Cell-Adapted Mutations and Antigenic Diversity of Influenza B Viruses in Missouri, 2019–2020 Season

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Abstract: Influenza B viruses (IBVs) are causing an increasing burden of morbidity and mortality, yet the prevalence of culture-adapted mutations in human seasonal IBVs are unclear. We collected 368 clinical samples from patients with influenza-like illness in Missouri during the 2019–2020 influenza season and recovered 146 influenza isolates including 38 IBV isolates. Of MDCK-CCL34, MDCK-Siat1, and humanized MDCK (hCK), hCK showed the highest virus recovery efficiency. All Missourian IBVs belonged to the Victoria V1A.3 lineage, all of which contained a three-amino acid deletion on the HA protein and were antigenically distant from the Victoria lineage IBV vaccine strain used during that season. By comparing genomic sequences of these IBVs in 31 paired samples, eight cell-adapted nonsynonymous mutations were identified, with the majority in the RNA polymerase. Analyses of IBV clinical sample–isolate pairs from public databases further showed that cell- and egg-adapted mutations occurred more widely in viral proteins, including the receptor and antibody binding sites on HA. Our study suggests that hCK is an effective platform for IBV isolation and that culture-adapted mutations may occur during IBV isolation. As culture-adapted mutations may affect subsequent virus studies and vaccine development, the knowledge from this study may help optimize strategies for influenza surveillance, vaccine strain selection, and vaccine development.

Keywords: influenza; respiratory diseases; influenza vaccines; influenza B virus; influenza viruses type B; antigenic drift; reassortment

1. Introduction
The seasonal influenza virus is responsible for 290,000–650,000 human deaths worldwide each year [1]. These annual epidemics are caused by the co-circulation of influenza A viruses (IAVs) and influenza B viruses (IBVs). The influenza A subtypes causing contemporary seasonal outbreaks are primarily composed of subtypes H1N1 and H3N2 viruses, which have been co-circulating in humans since 1977. The H1N1 subtype caused the 1918 pandemic and circulated in humans until 1957 [2], then re-emerged and caused seasonal outbreaks from 1977 through 2009 [3]. This was later replaced with a novel
swine-origin H1N1 virus, resulting in the 2009 pandemic [4,5]. The H3N2 subtype began circulating in 1968 and caused the 1968 pandemic [6,7]. Seasonal influenza B, identified in 1940 [8], comprises two lineages, B/Yamagata/16/88-like [Yamagata] and B/Victoria/2/87-like [Victoria], which have been co-circulating since at least 1983 [9,10]. While IAVs have been responsible for the majority of seasonal influenza complications, IBVs are causing considerably high burdens of morbidity and mortality [11–14].

Both genetic and antigenic drift facilitate the annual recurrence of influenza epidemics [15]. As an RNA virus, both IAVs and IBVs have a high mutation rate, ranging from $0.4 \times 10^{-3}$ to $2.0 \times 10^{-6}$ mutations per nucleotide per year, leading to genetic drift, because the RNA-dependent RNA polymerase (RdRp) lacks the 3′ to 5′ exonuclease proofreading capability. Antigenic drift is caused by evolutionary pressures that lead to an accumulation of mutations, particularly at the hemagglutinin (HA) and neuraminidase (NA) genes [16]. IBVs evolve more slowly compared with IAVs [17–19]. Of the IBV lineages, the Victoria lineage is undergoing more rapid evolution with greater positive selection pressure than the Yamagata lineage [17]. Structurally, the Yamagata virus binds to the host alpha-2,6-linked sialic acid receptors, whereas the Victoria virus binds to both alpha-2,3 and alpha-2,6 receptors, contributing to their differences in antigenic drift events [20,21].

Annual influenza vaccinations serve as a primary option for the prevention and control of influenza viruses [22,23]. Current vaccines are composed of two influenza A strains (H1N1 and H3N2) and either one (trivalent) or two (quadrivalent) influenza B strains [24]. Antigenic drift can cause low efficacy of seasonal influenza vaccines and necessitate annual vaccine reformulations. Due to antigenic drift, seasonal vaccine strains have been updated almost every year, especially during the past 10 years. Comparatively, IAVs have a much higher frequency of antigenic drift events than IBVs, as is reflected with the less frequent updates of the IBV component in the seasonal influenza vaccine.

Conventional serological tests such as hemagglutinin inhibition assays used in antigenic analyses for influenza vaccine strain selection require large virus quantities, and thus, the viruses often need to be isolated and propagated in Madin–Darby canine kidney (MDCK) cell lines or embryonated chicken eggs [25]. This can lead to cell- or embryonated egg-adapted mutations during viral propagation, particularly at the HA and NA surface glycoproteins [26–30]. Some prior studies have suggested that viruses propagated on MDCK cells have resulted in better antigenic matches than those grown on eggs [31]. During the seasons of 2012–2013 (VE = 49%, 43,000 deaths), 2016–2017 (VE = 40%, 38,000 deaths), and 2017–2018 (VE = 38%, 61,000 deaths), egg-adaptive mutations in the HA protein caused challenges in developing antigenically matched vaccine strain with circulating strains [32–36]. However, most of these studies on culture-adapted mutations were on IAVs, and the frequencies and implications of cell- or egg-adaptations in IBVs are yet to be fully understood.

