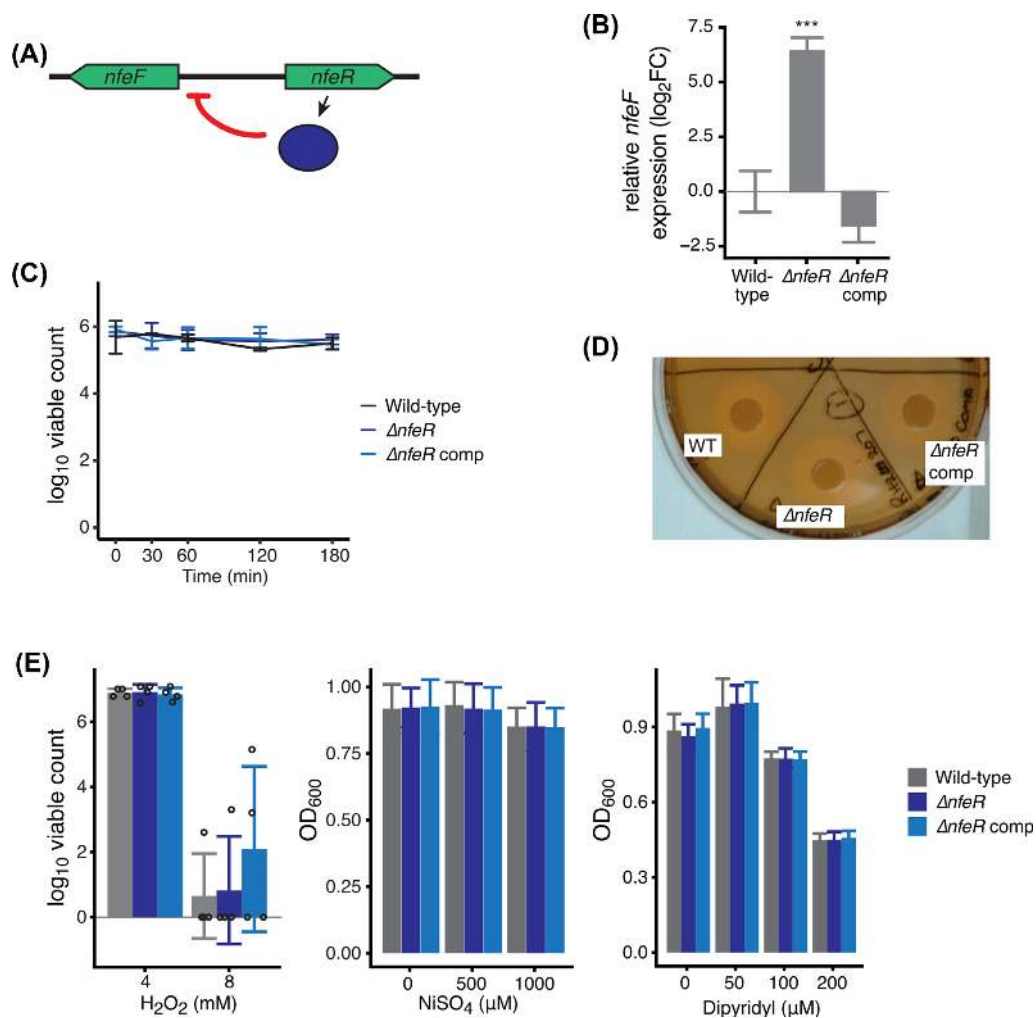


Figure 6. Investigation of NfeR function in RH201207. Mutation of *nfeR* resulted in a dramatic increase in *nfeF* expression. (A), Schematic of *nfeF* gene expression regulation by NfeR. (B), Deletion of *nfeR* leads to significant overexpression of *nfeF* as measured by qRT-PCR. Level of significance in comparison to RH201207 is indicated by asterisks (***, $P < 0.001$). (C), Serum survival of RH201207 $\Delta nfeR$. Late exponential phase bacteria were incubated in 66 % normal human serum for 3 hours, and viable counts measured over time. (D), Siderophore production by RH201207 $\Delta nfeR$, using chrome azurol S agar assay. Results shown are representative of two independent experiments, each comprising at least two biological replicates. (E), Sensitivity of RH201207 $\Delta nfeR$ to hydrogen peroxide, Nickel toxicity and dipyridyl-induced metal starvation. Stationary phase cells were diluted directly into H₂O₂-supplemented Mueller Hinton broth, and surviving cells enumerated after 100 min. For Nickel and dipyridyl toxicity, growth was measured after incubation for 18 hours at 37°C. Results shown are the mean and standard deviation of four (H₂O₂) or three biological replicates, which in the case of Ni and dipyridyl comprised of three technical replicates.

