Mutation E48K in PB1 polymerase subunit improves stability of a candidate live attenuated influenza B virus vaccine.

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Abstract: Influenza B virus (IBV) is a major respiratory pathogen of humans, particularly in the elderly and children and vaccines are the most effective way to control it. In previous work, incorporation of two mutations (E580G, S660A) along with the addition of a HA epitope tag in the PB1 segment of B/Brisbane/60/2008 (B/Bris) resulted in an attenuated strain that was safe and effective as a live attenuated vaccine. A third attempted mutation (K391E) in PB1 was not always stable. Interestingly, viruses that maintained the K391E mutation were associated with the mutation E48K.

To explore the contribution of the E48K mutation for stability of the K391E mutation, a vaccine candidate was generated by inserting both mutations along with attenuating mutations E580G and S660A in PB1 of B/Bris (B/Bris PB1att 4M). Serial passage of the B/Bris PB1att 4M vaccine candidate in eggs and MDCK indicated high stability. In silico structural analysis revealed a potential interaction between amino acids at positions 48 and 391. In mice, B/Bris PB1att 4M was safe and provided complete protection against homologous challenge. These results confirm the compensatory effect of mutation E48K to stabilize the K391E mutation, resulting in a safer, yet still protective, IBV LAIV vaccine.

Keywords: Influenza B; LAIV; PB1; vaccine stability

Introduction

Influenza B virus (IBV) is an enveloped virus from the Orthomyxoviridae family with a negative-sense single stranded RNA genome [1]. The IBV genome consists of 8 segments, encoding at least 11 proteins [2,3]. IBV causes seasonal respiratory disease epidemics in humans. Although IBV can affect all age groups, studies have shown that individuals under the age of 18 are more susceptible to infection [4]. IBV infection can be severe in children and has been associated with a high proportion of pediatric deaths related to influenza [5,6]. In most seasons, IBV infections are less common than influenza A virus (IAV) infections. However, the 2019-20 season showed an unusual increased number of IBV infections, particularly early in the season, accounting for approximately half of influenza infections in the U.S. (CDC Weekly U.S. Influenza Surveillance Report). IBV strains are classified into two antigenically distinct lineages, the B/Victoria (B/Vic) and the B/Yamagata (B/Yam) lineages [7, 8]. The two lineages co-circulate globally, show minimum to no antibody cross-reactivity between them and continue to evolve due to antigenic drift [7-9]. Thus, the development of effective vaccines against IBV is important to reduce disease burden, especially in vulnerable age groups.

Vaccines for influenza viruses mostly rely on antibody responses against the HA protein [10,11]. Most FDA-approved vaccines available against influenza are quadrivalent vaccines which incorporate strains from both IBV lineages along with 2 subtypes of IAV (H3N2, H1N1). Trivalent vaccines that only contain one of the IBV lineages are also available [7,12-14]. In the United States, seasonal vaccines are available in the form of either recombinant proteins, inactivated influenza (IIV) vaccines and/or live attenuated influenza virus (LAIV) vaccines. It is known that LAIVs can mimic a natural influenza infection, which can stimulate both mucosal and cellular immunity [15-17]. However, less than optimal vaccine effectiveness of the FluMist LAIV in the 2013-2014 and 2015-2016 seasons, particularly in young children, prompted CDC to temporarily recommend against its use...
in 2016 [18]. Although CDC has since reinstated the recommendation for use of FluMist, this incident highlights the importance of developing other LAIV alternatives that could be more efficacious.

