**Abstract**

Members of the *Mycoplasma mycoides* cluster represent important livestock pathogens worldwide. *Mycoplasma mycoides* subsp. *mycoides* is the etiologic agent of contagious bovine pleuropneumonia (CBPP), which is still endemic in many parts of Africa. We report the genome sequences and annotation of two frequently used challenge strains of *Mycoplasma mycoides* subsp. *mycoides*, Afadé and B237. The information provided will enable downstream ‘omics’ applications such as proteomics, transcriptomics and reverse vaccinology approaches. Despite the absence of *Mycoplasma pneumoniae* like cyto-adhesion encoding genes, the two strains showed the presence of protrusions. This phenotype is likely encoded by another set of genes.

**Keywords:** *Mycoplasma mycoides* subsp. *mycoides*, Challenge strain, Genome, Contagious bovine pleuropneumonia, Protrusion

**Introduction**

The ‘*Mycoplasma mycoides* cluster’ comprises five species/subspecies, *Mycoplasma mycoides* subsp. *mycoides*, *Mycoplasma leachii*, *Mycoplasma mycoides* subsp. *capri*, *Mycoplasma capricolum* subsp. *capripneumoniae* and *Mycoplasma capricolum* subsp. *capricolum* [1, 2]. Among them, *Mycoplasma mycoides* subsp. *mycoides*, the causative agent of contagious bovine pleuropneumonia (CBPP), is an economically very important bacterial bovine pathogen in sub-Saharan Africa. CBPP was first described in Europe already in 1773 [3], and the causative *Mycoplasma* was then cultivated and characterized in 1898 in Europe [4]. It has been shown that it spread from Europe to North America, Africa, Australia and Asia via livestock movements. Currently the disease is endemic and widespread in sub-Saharan Africa, ranging from western, central to eastern Africa. In Europe the last outbreaks were reported in Spain, Italy, Portugal and France in the 1980s and 1990s [5]. In comparison to other members of the ‘*Mycoplasma mycoides* cluster’, with the exception of *Mycoplasma capricolum* subsp. *capripneumoniae*, *Mycoplasma mycoides* subsp. *mycoides* shows limited sequence diversity, probably due to its recent emergence about 300 years ago [5, 6].

Currently the complete genomes of only three *Mycoplasma mycoides* subsp. *mycoides* strains have been deposited in GenBank, the type strain PG1 [7], which is often used in laboratories but which is considered to be avirulent, the Australian outbreak strain Gladysdale [8] and a European outbreak strain 57/13 [9]. PG1 has been shown to differ genetically and phenotypically from field stains of *Mycoplasma mycoides* subsp. *mycoides*, showing attenuated cytotoxicity and reduced adhesion to bovine epithelial cells [5, 10, 11], most likely because of the multiple *in vitro* passages this strain underwent before being deposited in the strain collections. In particular strain PG1 contains 2 large 24 kb repeats while 27 field strains isolated from three different continents only contain one [11]. Strain Gladysdale was isolated from Australia around 1953 [12]. Strain 57/13 was isolated in Italy in 1992. Neither of these three strains, therefore, represent virulent African strains. The genetic diversity of *Mycoplasma mycoides* subsp. *mycoides* strains has been reported to be highest in Africa.
[5] where the disease is present in many countries of sub-Saharan Africa [13]. We sequenced and annotated the genomes of two virulent African strains Afadé and B237, which are frequently used as challenge strains in animal experiments [14–18]. The strains have been re-isolated directly from experimentally infected animals and have not been exposed to subsequent passaging beyond filter-cloning to promote uniformity before genomic DNA was isolated for sequencing. The genomic sequence information from this work will contribute to comparative genomic analyses and therefore the characterization of the core and pan genome of the ‘Mycoplasma mycoides cluster’ and Mycoplasma mycoides subsp. mycoides in particular. The genomic information will also be useful for downstream ‘omics’ applications, such as proteomics, transcriptomics and reverse vaccinology approaches.

