In vitro propagation and genome sequencing of three ‘atypical’ Ehrlichia ruminantium isolates

Three isolates of Ehrlichia ruminantium (Kümm 2, Omatjenne and Riverside), the causative agent of heartwater in domestic ruminants, were isolated in Ixodes scapularis (IDE8) tick cell cultures using the leukocyte fraction of infected sheep blood. All stocks were successfully propagated in IDE8 cells, whereas initiation attempts using endothelial cell cultures were unsuccessful. Therefore, the new technique should be included in any attempt to isolate field strains of E. ruminantium to enhance the probability of getting E. ruminantium isolates which might not be initiated in endothelial cells. Draft genome sequences of all three isolates were generated and compared with published genomes. The data confirmed previous phylogenetic studies that these three isolates are genetically very close to each other, but distinct from previously characterised E. ruminantium isolates. Genome comparisons indicated that the gene content and genomic synteny were highly conserved, with the exception of the membrane protein families. These findings expand our understanding of the genetic diversity of E. ruminantium and confirm the distinct phenotypic and genetic characteristics shared by these three isolates.

Keywords: Ehrlichia ruminantium; heartwater; in vitro isolation; tick cell line; comparative genomics.

Introduction

The intracellular rickettsial agent Ehrlichia ruminantium causes a disease commonly known as heartwater or cowdriosis. It is an infectious, non-contagious disease which affects mainly cattle, sheep, goats and some wild ruminants. It is transmitted by ticks of the genusAmblyomma and has been reported from almost all African countries south of the Sahara, from the adjacent islands of the Indian Ocean and Atlantic Ocean (Uilenberg 1983) and from some Caribbean islands (Birnie et al. 1984; Perreau et al. 1980).

The method of choice for in vitro isolation of E. ruminantium is infection of endothelial cells (Bezuidenhout, Paterson & Barnard 1985), the cell type in which organisms occur in infected animals (Cowdry 1926). However, several isolates (Allsopp et al. 2007; Du Plessis & Kümm 1971; Steyn 2009) have failed to establish in endothelial cell cultures (Bezuidenhout & Brett 1992; Bezuidenhout et al. 1988). Besides endothelial cells, tick cell lines have been used to initiate E. ruminantium in in vitro cultures and have even allowed the establishment of infection directly from the leukocytes of sheep blood (Zweygarth, Josemans & Steyn 2008). Therefore, attempts were made to isolate ‘atypical’ E. ruminantium stocks in tick cells, atypical in the sense that they could not be initiated by using the classical ways of infecting endothelial cells (Bezuidenhout et al. 1985; Byrom et al. 1991).

Since the early reports of the E. ruminantium Omatjenne and Kümm stocks, it has been clear that these stocks share many phenotypical and genetic characteristics, but they differ from all other isolates (Allsopp et al. 1997, 2001; Du Plessis 1985, 1990). The Kümm stock was prepared from a goat, which was diagnosed with heartwater, from the Northern Province of South Africa, a heartwater endemic area (Du Plessis & Kümm 1971). Sheep injected with a lymph node suspension from goat developed heartwater symptoms. After more than 100 passages in mice it was still found to be pathogenic in mice, sheep and goats, but non-pathogenic to cattle (Du Plessis 1982). Many attempts were made to culture this organism in endothelial cells; however, it was only established in culture in 2002 using different monocyte cell lines (Zweygarth et al. 2002). It was observed that the Kümm stock comprised two 16S rRNA (16S ribosomal ribonucleic acid) genotypes, a 16S genotype typical of West African isolates (Kümm1) isolated in a canine macrophage-monocyte cell line (DH82) and a 16S genotype identical to E. ruminantium (Omatjenne) (Kümm2) isolated in a sheep blood mononuclear cell line (E2).
The Omatjenne genotype originated from the farm Omatjenne in the Otjiwarongo district of Namibia, a heartwater- and *Amblyomma*-free area (Du Plessis 1990). Healthy cattle on this farm reacted positively to *E. ruminantium* antigen using an indirect fluorescent antibody (IFA) test. Subsequently, ticks were collected from cattle on the farm and homogenates of individual ticks injected into mice. The serum of a single mouse, inoculated with homogenate prepared from a *Haemoloma truncatum* tick, tested positive in the IFA test. The original infective agent was non-pathogenic to mice, calves and sheep. Only after passing through three generations of *A. hebraeum*, it became fatal to sheep and mice (Du Plessis 1982). The organisms observed in brain smears of the sheep closely resembled those of the Kümm stock; fewer colonies of small size compared with those typically observed in animals infected with other *E. ruminantium* isolates.

