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# De novo genome assembly resolving repetitive structures enables genomic analysis of 35 European *Mycoplasma bovis* strains

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## Abstract

*Mycoplasma bovis* (*M.* *bovis*), the agent of mastitis, pneumonia, and arthritis in cattle, harbors a small genome of approximately 1 Mbp. Combining data from Illumina and Nanopore technologies, we sequenced and assembled the genomes of 35 European strains and isolate DL422\_88 from Cuba. While the high proportion of repetitive structures in *M. bovis* genomes represent a particular challenge, implementation of our own pipeline *MycoVista* (available on GitHub [www.github.com/sandraTriebel/mycovista](https://www.github.com/sandraTriebel/mycovista)) in a hybrid approach enabled contiguous assembly of the genomes and, consequently, improved annotation rates considerably. To put our European strain panel in a global context, we analyzed the new genome sequences together with 175 genome assemblies from public databases. Construction of a phylogenetic tree based on core genes of these 219 strains revealed a clustering pattern according to geographical origin, with European isolates positioned on clades 4 and 5. Genomic data allowing assignment of strains to tissue specificity or certain disease manifestations could not be identified. Seven strains isolated from cattle with systemic circular condition (SCC), still a largely unknown manifestation of *M. bovis* disease, were located on both clades 4 and 5. Pairwise association analysis revealed 108 genomic elements associated with a particular clade of the phylogenetic tree. Further analyzing these hits, 25 genes are functionally annotated and could be linked to a *M. bovis* protein, e.g. various proteases and nucleases, as well as ten variable surface lipoproteins (Vsps) and other surface proteins. These clade-specific genes could serve as useful markers in epidemiological and clinical surveys.

**Keywords** *Mycoplasma bovis*, de novo genome assembly, Oxford Nanopore Technologies, Illumina

## Introduction

Infections of the bovine pathogen *Mycoplasma bovis* (*M.* *bovis*) are of considerable economic importance due to their negative impact on animal health and production yields [1]. Pneumonia and mastitis are the most prominent clinical manifestations in cattle, but arthritis, genital disorders, and keratoconjunctivitis can also be caused by this agent [2–4]. Occasionally, *M. bovis* infections presenting as purulent fibrinous pleuropneumonia with sequestering were observed in adult cattle and had to be differentiated from other *Mycoplasma bovis* diseases, such as Contagious Bovine Pleuropneumonia. Recently, an

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increase of acute *M. bovis* infections leading to mastitis in medium-sized and large dairy herds was observed in the eastern and northern federal states of Germany. In some cases, a different clinical picture compared to previous outbreaks was noticed since, in addition to mastitis, affected animals showed symptoms of circulatory involvement manifesting as massive edema in the chest, abdomen, and leg area, similar to allergic reactions (see Fig. S1) [5, 6]. We will refer to this disease complex as Systemic Circular Condition (SCC).

While many aspects of *M. bovis* pathogenesis are still not fully understood, the agent's capability of subverting host immune response through surface antigen variation was characterized in several studies [7–10]. The central role in this process is played by members of a family of lipoproteins designated variable surface proteins (Vsps), which consist of a conserved N-terminal domain for membrane insertion and lipoprotein processing and a large stretch of repetitive tandem domains comprising up to 80 % of the entire Vsp molecule. As a result of spontaneous and non-coordinated deletions, insertions, and rearrangements in the *vsp* genomic locus, translated Vsps undergo variations in phase (on/off switching), size (varying number of tandem repeats), and/or surface presentation at high frequency [8].

In this context, sequencing and comparative genomics can be important tools to identify genetic features correlating with strain properties and/or disease symptoms [11]. High-quality genomes with precise and, if possible, complete annotations are needed to perform meaningful comparative studies. However, despite their small size (~ 1 Mbp), the assembly of *M. bovis* genomes is challenging due to the high rate of the above-mentioned repetitive structures. In the past, using state-of-the-art tools to assemble Illumina reads resulted in fragmented genomes because repetitive regions are not efficiently covered using short reads. Nevertheless, despite the lower contiguity of such short-read assemblies, the high sequence accuracy of short reads and the associated accurate annotation of open reading frames (ORFs) is essential for further downstream analysis. Currently, 588 assemblies of *M. bovis* strains are available in the NCBI database (Jan 16, 2023), of which 51 assemblies are marked as complete. Nine of them lack GenBank and/or RefSeq annotation entries. The current NCBI reference genome of *M. bovis* (strain 8790, GCF\_005061465.1) consists of 17 contigs and is incomplete. Sequencing via nanopores, e.g. the Oxford Nanopore Technologies (ONT) MinION device, has great potential to enable genome assemblies of higher contiguity since longer reads facilitate coverage of complex genomic regions [12, 13]. Recent studies have shown that the use of long and short reads is a suitable strategy for assembling bacterial genomes [14].

In the present study, we used a hybrid assembly approach combining the advantages of Illumina and Nanopore sequencing to obtain high-quality genome sequences of 36 *M. bovis* strains. The strain panel contains seven isolates associated with the SCC disease complex in Germany from the last ten years, strains from different geographical regions and from animals with other symptoms, as well as previous isolates. Genome sequence data were processed using our own *MycoVista* pipeline ([www.github.com/sandraTriebel/mycovista](http://www.github.com/sandraTriebel/mycovista)), which also includes improved annotation, to elucidate epidemiological and phylogenetic relationships.

## Material and methods

### Strains

The aim was to cover a broad spectrum of *M. bovis* field strains in the study, including isolates from animals with severe symptoms. Therefore, we selected 36 isolates from different geographical regions of Germany, other countries, and different tissue representing various disease manifestations. All German strains isolated in 2014 and later were collected especially for this study by diagnostic laboratories of several federal states and animal health services and incorporated in the strain collection of the Friedrich-Loeffler Institute, Germany. For comparison, the Cuban strain DL422\_88 was included. The basic characteristics of the included strains are given in Table 1. Most isolates were obtained as grown cultures. Further cultivation was done with a modified Hayflick medium containing 20 % horse serum [15] or with commercially available liquid medium containing a phenol-red pH indicator (Mycoplasma Experience Ltd, Bletchingley, UK) at 37 °C and 8 % CO<sub>2</sub> under static conditions for 2 to 4 days (until color change of the liquid broth to orange or yellow was observed). Only in one case, we obtained a tissue sample from the lung of a calf (16DD0054), which was cultured in liquid broth culture (Mycoplasma Experience Ltd, UK) containing Penicillin G (1000 IU/ml, WDT, Garbsen, Germany) to suppress other bacteria. Liquid cultures were stored in a -80 °C freezer and solid agar plates in at 4 °C.