**Organism information**

**Classification and features**

*Mycoplasma mycoides* subsp. *mycoides* is an obligate parasite, which resides in the respiratory tract of animals. It is a non-motile, non-sporulating bacterium. It lacks a cell wall and has a pleomorphic shape. Transmission electron microscopy images were generated for both Afadé and B237 strains (Fig. 1). Cell pellets were fixed in 150 mM HEPES, pH 7.35, containing 1.5 % formaldehyde and 1.5 % glutaraldehyde for 30 min at RT and at 4 °C overnight. After dehydration in acetone and embedding in EPON, ultrathin sections of 40 nm were mounted on formvar-coated coppergrids, poststained with uranyl acetate and lead citrate [19] and observed in a Morgagni TEM (FEI). Images were taken with a side mounted Veleta CCD camera.

Interestingly the transmission electron microscopy revealed protrusions resembling the attachment organelle observed in *Mycoplasma pneumoniae* [20–23]. The physiological function of these protrusions and branching phenotype needs to be defined in future studies. The general features of *Mycoplasma mycoides* subsp. *mycoides* strains Afadé and B237 are presented in Table 1 and Appendix: Table 6.

We previously confirmed that both strains Afadé and B237 are *Mycoplasma mycoides* subsp. *mycoides* using phenotypic growth characteristics, species-specific PCR and a Multi-Locus Sequence Typing (MLST) method [5, 6]. *Mycoplasma mycoides* subsp. *mycoides* strain Afadé originates from Northern Cameroon and was isolated at the Farcha laboratories in Tchad in 1966 [24]. It has since served for several experimental infections [14–18]. The filter-cloned strains used for this sequence analysis were re-isolated from experimentally infected cattle [14, 17] that showed severe clinical signs and pathomorphologic lesions typical of CBPP. *Mycoplasma mycoides* subsp. *mycoides* strain B237 was originally isolated in 1997 in Thika, Kenya, by the Kenya Agricultural Research Institute (KARI).

Figure 2 shows a phylogenetic tree of the 16S rRNA sequences. 16S rRNA gene sequences from *Mycoplasma mycoides* subsp. *mycoides* strains Gladysdale, 57/13 and PG1, *Mycoplasma mycoides* capri strains 95010 and GM12, *Mycoplasma capricolum* subsp. capricolum strain ATCC27343, *Mycoplasma capricolum* subsp. capripneumoniae strain M1601, *Mycoplasma leachii* strains 99/014/6 and PG50, *Mycoplasma feriruminatoris* strain G5847 (Accession numbers: CP002107, CP010267, NC_005364, NC_015431, NZ_CP001668, NC_007633, CM001150, NC_017521, ANFU0100033, NC_014751, respectively) were retrieved from GenBank. All *Mycoplasma* genome sequences retrieved from GenBank have two copies of 16S rRNA each, with the exception of *Mycoplasma feriruminatoris*, where two copies are present but are not resolved in the draft genome [25].

**Genome sequencing information**

**Genome project history**

The sequencing and quality assurance was performed at Lausanne Genomic Technologies Facility, Center for Integrative Genomics, University of Lausanne, Switzerland. The assemblies and finishing were done at the Institute for Genome Sciences and International Livestock Research Institute. Functional annotation was produced by the Institute for Genome Sciences Analysis Engine [26] (http://www.igs.umd.edu/research/bioinformatics/analysis/index.php). Table 2 presents the project information and its association with MIGS version 2.0 compliance [27].

**Growth conditions and genomic DNA preparation**

Both strains were grown in PPLO medium (Difco, Cat no. 255420) supplemented with 20 % heat-inactivated horse serum (Sigma, Cat. No. H1138), 0.5 % glucose, 0.03 % penicillin G, 20 mg/ml thallium acetate and 0.9 g/L yeast extract at 37 °C.

Liquid cultures of *Mycoplasma* were filter cloned using a 0.22 μm filter to disrupt possible cell aggregates. A serial dilution (1/10 - 1/10,000,000,000) was made immediately and 50 μl was plated on PPLO agar.

After 3–4 days of incubation at 37 °C, a single colony was picked and was used to inoculate 4 ml of PPLO medium which was aliquoted and stored at −80 °C.

