

Criibacterium bergeronii gen. nov., sp. nov., a new member of the family *Peptostreptococcaceae*, isolated from human clinical samples

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Abstract

A rod-shaped, motile anaerobic bacterium, designated CCRI-22567^T, was isolated from a vaginal sample of a woman diagnosed with bacterial vaginosis and subjected to a polyphasic taxonomic study. The novel strain was capable of growth at 30–42 °C (optimum, 42 °C), at pH 5.5–8.5 (optimum, pH 7.0–7.5) and in the presence of 0–1.5% (w/v) NaCl (optimally at 0.5% NaCl). The phylogenetic trees based on 16S rRNA gene sequences showed that strain CCRI-22567^T forms a distinct evolutionary lineage independent of other taxa in the family *Peptostreptococcaceae*. Strain CCRI-22567^T exhibited 90.1% 16S rRNA gene sequence similarity to *Peptoanaerobacter stomatis* ACC19a^T and 89.7% to *Eubacterium yurii* subsp. *schtitka* ATCC 43716. The three closest organisms with an available whole genome were compared to strain CCRI-22567^T for genomic relatedness assessment. The genomic average nucleotide identities (OrthoANIu) obtained with *Peptoanaerobacter stomatis* ACC19a^T, *Eubacterium yurii* subsp. *margaretiae* ATCC 43715 and *Filifactor alocis* ATCC 35896^T were 71.8, 70.3 and 69.6%, respectively. Strain CCRI-22567^T contained C_{18:1}ω9c and C_{18:1}ω9c DMA as the major fatty acids. The DNA G+C content of strain CCRI-22567^T based on its genome sequence was 33.8mol%. On the basis of the phylogenetic, chemotaxonomic and other phenotypic properties, strain CCRI-22567^T is considered to represent a new genus and species within the family *Peptostreptococcaceae*, for which the name *Criibacterium bergeronii* gen. nov., sp. nov., is proposed. The type strain of *Criibacterium bergeronii* is CCRI-22567^T (=LMG 31278^T=DSM 107614^T=CCUG 72594^T).

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Abbreviations: BV, bacterial vaginosis; CFA, cell fatty acid; CM, cytoplasmic membrane; Cryo-Et, cryo electron tomography; dDDH, digital DNA–DNA hybridization; EM, electron microscopy; OL, outer layer; PG, peptidoglycan; PYG, peptone yeast glucose; TSA, tryptic soy agar; TYGS, Type (Strain) Genome Server.

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The GenBank/EMBL/DBJ accession numbers for the 16S rRNA gene sequences of strains CCRI-22567^T and CCRI-24246 are NR_164617.1 and MN153816.1, respectively. The whole-genome shotgun project of strains CCRI-22567^T and CCRI-24246 have been deposited at DDBJ/EMBL/GenBank under the accession numbers MBEW00000000 and VJXW00000000, respectively. The versions described in this paper are MBEW00000000.2 and VJXW00000000.1.

Two supplementary tables and three supplementary figures are available with the online version of this article.

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Strain CCRI-22567^T was isolated from a vaginal clinical swab harvested from a woman diagnosed with bacterial vaginosis (BV; Nugent score of 9) in Spring 2014 at CHU de Québec-Université Laval (Quebec City, QC, Canada). Vaginal swabs requested by treating physicians are routinely submitted for BV testing using Nugent score at the clinical microbiology laboratory of the CHU de Québec-Université Laval (CHUL hospital). Swabs were streaked on sterile autoclaved glass slides so that swabs can be stored at 4 °C until the Nugent score report is generated. Anonymized swabs having Nugent scores between 7 and 10 were used for a research project to isolate BV-associated *Mobiluncus* species. This recovery of deidentified leftover clinical sample swabs was approved by the Research Ethics Committee of the 'CHU de Québec-Université Laval' (No. B13-08-1778, 26 September 2013). Swabs were used to inoculate Schaedler agar medium containing vitamin K and 5% sheep blood. An isolated translucent round colony growing to <0.5 mm after 2 days of incubation at 35 °C under anaerobic conditions was subjected to 16S rRNA gene sequencing for species identification. Comparative 16S rRNA gene sequence analysis showed that strain CCRI-22567^T represents a novel species of a new genus in the family *Peptostreptococcaceae*. The role of this new bacterium in BV remains unclear since *Gardnerella vaginalis*, *Bacteroides fragilis* and *Mobiluncus curtisii* (all species consistent with BV) were also isolated from the same vaginal swab. The comparison of the 16S rRNA gene sequence of strain CCRI-22567^T with microbiome projects available in the GenBank NR database suggests that the same novel species is also present on sebaceous parts of the skin [1]. Furthermore, strain CCRI-24246 (=L9284001) was isolated in 2018 from pus cultivated from the left armpit of an 81-year-old woman, suggesting that these bacterial isolates most probably represent a commensal inhabitant of both skin and vaginal discharge. *Anaerococcus prevotii*, another member of the family *Peptostreptococcaceae*, is also well known to be an inhabitant of these sites [2]. To determine the taxonomic position of these isolates, they were subjected to further characterization in accordance with the recommendations outlined by Kämpfer *et al.* and Tindall *et al.* [3, 4].