Both stocks were atypical in that they are highly pathogenic to mice, but apparently non-pathogenic to bovine and could not be cultured in endothelial cells. The Kümm stock was described as atypical in that it infected mouse peritoneal macrophages (Du Plessis 1982). Because of the differences in pathogenicity and anomalous behaviour in cell culture, it was questioned whether the Kümm stock belonged to the species *E. ruminantium* (Du Plessis 1982). Likewise Allsopp et al. (1997) suggested that *E. ruminantium* (Omatjenne) (then *Cowdria ruminantium* [Omatjenne]), not to be confused with *Ehrlichia* sp. (Omatjenne) later renamed *Anaplasma* sp. (Omatjenne), may belong to a different species because of its difference in vector specificity and virulence.

Phylogenetic studies revealed that all *E. ruminantium* stocks analysed routinely grouped into one of two major clades, a West African clade and a southern/eastern African clade, except for Kümm2 and/or Omatjenne that clustered either as a unique group or in one of the major clades (Allsopp & Allsopp 2007; Allsopp et al. 1997, 2001; Steyn 2009; Van Heerden et al. 2004). Even isolates from several other geographical areas of Africa, the Indian Ocean islands and the Caribbean cluster with the southern and eastern African isolates can be included in a worldwide clade (Cangi et al. 2016). All these studies, however, were conducted with a limited number of genes, which do not necessarily allow the identification of recombination events. Furthermore, only small numbers of *E. ruminantium* isolates with different genotypes have been isolated in cell culture, which limits studies to link variation in DNA sequence to phenotypic characteristics. Therefore, there is a need to establish more isolates in cell culture that would enable us to conduct experiments to determine virulence, cross-protection between isolates, and to produce whole genome sequences.

This study reports on the isolation of ‘atypical’ *E. ruminantium* in tick cell cultures. In addition, we generated draft genome sequences of all three ‘atypical’ isolates and determined the differences between them and previously characterised *E. ruminantium* genomes.

## Materials and methods

### Infective agents

Three stocks of *E. ruminantium* were used. The Kümm2 genotype (Zweygarth et al. 2002), which was derived from the Kümm stock (Du Plessis 1982), was originally isolated from a naturally infected goat in Rust de Winter (Limpopo Province, South Africa). The *E. ruminantium* Omatjenne isolate was isolated from a single *H. truncatum* tick collected from a heartwater- and *Amblyomma*-free area of Namibia (Du Plessis 1990). Its complete 16S rDNA (16S ribosomal RNA) sequence was determined and submitted to GenBank® (accession number C. *ruminantium* [Omatjenne] U03776) (Allsopp et al. 1997). The original inoculums from which the Kümm and Omatjenne stocks were isolated are no longer available and the complete history of the blood stabliles used in this study is unknown. The third isolate was derived from the blood obtained from a sick angora goat from the farm Riverside (26.83°E, -33.45°S; Grahamstown, Eastern Cape Province, South Africa) (Steyn 2009).

### Cell cultures

The tick cell line *Ixodes scapularis* (ID8), derived from *Ixodes scapularis* embryos (Munderloh et al. 1994) was used. It was propagated in L-15B medium (Munderloh & Kurtti 1989), which was supplemented with 5% heat-inactivated foetal bovine serum (FBS), 10% tryptose phosphate broth, 0.1% bovine lipoprotein concentrate (MP Biomedicals, Santa Ana, CA, United States [US]), 100 IU/mL penicillin and 100 μg/mL streptomycin. The pH was adjusted at approximately 7.0. Infected tick cell cultures were maintained in Dulbecco’s modified Eagle’s medium nutrient mixture Ham F-12 (DME/F-12; Sigma, St. Louis, MO, US; D 0547) containing 15 mM HEPES and 1.2 g/L sodium bicarbonate, further supplemented with 10% (volume per volume [v/v]) heat-inactivated FBS, 2 mM l-glutamine, 100 IU/mL penicillin and 100 μg/mL streptomycin, and are referred to as DF-12.

### Infection of *Ixodes scapularis* (ID8) cell cultures

Each of the *E. ruminantium* stocks was used to infect Merino sheep by intravenous injection of 5 mL of an infectious blood stablile. The sheep subsequently developed symptoms associated with *E. ruminantium* infection. The body temperature of each sheep was monitored daily and a blood sample was collected when it exceeded 41.0 °C. Blood collected at the peak febrile reaction was used to initiate cell cultures according to the method described by Zweygarth et al. (2008). Thereafter the sheep were treated with tetracycline.

Blood was collected by venipuncture into sterile Vac-u-test® tubes containing heparin (lithium heparin, 14.3 United States Pharmacopoeia [USP] per mL blood) as anticoagulant and stored in ice. The cooled blood was centrifuged (800 × g; 10 min; 4 °C) and theuffy coat collected and washed with cold physiological phosphate-buffered saline (PBS). Theuffy coat was re-collected, and the red blood cells were lysed for...
approximately 30 seconds in 18 mL sterile distilled water followed by the addition of 2 mL of 10-fold concentrated Hanks’ balanced salt solution to restore physiological toxicity. The lysate was centrifuged for 5 min at 290 × g at 18 °C. The resulting cell pellet was re-suspended in 5 mL DF-12 and distributed into 25 cm² culture flasks containing IDE8 cells. The cultures were incubated at 32 °C.