Previous work in our lab showed that incorporation of three specific mutations (K391E, D581G, A661T) in addition of an HA epitope tag (HA tag) at the C-terminus of the PB1 protein segment and a single mutation (N265S) into the PB2 protein segment of IAVs results in a temperature sensitive (ts) phenotype in vitro, attenuates (att) IAVs in vivo, and provides effective protection against aggressive virus challenge in a variety of mammalian and avian animal models [19-21]. Following a similar approach, we previously generated a novel IBV att strain by introducing analogous mutations in the PB1 (K391E, E580G and S660A with C-terminus HA tag) and PB2 (K267S) genes of the B/Brisbane/60/2008 (B/Bris) strain from the B/Vic lineage [22]. Although the PB2 K267S mutation could not be rescued and the PB1 K391E mutation was unstable, the remaining mutations in the PB1 (E580G, S660A, HA tag) and PB2 (K267S) genes of the B/Brisbane/60/2008 (B/Bris) strain from the B/Vic lineage [22]. Although the PB2 K267S mutation could not be rescued and the PB1 K391E mutation was unstable, the remaining mutations in the PB1 (E580G, S660A, HA tag) were stably maintained for at least 20 passages in Madin-Darby canine kidney (MDCK) cells or 15 passages in specific pathogen free (SPF) eggs. Furthermore, these modifications resulted in attenuation of the B/Bris in vitro and in vivo, and the resulting B/Bris PB1att strain protected mice against both homologous and heterologous challenge [22]. During stability tests, it was noted that the B/Bris PB1att that was serially passaged in MDCK cells retained the K391E mutation but showed an additional mutation in PB1, E48K. We hypothesized that E48K serves as a compensatory mutation to stabilize the K391E mutation while maintaining the attenuated phenotype.

To test this effect, we generated a new strain (B/Bris PB1att 4M) by incorporating E48K and K391E along the E580G, S660A mutations into the PB1 segment. The new B/Bris PB1att 4M strain, in which the HA tag was omitted, showed high stability in both MDCK cells and SPF eggs. In contrast, a strain containing the PB1 K391E (but not the E48K mutation, B/Bris PB1 3M) in the B/Bris PB1att background quickly acquired the E48K mutation after serial passage in eggs, further highlighting its importance as a compensatory mutation. Vaccination studies in mice showed that B/Bris PB1att 4M strain was safe, immunogenic, and provided complete protection against homologous challenge.

Material & Methods

Tissue culture cells and embryonated chicken eggs.

Human embryonic kidney 293T cells, Madin-Darby canine kidney (MDCK) cells, and specific pathogen free (SPF) embryonated chicken eggs (Charles River, Norwich, CT) were used for generation of viruses through reverse genetics, virus amplification and/or serial passages. Cells were cultured at 37°C at a 5% CO2 environment in Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% fetal bovine serum (FBS), 1% of L-glutamine and 1% of antibiotic/antimycotic solution (Sigma-Aldrich St. Louis, MO).

Cloning and site directed mutagenesis of PB1 segment.

The reverse genetics plasmid encoding the PB1 segment of the B/Bris att strain containing three amino acid mutations (K391E, E580G, and S660A) has been previously described [22]. Subsequently, the E48K mutation was introduced into the PB1 gene segment by site-directed mutagenesis with the QuikChange II XL kit (Agilent, Santa Clara, CA) to generate the PB1 4M plasmid. Plasmids were grown in One Shot TOP10 E. coli cells (Invitrogen, Carlsbad, CA) in the presence of ampicillin and stocks were prepared via the HiSpeed® Plasmid Maxi Kit (Qiagen, Germantown, MD). Plasmids were sequenced by Sanger sequencing with a specific set of primers (available upon request) to corroborate the mutations in the PB1 4M plasmid and to rule out spurious mutations.

B/Bris PB1att 4M strain generated by reverse genetics.

The B/Bris PB1att 4M (E48K, K391E, E580G, and S660A) virus was generated by reverse-genetics as previously described [22] in co-cultured 293T and MDCK cells. Transfected cells were incubated at 35°C for 24 h, and then the transfection media was replaced.
with Opti-MEM (Thermo-fisher Scientific, Waltham MA) containing 1 μg/ml tosylsulfonyl phenylalanyl chloromethyl ketone (TPCK)-trypsin (Worthington Biochemicals, Lakewood, NJ) and 1% of antibiotic/antimycotic solution (Sigma-Aldrich St. Louis, MO). The B/Bris PB1 3M (K91E, E580G, S660A, and the HA tag) virus was rescued in our previous study [22]. Rescued viruses were amplified once in MDCK cells at 33°C for 72h (passage 1 [P1]). The B/Bris PB1att 4M virus for the animal study was grown from the P1 stock in SPF eggs for 72 h at 33°C (egg passage 1 [E1]). Viral stocks were titrated by 50% tissue culture infectious dose (TCID₅₀) or 50% egg infectious dose (EID₅₀) using the Reed and Muench method [23], and sequenced by Sanger sequencing or next generation sequencing (NGS) to check for the presence of the inserted mutations and integrity of the strain.

