Klebsiella variicola and Klebsiella quasipneumoniae with capacity to adapt to clinical and plant settings

Esperanza Martínez-Romero, PhD,⁽¹⁾ Nadia Rodríguez-Medina, MSc,⁽²⁾ Marilú Beltrán-Rojel, Biol,⁽²⁾ Jeiry Toribio-Jiménez, PhD,⁽³⁾ Ulises Garza-Ramos, PhD.⁽²⁾

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Abstract

Objective. To compare the genetic determinants involved in plant colonization or virulence in the reported genomes of K. variicola, K. quasipneumoniae and K. pneumoniae. Materials and methods. In silico comparisons and laccard analysis of genomic data were used. Fimbrial genes were detected by PCR. Biological assays were performed with plant and clinical isolates. Results. Plant colonization genes such as cellulases, catalases and hemagglutinins were mainly present in K. variicola genomes. Chromosomal β -lactamases were characteristic of this species and had been previously misclassified. K. variicola and K. pneumoniae isolates produced plant hormones. Conclusions. A mosaic distribution of different virulence- and plant-associated genes was found in K. variicola and in K. guasipneumoniae genomes. Some plant colonizing genes were found mainly in K. variicola genomes. The term plantanosis is proposed for plant-borne human infections.

Keywords: bacterial infections; Gram-negative bacterial infections; Enterobacteriaceae infections Martínez-Romero E, Rodríguez-Medina N, Beltrán-Rojel M, Toribio-Jiménez J, Garza-Ramos U. Klebsiella variicola y Klebsiella quasipneumoniae con capacidad para adaptarse al ambiente clínico y a las plantas. Salud Publica Mex 2018;60:29-40. http://doi.org/10.21149/8156

Resumen

Objetivo. Comparar genes de colonización de plantas o de virulencia en los genomas reportados de K. variicola, K. quasipneumoniae y K. pneumoniae. Material y métodos. Se utilizaron análisis in silico y de Jaccard. Por PCR se detectaron genes de fimbrias. Se realizaron ensayos biológicos con aislados de plantas y clínicos. **Resultados.** Los genes de colonización de plantas como celulasas, catalasas y hemaglutininas se encontraron principalmente en genomás de K. variicola. Las β-lactamasas cromosómicas son características de la especie y en algunos casos estaban mal clasificadas. K. variicola y K. pneumoniae producen hormonas vegetales. **Conclusiones.** Se encontró una distribución en mosaico de los genes de asociación con plantas y de virulencia en K. variicola y K. quasipneumoniae. Principalmente en K. variicola se encontraron algunos genes involucrados en la colonización de plantas. Se propone el término plantanosis para las infecciones humanas de origen vegetal.

Palabras clave: infecciones bacterianas; infecciones de bacterias Gram-negativas; infecciones *Enterobacteriaceae*

(I) Centro de Ciencias Genómicas, Universidad Nacional Autónoma de México. Cuernavaca, Morelos, Mexico.

- Laboratorio de Resistencia Bacteriana, Centro de Investigación sobre Enfermedades Infecciosas, Instituto Nacional de Salud Pública. Cuernavaca, Morelos, México.
- (3) Laboratorio de Biotecnología y Genética Microbiana, Unidad Académica de Ciencias Químico Biológicas, Universidad Autónoma de Guerrero. Chilpancingo, Guerrero, México.

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Corresponding author. PhD. Ulises Garza-Ramos. Departamento de Diagnóstico Epidemiológico, Instituto Nacional de Salud Pública.

Av. Universidad 655 col. Santa María Ahuacatitlán. 62100, Cuernavaca, Morelos, Mexico.

E-mail: ulises.garza@insp.mx

Martínez-Romero E y col.

The taxonomy of the genus *Klebsiella* has been pe-I riodically revised. Currently, this genus includes K. pneumoniae subsp. pneumoniae, also known as KpI, and four novel species: Klebsiella quasipneumoniae, also known as KpII - with two subspecies: quasipneumoniae (KpII-A) and similipneumoniae (KpII-B); Klebsiella variicola (KpIII) and Klebsiella michiganensis.¹ K. variicola and K. quasipneumoniae are sister species of K. pneumoniae. K. variicola has been isolated from plant tissues,^{2,3} fungal gardens of leaf-cutter ant colonies,⁴ cotton disease vectors (specifically of the insect, Nezara viridula),⁵ human⁶⁻¹⁰ and animal infections¹¹ including bovine mastitis.¹² Unlike K. variicola, K. quasipneumoniae has been described exclusively in hospital settings.¹³ A recent genomic comparative study with a large number of Klebsiella genomes showed that the nif operon was detected in all the genomes of K. variicola, in half of those of K. quasipneumoniae, but only in one strain of K. pneumoniae.¹³ Additionally, K. pneumoniae and K. variicola shared some virulence determinants that cause infections in humans.14,15

Some *Klebsiella* isolates have been misclassified^{16,17} and consequently the taxonomic descriptions of their genome sequences in NCBI repositories are inaccurate. Here we used correctly identified *K. variicola* and *K. quasipneumoniae* genomes for the comparison of virulence and plant colonization determinants of this genus of enterobacteria.

