Complete Genome and Molecular Epidemiological Data Infer the Maintenance of Rabies among Kudu (Tragelaphus strepsiceros) in Namibia

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Abstract

Rabies in kudu is unique to Namibia and two major peaks in the epizootic have occurred since it was first noted in 1977. Due to the large numbers of kudu that were affected, it was suspected that horizontal transmission of rabies occurs among kudu and that rabies was being maintained independently within the Namibian kudu population – separate from canid cycles, despite geographic overlap. In this study, it was our aim to show, through phylogenetic analyses, that rabies was being maintained independently within the Namibian kudu population. We also tested, through complete genome sequencing of four rabies virus isolates from jackal and kudu, whether specific mutations occurred in the virus genome due to host adaptation. We found the separate grouping of all rabies isolates from kudu to those of any other canid species in Namibia, suggesting that rabies was being maintained independently in kudu. Additionally, we noted several mutations unique to isolates from kudu, suggesting that these mutations may be due to the adaptation of rabies to a new host. In conclusion, we show clear evidence that rabies is being maintained independently in the Namibian kudu population – a unique phenomenon with ecological and economic impacts.

Introduction

Rabies virus (RABV) is typically maintained within species of the mammalian families Carnivora and Chiroptera. The classical primary host species among carnivores is the dog (Canis familiaris), but several other carnivores are known to be able to maintain an independent cycle of RABV, for instance the red fox (Vulpes vulpes) in Europe [1] and raccoons (Procyon lotor) [2] and skunks (Mephitis mephitis) in North America, among others. With these classical trends, the emergence of an extensive rabies cycle in an herbivorous host is unusual. However, in 1977 a rabies epizootic in kudu emerged in Namibia [3] and, with several peaks and troughs in the numbers of observed cases through the decades, is still ongoing today. Due to the geographical extent and the numbers of animals affected, it was suspected that this rabies cycle was maintained within the kudu population.

For rabies, spill-over infections from carnivores to domestic and wild herbivores are known to occur regularly, but such spill-overs are invariably dead-end [4] and serve as an indicator of rabid carnivore (domestic or wildlife) activity [5]. Generally, there are several factors that are necessary for the adaptation and maintenance of rabies in a new host. One factor is the need for overlapping host ranges (sympathy) of a susceptible species with that of a species already maintaining a cycle of RABV [6]. It has also been suggested that the initial infection of a new host is aided by the similarity of the cellular and immunological traits of the new host with the donor, with the number of exposures (due to geographical overlap) being a secondary factor [6]. These factors would explain the relatively high rate of transmission of rabies to new canid hosts (e.g. fox, raccoon, skunk, bat-eared fox etc.) as opposed to non-canid hosts that share the same host range. Another factor that has been suggested is that of recipient host population densities [1,7].

A limited number of studies have aimed to determine whether specific mutations occur in the RABV genome due to the selective pressures caused by host switching and adaptation. One study noted mutations specific to certain fox species throughout Europe, specifically in the glycoprotein and nucleoprotein genes [1]. However, despite this phylogenetic clustering, different viruses predominantly clustered according to their geographical origin, suggesting that spatial and behavioral patterns of the host species play a greater role in the phylogenetic clustering of RABVs [1,8]. A second study showed evidence that two types of adaptation could occur: 1) Post-shift adaptation, where the virus mutates in order to better suit the new host; 2) Pre-shift adaptation, in which the virus adapts through convergent evolution to better suit the shift from one host to a new host [9]. It was shown that pre-shift
### Table 1. Brain samples from various species used in the phylogenetic analysis of partial RABV nucleoprotein gene sequences.