Sequencing.

Sanger sequencing (Psomagen, Rockville, MD) was performed between the 5’-UTR (Untranslated Region) and the portions of the PB1 segment (up to ~2000bp) which includes the 4 mutations. For Sanger sequencing, viral RNA was extracted from cell supernatants or allantoic fluids via the QIAamp Viral RNA Mini Kit (Qiagen, Germantown, MD) and the PB1 segment was amplified by RT-PCR using Super Script III reverse transcriptase (Thermo-fisher Scientific, Waltham MA) using specific sets of primers (available upon request) and PCR conditions as follows: 55°C for 2 min, 94°C for 2 min, followed by 35 cycles at 94°C for 30 sec, 50°C for 30 sec, 68°C for 3 min, and final elongation of 4 min at 68°C. For NGS, virus RNA samples were extracted via the MagNA Pure LC RNA Isolation Kit (Roche, Indianapolis, IN) and multi-segment RT-PCR (MS-RT-PCR) was conducted as previously described [22] with Superscript III high-fidelity RT-PCR kit (Thermo-fisher Scientific, Waltham MA). Libraries were prepared using the Nextera XT DNA library preparation kit (Illumina, Inc., San Diego, CA) according to manufacturer’s instructions. Full genome assembly was performed using a pipeline from previously published methods [24].

Genetic stability of viruses.

B/Bris PB1 3M or B/Bris PB1att 4M viruses (MDCK P1) were serially passaged in MDCK cells and SPF eggs at 33°C. Serial passages were performed until stability of the B/Bris PB1att 4M was confirmed compared to the B/Bris PB1 3M virus, resulting in 4 serial passages in MDCK (sP1-sP4) or in eggs (sE1-sE4). During each passage, viruses were inoculated at three different dilutions (10⁻¹, 10⁻², 10⁻³). Cell culture supernatants and allantoic fluids were harvested at 72 h post-inoculation (hpi). The highest dilution used for inoculation that showed the highest hemagglutination (HA) titers after 72 hpi was used for the subsequent passages. During each passage, virus from cell culture supernatants and allantoic fluids were subjected to Sanger sequencing and/or NGS as described.

Viral Growth Kinetics in vitro.

MDCK cells at approximately 90% confluency in 24 well plates were infected at a multiplicity of infection (MOI) of 0.01 of each virus in Opti-MEM (Thermo-fisher Scientific, Waltham MA) containing 1 μg/ml TPCK-trypsin (Worthington Biochemicals, Lakewood, NJ) and 1% of antibiotic/antimycotic solution (Sigma-Aldrich St. Louis, MO). Infected cells were incubated at 33°C, 35°C, and 37°C under a 5% CO2 environment. Supernatants were collected at 6, 12, 24, 48, and 72 h post-inoculation (hpi). For titration of the samples, 50% tissue culture infectious dose (TCID₅₀) titers were determined using the Reed and Muench method [23]. Viral growth kinetics were performed using B/Bris wild type (wt) strain as a control. The growth kinetics experiments were performed in triplicates.

Structural Analysis of the PB1 segment.

3-dimensional (3D) analysis was conducted on the PB1 protein of the PB1 3M and PB1att 4M predicted sequences. DNASTar (Madison, WI) was used to generate fasta files
for the PB1 nucleotide sequences for B/Bris PB1 3M and B/Bris PB1att 4M and open reading frames (ORF) were identified with ExPASy (SIB bioinformatics resources) to produce the protein sequences. I-TASSER (University of Michigan, Ann Arbor, MI) [25] and CHIMERAs 1.141 (University of California, San Francisco, CA) [26] were used to recreate the PB1 protein 3D structure and for analysis of amino acid distances.