Materials and methods K. variicola and K. quasipneumoniae genomes included in the study

A total of 41 *Klebsiella* genomes were included in the study, 31 of which corresponded to *K. variicola*, eight to *K. quasipneumoniae* and two to *K. pneumoniae* (table I). Of these genomes, 19 of *K. variicola* and six *K. quasipneumoniae* were originally misidentified.

In silico analysis of virulence, plant-associated determinants and efflux pumps, regulators, heavy metal resistance and β -lactamases

A set of 138 proteins was selected from previous reports^{18,19} and the Pasteur Institute webpage (http://bigsdb.web. pasteur.fr/klebsiella/klebsiella.html) for their putative roles in host interactions. Among these proteins, 84 were included for their involvement in virulence. Two efflux pump clusters, 11 efflux pump regulators, four heavy metal clusters, three β -lactamase family proteins (LEN-, SHVand OKP-type), as well as 35 plant-association proteins that included nitrogen fixation enzymes (encoded by the *nif* operon with 20 genes), were also included. All genes encoding these proteins/enzymes were further searched in 41 *Klebsiella* spp. genomes by BLAST20 (Genome BLASTp option with default values). A cluster analysis using the average linkage based on the Jacquard similarity coefficient and the respective dendrogram (constructed with UPGMA) were constructed using the DendroUPGMA program.²⁰ In the case of the LEN-, SHV- and OKP-type families of β -lactamases, a phylogenic reconstruction was performed using the maximum-likelihood approach based on the JTT matrix-based model and 100 bootstrap replications (Mega v6.06).²¹ In addition, phylogenetic analyses were carried out on the amino acid sequences of FimV fimbrial proteins (KVR801v1_60088) and NifH nitrogenase (KVR801v1_120019) proteins.

PCR screening of fimV gene

Specific oligonucleotides for the *fimV* gene (fimV-F 5'-TTTGCGGATACTGACCAGGG-3' and fimV-R 5'-GGTTACCACGGTCAGCGTAA-3') were designed. Twenty-one *K. variicola* isolates from plants and humans were analyzed by PCR as previously described.⁸

In vitro assays of plant growth promotion

The mechanisms involved in plant growth promotion, such as phytohormone production (auxin [indole-3-acetic acid] and gibberellins), phosphate solubilization, siderophore production and lytic enzyme activities were evaluated in six K. variicola isolated from plants, and in 15 K. variicola and eight K. pneumoniae clinical isolates. The production of auxin was analyzed according to Khalid and collagues²² with Azotobacter vinelandii and Salmonella enteritidis as positive and negative controls, respectively. Phosphate solubilization was evaluated using the methodology described by Mehta and colleagues²³ using Azotobacter vinelandii and E. coli DH5- α as bacterial controls. Siderophore production was evaluated using chrome azurol S according to Schwyn and Neilands²⁴ with Pseudomonas fluorescens and Staphylococcus aureus as controls. Activities of lytic enzymes, such as lipases, proteases, esterases and amylases, were evaluated using the methodology described by Malleswari and Bagyanarayana²⁵ using *Bacillus subtilis* and *E. coli* DH5- α as controls.

Ethical committee approval

This study is part of the *K. variicola* project that was revised and approved by the ethical commission from the *Instituto Nacional de Salud Pública* on the 21th of June 2011. The present study was carried out at the *Instituto*

Table I ORIGIN AND ACCESSION NUMBER DATA OF K. VARIICOLA, K. QUASIPNEUMONIAE AND K. PNEUMONIAE GENOMES INCLUDED IN THE STUDY

