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Heat Resistance Mediated by a New Plasmid Encoded Clp ATPase, ClpK, as a Possible Novel Mechanism for Nosocomial Persistence of *Klebsiella pneumoniae*

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**Abstract**

*Klebsiella pneumoniae* is an important opportunistic pathogen and a frequent cause of nosocomial infections. We have characterized a *K. pneumoniae* strain responsible for a series of critical infections in an intensive care unit over a two-year period. The strain was found to be remarkably thermotolerant providing a conceivable explanation of its persistence in the hospital environment. This marked phenotype is mediated by a novel type of Clp ATPase, designated ClpK. The *clpK* gene is encoded by a conjugative plasmid and we find that the *clpK* gene alone renders an otherwise sensitive *Escherichia coli* strain resistant to lethal heat shock. Furthermore, one third of a collection of nosocomial *K. pneumoniae* isolates carry *clpK* and exhibit a heat resistant phenotype. The discovery of ClpK as a plasmid encoded factor and its profound impact on thermal stress survival sheds new light on the biological relevance of Clp ATPases in acquired environmental fitness and highlights the challenges of mobile genetic elements in fighting nosocomial infections.

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**Introduction**

The Gram-negative enterobacterium *Klebsiella pneumoniae* is considered an important opportunistic pathogen frequently implicated in nosocomial infections such as urinary tract infection, pneumonia and sepsis [1]. As the cause of Gram-negative nosocomial sepsis, *K. pneumoniae* is second to only *Escherichia coli* [2], however, several recent pro- and retrospective studies have even placed *Klebsiella* as the predominant causal agent of Gram-negative bacteraemic cases, especially within the intensive care units (ICUs) [3–5]. The often immunocompromised patients suffering from severe underlying diseases in ICUs are frequently susceptible to infection with this opportunistic pathogen [6]. *K. pneumoniae* infections are associated with significant mortality depending on the site of infection i.e. systemic infection fatality rates between 20 and 50% are often reported, the rate even reaching 70% [4,7–9].

*K. pneumoniae* is ubiquitous in nature and a commensal of the human gastrointestinal tract. Hence, the gastrointestinal tract is considered the main reservoir from where the bacterium is spread in the nosocomial environment, and contaminated hospital equipment and person-person contact are primary routes of transmission [10]. Virulence factors associated with *K. pneumoniae* pathogenicity include capsular polysaccharide (CPS), lipopolysaccharide (LPS) and fimbrial adhesins of type 1 and type 3, although a comprehensive understanding of *K. pneumoniae* infection mechanisms remains elusive. *K. pneumoniae* strains of environmental origin are capable of expressing the same arsenal of virulence factors as clinical isolates [11] and have comparable virulence potential when tested in animal infection models [12]. Thus, the ubiquity of *K. pneumoniae* poses a constant threat to the immunocompromised host.

Bacterial resistance to adverse conditions depends on protein members of the heat shock response [13] comprising two major classes of proteins, chaperones and proteases, both being involved in protein homeostasis and regulatory functions [14,15]. The conserved Clp/Hsp100 ATPase protein family is traditionally divided into subfamilies based on the number of ATPase motifs as well as other structural motifs [16–18]. All Clp ATPases are expected to have intrinsic chaperone activity [16,19–21], whereas association with ClpP confer proteolytic activity and depends on specific signature motifs [22]. The expression of different Clp proteins and the phenotype associated with their presence is species-dependent, however, a protein-threading mechanism seems to be a shared feature [16].

In this study we have identified a recurrent clinical *K. pneumoniae* strain with a remarkably increased heat resistance. We show that the thermotolerance phenotype is attributable to a novel plasmid encoded Clp ATPase, designated ClpK, and that the presence of this *clpK* gene correlates positively with the thermotolerance phenotypes observed among clinical isolates.
Results and Discussion

The nosocomial strain *K. pneumoniae C132-98* exhibits a heat resistant phenotype

