Recovery of *Bordetella bronchiseptica* sequence type 82 and *B. pseudohinzii* from urban rats in Terengganu, Malaysia

Shih Keng LOONG¹, Nurul-Asma-Anati CHE-MAT-SERI¹, Osama ABDULRAZAK², Benacer DOUADI³, Siti-Noraisah AHMAD-NASRAH⁴, Jefree JOHARI¹, Siti-Nursheena MOHD-ZAIN³ and Sazaly ABUBAKAR¹,⁴*

¹Tropical Infectious Diseases Research & Education Centre, Faculty of Medicine, University of Malaya, 50603 Kuala Lumpur, Malaysia
²Microbial Evolutionary Dynamics Research Group, Department of Evolutionary Theory, Max Planck Institute for Evolutionary Biology, 24306 Plön, Germany
³Institute of Biological Sciences, Faculty of Science, University of Malaya, 50603 Kuala Lumpur, Malaysia
⁴Department of Medical Microbiology, Faculty of Medicine, University of Malaya, 50603 Kuala Lumpur, Malaysia

ABSTRACT. Rodents have historically been associated with zoonotic pandemics that claimed the lives of large human populations. Appropriate pathogen surveillance initiatives could contribute to early detection of zoonotic infections to prevent future outbreaks. *Bordetella* species are bacteria known to cause mild to severe respiratory disease in mammals and, some have been described to infect, colonize and spread in rodents. There is a lack of information on the population diversity of bordetellae among Malaysian wild rodents. Here, bordetellae recovered from lung tissues of wild rats were genotypically characterized using 16S rDNA sequencing, MLST and *nrdA* typing. A novel *B. bronchiseptica* ST82, closely related to other human-derived isolates, was discovered in three wild rats (n=3) from Terengganu (5.3333° N, 103.1500° E). *B. pseudohinzii*, a recently identified laboratory mice inhabitant, was also recovered from one rat (n=1). Both bordetellae displayed identical antimicrobial resistance profiles, indicating the close phylogenetic association between them. Genotyping using the 765-bp *nrdA* locus was shown to be compatible with the MLST-based phylogeny, with the added advantage of being able to genotype non-classical bordetellae. The recovery of *B. pseudohinzii* from wild rat implied that this bordetellae has a wider host range than previously thought. The findings from this study suggest that bordetellae surveillance among wild rats in Malaysia has to be continued and expanded to other states to ensure early identification of species capable of causing public health disorder.

KEY WORDS: bordetellae, infectious disease, MLST, *nrdA*, tropical
its zoonotic potential, no study has been conducted to investigate the diversity and clonal relationships among the bordetellae populations maintained in wild rodents. Reports of whooping cough and other pulmonary symptoms in humans living close to animals infected with *B. bronchiseptica* suggests that the infections originated from infected animals [12, 27, 32]. Even though there has not been any report of human *B. bronchiseptica* infection in Malaysia, all these past cases underscore the possibility of zoonotic transmission of bordetellae to humans or to other animals, and these animals can then serve as carrier in the transmission cycle [23]. Detection of *B. bronchiseptica* from livestock, domestic pets and wild animals in Africa, Asia, Europe and the U.S.A. have been documented, illustrating the global distribution of this species and the diversity of hosts it occupies [14, 25, 27, 33]. *B. pseudohinzii* also has a relatively wide geographical distribution evident by its recovery in Germany, Japan, Malaysia and the U.S.A., although this bacterium has only been isolated from laboratory-raised rodents [16].

Here, we recovered bordetellae from rodents captured in the wet markets and identified them to subspecies level using molecular tools. Multi locus sequence typing (MLST) and *nrdA* typing were utilized to determine the phylogenetic relationships among the different bordetellae isolates, which could shed useful insights into factors that shape the genetic structure of bordetellae. Sequence analysis of *nrdA*, encoding the ribonucleoside-diphosphate reductase alpha chain, has been demonstrated to reliably differentiate species within the *Bordetella* genus [21, 28]. In addition, the antimicrobial resistance and virulence profiles of the bordetellae isolates were also determined.

