Contrasting selective patterns across the segmented genome of bluetongue virus in a global reassortment hotspot

Maude Jacquot,1,*,†,††,‡‡ Pavuluri P. Rao,2,†† Sarita Yadav,3,§
Kyriaki Nomikou,4,§§ Sushila Maan,5 Y. Krishna Jyothi,6 Narasimha Reddy,7
Kalyani Putty,7 Divakar Hemadri,8 Karam P. Singh,9 Narender Singh Maan,5
Nagendra R. Hegde,2,** Peter Mertens,3,10 and Roman Biek1,*,***

1Institute of Biodiversity, Animal Health and Comparative Medicine, Boyd Orr Centre for Population and Ecosystem Health, College of Medical, Veterinary and Life Sciences, University of Glasgow, Glasgow, UK, 2Ella Foundation, Genome Valley Hyderabad, Hyderabad, Telangana, India, 3The Pirbright Institute, Pirbright, Woking, Surrey GU24 0NF, UK, 4MRC—University of Glasgow Centre for Virus Research, Glasgow, UK, 5College of Veterinary Sciences, LLS University of Veterinary and Animal Sciences, Hisar, Haryana, India, 6Veterinary Biological and Research Institute, Vijayawada, Andhra Pradesh, India, 7PVNR Telangana Veterinary University, Hyderabad, Telangana, India, 8ICAR—National Institute of Veterinary Epidemiology and Disease Informatics, Bengaluru, Karnataka, India, 9Centre for Animal Disease Research and Diagnosis, Division of Pathology, ICAR—Indian Veterinary Research Institute, Izatnagar, 243 122 Bareilly, Uttar Pradesh, India and 10The School of Veterinary Medicine and Science, University of Nottingham, Sutton Bonington, Leicestershire, UK

*Corresponding author: E-mails: maude.jacquot@gmail.com; roman.biek@glasgow.ac.uk
†Present address: Spatial Epidemiology Lab (SpELL), University of Brussels, Brussels, Belgium and UMR0346 Epidemiology of Animal and Zoonotic Diseases, INRA-VetAgro Sup, Saint-Genés-Champanelle, France.
‡Present address: Biovet Private Limited, Malur Taluk, Kolar District, Karnataka, India.
§Present address: ICAR—Central Institute for Research on Buffaloes, Hisar, Haryana, India.
**Present address: National Institute of Animal Biotechnology (NIAB), Gachibowli, Hyderabad, India.
***http://orcid.org/0000-0001-5945-4689
§§http://orcid.org/0000-0002-7013-1853
****http://orcid.org/0000-0003-3471-5357
††These authors contributed equally to this work.

Abstract

For segmented viruses, rapid genomic and phenotypic changes can occur through the process of reassortment, whereby co-infecting strains exchange entire segments creating novel progeny virus genotypes. However, for many viruses with segmented genomes, this process and its effect on transmission dynamics remain poorly understood. Here, we assessed the consequences of reassortment for selection on viral diversity through time using bluetongue virus (BTV), a segmented...
1. Introduction

RNA viruses have an extraordinary ability to diversify and evolve. The observed high evolution rate of RNA viruses is partly attributed to limited or no proof reading capacity of RNA-dependent RNA polymerases. In addition, for viruses with segmented genomes, rapid genomic and phenotypic changes can occur through the process of reassortment, whereby co-infecting different viral strains exchange entire segments (see McDonald et al. 2016 for a review). This has received considerable research attention in certain groups, such as influenza viruses, where reassortment has been linked to changes in host range and the emergence of novel strains (e.g. Li et al. 2004; Schweiger, Bruns, and Meixenberger 2006; Simonsen et al. 2007; Nelson et al. 2008; Rambaut et al. 2008; Smith et al. 2009; Trifonov et al. 2009; Wu et al. 2015; Pohlmann et al. 2017). However, for many segmented-RNA viruses, in particular those affecting non-human hosts, the consequences of reassortment in relation to selection and the maintenance of viral diversity at the genome scale remain poorly understood.

Bluetongue virus (BTV) belongs to the genus Orbivirus of the family Reoviridae. It is responsible for an economically important disease of ruminants, bluetongue disease (BT). The distribution of BTV overlaps with the distribution of BTV transmission-competent midges belonging to the genus Culicoides (Tabachnick 2004). BT is endemic in most parts of tropical and subtropical regions, and seasonal incursions of the disease are observed in parts of temperate regions during the summer (Mellor and Wittmann 2002; Purse et al. 2005; Mellor et al. 2008). Bluetongue outbreaks cause severe economic damage due to the direct effects on livestock, trade restrictions, and the cost of surveillance and control (e.g. Osburn 1994; Maclachlan et al. 2009; Velthuis et al. 2010; Piñior et al. 2015). In the USA for example, BTV is thought to cause losses in the order of US$125 million annually (Tabachnick 1996).

The genome of BTV consists of ten double stranded RNA (dsRNA) segments with each segment coding for at least one major protein. Some of these proteins form part of the three-layered structure of the virus whereas others are non-structural. The inner capsid consists of the three minor viral proteins (VP) required for genome transcription and replication, i.e. VP1 (RNA-dependent RNA polymerase), VP4 (RNA capping, guanylyltransferase, transmethylase 1 and 2), and VP6 (RNA helicase), coded by genome Segments 1, 4, and 9, respectively. These proteins, together with the ten genome segments, are encased by a shell composed of VP3 encoded by Segment 3, which is surrounded by the intermediate capsid protein layer composed of VP7 coded by Segment 7. The outer capsid is made up of VP2 and VP5, coded by Segments 2 and 6, respectively. VP2 and VP5 are involved in the virus binding to and entering into the mammalian host (and likely also insect vector) cells (Mertens and Diprose 2004; Loureno and Roy 2011). Variability in the sequence of VP2, and to some extent VP5, contributes to the specificity of differential neutralising antibody responses in infected animals, and forms the basis for classifying BTV into at least twenty-seven different serotypes. Five non-structural proteins (NS) are produced by BTV in infected cells. NS1 enhances VP synthesis and is involved in formation of tubules in the cytoplasm of infected cells (Boyce, Celma, and Roy 2012), while NS2 is a major component of viral inclusion bodies present in infected cells (Butan and Tucker 2010). These two proteins are coded by genome Segments 5 and 8, respectively. Segment 10 codes for NS3, a glycoprotein involved in egress of the virus (particularly from insect cells), and NS3a, a truncated version of NS3 produced by an alternate in-frame start codon (Wu et al. 1992; Han and Harty 2004; Wirblich, Bhattacharya, and Roy 2006; Celma and Roy 2009; Bhattacharya and Roy 2013). NS4 is a small peptide of 77–79 residues coded by Segment 9 from an alternate open reading frame (ORF) from that of VP6 (Belhouchet et al. 2011; Ratinier et al. 2011). A second alternate ORF was also detected in Segment 10 of different orbiviruses (including BTV) coding for a small peptide (50–59 aa), a putative NS5 protein (Stewart et al. 2015).