During the course of a surveillance study at a Danish hospital, we noticed a series of ICU infections caused by capsular serotype K28 *K. pneumoniae*. All isolates were resistant to gentamicin, netilmicin, tobramycin and sulphamamide. Additionally, two isolates exhibited an ESBL (extended-spectrum beta-lactamase) phenotype as revealed by clavulanic acid reversible ceftazidime resistance. These strains were isolated from at least 10 severe cases (nine cases being fatal) over a two-year period. The clonal relatedness of the isolates was confirmed by ribotyping using the restriction enzyme EcoRI (Fig. 1), which has a high discriminatory power for *K. pneumoniae* [23,24]. The recurring clone of *K. pneumoniae* is represented by strain C132-98. Source and route of transmission was never proven but, interestingly, infections caused by this clone ceased concomitantly with an increased focus on the cleaning/disinfection of reusable flexible endoscopes. We have previously shown that *K. pneumoniae* C132-98 is not more virulent compared to other clinical strains and isolates of environmental origin [12]. Thus, the repeated emergence of C132-98 was not directly attributable to a highly virulent phenotype.

Therefore, we initiated a search for increased resistance to environmental stressors that could explain the persistence of this specific strain in the nosocomial environment and found that C132-98 exhibits a markedly higher heat tolerance when compared to another clinical strain *K. pneumoniae* C3091 and the reference strain for *K. pneumoniae* capsule serotype K28. When shifted from 37°C to 55°C, strain C132-98 was detectable by plating for more than 30 minutes after the shift, whereas the other strains were undetectable already 15 minutes after the shift (Fig. 2). Even when exposed to 58°C or 60°C, C132-98 showed prolonged survival compared to the control strains (not shown).

Thus, the remarkable heat resistance of *K. pneumoniae* C132-98 may provide a plausible explanation for the repeated emergence of this specific strain within the given ICU, especially when taken into consideration that the standard disinfection procedure of flexible endoscopes is a combined thermo-chemical treatment below 60°C due to the thermo-lability of the endoscopes, combined with the extensive use of such endoscopes within ICUs. Our hypothesis is further supported by the virulence study [12] that excluded the possibility of explaining the clinical history of C132-98 by a specifically virulent phenotype.

Isolation and identification of clpK

A search for the genetic locus involved in the heat resistance of *K. pneumoniae* C132-98 was conducted. Plasmid profiling revealed that C132-98 possesses several plasmids (≥eight). Four of them are large plasmids equal to or larger than 100 kb (Fig. 3A). Thus, the thermo-protective genetic locus could be located on a plasmid. To examine this notion, we constructed a DNA library from C132-98 plasmid DNA (Fig. 3B) and, anticipating that the heat resistant property would be expressed in *E. coli*, a total of 384 library clones were pooled and subjected to thermal selection. Hereby, a specific plasmid, pMB58, was selected conferring increased heat resistance (Fig. 3B) also after reintroduction into a clean *E. coli* DH5α background (Fig. 3C). The plasmid carrying strain was detectable up to 45 min at 53°C compared to the vector control strain being undetectable after 20 min.

| Label | RiboGroup | RiboPrint(R) Pattern |
|-------|------------|----------------------|
| 1:B-8 | RIB01 58-S-6 | 1:RIB01 58-S-6 |
| 2:A-8 | RIB01 58-S-6 | 2:RIB01 58-S-6 |
| 1:H-3 | RIB01 58-S-6 | 1:RIB01 58-S-6 |
| 2:A-1 | RIB01 58-S-6 | 2:RIB01 58-S-6 |
| 1:F-8 | RIB01 58-S-6 | 1:RIB01 58-S-6 |
| 2:B-2 | RIB01 58-S-6 | 2:RIB01 58-S-6 |
| 2:B-1 | RIB01 58-S-6 | 2:RIB01 58-S-6 |
| 2:B-7 | RIB01 58-S-6 | 2:RIB01 58-S-6 |
| 1:F-5 | RIB01 58-S-6 | 1:RIB01 58-S-6 |
| 1:E-8 | RIB01 58-S-6 | 1:RIB01 58-S-6 |