Animal use compliance

The animal study was performed under animal biosafety level 2 (ABSL-2) containment and according to protocols approved by the University of Georgia’s Institutional Animal Use and Care Committees (IACUC). Mice had access to water and feed ad libitum. Mice that experienced significant weight loss (≥25%) or scored 3 or higher on a 4-point scale of disease severity were humanely euthanized according to IACUC approved protocol and AVMA guidelines.

Vaccination and Vaccine Safety Study Design.

The safety of B/Bris PB1att 4M was evaluated in DBA/2J mice (Charles River, Norwich, CT) similarly to as previously described [22]. Vaccination was performed using a prime-boost vaccination strategy 3 weeks apart. Vaccinations were carried out via the intranasal (I.N.) route after mice were anesthetized with isoflurane. Mice (7 weeks-old, n=36) were randomly divided into 3 groups (n=12/group, 1/2 females). The B/Bris PB1att 4M group received 50 μl containing 10^6 EID₅₀/mouse of B/Bris PB1att 4M and was boosted by the same route with the same dose at 21 days post-vaccination (dpv). The two other groups were mock vaccinated with 50 μl 1X phosphate buffered saline (PBS, pH 7.4). After prime and boost, mice were monitored daily for clinical signs and survival up until the day prior to virus challenge. Body weight was monitored for up to 12 days post-vaccination and 12 days post-boost. On day 20 post-boost (41 dpv), 4 mice/group (n=2 male, n=2 female) were bled for antibody titer determination.

Virus Challenge Study Design.

At 21 days post-boost (dpb), the mice in the B/Bris PB1att 4M vaccine group and the mice in one of the mock-vaccinated groups (mock-challenged) were challenged I.N. with a 50 μl inoculum containing 10^7 EID₅₀/mouse of the lethal strain B/Bris PB2-F406Y [22]. The remaining mock-vaccinated group served as negative control and mice were mock challenged with 50 μl 1X PBS (mock-mock). After challenge, mice were monitored daily for clinical signs, body weight changes and survival until 14 days post-challenge (dpc) at which time all remaining surviving mice were terminally bled via the facial vein under isoflurane anesthesia and humanely euthanized according to approved IACUC protocols and following AVMA guidelines.

Hemagglutination Inhibition assay.

Sera were treated with receptor destroying enzyme (RDE) as follows: 15 μl of serum were mixed with 30 μl of 1X PBS and 15 μl of RDE and incubated at 37°C overnight. The next morning, RDE was inactivated at 56°C for 30 min. The treated sera were then brought to a final dilution of 1:10 with 1X PBS. The HI assay was performed as described previously [27] using B/Brisbane/60/2008 as antigen and 0.5% suspension of turkey red blood cells.

Results

Mutations E48K and K391E remain stable in combination after serial passages in MDCKs and SPF Eggs.
The B/Bris PB1 3M and B/Bris PB1att 4M strains were serially passaged at 33°C in MDCK cells (sP1-sP4) and SPF eggs (sE1-sE4; Table 1). Sanger sequencing analysis of the PB1 segment confirmed that the B/Bris PB1att 4M virus stably retained all modifications introduced (E48K, K391E, E580G, S660A) after 4 serial passages in both MDCK cells and in SPF eggs. In contrast, two new mutations were identified in the PB1 of the B/Bris PB1 3M at sE4 passage virus (E48K and G161C) but not at sP1 or sP4 (MDCK cell passages).

**Table 1. Evaluation of genetic stability in the PB1 gene of B/Bris PB1 3M and B/Bris PB1att 4M strains after serial passage in MDCK cells and SPF eggs.**

| Strain            | SPF egg passage | MDCK passage |
|-------------------|-----------------|--------------|
|                   | sE1<sup>a</sup> | sP1<sup>a</sup> | sE4<sup>a</sup> | sP4<sup>a</sup> |
| B/Bris PB1 3M     | K391E, E580G, S660A | K391E, E580G, S660A | E48K, K391E, E580G, S660A, G161C | E48K, K391E, E580G, S660A |
| B/Bris PB1att 4M  | E48K, K391E, E580G, S660A | E48K, K391E, E580G, S660A |

<sup>a</sup> Mutations confirmed by Sanger sequencing.