Bacterial specie on GenBank	True bacterial specie*	Isolate	Origin of isolates	Accession number data base	PUBMED (PMID)	
K. pneumoniae	K. pneumoniae	NTUH-K2044	Human (blood)	AP006725.1	19447910	
K. pneumoniae	K. pneumoniae	MGH-78578	Human	CP000647.1	Unpublished	
K. pneumoniae	K. variicola	223/14	Human (pus)	JRTV0000000	Unpublished	
K. pneumoniae	K. variicola	BI	Plant (bitter gourd)	JSWX0000000	Unpublished	
K. pneumoniae	K. variicola	CH4	Plant (chili)	JSXA00000000	Unpublished	
K. pneumoniae	K. variicola	MGH20	Human (respiratory)	AYJK00000000	Unpublished	
K. pneumoniae	K. variicola	MGH40	Unknown (urine)	AYIX00000000	Unpublished	
K. pneumoniae	K. variicola	UCICRE10	Unknown	AYIF0000000	Unpublished	
K. pneumoniae	K. variicola	UCI18	Human (urine)	JCML0000000	Unpublished	
K. pneumoniae	K. variicola	BIDMC61	Human (urine)	JMWB0000000	Unpublished	
K. pneumoniae	K. variicola	MGH68	Human (urine)	JMZD0000000	Unpublished	
K. pneumoniae	K. variicola	MGH76	Human (bile)	JMZK0000000	Unpublished	
K. pneumoniae	K. variicola	MGH80	Human (urine)	JMZM0000000	Unpublished	
K. pneumoniae	K. variicola	BIDMC88	Human	LFBA0000000	Unpublished	
K. pneumoniae	K. variicola	BIDMC90	Human	LFBC0000000	Unpublished	
K. pneumoniae	K. variicola	MGH114	Human	LFAP00000000	Unpublished	
K. pneumoniae	K. variicola	MGH92	Human	LFAD0000000	Unpublished	
Klebsiella sp.	K. variicola	1.1.55	Human	ACXA00000000	Unpublished	
Klebsiella sp.	K. variicola	KTE92	Human	ASQN0000000	Unpublished	
K. pneumoniae	K. variicola	342	Plant (maize stems)	CP000964.1	18654632	
K. pneumoniae	K. variicola	KP5-I	Insect (Nezara viridula)	CP008700.1	25146146	
K. variicola	K. variicola	BZ19	Human (faeces)	JDWA0000000	25135672	
K. variicola	K. variicola	DX120E	Plant (banana roots)	CP009274	Unpublished	
K. variicola	K. variicola	DSM15968	Plant (banana roots)	CP010523	Unpublished	
K. variicola	K. variicola	CAG:634	Human (gut)	CBBA00000000	Unpublished	
K. variicola	K. variicola	801	Human (blood)	CDMV0000000	25886267	
K. variicola	K. variicola	8917	Human (sputum)	CEGG0000000	25858850	
K. variicola	K. variicola	06-268	Human (abscess)	CXOZ0000000	This work	
K. variicola	K. variicola	3	Plant (maize shoots)	CXOY0000000	26358599	
K. variicola	K. variicola	4880	Human (blood)	CXPB0000000	This work	
K. variicola	K. variicola	6A2	Plant (banana root)	CXPC0000000	26358599	
K. variicola	K. variicola	T29A	Plant (sugar cane stem)	CXPA0000000	26358599	
K. variicola	K. variicola	At-22	Insect (fungus matrix)	CP001891.1	19965433	
K. quasipneumoniae‡	K. quasipneumoniae	18A069	Human	CBZM00000000	24958762	
K. quasipneumoniae§	K. quasipneumoniae	07A044	Human	CBZR00000000	24958762	
K. pneumoniae	K. quasipneumoniae‡	UCICRE14	Unknown	AYIC00000000	Unpublished	
K. pneumoniae	K. quasipneumoniae§	12-3578	Human (blood)	AQOC0000000	Unpublished	
K. pneumoniae	K. quasipneumoniae§	ATCC 700603	Human	AOGO0000000	23723407	
K. pneumoniae	K. quasipneumoniae§	MGH44	Unknown (respiratory)	AYIV00000000	Unpublished	
K. variicola	K. quasipneumoniae§	HKUOPLA	Panda (feces)	CP012252	26472841	
K. pneumoniae	K. quasipneumoniae§	HKUOPLC	Panda (feces)	CP012300	26564041	

* The true bacterial specie was described previously by Chen and colleagues,¹⁶ and Martinez-Romero and colleagues.¹⁷ * Klebsiella quasipneumoniae subsp. quasipneumoniae

§ Klebsiella quasipneumoniae subsp. similipneumoniae

Nacional de Salud Pública and at the Centro de Ciencias Genómicas of UNAM, Cuernavaca, Morelos.

