Draft Whole-Genome Sequences of Five *Klebsiella pneumoniae* Isolates from the Subantarctic Islands of New Zealand

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**ABSTRACT** *Klebsiella pneumoniae* is a Gram-negative bacterium that can be found in the environment, as well as on mucosal surfaces of humans and animals. Here, we report the genome sequence of five *K. pneumoniae* isolates from substrate samples and bird feces collected in the Subantarctic Islands of New Zealand.

*Klebsiella pneumoniae* is an opportunistic pathogen causing nosocomial and community-acquired infections (1, 2). Over the past few decades, hypervirulent strains causing primary liver abscesses and septicemia have increasingly been documented (3, 4). Most hypervirulent strains have a hypermucoviscous phenotype (positive string test) and the *rmpA* and *rmpA2* genes (1, 3, 4). Hypervirulent strains have also been described in animals (5–7).

Hypervirulent *K. pneumoniae* infection has been reported as a cause of mortality events in New Zealand sea lion (NZSL) pups on Enderby Island, New Zealand, one of the Subantarctic Auckland Islands (7, 8). Annually, a large number of pups die from *K. pneumoniae* infection in the summer birthing season; however, reservoirs have not been investigated. Environmental substrates, birds that live on the island, and NZSL adults are possible reservoirs of this infection.

The use of animals and sample collection were completed under permit 39915-FAU from the New Zealand Department of Conservation and Massey University Animal Ethics Committee (approval 14/114). In this study, five *K. pneumoniae* isolates from cloacal swabs or voided feces from subantarctic skuas (*n* = 2) and a yellow-eyed penguin (*n* = 1) and substrate samples (water; *n* = 2) (using CHROMagar Orientation) were whole-genome sequenced. The NucleoSpin soil kit (Macherey-Nagel, GmbH & Co. KG, Germany) was used to extract genome-quality DNA from a single colony cultured on agar, which was sent to New Zealand Genomics Limited (Massey Genome Service, Massey University, Palmerston North, New Zealand). A fragment library was prepared using an Illumina TruSeq DNA library preparation kit v1 (Illumina, Inc., San Diego, CA). The five isolates from this study were *de novo* assembled using SPAdes v3.10 (in the “careful” mode) (9). The contigs of each isolate produced from SPAdes were annotated by Prokka v1.1.2, with default parameters (10). The sequence types (ST) and serotypes of bacterial isolates were determined using the Bacterial Isolate Genome Sequence Database (BIGIsdb) servers (http://bigisdb.pasteur.fr/klebsiella/klebsiella.html). Virulence genes were identified by mapping the reads of each isolate, along with 1,000-bp flanks on either side of the virulence gene sequence, using Bowtie 2 in both the “--very-sensitive-local” and “--very-sensitive” modes for local and global read mapping, respectively. Genome sizes, numbers of contigs, ST, sources, serotypes, and virulence genes are summarized in Table 1.

The virulence genes found in environmental and bird isolates were consistent with
<table>
<thead>
<tr>
<th>Isolate</th>
<th>GenBank accession no.</th>
<th>SRA accession no.</th>
<th>Source, location</th>
<th>Serotype</th>
<th>ST</th>
<th>String test result</th>
<th>Length (bp)</th>
<th>No. of contigs</th>
<th>Fold coverage</th>
<th>GC content (%)</th>
<th>Virulence genes</th>
</tr>
</thead>
<tbody>
<tr>
<td>E14_15_17Sa</td>
<td>QVNJ000000000</td>
<td>SRR7699496</td>
<td>Subantarctic skua, a</td>
<td>Non-K1/K2</td>
<td>2843</td>
<td>-</td>
<td>5,644,626</td>
<td>116</td>
<td>87</td>
<td>57.1</td>
<td>wabG, uge, ipi2, iucD, iutA, mrkD, rmpA</td>
</tr>
<tr>
<td>E14_15_42Sa</td>
<td>QVN100000000</td>
<td>SRR7699497</td>
<td>Subantarctic skua, a</td>
<td>K2</td>
<td>86</td>
<td>+</td>
<td>5,327,210</td>
<td>94</td>
<td>93</td>
<td>57.5</td>
<td>rmpA, wabG, uge, iroN, iip2, ybtS, mrkD</td>
</tr>
<tr>
<td>E14_15_53Ma</td>
<td>QVNH000000000</td>
<td>SRR7699498</td>
<td>Yellow-eyed penguin, a</td>
<td>K2</td>
<td>86</td>
<td>+</td>
<td>5,351,349</td>
<td>91</td>
<td>89</td>
<td>57.5</td>
<td>rmpA, wabG, uge, iroN, iip2, ybtS, mrkD</td>
</tr>
<tr>
<td>E13_14_10sub</td>
<td>QVNG000000000</td>
<td>SRR7699499</td>
<td>Water, Enderby Island</td>
<td>K2</td>
<td>86</td>
<td>+</td>
<td>5,332,787</td>
<td>85</td>
<td>98</td>
<td>57.5</td>
<td>rmpA, wabG, uge, iroN, iip2, ybtS, mrkD</td>
</tr>
<tr>
<td>C14_15_17sub</td>
<td>QVNF000000000</td>
<td>SRR7699495</td>
<td>Water, Campbell Island</td>
<td>K2</td>
<td>86</td>
<td>+</td>
<td>5,323,133</td>
<td>99</td>
<td>94</td>
<td>57.5</td>
<td>rmpA, wabG, uge, iroN, ybtS, mrkD</td>
</tr>
</tbody>
</table>

*This animal was apparently healthy. *K. pneumoniae* was isolated from cloacal swabs or voided feces from nonclinical, live animals.
those found in other studies on clinical human isolates (11), suggesting that the isolates have pathogenic potential. This further suggests that the environment and birds may be possible reservoirs of this pathogen. The ST of the environmental isolates from this study was different from that in previous reports on other environmental isolates (12), suggesting genetic diversity of this bacterium in the environment (13).

These are the first whole-genome sequences of \textit{K. pneumoniae} isolated from birds and environmental substrates from the Subantarctic Islands. The data from this study will provide information on genomomic relationships between isolates from animals and environmental substrate samples from the Subantarctic and help to inform future research on the role of \textit{K. pneumoniae} in harsh non-human-dominated island environments.

\textbf{Data availability.} The whole-genome shotgun sequences described here have been deposited in DDBJ/ENA/GenBank under the accession numbers listed in Table 1. Rare sequence reads have been deposited in the NCBI Sequence Read Archive under the accession numbers listed in Table 1.

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\textbf{REFERENCES}


