Metagenomic sequencing determines complete infectious bronchitis virus (avian \textit{Gammacoronavirus}) vaccine strain genomes and associated viromes in chicken clinical samples

Steven Van Borm\textsuperscript{1} \textsuperscript{a} · Mieke Steensels\textsuperscript{1} · Elisabeth Mathijs\textsuperscript{1} · Frank Vandebussche\textsuperscript{1} · Thierry van den Berg\textsuperscript{1} · Bénédicte Lambrecht\textsuperscript{1}

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Abstract
Infectious bronchitis virus (IBV, genus \textit{Gammacoronavirus}) causes an economically important and highly contagious disease in chicken. Random primed RNA sequencing was applied to two IBV positive clinical samples and one in ovo-passaged virus. The virome of a cloacal swab pool was dominated by IBV (82\% of viral reads) allowing de novo assembly of a GI-13 lineage complete genome with 99.95\% nucleotide identity to vaccine strain 793B. In addition, substantial read counts (16\% of viral reads) allowed the assembly of a near-complete chicken astrovirus genome, while lower read counts identified the presence of chicken calicivirus and avian leucosis virus. Viral reads in a respiratory/intestinal tissue pool were distributed between IBV (22.53\%), \textit{Sicinivirus} (Picornaviridae, 24\%), and avian leucosis virus (37.04\%). A complete IBV genome with 99.95\% nucleotide identity to vaccine strain H120 (lineage GI-1), as well as a near-complete avian leucosis virus genome and a partial \textit{Sicinivirus} genome were assembled from the tissue sample data. Lower read counts identified chicken calicivirus, \textit{Avibirnavirus} (infectious bursal disease virus, assembling to 98.85\% of segment A and 69.66\% of segment B closely related to D3976/1 from Germany, 2017) and avian orthoreovirus, while three avian orthoavulavirus 1 reads confirmed prior real-time RT-PCR result. IBV sequence variation analysis identified both fixed and minor frequency variations in the tissue sample compared to its in ovo-passaged virus. Metagenomic methods allow the determination of complete coronavirus genomes from clinical chicken samples while providing additional insights in RNA virus sequence diversity and coinfecting viruses potentially contributing to pathogenicity.

Keywords Metagenomics · \textit{Gallus gallus} · Coronaviridae · \textit{Igacovirus} · Astrovirus · Sicinivirus · Next-Generation Sequencing

Introduction
Infectious bronchitis virus (IBV) is a gammacoronavirus of the subgenus \textit{Igacovirus} (ICTV 2020, [1]) causing an acute, highly contagious upper respiratory tract disease with potential systemic complications in chickens posing a major economic burden on the poultry industry [2]. The virus replicates in the respiratory, digestive, excretory, and reproductive system, resulting in a variety of pathogenic effects and both respiratory and gastrointestinal viral excretions are possible (reviewed in [3, 4]). In addition to respiratory signs, decreased egg quality and production are common. Serological evidence suggests that poultry workers may develop anti-IBV antibodies following exposure to infected birds, although there is no evidence of active infection in humans [5]. The virus is worldwide in distribution, and there are many genetic and antigenic types that can co-circulate [2, 6], complicating control efforts by the available live attenuated and killed vaccines [7, 8]. Moreover, like other Coronaviridae, IBV is under constant evolution resulting from error prone genome copying that accumulates mutations and widely documented recombination events [9–11]. Laboratory differential diagnostic efforts are essential because of similarities to mild forms of disease caused by agents such as Newcastle disease virus, avian metapneumovirus, infectious laryngotracheitis virus, mycoplasmas, \textit{Avibacterium paragallinarum}, and \textit{Ornithobacterium rhinotracheale} and the co-circulation of these pathogens [12].

\textsuperscript{a} Steven Van Borm
Steven.Vanborm@sciensano.be

\textsuperscript{1} Department of Animal Infectious Diseases, Sciensano, Brussels, Belgium

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Coinfections with other poultry pathogens have been described to complicate the infectious bronchitis disease severity. For instance, coinfection with IBV has been documented in H9N2 avian influenza outbreaks in poultry [12] and may contribute to the disease outcome. Coinfecting respiratory pathogens including Mycoplasma gallisepticum, Mycoplasma synoviae, avian influenza virus, IBV, and avian metapneumovirus seem to frequently occur, as described in Algerian poultry flocks [13]. In addition, infectious bronchitis disease course has been documented to be influenced by infectious bursal disease virus and chicken anemia virus induced immunosuppression [12, 14].

Metagenomic methods, here defined as the random sequencing of all nucleic acids in a sample, are increasingly used since the wider availability of Next-Generation Sequencing (NGS) platforms. Potentially detecting all coinfecting pathogens, these methods proved a huge added value in a context where disease outcome is influenced by multiple infection (e.g., [15, 16]). Moreover, the hypothesis-free methodology allows the detection of unexpected pathogens or variants thereof, the investigation of diagnostic cases that have exhausted the available targeted assays (e.g., [17]), and provides a powerful tool for the discovery of novel viruses [18, 19]. Indeed, random metagenomic methods were critical in the identification of SARS-CoV2 as the etiological agent of the current COVID-19 pandemic [20] and novel emerging livestock diseases like Schmallenberg virus [21].