Genomic DNA of strains CCRI-22567^T and CCRI-24246 was extracted for 16S rRNA and whole genome sequencing as described previously [5]. PCR amplification of the 16S rRNA gene was performed with primers SSU27 and SSU1492 and sequencing using primers SSU27, SSU536, SSU685 and SSU926 as described by Sistek *et al.* [6]. 16S rRNA sequences were used for an initial BLAST against the GenBank NR database to identify the taxonomic neighbours of strains CCRI-22567^T and CCRI-24246. Based on this analysis, the closest cultured relatives of strain CCRI-22567^T were *Peptoanaerobacter stomatis* ACC19a^T (90.1%), *Eubacterium yurii* subsp. *schtitka* ATCC 43716 (89.7%) and *Eubacterium yurii* subsp. *margaretiae* ATCC 43715 (89.6%), respectively. These sequence identities were calculated based on nearly full-length sequences using the pairwise distance function of MEGA version 6.06 [7]. These identity values were below the accepted cut-off of 98.7% for species delineation, and also

under the cut-off of 95% for the differentiation of a new genus [4, 8, 9]. Therefore, strains CCRI-22567^T and CCRI-24246 meet the criteria of the description of a novel species and a new genus.

Subsequently, strains CCRI-22567^T and CCRI-24246 16S rRNA sequences were aligned with all type strain sequence of every species of the family *Peptostreptococcaceae* according to LPSN [10] using MUSCLE implemented in MEGA version 6.06 [7] to establish its relationship within the family *Peptostreptococcaceae*. Genera assigned to the families *Peptostreptococcaceae* and *Clostridiaceae* deemed to be closest to *Criibacterium* gen. nov. were discussed together here as inferred by Galperin *et al.* [11]. Organisms with names not yet validly published were also included to fully represent the family tree. The three subspecies of *Eubacterium yurii* were included in the analysis as they are members of *Peptostreptococcaceae* based on 16S rRNA phylogeny and are waiting to be transferred to another genus [12]. The 16S rRNA gene phylogenetic trees reconstructed using the maximum-likelihood method were carried out on a comparison of 1262 nucleotides using MEGA version 6.06 [7] (Fig. 1). Tree topology was confirmed using the minimum-evolution method and tree clustering structure with the neighbour-joining method (Fig. S1 and Fig. S2, available in the online version of this article). The maximum-likelihood tree reconstruction was similar to the one reported previously by Galperin *et al.* [11]. Strains CCRI-22567^T and CCRI-24246 showed strong bootstrap values, supporting their relationship within the cluster consisting of *Peptoanaerobacter stomatis*, *Eubacterium yurii* and *Filifactor* species. That cluster is already known to form a deep branch within the family *Peptostreptococcaceae* [13]. While *Filifactor villosus* was isolated from a subcutaneous abscess, *Peptoanaerobacter stomatis*, *Eubacterium yurii* and *Filifactor alocis* are oral pathogens [14–17]

Whole-genome sequencing of strain CCRI-22567^T was performed on an Illumina HiSeq 2500 using SBS version 4 to sequence a 126 bp paired-end library (Nextera XT, Illumina). A first genome assembly version using Ray software was described in Maheux *et al.* [5] (MBEW00000000.1). A second *de novo* assembly version (MBEW00000000.2) was performed using SPAdes software (v3.10.1) [18]. *De novo* assembly of the 7787976 reads yielded 79 contigs with an overall coverage of 450-fold. The genome length was 2176288 bp (N50 of 80728 bp) and the G+C content was 33.8mol%. The assembled genome was annotated using NCBI GenBank annotation pipeline (version 4.6) [19]. The genome of strain CCRI-22567^T had a total of 2174 identified features, including six rRNAs, 40 tRNAs, four non-coding RNA and 2124 protein-coding sequences from which 436 were assigned as hypothetical proteins. Whole-genome sequencing of strain CCRI-24246 was performed on an Illumina MiSeq using MiSeq Reagent Kit version 3 to sequence the 300 bp paired-end library. A total of 12281540 reads were assembled *de novo* in 139 contigs using SPAdes (version 3.13.0) [18] for an overall coverage of 400-fold. The total genome length was 2611389 bp (N50 of 115916 bp) with an average G+C content of 32.9mol%. The draft genome sequence was annotated using NCBI GenBank