The segmented nature of the BTV genome can lead to the generation of a mosaic of viruses through independent reassortment of different genome segments (Oberst et al. 1987; Samal et al. 1987; Batten et al. 2008; Rambaut et al. 2008; Shaw et al. 2013; Nomikou et al. 2015; Schirzinger et al. 2018). While this has the potential to change the phenotypic characteristics of the virus, as seen in other segmented viruses (e.g. Li et al. 2004; Schweiger, Bruns, and Meixenberger 2006; Simonsen et al. 2007; Nelson et al. 2008; Rambaut et al. 2008; Smith et al. 2009; Trifonov et al. 2009; Wu et al. 2015; Pohlmann et al. 2017), conclusive examples of this have been lacking for BTV. In vitro, BTV reassortment appears to be a flexible process that is largely unaffected by selective constraints (Shaw et al. 2013). In contrast, a recent analysis of BTV field strains from Europe revealed non-random associations between some segments and found a higher probability of amino acid substitutions on tree branches associated with reassortment events (Nomikou et al. 2015). While these latter results support a more important role for reassortment affecting BTV genome evolution under field conditions, there is a need for further studies aimed at assessing its significance for driving adaptive and epidemiological dynamics in vivo.

Reassortment is expected to be most common and biologically relevant in areas supporting a high diversity of BTV serotypes and strains. Serological data show that twenty-two BTV serotypes exist in India, making it a region with one of the
highest levels of BTV diversity in the world (Rao et al. 2016; Maan et al. 2017). Although most of these serotypes appear to be ‘native’ to the Indian sub-continent, there is evidence for the introduction of exotic BTV serotypes and strains belonging to the major western topotype, from Africa and the USA, linked to the use of live vaccines or the livestock trade (Maan et al. 2015; Krishnajyothi et al. 2016), and contributing to further diversity. In addition, the severity of BT outbreaks in Indian sheep has increased significantly in recent decades (Prasad et al. 2009; Chand et al. 2015; Rao et al. 2016), raising the possibility of selection of more virulent (possibly reassortant) strains and recent evolutionary changes in the virus driving rapid phenotypic changes. Together, this makes India an important area for studying BT evolution and its epidemiological implications. However, studies examining the evolutionary history of BTV in India based on whole genome data have begun only recently (Maan et al. 2015) and have so far been limited in scope.

Here, we triple the number of BTV genome sequences available from India (from 30 to >90), based on samples collected over the last four decades. Using these data, our aim was (1) to quantify the processes of reassortment and selection, and (2) to determine how these processes are interacting to shape the evolution of the BTV genome in a setting characterised by a high diversity of circulating strains.

2. Material and methods

2.1 Cells and virus isolates

Fifty-seven BTV isolates from India were newly sequenced for this study (Supplementary Table S1). These isolates were obtained from ethylenediamine tetraacetic acid blood or tissue samples (spleen) collected from outbreaks from animals (sheep, goat) showing clinical signs of BT between 1993 and 2015.

For forty-six isolates, Culicoides sonorensis cells (KC cells), an insect cell line derived from the embryo of C. sonorensis, BHK-21 (clone 13), and BSR cells (a sub-clone of BHK-21 cells) were used for isolation of BTV. KC cells were infected with washed and lysed red blood cells or homogenised tissues. KC viral isolates were propagated in BHK or BSR cell monolayers (175 cm² flasks) at 37°C in Dulbecco’s modified Eagle’s medium supplemented with 100 units/ml of penicillin and 100 μg/ml streptomycin until completion of cytopathic effects, generally between 48 and 72 h post-infection and passed 3–6 times in BHK or BSR cells before RNA isolation.

For eleven isolates, BHK-21 cells were infected with different isolates of BTV and the lysed cells were pelleted after completion of cytopathic effects.

2.2 RNA isolation

BTV dsRNA was extracted from pellets of virus-infected cell cultures using a guanidinium isothiocyanate procedure as previously described (Attoui, Billoir, and Cantaloube 2000). Total RNA was extracted from the infected cell pellet using TRIzol (Invitrogen, USA) as per manufacturer’s instructions. Cellular single stranded RNA was separated from the viral dsRNA by precipitating in 2 M lithium chloride (LiCl) at 4°C overnight. An equal volume of isopropanol and 0.25 volumes of 7.5 M ammonium acetate were added to precipitate dsRNA at −20°C for 2 h, pellets were washed with 70 per cent ethanol and finally suspended in 50 μl of nuclease free water. The isolated dsRNA was shipped to MRC-CVR, UK for deep sequencing. For the eleven BHK isolates, total RNA was isolated from the pellet by acid phenol method (Gauthami et al. 2015).

2.3 Sequencing

Full-length cDNA copies of BTV genome segments were synthesised and amplified in a sequence-independent manner using the ‘anchor spacer-ligation’ method (Maan et al. 2007). BTV dsRNA was DNase treated using Turbo DNase (Ambion Life Technologies), according to manufacturer’s instructions and ligated to an anchor primer (Maan et al. 2007). After purification of ligated RNA using the RNA Clean & Concentrator kit (Zymo research, USA), purified RNA was reverse transcribed to generate full length cDNA using SuperScript III first-strand synthesis system (Thermo Fisher Scientific). Amplification of the resulting cDNA was performed using KAPA HiFi Hot Start Ready Mix (Kapa Biosystems, UK) using the ‘5-15-1’ primer, which is complementary to the anchor primer followed by purification of dsDNA PCR products using the DNA Clean & Concentrator kit (Zymo research). Quantity and quality of PCR products were tested using Qubit Fluorometer 3 (Life Technologies Q32854) and Agilent 4200 TapeStation (Agilent Technologies, 5067-5584) and the concentration of each sample was adjusted to 0.2 ng/μl for library preparation using the Illumina Nextera XT DNA library preparation kit. The amplified PCR reaction products were purified using Ampure XP beads (Agencourt, Beckman Coulter). The concentration of library DNA was measured using Qubit Fluorometer 3 and Qubit dsDNA high sensitivity (HS) Assay Kit (Life Technologies, Q32854). The library sizing (quality and distribution) and quantification were determined using the High Sensitivity D5000 Screen Tape assay (Agilent, 5067-5584) in a 4200 TapeStation System (Agilent). The library pool (at 4 n mole) was loaded in the MiSeq reagent kit Version 2 cartridge (Illumina) and paired-end reads of 150 bp were produced on a MiSeq platform. The quality of the resultant fastq files were assessed using FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc/, accessed 1 July 2019). Primers, Illumina adapter sequences, low quality-bases on the 3’ end of reads with Phred-scaled quality less than 30 and reads with a length <50 bp were removed using Trim Galore (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/, accessed 1 July 2019).