The objectives of this study were (1) to validate metagenomic protocols for sequencing of avian coronavirus complete genomes from clinical samples; (2) the genetic characterization of selected IBV recently circulating in Belgium; and (3) the detection of viruses co-circulating with IBV.

**Materials and methods**

**Samples, pretreatment, and targeted testing**

One pooled sample of lung, trachea, and intestinal tissue from five broiler chickens (4439-PTLB) was submitted for Newcastle disease virus (NDV) diagnostics and differential diagnosis and tested weak positive (Cp 36.35) for avian Orthoavulavirus 1 (NDV), as well as Igacovirus (Infectious bronchitis virus of poultry, Cp 32.29) using routine real-time RT-PCR assays [22, 23]. The sample pretreatment included homogenization in sterile PBS (10% weight/volume) with antibiotics (penicillin: 10.106 U/L, streptomycin 10 g/L, gentamycin 0.25 g/L), followed by 10-min centrifugation at 2500 rpm at 4 °C and harvesting of the supernatant. A derived chorioallantoic fluid sample (4439_ECE) was obtained from the latter tissue sample following routine virus isolation procedures using two 5-day amplification rounds in the allantoic cavity of 9–11-day-old specific pathogen-free embryonated chicken eggs [24]. In addition, a pool of five chicken cloacal swabs (4134_PCS), in viral transport medium (BHI 37 g/L, supplemented with 107 U/L penicillin, 2 g/L streptomycin, 1 g/L gentamycin, and 650 mg/L kanamycin), testing positive for IBV (Cp 24.82) was included. Both clinical samples were investigated in a context of avian influenza and NDV exclusion diagnostics and differential diagnosis in farms showing increased mortality. The homogenate of the tissue sample pool, the pool of cloacal swabs, and the chorioallantoic fluid resulting from the in ovo passage of the former tissue sample pool were included in this metagenomic study.

**Sequencing**

The tissue homogenate, swab transport medium, and chorioallantoic fluid were centrifuged for 5 min at 10,000g in a precooled (4 °C) centrifuge prior to collection of the supernatant. RNA was extracted from the supernatant using a combination of the TRizol reagent (Thermo Fisher) and RNeasy mini kit (Qiagen), including on-column DNase treatment as previously described [25]. cDNA was synthesized using SuperScript IV reverse transcriptase (Thermo Fisher Scientific) and random hexamer primers, followed by double-strand cDNA synthesis using the NEBNext mRNA second-strand synthesis module (New England BioLabs). Sequencing libraries were prepared using the Nextera XT kit (Illumina) and standard Nextera XT index adapters (Illumina) and sequenced using a MiSeq reagent kit version 3 (Illumina) with 2x300-bp paired-end sequencing aiming for a minimum of 5-million read pairs per sample. Metagenomic NGS data were generated for 3 samples. The resulting fastq raw metagenomics datasets are publicly available in the Sequence Read Archive (SRA) under BioSample accession numbers SAMN19554998 (tissue sample pool), SAMN19554999 (in ovo passage resulting from tissue sample pool), and SAMN19555000 (cloacal swabs pool).

**Bioinformatic analysis**

Metagenomic read classification: raw NGS reads were trimmed using Trimmomatic v0.38 [26] to remove adapter sequences and low quality bases (setting the ‘ILLUMINACLIP:2:30:10’, ‘SLIDINGWINDOW:4:20’, and ‘MINLEN:50’ options). Only paired reads were retained for further analysis. Classification of trimmed reads was performed with Kraken2 (Galaxy version 2.0.7) as previously described [27]. A customized Kraken database was built using all available RefSeq “Complete Genome” sequences of six targeted taxonomic groups (archaea, bacteria, fungi, human, protozoa, and viral) downloaded from RefSeq Genome (ftp://ftp.ncbi.nlm.nih.gov/ genomes/refseq/) on 18/02/2019. The proportion of reads mapping to the host (Gallus gallus) was determined using a separate database of selected avian
genomes (Gallus gallus GCF-000002315; Columba livia GCF-00037935; Meleagris gallopava GCF-000146605; Anas Platyrhynchos GCF-000355855; Numida meleagris GCF-002078875). Read counts per taxonomic level classified by Kraken2 were further normalized as reads per million (RPM) total (trimmed) reads [27] to remove technical bias introduced by sequencing depth variation between samples. An arbitrary acceptance criterion of RPM > 1 for a significant mNGS finding was used as previously described [27].

