Identification of the Genome Segments of Bluetongue Virus Serotype 26 (Isolate KUW2010/02) that Restrict Replication in a Culicoides sonorensis Cell Line (KC Cells)

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Abstract

Bluetongue virus (BTV) can infect most ruminant species and is usually transmitted by adult, vector-competent biting midges (Culicoides spp.). Infection with BTV can cause severe clinical signs and can be fatal, particularly in naïve sheep and some deer species. Although 24 distinct BTV serotypes were recognized for several decades, additional ‘types’ have recently been identified, including BTV-25 (from Switzerland), BTV-26 (from Kuwait) and BTV-27 from France (Corsica). Although BTV-25 has failed to grow in either insect or mammalian cell cultures, BTV-26 (isolate KUW2010/02), which can be transmitted horizontally between goats in the absence of vector insects, does not replicate in a Culicoides sonorensis cell line (KC cells) but can be propagated in mammalian cells (BSR cells). The BTV genome consists of ten segments of linear dsRNA. Mono-reassortant viruses were generated by reverse-genetics, each one containing a single BTV-26 genome segment in a BTV-1 genetic-background. However, attempts to recover a mono-reassortant containing genome-segment 2 (Seg-2) of BTV-26 (encoding VP2), were unsuccessful but a triple-reassortant was successfully generated containing Seg-2, Seg-6 and Seg-7 (encoding VP5 and VP7 respectively) of BTV-26. Reassortants were recovered and most replicated well in mammalian cells (BSR cells). However, mono-reassortants containing Seg-1 or Seg-3 of BTV-26 (encoding VP1, or VP3 respectively) and the triple reassortant failed to replicate, while a mono-reassortant containing Seg-7 of BTV-26 only replicated slowly in KC cells.

Introduction

Arthropod-borne viruses (arboviruses) are transmitted between their vertebrate hosts (e.g. mammals or birds) by hematophagous arthropod vectors, especially mosquitoes, midges or ticks. Unlike other viruses, they require the ability to complete their replication cycle in two disparate host-species. The arboviruses are predominately RNA viruses [1] belonging to the
families Flaviviridae, Togaviridae, Bunyaviridae, Rhabdoviridae, and Reoviridae, along with a single DNA virus (African swine fever virus) belonging to the family Asfarviridae [1].

Bluetongue virus is the ’type species’ of the genus Orbivirus within the family Reoviridae [2]. The bluetongue virus (BTV) can infect all ruminant species, as well as camels and occasionally large carnivores [3–5]. Clinical signs of bluetongue disease (BT) are more severe in naïve animals, and are most commonly observed in sheep, and in white-tailed deer (e.g. in North America), although they are also seen (less frequently) in cattle and other species [6, 7]. The normal route of BTV transmission is via adults of vector-competent species of biting midge (Culicoides spp.), in which the virus also replicates. In addition, BTV can be transmitted via an oral route, or by vertical transmission in its ruminant hosts [8, 9]. Orbiviruses usually establish persistent infections in their vectors with no deleterious effects, and phylogenetic analyses indicate that they have evolved by co-speciation with their arthropod vectors [10]. Although Culicoides are regarded as the major vector for BTV, the virus can also replicate in cells of other arthropods including mosquitoes, drosophila and ticks [11–14].

BTV particles are composed of three concentric protein shells, surrounding a genome composed of 10 linear segments of double-stranded (ds) RNA [15, 16]. The genome segments range in size from 3954 to 822 bp, and are identified as segment 1 to 10 (Seg-1 to Seg-10) in order of decreasing molecular weight [2]. The BTV genome codes for 7 virus-structural proteins (VP1 to VP7) and 5 distinct non-structural (NS) proteins (NS1, NS2, NS3/NS3a, NS4 and S10-ORF2) [17–19].

Sequencing and phylogenetic comparisons show that Seg-2, and to a lesser extent Seg-6, are the most variable components of the BTV genome (encoding BTV VP2—’outer-capsid protein 1’; and VP5—outer-capsid protein 2, respectively). The sequences of BTV Seg-2 divide into distinct clades that correlate with the virus serotype, and can be used to ’type’ novel isolates by sequencing and/or type-specific RT-PCR assays [20–22]. The sequences of Seg-2 from different BTV serotypes can be grouped into ’nucleotypes’ (nucleotypes A to L), which also reflect the serological relatedness / cross-reactions between different serotypes [20, 22, 23].

Structural proteins VP3 and VP7 (encoded by Seg-3 and Seg-7), form the sub-core and core-surface layers of the BTV particle respectively. These proteins are more highly conserved between serotypes than the outer-capsid proteins [2, 8, 21, 22, 24–26]. The core surface protein VP7 has been identified as an immuno-dominant Orbivirus species / serogroup specific antigen and is therefore targeted by most serological diagnostic assays to detect BTV [27]. Earlier phylogenetic studies have shown that the conservation of Seg-3 sequences, allows them to be used to identify the members of individual Orbivirus species [28, 29].