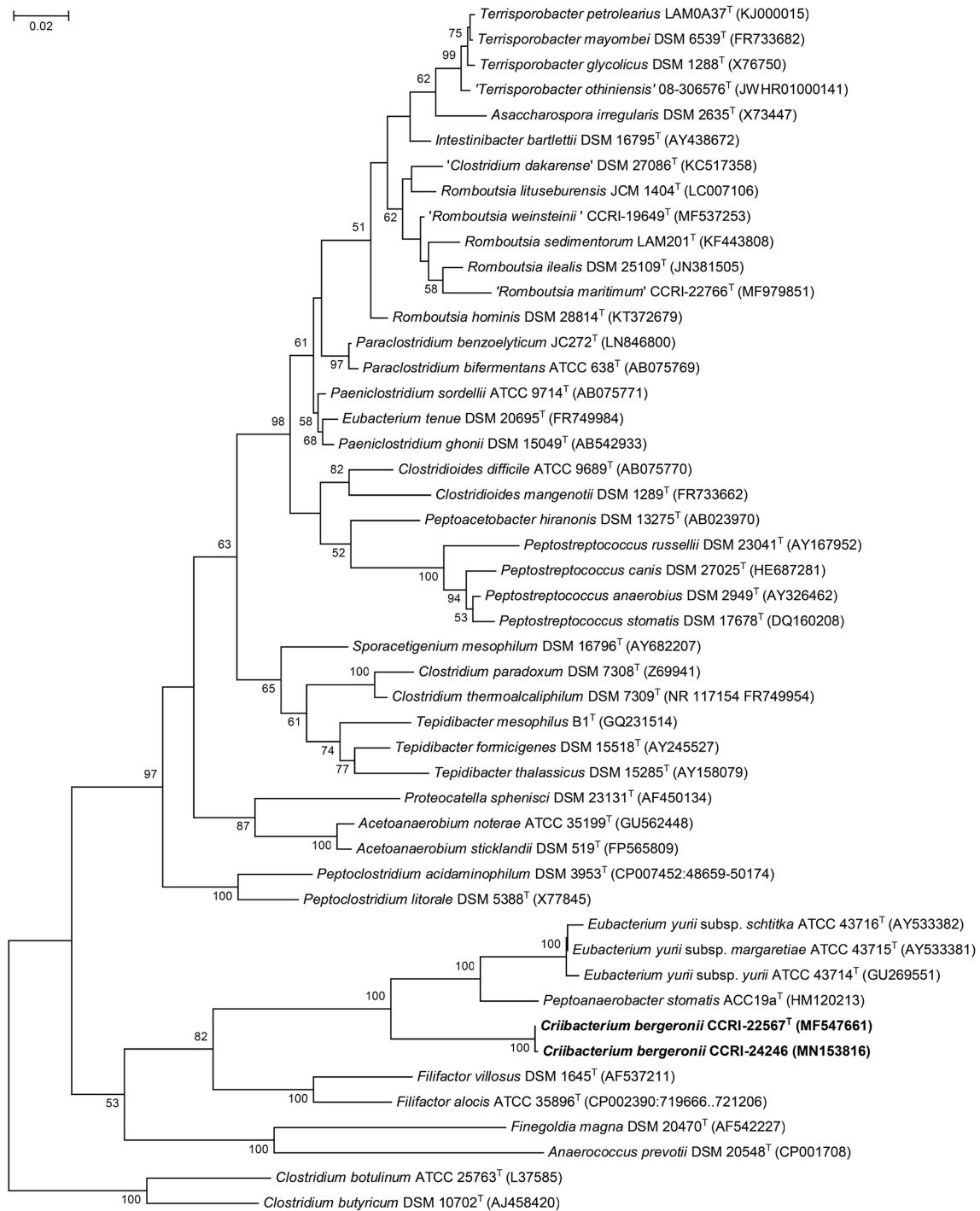


Fig. 1. Maximum-likelihood phylogenetic tree showing the relationship of *Criibacterium bergeronii* within the family *Peptostreptococcaceae*. The 16S rRNA sequences were aligned using MUSCLE [37]. The phylogenetic tree was reconstructed using the maximum-likelihood method with the Tamura–Nei substitution model implemented in MEGA version 6.0.6 [7]. Only bootstrap values >50% based on 1000 replications are shown at branching points. Organisms with names not validly published are presented with quotation marks. GenBank accession numbers for each 16S rRNA gene sequence are given in parentheses. Each bar indicates 0.02 substitutions per position. *Clostridium botulinum* ATCC 25763^T and *Clostridium butyricum* DSM 10702^T (*Clostridium sensu stricto*) were used as outgroups.

annotation pipeline (version 4.8) [19]. A total of 2554 features were identified, including seven rRNAs, 45 tRNAs, four non-coding RNA and 2497 putative coding sequences from which 516 were assigned as hypothetical proteins. As suggested by Chun *et al.* [20], 16S rRNA and *recA* genes sequences were extracted from genome assembly and compared to original Sanger sequences made on isolated strain CCRI-22567^T. The individual 16S rRNA and *recA* genes sequences were 100% identical (ambiguous bases were excluded) to sequences from the whole genome assembly, thus ensuring the authenticity of the whole genome data. Comparison of the proteins encoded by the two genomes using CD-HIT version 4.8.1 [21] revealed that strains CCRI-22567^T and CCRI-24246 have 1654 protein clusters in common (at 95% identity), while they have 441 and 820 unique protein clusters, respectively. Furthermore, the differences between the two genomes of strains CCRI-22567^T and CCRI-24246 are mainly found in regions of higher G+C content. The genes annotated in those parts belong to bacteriophages or transposons and thus appear to be related to mobile genetic elements.