De novo assembly and complete genome characterization

Raw sequence data was trimmed using Trim Galore! V0.5.0 (q = 25, l = 50, paired; https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/). The genome of IBV strain IBV/chicken/Belgium/4439_001_iPTLB/2020 was de novo assembled from a random paired subset (2 × 50,000 reads) of the reads resulting from allantoic fluid sample 4439_ECE using SPAdes v3.9.0 [28] and IVA v1.0.0. [29]. Relevant contigs (Igacovirus, as determined using a Blastn search of all contigs longer than 500 nt against the NCBI Nucleotide database) were joined using CAP3 [30]. The full trimmed dataset was mapped to the finished full genome length contig using BWA-0.6.2 [31]. Minimap2 [32], Samtools [33], and bamToFastq (https://bedtools.readthedocs.io/en/latest/content/tools/bamtofastq.html) were used to enrich on-target reads from datasets resulting from the clinical samples, using specific databases for Coronaviridae (NCBI:txid11118, all RefSeq entries on 14/08/2020, n = 66), Sicinivirus (NCBI:txid1755587, all NCBI nt entries with length > 2000 nt, n = 24), avian leucosis virus (NCBI:txid11864, all NCBI nt entries with length > 7000 nt, n = 200), and avian astrovirus (NCBI:txid249589, all NCBI nt entries with length > 7000 nt, n = 64). For data resulting from the cloacal swabs pool 4134_PCS, random paired subsets (2 × 50,000 reads) were used for the de novo assembly of Coronaviridae - and Avastrovirus-enriched reads, respectively, using the aforementioned strategy (IVA, SPAdes, CAP3), followed by mapping of all trimmed reads against the resulting contigs (Minimap2). All Coronaviridae, Sicinivirus, and avian leucosis virus reads enriched from the trimmed datasets resulting from tissue sample pool 4439_PTLB were used for de novo assembly using the aforementioned strategy. For Taxa reported at lower RPM in the Kraken2 analysis, all classified reads were extracted using Minimap2 and the reference genome(s) present in the Kraken2 classification database, followed by de novo assembly using SPAdes v3.9.0 [28] and similarity searches (Blastn) of resulting contigs or singletons.

To identify conserved protein domains, a RPS-BLAST search [34] against the Conserved Domain Database (CDD) was performed [35]. The annotation of assembled genomes and partial genomes was done using GATU [36] relative to references KF377577.1 and MN548287.1 with manual editing of the coronaviral conserved -1 ribosomal frameshift in ORF 1ab for the avian coronavirus genomes.

Identification of sequence variation in Igacovirus RNA population

The Bam files representing the mapping of all quality-trimmed data from the tissue sample 4439-PTLB or 4439_ECE to assembled genome IBV/chicken/Belgium/4439_001_iPTLB/2020 (consensus genome from 4439_ECE) were used to call RNA population frequency variation using Lofreq [37] using filtering settings min_dp_100, sb_fdr (applying a strand bias multiple testing correction using a fdr correction p value > 0.001), min_snv_qual_66, and min_indelqual_20. Only variants with a minimum allele frequency of 0.01 and a positional coverage of at least 100 were included.

Phylogenetic analysis

MAFFT v7.310 [38, 39] was used to align the complete S1 coding sequences of infectious bronchitis virus genomes IBV/chicken/Belgium/4439_001-iPTLB/2020 (genome assembled from chorioallantoic fluid 4439_ECE), IBV/chicken/Belgium/4439_001_PTLB/2020 (genome assembled from the tissue sample pool 4439_PTLB), and IBV/chicken/Belgium/4134_001/2019 (genome assembled from the cloacal swab pool, 4134_PCS) with the reference dataset proposed by Valastro and colleagues [6]. The final dataset contained 204 sequences of 1567 nucleotides long. After selection of the most suitable evolutionary model (lowest Bayesian Information Criterion score), a maximum likelihood phylogenetic tree was calculated in Mega X [40] (GTR + G + I [41]; partial deletion of missing data and gaps; 500 bootstrap replicates). For visual clarification, subtrees representing the identified lineages GI-1 and GI-13 were extracted from the full-dataset phylogenetic tree.

Due to low nucleotide sequence similarity with other siciniviruses, the amino acid sequence of the conserved RdRp domain was extracted from all Sicinivirus entries in the NCBI nucleotide database with length > 2000 (n = 15 after removal of duplicates and entries lacking the RdRp domain, NCBI:txid1755587) and aligned with the RdRp region of Sicinivirus strain Belgium/4439_001/2020 and two root sequences representing sister clades (Gallivirus NC018400 and Turdivirus NC014411) using MAFFT v7.310, followed by maximum likelihood phylogenetic analysis using Mega X (LG + G + I model; 500 bootstrap replicates).

The RdRp complete coding sequence as well as the Capsid partial coding sequence of chicken astrovirus strain Belgium/4134/2019 were aligned with 31, respectively, 68 coding sequences available in NCBI nucleotide database.
using MAFFT v7.310, followed by maximum likelihood phylogenetic analysis using Mega X (LG + G + I model; 100 bootstrap replicates).

Results

Next-generation sequencing raw data

Minimum 5.3 million paired reads were produced per sample (Table 1). As expected for tissues samples, 90% of the reads in sample 4439_PTLB corresponded to the host (chicken), while 1% of reads were viral and minimal bacterial contamination was seen (4%). The results for the corresponding chorioallantoic fluid, 4439_ECE, showed an excellent purity with 77% of the reads being viral and 18% avian, reflecting the viral growth in embryonated chicken eggs. The cloacal swab pool sample 4134-PCS showed a relatively high proportion (10%) of viral reads and as expected for cloacal samples, a high proportion of bacterial reads (56%), and relatively low proportion of host reads (13%).