BTV also encodes three minor enzymatic proteins, which are also highly conserved and are assembled within the central space of the sub-core particle. These include the RNA-dependent RNA polymerase—VP1; the capping enzyme -VP4; and the putative helicase VP6, encoded by Seg-1, Seg-4 and Seg-9 respectively [30].

Five non-structural proteins have been identified in BTV infected cells (the tubule protein—NS1; the viral inclusion body matrix protein—NS2; the virus-release protein—NS3/NS3a; and two recently discovered protein NS4 and S10-ORF2) [15, 17, 19, 31, 32]. These NS proteins are highly conserved across different BTV strains and serotypes [33, 34], although NS3/NS3a (encoded by Seg-10) can be more variable in other Orbivirus species, and represents the second most variable protein (after VP2) of AHSV [35, 36].

All of the BTV genome segments show significant nucleotide-sequence variations that at least partially correlate with the geographic origins of the virus. This suggests that the initial emergence of individual BTV serotypes was followed by a significant period of geographic isolation, allowing mutations to accumulate and generating geographically distinct virus lineages or ‘topotypes’ [20–22, 37].
Since 1998, multiple BTV types have emerged within Europe, events that have been linked to increased international trade, movement of insects and/or climate change in the region. This raised concerns about future threats posed by bluetongue and other related Culicoides transmitted diseases [8, 9, 38, 39]. During early 2008, an atypical BTV was detected in apparently healthy goats from the Toggenburg region of north eastern Switzerland [SWI/2008/01] [40]. Sequence analyses identified a novel BTV 'type' and showed that it did not belong to the 'major' eastern or western BTV topotypes previously identified [22]. Infection was experimentally confirmed in goats and the virus caused mild bluetongue-like disease in sheep, although subsequent attempts to propagate the virus in mammalian or insect cell cultures were unsuccessful [41, 42].

In February 2010, another orbivirus was successful isolated from sheep in Kuwait exhibiting clinical signs consistent with BT, by injection of washed blood into embryonated chicken eggs followed by passage onto BHK-21 cells (virus isolate KUW2010/02) [43]. Genomic dsRNA extracted from infected cells, generated an electrophoretic migration pattern in agarose-gels, typical for a BTV isolate. Identification of KUW2010/02 as BTV was confirmed by indirect antigen-sandwich ELISA, targeting the outer-core protein VP7. Whole genome sequence analysis of KUW2010/02 revealed 98% amino acid (aa) identity with BTV-25 in VP7, indicating a common ancestry, but only 81.2% identity in Seg-7, suggesting that these two viruses had diverged a long time ago and indicating high conservation-pressures on the aa sequence, structure and function of VP7 [44]. Comparisons of Seg-2 and Seg-6 sequences with those of other BTV serotypes, identified KUW2010/02 as a distinct 26th BTV 'type', within a 12th Seg-2 nucleotype (L) and a 9th Seg-6 nucleotype (I) [44]. Virus neutralization tests (VNT), using antisera against the existing 25 BTV types failed to neutralise KUW2010/02, confirming its identity as BTV-26 [43]. However, phylogenetic analyses showed a high level of divergence in most of the conserved genome-segments between most BTV strains and BTV-26 (KUW2010/02) and/or BTV-25 [SWI/2008/01], placing them as representatives of two novel and distinct BTV topotypes [44]. In addition, a novel BTV strain has recently been identified that represents a putative BTV serotype 27 [45].

The kinetics of BTV-26 infection in sheep and goats are similar to those of BTV-25. However, the virus can be horizontally transmitted to uninfected, in-contact goats, which subsequently seroconvert [46, 47]. This indicated that unlike BTV-25, BTV-26 can be transmitted horizontally by direct contact and replicates in mammalian cells (BHK-21, BSR and Vero cells) in vitro, but does not replicate in KC cells.

Reverse genetics was used to generate mono-reassortants between BTV-26 and the reference strain of BTV-1 (Western topotype—which replicates well in both mammalian and insect cell cultures) (see Results). The time course of replication was assessed for each reassortant strain in mammalian and insect vector cell lines, to identify individual genome segments of BTV-26 that restrict the ability of the virus to replicate in Culicoides cells.

Materials and Methods

Virus propagation and cells

The parental virus strains used were obtained from the dsRNA virus collection at IAH Pirbright (see: www.reoviridae.org/dsRNA_virus_proteins/ReoID/BTV-isolates.htm). BTV-26 [KUW2010/02] was originally isolated from the blood of a sheep from Kuwait as previously described [43]. BTV-1 [RSArrrrr/01] is the reference strain of BTV serotype 1, originally from South Africa.