### MATERIALS AND METHODS

**Rodent trappings**

Prior approvals were obtained from the local municipal councils to accompany their pest eradication teams to conduct rodent trappings within wet markets in several states in Peninsular Malaysia. Wet markets in four states were included; Johor (1.4927° N, 103.7414° E), Kedah (6.1248° N, 100.3678° E), Kelantan (6.1168° N, 102.2777° E) and Terengganu (5.3333° N, 103.1500° E) (Fig. 1). Rodent trappings were conducted over a period of 14 months, from November 2014 until December 2015. Steel wire traps (18 × 12 × 28 cm) were placed on floors in the wet market in the evening with baits such as dried fish, bread or peanut butter. The traps were collected early in the morning before the market was opened [3]. All rodents were captured alive and their species were determined before they were euthanized at the field laboratory according to protocols described elsewhere [1]. Lung tissues were harvested and homogenized, followed by inoculation onto Columbia agar according to previously published methods [22]. Cross contamination between tissue specimens from different individual rodents was prevented by washing the surgical tools with...
alcohol, bleach and water, in between surgical procedures as previously described [20]. This study was approved with the ethics reference no. ISB/31/01/2013/SNMZ (R) by the Institutional Animal Care and Use Committee and did not involve any endangered or protected species.

Antimicrobial susceptibility testing and amplification of virulence genes

All mixed cultures grown from the rodent lung homogenates were sub-cultured until pure cultures were obtained. Chromosomal DNA of all pure-cultured bacteria isolates was extracted using Nucleospin Tissue kit (Macherey-Nagel, Düren, Germany) according to the manufacturer’s instructions. Subsequently, 16S rDNA sequencing [18] was performed to identify the respective isolates’ species identity. Only members of the genus *Bordetella* were selected for this study and they were stocked in −80°C until further use. Antimicrobial susceptibility profiles were determined using M.I.C.Evaluator Strips (Thermo Fisher Scientific, Waltham, MA, U.S.A.) on Mueller-Hinton agar. The following antibiotics were utilized to determine the minimum inhibitory concentrations (MIC); amikacin, amoxicillin-clavulanic acid, ampicillin, cefazidime, ciprofloxacin, ceftriaxone, cefotaxime, erythromycin, gentamicin, imipenem and meropenem. Antimicrobial susceptibility results were interpreted according to Clinical and Laboratory Standards Institute breakpoints for *Enterobacteriaceae* [5, 24]. The presence of the virulence genes; dermonecrotic toxin (*dnt*), exogenous ferric siderophore receptor (*bfrZ*) and flagella (*fla*), were examined using nucleic acid amplification protocols previously described [29].

Multi locus sequence typing and *nrdA* typing

DNA fragments from seven housekeeping genes (*adk, fumC, glyA, tyrB, icd, pepA* and *pgm*) were amplified, trimmed to a standard length and compared with the *Bordetella* MLST database at http://pubmlst.org/bordetella/ [6]. A novel allele number was assigned to allele sequences that are not available in the database, resulting in the assignment of a novel ST. An additional *nrdA* typing method [28] was performed on bordetellae which failed MLST characterization [21]. This method was also applied to the other *Bordetella* isolates and sequences of all *nrdA* loci were extracted from the database, aligned and phylogenetically analyzed using MEGA 5.2.

Analysis of evolutionary relationships

The evolutionary relationships among bordetellae in this study, were analyzed and presented as a minimum spanning tree using the algorithm, PHYLOViZ [9]. The displayed minimum spanning tree was used to visualize the possible evolutionary relationships between *Bordetella* isolates based on their MLST allelic profile data. The minimum spanning tree firstly, links STs sharing single locus variants and the subsequent branches connect to STs with more than one locus variation. STs located multiple branches away from the core ST have the most loci variants.