Virome composition using Kraken2

A diverse virome was detected in pooled tissue sample 4439_PTLB (IBV and NDV RT-PCR positive tissue pooled sample), dominated by IBV (Gammaronavirus, representing 22.54% of viral reads), Sincivirus (Picornaviridae, 23.99% of viral reads), and avian leucosis virus (37.04% of viral reads) (Table 2). In addition, lower normalized read counts were detected for chicken astrovirus (0.26% of viral reads), while chicken calicivirus, IBDV (assembling to 98.85% of segment A and 69.66% of segment B of IBDV closely related to D3976/1 from Germany, 2017), and avian orthoreovirus were detected with normalized read counts just above our significance threshold. With only 3 reads detected, avian orthoavalivirus remained below our arbitrary significance level of RPM > 1.

Table 1  Distribution of metagenomics reads over (super)kingdoms

|         | U2004439PBLT | U2004439isolate | U1904134PCS |
|---------|--------------|----------------|------------|
| Archaea | 0.009%       | 0.001%         | 0.005%     |
| Bacteria| 4.000%       | 0.600%         | 56.000%    |
| No hits | 10.000%      | 8.000%         | 20.000%    |
| Fungi   | 0.000%       | 0.000%         | 3.000%     |
| Protozoa| 0.047%       | 0.026%         | 0.630%     |
| Galloanserae | 90.000% | 18.000% | 13.000% |
| Viruses | 1.000%       | 77.000%        | 10.000%    |

*The proportion of host reads was determined using a separate reference database with members of the Galloanserae

The IBV/chicken/Belgium/4439_001_iPTLB/2020 virus sequenced from the in ovo passage (4439_ECE) of the above sample showed a remarkable increase in the amount and proportion of coronaviral reads (99.80% of viral reads) (Table 2). About half of the avian coronavirus reads were attributed to IBV and half to turkey coronavirus, both being closely related species. Only a single Sincivirus read was detected in the chorioallantoic fluid sample (possibly carry-over between sequencing libraries), and only low read counts for avian leucosis virus (Retroviridae), while these taxa jointly contributed more than 60% of the viral reads in the clinical sample. Other viruses present in the clinical sample were not detected in its derived chorioallantoic fluid sample. Interestingly, not only gammacoronaviruses were classified by the Kraken2 approach (99.77% of viral reads) but also significant normalized read counts for betacoronaviruses and deltacoronaviruses. However, follow-up investigation (de novo assembly of Kraken2-classified beta- and deltacoronaviral reads followed by Blastn analysis of resulting contigs) showed that these reads mapped to conserved coronaviral sequences. Only gammacoronaviral contigs (IBV H120) were de novo assembled from reads classified by kraken2 as beta- or deltacoronaviral.

82.15% of viral reads from the pool of cloacal swabs 4134_PCS represented avian coronavirus (Gammacoronavirus), while additional alpha-, beta-, and deltacoronaviruses were present at lower normalized read counts (Table 2). About half of the avian coronavirus reads were attributed to IBV and half to turkey coronavirus, both being closely related species. Follow-up investigation (de novo assembly of Kraken2-classified beta- and deltacoronaviral reads followed by Blastn analysis of resulting contigs) showed that non-gamma coronaviral reads mapped to conserved coronaviral sequences. Only gammacoronaviral contigs were de novo assembled from reads classified by kraken2 as beta- or deltacoronaviral. In addition, 16.14% of viral reads represented chicken astrovirus, with some additional astroviral reads from related species. The low number of canine and porcine astrovirus reads represented bioinformatic artifacts (low complexity regions or database biases). Chicken calicivirus and pigeon picornavirus B were detected with significant normalized read numbers. However, follow-up investigation (de novo assembly of Kraken2-classified pigeon picornavirus B reads followed by Blastn analysis of resulting contigs) showed that the pigeon picornavirus B reads mapped to conserved picornaviral RdRp sequences. Avian leucosis virus was detected in the cloacal swab pool above our significance threshold, but with low normalized read numbers as expected because of the low amount of host cells in cloacal swabs.

Low normalized read counts for bacteriophage sequences were observed in the tissue sample and its derived in ovo passage, while a moderate RPM for bacteriophages was detected in the cloacal swab, as well as a few reads classifying as an environmental aquatic virus.
Table 2 RNA virome composition of chicken samples in the present study

| Sample          | 4439_PTLB | 4439_ECE | 4134PCS |
|-----------------|-----------|----------|---------|
| Dataset size (read pairs) | 6664056   | 6889556  | 5325938 |