Results and discussion

Virulence-associated determinants in K. variicola and K. quasipneumoniae genomes

In silico screening of 84 virulence-determinant proteins in reported genomes of *K. variicola, K. quasipneumoniae* and *K. pneumoniae* showed a mosaic distribution in different isolates (table I). Most (>98%, except for FimV) of the virulence determinants that were selected here were originally described in *K. pneumoniae* NTUH-K2044. From the 84 virulence determinants included in the *in silico* analysis, 20 were found in at least one *K. variicola* genome and 11 in at least one *K. quasipneumoniae* genome. The proteins coded by the urease (*UreA*) and fimbriae gene cluster *MrkABCDFHIJ* were present in all the isolates of *K. variicola* and *K. quasipneumoniae*. The gene coding glucuronic acid transferase (WabG) was also present in all *K. variicola* and *K. pneumoniae*, but was absent in a single *K. quasipneumoniae* genome.

Siderophores, such as salmochelin (IroN), aerobactin (IucA and receptor IutA), KfuABC cluster, enterobactin (EntB) and yersiniabactin cluster YbtAESTX, were unequally distributed among *Klebsiella*. The most prevalent siderophores were enterobactin and KfuABC, in 93.5% (29/31) of *K. variicola* and in 100% (8/8) of the *K. quasipneumoniae* isolates. The aerobactin (IucA) was not identified, however the receptor of aerobactin (IutA) was present in 87% (27/31) and salmochelin (IroN) in 39% (12/31) of *K. variicola* isolates; however, in the *K. quasipneumoniae* genomes, both aerobactin receptor and salmochelin siderophore were present in 100% of the isolates. The siderophore yersiniabactin YbtAESTX cluster was identified in only one *K. variicola* isolate (MGH20) and in none of the *K. quasipneumoniae* genomes.

The genes encoding enzymes from the glycerate pathway (Gcl, GlxK), GlxR, and Hyi) were identified in *K. variicola* isolates (41.9 to 45.1%). The allantoinase cluster *AllABCDRS* genes and the YlbE-F and YbbW were present in few *K. variicola* genomes at 3.2, 6.4 and 3.2%, respectively. The GlxK-R and Hyi encoding genes were present only in *K. quasipneumoniae* genomes, in 75% of them. The genes for the two-component system KvgAS proteins were identified in 12.9% (4/31) of the *K. variicola* genomes. The mucoviscosity-associated protein Wzy-K1 was identified in one *K. variicola* isolate (3.2%-1/31).

The Jacquard index analysis of the virulenceassociated determinants that are shared between the *K. variicola, K. quasipneumoniae* and *K. pneumoniae* genomes is shown in figure 1. Three main clusters were obtained, two of which were closely related and could be distinguished only by a single siderophore difference. Cluster 1 grouped sixteen K. variicola and two K. quasipneumoniae subsp. similipneumoniae genomes that shared the siderophores IroN (11/18 genomes); receptor IutA, KfuABC and EntB (15/18 genomes); WabG, Uge, the fimbriae cluster MrkABCDFHIJ, FimV (15/18 genomes) and urease UreA. Cluster 2 includes ten K. variicola genomes and mostly contains the virulence-determinants proteins of cluster 1, except for IroN. Genes associated with all of the K. variicola genomes from cluster 2 were those encoding Glc, GlxKR and Hyi proteins that are involved in the glycerate pathway. This pathway together with allantoinase cluster genes are involved in allantoin metabolism as a nitrogen source.²⁶ Cluster 3 consists of seven K. quasipneumoniae and one K. variicola genomes. All of the genomes of cluster 3 contain the Uge, WabG, IroN (except K. variicola KTE92 genome), IutA, KfuABC, EntB, UreA and MrkABCDFHIJ proteins. FimV protein was present in three (42.8%-3/7) of the K. quasipneumoniae genomes, corresponding to Klebsiella quasipneumoniae subsp. similipneumoniae (KpII-B).

K. pneumoniae NTUH-K2044 and *K. pneumoniae* MGH78578 each independently clustered together with *K. pneumoniae* BIDMC90 and *K. variicola* DSM15298, respectively (figure 1). The *K. pneumoniae* BIDMC90 genome shared with *K. pneumoniae* NTUH-K2044 fifteen virulence determinants. Finally, *K. pneumoniae* MGH78578 and *K. variicola* DSM15298 were different from the other clusters due to the absence of siderophore KfuABC, which was present in all of the *K. variicola*, *K. quasipneumoniae* and *K. pneumoniae* NTUH-K2044, and to the presence of the two-component system KvgAS, which was only present in the DX120E, MGH76 and UCICRE10 genomes (figure 1).