Amplification of the CRISPR-cas locus

A Type II-C CRISPR-Cas system was recently discovered in a newly described bordetellae, *B. pseudohinzii* [15]. CRISPR-Cas systems have not been identified in any other *Bordetella* species, making the amplification of genes associated to the Type II-C CRISPR-Cas system a defining feature for the identification of *B. pseudohinzii*. Nucleic acid amplification of the Type II-C CRISPR associated genes was performed on bordetellae which could not be characterized using MLST [15].

RESULTS

A total of four Malaysian states were included in this study (Johor, Kedah, Kelantan and Terengganu). Two types of rodent species were caught from the four states; *Rattus norvegicus* (n=177) and *Rattus rattus diardi* (n=100), with Kelantan having the most number of captured rats (n=143), followed by Terengganu (n=58), Kedah (n=57) and Johor (n=19). *Rattus norvegicus* dominated the number of captured rats in Kelantan (n=99), followed by Terengganu (n=38), Johor (n=19) and Kedah (n=21). The most number of *Rattus rattus diardi* was captured in Kelantan (n=44), followed by Kedah (n=36) and Terengganu (n=20). There was no *Rattus rattus diardi* trapped in Johor (Fig. 1). Several shrews (n=7) were trapped in Kelantan, however, they were omitted as this species was not included in the animal ethics application.

Analyses of the bacterial isolates cultured from rat lung homogenates using 16S rDNA sequencing revealed the isolation of 4 *Bordetella* isolates, yielding the prevalence of bordetellae at 1.44%. Based on the partial 16S rDNA sequences, the isolates were identified as *B. bronchiseptica* (n=3) and *B. hinzii* (n=1) (Table 1). These four bordetellae were isolated from Terengganu rats. *Rattus rattus diardi* harbored three isolates (2 *B. bronchiseptica* and 1 *B. hinzii*), while the other *B. bronchiseptica* was isolated from the lung of one *Rattus norvegicus* (Table 1). The *B. bronchiseptica* isolates (TRE151600, TRE150202 and TRE155202) were subjected to MLST and *nrdA* typing, whereas only the *nrdA* typing was performed on isolate TRE152202 (Table 1). Genotyping analyses uncovered that all of the *B. bronchiseptica* isolates as ST82 (novel ST) possessing *nrdA* locus 162 (Table 1). Isolate TRE152202 was shown to possess *nrdA* locus 189, corresponding to *Bordetella* genogroup 16 (Table 1). Upon obtaining this finding, the extracted DNA of isolate TRE152202 was used for the amplification and sequencing of the CRISPR-cas locus. Amplicons of the *cas1*, *cas2* and *cas9* were amplified (Fig. 2) and all displayed identical sequences as the CRISPR-associated genes of *B. pseudohinzii* HI4681 (accession no. CP016440) and BH370 [19]. Isolate TRE152202 was thus, redesignated as *B. pseudohinzii*.

The population structure among *B. bronchiseptica* isolated in this study was examined by building a minimum spanning tree

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using an established algorithm, PHYLOViZ. The ST82 isolates were shown to have evolved from the central ST6-lineage (Fig. 3). As depicted in Fig. 3, all of the STs were also members of the <em>B. bronchiseptica</em> complex I, which consists of predominantly animal-associated isolates. Based on the constructed phylogenetic tree using the trimmed 765-bp <em>nrdA</em> gene, we observed that the classical and non-classical bordetellae were distinctively separated into 3 clusters (Fig. 4); one belonging to the classical bordetellae, one belonging to non-classical bordetellae, of which has been partially characterized and one belonging to the recently discovered non-classical bordetellae (<em>B. bronchialis</em>, <em>B. flabilis</em> and <em>B. sputigena</em>). Two sub-clusters were noted among the classical bordetellae; one containing predominantly human isolates (<em>B. bronchiseptica</em> complex IV and <em>B. pertussis</em> complex II) and the other consists of predominantly animal isolates (<em>B. bronchiseptica</em> complex I and <em>B. parapertussis</em> complex III) (Fig. 4). The <em>B. bronchiseptica</em> isolates carrying <em>nrdA</em> locus 162 were clustered together with <em>B. bronchiseptica</em> complex I and <em>B. parapertussis</em> complex III. The <em>B. pseudohinzii</em> isolate carrying <em>nrdA</em> locus 189, however was clustered together with the non-classical bordetellae (Fig. 4).