| Sample Number | GenBank Number | Species | Year of Isolation | Origin - District | Origin - Region | Country | Reference |
|---------------|----------------|---------|-------------------|-------------------|----------------|---------|-----------|
| 3K08          | JQ691415       | Kudu    | 2008              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 8K08          | JQ691416       | Kudu    | 2008              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 18K08         | JQ691417       | Kudu    | 2008              | Omaruru           | Erongo         | Namibia | This study |
| 19K08         | JQ691418       | Kudu    | 2008              | Gobabis           | Omaheke        | Namibia | This study |
| 24K08         | JQ691420       | Kudu    | 2008              | Outjo             | Kunene         | Namibia | This study |
| 27K08         | JQ691421       | Kudu    | 2008              | Outjo             | Kunene         | Namibia | This study |
| 28K08         | JQ691422       | Kudu    | 2008              | Outjo             | Kunene         | Namibia | This study |
| 33K08         | JQ691423       | Kudu    | 2008              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 35K08         | JQ691424       | Kudu    | 2008              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 46K08         | JQ691426       | Kudu    | 2008              | Outjo             | Kunene         | Namibia | This study |
| 51K08         | JQ691427       | Kudu    | 2008              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 52K08         | JQ691428       | Kudu    | 2008              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 55K08         | JQ691429       | Kudu    | 2008              | Gobabis           | Omaheke        | Namibia | This study |
| 57K08         | JQ691430       | Kudu    | 2008              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 59K08         | JQ691431       | Kudu    | 2008              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 60K08         | JQ691432       | Kudu    | 2008              | Outjo             | Kunene         | Namibia | This study |
| 95K08         | JQ691437       | Kudu    | 2008              | Omaruru           | Erongo         | Namibia | This study |
| 130K09        | JQ691438       | Kudu    | 2009              | Outjo             | Kunene         | Namibia | This study |
| 131K09        | JQ691439       | Kudu    | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 142K09        | JQ691440       | Kudu    | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 143K09        | JQ691441       | Kudu    | 2009              | Windhoek          | Khomas         | Namibia | This study |
| 144K09        | JQ691442       | Kudu    | 2009              | Gobabis           | Omaheke        | Namibia | This study |
| 146K09        | JQ691443       | Kudu    | 2009              | Omaruru           | Erongo         | Namibia | This study |
| 147K09        | JQ691444       | Kudu    | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 152K09        | JQ691446       | Kudu    | 2009              | Okahandja         | Otjozondjupa   | Namibia | This study |
| 153K09        | JQ691447       | Kudu    | 2009              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 158K09        | JQ691448       | Kudu    | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 172K09        | JQ691449       | Kudu    | 2009              | Etosha National Park | Kunene | Namibia | This study |
| 190K09        | JQ691452       | Kudu    | 2009              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 191K09        | JQ691453       | Kudu    | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 201K09        | JQ691457       | Kudu    | 2009              | Omaruru           | Erongo         | Namibia | This study |
| 212K09        | JQ691459       | Kudu    | 2009              | Karibib           | Erongo         | Namibia | This study |
| 234K09        | JQ691460       | Kudu    | 2009              | Ondangwa          | Ohangwena      | Namibia | This study |
| 240K09        | JQ691462       | Kudu    | 2009              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 244K09        | JQ691463       | Kudu    | 2009              | Otavi             | Otjozondjupa   | Namibia | This study |
| 20J08         | JQ691419       | Jackal  | 2008              | Okahao            | Omausati       | Namibia | This study |
| 38J08         | JQ691425       | Jackal  | 2008              | Grootfontein      | Otjozondjupa   | Namibia | This study |
| 64J08         | JQ691433       | Jackal  | 2008              | Keetmanshoop      | Karas          | Namibia | This study |
| 67J08         | JQ691434       | Jackal  | 2008              | Etosha National Park | Kunene | Namibia | This study |
| 89J08         | JQ691435       | Jackal  | 2008              | Etosha National Park | Kunene | Namibia | This study |
| 93J08         | JQ691436       | Jackal  | 2008              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 151J09        | JQ691445       | Jackal  | 2009              | Walvis Bay        | Erongo         | Namibia | This study |
| 178J09        | JQ691450       | Jackal  | 2009              | Etosha National Park | Kunene | Namibia | This study |
| 179J09        | JQ691451       | Jackal  | 2009              | Etosha National Park | Kunene | Namibia | This study |
| 192J09        | JQ691454       | Jackal  | 2009              | Etosha National Park | Kunene | Namibia | This study |
| 193J09        | JQ691455       | Jackal  | 2009              | Etosha National Park | Kunene | Namibia | This study |
| 197J09        | JQ691456       | Jackal  | 2009              | Outjo             | Kunene         | Namibia | This study |
| 204J09        | JQ691458       | Jackal  | 2009              | Otjiwarongo       | Otjozondjupa   | Namibia | This study |
| 236J09        | JQ691461       | Jackal  | 2009              | Ondangwa          | Ohangwena      | Namibia | This study |
adaptation may have occurred in the introduction of RABVs from bats to skunks and foxes in the Flagstaff region of the USA [9]. As rabies in Namibia frequently affects canids, e.g. dogs and jackals [10], it remained unclear whether rabies cases in kudu are simply a result of a high rate of spillover from these primary hosts, or a truly independent cycle primarily driven by horizontal transmission among these herbivores. Experimental observations of RABV in kudu have suggested that transmission could occur via non-bite means and that the mucosal surfaces of kudu are particularly susceptible to infection [11]. An earlier epidemiological study did consider molecular phylogeny, but included limited sample numbers that did not allow for definitive conclusions about the independence of the kudu rabies cycle [12]. Thus, although the maintenance of an independent RABV cycle in Namibian kudu has been speculated, we intended to provide additional evidence - based on genome sequence properties - that horizontal transmission of rabies among Namibian kudu is indeed a common occurrence.