DNA preparation and sequencing

The E. ruminantium elementary bodies were isolated from the cell culture material on discontinuous Percoll density gradients (Mahan et al. 1995) and the bacterial DNA was extracted with the DNeasy Blood & Tissue kit (Qiagen, GmbH, Hilden, Germany). The genomes were sequenced using Illumina Nextera paired-end libraries on the Illumina MiSeq and/or HiSeq platforms (Illumina, San Diego, CA, US).

De novo assembly

Sequencing reads were processed and assembled using CLC Genomics Workbench version 7.0 (https://www.qiagenbioinformatics.com/). Default parameters were used for quality trimming, and adapter sequences were removed. Trimmed reads < 50 bp were discarded. Several de novo assemblies using different combinations of parameters were performed for each data set. The following stringent parameters were used in all assemblies: mismatch cost, insertion cost and deletion cost of three, and length fraction and similarity fraction = 0.9. We varied the minimum contig length, also whether to use global alignment and whether to perform scaffolding or not. Contigs <500 bp and <10-fold coverage were discarded; the remaining contigs were blasted with the National Center for Biotechnology Information’s (NCBI) nucleotide BLAST (blastn, ncbi.nlm.nih.gov/blast/executables/blast+/LATEST/); the tabular view option (n = 8) allowed visualisation of the alignments in the Artemis Comparison Tool (ACT) programme (Carver et al. 2005). In addition, BLAST Ring Image Generator (BRIG) (Alikhan et al. 2011) was used to compare whole genomes and subsets of genes. We used the locus tags for coding sequence (CDS) from the original annotation and publication (Collins et al. 2005) of the Welgevonden and Gardel sequences. Whole-chromosome alignments were performed locally with Blastall (ftp://ftp.ncbi.nlm.nih.gov/blast/executables/blast+/LATEST/); the tabular view option (n = 8) allowed visualisation of the alignments in the Artemis Comparison Tool (ACT) programme (Carver et al. 2005). In addition, BLAST Ring Image Generator (BRIG) (Alikhan et al. 2011) was used to compare whole genomes and subsets of genes. We used the locus tags for coding sequence (CDS) from the original annotation and publication (Collins et al. 2005) of the Welgevonden strain of E. ruminantium. The corresponding Reference Sequence database (RefSeq) locus tags created subsequently by the NCBI Prokaryotic Genome Annotation Pipeline are listed in Table 1-A1.

Ethical consideration

Experiments were performed in accordance with the stipulations of the animal ethics committee at ARC-Onderstepoort Veterinary Research and approved by the South African Department of Agriculture, Forestry and Fisheries under Section 20 of the Animal Disease Act of 1984.

Results

Infection of IDE8 cell cultures

Leukocytes isolated from the blood of infected sheep were used as infective inoculum. All three E. ruminantium stocks were established successfully in IDE8 cell cultures by this method (Table 1). The Kümm2 stock was detected in Giemsa-stained smears 28 days after initiation. Infection with the Omajenene stock was first demonstrated on day 47 post-infection. The Riverside isolate was detected in stained smears in IDE8 cell cultures of any of the three isolates using bovine endothelial cells as host cells (data not shown).
Genome assembly

The average coverage of all assemblies was greater than 100-fold, and the draft genome sequences of Kümm2, Omatjenne and Riverside isolates comprised 6, 7 and 9 contigs, respectively (Table 2). All contigs of these three genomes were successfully mapped to the reference genome (Figure 1-A1). The total length of the joined contig sequences ranged between 1448 megabases (Mb) and 1455 Mb. The remaining gaps were in repeat regions, including both tandem repeats and dispersed repeats with large repeat units.

In silico multi-locus sequence typing analysis

Multi-locus sequence typing was used to illustrate the relationship between different E. ruminantium isolates. The South African isolates (Welgevonden, Ball3, Mara87/7, Blaauwkrans and Kwanyanga), as well as the isolate from the Caribbean (Gardel), shared more than 99% identity across the eight genes selected for multi-locus sequence typing (MLST), whereas the West African isolates shared 98% identity with the South African isolates (Table 2-A1). The MLST sequences of the three ‘atypical’ isolates were identical, but had only 88% identity compared with other E. ruminantium isolates and were separated into a distinct clade in the phylogenetic tree (Figure 1). The other three species of Ehrlichia shared 88% – 89% identity (genetic distances 0.11–0.13), whereas different Anaplasma species are more diverse with 68.5% – 92.5% identity (distance of 0.08–0.15). In fact, the genetic distance between A. marginale and A. centrale (0.08, 92.5% identity) is considerably closer than that between the ‘atypical’ and other E. ruminantium isolates (0.12, 88% identity).