Stability of B/Bris PB1att 4M strain was confirmed after serial passages.

Deep sequencing of the viral genome of B/Bris PB1att 4M further confirmed that key mutations (E48K, K391E, E580G, S660A) were maintained after 4 passages in both MDCK cells and SPF eggs. As previously described for a similar IBV LAIV backbone [22], significant genomic stability was observed in the B/Bris PB1att 4M strain (Table 2). There was only one synonymous mutation in the PB2 (a1268g, nucleotide position).

**Table 2. Amino acid substitutions in all gene segments of B/Bris PB1att 4M after serial passage.**

| Segments | Predicted Mutations | Serially passaged in MDCKS (sP4) | Serially passaged in SPF eggs (sE4) |
|----------|---------------------|----------------------------------|-----------------------------------|
| PB2      | None                | 355G<sup>S</sup> (a1268g)       | 355G<sup>S</sup> (a1268g)         |
| PB1      | E48K (g163a)        | E48K (g163a)                     | E48K (g163a)                      |
|          | K391E (g1192a)      | K391E (a1192g)                   | K391E (a1192g)                    |
|          | E580G (a1760g)      | E580G (a1760g)                   | E580G (a1760g)                    |
|          | S660A (t1999g)      | S660A (t1999g)                   | S660A (t1999g)                    |
|          | Stop<sup>b,s</sup> (g2280a) | Stop<sup>b,s</sup> (g2280a)     | Stop<sup>b,s</sup> (g2280a)      |
| PA       | None                | None                             | None                              |
| HA       | None                | None                             | None                              |
| NP       | None                | None                             | None                              |
| NA       | None                | None                             | None                              |
| NB       | None                | None                             | None                              |
| BM1      | None                | None                             | None                              |
| BM2      | None                | None                             | None                              |
Mutations identified in the 3' untranslated region (UTR); Mutation identified at the stop codon of the open reading frame; Synonymous mutation; Higher case letters = amino acids; Lower case letters = nucleotides.

**B/Bris PB1att 4M virus shows a temperature-sensitive phenotype in vitro.**

To evaluate the temperature-sensitive (ts) phenotype of the B/Bris PB1att 4M strain, growth kinetics were evaluated in confluent MDCK cells at 33°C, 35°C, and 37°C and compared to the B/Bris wt virus. The B/Bris PB1att 4M and B/Bris wt viruses grew to similar titers at 33°C and 35°C (Fig. 1). The B/Bris wt strain displayed higher titers compared to the B/Bris PB1att 4M strain at 48 hpi at 35ºC and 37ºC and at 72 hpi in all temperatures. Overall, the B/Bris PB1att 4M strain showed a ts phenotype (Fig. 1).

**Figure 1.** B/Bris PB1att 4M displays a temperature-sensitive phenotype in growth kinetics in vitro. MDCKs cells were inoculated with an MOI of 0.01 with B/Bris PB1att 4M and B/Bris wildtype (wt) strains and incubated at 33°C (A), 35°C (B), and 37°C (C). Supernatants were collected at 6, 12, 24, 48, and 72 hours post infection and viral titers were quantified by TCID50 assays. The growth kinetics experiments were performed in triplicates and results represent an average of all experiments. Multiple t-test (Holm-Sidak method) was used to calculate P values. **p<0.01, ***p<0.001. Plotted data represent means ± standard errors.