withstand serum exposure (Fig. 6C), or its siderophore production (Fig. 6D). Finally, we examined three phenotypes that may be disrupted when metal homeostasis is altered: resistance to oxidative stress (hydrogen peroxide), nickel toxicity and sensitivity to iron starvation. Sensitivity to oxidative stress is influenced by cellular iron pools, and was previously shown to be growth-phase dependent for this reason (Touati 2000). Homologues of *nfeR* have been implicated in resistance to nickel toxicity in large-scale fitness screens (Price et al. 2018). It was also shown that some genes in the same pathway as *nfeF* are more sensitive to dipyrrolyl-mediated iron starvation (McHugh et al. 2003). However, *K. pneumoniae* RH201207 $\Delta nfeR$ did not show changes in any of these phenotypes. Thus, while $\Delta nfeR$ mutation increases *nfeF* expression and reduces virulence, its mechanism appears to be via subtle effects that are not replicated in vitro.

CONCLUSIONS

Despite the urgency of the public health threat posed by classical, multidrug-resistant *K. pneumoniae* strains, our understanding of their mechanisms of infection are still limited. *Galleria mellonella* is an increasingly popular alternative model for bacterial infections, which, unlike mice, is susceptible to infection by cKp. Here, we have performed the first high-throughput fitness profiling study of *K. pneumoniae* during *G. mellonella* infection.

Galleria mellonella had favourable infection parameters for high-throughput screening; the diversity of the highly saturated transposon mutant library was largely maintained through the experiment, and excellent reproducibility was achieved between biological replicates. Though the infection was examined over a short time course (up to 4 hpi and 6 hpi, respectively), similar to that used for *G. mellonella* TnSeq of *Acinetobacter baumannii* (Gebhardt et al. 2015), this still allowed identification of genes required for nutrient acquisition as well as defence against host immunity. Longer infections could be examined by using smaller mutant pools (~10 000 unique mutants) at a lower inoculum, and this may be of interest to future researchers. Infection-related genes identified showed high concordance with current knowledge of *K. pneumoniae* pathogenesis; genes for all of the major virulence factors (siderophores, capsule, O antigen) showed decreased mutant abundance, and many new virulence gene candidates were identified in cKp. This included the metal-dependent regulator *nfeR*, which did not contribute to virulence in the hvKp background. A limitation of our study, and indeed the majority of studies providing molecular detail on *G. mellonella*-pathogen interactions, is that these putative virulence factors have not been further examined in mammalian models.

Our results showed a substantially different fitness landscape for RH201207 and ATCC 43816 during *G. mellonella* infection. Fewer infection-related genes were identified in *K. pneumoniae* ATCC 43816 (hvKp). This finding may reflect masking of some relevant activities by dominant virulence factors; for example, the highly expressed K2 capsule of this strain may compensate for loss of other cell envelope components, such as multiple genes involved in RpoE signalling which were required in the cKp background. The complex interplay of virulence factors underscores the need to consider the phylogenetic diversity of *K. pneumoniae* when studying its pathogenesis.

We have demonstrated a simple, scalable method for virulence factor profiling in cKp, and used it to provide the first genome-scale view of a cKp infection and compare it to an hvKp strain. The capacity of the *G. mellonella* model to elucidate relevant virulence activities, and the ease with which it can be applied to new strains, opens the possibility for robust

species-wide comparisons of infection determinants in *K. pneumoniae* and other opportunistic pathogens.

SUPPLEMENTARY DATA

Supplementary data are available at [FEMSPD](https://www.femsdpd.com) online.

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