According to Chun *et al.* [20], the 16S rRNA gene comparison could have been sufficient to describe strains CCRI-22567^T and CCRI-24246 as a new species/new genus with regard to the established cut-off [9, 20]. To further support this description, the genomic average nucleotide identity was also calculated using OrthoANIu software [22] with the strains CCRI-22567^T and CCRI-24246 genome assemblies and the closest related whole genomes available at the time. The OrthoANIu identity obtained was 71.8% with *P. stomatis* ACC19a^T (NZ_AFZF00000000.2), 70.3% with *Eubacterium yurii* subsp. *margaretiae* ATCC 43715 (NZ_AEES00000000.1) and 69.6% with *F. alocis* ATCC 35896^T (NC_016630.1). A predicted digital DNA–DNA hybridization (dDDH) value was also determined using the GGDC 2.1 web service using the same whole genomes (<http://ggdc.dsmz.de>) [23]. The returned dDDH values were 27.8, 25.6, and 25.4%, respectively. These OrthoANIu and dDDH values were below the 95–96 and 70% accepted threshold for species delineation, respectively, thus supporting the *C. bergeronii* new species description [20, 22, 24, 25]. The OrthoANIu and digital DDH values between the two strains of *C. bergeronii* were 98.5 and 88.7%, respectively, indicating that these two strains were members of the same species according to the previously mentioned cut-off values.

The relationship of strains CCRI-22567^T and CCRI-24246 within the family *Peptostreptococcaceae* was investigated using available genome sequence data and the Type (Strain) Genome Server (TYGS) platform [26]. First, the 16S rRNA sequence of CCRI-22567^T was used to identify all close type strains with available genomes. Then, the whole genomes of CCRI-22567^T and CCRI-24246 were used to determine intergenomic distances between them. A minimum-evolution genome-based tree, supported by strong bootstrap values, showed the same clustering of isolates CCRI-22567^T and CCRI-24246 within the *P. stomatis*, *Eubacterium yurii* spp. and *F. alocis* cluster (Fig. S3). This strong branching also supporting a new genus and a new species identification.

Moreover, the two strains, CCRI-22567^T and CCRI-24246, were also defined as belonging to the same species using complete genomes on the TYGS platform. The same taxonomic assignment as a new genus and a new species was also supported using Genome Taxonomy Database (GTDB-Tk version 1.3.0) [27]

A more detailed phenotypic characterization was performed to support the species description. Strains CCRI-22567^T and CCRI-24246 were stored at –80 °C in Brucella broth (BD) liquid medium containing 15% glycerol until the characterization tests could be performed. The strains were sub-cultured three times on sheep blood agar. Culture using pre-reduced media and phenotypic identification tests were performed as described in the Wadsworth and VPI anaerobic manuals [28, 29].

Strain CCRI-22567^T did not grow in thioglycolate and tryptic soy broths. Thus, physiological characterization was carried out on tryptic soy agar with 5% sheep blood (blood agar) for atmosphere and temperature tests and tryptic soy agar (TSA) for pH and NaCl concentration tests, respectively. Growth was monitored by direct observation. Growth of the type strain was first tested on blood agar in an anaerobic chamber containing a gas mix of 10% H₂, 5% CO₂ and balance with N₂, and microaerophilic conditions, in an anaerobic jar containing a Campy GasPak EZ (gas generating system pouch, Becton Dickinson and Company), as well as under aerobic conditions, with or without 5% CO₂. Strain CCRI-22567^T grew only under anaerobic conditions. The determination of the temperature range for growth was tested at 25, 30, 35 and 42 °C on blood agar in an anaerobic atmosphere. Strain CCRI-22567^T was capable of growth at 30–42 °C, with optimum growth at 42 °C. The pH range for growth was tested at pH 4.0, 4.5, 5.0, 5.5, 6.0, 6.5, 7.0, 7.5, 8.0, 8.5, 9.0, 9.5, 10.0 and 10.5, at 35 °C, on TSA using different buffers within their optimal range, in an anaerobic atmosphere. We used four different buffer systems: pH 4.0, 4.5, 5.0 and 5.5 with sodium acetate buffer; pH 6.0, 6.5, 7.0, 7.5 and 8.0 with sodium phosphate buffer; pH 8.5 and 9.0 with Bis-Tris propane buffer; pH 9.5, 10.0 and 10.5 with sodium carbonate buffer. Strains CCRI-22567^T and CCRI-24246 were capable of growth at pH values from pH 5.5 to 8.5, with optimum growth at pH 7.0–7.5. The NaCl concentration range for growth was tested at 0, 0.5, 1.0, 1.5, 2.0, 2.5, 3.0, 3.5, 4.0, 4.5, 5.5 and 6.0% (w/v), at 35 °C, on TSA and pH 7.0, in an anaerobic atmosphere. Strain CCRI-22567^T was capable of growth at concentrations of 0–1.5% (w/v) NaCl, with optimum growth at 0.5% NaCl.