| Sample Number | GenBank Number | Species | Year of Isolation | Origin - District | Origin - Region | Country | Reference |
|---------------|---------------|---------|------------------|------------------|----------------|---------|-----------|
| RV1825        | DQ489835      | Bat-eared fox | 1990          | Gordonia         | Northern Cape | South Africa | [17]       |
| RV1826        | DQ489885      | Bat-eared fox | NK             | Etosha           | Kunene         | Namibia   | [17]       |
| RV1827        | DQ489886      | Bat-eared fox | NK             | Etosha           | Kunene         | Namibia   | [12]       |
| RV1828        | DQ489836      | Bat-eared fox | NK             | Etosha           | Kunene         | Namibia   | [17]       |
| RV1829        | DQ489887      | Jackal     | NK             | Etosha           | Kunene         | Namibia   | [17]       |
| RV1830        | DQ489888      | Dog        | NK             | NK               | NK             | Namibia   | [12]       |
| RV1831        | DQ489837      | Bat-eared fox | 1990          | Postmasburg      | Northern Cape | South Africa | [17]       |
| RV1857        | DQ489853      | Bat-eared fox | 1994          | Namaqualand      | Northern Cape | South Africa | [17]       |
| RAVMMGN       | M13215        | Pasteur virus |                |                  |                |          | [21]       |

NK - unknown information.

PLOS ONE | www.plosone.org 3 March 2013 | Volume 8 | Issue 3 | e58739

**Methods**

**Partial N gene sequencing**

Brain samples used in this study were received from the Central Veterinary Laboratory (CVL) in Windhoek, Namibia. The brain samples were obtained from dead, or suspected rabid animals. These samples were from kudu and jackal - from years 2008 and 2009 - that tested positive for rabies after being submitted for diagnostic testing (Table 1) via a network of farmers, interest groups and wildlife conservancies that was established for this study in Namibia. All samples obtained through this network were used in the study. These samples were from various regions throughout Namibia (Figure 1), although the majority originated from central Namibia where the game farming industry is predominant. In total, 49 fluorescent antibody test (FAT) positive samples were sequenced. RNA extraction was performed using the identification of unique amino acid (aa) changes in viruses recovered from kudu, as opposed to those from canid host species.
Figure 1. Kudu and jackal sample origins. Numbers and locations of all rabies virus samples used in the partial sequencing analysis of this study from kudu (green) and jackal (red). The size of the dot increases with the number of samples from each location.
doi:10.1371/journal.pone.0058739.g001

Table 2. Full genome RABV sequences from kudu and jackals constructed in this study.