Antimicrobial susceptibility testing performed on the <em>Bordetella</em> isolates showed that they had almost similar antimicrobial profiles. Resistance was displayed towards amoxicillin-clavulanic acid, ampicillin, ceftriaxone and cefotaxime. All of the <em>Bordetella</em> isolates were susceptible to amikacin, ceftazidime, ciprofloxacin, gentamicin, imipenem and meropenem (Table 1). The only observed antimicrobial disparity was on erythromycin, by which the <em>B. bronchiseptica</em> isolates displayed resistance while <em>B. pseudohinzii</em> was susceptible (Table 1). Additionally, the <em>B. bronchiseptica</em> isolates carried the virulence genes; <em>dnt</em>, <em>bfrZ</em> and <em>fla</em>, while <em>B. pseudohinzii</em> did not carry any (Table 1).

### Table 1. Antimicrobial susceptibilities, sequence types, <em>nrdA</em> loci and host diversity of bordetellae recovered from wild rodents

| Isolate name (Bordetella species) | Host | Virulence gene | ST | nrdA locus | MIC (µg/ml) |
|----------------------------------|------|----------------|----|------------|-------------|
| TRE151600 (<em>B. bronchiseptica</em>) | RR | dnt, bfrZ, fla | 82 | 162 | S (16) R (256) R (128) S (0.5) S (0.5) R (8) R (16) R (8) S (1) S (0.06) S (0.03) |
| TRE150202 (<em>B. bronchiseptica</em>) | RR | dnt, bfrZ, fla | 82 | 162 | S (4) R (128) R (64) S (0.06) S (0.25) R (16) R (16) R (8) S (2) S (0.5) S (0.5) |
| TRE155202 (<em>B. bronchiseptica</em>) | RN | dnt, bfrZ, fla | 82 | 162 | S (8) R (256) R (64) S (0.5) S (0.12) R (16) R (32) R (8) S (1) S (0.25) S (0.25) |
| TRE152202 (<em>B. pseudohinzii</em>) | RR na | na | 82 | 189 | S (8) R (128) S (0.03) R (8) R (16) S (0.5) S (0.5) S (0.06) S (0.06) |

RR, <em>Rattus rattus diardii</>; RN, <em>Rattus norvegicus</>; dnt, dermonecrotic toxin; bfrZ, exogenous ferric siderophore receptor; fla, flagella; ST, sequence type; na, not available; MIC, minimum inhibitory concentration; AMK, amikacin; AMC, amoxicillin-clavulanic acid; AMP, ampicillin; CAZ, ceftazidime; CIP, ciprofloxacin; CRO, ceftriaxone; CTX, cefotaxime; ERY, erythromycin; GEN, gentamicin; IMP, imipenem; MEM, meropenem; R, resistant; S, sensitive.