Morphological characteristics of colonies and cells of the type strain were determined using light microscopy (Leitz) and cryo electron tomography (cryo-ET). Cryo-ET is currently the only tool that preserves biological samples at or near the native, hydrated state. It allows for the three-dimensional imaging of whole cells to macromolecular resolution (3–4 nm). Thus, cryo-ET was the method used here to determine the cell envelope architecture [30]. Cryo-ET data was collected on frozen-hydrated bacterial samples. Briefly, cells were frozen on EM grids using a Vitrobot automated vitrification

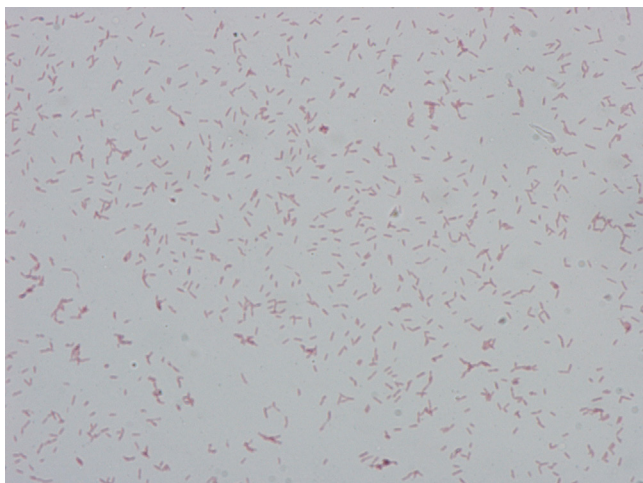


Fig. 2. Gram stain of *Criibacterium bergeronii* CCRI-22567^T after 96 h of incubation on sheep blood agar, under anaerobic atmosphere at 35 °C; ×1000 magnification.

device (ThermoFisher Scientific). Plunge-frozen samples were imaged on a Titan Krios 300kV transmission electron microscope equipped with a Falcon 2 direct electron detector (ThermoFisher Scientific). Tilt series were recorded from -60° to $+60^{\circ}$ with 1° increment oscillations, and a defocus of $-8\mu\text{m}$. The total final dose was $150\text{ e}/\text{\AA}$ [2]. Tilt series were recorded automatically using EPU Tomography software and tomographic reconstructions were calculated using the IMOD software package [31]. After 72 h of incubation on blood agar in an anaerobic chamber, colonies were grey-translucent, circular with entire edges, smooth and $\sim 0.6\text{ mm}$ in diameter. Colonies were non-adherent and no haemolysis was observed. Gram staining was performed using a four-step Gram stain kit (BD) on cells from 72 h blood agar culture. Cells stained Gram-variable (CCRI-22567^T stained Gram-negative, while CCRI-24246 stained Gram-variable, with red cells, purple cells and blue cells on the same glass slide) and both appeared as single cells or in pairs. Cells of strain CCRI-22567^T were rod-shaped ($\sim 630\text{ nm}$ wide and $\sim 1750\text{ nm}$ long) and non-spore-forming (Fig. 2). Although some genes of strains CCRI-22567^T and CCRI-24246 have been annotated as having homologies with known sporulation proteins, no genes presenting homologies with the Spo0A regulator protein, which is essential to sporulation initiation, has been identified. Gelatin hydrolysis was performed as described by De la Cruz and Torres [32]. CCRI-22567^T and CCRI-24246 did not grow on the nutrient gelatin medium. CCRI-22567^T and CCRI-24246 showed motility in wet mount and flagella were observed with cryo-ET. When CCRI-22567^T cells were imaged with cryo-ET (Fig. 3), they revealed a multi-layered cell envelope architecture reminiscent of typical Gram-negative bacteria with a relatively thin layer of peptidoglycan ($\sim 10\text{ nm}$). A cytoplasmic membrane (CM) and an S-layer, forming a continuous mesh on the surface of cells, were clearly visible (Fig. 3). An outer layer (OL), similar in appearance to the CM was observed right underneath the S-layer, suggestive of