General genome features

All three genomes have a low guanine-cytosine (GC) content (28.94% – 28.95%) and a low protein-coding capacity (65% (Table 3), which is comparable with the 27.5% GC and 63% – 64% coding capacity of other E. ruminantium genomes (Frutos et al. 2006; Nakao, Jongejan & Sugimoto 2016). In each genome, one set of rRNA genes, 36 transfer RNA (tRNA) genes, Transfer-messenger RNA (tmRNA) and two non-coding RNA (ncRNA) genes were identified.

The genomes generated in this study still have gaps because of dispersed repeats and large tandem repeats, and certain of the assembled tandem repeats are missing repeat units. These facts could contribute to the smaller genome size (1.45 Mb) of the draft genomes compared with 1.5 Mb reported for the Welgevonden and Gardel sequences (Collins et al. 2005; Frutos et al. 2006). It could also explain the slightly higher protein-coding capacity calculated for the assemblies because the repeat regions are mostly located in non-coding regions.

The gene content and genomic synteny were highly conserved between the three genomes as well as compared with previously sequenced genomes (Figure 2 and Figure 2-A1). Small inversions and rearrangements were found in the area around the shift in GC-skew value between 690 kilobases (kb) and 740 kb as well as other regions where duplications are found (at 630 kb, 1185 kb and 1248 kb) (Figures 1-A1 – 3-A1).

TABLE 1: Infection of IDE8 cell cultures using leukocytes from ovine blood isolated from sheep infected with three E. ruminantium stocks.

| E. ruminantium stock | First positive day/days to first subculture/days in culture | Passages |
|----------------------|-----------------------------------------------------------|----------|
| Kümm2                | 28/31/239                                                 | 9        |
| Omatjenne            | 47/72/169                                                 | 5        |
| Riverside            | 25/32/295                                                 | 10       |

TABLE 2: Assembly information.

| Assembly       | Kümm2 (bp) | Omatjenne (bp) | Riverside (bp) |
|----------------|------------|----------------|----------------|
| CLC assembly†  | 195        | 160            | 221            |
| Average x coverage | 109.6     | 104.6          | 134.9          |
| Number of contigs | 41        | 63             | 44             |
| Average contig length (bp) | 35 603 (11074–208 441) | 23 890 (1086–107 642) | 32 608 (1122–240 823) |
| Total length (bp) | 1 459 740 | 1 434 720       | 1 434 795       |
| GAP4 assembly   | 6          | 7              | 9              |
| Number of contigs | 242 475   | 207 235        | 160 924        |
| Average contig length (bp) | 1 454 859 | 1 450 642       | 1 448 329       |

bp, base pairs.† Information in table represents the CLC assembly with the largest average contig size and the smallest number of contigs.

FIGURE 1: Phylogenetic comparison of the multi-locus sequence typing. Unweighted Pair Group Method using Arithmetic averages tree constructed from the aligned concatenated sequences of the eight selected genes.
A high rate of DNA reorganisation in the terminus region is often observed between closely related bacteria, which may be associated with the mechanism of chromosome separation after replication (Hughes 2000).

Single nucleotide polymorphism (SNP) and insert or deletion (INDEL) analysis revealed that when the repeat regions were excluded, only four SNPs and four INDELs distinguished the Kümm2 and Omatjenne sequences. The Riverside sequence differed from either Kümm2 or Omatjenne by 328 SNPs and 16 INDELs.

Comparison of membrane protein families

Although gene content and genomic synteny are highly conserved, variation was observed between the genomes generated in this study and those reported previously. The variation was mainly limited to genes predicted to encode hypothetical proteins or membrane proteins and specifically the members of the four membrane protein families described previously (Collins et al. 2005). We identified the orthologous families in the Kümm2, Omatjenne and Riverside genomes, and found that their arrangement and number were identical in all the three genomes sequenced in this study. There were, however, variations compared with other E. ruminantium isolates. The major antigenic protein 1 (map1) family has been described in various isolates. The nucleotide sequences of the members of map1 family in Kümm2, Omatjenne and Riverside show a high degree of similarity to Welgevonden (Figure 3a), except for map1-5. In the syntenic locus of map1-5, two smaller open reading frames (ORFs) were identified in the ‘atypical’ genomes (Figure 3b).

Two other families described in Collins et al. (2005), here designated membrane family 1 (Erum2240–Erum2340; Erum2400; Erum2410) and membrane family 2 (Erum2750–Erum2800; Erum3600–Erum3630), were less conserved at nucleotide level (Figure 4a) and both of these families have one member less compared with the Welgevonden annotation (Figure 4b). Three of the genes in membrane family 2 are in opposite orientation in the ‘atypical’ strains as compared with Welgevonden.