Filter cloned *Mycoplasma* were grown overnight in 100 ml PPLO medium at 37 °C. Before entering the stationary growth phase the culture was centrifuged at 2,862 g for 1 h, and the pellet was resuspended in 2.5 ml of TNE buffer (0.01 M Tris–HCl, pH 8.0; 0.01 M NaCl; 0.01 M EDTA). Subsequently 50 μl SDS (10 %) and 50 μl Proteinase K (20 mg/ml) were added and the tubes were incubated at 37 °C for 2 h. After addition of 26 μl of 100 mM PMSF the tubes were incubated 15 min at room temperature, 25 μl of RNase A (10 mg/ml) was added, followed by incubation at 37 °C for 1 hr. Sodium acetate and Phenol Saturated Buffer...
was added (25 μl of NaOAc 1.5 M pH 5.2, and 2250 μl of Phenol), the solution was mixed by vortexing and centrifuged at 15,870 g for 10 min. The top phase was transferred to a new tube and mixed with Phenol:Chloroform:Isoamyl Alcohol Buffer (Phenol:Chloroform:Isoamyl Alcohol; 25:24:1) followed by another centrifugation at 15,870 g for 10 min and again the top phase was transferred to a new tube. Finally, the DNA was precipitated with isopropanol, washed with 70 % ethanol, dried and resuspended in 200 μl of 2 mM Tris, 0.2 mM EDTA.

**Genome sequencing and assembly**

The genome sequence of *Mycoplasma mycoides* subsp. *mycoides* strain Afadé was generated using a combination
of Pacific Biosciences R.S. (PacBio) sequencing (65,280 reads/2853 bp average read length) and Illumina MiSeq sequencing (7,078,010 reads/295 average read length) down-sampled to cover 50 times the expected genome size. The sequencing errors of the long PacBio single-molecule reads were corrected with the shorter, high accuracy Illumina reads using the Celera Assembler (CA) pacbio correction module PBcR (version 7.0, [28]). The resulting corrected PacBio reads were randomly sampled to 25 genome fold and assembled using CA (version 7.0, [29]) and yielded 18 contigs with a total size of 1,278,455 bp. Eight contigs comprised the draft genome of strain Afadé.

The whole genome sequence of Mycoplasma mycoides subsp. mycoides strain B237 was obtained using PacBio sequencing (59,775 reads/2674 average read length). PacBio reads were corrected with PBcR self-correction module. Corrected reads randomly sampled to 25 genome fold were assembled with CA and yielded 2 contigs with total size of 1,208,895 bp. One long contig comprises the entire genome and contained the other contig (5091 bp) in a repeat region. The final genome sequences had a 24-fold coverage for Afadé and 23-fold coverage for B237.

The contigs of both assemblies were aligned against the two Mycoplasma mycoides subsp. mycoides reference genomes of Gladysdale [8] and PG1 [7] available in Genbank (CP002107, NC_005364) using mummer [30] and we noticed that all small contigs (<15,000 bp) aligned to places

| Property       | Term                                      | Evidence code* |
|----------------|-------------------------------------------|----------------|
| Classification| Domain Bacteria                           | TAS [39]       |
|                | Phylum Firmicutes                         | TAS [40]       |
|                | Class Tenericutes                         | TAS [41–44]    |
|                | Order Mycoplasmatales                     | TAS [45, 46]   |
|                | Family Mycoplasmatace                      | TAS [46]       |
|                | Genus Mycoplasma                          | IDA            |
|                | Species Mycoplasma mycoides               | IDA [4]        |
|                | Subspecies Mycoplasma mycoides subsp. mycoides | IDA [4]      |
|                | Strains Afadé and B237                    |                |
| Cell shape     | Pleomorph                                 | IDA            |
| Motility       | Nonmotile                                 | IDA            |
| Sporulation    | Nonspore-forming                          | IDA            |
| Temperature range | 30–42 °C                                | IDA [42]       |
| Optimum temperature | 38.5 °C                                | IDA            |
| pH range, optimum | 6.5 – 8.5; 7.5                           | IDA            |
| Carbon Source  | Not determined since strains require complex media including serum for growth | -              |
| Energy Source  | Not determined since strains require complex media including serum for growth | -              |
| MIGS-6 Habitat | Respiratory tract                         | IDA            |
| MIGS-6.3 Salinity | 0.09 %, no growth was obtained at salinities ≥0.5 M NaCl | IDA [43]      |
| MIGS-22 Oxygen Requirement | Facultative anaerobe                      | [42]           |
| MIGS-15 Biotic relationship | Pathogen                                 | -              |
| MIGS-14 Pathogenicity | Etiological agent of Contagious Bovine Pleuropneumonia (CBPP) | -              |
| MIGS-4 Geographic location | Cameroon (Afadé), Kenya (B237)   | [3]            |
| MIGS-4.1 Sample collection time | 1965 (Afadé), 1997 (B237) | -              |
| MIGS-4.2 Longitude | N/A (Afadé) 37°05′E (B237) |              |
| MIGS-4.3 Depth | N/A                                       |                |
| MIGS-4.4 Altitude | N/A (Afadé), 1631 m (B237) |              |