a putative second membrane. A hidden Markov model search for outer membrane-related genes homologous to those of enterobacteria and *Negativicutes* failed to reveal any in the genomes of CCRI-22567^T and CCRI-24246 (supplementary results). Therefore, characterization of this OL will require further investigations to determine its presence and composition in members of *Peptostreptococcaceae*. The genome annotation of CCRI-22567^T revealed the presence of one gene annotated as S-layer protein and five S-layer homology domain-containing proteins. Additional ultrastructural features include the presence of flagella, flagellar motors and chemoreceptor arrays (Fig. 3b, d and e). Granules were also observed using cryo-ET (Fig. 3c), those are likely to be storage granules. Genome annotation revealed the presence of *glgA*, *glgB* and *glgD* genes known to code for glycogen storage granules as well as genes coding for polyphosphate storage *ppk1* and *ppx* in both strains CCRI-22567^T and CCRI-24246.

Special-potency discs (Oxoid) containing colistin ($10\mu\text{g}$), kanamycin ($1000\mu\text{g}$) and vancomycin ($5\mu\text{g}$) were used to group the anaerobic bacteria, as recommended in the Wadsworth manual [29]. Analysis using special potency discs showed that growth of strains CCRI-22567^T and CCRI-24246 was resistant to $10\mu\text{g}$ colistin and susceptible to $5\mu\text{g}$ vancomycin and $1000\mu\text{g}$ kanamycin.

The two novel isolates were examined (according to the manufacturer's instructions) using Rapid ID 32A, RapID ANA II and API ZYM systems (bioMérieux). Under the rapid ID 32A system, strains CCRI-22567^T and CCRI-24246 gave positive results for arginine dihydrolase, β -glucuronidase, indole production, arginine arylamidase, while it was negative for urease, β -galactosidase, β -galactosidase-6-phosphatase, α -glucosidase, β -glucosidase, α -arabinosidase, *N*-acetyl- β -glucosaminidase, mannose fermentation, raffinose fermentation, glutamic decarboxylase acid, α -fucosidase, nitrate reduction, proline arylamidase, leucyl glycine arylamidase, phenylalanine arylamidase, pyroglutamic acid arylamidase, tyrosine arylamidase, alanine arylamidase, glycine arylamidase and serine arylamidase. Discordant results were observed for α -galactosidase, leucine arylamidase, and histidine arylamidase where strain CCRI-22567^T gave positive results while strain CCRI-24246 gave negative results. Discordant results were also observed for alkaline phosphatase and glutamyl glutamic acid arylamidase where strain CCRI-22567^T gave negative results while strain CCRI-24246 gave positive results. Numerical profiles obtained using the Rapid ID 32A system were 6040212001 for strain CCRI-22567^T and 2040610000 for strain CCRI-24246, respectively.

Under the RapID ANA II system (Remel), both strains CCRI-22567^T and CCRI-24246 gave positive results for arginine- β -naphthylamide and indole production from tryptophan, while they were negative for urea, *p*-nitrophenyl- β -D-disaccharide, *p*-nitrophenyl- α -L-arabinoside, α -nitrophenyl- β -D-galactoside, *p*-nitrophenyl- α -D-glucoside, *p*-nitrophenyl- β -D-glucoside, *p*-nitrophenyl- α -D-galactoside, *p*-nitrophenyl- α -L-fucoside, *p*-nitrophenyl-*N*-acetyl- β -D-glucosaminide, *p*-nitrophenylphosphate, leucyl-glycine- β -naphthylamide,