Viruses were routinely propagated in BHK-21 cells (Baby hamster kidney, clone 13), obtained from the European Collection of Animal cell Cultures (ECACC– 84100501) in
Glasgow MEM medium (Invitrogen) with 10% heat-inactivated foetal calf serum (FCS). BSR cells, a clone of BHK-21 cells [48], were grown in Dulbecco’s modified Eagle medium (DMEM) containing 5% FCS. KC cells, derived from C. sonorensis midges [49], were grown in Schneider’s medium containing 10% FCS with penicillin (100 U/ml)/streptomycin (100 μg/ml) and fungizone (2 μg/ml). Aedes albopictus (mosquito) C6/36 cells [50] were grown in Leibovitz’s L15 medium (PAA) with 10% FCS, 10% tryptose phosphate broth solution (Sigma T8159), and penicillin/streptomycin. Mammalian cells were incubated at 37°C in air with 5% CO₂, and insect cells were incubated at 28°C.

**Preparation of viral dsRNA**

Viral dsRNA was extracted from virus-infected cell cultures (grown in T175 flasks) showing advanced cytopathic effect (CPE) using Trizol reagent (Gibco BRL) [51]. Briefly, dsRNA was separated from contaminating ssRNA by precipitating in 2M lithium chloride (LiCl). Then dsRNA in the supernatant was precipitated by addition of three volumes of 100% ethanol and 0.25 volumes of 7.5M ammonium acetate (final ammonium acetate concentration of 0.44 M), washed twice with 70% ethanol and re-suspended in RNase-free water. In addition, viral RNA was purified from small volumes of cell culture medium using a Kingfisher robot (Thermo Scientific) with a MagVet Universal Isolation Kit (Laboratoire Service International).

**Full-length amplification of cDNAs (FLAC)**

Full length cDNA copies of BTV genome segments were synthesised and amplified, after ‘anchor spacer–ligation’ as described previously [52].

**Preparation of cDNA clones and DNA sequence determination**

Complete genome segments were amplified from the cDNA of BTV-1 RSArrrr/01 and BTV-26 KUW2010/02 by PCR using the primers shown in Table 1. The upstream primers contained a T7 promoter sequence and a restriction site for unidirectional cloning, while the downstream primers contained restriction sites for cloning and for linearization at the 3’ end of the segment, as described [53]. PCR was carried out using a PCR extender system (5 Prime) and the PCR products were double digested with the appropriate restriction enzymes. These were ligated into pGEX-4T-2, digested with the same enzymes, and transformed into competent cells of E. coli STBL2 (Invitrogen). Colonies obtained on ampicillin-containing agarose plates were confirmed by plasmid purification (Qiagen) and DNA sequencing (using the BigDye Terminator v3.1 Cycle Sequencing Kit, Invitrogen). Alternatively, plasmid clones for some segments were synthesized by GeneArt (see results section for details).

**Reverse genetics and confirmation of recombinant viruses**

Capped positive sense transcripts were generated by linearization of plasmid DNA from each clone, by digestion with appropriate restriction enzymes, followed by in vitro transcription using a mMessage mMACHINE T7 Ultra Kit (Life Technologies) as described [53]. To rescue reassortant viruses, BSR cells at approximately 90% confluence were transfected with a mixture of the appropriate capped transcripts, in a two-step procedure [53]. First, transcripts from segments 1, 3, 4, 5, 8 and 9 (coding for VP1, VP3, VP4, NS1, NS2 and VP6) of the required reassortant were transfected (400 ng each in a 6 well format) using Lipofectamine 2000. After 18–20h, the cells were transfected with a mixture of RNAs containing all ten transcripts from the virus to be ‘rescued’ (400 ng of each). Transfected BSR cells were then incubated for 2–3 days, before passage onto fresh BSR cells (80–90% confluent) in T25 flasks as described previously.
These were examined daily for the appearance of CPE. For reassortants which failed to rescue in this way (those with segments 1–3 of BTV-26, see Results) attempts were made to modify the first transfection step. In these cases, expression plasmids, pCDNA3.1 containing open reading frames of Seg-1, 3, 4, 5, 8 and 9 from BTV-1 RSArrrr/01 were used in the first transfection step, instead of transcripts as described previously [55].

To confirm rescue of the required viruses, RNA was extracted from cultures showing advanced CPE using the Trizol method (see above), and cDNA was prepared from at least two

| Table 1. Sequences of cloning primers. |
|---------------------------------------|
| **Serotype, segment** | **Sequence (5′-3′)** |
| **BTV-1 [RSArrr/01]** | |
| Seg-1 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-1 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-2 forward | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-2 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-3 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-3 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-4 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-4 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-5 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-5 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-6 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-6 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-7 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-7 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-8 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-8 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-9 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-9 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-10 forward | TCTAGCCGATCTCTAATAGCACTCTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-10 reverse | GACGACGAGGCTAGTACATGATATATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| **BTV-26 [KUW2010/02]** | |
| Seg-2 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-2 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-4 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-4 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-5 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-5 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-6 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-6 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-7 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-7 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-8 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-8 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-9 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-9 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-10 forward | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |
| Seg-10 reverse | GTGATGGATCCTAATACGACTCACTATAGTTAAAATGCAATGGTCGCAATC |

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[54]. These were examined daily for the appearance of CPE. For reassortants which failed to rescue in this way (those with segments 1–3 of BTV-26, see Results) attempts were made to modify the first transfection step. In these cases, expression plasmids, pCDNA3.1 containing open reading frames of Seg-1, 3, 4, 5, 8 and 9 from BTV-1 RSArrr/01 were used in the first transfection step, instead of transcripts as described previously [55].