In this study, we collected clinical samples from Columbia, Missouri during the 2019–2020 influenza season, particularly in the influenza epidemic months of February and March of 2020, to study the genetic and antigenic evolution of IBVs in humans and to assess whether passage of clinical samples in cells induces adaptive mutations across viral genes. Specifically, we performed serological analyses to characterize antigenic properties of the circulating influenza virus strains during this period and genomic analyses to determine the genetic changes that occurred during the influenza outbreaks during the 2019–2020 season in Missouri and whether culture-adapted mutations occurred during virus isolation.

2. Materials and Methods

Clinical samples: Nasopharyngeal swab specimens ($N = 368$) were collected from patients with influenza-like illness, who visited any of three urgent care facilities in Columbia, Missouri between February and March 2020. Samples were collected using a BD™ Universal Viral Transport Kit (3 mL, BD, Franklin Lakes, NJ, USA) and stored at $-70^\circ\text{C}$. 


Nucleic acid extraction: RNA extraction was performed using the MagMAX™ Pathogen RNA/DNA kit (Thermo Fisher Scientific Baltics, Vilnius, Lithuania) following the manufacturer’s instructions. The nucleic acid was resuspended in 90 μL of elution solution.

Quantitative reverse transcription polymerase chain reaction (qRT-PCR): The commercial kit AgPath-ID™ One-Step RT-PCR (Ambion, Austin, TX, USA) and CDC influenza A/B primers were used for screening the samples for influenza viruses in a single reaction in a QuantStudio™ 6 Flex Real time PCR System (Life Technologies, Paisley, UK). A 25 μL reaction containing 12.5 μL of the 2X reaction buffer, 0.5 μL of each primer (50 μM stock), or probe (10 μM stock) was used to reach a final concentration of 1 μL of the 25X RT-PCR enzyme mix. Additionally, 1.67 μL of detection enhancer were used.

Cell lines: Humanized MDCK cells (hCK) were kindly provided by Dr. Yoshihiro Kawaoka from the University of Wisconsin-Madison, USA. The MDCK-Siat1 cells were kindly provided by Dr. Mikhail Matrosovich from Philipps University, Germany. The MDCK-CCL34 cells were obtained from the Biodefense and Emerging Infections Research Repository. All cells were maintained until use at 37 °C under 5% CO2 in Dulbecco’s modified Eagle medium (Gibco DMEM; Thermo Fisher Scientific, Waltham, MA, USA) supplemented with 10% fetal bovine serum (Gibco; Thermo Fisher Scientific Baltics, Vilnius, Lithuania). These three cell lines were selected, because hCK has overexpression of α-2,6-sialoglycans and almost completely deleted expression of α-2,3-sialoglycans, and hCK has been demonstrated to be a more efficient cell line for influenza B, A/H1N1pdm, and even more efficient for A/H3N2 virus isolation and propagation than other MDCK cells for influenza viruses [37]. MDCK-Siat1 has overexpression of α-2,6-sialyltransferase but a marginal decrease in α-2,3-sialylation [38]. Similar to hCK cells, Siat-1 has been shown to improve isolation rates over conventional MDCK cell lines (CCL34) in influenza B, A/H1N1, and especially in A/H3N2 viruses [39]. Finally, MDCK-CCL34 cells express higher levels of α-2,3-syaloglycans compared with α-2,6-syaloglycans and are widely used for influenza virus isolation and propagation, and they exhibit relatively low levels of both α-2,3- and α-2,6-sialic acids with slightly higher levels of α-2,3-sialylation compared with α-2,6-sialylation [40].

Virus isolation: Clinical samples were diluted 1:10 in Opti-MEM media containing a 1X final dilution of antibiotic–antimycotic solution for 30 min at 4 °C. A total volume of 0.5 mL was inoculated in T25 flasks containing confluent cells in Opti-MEM media with antibiotics–antimycotics at 35 °C for 3 to 4 days until 75% cytopathic effect (CPE) was observed. Supernatant was collected and centrifuged at 4000 revolutions per minute, and the hemagglutination capacity was tested using 0.5% solution of turkey erythrocytes.

Ferret antisera: The following sera were obtained from the Influenza Research Reagent Resources: FR-1617 B/Colorado/6/2017, FR-1081 B/Nevada/03/2011, FR-1305 B/Texas/02/2013, FR-392 B/Brisbane/60/2008, B/Washington/02/2019 (egg), FR-810 B/Wisconsin/1/2010.