**Coronaviridae**

- **All avian coronavirus** (Gammacoronavirus) 3134.6 (22.5% of viral reads) 771877.0 (99.8% of viral reads) 80425.5 (82.2% of viral reads)
- **Infectious bronchitis virus** (Gammacoronavirus) 1616.4
- **Turkey coronavirus** (Gammacoronavirus) 1477.8
- **Unassigned avian coronavirus** (Gammacoronavirus) 40.4
- **Bovine coronavirus** (Betacoronavirus)† –
- **Rat coronavirus Parker** (Betacoronavirus)† –
- **Rousettus bat coronavirus HKU19** (Betacoronavirus)† –
- **Thrush CoV HKU12-600** (Deltacoronavirus)† –
- **White-eye coronavirus HKU16** (Deltacoronavirus)† –
- **Unassigned orthocoronvirinae** –
- **Human coronavirus HKU1** (Betacoronavirus)† –
- **Bat Hp-betacoronavirus/Zheijang 2013** (Betacoronavirus)† –
- **Thrush deltacoronavirus HKU12-600** (Deltacoronavirus)† –
- **Wigeon coronavirus HKU20** (Deltacoronavirus)† –
- **Unassigned deltacoronaviridae** (Deltacoronavirus)† –
- **NL63-related bat coronavirus** (Alphacoronavirus)† –

**Picornaviridae**

- **Sicinivirus** (all) 3335.5 (24% of viral reads) 0.1 –
- **Sicinivirus A** 3112.4 0.2 –
- **Chicken Sicinivirus JSY** 210.1 – –
- **Unassigned Sicinivirus** 13.1 – –
- **Chicken picornavirus 1** 135.4 – –
- **Unassigned Picornaviridae** 9.6 – –
- **Pigeon picornavirus B†** – – 130.5

**Astroviridae**

- **Chicken astrovirus** 31.7 (0.2% of viral reads) – 15803.2 (16.1% of viral reads)
- **Goose astrovirus** – – 801.2
- **Duck astrovirus** – – 197.0
- **Unclassified avastrovirus** – – 8.4
- **Canine astrovirus** – – 2.4
- **Porcine astrovirus 3** – – 1.3

**Retroviridae**

- **Avian leukosis virus** 5150.5 (37% of viral reads) 112.9 (0.01% of viral reads) 21.2 (0.02% of viral reads)
- **Avian endogenous retrovirus** 18.3 – –
- **Y73 sarcoma virus** 636.4 11.6 4.1
- **Fujinami sarcoma virus** 457.4 4.1 3.4
- **Rous sarcoma virus** 414.8 5.1 1.3
- **Avian carcinoma virus** 101.0 2.5 0.9
- **Avian myelocytomatosis virus** 34.7 7.8 –
- **UR2 sarcoma virus** 100.8 0.3 –
- **Avian endogenous retrovirus EAV-HP** 18.3 – – 2.3
- **Other alpharetroviridae** 276.1 – –
- **Avian sarcoma virus CT10** – 1.7 0.6

**Caliciviridae**
Table 2 (continued)

| Sample                  | 4439_PTLB          | 4439_ECE          | 4134_PCS          |
|-------------------------|--------------------|-------------------|-------------------|
| Dataset size (read pairs) | 6664056           | 6889556           | 5325938           |
| Chicken calicivirus     | 11.7               | –                 | 170.5             |
| Paramyxoviridae         |                    |                   |                   |
| Avulavirus*             | 0.5                | –                 | –                 |
| (3 reads)               |                    |                   |                   |
| Birnaviridae            |                    |                   |                   |
| IBDV (avibirnavirus)    | 34.7               | –                 | –                 |
| Reoviridae              |                    |                   |                   |
| Avian orthoreovirus (Reoviridae) | 3.8           | –                 | –                 |
| All Caudovirales (bacteriophage)* | 5.7            | 1.6               | 31.2              |
| Birnaviridae            |                    |                   |                   |
| Enterobacteria phage MS2* | –                | –                 | 8.3               |
| Unassigned ss + RNA viruses | –                | –                 | 3.9               |
| Unclassified viruses    |                    |                   |                   |
| Pandoravirus salinus    | –                  | –                 | 1.3               |

The normalized read count (RPM) of detected taxa is reported for each sample

'–': not detected

*Avian virus detected below significance threshold (RPM > 1)

† Bioinformatic artifact invalidated by follow-up research

* Bacteriophage

Assembly of viral genomes and molecular characterization

Igacovirus

From the data resulting from the embryonated chicken egg-passaged sample 4439_ECE, a 27,648 bp contig was de novo assembled from Coronaviridae-enriched reads (using an enrichment database of all 66 complete coronavirus genomes in the RefSeq database and extraction of mapped reads). 6 million read pairs aligned to this contig. This complete genome sequence of IBV isolate IBV/chicken/Belgium/4439_001_iPTLB/2020 (submitted to Genbank under accession number MZ367367) has a 99.95% nucleotide identity to IBV strain H120 (MN548287.1), a widely used vaccine of the Mass serotype. Variant calling using Lofreq on the mapped coronaviral reads from the egg-passaged sample (using a minimum positional coverage of 100x as well as a minimum allele frequency of 0.10 indicated the presence of 5 additional genome positions with mixed populations (in the polyprotein 1ab as well as the envelope protein coding regions; Suppl. Table 1). Minor frequency variation (0.01 ≤ allele frequency ≤ 0.10) was detected in 40 additional genome positions scattered throughout the genome (Suppl. Table 1). All variant and minor frequency variant positions detected in the clinical sample were distinct from variant positions detected in the egg-passaged sample (Suppl. Table 1).