To confirm rescue of the required viruses, RNA was extracted from cultures showing advanced CPE using the Trizol method (see above), and cDNA was prepared from at least two
genome segments of the rescued viruses (including the heterologous segment from monoreassortants), using a Superscript III One-Step RT-PCR with Platinum Taq kit (Invitrogen) according to the manufacturer's instructions. Approximately 500–1000 bp regions were amplified by this method, and this was followed by DNA sequencing.

The identity of the rescued viruses was further verified by comparing migration of their purified viral dsRNA to the parental strains of BTV-1 and BTV-26, by electrophoresis on 11% SDS-PAGE gels [15].

**Plaque purification**

A volume of 0.5 ml of the suitable virus dilutions, prepared in DMEM with 5% FCS, were added to confluent monolayers of BSR cells in 12 well plates. After 90 min at 37°C with 5% CO₂, the diluted virus was removed and replaced with 3ml of MEM containing 3% FCS and 1.5% melted agarose type V11A (Sigma-Aldrich). Plates were incubated at 37°C with 5% CO₂ until plaques were visible. These were picked into DMEM/5% FCS, before passage on BSR cells.

**Virus titrations**

To determine the titre of stock viruses or supernatant samples from time-course experiments, DMEM (100µl) containing 1% FCS and penicillin / streptomycin was aliquotted into each well of 96 well plates. Virus (50µl) was added to the first well, then serially diluted across the plate to give half-log dilutions. Each virus was tested in 4 or 8 rows. The last well was a negative control with no virus. BSR cells in DMEM containing 5% FCS were then added to each well (3 x 10⁵ cells per well in 100µl). Plates were incubated for 6 days, and the number of wells with CPE was determined. TCID₅₀ values were calculated using the Karber formula [56]. For calculation of multiplicity of infection (MOI), a PFU/ml working estimate was calculated using the formula, one TCID₅₀ = 0.7 PFU/ml; [56]).

**Timecourse experiments**

Timecourse experiments were carried out for parent and reassortant in different cell lines. T25 flasks were seeded with 1.5 x 10⁶ BSR cells and after 24h (at approximately 80% confluence) were infected at an MOI of 0.04. After 30 mins at 37°C 250 µl samples (T₀ samples) of the medium were collected. The infected cells were further incubated at 37°C and samples of medium (250 µl) were taken at time intervals up to 96 h (at 8h, 24h, 32h, 48h, 56h, 72h and 96h). Insect cells were infected in suspension after determining the cell number using a haemocytometer. Infections were carried out at an MOI of approximately 0.04 and incubated for 30 mins at room temperature. T₀ samples were collected as for BSR cells and then the infected cells were transferred to T25 tissue culture flasks and incubated at 28°C. Samples of medium (250 microlitres) were collected at intervals up to 14 days post-infection.

The samples were stored at 4°C until the end of the experiment and then viral RNA was extracted from 100 µl volumes using a Kingfisher robot. Viral RNA was quantified using the segment 9-specific real-time RT-PCR assay [57] with the Superscript III One-Step qRT-PCR with Platinum Taq Kit (Invitrogen), and expressed as genome copies per reaction (each reaction contained RNA from 5 microlitres of tissue culture supernatant). To standardize the data, a dilution of BTV-1 [RSArrrr/01] viral RNA was included on every PCR plate. In one experiment in KC cells, infectious virus was also quantified in all samples by end-point titration on BSR cells to confirm that measurement of viral RNA replication gave comparable results to measurement of infectious virus, and was an appropriate method to quantify viral replication. The virus titers were expressed as tissue culture infective dose (TCID₅₀)/ml.
Results

Generation of plasmid clones

Most of the genome segments from BTV-1 [RSArrr/01] and BTV-26 [KUW2010/02] were successfully cloned as described in Material and Methods. The inserts were completely re-sequenced and those with the genome consensus sequence were selected where possible. A few clones had conservative changes in the coding regions (BTV-1 Seg-3, Seg-5 and Seg-8, and BTV-26 Seg-2 and Seg-8) but none had changes in the untranslated regions. Clones of Seg-1 and Seg-3 from BTV-26 were not obtained by this method. The relevant cDNA copies with upstream T7 promoters were therefore synthesized and subcloned (Gene Art) and plasmids containing the synthetic inserts were used directly for \textit{in vitro} transcription.