Hemagglutination inhibition (HAI) assays: Isolated viruses were characterized antigenically following previously described protocols [41].

Antigenic cartography: HAI results were visualized by AntigenMap (https://sysbio.missouri.edu/software/AntigenMap/, Accessed 19 July 2021) [42–44]. A titer of 20 was used as the low reactor threshold and a low rank (rank = 2) matrix completion was used to minimize noise from the HAI data. A single unit on the antigenic map represents a log₂ unit in HAI titers.

Next generation sequencing: Whole genome RT-PCR amplification was conducted using 12 pairs of primers [45–47]. Amplicon libraries were prepared using the Nextera DNA Flex Library Prep kit, then sequenced with MiSeq Reagent Kit v3 (600 cycles) and MiSeq sequencing system (Illumina, San Diego, CA, USA). The resulting paired-end reads were analyzed for quality, mapped to a reference genome (B/Utah/17/2020, accession: MT499571-MT499578), and assembled using CLC Genomics Workbench v21.0.3 (Qiagen,
Pairwise analyses were conducted using CutAdapt v3.4 (National Bioinformatics Infrastructure Sweden, Uppsala, Sweden) [48], Bowtie2 v2.4.4 (Johns Hopkins University, Baltimore, Maryland, USA) [49], and samtools v1.13 (https://github.com/samtools/, Accessed 4 May 2021), and polymorphisms were identified using pysamstats v1.1.2 (https://github.com/alimanfoo/pysamstats, Accessed 4 May 2021) and confirmed using the Integrated Genomics Viewer (https://software.broadinstitute.org/software/, Accessed 10 June 2021). Cell-adapted mutations were verified manually using BioEdit v7.2.5 [50]. Total reads ranged from 142,724 to 2,161,170 for each sequence, with an average of 1,092,806 reads (standard deviation (SD) = 751,073). The average number of reads for clinical samples was 1,725,216 (SD = 239,826), and the average number of reads for the isolates was 460,396 (SD = 520,405). GenBank accession numbers for these 31 IBV isolates are MZ951177–MZ951671.

Phylogenetic analyses: The basic local alignment search tool (BLAST) was used to identify sequences with the closest identities to each sample, and the top 20 sequences across all segments were included in each phylogeny. Reference genomes from the Victoria lineage (B/Rhode Island/01/2019, EPI_ISL_409247; B/Nigeria/3352/2018, EPI_ISL_330532; B/Missouri/12/2018, EPI_ISL_309120; B/Hong Kong/286/2017, EPI_ISL_276541; B/Florida/78/2015, EPI_ISL_207237; B/Brisbane/60/2008, EPI_ISL_246494; B/Brisbane/46/2015, EPI_ISL_219054; B/Hong Kong/269/2017, EPI_ISL_276540; B/Florida/103/2016, EPI_ISL_248973; B/New Jersey/1/2021, EPI_ISL_246881; B/Colorado/6/2017, EPI_ISL_277231; B/Nevada/03/2011, EPI_ISL_94747; B/Florida/78/2015, EPI_ISL_207237; B/Texas/02/2013, EPI_ISL_139913; B/Washington/02/2019, EPI_ISL_341131) and the Yamagata lineage (B/Phuket/3073/2013, EPI_ISL_517766; B/Texas/81/2016, EPI_ISL_237464; B/New Hampshire/01/2018, EPI_ISL_296628) were also included in the phylogenetic trees for each segment. These sequences were downloaded from the Global Initiative on Sharing Avian Influenza Data (GISAID) consortium EpiFlu database (http://www.gisaid.org, Accessed 2 July 2021).

All selected sequences were aligned using MUltiple Sequence Comparison by Log-Expectation (MUSCLE8, European Bioinformatics Institute, Cambridgeshire, United Kingdom). Phylogenetic analyses were performed with BEAST2 (https://www.beast2.org/, Accessed 22 July 2020) with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Tracer v1.7.1 was used to evaluate phylogenetic quality using a cutoff of 200 for the effective sample size (ESS) to assess convergence. The consensus tree was generated using TreeAnnotator v2.6.3.0 and visualized with FigTree v1.4.4 (http://tree.bio.ed.ac.uk/software/figtree/, Accessed 2 February 2021).