| Sample Number | GenBank accession no. | Species\(^\text{a}\) | Year of Isolation | Origin – District | Origin - Region | Country |
|---------------|-----------------------|----------------------|-------------------|------------------|-----------------|---------|
| 239K09        | JX473840              | Kudu                 | 2009              | Windhoek         | Khomas          | Namibia |
| 240K09        | JX473841              | Kudu                 | 2009              | Grootfontein     | Otjozondjupa    | Namibia |
| 178J09        | JX473838              | Jackal               | 2009              | Etosha National Park | Kunene         | Namibia |
| 192J09        | JX473839              | Jackal               | 2009              | Etosha National Park | Kunene         | Namibia |

\(^{a}\)Kudu – *Tragelaphus strepsiceros*; Jackal – *Canis mesomelas.*
doi:10.1371/journal.pone.0058739.t002
| Virus isolate | Country and area of isolation | Host species* | Year of isolation | Laboratory reference number | GenBank accession numbers |
|---------------|------------------------------|---------------|------------------|----------------------------|--------------------------|
| RABV (canid variant) Sibasa, South Africa | Dog | 2006 | 262/06 | HM179504 (N), HQ266628 (P), HQ266609 (M), HQ266620 (G). |
| RABV (canid variant) emKhondo, formerly Piet Retief, South Africa | Dog | 2004 | 567/04 | HM179505 (N), HQ266626 (P), HQ266607 (M), HQ266618 (G). |
| RABV (canid variant) Thabazimbi, South Africa | Dog | 1996 | 479/96 | HM179506 (N), HQ266625 (P), HQ266610 (M), HQ266621 (G). |
| RABV (canid variant) Soutpansberg, South Africa | Black-backed jackal | 2005 | 819/05 | HM179507 (N), HQ266629 (P), HQ266611 (M), HQ266622 (G). |
| RABV (canid variant) Umtata, South Africa | Bat-eared fox | 2005 | 31/05 | HM179508 (N), HQ266627 (P), HQ266608 (M), HQ266619 (G). |
| RABV (canid variant) Japan | Laboratory strain | 1915 | Nishigihara | AB044824 |
| RABV (canid variant) USA | Silver-haired bat | 1983 | SBBRV-18 | AY705373 |
| RABV (canid variant) Japan | Nishigahara derivative | 1918 | RC-HL | AB009663 |
| RABV (canid variant) China | Human | Flury-LEP | FJ577895 |
| RABV (canid variant) USA | LEP-Fury derivative | 1939 | HEP-Flury | AB085828 |
| RABV (canid variant) Vaccine | PV | M13215 |
| RABV (mongoose variant) Rusape, Zimbabwe | Slender mongoose | 1994 | 22107 | FJ392391 (N), HQ266633 (P), HQ266615 (M), FA65408 (G). |
| RABV (mongoose variant) Grootgewaagd, South Africa | Yellow mongoose | 1990 | 669/90 | FJ392385 (N), HQ266616 (M), FA65402 (G). |
| RABV (mongoose variant) Kroonstad, South Africa | Yellow mongoose | 1995 | 767/95 | FJ392388 (N), HQ266630 (P), HQ266617 (M), FA65405 (G). |
| RABV (mongoose variant) Uitenhage, South Africa | Yellow mongoose | 1996 | 364/96 | FJ392379 (N), HQ266632 (P), HQ266614 (M), FA65397 (G). |
| RABV (mongoose variant) Beaufort West, South Africa | Water mongoose | 1991 | 113/91 | FJ392372 (N), HQ266631 (P), HQ266613 (M), FA65390 (G). |
| LBV Durban, South Africa | Wahlbergs Epauletted Fruit bat | 2008 | LBVSA2008 | HM179509 (N), HQ266634 (P), HQ266612 (M), HQ266623 (G). |
| LBV Amanzimtoti, South Africa | Wahlbergs Epauletted Fruit bat | 2006 | LBVSA2006 | EFS47452 (N), EFS47414 (P), EFS47435 (M), EFS47422 (G). |
| LBV Exported to France from an unknown African origin | Egyptian fruit bat | 1999 | LBVAFR1999 | EFS47447 (N), EFS47418 (P), EFS47445 (M), EFS47432 (G). |
| LBV Lagos Island, Nigeria | Straw-coloured fruit bat | 1956 | LBVNIG1956 | EFS47459 (N), EFS47407 (P), EFS47444 (M), EFS47431 (G). |
| LBV Durban, South Africa | Wahlbergs Epauletted Fruit bat | 2004 | LagSA2004 | EFS47458 (N), EFS47415 (P), EFS47440 (M), EFS47428 (G). |
| LBV Westville, South Africa | Slender mongoose | 2004 | Mongoose2004 | EFS47453 (N), EFS47409 (P), EFS47438 (M), EFS47423 (G). |
| MOKV Bulawayo, Zimbabwe | Cat | 1981 | 12341 | FA65417 (N), GQ861350 (P), GQ472991 (M), GQ473003 (G). |
| MOKV East London, South Africa | Cat | 1995 | 543/95 | FA65415 (N), GQ500116 (P), GQ472992 (M), GQ500110 (G). |
| MOKV Pinetown, South Africa | Cat | 1997 | 252/97 | Unpublished (N) AF369376 (P), GQ472997 (M), GQ500112 (G). |
Trizol (Invitrogen) according to the manufacturer’s instructions. The RT-PCR reaction targeted a region (bases 16-646 according to the Pasteur virus rabies genome, GenBank accession number: M13215) of the nucleoprotein gene approximately 602 bp in size [13]. PCR amplicons were first confirmed by agarose gel electrophoresis, followed by PCR purification using the Wizard® SV Gel and PCR Clean-Up System (Promega), according to the manufacturer’s instructions.