For the BHK-21 isolates, genomic libraries were prepared, sequenced, and processed as described in Rao et al. (2013).

2.4 Genome sequence reconstruction

Whole genomes were generated using de novo assembly and reference-based mapping methods.

Briefly, quality-controlled reads were de novo assembled using the SPAdes genome assembler version: 3.10.1 (Bankevich et al. 2012). Resulting contigs were grouped by segment and in each case, the one covering the largest portion of the segment was retained. For reference-based mapping, a unique reference was selected for each sample based on the best match from a BLAST search for each segment. Quality-controlled reads were mapped against the reference sequences using Tanoti (http://bioinformatics.cvr.ac.uk/tanoti.php, accessed 1 July 2019) and Bowtie2 (Langmead and Salzberg 2012). Assemblies were viewed using UGENE (Okonechnikov, Golosova, and Fursov 2012).

Summary statistics of mapped and unmapped reads, depth and breadth of coverage across all genome segments for all samples are shown in Supplementary Table S2. Consensus sequences were generated using SAMtools (Li et al. 2009). Sequences
generated in this study have been submitted in GenBank under accession numbers listed in Supplementary Table S1.

2.5 Sequence alignments

The consensus sequences generated for individual genome segments were combined with sequences from one Pakistani and other Indian BTV genomes available from GenBank in November 2017 (and for which metadata was sufficient), bringing the total number of genomes used in this study to ninety-two. Spatial distribution of samples and other metadata are summarised in Supplementary Fig. S1 and Table S1.

For each segment, the main coding region—VP1, VP2, VP3, VP4, NS1, VP5, VP7, NS2, VP6, and NS3 for the segments 1–10, respectively—was aligned using PAL2NAL (Suyama, Torrents, and Bork 2006) according to protein sequences aligned with Clustal Omega (Sievers et al. 2011). For the remainder of the manuscript, these coding regions are referred as Seg-1 to Seg-10. Alignments were also obtained for NS4 and NS5 (alternate ORFs on Seg-9 and Seg-10) but were only used for selection pattern analyses, as described below.

2.6 Phylogenies and evolutionary rate estimates

For each BTV segment, we evaluated the temporal signal of the heterochronous sequences with TempEst v1.5.1 (Rambaut et al. 2016) and time-scaled Bayesian phylogenetic trees were estimated in BEAST 1.8.4 (Drummond et al. 2012) using the SDR06 codon substitution model for the coding sequence (Shapiro, Rambaut, and Drummond 2006) under a relaxed molecular clock with a lognormal distribution of rates. A Gaussian Markov Random Field Bayesian skyline coalescent model was used as the tree prior (Minin, Bloomquist, and Suchard 2008). For samples for which exact dates of collection were unavailable, approximate dates were used and uncertainty added to the tips (Precision column in Supplementary Table S1). For each segment, a Monte Carlo Markov chain (MCMC) was run for the number of generations needed to achieve stationary distributions and convergence (5 × 105–5 × 106 generations) and sampling frequency adjusted accordingly to yield 10,000 samples from the posterior. We used Tracer v1.6 (Rambaut et al. 2013) to visualise the posterior distribution of each parameter and to obtain an estimate of the effective sample size (ESS) after removal of the initial 10 per cent burn-in. We assumed a run had reached sufficient mixing if the ESS of all the parameters was above 200. For each BTV segment, a maximum clade credibility (MCC) tree was produced after removal of the initial 10 per cent burn-in based on common ancestor heights method (Heled and Bouckaert 2013) using the auxiliary program TreeAnnotator included in the BEAST package.

2.7 Sequence clustering and reassortment

In order to quantify functional genetic diversity present in Indian BTV genomes and its changes through time, we used the Bayesian clustering algorithm provided by the program K-Pax2 (Pessia et al. 2015). This approach overcomes problems encountered with standard clustering tools relying on arbitrary criteria that are challenging to calibrate in cases where evolutionary histories and levels of divergence may vary considerably across the genome. For each segment, clusters of sequences based on shared amino acid changes were defined via K-Pax2. Ten independent runs were carried out from different starting configurations under the default prior settings and upper bound values for the number of clusters corresponding to the number of samples (n = 92). The optimal clustering of each segment was identified using the log posterior scoring function of the method. Delineated clusters were highlighted on the plotted phylogenies using the R package ggtree (Yu et al. 2017).

We used multi-dimensional scaling (MDS) plots summarising tree-to-tree variation in branch lengths to assess how reassortment might have broken up associations among segments (Rambaut et al. 2008; Bahl et al. 2013). For each BTV segment, 500 trees sampled from the MCMC chain were used to determine the time to most recent common ancestor (TMRCA) for any pair of Indian isolates sampled within the same year (from 2003 onwards). The correlation coefficient of TMRCA estimates across all pairwise comparisons of trees was calculated and used to estimate the tree-to-tree distance (inverse). The matrix of tree-to-tree distances was then plotted in two dimensions using MDS. In this analysis, the spread of each point cloud represents the statistical uncertainty in the phylogenetic history of each BTV segment, where overlapping point clouds represent cases for which the null hypothesis of no reassortment cannot be rejected.

Reassortment events were further identified by the graph incompatibility-based reassortment finder (GiRaF) program originally developed for influenza viruses (Nagarajan and Kingsford 2011). Fifty repeats of the following two-steps procedure were performed. First, 1,000 unrooted candidate trees were inferred for each segment with MrBayes v3.2.7 (Ronquist and Huelsenbeck 2003; Ronquist et al. 2012) using a GTR+I+G substitution model, a burn-in of 50 per cent (100,000 iterations), and sampling every 200 iterations. This tree sample was used to account for uncertainty in the phylogenetic inference for each segment. GiRaF, which identifies sets of taxa with different phylogenetic placement across the trees, was applied to each tree set using default settings across all forty-five possible combinations of segments to comprehensively catalogue reassortment events.

We also used Recombination Detection Program (RDP) version 4.97 (Martin et al. 2015) with default settings, except that we invoked the `scan for reassembly and recombination' setting to identify any reassortment between the gene segments of Indian BTV strains. As evidence for reassortment, we took events detected by at least one of seven different recombination detection methods implemented in RDP: RDP, MAXCHI, and GENECONV methods in primary scanning mode and the Bootscan, CHIMAERA, SisScan, and 3SEQ methods in secondary scanning mode, each with a Bonferroni-corrected P-value cutoff of 0.05. Distribution of inferred breakpoints of events detected by at least five methods was also analysed.

Finally, linkage disequilibrium measures between pairs of single-nucleotide polymorphisms (SNPs) (within and among segments) were obtained using the D’ statistic implemented in DnaSP v6 (Librado and Rozas 2009).

2.8 Selection patterns

We used three complementary approaches to examine the BTV genome data for evidence of selection.