Figure 1. Verification of strain clonality among clinical *Klebsiella pneumoniae* isolates. 10 independent ICU isolates collected over a two-year period are positioned within the same RiboGroup as shown by identical EcoRI RiboPrint patterns. doi:10.1371/journal.pone.0015467.g001
DNA sequencing of pMB58 revealed an 8187 bp fragment (GenBank accession FJ042668) covering seven putative open reading frames (ORFs). The segment shows a high overall similarity to a chromosomal region from the *Methylobacillus flagellatus* genome (GenBank accession NC_007947). Three of the ORFs are also homologous to a segment of a plasmid encoded sequence (pKPN3) found in *K. pneumoniae* MGH78578 (GenBank accession NC_009649) (Fig. 3D). The cloned segment covers genes corresponding to, among others, putative small heat shock proteins (sHSP) and a Clp ATPase. In order to identify the gene responsible for the heat resistant phenotype, in vitro transposon mutagenesis of pMB58 was performed and a mutant exhibiting a thermal tolerance comparable to *E. coli* carrying the empty vector was identified (Fig. 3C). The mutagenesis revealed that the ORF corresponding to the putative Clp ATPase gene was responsible for the heat resistance mediated by pMB58 in *E. coli* DH5α. The protein corresponding to the Clp ATPase gene was designated ClpK (K for *Klebsiella*).

The implication of the *clpK* gene in thermal protection was further confirmed by expressing ClpK (pClpK) in *E. coli* leading to survival of approximately 10% of the cells after 20 min of heat shock at 53°C compared to a 100-fold reduction (0.1%) in the absence of ClpK (Fig. 4).

**ClpK is necessary for the heat tolerance of *K. pneumoniae***

A Δ*clpK* mutation was constructed in *K. pneumoniae* C132-98 by allelic replacement. When exposed to heat, viable *K. pneumoniae* were detectable for at least 50 min at 53°C, whereas the CFU of the Δ*clpK* mutant was below the detection limit after 20 min exposure (Fig. 5). The reduced heat tolerance of the C132-98 Δ*clpK* mutant strain could be complemented in part by plasmid pClpK expressing ClpK (Fig. 5) and therefore, we conclude that ClpK is involved in the heat resistance of *K. pneumoniae* C132-98. Since the Δ*clpK* mutant was incompletely complemented by pClpK, a polar effect due to the deletion was considered. Indeed, transformation of the entire heat resistance locus (cf. Fig. 3D) by plasmid pMB58-sub fully restored the thermal tolerance of C132 Δ*clpK* (Fig. 5) Therefore, obstruction of *clpK* in C132 most likely lead to malfunction of co-localized genes involved in heat resistance. Interestingly, genes encoding small heat shock proteins (sHSP) are present up- and down-stream *clpK*. In *E. coli*, the sHSPs IbpA/IbpB (inclusion body-associated proteins) bind intracellular proteins aggregated by heat shock and allow refolding in a process involving the ClpB ATPase [25–29]. A similar collaboration between ClpK and the products from these co-localized sHSP genes in stress alleviation is possible. At least, given their co-localization, it is tempting to speculate that these genes may be functionally related. However, the thermoprotective activity of ClpK is most likely not strictly dependent on any co-localized genes since *clpK* alone exerts a heat resistance phenotype when transformed into *E. coli* and significantly enhances the heat resistance of C132 Δ*clpK*. Further studies are needed to identify the specific genes involved in ClpK-mediated heat resistance and to assess any functional collaboration.

No other specific phenotypes, e.g., sensitivity to high salt (NaCl) concentrations, impact of H₂O₂ on growth, and survival under desiccative conditions were observed for the Δ*clpK* mutant strain when compared to the wild type. Moreover, the maximum temperature allowing growth (46°C) was unaffected by the Δ*clpK* mutation (not shown). Thus, the *clpK* locus seems to specifically enhance the ability of C132-98 to survive an otherwise lethal temperature.