### Description of the disease complex

After calving, the Systemic Circular Condition (SCC) affected cows in medium-sized and large dairy herds. Animals suffered from painful, swollen joints that partially ruptured, edema mainly in the legs and chest area, shortness of breath, and general nervousness while excreting glassy-serous saliva, usually without any fever. Signs of severe mastitis also typically occurred, with udders hardened to a rubbery consistency, milk becoming creamy and sandy and milk yield dropping sharply. Affected udder quarters either became permanently

**Table 1** Basic parameters of the *M. bovis* strains included in this study. CU – Cuba; DE – Germany; PL – Poland; IE – Ireland; BB – Brandenburg; BW – Baden-Württemberg; BY – Bavaria; HE – Hesse; SL – Saarland; MV – Mecklenburg Western Pomerania; NI – Lower Saxony; NRW – Northrhine-Westphalia; RP – Rhineland Palatinate; SN – Saxony; ST – Saxony-Anhalt; TH – Thuringia; SCC – Systemic Circulatory Condition

Strain	Source	Pathology	Country, State	Year of isolation	Supplier
11DD0261	Milk, cow	unknown	DE, BY	2011	Landesuntersuchungsamt Oberschleissheim
13DD0918	Nasal swab, cow	SCC	DE, BW	2013	Rindergesundheitsdienst Fellbach (J. Mandl)
14DD0147	Nasal swab, cow	unknown	DE, BW	2014	A1: Staatl. Tierärztl. Untersuchungsamt Aulendorf (I. Holst)
14DD0148	Joint, bull	unknown	DE, BW	2014	A1
14DD0156	Nasal swab, calf	unknown	DE, BW	2014	A1
14DD0475	Milk	Mastitis	DE, NI	2014	W: Milchherden-Betreuungs- & Forschungsgesellschaft Wunstorf (M. Entorf)
15DD0123	Milk	SCC	DE, MV	2015	SD: B. Schwagerick (via Dr. Felgenträger & Co. Dessau-Rosslau, T. Forbrig)
15DD0140	Milk	Mastitis, Arthritis, Abortion in herd	DE, BB	2015	LKV: Landeskontrollverband Berlin-Brandenburg Waldsieversdorf (U. Nebel)
15DD0141	Milk	Mastitis, Arthritis, Abortion in herd	DE, BB	2015	LKV
15DD0160	Lung, calf	Bronchopneumonia	DE, ST	2015	St: Landesamt für Verbraucherschutz Stendal (A. Schliephake)
15DD0161	Lung, cow	Bronchopneumonia	DE, HE	2015	Landesbetrieb Hessisches Landeslabor, Gießen (T. Eisenberg)
15DD0163	Lung, calf	Bronchopneumonia	DE, ST	2015	St
15DD0164	Milk	unknown	DE, TH	2015	J: Tiergesundheitsdienst, Jena (K. Klengel)
15DD0165	Milk	none	DE, BB	2015	LKV
15DD0207	Milk	SCC	DE, MV	2015	SD
15DD0210	Milk	(poor milk hygiene)	DE, BB	2015	LKV
15DD0218	Milk	(elevated cell count in milk)	DE, TH	2018	J
15DD0228	Milk	Mastitis	DE, BB	2015	LKV
15DD0233	Milk	Mastitis	DE, MV	2015	SG: B. Schwagerick (via MQD-Qualitätssprüfungs- & Dienstleistungsgesellschaft Mecklenburg-Vorpommern, Güstrow)
15DD0234	Lung, cow	SCC	DE, MV	2015	SR: B. Schwagerick (via Landesamt für Landwirtschaft, Lebensmittelsicherheit & Fischerei, Rostock)
15DD0238	Lung or joint, dead cow	SCC	DE, MV	2015	SR
15DD0261	Nasal swab, calf	unknown	DE, BW	2015	A2: Staatl. Tierärztl. Untersuchungsamt Aulendorf (S. Bracknies)
15DD0263	Joint puncture, cattle	Arthritis	DE, BW	2015	A2
15DL0124	Milk	SCC	DE, MV	2016	B. Schwagerick (via IDT Biologika GmbH, Dessau-Rosslau)
16DD0001	Milk	unknown	DE, SN	2016	Landesuntersuchungsanstalt f. Gesundheits- & Veterinärwesen Sachsen, Dresden (C. Kruppe)
16DD0054	Pulmonary tissue, calf	Bronchopneumonia	DE, NRW	2016	Chemisches u. Veterinäruntersuchungsamt Münsterland-Emscher-Lippe (A. Nagel)
16DD0100	Milk	SCC	DE, MV	2016	SG
16DD0186	Milk	Mastitis, Pneumonia	DE, MV	2016	SG
16DL0615	Milk	unknown	DE, NI	2016	W
17DD0007	Milk	unknown	DE, NI	2017	W

**Table 1** (continued)

Strain	Source	Pathology	Country, State	Year of isolation	Supplier
17DD0020	Pulmonary tissue, cow	Bronchopneumonia	DE, RP	2017	Landesuntersuchungsamt Koblenz (A. König-Mozes)
15DD0240	Nasal swab, calf	unknown	PL	2014	PL: Dept. Cattle and Sheep Diseases, The National Veterinary Research Institute, Pulawy, Poland (E. Scacawa)
15DD0249	Nasal swab, calf	unknown	PL	2014	PL
15DD0250	Nasal swab, calf	unknown	PL	2014	PL
DL422_88	Lung, calf	Pneumonia	CU	1980	Centro Nacional de Sanidad Agropecuaria, S. José de las Lajas, Cuba (A. Fernández)
DL81_99	Milk	Mastitis	IE	1999	Bacteriology Branch Veterinary Sciences Division, Belfast (H. Ball)

atrophic or their milk production was fully restored after recovery. In the course of the disease, milk cell counts were significantly elevated. Within a few days, some animals could not stand up and finally died from cardiovascular failure. Treatment with antibiotics had no effect while pain-relieving, anti-inflammatory therapy brought some improvement. Feed consumption was only temporarily reduced. Usually, several animals in the herd fell ill one after another with the same symptoms. For example, on a dairy farm with 450 cows, 60 animals showed the same systemic symptoms, of which 20 died. *M. bovis* was detected in up to 50 % of the diseased cows in a herd. Using commercial ELISA tests, specific antibodies against *M. bovis* were detected in all diseased animals.