Generation of reassortant viruses using reverse genetics

Seven of the ten possible mono-reassortant viruses (containing a single BTV-26 genome-segment and nine BTV-1 segments) were obtained by the original procedure (see Materials and Methods). These reassortants contain segments 4 to 10 of BTV-26. Their identities were confirmed by partial DNA sequencing, and they are designated as BTV-1\textsubscript{26-S4}, etc, where the uppercase section of the name refers to the BTV-1 [RSArrr/01] backbone, with subscript referring to the added BTV-26 segment. Attempts to rescue mono-reassortants containing Seg-1 or Seg-3 of BTV-26 (BTV-1\textsubscript{26-S1} and BTV-1\textsubscript{26-S3}), using lipofection of RNA transcripts were unsuccessful. However, the required mono-reassortants were subsequently generated by use of ‘expression plasmids’ in the first transfection step. With both these reassortants, there was also a longer lag before CPE was visible (4–5 days after passage of the transfected wells) than for the other reassortants (approximately 2 days after passage).

Several attempts to rescue BTV-1\textsubscript{26-S2} were unsuccessful, despite using either transcripts or expression plasmids for the first transfection step. It was considered possible that VP2 of BTV-26 was incompatible with proteins VP5 and VP7 of BTV-1 (encoded by Seg-6 and Seg-7), with which it would interact in the virus particle [58, 59]. A triple reassortant containing all three segments from BTV-26 (BTV-1\textsubscript{26-S2}, S6, S7) was successfully rescued, although attempts to generate double reassortants BTV-1\textsubscript{26-S2}, S6 or BTV-1\textsubscript{26-S2}, S7 were unsuccessful. Generated reassortants are summarized in Table 2.

In addition, BTV-1 [RSArrr/01] virus was rescued by the original procedure, and this virus was used for all the subsequent analyses. However, despite many attempts, BTV-26

\begin{table}[h]
\centering
\caption{The genome-segment composition of reassortants generated between BTV-1 [RSArrr/01] and BTV-26 [KUW2010/02].}
\begin{tabular}{|c|c|c|c|c|c|c|c|c|c|}
\hline
\textbf{Virus} & \textbf{Seg-1} & \textbf{Seg-2} & \textbf{Seg-3} & \textbf{Seg-4} & \textbf{Seg-5} & \textbf{Seg-6} & \textbf{Seg-7} & \textbf{Seg-8} & \textbf{Seg-9} & \textbf{Seg-10} \\
\hline
BTV-1\textsubscript{26-S1} & - & 26 & - & - & - & - & - & - & - & - \\
BTV-1\textsubscript{26-S2,S6,S7} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S3} & 26 & - & - & - & - & - & - & - & - & - \\
BTV-1\textsubscript{26-S4} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S5} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S6} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S7} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S8} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S9} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
BTV-1\textsubscript{26-S10} & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 & 26 \\
\hline
\end{tabular}
\end{table}

26 indicates those genome-segments derived from BTV-26. Other segments are from BTV-1.

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[KUW2010/02] has so far not been rescued, and the originally isolated virus was therefore used throughout.

Most of the genome segments from the two parent viruses (BTV-1 and BTV-26) migrate at different rates during SDS-PAGE, with the exception of Seg-1 which could not be clearly differentiated. The parental and reassortant viruses that were generated were verified by SDS-PAGE of their dsRNA segments (Fig 1). The reassortants all showed the expected migration patterns, with the exception of BTV-126-S1 which had an extra band below the segment 3 band (indicated by an asterisk, Fig 1). This band was purified from an agarose gel and sequenced, which showed it was a truncated form of Seg-3. This virus was therefore plaque purified before further analysis (so that it contained only the full-length Seg-3—data not shown).

**Replication kinetics of reassortant BTV strains**

In BSR cells, the parental strains of BTV-1 [RSArrrr/01] and BTV-26 [KUW2010/02] both replicated well, reaching equivalent final genome copy numbers per reaction, although BTV-26 was somewhat slower with lower RNA copies at 24 to 72 hours post infection (h.p.i., Fig 2). Most of the reassortants also replicated well in these mammalian cells, with growth kinetics intermediate between the two parent viruses, although mono-reassortants containing Seg-3, 5, 6 or 7 of BTV-26, matched the slower growth rate of BTV-26. Only reassortant strain BTV-126-S8 (expressing NS2 of BTV-26) had a markedly slower replication rate, with a final genome copy number of over a log lower than either parental strain at 48 to 96 h.p.i.

The BTV-26 parental strain failed to replicate in KC cells, showing a steady, gradual decline in genome copy number per reaction (Fig 3). In contrast the BTV-1 parental strain did replicate but achieved a genome copy number of only ~ $10^7$ per 5 microlitres of tissue culture supernatant (by 7 days post infection (d.p.i.)), compared with over $10^6$ in BSR cells (by 4 d.p.i.) (Fig 3). Several of the reassortant strains replicated with similar kinetics to BTV-1 in KC cells, these included BTV-126-S8, BTV-126-S5, BTV-126-S6, BTV-126-S9 and BTV-126-S10. However, there was an initial lag in the replication (RNA synthesis) of BTV-126-S8 in KC cells (by approximately 3 days), which was not observed in BSR cells or with the parental BTV-1 strain, in either cell line. There was a significant delay (an eclipse period of at least 2 days, ~four days for BTV-126-S8) in the rise in infectivity of all of the strains that replicated in KC cells, (including the BTV-1 parental strain), with the same final titre achieved by BTV-1 and BTV-126-S8 at 6 d.p.i. in KC cells (Figs 3 and 4).