**clpK is encoded on a conjugative plasmid**

To confirm that the *clpK* gene is present on a plasmid we purified plasmid DNA from C132-98 Δ*clpK*, in which the *clpK* gene is substituted with a tetracycline encoding cassette; transformed *E. coli* and selected a tetracycline resistant transformant that harbored only the largest 150 kb plasmid (not shown). The presence of the *clpK* deletion on the plasmid was confirmed by PCR (not shown), demonstrating that *clpK* is indeed encoded by the largest plasmid in *K. pneumoniae* C132-98. Consequently, we wished to establish whether *clpK* and the heat resistance phenotype was transferable by conjugation. Indeed, by simple plate mating and selection for gentamicin resistance we isolated a transconjugant of *Escherichia coli*
MG1655 (Fig. 6A) with a notably heat resistant phenotype. In fact, the recipient strain exhibited an approximately 100-fold improved survival after 50 minutes at 53°C (Fig. 6B). A few plasmid encoded Clp ATPases have previously been identified; however, no significant phenotypes were associated with their presence or inactivation [30,31]. We believe that the present study provides the first reported example of a plasmid encoded Clp ATPase having a clear phenotypic effect demonstrated by an improved tolerance to thermal stress.

ClpK defines a novel subclass of Clp ATPases

Alignments of ClpK with established members of the Clp ATPase family revealed that ClpK displays primary amino acid similarity to the class I Clp ATPases carrying two nucleotide binding domains (cf. Fig. 7). Uniquely, ClpK carries a 100 amino acid N-terminal extension not observed in other Clp proteins, and the linker domain separating the nucleotide-binding domains is of intermediate size resembling ClpC and its relatives, ClpE and ClpL, that are considered Gram-positive Clp ATPases only. ClpK is clearly distinct from the previously recognized Gram-negative Clp ATPases, ClpA and ClpB, carrying small and large linker domains, respectively. Thus, this is the first report on a Gram-negative Clp ATPase having a linker domain of intermediate size.

The amino acid sequence of ClpK shares extensive identity with uncharacterized Clp proteins in completed genomes from *E. coli* (98%), *Enterobacter cloacae* (98%), *M. flagellatus* (95%), *Desulfovibrio desulfuricans* (94%), *Comamonas testosteroni* (93%), *Marinobacter aquaeolii* (92%), *Pseudomonas mendocina* (90%) and *Ochrobactrum anthropi* (73%) among others. Most intriguingly, these bacteria represent species from all proteobacterial subdivisions (α, β, Δ and γ) except division ε. Thus, ClpK is more likely to have evolved by horizontal transfer rather than vertical inheritance. Of additional interest, ClpK homologous proteins are identified in species such as *Silicibacter lacuscaerulensis* and *Dictyoglomus thermophilum* considered moderately thermostolerant mesophile and bona fide thermophile species, respectively. We did not identify ClpK homologues in any Gram-positive species, suggesting that this ATPase is unique to Gram-negative bacteria.

By comparing the available amino acid sequences of ClpK homologues, other distinct features were identified. These include the absence of a characteristic tripeptide that is proposed to be required for the Clp ATPase partners to interact with the

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Figure 3. Isolation and identification of a Clp ATPase involved in the heat resistance of *K. pneumoniae* C132-98. (A) Plasmid profile of C132-98 compared to reference molecular marker of known molecular mass. (B) C132-98 plasmid DNA library construction and thermal selection. Verification of clonal diversity of library constructs by EcoRI restriction analysis of 5 representative clones (cut and uncut plasmids, respectively) prior to thermal selection. Selection of a specific clone (pMB58) as shown by similar EcoRI restriction patterns of 5 representative colonies (uncut and cut plasmids, respectively) obtained from the pooled library following repeated heat shock treatments. (M) Molecular marker ~10 kb, (v) empty vector. (C) Assessment of thermal protection of *E. coli* DH5α at 53°C conferred by plasmid pMB58 (A) compared to the empty vector background pBR322 (B). A derivative of pMB58 subjected to in vitro transposon mutagenesis, pMB58_Tn (C), shows basal thermal protection comparable to the empty vector. One representative experiment out of several is shown. (D) Seven putative ORFs identified on the cloned fragment (pMB58) from C132-98 plasmid DNA. Identities of deduced amino acids to a chromosomal fragment of *M. flagellatus* and a locus on plasmid pKPN3 from *K. pneumoniae* MGH78578 are given. The position of the transposon insertion in pMB58_Tn within the Clp ATPase is indicated.