Orthologs of four predicted membrane proteins (Erum7990, Erum8000, Erum8010 and Erum8020; membrane family 3)
were identified in the same relative location in the three newly sequenced genomes (Figure 4-A1). In all three genomes at this location, however, membrane family 3 expanded to seven members and five more members were identified at 82 Mb upstream (Figure 4-A1, Table 3-A1).

**Discussion**

*In vitro* isolation of heartwater organisms in cell culture is undoubtedly the most valuable aid for research in heartwater. For years, *E. ruminantium* was isolated from the blood or plasma of reacting animals by cultivation in ruminant endothelial cells, although it was recognised quite early that it was not possible to initiate *in vitro* cultures of some strains using this approach (Bezuidenhout & Brett 1992; Bezuidenhout et al. 1988).

It was shown in previous experiments that leukocytes obtained from four sheep each of which was infected with a different stock of *E. ruminantium* were able to infect IDE8 tick cell cultures and endothelial cell cultures simultaneously (Zweygarth et al. 2008). We applied this technique to the three ‘atypical’ strains that could not be isolated in endothelial cells, the Kümm (Du Plessis 1982), Omatjenne (Du Plessis 1990) and Riverside (Steyn 2009) isolates. Here we show that all three isolates could be established in IDE8 tick cell cultures and continuously propagated for 169 days or more. These cultures provided excess material for additional molecular characterisations of the isolates.

The most distinctive characteristic of the ‘atypical’ isolates is the fact that they cannot be initiated in ruminant endothelial cell cultures. However, some of the reported anomalies may
FIGURE 4: Genomic comparison of membrane families (1) (Erum2240–2410) and (2) (Erum2750–3630). (a) BRIG alignments of the three isolates against selected *E. ruminantium* Welgevonden gene sequences. (b) Linear representation of the alignments indicating members of the families in black with adjacent genes in grey. The locus tag prefixes for Kümm2 (EDL79), Riverside (EDL81) and Omatjenne (EDL80) were omitted to simplify the image.

kbp, kilobase pairs; bp, base pairs.
be contributed to other experimental factors. The experiments were conducted using uncharacterised inoculums passaged many times in different animals during the pre-PCR era when it was difficult to detect pathogen-free animals. Hence, the original Kümm stock comprised two organisms or one of them was introduced over the years during passaging. There is also a possibility that the initial Omatjenne agent is not the same as the organism we have cultured in this study. The pathogenicity and vector specificity of the cultured organisms need to be verified experimentally.

The sequences of the concatenated MLST loci of the three ‘atypical’ isolates were identical; however, they formed a distinct clade in tree topology. These results were confirmed by genome sequences. Although these three sequences only differed by a few SNPs and INDELs, as well as variation and small gaps in repeat regions that may be ascribed to assembly errors, they are markedly different from other E. ruminantium genomes. The 16S rRNA and map1 gene sequences identified the Omatjenne agent, and later Kümm2, as E. ruminantium in the southern African clade (Allsopp et al. 1997; Van Heerden et al. 2004). In contrast, tree topology and pairwise comparison of eight genes presented in this study may support an argument for the ‘atypical’ isolates to be classified as a separate species. The genetic distances and identity shared between the ‘atypical’ isolates and other E. ruminantium isolates are similar to the distances and percentage identity between the different species of Ehrlichia and Anaplasma. Whether these three isolates indeed represent a unique species needs to be validated.

All the E. ruminantium genomes sequenced thus far are syntenic (Frutos et al. 2006; Nakao Jongejan & Sugimoto 2016), and it is known that Anaplasma spp. and Ehrlichia spp. share conserved gene order (Dunning Hotopp et al. 2006; Pierlé et al. 2012). The synteny is also observed for the three isolates sequenced in this study with a few exceptions in the membrane protein families. Of the four families analysed, the map1 family was the most conserved one across all E. ruminantium genomes. The paralogs are maintained in the same order in all genomes, but in place of map1-5, two small ORFs were detected in the genomes presented here. The map1-5 gene is truncated compared with the other paralogs in all isolates analysed thus far. It was also identified as one of the paralogs that undergoes balancing selection, a type of selection that is reported to maintain genetic variation in genes that are involved in evasion of host immune response (Salim et al. 2019).

Most variation was detected in membrane family 1 and membrane family 2. The nucleotide sequences differed significantly between orthologous genes, and the number, order and, in some cases, orientation of the paralogs were different in new annotations. In membrane family 3, the nucleotide sequences between orthologs were conserved, but this family was expanded from four to 12 members in the ‘atypical’ genomes. At present, the function of these putative proteins is unknown, and it has not been shown that all members of these two protein families are expressed. Therefore, it is not known whether these variations have an effect on the expression or function of the predicted proteins encoded by these ORFs. It is known that Anaplasma and Ehrlichia spp. present a wide range of paralogous genes encoding various functions that ensure survival in diverse host and vector environments (Dunning Hotopp et al. 2006).