A near-complete avian coronavirus genome (27,600 bp contig, submitted under Genbank accession number MZ367369) was de novo assembled from Coronaviridae-enriched reads from the pooled swabs 4134_PCS. About 1.2 million reads aligned to this contig. This near-complete genome sequence obtained from pooled cloacal swabs showed a 99.98% nucleotide identity to IBV strain ck/CH/LLN/130102 (KP118888.1), a 793B genotype strain isolated...
in China in 2013, and a 99.95% nucleotide identity to IBV vaccine strain 4/91 (KF377577.1, a 793B-type strain).

Phylogenetic classification of the complete S1 coding sequences from the three samples in comparison with 201 reference sequences confirmed the circulation of IBV with close similarity to H120 (4439_PTLB) and 793B B genotype (4134_PCS) vaccine-like strains of lineages GI-1 and GI-13, respectively (Fig. 1), following the classification criteria and nomenclature by Valastro and colleagues [6]. The genomes assembled from the sample 4439_PTLB and the corresponding in ovo-passage IBV/chicken/Belgium/4439_001_iPTLB/2020 from 2020 shared an identical S1 complete coding sequence.

A partial Sicinivirus genome was de novo assembled from Sicinivirus-enriched reads originating from the pooled tissue sample 4439_PTLB (using an enrichment database containing all 24 Sicinivirus in nt database longer than 2000 nucleotides and extraction of mapped reads). Two contigs were assembled, corresponding to the 5′ end (1347 nt) of MN873047 and the 3′ end (7719 nt) of MN873045, both from chicken fecal samples, USA 2003–2004. These partial sequences were submitted to Genbank under accession numbers MZ367370 and MZ367371. The largest contig encoded the anterior 2573 amino acids of the polyprotein and contains two picornaviral capsid (rhv), an RNA helicase, and an RNA-dependent DNA polymerase (RdDp)-conserved domain. The maximum likelihood amino acid phylogenetic analysis of RdDp conserved region of 28 Sicinivirus sequences (Supplementary Fig. 1) confirmed grouping with other Sicinivirus genomes and lack of geographical clustering which may indicate widespread global circulation of Sicinivirus.

The near-complete genome of an endogenous avian leukosis genome (8532 nt contig, Genbank accession number MZ367376) was de novo assembled from avian leucosis virus-enriched reads from the tissue sample pool 4439_PTLB (enrichment database containing all nt entries longer than 7000 nt). The 8532 nt contig showed 99.95% nucleotide identity to avian leukosis virus genome ev21 (KY235336.1), an avian leukosis subgroup E endogenous virus sequenced from a chicken in China in 2015.

A partial genome (covering positions 8-6478 and 6714-7497, Genbank accession number MZ367372) of a chicken astrovirus was de novo assembled from avian

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Fig. 1 Maximum likelihood phylogenetic analysis of the complete S1 coding sequence of 204 infectious bronchitis virus strains. A GTR + G + I evolutionary model was used with partial deletion of missing data and gaps and 500 bootstrap replicates to evaluate the significance of nodes. For visual clarification, subtrees representing the identified lineages GI-1 and GI-13 were extracted from the full-dataset phylogenetic tree: sequence determined in the present study. IBV/chicken/Belgium/4439_001_PTLB/2020 and IBV/chicken/Belgium/4439_001_iPTLB/2002 correspond to the genome sequences determined from the tissue sample pool and the derived embryonated egg-passaged virus, respectively, while Infectious bronchitis virus strain IBV/chicken/Belgium/4134_001/2019 corresponds to the sequence determined from the cloacal swabs pool.
Discussion

Since 2006, an enhanced surveillance has been put in place in the Belgian poultry sector for avian influenza. Therapeutic treatment of poultry may only be initiated after the following findings have been submitted to the regional laboratories for examination: a reduction of more than 20% in normal feed and water consumption, a mortality of more than 3% per week, a reduction in egg laying of more than 5% for more than 2 days, or clinical signs or post-mortem lesions suggestive of influenza. Samples analyzed in this study originated from this surveillance and the investigated flocks did not show any specific clinical signs besides increased mortality rates. In addition, vaccination against NDV is compulsory in Belgium and live vaccines are used at young age. Moreover, vaccination against IBV and IBDV is also widely practiced in parallel using live vaccines harboring various residual pathogenicity. Therefore, a booster spray vaccination with a NDV live-attenuated vaccine 4 weeks prior to obtaining sample 4439_PTLB.

The comparative metagenomic sequencing of a clinical tissue sample (4439_PTLB) and its derived embryonated chicken egg-passaged sample (4439_ECE) demonstrates a remarkable loss of viral taxa (from the families Picornaviridae, Astroviridae, Caliciviridae, Birnaviridae, and Reoviridae) during routine virus isolation procedures. These viruses are either not easily cultivated or need different biological systems from embryonated chicken eggs routinely used to grow notifiable poultry viruses.