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proteolytic subunit ClpP in *E. coli* and other bacteria [32] and the presence of three conserved cysteine and a single histidine residue in the amino terminal domain possibly forming a zinc finger motif or a reminiscent thereof. Zinc finger motifs are found in ClpX and in ClpE where it is functionally important [33,34]. The significance of the identified motif in ClpK remains to be elucidated, however, the high interspecies amino acid similarity within the N-terminal 250 amino acids of ClpK proteins (>85%) suggests that this domain exerts functional properties essential and/or unique to ClpK.

Defining features of ClpK compared to established subclasses of Clp proteins are depicted in Figure 7. We named the protein ClpK as it constitutes a separate subclass within the Clp ATPase family and *K. pneumoniae* ClpK is the prototype of this novel subclass.

Prevalence of clpK in clinical *K. pneumoniae* strains and correlation with heat resistant phenotype

In order to determine the prevalence of clpK among *K. pneumoniae* strains, a collection of 105 clinical isolates of *K. pneumoniae* was tested using a nucleotide probe targeting the conserved linker domain sequence unique to clpK. By this screening, we detected clpK in 31 (30%) of the strains (not shown). Hence, clpK is not unique to strain C132-98, but the gene is present in a subset of clinical *K. pneumoniae* isolates. Consequently, the thermotolerance of a randomly chosen representative collection of clpK positive and negative *K. pneumoniae* strains was compared (Fig. 8) and a positive correlation between having the clpK gene and expressing a thermotolerant phenotype was established. Of the strains tested, all strains devoid of clpK were undetectable after 20 min of heat shock at 53°C, whereas all clpK
positive strains exhibited detectable survival at 40 min and beyond. A marked strain-to-strain difference in heat resistance was observed, which may be due to the presence/absence of other genes coexisting with clpK, such as the downstream sHSP found in C132-98 which are absent in MGH78578 (cf. Fig. 3D). Indeed, our complementation experiment (Fig. 5) showed that other factors encoded at the clpK locus are of importance in resistance toward thermal stress. Thus, high level heat resistance requiring more than the ClpK protein alone seems a plausible reason for observed differences in resistance phenotypes among isolates. Alternatively, ClpK expression may also differ between strains. Nevertheless, a clear association between the presence of the clpK gene and elevated thermal tolerance was established for K. pneumoniae.

Concluding remarks

A K. pneumoniae strain being considerably resistant towards thermal stress was identified in a hospital setting. Molecular cloning and phenotypic studies allowed identification of a novel plasmid encoded ClpK ATPase mediating the increased heat resistance. In previous studies, chromosomally encoded Clp ATPases have been demonstrated to be required for growth or survival at high temperature such as the ClpB protein in E. coli [35] and Staphylococcus aureus [36]; and the ClpC in Listeria monocytogenes [37] and S. aureus [36] among others. This study is the first to report on a novel Clp ATPase that, as an accessory factor expressed from a plasmid, enhances the heat tolerance of the host strain and greatly increases survival under conditions that are found in the hospital setting. The clpK gene as a predictor of elevated thermal tolerance in K. pneumoniae is an interesting concept that may be determining in the ability of this opportunistic pathogen to survive and persist in niches of the nosocomial environment e.g. in reusable flexible endocopes undergoing thermo-chemical processing in which temperature is a critical parameter [38]. It can be speculated that the nosocomial persistence of K. pneumoniae C132-98 (and maybe other clinical isolates) was facilitated by acquisition of clpK.

**Figure 6. A plasmid encoded thermotolerance locus from K. pneumoniae C132-98 is transferable by conjugation rendering E. coli MG1655 heat resistant.** (A) Plasmid profile. Lanes 1-3 represent K. pneumoniae C132-98, recipient E. coli MG1655, and a gentamicin resistant transconjugant of E. coli MG1655, respectively, and lane M represents E. coli 39R861 harboring four plasmids as a molecular weight marker. (B) Heat shock survival at 53°C of E. coli MG1655 transconjugant (●) compared to the parental MG1655 recipient strain (○). Means and standard error of the means from three independent experiments are shown. doi:10.1371/journal.pone.0015467.g006