Several studies have reported a high level of genetic diversity among E. ruminantium isolates (Allsopp & Allsopp 2007; Cangi et al. 2016; Nakao et al. 2011). In contrast, here we found that the genome sequences of the three ‘atypical’ isolates from distant geographical areas and diverse habitats are almost identical. Excluding repeat regions, Kümm2, from the Limpopo Province of South Africa, and Omatjenne, from the much drier and heartwater-free Otjiwarongo district of Namibia, differ by four SNPs and four INDELs only. In fact, there were more substitutions and small deletions detected between the parental Welgevonden strain and its daughter strain after 11–13 passages in a different cell culture environment (Frutos et al. 2006).

Although the current results have not connected any genetic variation to the phenotypes that distinguish these isolates, variations in the membrane protein families may contribute to the ability of these organisms to infect different cells. A comprehensive SNP analysis, including all genomes sequenced, may elucidate the determinants of diversity. The synteny conservation in E. ruminantium genomes suggests that at least some of the phenotypes are associated with small polymorphisms. In view of this, we are in the process of generating whole genome sequences of all the E. ruminantium isolates we have in cell culture. In addition, we need to establish more field isolates in culture to conduct phenotypic and genotypic analyses in the future work. To date, no standard cell line has been designated for isolation of E. ruminantium, and it is clear that the ‘atypical’ isolates cannot be easily isolated in bovine endothelial cells. We therefore recommend the use of both ruminant endothelial cells and tick cell cultures concurrently. The two methods complement each other and should be used when isolating field strains of Ehrlichia spp.

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Author’s contributions
J.L. and E.Z. conceptualised and wrote the article. E.Z. performed the cell culture experiments and interpreted results. J.L. assembled and analysed the sequencing data. H.C.S. performed experimental work involving animals and contributed to the editing of the article. A.I.J. and E.F.
contributed to experimental laboratory work and reviewed the article. All authors read and approved the final article.

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**Data availability statement**

The nucleotide sequences and annotation of the three genomes are available from NCBI (https://www.ncbi.nlm.nih.gov/genome/microbes/). The accession numbers are listed in Table 3.

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The views expressed in this article are those of the authors and do not necessarily reflect the official policy or position of any affiliated agency of the authors or funders.