In silico analysis and prevalence of Fimbrial FimV protein

Previously a total of 114 unique proteins were identified in *K. variicola* genomes.⁸ Excluding transposons and *tra*-genes that are involved in horizontal transfer and selecting those that are involved in metabolism and cellular structure, 13 structural proteins were found to be unique for *K. variicola*, from which a fimbrial protein was chosen for further analysis (fimbria have been shown to mediate host interaction). The fimbrial FimV protein was identified in 90.3% (28/31) of the *K. variicola* (except for KTE92, B1 and BIDMC90) and in 37.5% (3/8) of *K. quasipneumoniae* but not in *K. pneumoniae* NTUH-K2044 and MGH78578 genomes. An additional BLASTp search in GenBank



FIGURE 1. JACQUARD INDEX OF THE VIRULENCE-DETERMINANT PROTEINS OF THE K. VARIICOLA (KV), K. QUASI-PNEUMONIAE (KQ) AND K. PNEUMONIAE (KPN) GENOMES

using a non-redundant protein sequence (nr-BLASTp protein-protein BLAST) identified the FimV protein in *K. variicola* genomes described in GenBank (except for KTE92, B1 and BIDMC90) and in the misclassified *K. pneumoniae* genomes that actually corresponded to *K. variicola* species (table I). The phylogenetic analysis of FimV proteins generally showed high amino acid conservation in *K. variicola* (>98%) (data not shown);

however, in a few *K. variicola* strains, for example the wrongly named "*K. pneumoniae* 342", a lower amino acid identity was observed (92 to 94%). Conversely, *K. quasipneumoniae* subsp. *similipneumoniae*, 07A044, 700603 and MGH44 FimV amino acid sequence had 91% similarity. FimV protein sequence was not found in *bona fide K. pneumoniae*; nevertheless it was identified in related *Enterobacteriaceae* such as *Citrobacter koseri*,

Citrobacter freundii, Enterobacter, Salmonella enterica and *Escherichia albertii,* all with an amino acid similarity less than 53%. The FimV protein is also widely distributed in *Escherichia coli* and *Shigella sonnei,* with an amino acid identity of 33% to that from *K. variicola.*

The PCR screening of the *fimV* gene was carried out in a collection of twenty-one *K. variicola* isolates (see Materials and methods). The *fimV* gene was identified in 86.9% (20/23) of the *K. variicola* isolates. This prevalence is similar to that from *in silico* analysis (90.3%). This indicates that the *fimV* gene is not universally encoded in *K. variicola* genomes.

Plant-associated proteins in the K. variicola, K. quasipneumoniae and K. pneumoniae genomes

Plant-associated proteins were identified in K. variicola and K. quasipneumoniae isolates. Twenty-seven plantassociated determinants were found in at least one K. variicola or K. quasipneumoniae isolate. In general, 21 plantassociated determinants were contained in all of the K. variicola, K. quasipneumoniae and K. pneumoniae genomes that were included in the study. The NifI-NifO nitrogen fixation cluster was identified in all of the K. variicola genomes, in 62.5% (5/8) of K. quasipneumoniae genomes and was absent from the K. pneumoniae NTUH-K2044 and MGH78578 genomes. The NifH protein has high amino acid identity among all K. variicola genomes and in a phylogenetic analysis it is grouped in a cluster different from the corresponding sequence from the K. quasipneumoniae or other species. Holt and colleagues¹³ identified only one K. pneumoniae genome that contained the nifJ-nifQ operon. In this work, we identified another K. pneumoniae genome (from KPNIN29) that contains the nif operon in addition to that described by Holt and colleagues.¹³ Hazen and col*leagues*²⁷ identified the absence of *nifJ-nifQ* gene cluster in K. pneumoniae NTUH-K2044, MGH-78578, 1162281, JH1, MS 92-3, 1191100241, ATCC13884 and KCTC-2242 genomes. Here we identified a conserved genetic context and gene synteny of *nifJ-nifQ* operon both in K. variicola and K. quasipneumoniae genomes (data not shown). Our results confirm that nitrogen fixation seems to be a characteristic trait of K. variicola.

Other differences identified in this work between the *K. variicola, K. quasipneumoniae* and *K. pneumoniae* genomes are genes encoding cellulases (CelK and BglH), catalases (KPK_4954 and KPK_2333) and hemagglutinins (HecA). While BglX cellulose is present in all *K. variicola, K. quasipneumoniae* and *K. pneumoniae* isolates, CelK and BglH were absent from *K. pneumoniae* NTUH-K2044 and MGH78578 and from 87.5% of the *K. quasipneumoniae* isolates. The BglH enzyme, which has specificity towards 1,4-b glucosidic bonds and that most likely acts by hydrolyzing short cello-oligosaccharides, was present in 61.2% (19/31) of *K. variicola* isolates. The CelK gene encoding an enzyme for the decomposition of highly ordered forms of insoluble cellulose was present in *K. variicola* strains obtained from three different plants 342, T29A and 6A2 representing 10% (3/31) of the isolates. KPK_2233 (catalase) gene was present in 22.5% of the *K. variicola* genomes (7/31). KPK_4954 gene encoding a cyclic beta 1-2 glycan synthase possibly playing a role in osmotic adaptation was present in 16.1% of the *K. variicola* isolates (5/31).