Four of the reassortant strains showed much reduced, or no replication in the *Culicoides* cells. BTV-126-S1 failed to replicate, showing a similar gradual reduction in viral RNA copies to
the BTV-26 parent strain, over 14 days (Fig 3). Titration results confirmed that virus infectivity also decreased throughout the incubation period (Fig 4). Reassortant BTV-1_{26-S3} and triple reassortant BTV-1_{26-S2, S6, S7} also failed to replicate productively in KC cells, with viral RNA levels and infectious virus titres that either showed a small decrease, or for strain BTV-1_{26-S2, S6, S7},
S7 were unchanging over the two week incubation period (Figs 3 and 4). In contrast, BTV-126-S7 was able to replicate in KC cells, but it did so at a substantially reduced rate (as assessed by both viral RNA levels and infectious virus titre) compared to the BTV-1 parent strain. The infectivity of BTV-126-S7 only increased by about 10 fold, in contrast to BTV-1 [RSArrr/01] which increased by ∼100 fold at its peak (Fig 4).
A time course for BTV-1 replication in C6/36 cells (Fig 5) showed levels of viral RNA synthesis comparable to those achieved in BSR cells (Fig 2), generating considerably larger amounts of RNA than produced in KC cells (Fig 3). Although BTV-26 also replicated in C6/36 cells, it was significantly slower than BTV-1, with levels of viral RNA only starting to increase at 4 days post-infection, and reaching a lower final titre at 14 days post-infection (Fig 5).
levels of BTV-26 replication in mosquito cells are comparable to those achieved by BTV-1 in KC cells.

All of the reassortants also replicated in C6/36 cells (Fig 5). Those strains that showed no significant increase, or even a decrease in virus titre in KC cells (BTV-126-S1; BTV-126-S2, S6, S7 and particularly BTV-126-S3) also replicated more slowly in C6/36 cells, although viral RNAs...
continued to rise up to 14 d.p.i. The other reassortants generated similar levels of viral RNA to BTV-1 by day 14, although with some initial delays. In particular BTV-1_{26-S8}, which replicated to lower levels in BSR cells and showed a significant delay in KC cells, also showed a delay of ~3 days, during the early stages of replication in C6/36 cells. BTV-1_{126-S7}, which showed a large delay in replication in KC cells, showed only a minor delay in replication in C6/36 cells.

Discussion

BTV-26 [KUW2010/02] replicates in mammalian cells (BHK or BSR cells) in vitro but unlike BTV-1 [RSArrrrr/01] and most other BTV strains that have been tested, does not replicate productively in KC cells, which are derived from the BTV-competent vector, *C. sonorensis*. A series of reassortant strains containing individual genome segments (or a selected combination of segments) from BTV-26, were generated within a BTV-1[RSArrrrr/01] genetic 'backbone', making it possible to identify individual genome segments that restrict replication in KC cells.

Initial attempts to rescue Seg-2 of BTV-26, as a mono-reassortant within the BTV-1 backbone, were unsuccessful. However, this segment was recovered as a component of BTV-1_{126-S2, S6, S7}, a triple reassortant that has both outer capsid proteins (VP2 and VP5) and the outer-core protein (VP7) from BTV-26. This suggests that physical interactions that are known to exist between VP2, VP5 and VP7 [58, 59] allow VP2 of BTV-26 to be incorporated and fully functional within the virus particle. BTV-1_{126-S2, S6, S7} replicates ‘normally’ in BSR cells, but failed to replicate in KC cells. Mono-reassortant BTV-1_{126-S6}, replicated in KC cells with similar kinetics to the BTV-1 parental strain, while BTV-1_{126-S7} replicates substantially slower in KC cells than BTV-1. This indicates that both VP2 and VP7 of BTV-26 can either restrict or reduce infection and/or replication in KC cells.

Previous reverse genetics studies between the more closely related Western-topotype strains of BTV-1 and BTV-8, showed that all possible mono-reassortants could be rescued by reverse genetics [60–62]. Further investigations will be needed to determine the relative importance of the major BTV topotypes in reassortment compatibility groups. Our findings match studies of natural reassortants of a closely related orbivirus species, epizootic hemorrhagic disease virus (EHDV), showing that the genome segments encoding VP2, VP5 and VP7 from serotype 1 and serotype 2, preferentially remain within the homologous serotype during co-infections, forming a reassortment ‘linkage group’ [63]. This is further supported by evidence that VP2 and to a lesser extent VP5, both show sequence variations that correlate with BTV serotype. Interestingly, in a recent study, a double reassortant containing just the genome segments for VP2 and VP5 of BTV-26 in the BTV-1 backbone was successfully rescued [64].