First, data were subjected to a neutrality test using Tajima’s D (Tajima 1989) with MEGA v7.0 (Kumar, Stecher, and Tamura 2016). Tajima’s D statistic measures the difference between two statistics describing the diversity within the alignment (θI, based on the average number of pairwise differences among sequences and θH based on the number of segregating sites), standardised by the variance of this difference. Under the neutral evolutionary model, which assumes that polymorphisms segregate at mutation–drift equilibrium, Tajima’s D is expected to be null. Positive Tajima’s D values indicate that there is an excess
of polymorphisms at intermediate frequency and is typically associated with diversifying selection. Conversely, negative Tajima’s D values can be the result of purifying selection or a selective sweep (a beneficial allele has recently reached fixation due to strong positive selection), which both result in an excess of weakly divergent alleles. However, both positive and negative Tajima’s D values can also result from demographic and epidemiological processes.

Second, we tested for selection acting on particular amino acid sites by comparing the rates of non-synonymous (dN) and synonymous substitutions (dS), where dN/dS ratios are described by the parameter ω. Values of ω > 1 indicate positive selection, whereas a ω < 1 indicates the operation of purifying selection (Yang and Bielawski 2000). Analysis was performed with the program CODEML, which is part of the PAML package (Yang 1997). Likelihood ratio tests were computed to compare pairs of models: M0 (single ratio) with M3 (discrete), M1a (nearly neutral) with M2a (positive selection), and M7 (beta distribution) with M8 (beta distribution + positive selection) (Wong et al. 2004; Yang, Wong, and Nielsen 2005). The M0–M3 comparison is used as a test of variable ω among sites and M1a–M2a and M7–M8 comparisons are used as tests of positive selection (Yang 1997).

Finally, selection pressures were quantified for all codon alignments for each segment in a maximum likelihood framework. To determine the degree of natural selection acting on all protein coding regions, the average dN/dS ratio was estimated for the entire tree by using single likelihood ancestor counting (SLAC) (Kosakovsky Pond and Frost 2005). Positively selected codons along internal branches were detected with the fixed-effects likelihood (FEL) method (Kosakovsky Pond et al. 2006). Codons subject to episodic diversifying selection were identified with the mixed-effects model of evolution (MEME) method (Kosakovsky Pond et al. 2013), using a significance threshold of P < 0.05. Fast unconstrained Bayesian approximation (FUBAR) was used to rapidly detect negative and positive selection by Bayesian MCMC analyses to robustly account for parameter estimation errors (Murrell et al. 2013). Codons with posterior probabilities ≥0.9 were reported as being either negatively or positively selected. All methods were implemented in the HyPhy package (Kosakovsky Pond, Frost, and Muse 2005), combined with the best-fit model of nucleotide substitution based on Akaike information criterion. All these analyses were accessed through the Datamonkey webserver (Delport et al. 2010).

3. Results

3.1 TMRCAs and evolutionary rates

Regression of root-to-tip genetic distance against sampling time (Supplementary Fig. S2) exposed sufficient temporal signal in each dataset (segment) to proceed with phylogenetic molecular clock analyses. BEAST (Drummond et al. 2012) estimates of the coefficients of variation (CoV) were consistently >2 for each BTV segment (Table 1), indicating that strict clock-like evolution could be rejected and validating our choice to use a relaxed molecular clock. However, due to this high degree of clock rate variability seen, we examined whether TM RCA estimates were affected by this. Exploratory analyses revealed that three isolates of BTV-10 (IND2014/BTV1_2014-12-06, IND2014/ BTV2_2014-12-06, and BTV10IND2003K3_2003-08-01) in particular contributed to the high CoVs observed (MCC trees available at doi: 10.6084/m9.figshare.8275424). These three genomes exhibited >99 per cent similarity to an attenuated BTV strain, which is part of a live vaccine and thus might have been introduced into India through vaccinated animals or unauthorised use of American BTV-vaccines in the country (Maan et al. 2015). Repeating the BEAST analysis after excluding the BTV-10 sequences brought the CoV for the clock rate down for five of the ten segments but this had a negligible effect on their TRMCA estimates (Supplementary Table S3). For these segments, temporal calibration of BTV phylogenies was therefore not sensitive to clock rate variation. For the remaining segments, TMRCAs should be treated with more caution. However, given that TRMCA estimates showed limited variation across all ten segments, we conclude that clock rate variation was unlikely to have caused biases in our temporal estimation.

For Seg-2 and Seg-6, the mean TMRCAs of the sampled Indian BTV isolates were estimated to have existed in the late 1930s or early 1940s. The mean TMRCAs for other segments placed their ancestry in the 1950s (Table 1). Also taking the uncertainty of these estimates into account, these results indicate that the present genetic diversity seen in Indian BTV isolates has arisen within the past century.

| Table 1. Rates of nucleotide substitution and root height of each segment of BTV. |
|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
|                  |                  |                  |                  |                  |                  |                  |                  |                  |                  |
|                  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  |
|                  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  | Mean  | Median | 95% HPDc lower  | 95% HPDc upper  |
| Seg-1            | 3.12  | 3.13   | 2.37             | 4.12             | 1.14E-03 | 1.13E-03 | 7.99E-04          | 1.47E-03          | 1953  | 1954   | 1941               | 1961               |
| Seg-2            | 4.60  | 4.59   | 3.98             | 5.26             | 1.46E-02 | 1.45E-02 | 1.06E-02          | 1.86E-02          | 1937  | 1938   | 1918               | 1954               |
| Seg-3            | 4.43  | 4.31   | 2.64             | 6.26             | 1.34E-03 | 1.33E-03 | 9.55E-04          | 1.76E-03          | 1956  | 1957   | 1949               | 1961               |
| Seg-4            | 3.04  | 2.96   | 2.04             | 4.21             | 1.28E-03 | 1.27E-03 | 9.09E-04          | 1.63E-03          | 1956  | 1957   | 1947               | 1961               |
| Seg-5            | 3.88  | 3.82   | 2.64             | 5.05             | 1.13E-03 | 1.12E-03 | 8.50E-04          | 1.41E-03          | 1955  | 1955   | 1947               | 1961               |
| Seg-6            | 4.95  | 5.03   | 3.57             | 5.79             | 9.40E-03 | 9.27E-03 | 5.96E-03          | 1.31E-02          | 1946  | 1946   | 1927               | 1961               |
| Seg-7            | 5.36  | 5.30   | 3.97             | 6.79             | 2.28E-03 | 2.25E-03 | 1.54E-03          | 3.11E-03          | 1954  | 1956   | 1944               | 1961               |
| Seg-8            | 3.05  | 2.97   | 1.98             | 4.21             | 1.22E-03 | 1.21E-03 | 8.54E-04          | 1.63E-03          | 1956  | 1957   | 1949               | 1961               |
| Seg-9            | 3.45  | 3.39   | 2.44             | 4.68             | 7.81E-04 | 7.77E-04 | 5.29E-04          | 1.05E-03          | 1950  | 1951   | 1934               | 1961               |
| Seg-10           | 2.28  | 2.21   | 1.45             | 3.17             | 1.06E-03 | 1.04E-03 | 7.31E-04          | 1.41E-03          | 1957  | 1957   | 1950               | 1961               |