**Structural analysis of PB1 from B/Bris PB1att 4M suggests molecular interactions between amino acid residues at positions 48 and 391.**

To visualize molecular interactions between the individual mutations/residues, the predicted 3D structure of the PB1 protein was recreated (Fig. 2). The amino acid in position 48 is located on one of the alpha helixes facing the amino acid in the 391 position. The wild type B/Bris PB1 protein structure showed potential interaction between the amino acids 48E and 391K via a single hydrogen bond or salt bridge with a predicted distance between the two residues of 1.856 Å. In contrast, in the PB1 3M predicted structure, no hydrogen bonds or any notable interactions between the side chain of the wild type amino acid 48E and the 391E mutation were noted. In the PB1 3M protein, the length between the most proximate hydrogen and oxygen groups capable of forming salt bridges was 5.714 Å. However, in the PB1att 4M, there is indication of potential interaction of 48K and 391E via a single hydrogen bond/salt bridge and the 1.809 Å distance between the two resembles the distance between the E48 and K391 in the wild type PB1, suggesting a stronger interaction and potential effect on protein stability.

**Figure 2.** Predicted 3D structure of PB1 showing mutations in residues 48 and 391 and respective molecular interactions between them. 3D structure predictions using iTASSER of the B/Bris PB1 protein sequence and potential interactions at the interface of amino acid 48 and 391. Blue lines represent presence of hydrogen bonds. Dotted yellow lines indicate predicted distances in
Amstrongs (Å). (A) Potential interaction via a single hydrogen bond and a predicted distance of 1.856 Å between amino acids 48E and 391K in the wt PB1. (B) Lack of predicted hydrogen bonds and a predicted distance of 5.714 Å between amino acids 48E and 391E in the PB1 3M protein. (C) Amino acids 48K and 391E restore the predicted single hydrogen bond interaction and a distance of 1.809 Å between these two amino acids in the PB1att 4M protein.

**B/Bris PB1att 4M is safe and immunogenic as a potential vaccine candidate**

To test the safety and immunogenicity of the B/Bris PB1att 4M vaccine strain in vivo, DBA/2J mice were vaccinated with either the B/Bris PB1att 4M virus (n=12) or mock-vaccinated (1X PBS; n=24) by intranasal inoculation in a prime-boost regimen, 20 days apart. Mice vaccinated with B/Bris PB1att 4M showed minor weight loss starting at 7 dpv, but quickly recovered (Fig. 3A). No body weight loss was observed after boost. No other clinical signs were observed, and all mice survived the vaccination regimen. HI titers were measured at 20 dpb and vaccinated mice showed titers ≥160, predictive of protection against clinical disease (Fig. 3B).
Figure 3. Safety and immunogenicity of the B/Bris PB1att 4M virus. DBA/2J 4-week-old, male and female mice were inoculated I.N. with either B/Bris PB1att 4M or PBS (Mock groups). (A) Percentage in body weight changes was evaluated daily for 12 days following vaccination. (B) Immunogenicity of the B/Bris PB1att 4M virus was measured by HI assay in mouse sera at 20 days post boost (dpb). Plotted data represent means ± standard errors.

B/Bris PB1att 4M vaccine candidate successfully protects mice against lethal challenge

In order to assess the efficacy of the B/Bris PB1att 4M vaccine strain, mice were challenged at 21 dpb with a lethal dose (10^7 EID50/mouse) of the B/Bris PB2-F406Y mutant strain [22]. A group of mock-vaccinated mice remained unchallenged throughout the study (mock-mock). Mice in the B/Bris PB1att 4M-vaccinated group showed no apparent signs of disease with minimum weight loss and all survived the virus challenge (Fig. 4A-B). Mice from the mock-challenge group showed rapid and severe weight loss along with severe clinical signs and approximately 87% of the mice in this group succumbed or had to be humanely euthanized by 8 dpc (Fig. 4A-B). There was one survivor out of eight, a female mouse, that also showed severe weight loss but started to recover at 7 dpc, showing a rapid recovery starting at 8 dpc (Fig. 4A-B). As expected, HI antibody titers against B/Bris PB1att 4M remained high in vaccinated mice at 14 dpc (Fig. 4C). The survivor mouse from the mock-challenge group seroconverted as well (Fig. 4C).