**Figure 7. Major defining features separating the novel ClpK protein from established Clp ATPase families.** ClpK is characterized by a unique extended N-terminal, the presence of a putative zinc finger motif (Zn), having a linker domain of intermediate length, and lacks a ClpP interaction tripeptide (P). doi:10.1371/journal.pone.0015467.g007
ClpK homologues were found in genomes from *E. cloacae* and *E. coli* among others. To date, clpK was observed only in one genome from all the *E. coli* genomes available at NCBI, *Escherichia coli* MS 115-1, and in genome *Escherichia* sp. 1_1_43. Thus, as for *K. pneumoniae*, clpK is not ubiquitous in *E. coli* but present only in a subset of isolates. Since we observed that transformation of laboratory *E. coli* strains with clpK confer thermal resistance, it is a reasonable anticipation that ClpK exerts a heat resistant phenotype in this species, and the clpK gene may very well be of importance in acquired environmental fitness across several clinically relevant species.

We observed that clpK was transferable by conjugation which also endowed gentamicin resistance. Intriguingly, the endemic cluster from which C132-98 was isolated included clonal isolates that had acquired ESBL producing capabilities. Multiple drug resistant *K. pneumoniae* producing ESBLs are of emerging clinical significance [39] and of major concern due to poorer prognosis [40,41]. Studies have identified mobile environmental loci as source and route of infection [42,43] and studies have established a link between inadequately decontaminated endoscopes and a source and route of infection [42,43] and studies have established a link between inadequately decontaminated endoscopes and a source and route of infection [42,43] and studies have established a link between inadequately decontaminated endoscopes and a source and route of infection [42,43]. In this respect, the combination of multiple antibiotic resistance and bacterial factors increasing environmental fitness may prove to be of significant importance in the prevention of infections and spreading of clinically relevant pathogens. Bacterial factors of clinical relevance, such as virulence genes and antibiotic resistance genes, are often situated on plasmids. Thus, a compelling scenario is the probable co-transfer of such genes with genes enhancing environmental fitness which would be highly advantageous to clinical strains residing in the hospital setting.

**Materials and Methods**

**Strains and growth conditions**

The clinical *K. pneumoniae* strains used in this study were all obtained from The International Escherichia and Klebsiella Reference Centre (WHO), Statens Serum Institut. 105 bacteremic isolates previously serotyped according to CPS antigens by counter-current immuno-electrophoresis [9] and the ICU blood isolate *K. pneumoniae* C132-98 [12] were collected from Hvidovre Hospital, Denmark. The clinical strain *K. pneumoniae* C3091 [46], *K. pneumoniae* strain MGH78578 (ATCC700721) and the serotype K28 type strain 5758 [47] were included as references. *E. coli* 39R36 [48] was used as a standard reference for plasmid size estimation. *E. coli* DH5α was used for cloning purposes. All strains were grown in Luria-Bertani (LB) medium at 37°C. Transformation of *E. coli* and *K. pneumoniae* was performed by electroporation followed by selection on LB plates supplemented with appropriate antibiotics at the following concentrations: ampicillin (100 μg ml⁻¹), apramycin (30 μg ml⁻¹), chloramphenicol (12.5 μg ml⁻¹), gentamicin (20 μg ml⁻¹), and tetracycline (8 μg ml⁻¹).

**DNA manipulations and plasmid constructions**

Molecular typing of isolates was carried out by using a RiboPrinter™ and restriction enzyme EcoRI. Extraction of *K. pneumoniae* C132-98 endogenous plasmids was performed by a procedure in essence as previously described [49]. Purification of large endogenous plasmids for cloning purposes was performed with Plasmid Mini AX DNA (DNA-Gdańsk). Cloned plasmid DNA was purified using the Qiaprep Spin Miniprep Kit (Qiagen). DNA restriction enzyme digestions and phosphatase treatment (Antarctic Phosphatase) were performed according to the manufacturer’s instructions (New England Biolabs). Ligation was performed by the Fast-Link DNA Ligation Kit (EPICENTRE Biotechnologies). In vitro transposon mutagenesis was carried out by the EZ-Tn5™<sub>TM</sub> <i>oriV</i>/KAN-2™ Insertion Kit as recommended (EPICENTRE Biotechnologies). Polymerase chain reactions (PCR) were conducted with Expand High Fidelity PCR Kit and PCR DIG Probe Synthesis Kit (both Roche) under standard conditions according to the manual. Primers were obtained from and sequencing carried out at MWG Biotech AG.