#### DNA extraction & sequencing

DNA extraction for sequencing with Illumina was done using High Pure PCR Template Preparation Kit from Roche (Mannheim, Germany) according to the manufacturer's instructions. DNA extraction for Nanopore MinION sequencing was performed using phenol-chloroform extraction as described by Sambrook and Russell [16]. The final DNA extracts were recovered in 50 µl buffer. For the approval of quality and quantity, the Nanodrop spectrophotometer (Thermo Fisher Scientific, Madison, USA) and QUBIT 2.0 fluorometer (Life Technologies Holdings PTE Ltd, Singapore), as well as agarose gel electrophoresis, were used. Illumina sequencing. Genomic DNA (2–10 µg) of each isolate was sent to GATC/Eurofins Genomics (Konstanz, Germany) for genomic library preparation and Illumina MiSeq (2 x 125 bp paired-end or 2 x 150 bp paired-end sequencing) with 5 Million read pairs resulting in an average coverage of around 750x. Raw sequencing data were quality-controlled using FastQC v0.11.8. We used four different MinION flow cells (R9.4.1) numbered 1–4 in Table S1. Nanopore sequencing. Library preparations

were done using the 1D genomic DNA by ligation kits (SQK-LSK 108 and SQK-LSK 109) and the native barcoding expansion kits (EXP-NBD103, EXP-NBD104, and EXP-NBD114). In response to an update by ONT, we transitioned to the newer ligation kit SQK-LSK 109 for our nanopore sequencing experiments since the second sequencing run (see Table S1 column F). Briefly, size selection and DNA clean-up were performed using Agencourt AMPure XP beads (Beckman Coulter GmbH, Krefeld, Germany) at a ratio of 1:1 (w:v) before library preparation. Potential nicks in DNA and DNA ends were repaired in a combined step using NEB Next FFPE DNA Repair Mix and NEB Next Ultra II End repair/dA-tailing Module (NewEngland Biolabs, Ipswich, USA) by tripling the incubation time to 15 min at 50 °C and 15 min at 65 °C. The ligation of sequencing adapters followed a subsequent second AMPure bead purification onto prepared ends and a third clean-up step with AMPure beads. Additional barcoding and clean-up steps were performed before adapter ligation. Sequencing buffer and loading beads were added to the library. At the start of sequencing, an initial quality check of the flow cells showed 1571, 1379, 1761, and 1236 active pores. Genomic DNA samples used for loading comprised around 50–150 ng per strain (measured by Qubit 4 Fluorometer; ThermoFisher Scientific, Waltham, USA). The sequencing ran for 48 hrs using the MinKNOW software versions 2.2, 18.12.4, 19.05, 19.10. MinION signals are base-called and demultiplexed using guppy v6.1.2 model R9.4.1 (only available to ONT customers <https://community.nanoporetech.com>).

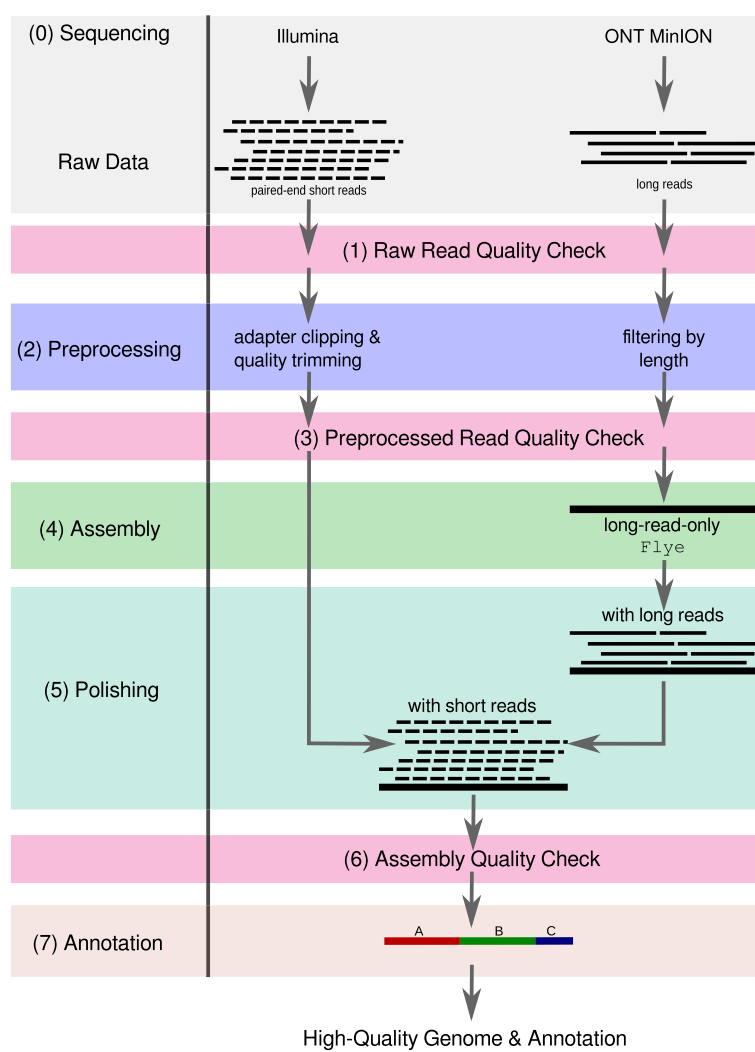
#### Hybrid *de novo* assembly pipeline

We developed Mycovista, a pipeline to assemble highly repetitive bacterial genomes such as *M. bovis* in a hybrid or long-read-only sequencing approach. Our pipeline follows the state-of-the-art *de novo* genome assembly

workflow by incorporating preprocessing of input reads, assembly, polishing, and genome annotation [14]. As input, demultiplexed (nanopore) long reads are required. Illumina paired-end short reads can be used in the hybrid assembly mode. The rapid development of ONT allows now the generation of high-quality assemblies by long read-only data, which is why our pipeline has short-read data as an optional input [17–19]. Mycovista returns the assembled genome, a suitable annotation, and general assembly statistics. The pipeline is automated using the workflow management system `snakemake v7.3.8`

[20] in combination with `conda v22.9.0` for reproducible results via stored tool versions in corresponding environment files [21]. All code is publicly available on GitHub ([www.github.com/sandraTriebel/mycovista](http://www.github.com/sandraTriebel/mycovista)). We ran Mycovista in release version 1.0.

The pipeline consists of seven steps, see Fig. 1: Before using Mycovista, sequencing and basecalling has to be done by the user to provide the input reads. (0) *Sequencing and Basecalling*: The assembly pipeline can be started in two modes: *long* requiring only ONT long reads and *hybrid* where ONT long and Illumina paired-end short



**Fig. 1** Mycovista – a *de novo* assembly pipeline. Mycovista can be used in two assembly modes: long and hybrid, requiring only long reads or additionally paired-end short reads, respectively. (0) Basecalled long-read sequencing data (e.g. from ONT) is required. Illumina paired-end short reads can be used as additional input. (1) A raw read quality check (FastQC and Nanoplot) is followed by (2) preprocessing of the input reads (Filterlong, fastp, Trimmomatic) which are then (3) checked regarding their quality. (4) Then, a long-read-only assembly is generated by Flye. (5) Afterwards, the contigs are polished with the preprocessed long reads in several steps (Racon, minimap2, medaka). The assembly can be further postprocessed with short reads. Finally, the (6) quality of the final assembly is assessed (QUAST) followed by a (7) gene annotation (Prokka)