Besides high numbers of gammacoronaviral sequences corresponding to Igaviruses, our original Kraken2 classification identified alpha-, beta-, and deltacoronavirus reads in two samples. However, careful follow-up investigation (extraction of reads classified as alpha beta delta + de novo assembly + blastn similarity searches against the entire nt database) showed that these reads and their resulting contigs aligned to conserved regions of IBV strain H120 genome and hence do not allow a classification below the Coronaviridae family level. Similarly, reads classifying as pigeon picornavirus mapped to a conserved picornaviral sequence (RdRp motif) and thus most likely reflect RdRp sequences from Siciniviruses. The validation of the NDV detection based on the presence of NDV vaccine strain traces vs. circulating field viruses. Of note, the poultry flock received a booster spray vaccination with a NDV live-attenuated vaccine 4 weeks prior to obtaining sample 4439_PTLB.

The 908 calicivirus reads identified in the cloacal swab pool were dispersed unequally over the reference (NC033081) genome, resulting in a total coverage breadth of 65.76% of the genome. However, the unequal distribution of reads across the genome only allowed a single contiguous > 1000 bp to be de novo assembled (1169 nt, coverage 52.99x, Genbank accession number MZ367375) showing an 89.56% nucleotide identity to the partial polyprotein and VP2 coding sequence of a calicivirus from Korean poultry (Supplementary Fig. 2).

Although only a limited normalized read count was reported, two contigs were de novo assembled from Infectious bursal disease virus-classified reads originating from tissue sample pool 4439_PTLB corresponding to 98.85% of segment A and 69.66% of segment B of an IBDV closely related to recombinant strain D3976/1 from Germany, 2017 (98.63% identity to MN786767.1 segment A and 99.29% identity to MN786769.1 segment B). These partial genome sequences of IBDV/Belgium/4439_001/2020 were submitted to Genbank under accession numbers MZ367373 and MZ367374.

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on only 3 reads and the invalidation of relatively high normalized “non-gamma” coronaviral read counts as conserved coronaviral domains illustrate the critical importance of expert interpretation and follow-up of metagenomic results as was also highlighted in a recent proficiency study [43].

The sequenced Igacovirus genomes from 2019 (pool of cloacal swabs) and 2020 (respiratory and intestinal tissue sample pool) confirm the detection of IBV vaccine strains in Belgium including 4/91 (lineage GI-13) and H120 (lineage GI-1), respectively. However, the limited sampling size does not allow any conclusions about potential circulation of other field and vaccine strains. IBV control relies on a combination of routine vaccination and biosecurity measures [44]. Vaccination programs are complicated by the high genetic variability of the avian coronaviruses, powered both by mutation and recombination events, reducing cross-protection conferred by vaccines toward new variants [45]. Widespread circulation of IBV vaccine strains has been previously documented [46]. A recent detailed survey based on partial S1 gene sequences of IBV strains circulating in Poland suggested that vaccine spreading and persistence seem to occur commonly, stressing the need to further study the epidemiological consequences of the extensive use of live vaccines [46]. In Belgium, a recent study indicated the circulation of at least 7 IBV types, predominated by 4/91-793B and QX-D388, and that a large proportion of the detected IB viruses were vaccine strains [47], confirming a previous study documenting high proportions of vaccine strains as well as a predominant circulation of QX-like field virus [48]. In addition, the genome of a nephropathogenic IBV reference strain isolated in Belgium in 1984 was recently fully sequenced [49].

In Belgium, both H120 (Mass-type) and 4/91 (793B-type) strain-based live-attenuated vaccines are routinely used in poultry [50] and their use in combination is known as “protectotype” approach [8]. Both sampled farms were vaccinated with live-attenuated IBV vaccines according to common practices in Belgium. Unfortunately, the vaccination records of the affected flocks were not able to allow the identification of the exact vaccine used, nor whether the detection of high IBV vaccine strain genome loads in the clinical samples was linked to a recent vaccination event or, alternatively, was a demonstration of circulation or persistence of live-attenuated vaccines. A recent survey in vaccinated poultry in Poland identified one of the applied vaccines in 46.3% of IBV-positive samples [46], while extended persistence of vaccine IBV has been documented previously, especially for 793B vaccines [51, 52].

Comparing the full genome resulting from tissue sample pool 4439_PTLB and 4439_ECE, the corresponding chorioallantoic liquid sample obtained through inoculation, and a subsequent passage in embryonated chicken eggs, only a single consensus level mutation was observed in the envelope small membrane protein coding sequence. This is within the expected range, as previous studies comparing full genome sequences of wild type and egg-adapted IBV focused on high passage attenuation histories (typically 75–100 passages) and documented up to 44 consensus level mutations [53, 54], with mutation hot spots in the polyprotein [53], spike [53, 54], nucleocapsid, and 3’UTR regions [54]. The full IBV genomes were sequenced to a sufficient depth to document virus RNA population differences between clinical sample 4439_PTLB and the corresponding chorioallantoic liquid sample 4439_ECE. Remarkable, a single variant from the clinical sample got fixed in the ECE sample, while all other variants (minimum allele frequency 0.10) and minor frequency variants present in the clinical sample got lost from the RNA population. Moreover, the egg passage introduced 2 new variant sites and 24 new minor frequency variants in the RNA population of 4439_ECE. This indicates a loss of viral diversity from the original population in combination with the generation of novel variation introduced by the passage in embryonated chicken eggs.