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### Appendix 1

**TABLE 1-A1:** The RefSeq locus tags of the members of the membrane protein families.

| Membrane protein family | ORF ID   | Gene name | Original annotation | Functional classification | RefSeq locus tags   |
|-------------------------|----------|-----------|---------------------|---------------------------|--------------------|
| Membrane family 1       | Erum2230 | trmU      | tRNA (5-methylaminomethyl-2-thiouridylate)-methyltransferase    | ERUM_RS01235              |
|                         | Erum2240 | -         | membrane protein     | ERUM_RS01240              |
|                         | Erum2250 | -         | membrane protein     | ERUM_RS01245              |
|                         | Erum2260 | -         | membrane protein     | ERUM_RS01250              |
|                         | Erum2270 | -         | membrane protein     | ERUM_RS01255              |
|                         | Erum2280 | -         | membrane protein     | ERUM_RS01260              |
|                         | Erum2290 | -         | membrane protein     | ERUM_RS01265              |
|                         | Erum2300 | -         | membrane protein     | ERUM_RS01270              |
|                         | Erum2310 | -         | membrane protein     | ERUM_RS01275              |
|                         | Erum2320 | -         | membrane protein     | ERUM_RS01280              |
|                         | Erum2330 | -         | membrane protein     | ERUM_RS01285              |
|                         | Erum2340 | -         | membrane protein     | ERUM_RS01290              |
|                         | Erum2400 | -         | membrane protein     | ERUM_RS01320              |
|                         | Erum2410 | -         | membrane protein     | ERUM_RS01325              |
|                         | Erum2420 | gyrA      | DNA gyrase subunit A | ERUM_RS01330              |
| Membrane family 2       | Erum2740 | -         | integral membrane transport protein                       | ERUM_RS01510              |
|                         | Erum2750 | -         | membrane protein     | ERUM_RS01515              |
|                         | Erum2760 | -         | membrane protein     | ERUM_RS01520              |
|                         | Erum2770 | -         | membrane protein     | ERUM_RS01525              |
|                         | Erum2780 | -         | membrane protein     | ERUM_RS01530              |
|                         | Erum2790 | -         | membrane protein     | ERUM_RS01535              |
|                         | Erum2800 | -         | membrane protein     | ERUM_RS01540              |
|                         | Erum3600 | -         | membrane protein     | ERUM_RS01960              |
|                         | Erum3610 | -         | membrane protein     | ERUM_RS01965              |
|                         | Erum3620 | -         | membrane protein     | ERUM_RS01975              |
|                         | Erum3630 | -         | membrane protein     | ERUM_RS01980              |
|                         | Erum3640 | -         | unknown              | ERUM_RS01985              |
| Membrane family 3       | Erum7570 | -         | NADH-ubiquinone oxido-reductase                             | ERUM_RS04090              |
|                         | Erum7580 | -         | integral membrane transport protein                       | ERUM_RS04095              |
|                         | Erum7980 | -         | type IV secretion system protein                           | ERUM_RS04310              |
|                         | Erum7990 | -         | integral membrane protein                                 | ERUM_RS04315              |
|                         | Erum8000 | -         | integral membrane protein                                 | ERUM_RS04320              |
|                         | Erum8010 | -         | integral membrane protein                                 | ERUM_RS04325              |
|                         | Erum8020 | -         | integral membrane protein                                 | ERUM_RS04330              |
|                         | Erum8030 | hflK      | FtsH protease activity modulator                          | ERUM_RS04335              |
| map1 family             | Erum8580 | -         | transcriptional regulator                                 | ERUM_RS04630              |
|                         | Erum8590 | map1-14    | outer membrane protein                                    | ERUM_RS04635              |
|                         | Erum8600 | map1-13    | outer membrane protein                                    | ERUM_RS04640              |
|                         | Erum8610 | -         | exported protein                                            | ERUM_RS04645              |
|                         | Erum8620 | map1-12    | outer membrane protein                                    | ERUM_RS04650              |
|                         | Erum8630 | map1-11    | outer membrane protein                                    | ERUM_RS04655              |
|                         | Erum8640 | map1-10    | outer membrane protein                                    | ERUM_RS04660              |
|                         | Erum8650 | map1-9     | outer membrane protein                                    | ERUM_RS04665              |
|                         | Erum8660 | map1-8     | outer membrane protein                                    | ERUM_RS04670              |
|                         | Erum8670 | map1-7     | outer membrane protein                                    | ERUM_RS04675              |
|                         | Erum8680 | map1-6     | outer membrane protein                                    | ERUM_RS04680              |
|                         | Erum8690 | map1-5     | outer membrane protein                                    | ERUM_RS04685              |
|                         | Erum8700 | map1-4     | outer membrane protein                                    | ERUM_RS04690              |
|                         | Erum8710 | map1-3     | outer membrane protein                                    | ERUM_RS04695              |
|                         | Erum8720 | map1-2     | outer membrane protein                                    | ERUM_RS04700              |
|                         | Erum8730 | map1-1     | outer membrane protein                                    | ERUM_RS04705              |
|                         | Erum8740 | map1       | outer membrane protein                                    | ERUM_RS04710              |
|                         | Erum8750 | map1+1     | outer membrane protein                                    | ERUM_RS04715              |
|                         | Erum8760 | -         | unknown                                                      | ERUM_RS04720              |

Note: Adjacent genes in bold.
| TABLE 2-A1: Pairwise comparison table of the multi-locus sequence typing alignment. The upper comparison shows percentage of identical bases in alignment positions between two isolates and the lower comparison depicts the Jukes-Cantor distance between sequences. |
|---|
| **E. ruminantium isolate** | No. 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 |
| Welgevonden | 1 | - | 99.56 | 99.26 | 99.36 | 99.35 | 99.36 | 97.67 | 97.70 | 88.40 | 88.40 | 88.40 | 82.29 | 81.85 |
| Ball3 | 2 | 0.00 | - | 99.28 | 99.39 | 99.48 | 99.36 | 97.74 | 97.84 | 88.41 | 88.41 | 88.41 | 82.25 | 81.91 |
| Mara | 3 | 0.01 | 0.01 | - | 99.69 | 99.60 | 99.26 | 97.74 | 97.77 | 88.49 | 88.49 | 88.49 | 82.24 | 81.83 |
| Blauwkrans | 4 | 0.01 | 0.01 | 0.00 | - | 99.90 | 99.11 | 97.69 | 97.84 | 88.41 | 88.41 | 88.41 | 82.27 | 81.83 |
| Kwanyanga | 5 | 0.01 | 0.01 | 0.00 | 0.00 | - | 99.12 | 97.72 | 97.77 | 88.44 | 88.44 | 88.44 | 82.28 | 81.88 |
| Gardel | 6 | 0.01 | 0.01 | 0.01 | 0.01 | 0.01 | - | 97.87 | 97.88 | 88.45 | 88.45 | 88.45 | 82.30 | 81.89 |
| Senegal | 7 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | - | 99.94 | 88.46 | 88.46 | 88.46 | 82.27 | 81.71 |
| Sankat | 8 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.00 | - | 88.46 | 88.46 | 88.46 | 82.27 | 81.70 |
| Kümm2 | 9 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | - | 100 | 100 | 82.39 | 82.11 |
| Omatjenne | 10 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.00 | - | 100 | 82.39 | 82.11 |
| Riverside | 11 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.12 | 0.00 | 0.00 | - | 82.39 | 82.11 |
| E. canis | 12 | 0.19 | 0.02 | 0.02 | 0.19 | 0.19 | 0.19 | 0.19 | 0.19 | 0.19 | 0.19 | 0.19 | - | 87.86 |
| E. chaffeensis | 13 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.02 | 0.13 | - |