Figure 4. Protective efficacy of B/Bris PB1att 4M vaccine against homologous challenge. Vaccinated (B/Bris PB1att 4M) or unvaccinated (mock-challenged) DBA/2J mice were challenged with the homologous lethal strain B/Bris PB2-F406Y 21 days post boost. A non-vaccinated, non-challenged control group was included (mock-mock). (A) Percentage in body weight change and (B) survival rate were evaluated daily for 12 days after challenge. (E) The serum antibody response against the homologous (B/Bris wt) virus was measured by HI assay at 14 days post-challenge (dpc). Plotted data represent means ± standard errors.

Discussion

LAIVs are thought to generally elicit superior immunity and cross-protection against influenza viruses compared to inactivated products due to the perceived notion that they can better stimulate cell-mediated and mucosal immune responses in addition to humoral systemic immunity [28,29]. However, in recent years the overall low performance of
LAIVs in some age groups highlight the need for the continued development of more efficacious alternative LAIV technologies. We have developed a LAIV strategy by incorporating ts mutations on the PB1 that can be applied similarly for both IAV and IBV [19,21,22]. This strategy has been shown to be safe and effective in mice, pigs and chickens, stimulating cross-protective immunity against antigenically distinct viruses and can be used in a quadrivalent formulation [19,20,22,30-32]. In a previous study using an IBV LAIV candidate with the PB1 gene segment carrying three ts mutations K391E, E580G, S660A and a C-terminus HA tag, the K391E mutation reverted to its wild type form (E391K) during the initial passages in SPF eggs [22]. In contrast, passage in MDCK cells led to the quick emergence of the E48K mutation and retention of the K391E. Compensatory mutations are thought to emerge when the initial fitness of the mutant strain is reduced [33]. Our initial observations suggested that the E48K mutation could stabilize the ts K391E mutation in the vaccine candidate strain and therefore could contribute to a more stable and safe attenuated vaccine candidate. In this report, we found that indeed the K391E mutation was maintained in the presence of the E48K mutation, and the resulting B/Bris PB1att 4M strain was attenuated and safe, resulting in efficacious protection against homologous challenge.

Genomic analysis of the B/Bris PB1att 4M strain after 4 passages in eggs revealed the stability of the E48K, K391E, E580G, S660A mutations. In contrast, the B/Bris PB1 3M strain passaged 4 times in eggs showed the presence of not only the intended mutations (K391E, E580G, S660A) but also the acquisition of the potentially compensatory mutation E48K and the additional G161C mutation whose role remains unknown. Next generation sequencing (NGS) revealed all the target mutations were maintained in the PB1 of B/Bris PB1 4M strains, and there were no significant mutations in other segments. There was one synonymous mutation (355G) in the PB2 of both strains, in which the codon was changed from 1268GGA to 1268GGG. This shows the viral genome of the B/Bris PB1 4M strain is stable even after several passages in MDCKs and SPF eggs, indicating the introduction of the compensatory mutation E48K does not significantly alter the overall genome structure.

Compensatory mutations may contribute to improved virus fitness by either inducing conformational changes towards either a more energetically favorable form of the protein and/or by altering molecular bonds [34,35]. Forming and/or increasing hydrogen bonds among key amino acid residues is a known effector of protein stability [36,37]. In addition, E and K amino acid residues can form salt bridges if the distance between their side chains is <4 Å. 3D structure prediction analysis of the IBV PB1 protein suggests that the 48E-391K pair are part of opposite alpha-helices occupying a space that would favor formation of a single hydrogen bond and/or salt bridge (~1.8 Å). This is not the case for the 48E-391E pair in the PB1 3M protein (distance > 5 Å); however, the potential for a hydrogen bond/salt bridge is restored in the 48K-391E pair in the PB1att 4M protein (~1.8 Å). The stability of the B/Bris PB1att 4M strain in either eggs or MDCK cells and the acquisition of the E48K mutation in the egg-passaged B/Bris PB1 3M is consistent with the notion of a stabilizing role of the E48K that maintained the K391E mutation.