Fig. 2. Nucleic acid amplification products of <em>cas9</em>, <em>cas1</em> and <em>cas2</em> of <em>B. pseudohinzii</em> TRE152202. Lane 1, <em>B. pseudohinzii</em> BH370 (positive control); Lane 2, <em>B. pseudohinzii</em> TRE152202; Lane 3, <em>B. bronchiseptica</em> TRE151600; Lane 4, <em>B. bronchiseptica</em> TRE150202; Lane 5, <em>B. bronchiseptica</em> TRE155202.
Even though rats were implicated as the host for *B. bronchiseptica* [2], the prevalence of this species in the Malaysian wild rat population is unknown. The overall prevalence of bordetellae in rodents from the four Malaysian states was 1.44% (4/277), slightly higher as compared to the findings by Jiyipong *et al.* (2013) where only one bordetellae was cultured out of 206 rodent specimens yielding a prevalence of 0.49% [17]. This was also consistent with the reported low prevalence of *B. bronchiseptica* carriers in rodent populations [2] and a well-fed population of rats in the market would have developed stronger immunological defenses, reducing *B. bronchiseptica* infection risk [8]. The isolation of bordetellae from the lung tissues of *Rattus norvegicus* and *Rattus rattus diardii* suggests that these two rodent species were competent hosts and indicated the propensity of this bacterial genus to the respiratory tract, similar to other reports [4, 30]. Only rats from the state of Terengganu harbored bordetellae and this was to be expected due to the low prevalence of this bacterium in the host [2] coupled with the limited number of study sites. The absence of *Rattus rattus diardii* in Johor probably reflected the sampling limitation in this study whereby trappings were conducted for only one night and rat species with low abundance was not captured. Identification of bordetellae using a three-pronged approach (16S rDNA sequencing, *nrda* typing and MLST) ensures resolution to the species level, at least for the classical bordetellae [21], owing to the close phylogenetic relationships within the genus [28]. It has been reported that rodents are unnatural hosts for *B. bronchiseptica* although they can be colonized and shed the bacteria [30]. Rats have been shown to demonstrate strong resistance towards clinical infection by *B. bronchiseptica*, however they become asymptomatic carrier for life after exposure [2], similar to our observation of the seemingly healthy captured rats. *B. bronchiseptica* can remain within the respiratory epithelial cells to evade the host’s immune system [10] contributing to life-long disease persistence in infected rats.

The *B. bronchiseptica* isolates recovered from *Rattus norvegicus* and *Rattus rattus diardii* (Table 1) displayed identical genotype (ST82; *nrda* locus 162), suggesting inter-species *B. bronchiseptica* transmission between rats in that particular area. This contradicts the earlier suggestion that *B. bronchiseptica* transmission does not occur between wild type host [30], although colonization of hosts can be affected by the different physiological responses between mice and rats [20] and/or *B. bronchiseptica* strain variations [4]. Alternatively, *B. bronchiseptica* recovered from rats in this study could be transmitted by other animals in the market, as the bacterium is highly contagious among poultry and livestock [30]. This species can also survive in the environment and on inanimate surfaces [2, 30], possibly aiding the spread and transfer to other mammals. *nrda* typing identified isolate TRE152202 as *Bordetella* genogroup 16 possessing *nrda* locus 189 after the inability of 16S rDNA sequencing and MLST to resolve its species identity (Table 1). We have previously isolated a similar bordetellae with *nrda* locus 189 [21] and this isolate was recently classified as *B. pseudohinzii* [19]. This species can be differentiated from the other bordetellae by the amplification of the Type II-C CRISPR-Cas system (Fig. 2), exclusively found in *B. pseudohinzii* [15]. Initially found only in laboratory mice.
we have now established that *B. pseudohinzii* also reside in the lung of wild rats, although the clinical significance remains unclear.

The *nrdA*-based phylogeny [28] (Fig. 4) displayed a congruent clustering of *Bordetella* complexes with the MLST-based phylogeny. The first cluster comprised *B. bronchiseptica* complex IV and *B. pertussis* complex II, and the second cluster comprised *B. bronchiseptica* complex I and *B. parapertussis* complex III. The bordetellae population structure inferred from *nrdA* typing concurred with the MLST-based phylogeny, affirming that *B. parapertussis* and *B. pertussis* evolved from *B. bronchiseptica* complex I and IV, respectively [6]. In addition, the non-classical bordetellae formed a separate cluster, demonstrating excellent *Bordetella* subspecies demarcation using the *nrdA*-based phylogeny [28] (Fig. 4).