HecA/B hemolysin/hemagglutinin secretion protein is involved in plant attachment in *Erwinia chrysanthemis*. A *hecA* gene mutant had reduced attachment, cell aggregate formation, and virulence in its host. HecA protein was present in 64.5% (20/31) of the *K. variicola* isolates. In *K. quasipneumoniae*, HecA protein is present only in HKUOPLA²⁸ and HKUOPLC²⁹ isolates obtained from giant panda feces.

The Jacquard index analysis showed two main clusters (A and B) that were grouped by the absence or presence of HecA, CelK or BglH protein (figure 2). Cluster A included the *K. variicola* and *K. quasipneumoniae* genomes that did not contain the HecA and CelK proteins. The *K. quasipneumoniae* 18A069 genome is the unique genome of this species that does not contain the YqeF (acetyl-CoA acetyltransferase), induce plant colonization and Ada (a regulatory protein of adaptive response) and DinF (DNA-damage-inducible protein F) putative stress response proteins.¹⁸

Efflux pump, regulators, heavy metal resistance and chromosomal β -lactamase proteins in K. variicola, K. quasipneumoniae and K. pneumoniae

The OqxABR efflux pump gene clusters are present in all *K. variicola* and *K. quasipneumoniae* genomes examined. The AcrABR protein efflux was identified in 58.0% (18/31) and 100% (8/8), of *K. variicola* and *K. quasipneumoniae*. All of the protein regulators were identified (MarAR, SoxSR, RamAR, Rob, SdiA, Fis, EnvR, and RarA) in >98% and 100%, of the *K. variicola* and *K. quasipneumoniae* isolates. All efflux pumps and efflux pump regulator proteins were identified in the *K. pneumoniae* NTUH-K2044 and MGH78578 genomes.

The copper (PcoABCDERS), silver (SilCERS) and tellurium (TerABCDEWXYZ) protein clusters for heavy metal resistance were analyzed. The PcoABCDERS and SilCERS protein clusters were identified together in 22.5% (7/31) and 37.5% (3/8) of the *K. variicola* and



FIGURE 2. JACQUARD INDEX OF THE PLANT-DETERMINANT PROTEINS OF THE K. VARIICOLA (KV), K. QUASIPNEU-MONIAE (KQ) AND K. PNEUMONIAE (KPN) GENOMES

K. quasipneumoniae genomes, respectively, while the *K. variicola* Bz19 genome contained only the PcoABCDERS protein cluster.

Tellurite resistance is conferred by the TerABC-DEWXYZ protein cluster and seemingly is strongly associated with the hypervirulent clonal groups.³⁰ Tellurite resistance is needed to colonize macrophages.³¹ The TerABCDEWXYZ proteins cluster was identified in 12.9% (4/31) and 12.5% (1/8), of K. variicola and K. quasipneumoniae, all of human origin. K. variicola 8917 isolate was described as hypermucoviscous.¹⁵

In the analysis of chromosomal β -lactamase, we found that all the *K. pneumoniae* genomes that were identified as *K. variicola* contained the LEN-type β -lactamases

allele. The *K. pneumoniae* genomes that were correctly identified as *K. quasipneumoniae* contained the OKP-type alleles, while the *K. pneumoniae* NTUH-K2044 and MGH78578 genomes contained the SHV-type alleles. The chromosome-encoded β -lactamases corresponded to constitutive genes of these bacterial species (figure 3A). The OKP-A1 and OKP-B1 β -lactamases is characteristic of *K. quasipneumoniae* subsp. *quasipneumoniae* (KpII-A) and *K. quasipneumoniae* subsp. *similipneumoniae* (KpII-B), respectively.