Purified PCR products were sequenced using the BigDye Terminator v1.1 Kit cycle sequencing protocol (Applied Biosystems) at the University of Pretoria on an ABI3130 DNA sequencer. Sequences were edited using CLC Main Workbench v6.0 (CLCBio). For phylogenetic analysis, the newly obtained sequences - as well as various relevant sequences from previous studies (Table 1) - were aligned using the Clustal X function in BioEdit [14]. A Neighbour-joining phylogenetic tree was constructed with the use of the Kimura 2-parameter model with 1000 bootstrap replications in MEGA 5.0 [15].

Full genome sequencing

RNA extraction... RNA was extracted from two kudu and two jackal brain samples (Table 2) using a protocol combining Trizol (Invitrogen) and RNAeasy mini kit (Qiagen) according to the manufacturer’s instructions.

Reverse-Transcription Polymerase Chain Reaction (RT-PCR). A One-Step Reverse Transcription PCR was performed using the SuperScript® III One-Step RT-PCR kit (Invitrogen) for products less than 1500 base pairs in length, according to the manufacturer’s instructions with several primer pairs (Table S1). The following cycle conditions were used on a model 2720 thermocycler (Applied Biosystems): 50°C for 30 minutes; 93°C for 2 minutes and 42 cycles of 95°C for 30 seconds, 55°C for 30 seconds, 68°C for 60 seconds; and a final extension step for 5 minutes at 68°C.

A long range One-Step RT-PCR was performed using the SuperScript® One-Step RT-PCR for Long Templates (Invitrogen), according to the manufacturer’s instructions with the following conditions: 50°C for 30 minutes, 95°C for 2 minutes followed by 42 cycles of 95°C for 30 seconds, 55°C for 30 seconds, 68°C for 1 minutes/kb. A final extension step was performed for 5 minutes at 68°C.

PCR products of sizes less than 1500 bp were analysed by 1.5% agarose gel electrophoresis using 1 x TAE buffer (1.6 M Tris-acetate, 40 mM EDTA). For products larger than 1500 bp, a 1% agarose gel was prepared. A 100 bp DNA molecular weight marker (Promega) was included to identify the size of the amplicons.