| TABLE 3-A1: Membrane family 3 of the Welgevonden isolate compared to Kümm2, Omatjenne and Riverside. |
|---|
| **Welgevonden** | **Kümm2** | **Omatjenne** | **Riverside** |
| 1 377 584…1 387 657† | 1 319 020…1 329 752 | 1 315 503…1 326 234 | 1 312 770…1 323 594 |
| Erum7980 (R‡) | EDL79_04495 (R) | EDL80_04485 (R) | EDL81_04475 (R) |
| | EDL79_04500 (R) | EDL80_04490 (R) | EDL81_04480 (R) |
| Erum7990 (R) | EDL79_04505 (R) | EDL80_04495 (R) | EDL81_04485 (R) |
| Erum8000 (R) | EDL79_04510 (R) | EDL80_04500 (R) | EDL81_04490 (R) |
| Erum8010 (R) | EDL79_04515 (R) | EDL80_04505 (R) | EDL81_04495 (R) |
| Erum8020 (R) | EDL79_04520 (R) | EDL80_04510 (R) | EDL81_04500 (R) |
| | EDL79_04525 (F) | EDL80_04515 (F) | EDL81_04505 (F) |
| | EDL79_04530 (R) | EDL80_04520 (R) | EDL81_04510 (R) |
| Erum8030 (hflK) (F) | EDL79_04535 (hflK) (F) | EDL80_04525 (hflK) (F) | EDL81_04515 (hflK) (F) |
| 1 294 686…1 296 536 | 1 237 185…1 242 924 | 1 233 416…1 239 155 | 1 230 696…1 236 436 |
| Erum7570 (F) | EDL79_04255 (F) | EDL80_04245 (F) | EDL81_04235 (F) |
| | EDL79_04260 (R) | EDL80_04250 (R) | EDL81_04240 (R) |
| | EDL79_04265 (R) | EDL80_04255 (R) | EDL81_04245 (R) |
| | EDL79_04270 (R) | EDL80_04260 (R) | EDL81_04250 (R) |
| | EDL79_04275 (R) | EDL80_04265 (R) | EDL81_04255 (R) |
| | EDL79_04280 (R) | EDL80_04270 (R) | EDL81_04260 (R) |
| Erum7580 (F) | EDL79_04285 (F) | EDL80_04275 (F) | EDL81_04265 (F) |

Note: The orthologs flanking the members of the membrane protein family are set in bold. F, forward strand; R, reverse strand.
†, base location in genome sequence; ‡, orientation of open reading frame (ORF).
FIGURE 1-A1: Mauve alignment of the contigs of the three draft genome sequences against the linearised genome sequence of *E. ruminantium* Welgevonden (top panel). The second, third and bottom panels represent the ordered contig sequences, separated with vertical red lines, of Kümm2, Omatjenne and Riverside respectively. Each coloured block represents a segment of the genome sequence that aligned to a homologous part of another genome without any genome rearrangements. Blocks above the centre line of each panel is in the forward orientation relative to the reference genome sequence, while those below the centre line indicate regions that align in the inverse orientation. The height of the similarity profile inside a block corresponds to the average level of conservation in that region.
FIGURE 2-A1: ACT alignment of the three draft genome sequences against the whole genome sequence of *E. ruminantium* Welgevonden. The coloured lines drawn between two adjacent linearised chromosomes show the location of homologous genes and indicate the same (red) or opposite (blue) orientation relative to the chromosome immediately above.

Source: Carver, T., Thomson, N., Bleasby, A., Berriman, M. & Parkhill, J., 2009, 'DNAPlotter: Circular and linear interactive genome visualization', Bioinformatics 25(1), 119–120. https://doi.org/10.1093/bioinformatics/btn578

FIGURE 3-A1: Circular representations of the genomes generated with DNAPlotter. In each circle the tracks from outside to inside represent: (a) coding sequences on the forward (blue) and (b) reverse strands (red), % GC plot, and (c) GC skew ([GC]/[G+C]).
FIGURE 4-A1: ACT alignment of the *E. ruminantium* Kümm2 sequence (bottom sequence in each panel) against *E. ruminantium* Welgevonden (top sequence in each panel) to depict the expansion of membrane family 3 in the Kümm2 isolate. The coloured lines drawn between the sequences show the location of homologous genes in the same (red) or opposite (blue) orientation. Top (a) Four *E. ruminantium* Welgevonden open reading frames matching open reading frames in two locations (yellow lines) in the *E. ruminantium* Kümm2 annotation. Middle (b) The syntenic matches of the four Welgevonden ORFs to seven Kümm2 open reading frames. Bottom (c) The same four Welgevonden open reading frames matching five Kümm2 open reading frames upstream.