 β -lactamase genes were present in most of the *K*. *variicola* and *K*. *quasipneumoniae* genomes and were previously annotated as "Beta-lactamase or class A

beta-lactamase". In *K. pneumoniae* B1 and *K. variicola*, BZ19 the corresponding sequence was annotated as "Beta-lactamase TEM" and had an amino acid identity of 99 and 100%, respectively, with LEN-2 β -lactamase. In the *Klebsiella* sp. 1.1.55 genome, β -lactamase was annotated as "LEN family Class A β -lactamase" and had a 99% of amino acid identity with LEN-2. Figure 3B shows the phylogenetic analysis of *K. quasipneumoniae* chromosomal β -lactamase genes. The β -lactamases were annotated as class A beta-lactamase, or SHV-1 or TEM proteins. However, all these proteins corresponded to OKP-A1 and OKP-B1 β -lactamases, respectively, in subspecies of *K. quasipneumoniae* (KpII-A, KpII-B)



FIGURE 3. PHYLOGENETIC ANALYSIS OF CHROMOSOMAL B-LACTAMASES. A) LEN-, SHV- AND OKP-TYPE B-LACTAMASES THAT WERE IDENTIFIED IN THE K. VARIICOLA, K. QUASIPNEUMONIAE AND K. PNEUMONIAE GENOMES. B) CHROMOSOMAL B-LACTAMASE PROTEIN THAT WAS ANNOTATED IN K. QUASIPNEUMONIAE GENOMES AND THAT CORRESPONDED TO THE OKP FAMILY. HPLA AND HPLC, RESPECTIVELY, CORRESPONDS TO K. QUASIPNEUMONIAE SUBSP. SIMILIPNEUMONIAE HKUOPLA AND HKUOPLC

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(figure 3B). Chromosomal LEN- and OKP- β -lactamase proteins were not properly annotated and could have contributed to the bacterial species misclassification, if these chromosomal β -lactamase proteins are considered for *Klebsiella* classification.

In vitro assays of plant growth-promoting mechanisms in K. variicola and K. pneumonia

Indole acetic acid (an auxin) has the ability to regulate root growth, differentiation of vascular tissues, elongation, apical dominance, initiation of lateral roots and maturation.³² This molecule also functions as an important signaling molecule in the regulation of plant growth, expansion, cell differentiation and cell division regulation.³³ *K. variicola* and *K. pneumoniae* isolates were capable of producing indole acetic acid (table II).

Gibberellins (GA) are involved in seed germination, seedling emergence, stem and leaf growth, flower induction, vegetative regulation of reproductive dormancy buds and fruit growth.^{34,35} Almost all of the *K. variicola* and *K. pneumoniae* isolates (except for two *K. pneumoniae* isolates) were capable of producing gibberellins.

In addition to nitrogen, phosphorus is a nutrient that is required by plants.³⁶ K. variicola and K. pneumoniae isolates were not able to solubilize phosphate (table II), which is a common characteristic among plant associated bacteria. Klebsiella genus, unlike other plant growth promoting bacteria, did not produce chitinases or proteases that may help fighting pathogenic fungi of plants (table II). Siderophores are synthesized mainly by Gram-negative bacteria, fungi, yeast and some plants (phytosiderophores) and act as specific chelating agents. They are soluble in aqueous solutions at neutral pH and are considered as secondary metabolites.³⁷ Siderophores may be important in the process of colonizing plant and human tissues.³⁸ Different siderophores were identified in silico in the genome of K. variicola, K. quasipneumoniae and K. pneumoniae. In agreement with these findings, the in vitro siderophore assay was positive for all of the K. variicola and K. pneumoniae isolates that were tested.

The name *K. variicola* means from various places and novel reports confirmed the adequacy of its name. This species has now been identified in diverse plants, such as bitter gourd and chili, as well as in insects such as *Nezara viridula*. It is also significant that isolates have been obtained from different human samples, such as urine, pus and bile.¹⁷

K. variicola isolates were additionally obtained from the fecal microbiota of two cohorts of Malawian infants/ children, but their genomes could not be included in this work because they are fragmented into a high number of contigs. Of note, the *K. variicola* strain has been identified in giant panda feces,²⁸ and corresponds to other example of misclassification and corresponds to *K. quasipneumoniae*.¹⁷ The first OXA-181 (carbapenemase)-producing *K. variicola* isolate was identified (by *rpoB* gene analysis) in fresh vegetables that were commercialized between different continents (Asia-Europe).³⁹ This work showed the fist carbapenem-resistance *K. variicola* strain and may suggest possible emergence paths of resistance in *Enterobacteriaceae*.

Gene flow may occur among different related *Klebsiella* species, mediated by plasmid transfer. The plasmid of *K. variicola* pBz19 showed high identity to pl9 (both associated with IncN incompatibility) from *K. pneumoniae* isolate.¹⁴ *K. variicola* DX120E contains pKV1 and pKV2 plasmids. The pKV1 plasmid is very similar to the pKp5-1 plasmid that was identified in *K. pneumoniae* KP5-1.¹⁹ Plasmid pKV2 is most similar to plasmid pKOXM1C from *K. oxytoca* strain M1, suggesting plasmid exchange between these bacterial species. Additional analyses of plasmids are required specially in *K. variicola* from different niches to further understand their role in host interaction.