Plasmid pBR322 was used for the construction of plasmid DNA library in *E. coli* DH5α. Purified plasmid DNA from *K. pneumoniae* C132-98 was partially digested with Sau3AI and size-fractionated by agarose electrophoresis from which 4–10 kbp fragments were

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**Figure 8. The presence of clpK correlates with a heat shock resistant phenotype of clinical *K. pneumoniae*.** Survival potential during heat shock at 53°C evaluated on nine clinical *K. pneumoniae* strains, including C132-98 (△) and MGH78578 (□), being positive for clpK (determined by colony blot hybridization) compared to nine clpK negative strains. The two groups are separated by the discontinuous line. doi:10.1371/journal.pone.0015467.g008
purified. Following ligation with a BamHI linearized dephosphorylated pBR322 vector and transformation of electro-competent E. coli, a total of 384 library clones were picked and held in microtiter trays. Plasmid pACYC184 was used for subcloning and expression in E. coli and K. pneumoniae. The entire heat resistance locus from pMB58 (pBR322) was cloned into pACYC184, generating pMB58-sub, using enzymes NheI and Sall to allow transformation of K. pneumoniae (intrinsically ampicillin resistant). The clpK encoding region was cloned by PCR using primer combination F_clpk (5'-GGGCGACACCTT-3') and R_clpk (5'-CTCCAAACACGTCAACT-3'), respectively. At their 5'-ends, primers R_upclpK and F_dwnclpK were equipped with 18 bp homologous to the extremities of the ClpK gene in K. pneumoniae C132-98 (5'-CAAGAATTGCCGGCGGAT-3') and 20 bp regions homologous to the extremities of the ClpK gene by a modification of the lambda Red mediated recombination procedure [50] previously applied to chromosomal gene replacement in K. pneumoniae [51]. Alleric replacement was aided by the thermo-sensitive helper plasmid pKOBEGApra, an apramycin resistant derivative of pKOBEG [52], encoding lambda Red recombination functions. Plasmid pKOBEGApra bearing C132-98 was grown at 30°C, induced by addition of 0.2% arabinose (Sigma), and cured of the plasmid at 37°C. Initially, the tetracycline resistance encoding cassette was amplified from pAR82 using the primer pair Ucas (5'-CAAGAATTGCCGGCGGAT-3') and Dcas (5'-GGGCGACACCTT-3'). At their 5'-ends, primers R_upclpK and F_dwnclpK were equipped with 18 bp and 20 bp regions homologous to the extremities of the tetracycline resistance cassette. Subsequently, the deletion fragment was generated by PCR using the primer combination F_upclpK/R_dwnclpK and equimolar amounts of all three fragments as template. The purified PCR product was introduced to electro-competent C132-98 harboring pKOBEGApra followed by tetracycline selection and temperature-induced curing of pKOBEGApra. Correct allelic replacement was verified by PCR.