reads are needed. MinION signals need to be basecalled and demultiplexed using *guppy*. We applied *guppy* v6.1.2 model R9.4.1. using the following parameters: `-c dna_r9.4.1_450bps_sup.cfg, --trim_strategy dna, --disable_pings, --disable_qscore_filtering, --calib_detect, --barcode_kits <barcode kits>`. (1) *Raw Read Quality Check*: FastQC v0.11.8 [22] is incorporated for a first quality check of the raw Illumina data. The ONT raw read quality is visualized with Nanoplot v1.41.0 [23]. (2) *Preprocessing*: Only reads longer than 1,000 nt are considered for the assembly step (filtered via *Filtlong* v0.2.0 [24]), as *M. bovis* genomes are known to contain repeats. Illumina raw reads are preprocessed with *fastp* v0.20.0 [25] for adapter clipping and *Trimmomatic* v0.39 [26] (sliding window size 4, Phred score quality cut-off of 28, minimum read length 20) to perform quality trimming. However, using *Trimmomatic* alone is not sufficient to remove all adapter sequences, and thus *fastp* is included. (3) *Preprocessed Read Quality Check*: FastQC and Nanoplot are again applied to examine the quality of preprocessed reads and to make a comparison to step (2) possible. (4) *Assembly*: The filtered long reads are assembled with *Flye* v2.6 [27]. Optional Illumina reads are not used for assembly but for polishing. (5) *Polishing*: Post-processing of the long-read assemblies consisted of three steps: (i) four polishing runs with *Racon* v1.3.2 [28] in combination with *minimap2* v2.17 [29] using long reads followed by (ii) one long read-based polishing with *medaka* v0.11.4 [30] and finally (iii) four polishing runs with *Racon* and *minimap2* using short reads. (6) *Assembly Quality Check*: Finally, the quality of the so-produced hybrid assemblies is validated by *QUAST* v5.0.2 [31]. (7) *Annotation*: All assemblies are annotated using *Prokka* v1.14.5 [32] with the parameter `--gcode 4` to use the required codon table matching characteristics of *M. bovis*.

### Genome synteny analysis

We used *Mauve* v1.2.0 [33] to detect recombinations in the genome assemblies, such as gene loss, duplication, rearrangement, and horizontal transfer. In order to make our *M. bovis* genome assembly panel comparable regarding genome rearrangements, we designated the first base of the gene *dnaA* to be the first base of the genome as described in the study by Mackiewicz et al. [34].

### Pangenome analysis and virulence genes

We used *PPanGGOLiN* v1.2.74 [35] for pangenome analysis of our *M. bovis* genome assembly panel. *PPanGGOLiN* generates a core gene set based on a

Partitioned Pangenome Graph (PPG), which integrates information about protein-coding genes and their genomic neighborhood. The input genomes are annotated by *ppanggolin* *annotate*. Their specific genetic code 4 for *Mycoplasmopsis* was set with `--translation_table 4`. Subsequently, clustering was performed using *ppanggolin* *cluster*, followed by the graph construction and partitioning (*ppanggolin* *graph*, *ppanggolin* *partition*). The core genes of each genome were aligned using *ppanggolin* *msa* with the parameters `--source dna --translation_table 4` and concatenated in one multiple sequence alignment [35], which serves as input for the subsequent reconstruction of the phylogenetic tree. After generating our 36 assemblies, we compared our data with the study by Yair et al. [36], which included 175 assemblies (whole-genome shotgun project PRJNA564939), as well as seven NCBI RefSeq genomes of *M. bovis* (PG45 NC\_014760.1, Hubei-1 NC\_015725.1, HB0801 NC\_018077.1, CQ-W70 NZ\_CP005933.1, 08M NZ\_CP019639.1, Ningxia-1 NZ\_CP023663.1, JF4278 NZ\_LT578453.1) [37–41] and one genome provided by the NCBI (NM2012 CP011348.1).

To compare the gene composition of the 35 European strains concerning phylogenetic clustering, we performed statistical association tests (Fisher's exact test) for the 35 strains using an R script [42] available in our *Mycovista* GitHub ([www.github.com/sandraTriebel/mycovista/tree/master/scripts/gwas.R](http://www.github.com/sandraTriebel/mycovista/tree/master/scripts/gwas.R)). Moreover, we focused on potential virulence factors (such as *Vsps*) or genes important for the life cycle [10, 43–45]. We listed the presence of these genes based on reference-based annotations. Only genes that were assigned with the correct name for the potential virulence factors were taken into account. Multiple occurrences of a gene are possible due to gene duplication or fragmentation. It is noteworthy that *Prokka* could not annotate *vsp* genes in our assemblies without a reference genome. This may be due to the absence of those sequence annotations in the database or the very diverse structure of *vsp* genes.

### Phylogenetic tree reconstruction

Phylogenetic trees were reconstructed using *IQ-TREE* v2.0.3 [46, 47] with a generalized time-reversible (GTR) model based on the alignment of the core gene set determined in pangenome analysis. For this purpose, we included *M. agalactiae* PG2 (NC\_009497.1) as an outgroup [48]. The tree reconstruction of our 36 assemblies was done with 1000 bootstrap runs. For comparison among the 219 *M. bovis* strains, we reduced the number of bootstraps to 500. Trees and metadata were visualized using *iTOL* v6.6 [49].



## Results

### Mycovista in comparison to other tools

In this study, we developed and evaluated a comprehensive *de novo* genome assembly pipeline to obtain the most accurate and high-quality genomic sequences. The Mycovista pipeline encompassed various stages, including data preprocessing, assembly, polishing, quality assessment, and annotation. To identify the optimal combination of tools and algorithms, we performed a comparison of different state-of-the-art software and methodologies. We assessed the performance of *de novo* genome assembly tools by analyzing a strain of our dataset (see Table S3). The evaluation criteria included general assembly and annotation statistics. In terms of genome contiguity, Flye [27] performed best with 3 contigs in our benchmark dataset, where the longest contig covered  $\sim 1$  Mb, which is the known genome size of *M. bovis*. Unicycler [50] and Canu [51] failed to assemble the input reads into a contig covering the expected genome size. While Canu resulted in a small number of contigs, assembly with Unicycler resulted in a more fragmented genome. Thus, we integrated Flye into the Mycovista workflow. For genome annotation, we compared Prokka [32] and Bakta [52]. At the time of the comparison, Bakta annotated twice as many CDSs as Bakta, with many fragmented genes (see Table S4). Prokka, on the other hand, was able to annotate about as many CDS as could be expected for *M. bovis*. Therefore, we use Prokka for genome annotation. Based on our analyses, we identified the most efficient and reliable tools, which collectively produced highly contiguous and annotated genome assemblies.

### Hybrid assembly approach generates contiguous and accurate genomes

Illumina sequencing produced an average of 9,359,418 paired-end reads with a length of 125–150 nt, whereas ONT sequencing produced 317,908 reads on average with a length of about 3,726.8 nt (longest read: 461,771 nt), see Table S1 Illumina & ONT Information.