*Evidence codes - IDA: Inferred from Direct Assay; TAS: Traceable Author Statement (i.e., a direct report exists in the literature); NAS: Non-traceable Author Statement (i.e., not directly observed for the living, isolated sample, but based on a generally accepted property for the species, or anecdotal evidence). These evidence codes are from the Gene Ontology project [47]
already covered in other bigger contigs. On closer inspection, most of these contigs aligned to a previously characterized 26 kb region [11], consisting of a tandem repeat of three 8 kb segments, interspersed with transposon elements. Due to its repetitive nature, this 26 kb region was not clearly resolved during the assembly process. In order to resolve part of it, we were able to design unique primer pairs and amplify two long-range PCRs fragments of 4,800 and 5,200 bp respectively. For each genome, both Sanger derived sequences were aligned to the assembled genomes.

| Table 2 | Project information |
|---------|---------------------|
| **MIGS ID** | **Property** | **Term** | **Property** | **Term** |
| MIGS-31 | Finishing quality | High-quality draft | Illumina MiSeq, Pacific Biosciences R.S. | Illumina MiSeq, Pacific Biosciences R.S. |
| MIGS-28 | Libraries used | 1. Illumina Paired End 7,078,010 reads; Average read length 295 bp; Average insert size 725 bp. 2. PacBio 65,280 reads, 2853 bp average read length; | 1. PacBio 59,775 reads; Average read length 2674 bp |
| MIGS-29 | Sequencing platforms | Illumina MiSeq, Pacific Biosciences R.S. | Celeria Assembler v.7 |
| MIGS-31.2 | Fold coverage | 24X | 23X |
| MIGS-30 | Assemblers | Celeria Assembler v.7 | Celeria Assembler v.7 |
| MIGS-31.2 | Gene calling method | Prodigal | Prodigal |
| Genbank ID | LAEX00000000 | LAEW000000000 |
| Date of Release | 20-Mar-15 | 20-Mar-15 |
| BIOPROJECT | PRJNA272775 | PRJNA272775 |
| MIGS 13 | Source Material Identifier | ILRI_Aziz Biobank Strain Afadé | ILRI_Aziz Biobank Strain B237 |
| Project relevance | Challenge strains of CBPP | Challenge strains of CBPP |
before and after polishing with multiple iterations of the PacBio Quiver algorithm (version 0.9.0 [31]). We verified that in the regions covered by the Sanger sequences, all substitution mismatches were resolved by Quiver, however we manually fixed a few indels present in the post polishing alignment, which were not corrected by Quiver.

**Genome annotation**

Open reading frames (ORFs) were predicted using Prodigal 2.50 [32]. Functional annotation was produced by the Institute for Genome Sciences Analysis Engine [26].

We annotated the small contigs overlapping bigger ones described above separately and noticed that these contigs had more ambiguous characters and ORFs that were on average half the size of the corresponding ORFs in larger contigs (498 nt versus 920 nt). This was due to insertions and deletions. We therefore excluded the small contigs from the assemblies and report 1 contig for *Mycoplasma mycoides* subsp. *mycoides* strain B237 and 8 contigs for *Mycoplasma mycoides* subsp. *mycoides* strain Afadé.

We also reannotated the genomes of *Mycoplasma mycoides* subsp. *mycoides* strain PG1, *Mycoplasma mycoides* subsp. *mycoides* strain Gladysdale and *Mycoplasma mycoides* subsp. *mycoides* strain 57/13 using the same Engine, for ease of comparison.