### 3. Results

#### 3.1. Virus Isolation in MDCK-hCK Cell Lines Showed the Highest Virus Recovery Rate

Of the 368 samples, 146 (39.7%) tested positive for influenza with 38 samples (10.3%) positive for IBV and 108 (29.3%) for IAV by qRT-PCR. Thirty-one of the IBV samples were positive to virus isolation in hCK cells. Other MDCK cell lines such as CCL34 and SIAT1 were also used to compare cell sensitivity to virus isolation. The MDCK-hCK cell lines showed the highest performance for both IAV and IBV following the first passage of the viruses with an isolation rate of 87.96% and 81.58%, respectively, compared with the CCL34 (42.12% and 57.89%, respectively) and SIAT1 (71.30% and 57.89%, respectively) cell lines (Table 1).

| Type      | Samples | CCL34 P1 | CCL34 P2 | SIAT1 P1 | hCK P1   |
|-----------|---------|----------|----------|----------|----------|
| IAV (H1)  | 108     | 39 (42.12%) | 76 (70.37%) | 77 (71.30%) | 95 (87.96%) |
| IBV       | 38      | 22 (57.89%) | 24 (63.16%) | 22 (57.89%) | 31 (81.58%) |
| Total     | 146     | 61 (41.78%) | 100 (68.49%) | 99 (67.80%) | 127 (86.98%) |
3.2. Culture-Adapted Mutations Were Identified in IBVs

To identify culture-adapted mutations, we sequenced 38 IBVs from the IBV-positive clinical swab samples. Of the 38 paired swab and isolate (hCK derived; first passage) samples subjected to MiSeq sequencing, 31 paired samples yielded complete genomic data and were included in the analysis to determine hCK cell-adapted mutations and polymorphisms. The genomes for the other seven pairs were either incomplete or with a low coverage and thus not included in our subsequent analyses. From these 31 paired samples, we identified 13 cell-adapted mutations in total among nine viruses, ranging from one to three mutations in each virus, and each of these mutations was unique to each virus. We detected eight nonsynonymous mutations among these paired samples that have not been previously published (HA-R581C; PB2-T46S, PB2-V98G; PB1-E112G, PB1-R350G, PB1-V501A, PB1-L674Q; and NP-E140A), six of which were from the RNA polymerase complex (Figure 1) and five synonymous mutations (Table 2). Polymorphisms were also identified in PB2-T293G, PB2-T2097C, PB1-A255G, PB1-A1048G, PB1-T1502C, and NP-A419C (Table 2).

![Figure 1](image_url). Cell-adapted mutations in the RNA polymerase complex of influenza B virus isolates (2019–2020). Mutations at the RNA polymerase complex (PB1, PB2, PA) are visualized in PyMOL using the 6QCT protein structure from the Protein Data Bank (PDB, https://www.rcsb.org/, Accessed 27 June 2021). Green represents PA, blue represents PB1, purple represents PB2.

| Sample | Protein | Position | Clinical Swab (Nucleotide) | Isolate (Nucleotide) | Amino Acid |
|--------|---------|----------|---------------------------|---------------------|------------|
|        |         |          | Swab (Proportion Reads) x | Reads (Total) b     | % Reads at Position c | Isolate (Proportion Reads) x | Reads (Total) b | % Reads at Position c |
| CS20N01| PB2     | 136      | A (0.99)/T (0.01)          | 1,574,718           | 26224 1.67  | T (0.99)          | 1,149,216       | 17,538 1.53  |
|        | PB2     | 2181     | T (0.98)/C (0.02)          | 1,574,718           | 7075 0.45  | C (0.99)/T (0.01) | 1,149,216       | 54,788 4.77  |
|        | PB1     | 2160     | G (1.00)                   | 1,574,718           | 47919 3.04  | A (1.00)         | 1,149,216       | 43,524 3.79  |
| CS20N02| HA      | 1741     | C (0.79)/T (0.16)/N (0.05)| 1,588,004           | 45 0.00  | T (1.00)         | 182,972         | 14,112 7.71  |
|        | PB1     | 2016     | T (0.97)/A (0.03)          | 1,940,088           | 39 0.00  | C (0.50)/T (0.46)/A (0.04) | 1,468,024       | 42,152 2.87  |
|        | PB1     | 2020     | T (1.00)                   | 1,940,088           | 41 0.00  | A (0.53)         | 1,468,024       | 43,968 3.00  |
| CS20N09| NP      | 419      | A (0.96)/N (0.03)/T (0.01)| 2,161,170           | 1615 0.07  | C (0.76)/A (0.24) | 177,412         | 4283 2.41   |

Table 2. Cell-adapted mutations in influenza B virus isolates (2019–2020).

* indicates a nonsynonymous substitution.
In addition to our samples, we analyzed all Victoria lineage IBV samples available on GISAID from an original specimen and at least one cell or egg sequence to determine whether culture-adapted mutations also occurred among samples submitted to a public database. Passage information is not consistently reported in the GISAID strains and, thus, was not included in the analysis. Of the initial 91,884 distinct and complete Victoria lineage sequences on GISAID as of 2 July 2021, 539 strains contained at least one clinical sample–isolate pair. Pairwise comparison of these strains revealed multiple additional cell- and egg-adapted mutations (Table 3). Three distinct mutations were identified at PB2, one at PB1, two at PA, two at NS1, two at NP, two at NEP, one at NB, 11 at NA, and eight at HA. Notably, on the HA protein, 14 strains contained a cell- or egg-adapted mutation at position 212 and six strains contained a mutation at position 214 (Figure 2). Additionally, of the complete Victoria IBV sequences with paired isolates, 10.3% contained culture-adapted mutations (6.7% cell, \( N = 537 \); 38.3% egg, \( N = 60 \)).