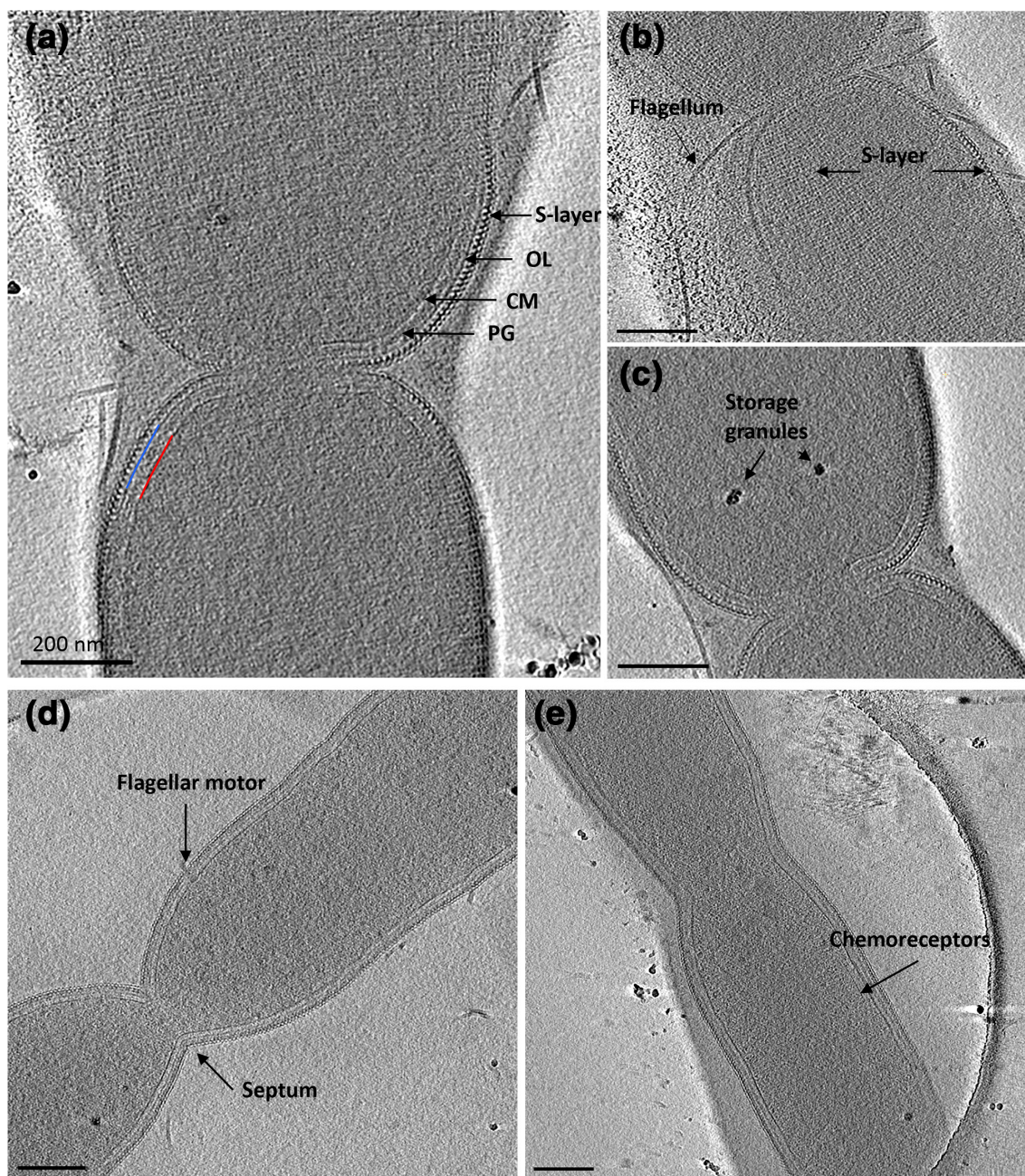


Fig. 3. Ultrastructure of *Criibacterium bergeronii* CCRI-22567 ($n=7$) revealed by cryo electron tomography. (a) A 20 nm thick tomographic slice reveals the presence of a cytoplasmic membrane (CM), thin peptidoglycan (PG) layer, an outer layer (OL) and a proteinaceous S-layer. The CM is also depicted in red while the OL is shown in blue. (b-e) Structural details reveal the presence of flagella and flagellar motors, chemoreceptor arrays and dense storage granules (likely polyphosphate). Scale bar, 200 nm.

glycine- β -naphthylamide, proline- β -naphthylamide, phenylalanine- β -naphthylamide, serine- β -naphthylamide and pyrroindonyl- β -naphthylamide. Microcode profiles obtained using the RapID ANA II system were 000044 for both strains.

Under the API ZYM system, both strains CCRI-22567^T and CCRI-24246 gave positive results for phosphatase acid, naphthol-AS-BI-phosphohydrolase and β -glucuronidase,

while they were negative for alkaline phosphatase, esterase, esterase lipase, lipase, valine arylamidase, cystine arylamidase, trypsin, α -chymotrypsin, β -galactosidase, α -glucosidase, β -glucosidase, *N*-acetyl- β -glucosaminidase, α -mannosidase and α -fucosidase. Discordant results were observed for leucine arylamidase and α -galactosidase where strain CCRI-22567^T gave positive results while strain CCRI-24246 gave negative results.

For the determination of the whole-cell fatty acid (CFA) composition, strains were extracted after 16 h growth at 35 °C in pre-reduced peptone–yeast–glucose (PYG) broth (Anaerobe Systems), using the Sherlock System (MIDI) as described previously [33] except that operating system version 4.5 was used. CFAs were identified using the peak-naming table found in MIDI (Moore 6.0). CFAs for strain CCRI-22567^T were deemed to be qualitatively similar to other genera found in the families *Clostridiaceae* and *Peptostreptococcaceae* [34]. The main fatty acids in the cell membrane of strain CCRI-22567^T comprised C_{18:1} ω9c and C_{18:1} ω9c DMA. The presence of a large quantity of these fatty acids appears to be specific to the type strain of this species. Furthermore, it appears that CFA C_{18:1} ω9c DMA (ECL 18.224) is not found in other taxa of the family *Peptostreptococcaceae* tested in this study (Table S1).