Purification and sequencing. PCR amplicons generated were purified using the QIAquick Gel Extraction Kit (Qiagen), according to the manufacturer’s instructions. PCR products were sequenced using the BigDye Terminator v1.1 Kit cycle sequencing protocol (Applied Biosystems) with slight modifications. One microlitre 5 x sequencing buffer, 1 μl 5 pmol/μl primer (Table S1), 5 μl [15 ng/μl] template, 1 μl nuclease free molecular grade water (Qiagen) and 2 μl BigDye Terminator mix v1.1 was added to a final volume of 10 μl. The reaction was performed using the following profile: an initial denaturation step at 96°C for 1 min; then 26 cycles of 96°C for 15 seconds; 53°C for 10 seconds; and 60°C for 4 minutes.

Following the sequencing reaction, sequence products were purified using Sigma Spin Sequencing Reaction Clean-up Post Reaction Purification Columns (Sigma) according to the manufacturer’s instructions. Fifteen microlitres of the eluate were added to 15 μl of Hi-Di-Formamide and subsequently analysed using an ABI 3130 DNA sequencer (Applied Biosystems).

Genetic analyses. Sequences were assembled, translated and annotated using CLC Main Workbench v6 (CLCBio) and the ClustalX function of BioEdit [14]. Multiple alignments including sequences from the public domain (Table 3) were then manually screened for nucleotide sequence differences and those that also resulted in aa changes were identified. To identify any known viruses with the same aa in the variable positions identified, we used a blast analysis, screening the protein data bases in Genbank with 7 aa sequence segments that were composed of the variable aa and the 3 aa immediately to the left and right of this position. Finally, phylogenetic analysis was performed with the inclusion of the full genome sequences presented in Table S2 and according to the methodology described for partial N gene sequences, above.

Rapid Amplification of cDNA Ends (RACE). Rapid Amplification of cDNA Ends (RACE) determined the 5'- and 3'-termini of viral RNA from samples 239K09, 240K09, 178J09 and 192J09. RACE was performed with freshly extracted RNA using the 5’- and the 3’-RACE system from Invitrogen (Invitrogen,
Carlsbad, USA) according to the manufacturer’s recommendations. Due to a lack of a poly-A tail at the 3’-terminus of the viral RNA, a poly-C tailing was done prior to 3’-RACE. Some modifications were introduced to the 3’-RACE procedure because of the poly-C tailing: for first strand cDNA synthesis, amplification of the cDNA and subsequent hemi-nested PCR of the abridged anchor primer (AAP) from the 5’-RACE kit had to be used instead of the recommended adapter primer (AP, designed for annealing to a poly-A tail). As gene specific primers (GSPs) for the 3’-RACE system RAB-JACKAL-150R (amplification of cDNA) and RAB-JACKAL-100R (hemi-nested PCR) and for the 5’-RACE procedure RAB-JACKAL-12235F (cDNA synthesis) and RAB-JACKAL-12265F (amplification of cDNA) were used for all four samples (Table S1). Amplification products were visualized on a 2% agarose gel, purified and subsequently sequenced (see Purification and sequencing).

Results

Phylogenetic analyses

A few previous studies have included partial nucleoprotein gene sequencing of Namibian and other southern African RABV isolates and a sizeable collection of sequences can be found in Genbank [12,16,17]. We therefore chose to continue with this sequence region in our analyses of the 49 new isolates from kudu and jackal (35 kudu and 14 jackal) (Table 1) from regions throughout Namibia (Figure 1). Subsequent phylogenetic analysis depicted that 42 from 43 kudu RABV isolates belonged to a single clade that were distinct from any other RABV isolates from Namibia, or elsewhere in southern Africa (Figure 2). The single exception was the kudu specimen 190K09 from Grootfontein, which bundled with a group consisting of 7 jackal isolates from Etosha National Park. The majority of RABV from jackals sequenced were from Etosha National Park (7/14) and these all grouped together; separate from a rabies cycle in bat-eared foxes – also from the Etosha National Park (Figure 2). Samples 151J09 and 204J09 grouped with jackal and dog RABV sequences from Botswana and central Namibia. Sample 64J08, originating from Keetmanshoop in southern Namibia, grouped closely with other RABV sequences from a South African bat-eared fox rabies cycle from areas bordering Namibia to the south. One eland (Tragelaphus oryx) sample from a previous study (RV1518) also grouped with the RABV sequences from kudu.