Table 3. Cell- and egg-adapted amino acid mutations of Victoria lineage IBVs from the paired clinical isolate samples available at the GISAID.

| Protein | Strain | Position | Original | Cell | Egg |
|---------|--------|----------|----------|------|-----|
| HA      | B/North Carolina/17/2019/EPI1693897 | 143 | E | K |
|         | B/Florida/78/2015/EPI721064 | 156 | G | G |
|         | B/Mozambique/413/2016/EPI854592 | 179 | D | Y |
|         | B/Pennsylvania/19/2016/EPI807677 | 212 | N | N |
|         | B/Michigan/18/2016/EPI807685 | 212 | N | T |
|         | B/Indiana/08/2016/EPI807669 | 212 | N | N |
|         | B/Florida/94/2016/EPI872715 | 212 | N | N |
|         | B/Hawaii/26/2018/EPI1315023 | 212 | N | D |
|         | B/Florida/79/2015/EPI745107 | 212 | N | Y |
|         | B/Utah/18/2018/EPI11249578 | 212 | N | T |
|         | B/Montana/34/2018/EPI1386557 | 212 | N | D |
|         | B/Iowa/14/2017/EPI1025486 | 212 | T | T |
|         | B/Guatemala/35/2018/EPI11249639 | 212 | N | S |
|         | B/Guatemala/109/2018/EPI1312932 | 212 | N | S |
|         | B/Florida/97/2017/EPI1138021 | 212 | N | K |
| B/Alabama/02/2017/EPI980739 | 212 | N | N | S |
| B/Lebanon/16/2020/EPI1840705 | 212 | N | N | S |
| B/Pennsylvania/60/2016/EPI892343 | 214 | T | I | |
| B/Hawaii/55/2017/EPI1089741 | 214 | T | N | |
| B/Costa Rica/5521/2016/EPI908149 | 214 | T | A | |
| B/El Salvador/697/2018/EPI1357148 | 214 | T | A | |
| B/Louisiana/16/2019/EPI1574236 | 214 | T | T | N |
| B/Hong Kong/269/2017/EPI1141721 | 214 | T | T | I |
| B/Minnesota/39/2017/EPI1026300 | 236 | I | T | |
| B/Ulaabaatar/1868/2017/EPI1089764 | 433 | N | D | |
| B/Ulaabaatar/1834/2017/EPI1053102 | 433 | D | N | |
| B/Ulaabaatar/1767/2017/EPI1053094 | 433 | N | D | |
| B/Wisconsin/22/2018/EPI1312153 | 528 | D | G | |

| NA | B/North Carolina/17/2019/EPI1693896 | 19 | L | I |
| B/North Carolina/16/2019/EPI1664425 | 19 | I | L |
| B/Wisconsin/45/2016/EPI834520 | 154 | L | K |
| B/North Carolina/17/2019/EPI1693896 | 284 | S | G |
| B/North Carolina/16/2019/EPI1664425 | 284 | G | S |
| B/Ulaabaatar/1868/2017/EPI1089763 | 310 | T | S |
| B/Nevada/23/2019/EPI1688225 | 321 | D | N |
| B/Colombia/9962/2016/EPI908031 | 391 | D | E |
| B/Bangladesh/8018/2016/EPI880659 | 395 | A | D |
| B/North Carolina/17/2019/EPI1693896 | 400 | V | I |
| B/Arizona/83/2017/EPI1147826 | 401 | M | V |
| B/California/18/2017/EPI1009716 | 435 | G | R |
| B/Guyane/310/2020/EPI1807176 | 450 | M | M | V |

| NB | B/North Carolina/17/2019/EPI1693896 | 21 | I | N |
| B/North Carolina/16/2019/EPI1664425 | 21 | N | I |

| NEP | B/Oregon/11/2020/EPI1840666 | 30 | S | S | L |
| B/Iowa/14/2017/EPI1011596 | 79 | D | D | N |

| NP | B/Bangladesh/5006/2017/EPI1094384 | 33 | R | K |
| B/Ulaabaatar/1868/2017/EPI1089757 | 239 | A | T |

| NS1 | B/North Carolina/17/2019/EPI1693891 | 62 | T | A |
| B/North Carolina/16/2019/EPI1664420 | 62 | A | T |
| B/Ulaabaatar/1868/2017/EPI1089758 | 115 | T | M |
| B/Ulaabaatar/1834/2017/EPI1053096 | 115 | M | T |
| B/Ulaabaatar/1767/2017/EPI1089797 | 115 | T | M |

| PA | B/Bangladesh/8018/2016/EPI880656 | 57 | E | K |
| B/New Jersey/09/2017/EPI1011540 | 393 | I | I | T |

| PB1 | B/North Carolina/17/2019/EPI1693895 | 378 | L | S |
| B/North Carolina/16/2019/EPI1664424 | 378 | S | L |

| PB2 | B/Florida/97/2017/EPI1138018 | 263 | V | I |
| B/Macedonia/421/2020/EPI1806732 | 379 | S | S | C |
| B/Bangladesh/5006/2017/EPI1094388 | 391 | A | T |