Scores from in-house generated mass spectrometry profiles [35] made using the two strains suggested that MALDI-TOF could be used to unambiguously identify *C. bergeronii* and had no significant match (<1.7) towards all other entries in the Bruker Biotyper library (version 7854). Genomic, morphological and chemotaxonomic characteristics of the 16 genera of *Peptostreptococcaceae*, including *Criibacterium*, are summarized in Table S2 [11, 36].

On the basis of the data from the polyphasic taxonomic analysis presented here, strain CCRI-22567^T is considered to represent a new genus and species within the family *Peptostreptococcaceae*, for which the name *Criibacterium bergeronii* gen. nov., sp. nov., is proposed.

DESCRIPTION OF *CRIBACTERIUM* GEN. NOV.

Criibacterium (Cri.i.bac.te'ri.um. N.L. neut. n. *bacterium* a small rod; N.L. neut. n. *Criibacterium* a small rod isolated at CRI-Centre de Recherche en Infectiologie.)

Members of the genus grow under anaerobic conditions. Cells are Gram-variable with a cytoplasmic membrane, a relatively thin layer of peptidoglycan, an additional outer layer of yet-uncharacterized composition and an S-layer (ultrastructural data from type strain only), rod-shaped with rounded ends, and non-spore-forming. Cells are motile. Colonies on blood agar are grey-translucent, circular with entire edges, smooth and raised. The predominant cellular fatty acids detected are C_{18:1} ω9c and C_{18:1} ω9c DMA. Phylogenetically, this genus is a member of the *Peptostreptococcaceae* and the type species is *Criibacterium bergeronii*.

DESCRIPTION OF *CRIBACTERIUM BERGERONII* SP. NOV.

Criibacterium bergeronii [ber.ge.ro'ni.i. N.L. masc. gen. n. *bergeronii* of Bergeron, named after Professor Michel G. Bergeron (CM, OQ, M.D., FRCPC, FCAHS), infectious diseases specialist, serial entrepreneur, founder, and former director of Centre de Recherche en Infectiologie de l'Université Laval in Québec City (Québec), Canada].

C. bergeronii sp. nov. has the following characteristics in addition to those given for the genus. Cells are ~630 nm wide by ~1750 nm long and occur isolated or in pairs. Growth occurs on TSA with 5% sheep blood under anaerobic conditions at 35 °C, but no growth occurs following subculture in thioglycolate and tryptic soy broths. After 72 h incubation under anaerobic conditions colonies are flat, translucent and <1.0 mm in diameter. Cells grow at 30–42 °C (optimum, 42 °C), at pH 5.5–8.5 (optimum, pH 7.0–7.5) and in the presence of 0–1.5% (w/v) NaCl (optimally at 0.5% NaCl). Using Rapid ID 32A, RapID ANA II and API ZYM systems, positive results are obtained for L-arginine, β-glucuronidase, indole production, arginine arylamidase, phosphatase acid, naphthol-AS-BI-phosphohydrolase and arginine-β-naphthylamide. Negative results are obtained for urea, β-galactosidase, β-galactosidase-6-phosphatase, α-glucosidase, β-glucosidase, α-arabinosidase, N-acetyl-β-glucosaminidase, mannose fermentation, raffinose fermentation, glutamic decarboxylase acid, α-fucosidase, α-mannosidase, nitrate reduction, proline arylamidase, leucyl glycine arylamidase, pyroglutamic arylamidase acid, tyrosine arylamidase, alanine arylamidase, cysteine arylamidase, glycine arylamidase, valine arylamidase, serine arylamidase, *p*-nitrophenyl-β-D-disaccharide, *p*-nitrophenyl-α-L-arabinoside, α-nitrophenyl-β-D-galactoside, *p*-nitrophenyl-α-D-glucoside, *p*-nitrophenyl-α-D-galactoside, *p*-nitrophenyl-α-L-fucoside, *p*-nitrophenyl-N-acetyl-β-D-glucosaminide, leucyl-glycine-β-naphthylamide, glycine-β-naphthylamide, proline-β-naphthylamide, phenylalanine-β-naphthylamide and pyrroldonyl-β-naphthylamide. The *C. bergeronii* type strain G+C content is 33.8mol%, which is consistent with the other members of family *Peptostreptococcaceae* (Table S2) [11, 36]

The type strain, CCRI-22567^T (=LMG 31278^T=DSM 107614^T=CCUG 72594^T), was isolated from a woman diagnosed with bacterial vaginosis while strain CCRI-24246 was isolated from pus of the left armpit of an 81-year-old woman. The GenBank/EMBL/DDBJ accession numbers for the 16S rRNA gene and draft whole genome sequences of strain CCRI-22567^T are NR_164617.1 and MBEW00000000.2, respectively.

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Conflicts of interest

The authors declare that there are no conflicts of interest.

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