Construction of an isogenic clpK deletion mutant

The clpK gene in K. pneumoniae C132-98 was deleted by allelic exchange with a PCR synthesized cassette encoding tetracycline resistance flanked by regions homologous to sequences up- and down-stream the plasmid encoded clpK gene by a modification of the lambda Red mediated recombination procedure [50] previously applied to chromosomal gene replacement in K. pneumoniae [51]. Allelic replacement was aided by the thermostable helper plasmid pKOBEGApra, an apramycin resistant derivative of pKOBEG [52], encoding lambda Red recombination functions. Plasmid pKOBEGApra bearing C132-98 was grown at 30°C, induced by addition of 0.2% arabinose (Sigma), and cured of the plasmid at 37°C. Initially, the tetracycline resistance encoding cassette was amplified from pAR82 using the primer pair Ucas (5'-CAAGAATTGCCGGCGGAT-3') and Dcas (5'-GGGCGACACCTT-3') [53]. In addition, 528 bp and 569 bp regions flanking the clpK gene were amplified from C132-98 plasmid DNA by use of primer pair F_upclpK (5'-GGGCGGCTGAGCACGACCT-3')/R_upclpK (5'-ATCCGCCGGAATTTCTTCGCGGTCGATCCCAGTATCCAATCTTCCAAAACAAGGGGGAT-3') and F_dwnclpK (5'-GGGCGGCTGAGCACGACCT-3')/Dcas (5'-GGGCGGCTGAGCACGACCT-3') [53]. At their 5'-ends, primers R_upclpK and F_dwnclpK were equipped with 18 bp and 20 bp regions homologous to the extremities of the tetracycline resistance cassette. Subsequently, the deletion fragment was generated by PCR using the primer combination F_upclpK/R_dwnclpK and equimolar amounts of all three fragments as template. The purified PCR product was introduced to electro-competent C132-98 harboring pKOBEGApra followed by tetracycline selection and temperature-induced curing of pKOBEGApra. Correct allelic replacement was verified by PCR.

Heat shock assays

All heat shock assays were conducted by dilution of stationary phase bacterial cultures 1/10 in pre-heated 0.9% saline (NaCl) solutions at indicated temperatures until sample withdrawal at relevant time intervals. Selection of the Sall3A1 generated plasmid DNA library was performed by 5 min of heat shock of the pooled library clones (384 individual clones) at 58°C followed by reinoculation into fresh LB media. Following outgrowth of the pooled bacteria the combined heat shock and outgrowth procedure was repeated three times. Clonal selection was verified by plasmid restriction analysis of representative randomly picked colonies after growth on LB plates. Heat shock survival curves of individual strains or clones at indicated temperatures were generated by sample withdrawal at appropriate time intervals and the survival rates were estimated by determination of CFU relative to the bacterial counts prior to heat shock.

Hybridization experiments

105 bacteremic isolates of K. pneumoniae were screened for presence of the clpK gene by colony blot hybridization on Amersham Hybond11M-N+ membranes (GE Healthcare). A 437 bp digoxigenin (DIG)-labeled probe was PCR synthesized from K. pneumoniae C132-98 plasmid DNA using primers F_clpkprobe (5'-GGGGGCCCTG-GCGCAACGACCTT-3') and R_clpkprobe (5'-TTTCCGGGCTCT-TCCACCGTGCAACT-3') directed against the sequence corresponding to the unique and conserved linker domain of clpK. Stringent hybridization conditions were applied and immunological detection of clpK positive isolates was performed by the DIG Nucleic Acid Detection Kit according to the directions of the manufacturer (Roche). K. pneumoniae C132-98 and K. pneumoniae C132-98 ΔclpK were used as positive and negative controls, respectively.

Database comparisons and sequence analysis

The nucleotide sequence data reported in this work have been deposited in the GenBank database under accession number FJ042668. Nucleotide sequence analysis was performed by using the Lasergene v. 5.07 software (DNAStar, Inc.). NCBI BLAST searches were performed applying default parameter values [54]. Multiple sequence alignments were conducted by the ClustalW2 algorithm provided by EMBL-EBI [55].

Accession numbers of sequences and proteins mentioned in the text

Escherichia coli ClpK, EFJ96502; Escherichia sp. ClpK, EEH72429; Enterobacter cloacae ClpK, AD63230; Klebsiella pneumoniae C132-98 heat resistance locus, FJ042668; K. pneumoniae MGH78578 plasmid pKN3, NC_009649; Myxobolus volcanius flagellate genome, NC_007947; K. pneumoniae MGH78578 ClpK, ABR80278; M. flagellum ClpK, ABE49438; Desulfovibrio desulfuricans ClpK, ABB40206; Comamonas testosteroni ClpK, EDD5962; Marinobacter aquaeolei ClpK, ABM20464; Pseudomonas mendocina ClpK, ABP85128; Ochrobactrum anthrophi ClpK, ABS17166.

Author Contributions

Conceived and designed the experiments: MSB CS HI DSH KAK. Performed the experiments: MSB CS. Analyzed the data: MSB CS HI DSH KAK. Wrote the paper: MSB CS HI DSH KAK.

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