Based on the MLST *Bordetella* phylogeny (Fig. 3), the novel ST82 clustered together with other clones from complex I [6], which included isolates from humans and animals. A high possibility of zoonotic transmission can be anticipated with ST82 isolates as they are closely related to other STs (ST4, ST7, ST11, ST12, ST23, ST27 and ST55) (Fig. 3) which were recovered from humans [4, 6, 27]. Transmission from infected animals to humans can occur through inhalation of aerosols or by contact with respiratory secretions [10]. Animals may not show apparent clinical symptoms of *B. bronchiseptica* infection however, human-adapted strains may modulate their virulence differently and lead to respiratory illnesses [4], augmenting the risk to humans. ST27
from a household cat, for example, was strongly suspected to be the source of B. bronchiseptica infection in a child [27] and a B. bronchiseptica strain originating from companion animals was suggested to infect and, transmit between humans who were in close contact with each other [14]. Human to human transmission, hence, is possible via contact with respiratory droplets [10].

Antimicrobial susceptibility testing revealed that B. bronchiseptica and B. pseudohinzii had almost similar antimicrobial profiles, reflecting the close phylogenetic relationships between the two bordetellae. The only antimicrobial susceptibility variance between B. bronchiseptica and B. pseudohinzii was observed using the macrolide; erythromycin, where it was ineffective against B. bronchiseptica although it was reported to be effective for the treatment of most Bordetella clinical infections [31], including B. pseudohinzii in this study. Resistance to erythromycin could be an innate phenotype of B. bronchiseptica as this was also observed among B. bronchiseptica isolates from pigs with respiratory diseases in China [33]. The anti-pseudomonal cephalosporin; ceftazidime, was effective for the eradication of B. bronchiseptica and B. pseudohinzii, similar to other reports [21, 32]. The isolates were however, susceptible to the third generation cephalosporins, cefotaxime and ceftriaxone, in agreement with a previous case of suspected B. bronchiseptica transmission from a kitten [26] and B. pseudohinzii BH370 recovered from ICR mouse [21]. Both bordetellae appeared to be susceptible to aminoglycosides (amikacin and gentamicin), carbapenem (imipenem and meropenem) and fluoroquinolone (ciprofloxacin), comparable to B. bronchiseptica human infections [11, 14, 32] and B. pseudohinzii BH370 [21]. In accordance with a previous study on human clinical isolates [32], our B. bronchiseptica isolates displayed identical resistance towards ampicillin and amoxicillin-clavulanic acid. B. pseudohinzii BH370 [21], 8-296-03 16 [16] and TRE152202 (this study) exhibited analogous resistance to ampicillin and even though 8-296-03 3 was not tested against amoxicillin-clavulanic acid [16], it was expected to display resistance similar to BH370 and TRE152202, suggesting that the difference in hosts did not influence antimicrobial properties. The nucleic acid amplification of dnaT, bfrZ and fla from B. bronchiseptica but not B. pseudohinzii may reflect the roles of these genes in host colonization [29], catering to the broad host range for B. bronchiseptica.

Considered together, the much overlooked bordetellae surveillance despite the low prevalence among wild rats in Malaysia is a cause for concern as bordetellae such as B. bronchiseptica can be highly contagious in nature [30], persistent [12] and has a wide host range [2]. The persistence of this species in the host predisposes it to progressively acquire antimicrobial resistance [11] that could exacerbate pre-existing clinical conditions [27]. Upon browsing the Bordetella MLST database at http://pubmlst.org/bordetella/, we noticed that the novel B. bronchiseptica ST82 was the sole complex I representative from Asia, reflecting the neglected state of research on this species. The significant finding of B. pseudohinzii in wild rat is important as this species was previously thought to specifically infect laboratory mice [16]. Although the clinical significance remains unknown and prevalence is low, the potential for pathogen transmission from the wild to the laboratory is possible and this could have serious implications for studies using these infected animals. It is therefore important to study the genetic composition and prevalence of bordetellae in urban rats for comparison with other geographical regions to ensure early identification of pathogens capable of causing public health disorder.

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