**Genome properties**

The genomes of *Mycoplasma mycoides* subsp. *mycoides* strain Afadé and B237 have a total size of 1,190,241 bp and 1,203,804 bp, respectively. The GC-content of both genomes is 23.9 %. Both strains have two copies of the 12 kb and 13 kb repeat described in [11], the difference in size between the two genomes is therefore not due to a missing copy in Afadé.

A total of 1,124 ORFs as well as 30 tRNA and 2 copies of the 23S, 16S and 5S rRNA operons were predicted. The average gene length is 920 bp and 927 bp for Afadé and B237, respectively. The coding density of the genome is 86.7 %. Signal peptides were detected using pSortb v3.0 [33] and LipoP v1.0 [34]. Transmembrane helices were detected with the TMHMM server v2.0 [35, 36]. CRISPR repeats were searched with the CRISPR Finding program online. The properties and the statistics of both genomes are summarized in Tables 3, 4, 5.

**Insights from the genome sequence**

The genomes of the two African strains *Mycoplasma mycoides* subsp. *mycoides* Afadé and B237 were compared to the three previously sequenced *Mycoplasma mycoides* subsp. *mycoides* strains Gladysdale, PG1 and 57/13 using CloVR and Sybil [37, 38]. Figure 3 shows a synteny gradient of the aligned genomes. Although there is a high number of transposable elements in all genomes, no major rearrangements have been observed. These results fit well with the very recent emergence of the pathogen, estimated to be as young as 300 years, and the narrow host specificity of *Mycoplasma mycoides* subsp. *mycoides* [5].

The core genome length is 1,148,950 bp. A total of 773 SNPs were identified when comparing the five core genomes. Only 72 SNPs distinguish B237 from Afadé. Two hundred and sixty six SNPs separate the Australian and European strains Gladysdale and 57/13. PG1 is the most distant from the other four genomes with 399, 483, 465 to 425 SNPs when compared to Afadé, Gladysdale, 57/13 and B237, respectively. This confirms previous reports [5].

We looked for homologs to the Cytadhesin proteins P1, P30, P40. P65, P90, HMW1 and HMW3 from *Mycoplasma pneumoniae* in the Afadé and B237 proteomes using blastp. No significant hits were found for any of the proteins. Other proteins might be involved in the adhesion process and will need to be identified and characterized.

| Table 3 Summary of the B237 and Afadé genomes: one circular chromosome |
|---------------------------------------------------------------|
| **Strain** | **Size (Mb)** | **Topology** | **INSDC identifier** |
|-------------|---------------|--------------|----------------------|
| Afadé       | 1,190,241     | 8 contigs    | LAEX000000000000     |
| B237        | 1,203,804     | Circular     | LAEW000000000000     |

| Table 4 Nucleotide content and gene count levels of the genome |
|---------------------------------------------------------------|
| **Attribute** | **Afadé** | % of total | **B237** | % of total |
|----------------|-----------|------------|-----------|------------|
| Genome Size (bp) | 1,190,241 | 100.00     | 1,203,804 | 100.00     |
| DNA coding (bp)  | 1,032,189 | 86.70      | 1,043,698 | 86.70      |
| DNA G + C (bp)   | 284,536   | 23.90      | 287,709   | 23.90      |
| DNA scaffolds    | na        | na         | na        | na         |
| Total genes      | 1,156     | 100.00     | 1,157     | 100.00     |
| Protein-coding genes | 1120   | 96.89      | 1121      | 96.89      |
| rRNA genes       | 6         | 5.19       | 6         | 5.19       |
| Pseudogenes      | 0         | 0.00       | 0         | 0.00       |
| Genes in internal clusters | na | na         | na        | na         |
| Genes with function prediction | 687 | 59.43      | 693       | 59.90      |
| Genes assigned to COGs | 681 | 58.71      | 693       | 59.90      |
| Genes with Pfam domains | 389 | 33.65      | 355       | 30.68      |
| Genes with signal peptides | 74 | 6.40       | 74        | 6.40       |
| Genes with transmembrane helices | 234 | 20.24      | 241       | 20.83      |
| CRISPR repeats   | 0.00      | 0.00       | 0.00      | 0.00       |