Interestingly, similar number of passages in MDCK cells did not affect the stability of the PB1 gene segment in the B/Bris PB1 3M strain, and the E48K mutation was not observed. This finding was unexpected considering prior observations with our original IBV LAIV candidate in which the E48K mutation was initially identified after the first serial passage in MDCK cells [22]. Another unexpected finding was the E48K mutation emerging and stabilizing the K391E mutation during serial passages in SPF eggs. A key difference between the previous results and those in this report is that the virus stocks used for serial passages here were amplified in MDCK cells, while in the previous study they were initially amplified in MDCK cells followed by amplification in eggs. Another difference is the absence of the HA tag at the C-terminus of the PB1 protein gene segment in the B/Bris PB1att 4M strains. It is important to note that in our previous work, a virus rescued without the HA tag also showed the PB1 E48K mutation that seemed to stabilize the K391E mutation after 9 serial passages in eggs [22]. It is tempting to speculate that the C-terminal
HA tag further impairs the polymerase function of the PB1 subunit protein forcing the K391E mutation to quickly revert to wild type when passaged in eggs. Regardless of the potential bottleneck effect of the HA tag for PB1 activity and the effect of initial virus stock amplification in eggs, our results strongly suggest a functional link between amino acids 48 and 391 protein and significant stability of the other two ts mutations (E580G, S660A) in the PB1 protein of IBV [22]. The E48K mutation is unique to the B/Bris PB1att 4M strain as revealed by the analysis of 662 unique IBV PB1 protein sequences deposited in the Influenza Research Database (IRD; www.fludb.org). Furthermore, the B/Bris PB1att 4M strain showed remarkable genomic stability in the rest of the genome (Table 2), a finding that is consistent with our previous study [22].

In vitro kinetics studies confirmed that the B/Bris PB1att 4M strain retained its ts phenotype despite the E48K mutation. In vivo in DBA/2J mice, the B/Bris PB1att 4M was attenuated and provided protection against homologous challenge. DBA/2J mice primed with a high dose of the B/Bris PB1att 4M strain showed minor weight loss between 3-8 days post-vaccination and quickly recovered, confirming its attenuation and safety in vivo. Female and male mice were equally protected against a lethal dose of the antigenically homologous B/Bris/PB2 F406Y strain, exhibiting no weight loss and no mortality. In contrast, the non-vaccinated B/Bris/PB2 F406Y-challenged mice showed significant signs of disease, weight loss, and mortality. While none of the non-vaccinated male mice survived the virus challenge, only one female survived. In conclusion, we provide evidence of the molecular basis for a compensatory mutation in the PB1 protein of an IBV LAIV candidate that improves its stability while retaining the safety and efficacy profiles expected in an attenuated influenza vaccine candidate.

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Author Contributions
Conceptualization, Jongsuk Mo, Stivalis Cardenas-Garcia, C. Joaquín Cáceres, Daniel Perez and Daniela Rajao; Formal analysis, Jongsuk Mo; Funding acquisition, Daniel Perez; Investigation, Jongsuk Mo, Stivalis Cardenas-Garcia, Lucas Ferreri, C. Joaquín Cáceres and Ginger Geiger; Methodology, Jongsuk Mo, Stivalis Cardenas-Garcia, Jefferson Santos, Lucas Ferreri, C. Joaquín Cáceres, Ginger Geiger, Daniel Perez and Daniela Rajao; Project administration, Jongsuk Mo, Stivalis Cardenas-Garcia and C. Joaquín Cáceres; Supervision, Daniel Perez and Daniela Rajao; Writing – original draft, Jongsuk Mo and Daniela Rajao; Writing – review & editing, Jongsuk Mo, Stivalis Cardenas-Garcia, Jefferson Santos, Lucas Ferreri, C. Joaquín Cáceres, Ginger Geiger, Daniel Perez and Daniela Rajao.

Institutional Review Board Statement
The study was conducted according to the guidelines of the Guide for the Care and Use of Laboratory Animals and approved by the University of Georgia’s Institutional Animal Use and Care Committee (IACUC; protocol number A2019 03-032-Y2-A10) on December 13, 2019.

Conflicts of Interest
The authors declare no conflicts of interest.

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