Based on our hybrid approach, Mycovista achieved high genome contiguity: All 36 hybrid assemblies are comprised of one to maximal three contigs (with the notable exception of 16 contigs in 17DD0020) and have an average N50 of 1,050,036. This is a remarkable improvement given the challenges posed by multiple sequence repeats in the genome [53]. The average genome size of the assemblies in this study was 1.063 Mbp, with the largest genome size being 1.166 Mbp (15DD0218) and the smallest being 0.952 Mbp (15DD0240). Out of 36 *M. bovis* strains, 23 were assembled into single contigs representing the full chromosome. No contigs of

plasmid-origin were found by PlasmidFinder [54], which is in line with other studies of *M. bovis* [55]. The GC content per assembly ranged from 29.11 % to 29.45 % which is in accordance with previously published *M. bovis* genomes.

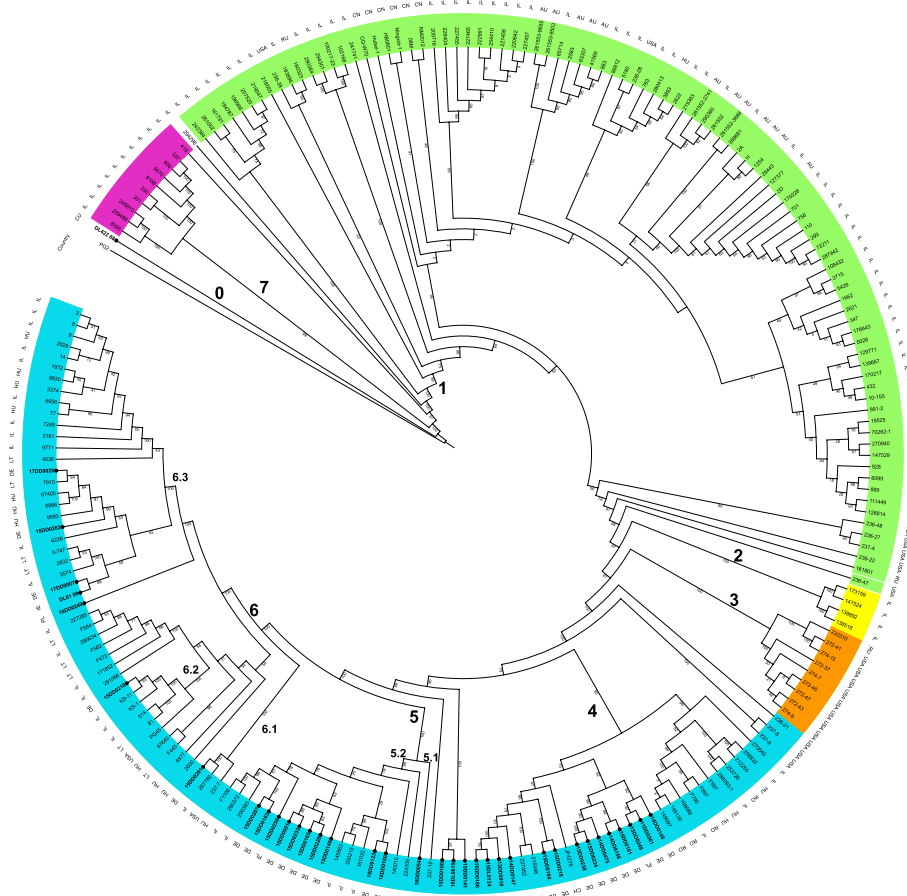
The number of annotated protein-coding genes in our European strain panel varied between 797 and 962, see Table S1, which is in accordance with the expected amount of CDS in *M. bovis* genomes. We observed varying CDS numbers in strains located at different clades of the phylogenetic tree. In the following, we refer to the assemblies according to their association with a particular clade, see Fig. 3, consistent with the literature [36]. On average, clade 4 shows 50 more CDS than clades 5 and 6 (clade 4: 910, clade 5: 860, clade 6: 856). Almost every genome consists of six rRNA operons (16S, 5S, and 23S rRNA). The number of tRNAs is 34, whereas the annotation displays two more tRNAs with codons for Arg (CCT, TCT) in 15DD0240. Strain 15DD0263 shows four extra tRNAs with codons for Thr (GGT, CGT), Lys (CTT), and Trp (TCA), respectively. Once per *Mycoplasma* genome, a tmRNA is present, which is representative of other housekeeping ncRNAs illustrating the general quality of the assemblies. Basic statistics about the ONT sequencing, assembly, and annotation results are given in Table S1.

### Comparison with global strain panel shows clustering according to geographical origin

To put our assemblies in a global context, we compared the 36 genomes with 175 assemblies from the study by Yair et al. [36] and eight complete assemblies of *M. bovis* available from the NCBI database (PG45, Hubei-1, HB0801, CQ-W70, NM2012, 08M, Ningxia-1, JF4278). The phylogenetic tree (see Fig. 2) reconstructed from the core gene set (*M. agalactiae* PG2 included as outgroup) indicates clustering according to geographical origin and is in agreement with the findings of Yair *et al.* With the exception of the Cuban strain DL422\_88 (marked as clade 0), all of our *de novo* assembled genomes are located in the European cluster (clades 4, 5, and 6), which is consistent with their origin.

For a more detailed analysis of our *de novo* assembled strain panel, we reconstructed a phylogenetic tree based on the genomes (Fig. 3). The tree reveals the same clustering pattern as observed in the global phylogenetic tree in Fig. 2. Therefore, we will use the same clade designations 0, 4, 5, and 6 when referring to members of our strain panel.

We observed no enrichment of a single clade for isolation source or geographical parameters. Concerning disease manifestations, it is noteworthy that the seven strains associated with SCC were located in two clades:



**Fig. 2** Phylogeny of 219 *M. bovis* isolates based on the alignment of the core gene set (provided by PPaNGGOLiN). We combined our *de novo* assembled strain panel with 175 assemblies deposited at the whole genome shotgun project PRJNA564939 [36] and eight complete genomes of *M. bovis* available at the NCBI database. *M. agalactiae* PG2 was included as an outgroup. The clustering of genomes reflects the geographical origin of strains, with 35 of our own strain panel situated in clades 4, 5, and 6 of the European cluster (cyan), and Cuban strain DL422\_88 standing outside. Clade numbers and colors are the same as in the SNP-based phylogenetic tree of the paper by Yair et al. [36]

5.2 and 4. As shown in the right-hand part of Fig. 3, the *vsp* gene content varies considerably among the *M. bovis* strains. We used two different sets of annotated variable surface proteins based on strains 8790 (GCF\_005061465.1) and PG45 (NC\_014760.1), respectively, to search for homologs in our strains (see Fig. S2).