*The total is based on either the size of the genome in base pairs or the total number of protein coding genes in the annotated genome*
Conclusions
The genomes of the two African strains as expected differ from the laboratory type strain PG1, the European outbreak strain 57/13 and the Australian outbreak strain Gladysdale. Therefore these genome sequences should be included in subsequent genome comparisons and ‘omics’ studies. The presence of protrusions and branching phenotypes in these two Mycoplasmas but the absence of protein encoding genes similar to the ones characterized in *Mycoplasma pneumoniae* indicates that other/novel proteins in the *Mycoplasma* genomes encode the development of protrusions and branching.

### Table 5 Number of genes associated with the 25 general COG functional categories

| Code | Value Strain Afadé | % of totala | Value Strain B237 | % of totala | Description |
|------|-------------------|-------------|-------------------|-------------|-------------|
| J    | 141               | 12.19       | 139               | 12.01       | Translation, ribosomal structure and biogenesis |
| A    | 0                 | 0.00        | 0                 | 0.00        | RNA processing and modification |
| K    | 34                | 2.94        | 34                | 2.94        | Transcription |
| L    | 50                | 4.32        | 50                | 4.32        | Replication, recombination and repair |
| B    | 0                 | 0.00        | 0                 | 0.00        | Chromatin structure and dynamics |
| D    | 9                 | 0.78        | 8                 | 0.69        | Cell cycle control, Cell division, chromosome partitioning |
| Y    | 0                 | 0.00        | 0                 | 0.00        | Nuclear structure |
| V    | 12                | 1.04        | 13                | 1.12        | Defense mechanisms |
| T    | 15                | 1.30        | 15                | 1.30        | Signal transduction mechanisms |
| M    | 27                | 2.34        | 33                | 2.85        | Cell wall/membrane biogenesis |
| N    | 8                 | 0.69        | 9                 | 0.78        | Cell motility |
| Z    | 0                 | 0.00        | 0                 | 0.00        | Cytoskeleton |
| W    | 0                 | 0.00        | 0                 | 0.00        | Extracellular structures |
| U    | 5                 | 0.43        | 6                 | 0.52        | Intracellular trafficking and secretion |
| O    | 26                | 2.25        | 25                | 2.16        | Posttranslational modification, protein turnover, chaperones |
| C    | 29                | 2.51        | 28                | 2.42        | Energy production and conversion |
| G    | 71                | 6.14        | 70                | 6.05        | Carbohydrate transport and metabolism |
| E    | 44                | 3.81        | 42                | 3.63        | Amino acid transport and metabolism |
| F    | 32                | 2.77        | 32                | 2.77        | Nucleotide transport and metabolism |
| H    | 30                | 2.60        | 29                | 2.51        | Coenzyme transport and metabolism |
| I    | 14                | 1.21        | 14                | 1.21        | Lipid transport and metabolism |
| P    | 39                | 3.37        | 48                | 4.15        | Inorganic ion transport and metabolism |
| Q    | 1                 | 0.09        | 1                 | 0.09        | Secondary metabolites biosynthesis, transport and catabolism |
| R    | 45                | 3.89        | 45                | 3.89        | General function prediction only |
| S    | 6                 | 0.52        | 6                 | 0.52        | Function unknown |
| -    | 101               | 8.74        | 105               | 9.08        | Other COG categories |
| -    | 442               | 38.24       | 431               | 37.25       | Not in COGs |

*a The total is based on the total number of protein coding genes in the annotated genome

**Fig. 3** (quarter page, two columns): Synteny gradient display for the four available *Mycoplasma mycoides* subsp. *mycoides* genomes, using PG1 as a reference. A white bar in the reference denotes a region with no gene annotation. The matching genes are colored based on the relative position in their respective genomes (yellow for the beginning and blue for the end). Genes shown in black are part of a paralogous cluster in their respective genome and therefore do not have a single native location. The GC-content in % is plotted for the reference genome.
## Appendix