Blank cells under cell and egg mutation columns indicate that no sequence was available for analysis.
Figure 2. Cell- and egg-adapted mutations at the 212 and 214 positions within the receptor binding domain and antibody binding site on the HA protein visualized in PyMOL using the 4FQM protein structure from the Protein Data Bank (PDB, https://www.rcsb.org/, Accessed 27 June 2021).

3.3. Antigenic and Genetic Diversity of Missourian IBV in the 2019–2020 Influenza Season

Phylogenetic analyses showed that the IBVs circulating during the 2019–2020 influenza season belonged to the Victoria lineage, all in the clade V1A.3-3DEL. Additionally, all samples contained three amino acid deletions (3DEL) at positions 162–164 of the HA protein sequences when compared with B/Brisbane/60/2008 (EPI_ISL_246494) as shown in Figure 3. Of note, the IBV vaccine strain used in the 2019–2020 influenza season (B/Colorado/06/2017) had two amino acid deletions (2DEL).
Figure 3. Phylogenetic tree of the HA segment of IBV. Missouri samples from the 2019–2020 season that were collected for this study are labeled in red. Recommended vaccine strains are labeled in blue. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability ≥0.70 were labeled. The nomenclature of the genetic clades is adapted from that reported by the World Health Organization [51].
### 3.4. Antigenic Diversity of Missourian IBVs in the 2019–2020 Influenza Season

To test the antigenic diversity of the IBVs, hemagglutination inhibition assays and antigenic cartography were performed for 32 Missouri isolates and 11 reference strains against seven ferret sera for six representative IBVs (B/Colorado/6/2017, B/Nevada/03/2011, B/Texas/02/2013, B/Brisbane/60/2008, B/Washington/02/2019, and B/Wisconsin/1/2010). The sera for egg and cell isolates of B/Washington/02/2019 were included (Table 4). Antigenic analyses suggested all Missourian isolates reacted well to the Victoria lineage IBV-derived ferret sera but poorly with the Yamagata lineage IBV-derived ferret sera. Of all ferret sera, all Missourian isolates had the highest HI titer against B/Washington/02/2019, which also has the 3DEL, but 4- to 8-fold lower titers against B/Colorado/06/2017, which is the vaccine strain used during the 2019–2020 season and contains the 2DEL [51]. Of note, these Missouri isolates reacted better to the ferret antisera against the cell isolate than the egg isolate of B/Washington/02/2019.

### Table 4. Hemagglutination inhibition results of influenza B virus isolates (2019–2020) using ferret antisera.

| Virus * | Clade b | Ferret Reference Sera c |
|---------|---------|-------------------------|
|         | [B/Nevada/03/2011] | [B/Texas/02/2013] | [B/Brisbane/60/2008] | [B/Colorado/06/2017] | [B/Washington/02/2019 (egg)] | [B/Washington/02/2019 (Cell)] | [B/Wisconsin/1/2010] |
| B/Nevada/03/2011 | V1A | 1280 | 320 | 640 | 20 | 160 | 80 | 10 |
| B/New Jersey/01/2012 | V1A | 1280 | 640 | 320 | 40 | 40 | 40 | <10 |
| B/Texas/02/2013 | V1A | 320 | 320 | 160 | 40 | 40 | 40 | <10 |
| B/Florida/78/2015 | V1A | 1280 | 1280 | 640 | 80 | 80 | 80 | 20 |
| B/Brisbane/60/2018 | V1A | 640 | 320 | 1280 | 40 | 160 | 40 | 10 |
| B/Colorado/06/2017 | V1A-2DEL | V1A-3DEL | 40 | 20 | 0 | 320 | 40 | 160 | <10 |
| B/Hong Kong/286/2017 | V1A-3-3DEL | V1A-3-3DEL | 320 | 160 | 160 | 80 | 320 | 160 | 10 |
| B/Washington/02/2019 (egg) | V1A-3-3DEL | V1A-3-3DEL | 320 | 80 | 80 | 80 | 640 | 640 | <10 |
| B/Washington/02/2019 (cell) | V1A-3-3DEL | V1A-3-3DEL | 80 | 40 | 10 | 80 | 320 | 640 | <10 |
| B/Rhode Island/01/2019 | V1A-3-3DEL | V1A-3-3DEL | 320 | 160 | 320 | 40 | 320 | 320 | 10 |
| B/Phuket/23/2013 | Y3 | 10 | <10 | <10 | <10 | <10 | <10 | 1280 | |