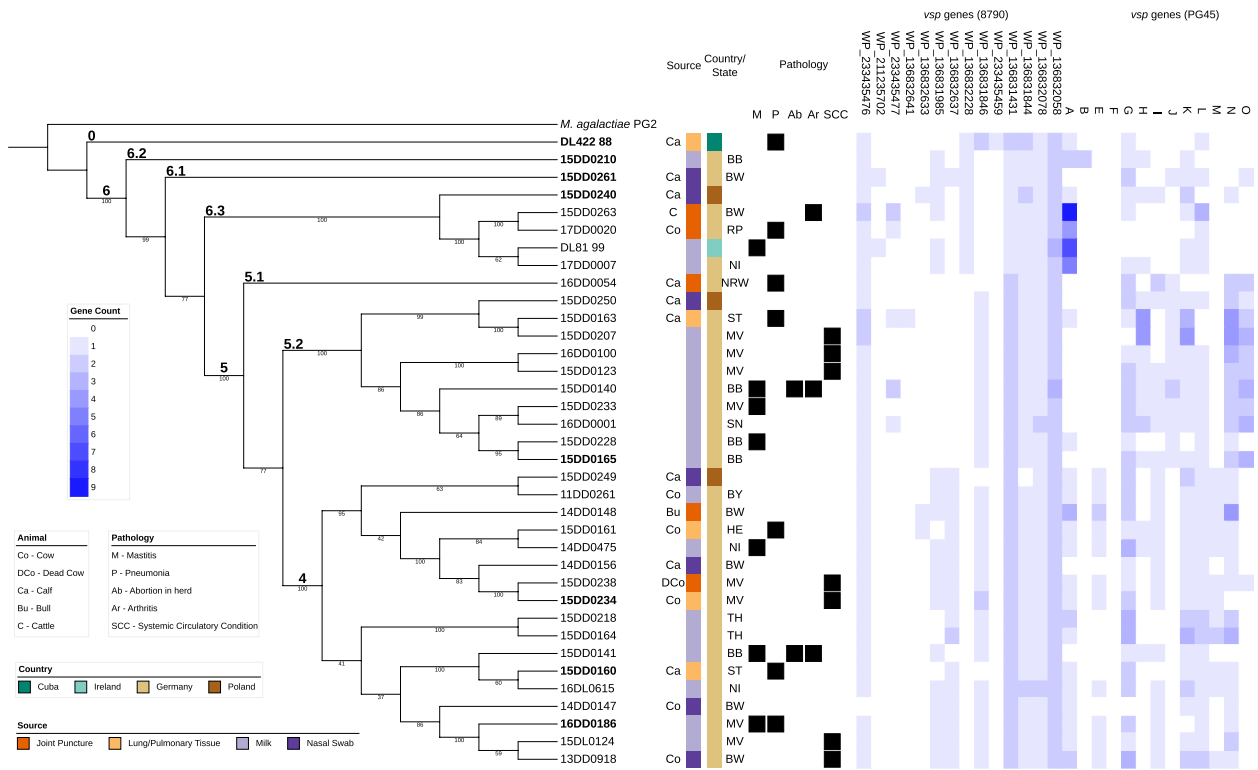
Analysis of the reference-based annotations revealed three general patterns characterizing the occurrence of individual *vsp* genes in the European strain panel: *i*) abundant *vsps* (occurring in more than 30 strains), *ii*) rare *vsps* (occurring in less than 5 strains and not clade-specific or absent altogether), as well as *iii*) those having a preference for one or two clades. Five of the strain 8790-based *vsp* genes were classified as abundant (WP\_233435476, WP\_136831431, WP\_136831844, WP\_136832078, WP\_136832058), four as rare (WP\_211235702, WP\_136832641, WP\_136832633, WP\_233435459), and the following genes showed a clade preference:

WP\_233435477 (for clades 5 and 6), WP\_136831985 (4 and 6), WP\_136832637 (4), WP\_136832228 (6), and WP\_136831846 (5). When looking for *vsp* homologs of strain PG45, *vsps* G, K, and L were present in more than 30 genomes in our strain panel and in all clades, respectively. Members of subclade 6.3 were found to harbor several *vspA* gene copies in their genome. The *vspB* gene was identified only in one strain, while *vspF* was absent in all 36 strains. Among *vsps* with clade preference, *vspE* and *vspM* genes seemed to be confined to clade 4, whereas *vspJ* and *vspO* were only encountered in clade 5. Total count of genes are shown in Table S2.

**Comparison of genome synteny among the European strain panel**

Analysis of the global organization of the 36 *de novo* assembled genomes revealed rearrangements when comparing clusters or subclusters. Notably, there are





**Fig. 3** Phylogenetic tree of our *de novo* assembled strain panel based on the alignment of the core gene set (provided by PpanGOLin). The tree can be divided into three major clusters (4, 5, and 6) and one outlier (Cuban strain DL422\_88). We included *M. agalactiae* PG2 as an outgroup. Biological information such as geographical origin (country, state), source (animal, tissue), pathology, and the presence/absence of *vsp* genes are shown. Strain designations in boldface denote representative sequences of the clusters used for genome synteny analysis shown in Fig. 4. In the right-hand part, *vsp* genes in each strain annotated according to NCBI RefSeq strains 8790 and PG45 are depicted. Clade numbers are labeled according to the study by Yair et al. [36]. Bio Information – Biological Information: Central columns show source of isolation (animal, tissue), geographical origin (country, state), and pathology (if available); Co – Cow; DCo – Dead Cow; Ca – Calf; Bu – Bull C – Cattle; BB – Brandenburg; BW – Baden Wurttemberg; BY – Bavaria; HE – Hesse; SL – Saarland; MV – Mecklenburg Western Pomerania; NI – Lower Saxony; NRW – Northrhine-Westphalia; RP – Rhineland Palatinate; SN – Saxony; ST – Saxony-Anhalt; TH – Thuringia; M – Mastitis; P – Pneumonia/ Bronchopneumonia; Ab – Abortions; Ar – Arthritis; SCC – Systemic Circulatory Condition

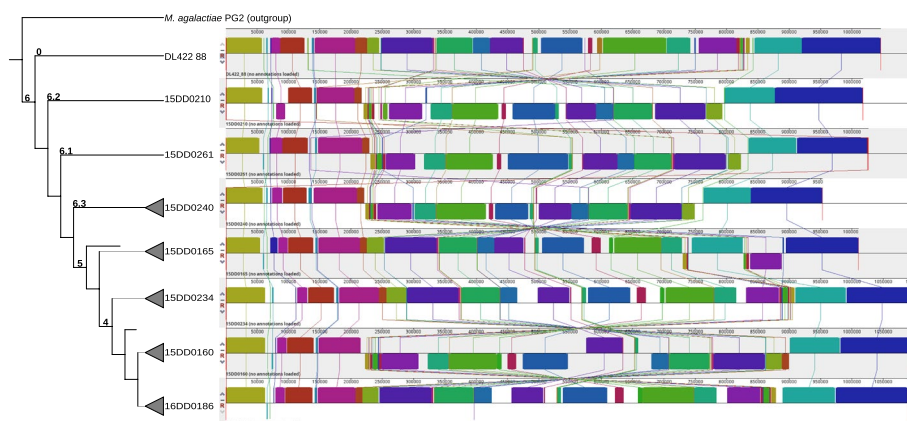
characteristic features that are consistent within the genomes of a cluster, which also indicates that they are not the result of assembly errors. To illustrate clade-specific changes in genome synteny, we selected eight strains representing the major clades for calculating an alignment using Mauve, see Fig S2.