Parameters were estimated based on ninety-two time-stamped Indian isolates in program BEAST v1.8.4 (Drummond et al. 2012) under a relaxed molecular clock and a flexible skygrid tree prior. Date of MRCA was deduced from the estimated mean root height.

aCoefficient of variation.

bMost recent common ancestor.

cHighest posterior density.
3.2 Sequence clustering and reassortment

Time-scaled phylogenies and K-Pax2 clustering (Pessia et al. 2015) distinguished ten phylogenetic groups at the protein level for Seg-2 and nine groups for Seg-6 (Fig. 1, Supplementary Fig. S3). On Seg-2, K-Pax2 clustering was perfectly consistent with the serotype delineation (neighbour-joining), with the exception of one published BTV-3 sequence (BTV03/08) which grouped with BTV-16 sequences (Fig. 1). The serotypes dominating during each decade of sampling changed through time with Serotypes 1 and 9 being common early on whereas the most recent samples were almost exclusively comprised of Serotypes 5 and 12. During each decadal period, between three and eight clusters (i.e. serotypes) were co-circulating (Fig. 2).

For other segments, time-scaled phylogenies along with the K-Pax2 clustering revealed a high turnover rate involving on average the simultaneous circulation of 3.8 clusters per decade (Fig. 3, Supplementary Figs S4–S10). Notably, results for Seg-5 indicated a strong gradual reduction in the number of circulating variants through time (Fig. 3). The observed patterns suggest that a single variant has become completely dominant by 2015 and is associated with the full spectrum of genomic backgrounds found in Indian BTV strains currently circulating.

We used MDS of TMRCAs (Bahl et al. 2013) to quantify the degree to which BTV segments had remained associated with each other through time and in the face of common genome reassortment. The analysis clearly indicated variable levels of reassortment for different segments (Fig. 4). Overlapping point clouds were seen for Seg-2, Seg-6, Seg-1, and Seg-7 as well as for Seg-3, Seg-8, Seg-9, and Seg-10, suggesting high level of linkage among segments within each of these groups. In contrast, Seg-4 and Seg-5 failed to show any clear association with other segments indicating that their evolutionary histories were more independent from those of other segments due to reassortment having disrupted these linkages.

We also found a high number of well-supported incompatibilities between trees, indicating reassortment, using GiRaF (Fig. 5). This was seen in all ten segments, although the frequency of reassortment and the number of sequences impacted varied among them (Fig. 5, Supplementary Fig. S11). Tree topology inconsistencies were found for strains of all serotypes except BTV-12 which did not exhibit admixture at the amino-acid level either (Fig. 1).

Explicitly testing for intra-segment recombination and reassortment using RDP4, yielded no evidence of intracomponent recombination in any virus but strong evidence of reassortment. Indeed, we detected fifty-eight reassortment/reassortment events by at least one of the seven methods used, forty-six being supported by five or more methods and inferred breakpoints.

Indian BTV viruses exhibited variable rates of evolutionary change across segments, ranging from 7.81E-04 (Seg-9) to 1.46E-02 (Seg-2) nucleotide substitutions per site per year (subs/site/year) with Seg-2 and Seg-6 showing higher rates than other segments (Table 1).

Figure 1. Phylogeny of Indian BTV Seg-2 and amino-acid based clusters of all genomic segments. Shown is a time-scaled maximum clade credibility (MCC) tree based on Seg-2 for ninety-two Indian BTV isolates, obtained using Bayesian phylogenetic inference in program BEAST (Drummond et al. 2012). Seg-2 sequences were assigned to amino-acid based clusters (Pessia et al. 2015), as highlighted on the tree, which corresponded almost perfectly to serotype (shown on the very right). The same clustering approach was used for all other segments in the genome, for which cluster memberships is shown as vignettes on the right side of the tree. Colours chosen to represent clusters are used consistently across all figures of the manuscript.
always fell on or very close to concatenation sites between segments (Supplementary Fig. S12A). These results confirm that reassortment has been frequent in our data. Consistent with this conclusion, we found mean linkage disequilibrium between SNPs of the different segments to be low (Supplementary Fig. S12B).

3.3 Selection patterns

The population genetics analysis showed that, at both the nucleotide and the amino acid level, Seg-2, Seg-6, and Seg-7 presented positive Tajima’s D values which indicates an excess of polymorphisms at intermediate frequency and suggests positive diversifying selection is acting on these segments (Tajima 1989). In contrast, all other segments showed negative Tajima’s D values (Table 2) suggesting purifying selection and/or a recent population expansion. The single exception to this was Seg-9 which showed a Tajima’s D close to zero.

No evidence for positive selection of amino acid replacements was found in the data with codeml (Yang 1997). While nine out of ten BTV segments supported the hypothesis of variable \( \omega \) among sites with significant better support for M3 compared to M0, a high proportion of sites was inferred to correspond to \( \omega < 1 \) (Table 2). The model M2a (positive selection) did not provide a significantly better fit for any segment than the nearly neutral model (M1a) which assumes a combination of purifying selection and neutral sites (\( P \)-values > 0.9). Similarly, the model M8 did not provide better fit than M7 (\( P \)-values > 0.09) (Supplementary Table S4). Further support for the BTV genomic segments being predominantly subject to purifying selection came from the mean \( d_{s}/d_{a} \) ratios, which ranged between 0.01 and 0.31 across the ten ORF (Table 2). Notably, Seg-10 alternate ORF, NS5, showed a higher \( d_{s}/d_{a} \) ratio of 6.56 (Table 2).

Positively and negatively selected codons were estimated by the FUBAR, FEL, and MEME methods of the HyPhy suit (Kosakovsky Pond, Frost, and Muse 2005). Only a single site on Seg-9 was inferred to be under positive selection by the FUBAR method (Table 3). FEL detected three more positively selected codons on Seg-2. MEME, which identifies fixed amino acid substitutions as well as codons under more sporadic positive selection, found a higher number of sites compared to other methods but these numbers remained low: six codons for Seg-1 and Seg-9, five for Seg-4, four for Seg-2, and one or two for the remaining segments (Table 3). Consistent with previous results, all segments contained an abundance of negatively selected codons ranging from 23 to 92 per cent for the FUBAR method and from 3 to 85 per cent for the FEL method. The alternative ORF for NS5 did not show evidence of negatively selected codons.

Motivated by the striking observation that a single variant of Seg-5 appeared to have replaced all other variants, we examined codon substitutions of the NS1 gene ORF (Seg-5) that could be responsible for conferring higher viral fitness to the dominant variant using DnaSP v6 (Librado and Rozas 2009). We were able to identify a single non-synonymous change at the tree node delineating the emerging cluster which involved the replacement of an aspartic acid by a glutamic acid at codon position 322 (nucleotide position 966) (Fig. 3). This codon was not among the sites inferred to be under either positive of negative selection in our previous analysis (data not shown).