* = Virus isolate.
  
b = Clade (3DEL, 2DEL, V1A).
Antigenic cartography showed that these Missouri isolates have an average antigenic distance of $1.60 \pm 0.22$ units from B/Colorado/06/2017 (Figure 4), suggesting these isolates are antigenically distinct from the vaccine used in this season. On the other hand, the average antigenic distances within Missourian isolates were $0.40 \pm 0.21$ units (Figure 4). Each unit represents a 2-fold change in hemagglutination inhibition titers.

![Antigenic cartography of hemagglutination inhibition results of influenza B virus isolates (2019–2020). Each grid represents one unit, a 2-fold change in hemagglutination inhibition titers. The vaccine component strains are labeled in cyan. Strains from the 2019–2020 season in Missouri are shown in red. Additional reference strains are blue. Antigenic cartography was constructed using AntigenMap (https://sysbio.missouri.edu/software/AntigenMap/, Accessed 19 July 2021) [42–44].](image)

**Figure 4.** Antigenic cartography of hemagglutination inhibition results of influenza B virus isolates (2019–2020). Each grid represents one unit, a 2-fold change in hemagglutination inhibition titers. The vaccine component strains are labeled in cyan. Strains from the 2019–2020 season in Missouri are shown in red. Additional reference strains are blue. Antigenic cartography was constructed using AntigenMap (https://sysbio.missouri.edu/software/AntigenMap/, Accessed 19 July 2021) [42–44].

### 4. Discussion

Influenza B viruses are endemic around the world, leading the World Health Organization (WHO) to recommend the inclusion of Yamagata and Victoria strains in the seasonal influenza vaccine [24]. While the use of clinical swab samples for vaccine development would be ideal in minimizing cell- and egg-adapted mutations, this is not yet feasible. Previous studies have identified MDCK cells to promote the most viral growth when compared with Vero cells, both of which have been used frequently in influenza studies [25]. We extended this finding by testing our sample growth on various MDCK cell lines, including hCK, CCL34, and SIAT1 and found that the hCK cell lines showed the highest performance for both IAV and IBV following the first passage, consistent with findings by

| Strain       | HA Titer | NA Titer | HA Titer | NA Titer | HA Titer | NA Titer | HA Titer | NA Titer |
|--------------|----------|----------|----------|----------|----------|----------|----------|----------|
| CS20N294     | 80       | 40       | 10       | 80       | 320      | 640      | <10      |          |
| CS20N349     | 40       | 20       | 10       | 80       | 320      | 640      | <10      |          |
| CS20N351     | 80       | 40       | 10       | 160      | 640      | 1280     | <10      |          |

*The viruses underlined are the vaccine strains used in the 2019–2020 influenza season. Passage 2 viruses for all Missouri isolates were used in the serological assays. The 3DEL noted in clades occurs at positions 162–164 on the HA protein. The homologous hemagglutination inhibition titers were marked in bold.*
Takada et al. [37]. Higher replication efficiency results in fewer required passages and likely fewer cell adaptations. This finding suggests hCK cell lines are an effective platform for virus isolation and propagation and potentially allow for fewer culture adaptations during vaccine development.

During the 2019–2020 influenza season in the US, influenza caused 400,000 hospitalizations and 22,000 deaths [52]. Notably, influenza B and predominantly the B/Victoria lineage (>99% of IBVs) emerged unusually early in the season and made up approximately half of all influenza cases compared with 10–30% in prior years [53–55]. Of the IBVs, 97% belonged to subclade V1A.3 as compared with the B/Victoria vaccine reference strain (B/Colorado/06/2017) recommended by the WHO for the 2019–2020 season [54]. The B/Colorado/06/2017-like strain recommended by the WHO contains 2DEL at 162–163, which confers reduced hemagglutinin inhibition cross-reactivity against viruses with 3DEL [56]. However, most of the circulating strains (V1A.3 subclade) contained three amino acid deletions at positions 162–164 of the HA protein sequence [57]. The overall VE for the 2019–2020 season was 46% [57]. Similarly, our study showed that all influenza B strains that were collected in Columbia, Missouri belonged to the Victoria lineage, contained the 3DEL, and were antigenically distinct from the vaccine strain used during this season (Figure 4). As the primary influenza epidemic occurred between February and March of 2020 in this influenza season, the sampling period was somewhat limited, and the inclusion of samples outside of the influenza peak time period may have provided additional insights to the antigenic and genetic evolution of IBVs in this season.