Conclusions

This work provides a new molecular framework for distinguishing different organisms of the Klebsiella genus. The in silico analyses showed that the genomes of K. variicola and K. quasipneumoniae shared a set of virulence-associated genes encoding mainly siderophores (aerobactin and enterobactin KfuABC cluster), urea metabolism, lipopolysaccharide biosynthesis enzymes (WabG and Uge), and fimbriae genes encoding the MrkABCDFHIJ gene cluster. The in vitro analysis indicated the presence of different siderophores in distinct isolates and species. Siderophores may have redundant functions given that they all are involved in bacterial iron acquisition. The number of siderophores in a single cell and their type however, may be important to regulate the flow of iron incorporation into the bacterial cell, a process that is linked to invasion of host tissues and thus virulence.

FimV, on the other hand, could be considered a common character in *K. variicola* strains distinguishing them from those of *K. pneumoniae*; however, not all *K. variicola* genomes contained this gene, as shown by both *in silico* and PCR assays. To explain the mosaic distribution of putative host-interaction genes in *Klebsiella* genomes, we envisage the possibility that *Klebsiella* cells may acquire virulence genes by horizontal transfer from closely related bacteria. However, this possibility is tempered by the alternative process of gene loss in bacterial evolution, which could be rather significant in the conformation of

laslates		Indirect mechanism			
isolates	Indole acetic acid (mg/ml)*	Solubilization of phosphorus index (mm)	Gibberellins	Siderophores	Lytic enzymes
K. variicola					
F2R9 [⊤]	17.19 ± 0.740	1.83	+	+	-
CFNE 2600	20.63 ± 0.191	1.63	+	+	-
T29A	19.96 ± 2.014	1.99	+	+	-
VI	21.81 ± 1.155	2.22	+	+	-
3	19.47 ± 2.532	2.1	+	+	-
6A2	20.03 ± 0.172	1.3	+	+	-
7	18.09 ± 0.871	1.46	+	+	-
1258	20.01 ± 0.827	2.04	+	+	-
4880	18.58 ± 1.022	2.18	+	+	-
8917	19.63 ± 0.894	1.88	+	+	-
9326	18.28 ± 0.806	1.68	+	+	-
9351	19.95 ± 1.375	2.12	+	+	-
9352	18.65 ± 0.841	2.4	+	+	-
9353	19.17 ± 0.477	1.81	+	+	-
9387	17.27 ± 0.376	1.59	+	+	-
9388	20.22 ± 0.543	2.63	+	+	-
9635	18.00 ± 0.455	1.48	+	+	-
9925	19.63 ± 0.463	0	+	+	-
K. pneumoniae					
6419	17.63 ± 2.13	1.6	-	+	-
6421	17.09 ± 2.57	1.2	+	+	-
9018	16.71 ± 1.56	1.9	-	+	-
1610	16.32 ± 0.73	2.2	+	+	-
06-208	16.88 ± 2.53	2.1	+	+	-
01-250	15.90 ± 0.97	1.5	+	+	-
2989	18.12 ± 1.54	2.6	+	+	-
3407	15.80 ± 1.19	1.4	+	+	-
Control strains					
Azotobacter vinellandii	5.65 ± 0.342	5.8	+	+	NA
Salmonella enteritidis	0.002 ± 0.00	NA	-	NA	NA
Escherichia coli DH5-a	NA	0.0	NA	NA	-
Staphylococcus aureus	NA	NA	NA	-	NA
Pseudomonas fluorecens	NA	NA	NA	+	NA
Bacillus subtilis	NA	NA	NA	NA	+

Table II In vitro assays of plants growth-promoting mechanisms in K. variicola and K. pneumoniae isolates

* The in vitro assay was carried by triplicated and the standard errors are shown

NA; not applied, +; positive, -; negative

Klebsiella genomes as has been shown in other bacteria.⁴⁰ *K. variicola* and *K. quasipneumoniae* strains with different genetic repertoires have the capacity to adapt to plant and clinical settings. Previously, we suggested that a different epidemiological dynamics occurred with *K.*

variicola in comparison to *K. pneumoniae* with *K. variicola* coming from the environment,⁴¹ we further refine here that the environment refers most probably to the plant environment. Plants can be a reservoir for *K. variicola* isolates that may opportunistically infect humans or

animals. *K. variicola* infection in humans thus seems to be a case of "phytonosis", a term we suggest for symbiotic bacteria plant-borne, parallel to the term zoonosis bacteria pathogen animal-borne.

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 $\ensuremath{\textit{Declaration}}$ of conflict of interests. The authors declare that they have no conflict of interests.

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