4. Discussion

Previous studies based on field isolates of BTV have shown that reassortment is common where multiple distinct strains of the virus circulate and/or live attenuated vaccines have been used (Heidner et al. 1991; Nomikou et al. 2015; Schirtzinger et al. 2018). Our findings confirm that genetic exchange by reassortment has had a major impact on the genomic composition of Indian BTV strains. The current work further indicates that many of these reassortment events must have occurred in recent decades given that our molecular clock estimates for all ten segments point to most recent common ancestors of Indian isolates dating to the middle of the 20th century, coinciding with dates of the first reports of BT disease on the Indian subcontinent in the 1950s and 1960s (Sarwar 1962; Sapre 1964). This suggests that most BTV strains currently circulating in India might have been introduced during that period. However, we cannot exclude the possibility that other strains had been present previously but that their diversity during that time was reduced to a single lineage or that these earlier strains might have since become extinct (Holmes 2003), resulting in an underestimate of the real origin of segments.

Examining the correlation among ages of most recent common ancestors using MDS suggests that two core sets of segments (Seg-1, Seg-2, Seg-6, and Seg-7 on one hand and Seg-3, Seg-9, and Seg-10 on the other hand) are less permissive to be broken up by reassortment than a third set comprising Seg-4 and Seg-5. These groups of associated segments inferred for Indian BTV virus segments are noticeably different from what was observed for BTV in Europe (Nomikou et al. 2015) where Seg-1, Seg-3, Seg-4, Seg-5, Seg-8, and Seg-9 grouped together while Seg-7 and Seg-10 failed to show any clear association with other segments and Seg-2 and Seg-6 showed weak evidence for a connection. Whether these differences are due to biological reasons or chance will require further investigation.

While reassortment in segmented-RNA viruses appears to often be a selectively neutral process (Marshall et al. 2013; Shaw et al. 2013), a specific combination of viral segments might quickly rise to dominance if it results in a fitness advantage for the virus (McDonald and Patton 2011; Shaw et al. 2013; Anbalagan et al. 2014; Lu, Lyckett, and Leigh Brown 2014; Lago et al. 2017). Our results show that a particular BTV Seg-5 variant has strongly increased in frequency over the past decades, consistent with this kind of selective sweep scenario. We found Indian BTV strains to have highly admixed genomes and our MDS plot analysis shows that Seg-5 does not share its evolutionary history with any other segment. A previous study had suggested that the now dominant variant of Seg-5 was introduced into India as part of an exotic western virus strain or
It is therefore plausible that frequent reassortment events have allowed the successful Seg-5 variant to become associated with a large spectrum of genomic backgrounds in India. Indeed, all but three of the sixty-three BTV viruses collected after 2008 contained this variant (Fig. 3). Unexpectedly, the number of tree incompatibilities detected (and thus the number of reassortment events) involving Seg-5 is quite low compared to other segments (Fig. 5, Supplementary Fig. S11). One possible explanation is that the proposed takeover by the dominant variant occurred very rapidly and, by replacing all other strains, has limited the opportunity to detect any further reassortment events involving Seg-5.

While a genomic variant can also increase in a population as a consequence of positive selection, we found no indication for this in Seg-5. Instead, the observed increase in nucleotide diversity in this segment over time appears to be due to a population expansion following a bottleneck. Indeed, to understand the role of positive selection on evolution of Seg-5 of Indian BTV strains, we analysed sequences using a suite of methods (FUBAR, SLAC, and FEL), all of which failed to identify any positively selected codons. The MEME method identified only one site with evidence of episodic diversifying selection (codon 521). In contrast, most codons were identified to be under negative/purifying selection. This indicates strong structural and functional constraints for NS1 to evolve and suggests that the putative fitness benefits conferred by the invading Seg-5 variant applied regardless of the genotypic makeup of the other segments and did not require any further mutational changes of

Figure 3. Changes in the phylogenetic diversity of Indian BTV Seg-5 sequences across time and space indicates that a single variant has risen to dominance in recent decades. (A) Seg-5 time-scaled MCC tree of the ninety-two Indian BTV isolates, which was obtained using Bayesian phylogenetic inference in program BEAST (Drummond et al. 2012). Sequences could be divided into four amino-acid based clusters (Pessia et al. 2015) represented by different colours on the tree. According to ancestral state reconstruction of Seg-5 amino acid sequences, the branch defining the largest cluster (shown in red) only includes a single unique non-synonymous change in codon 322 (Asp → Glu). The inferred amino acid states in this position across other branches of the phylogeny are also shown. (B) Changes in the proportions of samples belonging to each of the four Seg-5 clusters over the past four decades, documenting the continuous rise in frequency of the red cluster, which has become the only variant found among samples collected since 2014. (C) The spatial distribution of Seg-5 clusters before and after 2005, confirming that the recent dominance of the red cluster applies across the Indian sub-continent.
This supports the hypothesis that the invasive variant of Seg-5 underwent a recent selective sweep based on pre-existing genotypic and phenotypic characteristics.

The Seg-5 phylogeny of Indian BTV isolates revealed a single non-synonymous change in the emerging variant at the internal node corresponding to the basal branch of the cluster (codon 322, amino acid change: aspartic acid -> glutamic acid, Fig. 3). Since this replacement is the only one that separates this variant from the other Seg-5 variants it has replaced, one might expect that this change must be responsible for the apparent selective advantage. However, more basal branches in the Seg-5 phylogeny and some of the older sequences of BTV-1 and BTV-16 also harboured a nucleotide sequence coding for a glutamic acid at this codon position (green cluster in Fig. 3). At least two non-exclusive hypotheses might explain this. First, the selective environment in which BTV circulates could have changed during the last two decades (e.g. host immune status changes, increase of the viral diversity, or modification of the vector distribution and activity patterns) and a specific amino-acid could have become advantageous in this new environment when it had not been before. Second, it is possible that the variant success observed in recent years might be due to epistasis, i.e. to specific interactions with concurrent substitutions in other BTV proteins. Epistatic effects play a central role in shaping the path of viral evolution in fitness landscapes of RNA viruses (Michalakis and Roze 2004; Sanjuan, Moya, and Elena 2004; Elena, Solé, and Sardanyés 2010) but are generally difficult to establish. Further in vitro experiments are required to elucidate the specific mechanisms underlying the differences in fitness and selection across Indian BTV genomes and high-throughput genetics approaches could be used to identify potentially relevant epistatic interactions involved (Wu et al. 2016).