A primary function of outer-capsid protein VP2 is in cell attachment during initiation of infection (51). Analyses of its replication time-course suggest that BTV-26 may be unable to bind to or initiate infection in KC cells. If the virus failed to enter KC cells, and consequently did not ‘uncoat’ during the early stages of infection, it would presumably remain intact in the tissue culture media, which may explain the stable RNA signal observed throughout the BTV-1_{126-S2, S6, S7} time course in KC cells.

Although the triple reassortant replicates slowly in C6/36 mosquito cells, it was faster than the BTV-26 parent, suggesting that VP2 of BTV-26 can mediate attachment to C6/36 cells but possibly less efficiently than to BSR cells. The incorporation of VP5 or VP7 of BTV-26 reduced the initial replication rate of BTV-1_{126-S6} or BTV-1_{126-S7} in C6/36 cells, although both viruses reach the same plateau level as BTV-1 by day 14 p.i.

Although the surface of BTV core-particles is composed entirely of VP7 (25), cores can still infect either BHK-21 or C6/36 cells with low efficiency, but show a much higher specific infectivity, which is comparable to that of intact virus particles, in KC cells, indicating that VP7 can...
mediate cell attachment and penetration in these Culicoides cells [14]. The RGD motif present in VP7 has been reported as essential for cell-binding/entry by viral cores [65]. Our results show that although VP7 of BTV-26 still contains the RGD motif, it is associated with a reduced rate of replication in KC cells. It is unclear if this reduction reflects a reduced efficiency in the initiation of infection, a delay in replication, or a reduced efficiency of virus assembly within the host cell cytoplasm. Comparison of VP7 from the two parental strains of BTV-1 and BTV-26, show only 19 aa differences (94.6% amino acid identity: Table 3). The significance of these differences in the reduced rate of replication of BTV-126-S7 in KC cells is currently being investigated.

Two mono-reassortants, BTV-126-S1 and BTV-126-S3 were unable to replicate in KC cells. Since both reassortants have functional cell entry proteins (VP2, VP5 and VP7) of BTV-1 and can infect BSR cells efficiently, it appears likely that they can also initiate infection in KC cells. However, during time course experiments, the viral RNA concentration gradually decreased over the 14 day incubation period, indicating that they did not replicate and their genomes were gradually being degraded. Interestingly, these two viruses showed quite different replication kinetics in C6/36 cells, which provides some clues to the mechanisms involved. BTV-126-S3 showed no replication in C6/36 cells for the first 7 days, whereas BTV-126-S1 replicated relatively well in these cells. This suggests that the phenotype of BTV-126-S3 might not be specific to KC cells, but could be a result of the low temperature used for both insect cell lines compared to the BSR replication experiments. Since Seg-3 codes for the inner core protein, VP3, a protein essential for core assembly, it is possible that viral assembly is negatively affected at low temperature. In contrast, the phenotype of BTV-126-S1, which has the BTV-26 polymerase, is more specific to KC cells. This is difficult to explain since VP1 functions within cores. One hypothesis is that this mono-reassortant is more susceptible to the antiviral pathways in Culicoides cells. Further investigations will be required to elucidate the reasons for the replication-negative phenotypes of these viruses in KC cells.

It was interesting that both BTV-126-S1 and BTV-126-S3 were more difficult to rescue than the other mono-reassortants, although these rescued viruses subsequently replicated normally in BSR cells, suggesting they had normal protein functions. Similarly, it has been reported that the rescue of BTV containing particular mutations or deletions in Seg-10 was delayed [54, 66]. These studies showed that despite the successful rescue of deletions in the Seg-10 ORF, RNA inserts from several genome segments were found in Seg-10 very quickly after virus rescue.

| Segment          | % nucleotide identity | % amino acid identity |
|------------------|-----------------------|-----------------------|
| Seg-1 (VP1)      | 75.9                  | 88.3                  |
| Seg-2 (VP2)      | 50.6                  | 38.5                  |
| Seg-3 (VP3)      | 76.0                  | 88.5                  |
| Seg-4 (VP4)      | 73.7                  | 80.4                  |
| Seg-5 (NS1)      | 73.3                  | 80.3                  |
| Seg-6 (VP5)      | 67.4                  | 73.2                  |
| Seg-7 (VP7)      | 79.2                  | 94.6                  |
| Seg-8 (NS2)      | 72.1                  | 69.8                  |
| Seg-9 (VP6, NS4) | 71.4                  | 60.9 (VP6), 76.7 (NS4) |
| Seg-10 (NS3/3A)  | 79.2                  | 85.6                  |

Sequences for BTV-1 [RSArrrr/01] linked to Genbank accession numbers FJ969719 to FJ969728, and for BTV-26 [KUW2010/02] accession numbers JN255156 to JN255162 and HM590642 to HM590644.