While the first three blocks 5' and the last two blocks 3' are highly conserved among all strains in terms of synteny, the central genomic region, which contains more than 10 locally collinear blocks, displays the highest divergence. As expected, the Cuban strain DL422\_88 is distinct from all other European isolates and has a markedly different synteny compared to the rest of the strain panel. A major characteristic of clade 6 strains (15DD0210, 15DD0261, 15DD0240) is the inverted central part of the genome, in which the blocks appear in the reverse complement orientation relative to the other genomes. This region displays the highest degree of divergence among clade members in comparison to clades 4

and 5 (Fig. S2). Strain 15DD0160 (and two other strains from clade 4, see Fig. S2) also appears to have an inverted central block, similar to the strains of clade 6. In contrast, the remaining representatives of clades 4 (15DD0234 and 16DD0186) and 5 (15DD0165) share the central genomic blocks arranged in the forward orientation, see Fig. 4. There is also variation among individual strains within the clade (see Fig. S2). The genomes of clade 4 are 50,000 to 100,000 bp larger than those of clades 5 and 6, which explains the higher number of CDSs in clade 4 strains. These genomes contain a number of segments depicted as blank spaces that have no homologs in the other strains (Fig. 4 and S2).

**Pangenome analysis reveals cluster-associated genes including virulence factors**

We identified a pangenome of our 36 assemblies consisting of 1,143 gene families, composed of 598 core genes (i.e., genes present in all 36 genomes), 105 shell genes



**Fig. 4** Multiple genome alignment of eight strains representing the major clades using Mauve. Each genome was linearized and normalized, with homologous segments (locally collinear blocks) shown as colored rectangles. Inverted regions are set below those that match the neighboring genome in the forward orientation. Lines collate aligned segments between genomes. The alignment of all 36 assemblies done in this study is shown in Fig. S2

present in ~ 55 % genomes, and 351 cloud genes present at low frequency (~ 8 %). Gene association analysis revealed 108 to be significantly associated with at least one cluster (4, 5, 6), i.e. are either significantly enriched or depleted in the strains of the analyzed cluster (see Fig. S3). Of those genes, 25 could be linked to a gene product or at least a class of products. These clade-specific genes include various enzymes, e.g. proteases and nucleases, as well as ten variable surface lipoproteins and other surface proteins (see Fig. 5). In particular, some *vsp* genes show correlations to phylogenetic clusters. In Fig. 3, we can observe *vsp* WP\_136832228 to be only present in clade 6 and Cuban strain DL422\_88 (clade 0). The genes *vspJ* and *vspO* are only present in clade 5. Genes *vsp* WP\_136832637 and *vspE* are associated with clade 4.

As shown in Fig. 5, genes encoding iron-sulfur cluster carrier protein, tyrosine recombinase XerD, conjugal transfer protein TraE, ATP-dependent zinc metalloprotease FtsH, transglutaminase domain-containing protein, and DNA cytosine methyltransferase seem to be only annotated in genomes of clade 4. Gene *xerC* coding for tyrosine recombinase XerC is annotated three to four times in clades 5 and 6, but only two times in clade 4. We observed these differences in the annotation files as well as in the gene association analysis in which one annotation of *xerC* was detected as absent in clade 4. The gene *vsp* WP\_136831846 is absent in clade 6. Genes *vspN* were annotated multiple times in some genomes. One of them occurred only in clade 5 strains, and another one was only present in clade 4. We observed similar results for *vsp* WP\_136831985, see Fig. 5, which was never annotated in group 5 genomes, but found once in each of clades 4 and 6, respectively. The gene *vsp* WP\_136831431 was annotated twice in clades 4 and 5 and in Cuban strain

DL422\_88 (clade 0), but only once in clade 6 as revealed by Fisher’s exact test.

**Discussion**

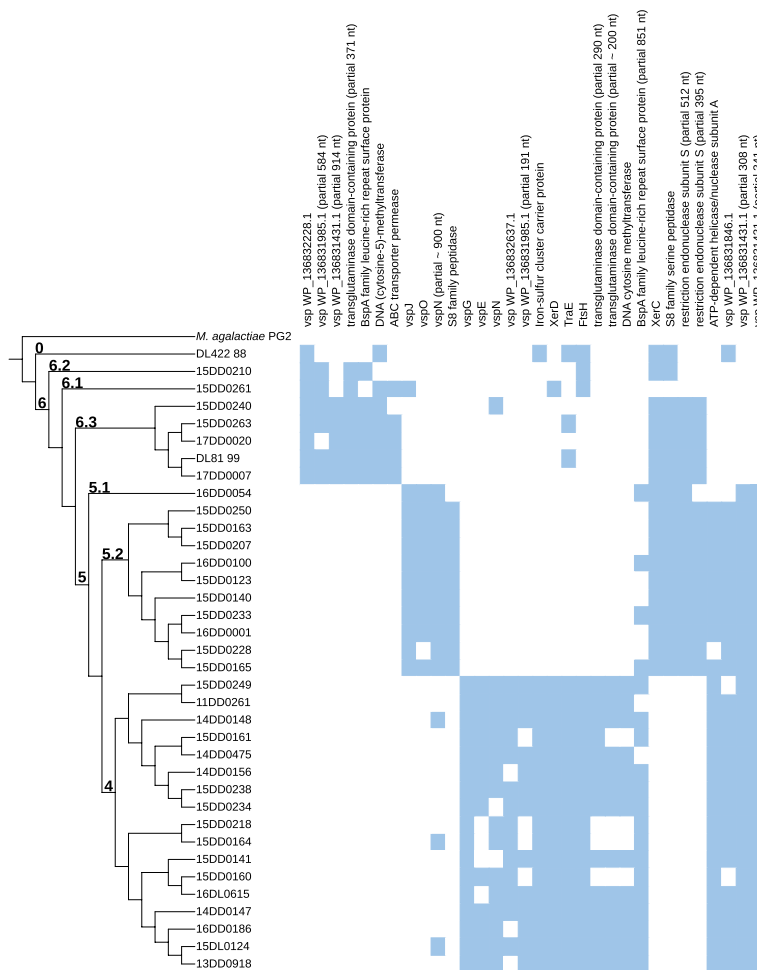
**De novo assembly**

Mycovista assembles bacterial genomes using a hybrid approach to tackle the problem of resolving highly repetitive regions. The use of long reads (ONT used here; in theory PacBio too, but not tested) in the initial assembly is beneficial for generating contiguous genomes, and the polishing steps involving short reads improve the sequence accuracy of the genomes. This aligns with state-of-the-art studies and current best practices in assembling high-quality bacterial genomes [14]. General assembly statistics and genome annotation are additional steps in the pipeline for further analysis. With our method, we were able to assemble 36 *M. bovis* strains examined in this study into contiguous genomes and to ensure high sequence accuracy through postprocessing of the assemblies, resulting in suitable annotations.