An earlier study had reported that the estimated substitution rates for global isolates of BTV were comparable across segments, except for an order of magnitude lower rate for Seg-6 (5.22E-5) (Carpi, Holmes, and Kitchen 2010). Our estimates for BTV in India indicate that genes encoding for proteins interacting with host immunity (outer capsid) on Seg-2 and Seg-6 evolve at higher mean rates (1.47E-2 and 9.36E-3, respectively) and have longer TMRCA’s than genes on other segments (2.25E-3, Table 1). Whereas variants of outer capsid proteins will be maintained by balancing selection, those of inner capsid and non-structural genes are likely to be replaced continuously by new introduced variants through serial hitchhiking. This might result in an allele changing in frequency not because it is under natural selection itself, but...
because it is associated with another gene that is undergoing a selective sweep- and genetic drift.

Deviation from neutrality tests also showed two contrasting patterns; Seg-2, Seg-6 as well as Seg-7 (which encodes for the intermediate capsid protein VP7) displayed positive $D$ values, whereas other segments displayed negative $D$ values or values close to 0. Combined with other selection detection methods, our results provide evidence for differences in the predominant mechanisms of evolution of these genes. Outer capsid protein genes may be subject to positive and balancing selection leading to the generation of diverse serotypes and their variants, whereas evolution of all other genes appears to be driven by a combination of neutral and purifying selection. This is consistent with previous studies where purifying selection was reported to be one of the major evolutionary mechanisms in viruses including BTV (e.g. Balasuriya et al. 2008; Worobey, Han, and Rambaut 2014).

In conclusion, our comprehensive analysis of BTV evolution in India, a global hotspot of BTV diversity, provides new insights into the role of reassortment in driving rapid evolutionary change in segmented viruses. Significantly, we present strong evidence for a selective sweep event that has allowed a single
variant of NS1 to invade all BTV serotypes currently circulating in India through reassortment, though the basis for this apparent selective advantage remains yet unclear. The spread of this variant throughout the Indian BTV population may be related to the observation of more virulent BTV forms in India (Prasad et al. 2009; Chand et al. 2015; Rao et al. 2016). Our work has thus laid the groundwork for future studies that should focus on characterising the mechanisms underlying differences in fitness and selection across the BTV genome.

**Ethics**

This research project did not require any ethical approval.

**Acknowledgements**

We thank Dr Y. Krishnamohan Reddy (Tamil Nadu, India) for helping collecting samples from Tamil Nadu and Prof. Massimo Palmarini for his support and constructive comments.
Funding
This work was supported by BBSRC/DBT grant (BB/L004690/1 to P.M.) and by the Government of India grant (BT/IN/Indo-UK/FADH/46/SM/2013 to S.M.). S.Y. received support from an ICAR International Fellowship awarded by Government of India. Work at MRC-CVR, UK was supported by Wellcome Trust, grant (092806/B/10/2). Interactions between co-authors were also supported by the EU H2020 PALE-Blu grant (project No: 727393-2).

Data availability
Samples information and GenBank accession numbers of the ninety-two genomes used in this study are summarised in Supplementary Table S1.

Supplementary data
Supplementary data are available at Virus Evolution online.

Conflict of interest: None declared.

References
Anbalagan, S. et al. (2014) 'Whole Genome Analysis of Epizootic Hemorrhagic Disease Virus Identified Limited Genome Constellations and Preferential Reassortment’, The Journal of General Virology, 95: 434–41.
Attoui, H., Billoir, F., and Cantaloube, J. F. (2000) 'Strategies for the Sequence Determination of Viral dsRNA Genomes’, Journal of Virological Methods, 89: 147–58.
Bahl, J. et al. (2013) ‘Influenza A Virus Migration and Persistence in North American Wild Birds’, PLoS Pathogens, 9: e1003570.
Balasuriya, U. B. R. et al. (2008) ‘The NS3 Proteins of Global Strains of Bluetongue Virus Evolve into Regional Topotypes through Negative (Purifying) Selection’, Veterinary Microbiology, 126: 91–100.
Bankevich, A. et al. (2012) ‘SPAdes: A New Genome Assembly Algorithm and Its Applications to Single-Cell Sequencing’, Journal of Computational Biology, 19: 455–77.
Baten, C. A. et al. (2008) ‘A European Field Strain of Bluetongue Virus Derived from Two Parental Vaccine Strains by Genome Segment Reassortment’, Virus Research, 137: 56–63.
Belhouchet, M. et al. (2011) ‘Detection of a Fourth Orbivirus Non-Structural Protein’, PLoS One, 6: e25697.
Bhattacharya, B., and Roy, P. (2013) ‘Cellular Phosphoinositides and the Maturation of Bluetongue Virus, a Non-Enveloped Capsid Virus’, Virology Journal, 10: 1–11.
Boyce, M., Celma, C. C. P., and Roy, P. (2012) ‘Bluetongue Virus Non-Structural Protein 1 Is a Positive Regulator of Viral Protein Synthesis’, Virology Journal, 9: 178.
Butan, C., and Tucker, P. (2010) ‘Insights into the Role of the Non-Structural Protein 2 (NS2) in Bluetongue Virus Morphogenesis’, Virus Research, 151: 109–17.
Carpi, G., Holmes, E., and Kitchen, A. (2010) ‘The Evolutionary Dynamics of Bluetongue Virus’, Journal of Molecular Evolution, 70: 583.
Celma, C. C. P., and Roy, P. (2009) ‘A Viral Nonstructural Protein Regulates Bluetongue Virus Trafficking and Release’, Journal of Virology, 83: 6806–16.
Chand, K. et al. (2015) ‘Bluetongue in India: A Review’, Advances in Animal and Veterinary Sciences, 3: 605–12.
Delport, W. et al. (2010) ‘Datamonkey 2010: A Suite of Phylogenetic Analysis Tools for Evolutionary Biology’, Bioinformatics, 26: 2455–7.
Drummond, A. J. et al. (2012) ‘Bayesian Phylogenetics with BEAUti and the BEAST 1.7’, Molecular Biology and Evolution, 29: 1969–73.
Elena, S. F., Solé, R. V., and Sardanyés, J. (2010) ‘Simple Genomes, Complex Interactions: Epistasis in RNA Virus’, Chaos: An Interdisciplinary Journal of Nonlinear Science, 20: 026106.
Gauthami, S. et al. (2015) ‘MPGIT: A Concentrated Monophasic Phenol-Guanidinium Isothiocyanate Reagent for Isolation of Viral RNA’, Indian Journal of Biotechnology, 14: 376–81.
Han, Z., and Harty, R. N. (2004) ‘The NS3 Protein of Bluetongue Virus Exhibits Viroporin-like Properties’, The Journal of Biological Chemistry, 279: 43092–7.
Heidner, H. W. et al. (1991) ‘Genetic Variation and Evolutionary Relationships Amongst Bluetongue Viruses Endemic in the United States’, Virus Research, 21: 91–109.
Heled, J., and Bouckaert, R. R. (2013) ‘Looking for Trees in the Forest: Summary Tree from Posterior Samples’, BMC Evolutionary Biology, 13: 221.
Hemadri, D. et al. (2017) ‘Dual Infection with Bluetongue Virus Serotypes and First-Time Isolation of Serotype 5 in India’, Transboundary and Emerging Diseases, 64: 1912–7.
Holmes, E. C. (2003) ‘Molecular Clocks and the Puzzle of RNA Virus Origins’, Journal of Virology, 77: 3893–7.
Kosakovsky Pond, S. L., and Frost, S. D. W. (2005) ‘Not So Different after All: A Comparison of Methods for Detecting Amino Acid Sites under Selection’, Molecular Biology and Evolution, 22: 1208–22.
Kumar, S., Stecher, G., and Tamura, K. (2016) ‘MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for Bigger Datasets’, Molecular Biology and Evolution, 33: 1870.
Lago, M. et al. (2017) ‘In Vitro Reassortment between Infectious Pancreatic Necrosis Virus (IPNV) Strains: The Mechanisms Involved and Its Effect on Virulence’, Virology, 501: 1–11.
Kumar, S., Stecher, G., and Tamura, K. (2016) ‘MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for Bigger Datasets’, Molecular Biology and Evolution, 33: 1870.
Librado, P., and Rozas, J. (2009) ‘DnaSP v5: A Software for Comprehensive Analysis of DNA Polymorphism Data’, Bioinformatics (Oxford, England), 25: 1451–2.
Lourenco, S., and Roy, P. (2011) ‘In Vitro Reconstitution of Bluetongue Virus Infectious Cores’, Proceedings of the National Academy of Sciences of the United States of America, 108: 13746–51.
Liang, Y., Lyczek, S. J., and Leigh Brown, A. J. (2014) ‘Reassortment Patterns of Avian Influenza Virus Internal Segments among Different Subtypes’, BMC Evolutionary Biology, 14: 16–5.
Sarwar, M. M. (1962) ‘A Note on Bluetongue in Sheep in West Pakistan’, Pakistan Journal of Animal Sciences, 1: 1–2.