An important complementary value of metagenomic methods to targeted diagnostic assays is demonstrated here by documenting the full virome of co-infecting-known poultry viruses, including their high-resolution molecular characterization. In the present study this resulted in (near)-complete genome sequences of a chicken astrovirus of serotype B, an endogenous avian leukosis virus of subtype E, and a Sicinivirus.

Chicken astrovirus is an enteric poultry virus that was identified only in 2004 [55] as a separate species of avian astrovirus. It has been documented globally as an enteric virus in poultry with studies from Poland [56], Jordan [57], India [58, 59], Nigeria [60], China [61], Korea [62], and Brazil [63]. Although its contribution to pathology is often unclear, chicken astrovirus (CAstV) has been associated with poor growth of broiler flocks, enteritis, and diarrhea and is a candidate pathogen in cases of running stunting syndrome. More recently chicken astrovirus has been implicated in cases of two other diseases of broilers as the sole etiological agent, namely severe kidney disease of young broilers with visceral gout and the “White Chicks” hatchery disease [64]. Our phylogenetic analysis of the Capsid and RdRp coding sequences classifies this first Belgian near-complete chicken astrovirus genome as a member of the predominant serotype CAstV-B [65]. We identified a high normalized astroviral read count in a pool of cloacal swabs, suggesting a high viral load. However, different methodological biases prevent a correlation of normalized read counts and viral load, as some virus families may be preferentially sequenced or identified in bioinformatical workflows.

The present metagenomics study further confirmed the circulation of two recently discovered uncultivable intestinal poultry viruses in Belgium. Sicinivirus was discovered as a new picornavirus with unknown clinical manifestation of
poultry in 2012 in Ireland [66] and currently has a worldwide documented distribution [67–70], while a retrospective metagenomic analysis showed its presence in samples dating back to 2003 [71]. Our study identified a significant normalized *Sicinivirus* read count in a pool of respiratory and intestinal tissue samples, suggesting a relatively high viral load.

Chicken calcivirus was discovered as an intestinal virus on chicken farms in Germany and in the Netherlands in 2012 [72] and has been confirmed in Brazil, Korea, and the USA (unpublished sequence information cf. NCBI nt database txid1172196 including retrospective identification in a USA sample from 2003). Our study identified a significant but relatively low normalized calcivirus read count in a pool of cloacal swabs, suggesting low viral load.

The identified avian leucosis virus in the tissue sample pool belongs to subgroup E, typically representing endogenous viruses integrated in the host genome [73]. The high normalized read counts for this endogenous virus reflect the high host genomic content (90% of trimmed reads) of the pooled tissue sample. Although only reported with a moderate normalized read count, the *Avibirnavirus* reads detected in the tissue sample pool assembled to a near-complete segment A and 77% of segment B of infectious bursal disease virus, confirming the previously documented circulation in Europe of recombinant strains with a segment A from very virulent and segment B from attenuated IBDV strains [74]. This finding may be of particular relevance to the observed high IBV genome loads, as IBV disease course can be influenced by IBDV-induced immunosuppression [12, 14].

A limited number of reads (RPM = 3.8) classified as avian orthoreovirus, a globally distributed avian virus, of which some strains cause poultry disease inducing lameness, gastrointestinal, and respiratory and neurological signs, making it relevant in differential diagnosis for notifiable poultry diseases such as avian influenza and Newcastle disease. The reads are mostly distributed along the L3 segment of the single avian orthoreovirus genome included in our Kraken classification database and failed to assemble in meaningful contigs de novo (applying a minimum coverage threshold of 5× and a minimum contig length of 700) [75].

In conclusion, in addition to providing complete genome sequences of viruses present with sufficient viral load in clinical samples—here exemplified by avian gammacoronaviruses-, metagenomic methods allow the detection and characterization of co-circulating viruses that may potentially contribute to the severity of the disease. Moreover, they may facilitate differential diagnosis as exemplified here by the detection of IB, IBDV, NDV, *Sicinivirus*, *Orthoreovirus*, and calcivirus in samples originally submitted to the diagnostic laboratory for NDV exclusion. In this sense, these open scope genome detection methods are an important addition to targeted molecular diagnostic assays to unravel clinical issues in livestock. However, a careful follow-up and interpretation of metagenomics findings are essential and reproduction of the disease in controlled conditions might be required.

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**Author contributions** SVB, FV, EM, and MS performed experiments; SVB, EM, and FV performed data analysis; SVB, MS, TVDB, and BL designed the experiment and wrote the manuscript.

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**Availability of data and material** Raw NGS metagenomic data are publicly available in the Sequence Read Archive (SRA) under BioSample accession numbers SAMN19554998-SAMN19555000. Viral genome sequences and partial genome sequences are available under Genbank accession numbers MZ367367-MZ367376.

**Code availability** Not applicable.

**Declarations**

**Conflict of interest** The authors declare that they have no conflict of interest.

**Informed consent** Not applicable.

**Consent for publication** All contributing authors have read and approved of the final version of the manuscript.

**Research involving human participants and/or animals** As only bio-banked clinical animal specimens were used, no approval from the ethical committee for animal experiments was needed. Laboratory work and data analysis at Sciensano were not subjected to ethical approval.

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