**Phylogenetic clades**

Comparison in a global context showed clustering according to geographical origin, with the newly assembled genomes positioned in the European cluster (comprising clades 4, 5, and 6) of the tree. In contrast, the genome of the Cuban strain DL422\_88 proved phylogenetically more distant and did not cluster with any other genome.

The geographic clustering, however, is not of high resolution, i.e. not country-specific, which is probably due to the extensive international animal trade throughout Europe. The fact that the European cluster also includes strains from the US (e.g. type strain



**Fig. 5** Gene association analysis (Fisher’s exact test) of our European assembly panel in correlation to phylogeny. We detected 108 genes to be significantly associated with at least one cluster. Of these genes, 25 are linked to a gene product or at least a class of products, including various enzymes, e.g. proteases and nucleases, as well as ten variable surface lipoproteins and other surface proteins. Some genes are listed more than once when truncated versions are also associated with a cluster

PG45) could be seen as an indication that phylogenetic processes in *M. bovis* occur at a slow pace.

Analysis of our assemblies also revealed cluster association of several genes, including the *vsps*, whose products are presumed virulence factors. Other correlations between phylogeny and strain properties were not observed. Genomic data supporting tissue specificity of strains or association with specific disease manifestations have not been provided in the literature so far. The identification of a mastitis-dominant lineage of *M. bovis* strains by Yair et al. [36] was certainly due to the relatively small size of the cattle population in Israel, where calves were imported from five to seven countries only. Nevertheless, strains associated with mastitis from that study did not cluster on a single clade, which indicates that genomic factors alone are

probably insufficient to characterize a strain’s association with a particular clinical manifestation.

**Genome, core genome, pangenome**

Genomes of *Mycoplasmopsis spp.* are the smallest among bacteria. With an average length of 1.063 Mbp, our panel of 36 strains has genomes of the expected size, with a core genome of 598 genes and a pangenome of 1,143 (Table S1). The average number of annotated protein-coding genes of 880 (Table S1) is higher than in other studies, probably due to the hybrid assembly approach, which allowed complete genome assembly in most cases and improved annotation. Kumar et al. analyzed a large set of 250 *M. bovis* strains, mainly from the US and Australia, appeared to be phylogenetically more heterogeneous than our European panel. These authors used Illumina sequence data and reported an average CDS number of

770, core and pangenome of 283 and 1,186, respectively [56]. The data of our own study comprising 219 strains are comparable: 327 core genes and 1,623 genes in the pangenome (the latter due to annotation, see above).

### Essential vs. dispensable genes

From a formal point of view, the core genome elements are considered indispensable for the organism's survival [57]. From a biological viewpoint, genes encoding replication and translation factors, as well as elements of metabolic pathways, are considered essential because they belong to the fundamental cellular machinery of bacteria. Josi et al. [58] identified 352 out of 900 *M. bovis* genes as essential, most of them involved in nucleotide metabolism or biosynthesis of secondary metabolites and amino acids. Using this definition, the clade-specific genes identified in our study (partly given in Fig. 5) would mainly be classified as non-essential.

### *vsp* locus

No precise figures on the varying size of the entire *vsp* locus in *M. bovis* genomes are available. Early data from Lysnyansky et al. [7, 8] indicate that the size of all 13 sequences coding for Vsp polypeptide chains in strain PG45 is about 11 kbp, to which the promoter (150 bp each) and signal peptide (75 bp) regions have to be added (total approx. 3 kbp). This means that 1.4 % of the genome would be used to encode Vsp family members. The proportion could be higher in strains equipped with multiple *vsp* gene copies and lower in strains lacking individual *vsp* genes.

Josi et al. classified *vsp* genes as non-essential for the organism [58]. This may be due to the software used in their study, which could not assign reads from highly repetitive regions to a specific position in the genome, and, therefore, did not consider *vsp*s. From the biological perspective, it seems more likely that the *vsp* locus is not dispensable since individual members were suggested to play a role in cytoadhesion and evasion of the host immune response [10, 59]. The fact that *M. bovis* strains can swiftly alter their Vsp repertoire and protein chain length in the face of host or environmental challenges could also indicate that the underlying genes are essential for the organism.

When Kumar et al. used the sequences of the 13 *vsp* genes of strain PG45 as BLAST queries they found that none of their strains harbored the complete *vsp* gene set [56]. While it is known that the number of *vsp* family members can be reduced in certain *M. bovis* strains the high degree of sequence variation in the *vsp* genes may have contributed to the low number of hits in that study.

### SCC as a largely unknown manifestation of *M. bovis* infection

The symptomatology observed in SCC has not been described before and represents an additional clinical picture that might be associated with *M. bovis* infection. The signs of this new condition primarily include severe edema, particularly in the thoracic and abdominal regions, as well as arthritis, even in adult animals (see Fig. S1). The formation of edema indicates involvement of the cardiovascular system. We observed several fatalities in dairy cows with SCC in conjunction with *M. bovis* isolation. Our hypothesis was that *M. bovis* strains isolated from animals presenting with SCC carry genomic traits responsible for higher virulence. However, using our comparative genomic approach involving genome-wide association analysis, such traits could not be identified. Therefore, the question of whether *M. bovis* contributes directly or indirectly to SCC pathogenesis remains open. In addition to SCC, several cases have been reported, in which *M. bovis* was not only implicated in Bovine Respiratory Disease (BRD) in calves but also caused fibrinous bronchopneumonia in adult animals reminiscent of Contagious Bovine Pleuropneumonia (CBPP) [5, 6]. This underlines the importance of *M. bovis* as a differential diagnosis for the reportable CBPP, which should be conducted by means of molecular identification tools.

### Supplementary Information

The online version contains supplementary material available at <https://doi.org/10.1186/s12864-023-09618-5>.

Additional file 1.

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We thank all the colleagues mentioned in Table 1, who kindly provided strains for this study. Susann Bahrmann is acknowledged for excellent technical assistance.

### Authors' contributions

KS, MHö, and MM designed the study. KS did conventional sequence assemblies at the initial stage. MHö and ST developed the new assembly pipeline. ST conducted the assembly and processing of raw sequence data. MHe and CS provided the strains and submitted them for sequencing. CD did Nanopore sequencing and wetlab work. MW conducted pairwise association analysis. ST, KS, MHö, and MM wrote the manuscript. All authors read and approved the final version of the manuscript.

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### Availability of data and materials

*Mycovista* is available on GitHub <https://github.com/sandraTriebl/mycovista>. Our *de novo* assembly panel (36 assemblies) is provided in the NCBi BioProject PRJNA954308.

## Declarations

### Ethics approval and consent to participate

Samples were collected by attending veterinarians as part of diagnostic investigations and as no additional pain, suffering or harm was inflicted on the animals, no approval from an ethics committee was required under national law. Written consent was obtained from the animal owners.

### Consent for publication

Not applicable.

### Competing interests

MHö is a co-founder of nanozoo GmbH and holds shares in the company. The authors declare that they have no competing interests.

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