Schirtzinger, E. E. et al. (2018) ‘Recent US Bluetongue Virus Serotype 3 Isolates Found Outside of Florida Indicate Evidence of Reassortment with Co-Circulating Endemic Serotypes’, The Journal of General Virology, 99: 157–68.

Schweiger, B., Bruns, L., and Meixenberger, K. (2006) ‘Reassortment between Human A(H3N2) Viruses Is an Important Evolutionary Mechanism’, Vaccine, 24: 6681–90.

Shapiro, B., Rambaut, A., and Drummond, A. J. (2006) ‘Choosing Appropriate Substitution Models for the Phylogenetic Analysis of Protein-Coding Sequences’, Molecular Biology and Evolution, 23: 7–9.

Shaw, A. E. et al. (2013) ‘Reassortment between Two Serologically Unrelated Bluetongue Virus Strains Is Flexible and Can Involve Any Genome Segment’, Journal of Virology, 87: 543–57.

Sievers, F. et al. (2011) ‘Fast, Scalable Generation of High-Quality Protein Multiple Sequence Alignments Using Clustal Omega’, Molecular Systems Biology, 7: 539.

Simonsen, L. et al. (2007) ‘The Genesis and Spread of Reassortment Human Influenza A/H3N2 Viruses Conferring Adamantane Resistance’, Molecular Biology and Evolution, 24: 1811–20.

Smith, G. J. D. et al. (2009) ‘Origins and Evolutionary Genomics of the 2009 Swine-Origin H1N1 Influenza A Epidemic’, Nature, 459: 1122–5.

Stewart, M. et al. (2015) ‘Characterization of a Second Open Reading Frame in Genome Segment 10 of Bluetongue Virus’, The Journal of General Virology, 96: 3280–93.

Suyama, M., Torrents, D., and Bork, P. (2006) ‘PAL2NAL: Robust Conversion of Protein Sequence Alignments into the Corresponding Codon Alignments’, Nucleic Acids Research, 34: W609–12.

Tabachnick, W. J. (1996) ‘Culicoides Variipennis and Bluetongue Virus Epidemiology in the United States’, Annual Review of Entomology, 41: 23–43.

(2004) ‘Culicoides and the Global Epidemiology of Bluetongue Virus Infection’, Veterinaria Italiana, 40: 144–50.

Tajima, F. (1989) ‘Statistical Method for Testing the Neutral Mutation Hypothesis by DNA Polymorphism’, Genetics, 123: 585–95.

Trifonov, V. et al. (2009) ‘The Origin of Recent Swine Influenza A (H1N1) Virus Infecting Humans’, Eurosurveillance, 14: 191–3.

Velthuis, A. G. J. et al. (2010) ‘Financial Consequences of the Dutch Bluetongue Serotype 8 Epidemics of 2006 and 2007’, Preventive Veterinary Medicine, 93: 294–304.

Wirblich, C., Bhattacharya, B., and Roy, P. (2006) ‘Nonstructural Protein 3 of Bluetongue Virus Assists Virus Release by Recruiting ESCRT-I Protein Tsg101’, Journal of Virology, 80: 460–73.

Wong, W. S. W. et al. (2004) ‘Accuracy and Power of Statistical Methods for Detecting Adaptive Evolution in Protein Coding Sequences and for Identifying Positively Selected Sites’, Genetics, 168: 1041–51.

Worobey, M., Han, G. Z., and Rambaut, A. (2014) ‘A Synchronized Global Sweep of the Internal Genes of Modern Avian Influenza Virus’, Nature, 508: 254–7.

Wu, D. et al. (2015) ‘Poultry Farms as a Source of Avian Influenza A (H7N9) Virus Reassortment and Human Infection’, Scientific Reports, 5: 1–8.

Wu, N. C. et al. (2016) ‘Coupling High-Throughput Genetics with Phylogenetic Information Reveals an Epistatic Interaction on the Influenza a Virus M Segment’, BMC Genomics, 17: 1–15.

Wu, X. et al. (1992) ‘Multiple Glycoproteins Synthesized by the Smallest RNA Segment (S10) of Bluetongue Virus’, Journal of Virology, 66: 7104–12.

Yang, Z. (1997) ‘PAML: A Program Package for Phylogenetic Analysis by Maximum Likelihood’, Cabios Applications Note, 13: 555–6.

—, and Bielawski, J. P. (2000) ‘Statistical Methods for Detecting Molecular Adaptation’, Trends in Ecology & Evolution, 15: 496–503.

—, Wong, W. S. W., and Nielsen, R. (2005) ‘Bayes Empirical Bayes Inference of Amino Acid Sites under Positive Selection’, Molecular Biology and Evolution, 22: 1107–18.

Yu, G. et al. (2017) ‘GGTREE: An R Package for Visualization and Annotation of Phylogenetic Trees with Their Covariates and Other Associated Data’, Methods in Ecology and Evolution, 8: 28–36.