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Although the reasons for the delayed rescue of BTV-126-S1 and BTV-126-S3 remain unknown, it is possible that there are changes on other segments which might be relevant, for example, the additional Seg-3 band that was observed upon rescue of BTV-126-S1 (Fig 1) might be important. Full genome sequencing might reveal other changes that could help explain the difficulties in virus rescue.

The mono-reassortant containing NS2 from BTV-26 (BTV-126-S8) replicated more slowly than BTV-1 [RSArrrr/01] in all three cell types, suggesting that the heterologous NS2 caused a general loss of fitness. This may result from a partial mismatch between NS2 of BTV-26 and the other viral components from BTV-1. NS2 of BTV-1 [RSArrrr/01] and BTV-26 have 107 amino acid changes, which could be responsible for such a mismatch. Another possibility is that interactions between BTV-26 NS2 and cellular components are less effective than with BTV-1 NS2. Substitution of Seg-8 in mono-reassortants between BTV-1 and BTV-8 also generated consistently smaller plaques than those of the two parental viruses [60].

The finding that the BTV-26 parental virus could not be rescued could be a result of its slightly slower replication in BSR cells compared to the BTV-1 parent (Fig 2). Alternatively, it might be a result of the cumulative negative effect of the conservative differences in some of the BTV-26 plasmid clones that together result in failure to rescue BTV-26.

Although BTV-26 and other BTV serotypes can replicate in C6/36 cells [13] the epidemiological significance is unclear, as mosquitoes are not usually regarded as an important vector for BTV. Initial studies indicate that BTV-26 cannot infect adult C. sonorensis, although mosquito infection studies are still in progress. The failure of BTV-26 to replicate in KC cells, or infect C. sonorensis midges (unpublished), suggests that it may have lost the ability to be transmitted by adult Culicoides, although we cannot exclude the possibility that other Culicoides spp. (for example those indigenous to Kuwait) might act as vectors. However, BTV-26 appears to have an increased ability compared to other BTV strains, to transmit horizontally, via direct animal to animal contact [46, 47]. It is possible that proteins responsible for the inability of BTV-26 to replicate in KC cells (VP1, VP2, VP3 and VP7) are also associated with the apparent increase in efficiency of contact transmission.

In summary, a reverse genetics approach was used to generate reassortants between BTV-1 and BTV-26, to investigate their replication in BSR, KC and C6/36 cells. Although, most genome segments of BTV-26 could be rescued as mono-reassortants in a BTV-1 backbone, there is evidence of reassortment-groups that reflect interactions between the different components of the virus-particle (for example VP2, VP5 and VP7). The significance of sequence differences in the RNAs and proteins involved in these groupings, between different serotypes or topotype, still need to be determined.

The inability of BTV-26 to infect/replicate in KC (C. sonorensis) cells was shown to be associated with differences in Seg-1/VP1, Seg-2/VP2, Seg-3/VP3 and Seg-7/VP7. This suggests that although BTV cell-attachment / entry mechanisms (which involve VP2), are effective for BTV-26 in BSR cells and C6/36 cells, they are unable to mediate infection in C. sonorensis cells, suggesting the absence of an appropriate receptor. The mechanisms that restrict replication of BTV reassortants containing VP1, VP3 or VP7 of BTV-26 in KC cells, have not been determined. It is possible that some component or mechanism present in BSR cells is essential for BTV replication, but is missing in KC cells. Alternatively some component or mechanism present in KC cells may be triggered by or target these RNAs or proteins (such as an innate immune responses, or silencing) restricting replication.

The identification of individual genome segments/proteins that restrict the replication of BTV-26 in KC cells, or in adult C. sonorensis (a known BTV-vector species), shows that vector-competence can vary dramatically for different BTV strains within an individual Culicoides species. It is also possible that the transmission efficiency of an individual BTV strain varies...
between different Culicoides species. This could be better explored if cell lines and colonized insects, were available for other Culicoides species. Variations in their transmission efficiency could potentially explain why the spread of different BTV strains/serotypes is not uniform, with certain strains/serotypes apparently restricted to a limited geographic range (e.g. the spread of BTV-2, 4, 9 and 16 in southern Europe), while others are more widely distributed (e.g. BTV-1, 8, and 14 in northern Europe). Some BTV strains/serotypes may be better adapted to different episystems [67] containing different Culicoides species or populations, or like BTV-26 may be capable of horizontal transmission in the absence of vector insects. Further analyses of the interactions between viral RNAs/proteins and components of the host cell, will help to elucidate the mechanisms involved in vector transmission and competence, as well as the viral proteins and mechanisms that can enhance horizontal transmission of certain BTV strains.

**Author Contributions**

Conceived and designed the experiments: GDP KN MB HA PPM. Performed the experiments: GDP MGB. Analyzed the data: GDP MGB. Contributed reagents/materials/analysis tools: KN MB HA. Wrote the paper: GDP MB HA PPM.

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