In addition to the antigenic mismatch between the vaccine and circulating influenza B strain due to an unusually early emergence and predominance of IBVs, we demonstrated that 13 hCK cell-adapted mutations were present in the first passage of Missouri strains during the 2019–2020 season. In our samples, no mutations were identified on the antibody binding sites of HA and NA proteins, although a single adaptation was identified at the HA2 protein. However, multiple adaptations in the RdRp were generated during the isolation and propagation of the original wild-type viruses. The effects of mutations in the RdRp have been shown to contribute towards influenza A pathogenicity [58]. However, this process is not well studied in influenza B. The mutations that were identified in the RdRp have not been previously reported, and, as future studies, we plan to explore the antigenic and pathogenic implications of cell-adapted mutations in the RdRp. Further exploration the IBV genomic sequences for the isolates in the GISAID database revealed no or low frequencies for the nonsynonymous mutation site mutations that we observed in the hCK-derived isolates (for example, NP-140 (2 of 11,110 GISAID sequences; 2/11,110), PB2-98 (9/11,121), PB1-350 (1/11,122), and HA-581 (0/11,132)).

One potential limitation for this study is that we performed analyses for the paired swab only with hCK-derived isolates but not Stat1-, CCL34-, or egg-derived isolates. In addition, the number of paired swab clinical samples \( n = 31 \) included in this study is relatively small. Nevertheless, through analyses with those paired samples in the public database, when the entirety of the available Victoria lineage pairs from GISAID were analyzed, cell- and egg-adapted mutations were identified in the majority of protein segments (Table 3; Figure 2). Egg-adapted mutations occurred more frequently than cell-adapted mutations. Multiple reoccurrences appeared at the HA 212 and 214 positions (Figure 2), of which site 212 has been identified as a site of N-glycosylation that undergoes positive selection and frequent mutation (PDB 4FQM) [59]. Cell-adapted mutations at position 212 on HA have also been shown to decrease acid stability in H3N2 viruses in ferret models [60]. Four antigenic sites have been identified in IBVs and are located on the 120-, 150-, and 160-loops and the 190-helix [61]. The 212 and 214 positions reside on the receptor binding domain (RBD) and 190-helix of HA (Figure 2). Additionally, one sequence contained a cell-adapted mutation in the 120-loop (E143K) (Appendix A). Of interest, this observation is different from the mutations in the Missouri swab and hCK samples, which were only present in the internal genes, particularly in the RNA polymerase complex. The
HA protein plays a major role in host receptor binding and membrane fusion, and mutations at the RBD and antibody binding sites may affect the antigenic characteristics of and immune responses against IBVs. Culture-adapted mutations during vaccine development at this gene may lead to antigenic variants from the circulating virus.

Analysis of the GISAID database confirms our findings that cell culture-adapted mutations occur in Victoria lineage IBVs, though infrequently. We also found that egg adaptations occurred 5.7 times more often than cell-adapted mutations, and such adapted mutations were reported to be under positive selection due to egg culture [62]. The adapted mutations in HA, particularly those in or close to receptor binding sites, may alter virus receptor binding properties to increase replication in eggs [63]. Similar to our findings, Park et al. showed that H3N2 viruses cultured in eggs had more frequent antigenic mutations in the HA protein than in cells, where they found no antigenic mutations [64]. These culture-adapted mutations have been suggested to compromise vaccine effectiveness for IAVs in humans during multiple influenza seasons [32–36] and remain a concern. Nevertheless, our study showed that using cell culture can help mitigate these challenges.

In summary, our study suggests that cell culture generates less frequent culture-adapted mutations than egg culture for isolating IBVs and that, of the commonly used MDCK cell lines tested, hCK is the most effective platform for IBV isolation. Additionally, culture-adapted mutations may occur in IBVs even after a single passage in hCK cells, and these mutations could generate biases on our understanding of the viral characteristics in the human population. As culture-adapted mutations may affect subsequent virus studies and vaccine development, the knowledge from this study may help optimize strategies for influenza surveillance, vaccine strain selection, and vaccine development.

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Informed Consent Statement: Patient consent was waived due to the de-identified nature of the aggregated descriptive data.

Data Availability Statement: The data presented in this study are openly available in the GenBank (accession numbers: MZ951177–MZ951671).

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Appendix A

Full phylogenetic trees of each IBV protein segment. Missouri 2019–2020 sequences are shown in red and vaccine strains are shown in blue. Each cluster is grouped within colored boxes. The phylogenetic analyses were performed by using BEAST2 with a Ha-
Figure A1. PB2 Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A2. PB1 Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A3. PA Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A4. HA Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A5. NP Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A6. NA Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A7. MP Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
Figure A8. NS Tree. The phylogenetic analyses were performed by using BEAST2 with a Hasegawa–Kishino–Yano (HKY) substitution empirical model, strict clock model, and coalescent constant population prior. Branches with a posterior probability >0.70 were labeled.
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