### Table 6 Associated MIGS record

| MIGS-ID | field name | description | description |
|---------|------------|-------------|-------------|
| Strain  |            |             |             |
| MIGS-1  | Submit to INSDC/Trace archives | LAEX00000000 | LAEW00000000 |
| 1.1     | PID        | PRJNA272471 | PRJNA272775 |
| 1.2     | Trace Archive |            |             |
| MIGS-2  | MIGS CHECK LIST TYPE |            |             |
| MIGS-3  | Project Name | High quality draft genomes of the *Mycoplasma mycoides* subsp. *mycoides* challenge strains Afadé and B237 | High quality draft genomes of the *Mycoplasma mycoides* subsp. *mycoides* challenge strains Afadé and B237 |
| MIGS-4  | Geographic Location | Cameroon | Kenya |
| 4.1     | Latitude | not reported | 01°03′S |
| 4.2     | Longitude | not reported | 37°05′E |
| 4.3     | Depth | na | na |
| 4.4     | Altitude | not reported | 1631 m |
| MIGS-5  | Time of Sample collection | not reported | not reported |
| MIGS-6  | Habitat (EnvO) | Respiratory tract | Respiratory tract |
| 6.1     | temperature | 38.5 | 38.5 |
| 6.2     | pH | 6.5–8.5 | 6.5–8.5 |
| 6.3     | salinity | 0.09 % | 0.09 % |
| 6.4     | chlorophyll | na | na |
| 6.5     | conductivity | na | na |
| 6.6     | light intensity | na | na |
| 6.7     | dissolved organic carbon (DOC) | na | na |
| 6.8     | current | na | na |
| 6.9     | atmospheric data | na | na |
| 6.10    | density | na | na |
| 6.11    | alkalinity | na | na |
| 6.12    | dissolved oxygen | na | na |
| 6.13    | particulate organic carbon (POC) | na | na |
| 6.14    | phosphate | na | na |
| 6.15    | nitrate | na | na |
| 6.16    | sulfates | na | na |
| 6.17    | sulfides | na | na |
| 6.18    | primary production | na | na |
| MIGS-7  | Subspecific genetic lineage | strain | strain |
| MIGS-9  | Number of replicons | 1 | 1 |
| MIGS-10 | Extrachromosomal elements | none | none |
| MIGS-11 | Estimated Size | 1.2 MB | 1.2 Mb |
| MIGS-12 | Reference for biomaterial or Genome report | primary genome report | primary genome report |
| MIGS-13 | Source material identifiers |            |             |
| MIGS-14 | Known Pathogenicity | Contagious Bovine Pleuropneumonia | Contagious Bovine Pleuropneumonia |
| MIGS-15 | Biotic Relationship | obligate parasite | obligate parasite |
Abbreviations

CBPP: Contagious bovine pleuropneumonia.

Competing interests

The authors declare that they have no competing interests.

Authors’ contributions

AF, ISC, HG, JW, ML, SN analyzed the data. ES, RAM, JJ, JH, JM, JF performed laboratory work. HW provided reagents. SV provided tools and protocols. AF, JJ drafted the manuscript. All authors read and approved the final manuscript.

Acknowledgments

This work was funded by the German Federal Ministry for Economic Cooperation and Development (contract 81121408, project No 09.7860.1 - 001.00). The Center of International Migration (CIM) supported Anne Fischer. Elise Schieck was supported by BMZ (grant project No.: 09.7860.1-001.00). Joerg Jores and Sanjay Vashee were supported partly by the National Science Foundation under Grant No. IOS-1110151. Infrastructure of PacBio sequencing was financed by the Fonds de la Loterie Romande. The functional annotation was conducted using the IGS Annotation Engine, University of Maryland School of Medicine. We thank Gerhard Preiss for excellent maintenance and help with electron microscopes and Andrea Kofink-Germershausen and Sabine Fiedler for excellent technical assistance. We thank Cecilia Muriuki for her help in determining the growth temperature and Herve Tettelin and Sonia Agrawal for guidance on the use of cloVR. All authors read and approved the manuscript.

Nucleotide sequence accession numbers

This Whole Genome Shotgun projects for Afadé and B237 have been deposited at DDBJ/EMBL/GenBank under accession numbers LAEX00000000, LAEW00000000 respectively. The versions described in this paper are the first versions.

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Received: 9 April 2015 Accepted: 16 September 2015

Published online: 29 October 2015

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