We also wanted to compare and evaluate phylogeny based on full genome sequences, where the resultant tree (Figure 3) supported the results obtained from the partial N gene sequences shown above. The 4 isolates that were sequenced in full grouped separately from any other known sequences, and the 2 isolates from jackal (178J09 and 192J09) grouped separately from the 2 isolates from kudu (239K09 and 240K09) with 100% bootstrap support. The next closest strains were both vaccine strains from Russia and Japan respectively. The separate grouping of the viruses sequenced in this study was expected as no other full genome sequences of African RABV isolates were available, and the distinction between kudu and jackal isolates was consistent
Amino acid sequence analysis of full genomes

The full genomes of the 2 RABV isolates from jackal were found to be identical with respect to aa sequence for all five genes, while the 2 isolates from kudu differed from one another in only one aa position, in the L gene. However, there were 13 aa differences between the kudu and the jackal RABV isolates. Three aa variations in the G gene were unique to the kudu RABV isolates and a further variation unique to both kudu and jackal RABV (L gene), even after comparing these aa to all other lyssavirus G and L sequences known (non-redundant protein and Swiss-Prot databases, Table 4).

Discussion

RABV is typically maintained within carnivorous hosts and maintenance within an herbivorous host population has never been proven. The apparently independent cycling of a RABV variant within Namibian kudu, known to be a non-aggressive herbivore, is therefore truly unique. In this study, we aimed to determine whether the RABVs isolated from Namibian kudu showed characteristics of host adaptation and whether they represented a distinct cycle of the disease. In general, phylogenetic analyses have shown clustering of RABVs to occur according to the geographical origin of the viruses, despite the fact that the viruses were isolated from several different species [1]. However, some exceptions have been noted, more specifically and predominantly, in bat species in the Americas [18]. Using molecular analyses, we also intended to determine whether unique mutations had occurred in the cross-species transmission from carnivores to kudu in order to: 1) further support/reject the notion of an independent cycle and; 2) determine whether the particular susceptibility of kudu to RABV infection is due to the adaptation of the virus to a new host. Historical epidemiological data is clear on the directionality of original transmission pathways for rabies in Namibia. Endemic cycles in jackal were firmly established at the turn of the 20th century, with the first recorded cases in kudu appearing several decades later [5,7]. With the advent of molecular genetics, the close relationship between the viruses in jackal and kudu was shown – and the establishment of an independent kudu cycle is demonstrated in the present study.

Phylogenetic analyses of partial N gene sequences from 35 kudu and 14 jackal RABVs showed a significant clustering of all of the sequences from kudu RABVs separately from any other sequences from canid RABVs, with high bootstrap confidence. As the samples were all taken within the same temporal and similar spatial range, the divergence seen between RABV from canids and kudu can only be explained by the existence of a RABV cycle that is being maintained within the kudu population, separately to other RABV cycles in canid populations. An exception to the general clustering of RABV from kudu is sample 190K09, which grouped with RABV from jackals from Etosha National Park. This can be explained by a spill-over infection from a jackal to that kudu, as is commonly seen in cases of bovine infections from rabid jackals [10,19]. It is a foregone conclusion that rabies was first introduced into the kudu population in this manner and that sporadic subsequent spillovers could be expected, but not on a scale that would account for the numbers of kudu rabies cases recorded since 1977. In addition, if a separate rabies cycle was not being maintained within the kudu population, several RABVs from canids would be interspersed within the kudu group with the assumption that one jackal could infect several kudu. It was also noted that several samples from jackals from the central regions of Namibia (where the majority of samples from kudu were taken), grouped in separate RABV cycles with other RABVs from jackals and dogs from the central region (Figure 2). This lends further support to the observation that RABV is being maintained in a separate cycle in kudu.
Table 4. Amino acid variations unique to kudu and jackal RABV genomes when compared to all known lyssavirus sequences.

| Gene             | Host species | Position | Variable amino acid | Amino acid in other RABV (Table 3) | Charge change |
|------------------|--------------|----------|---------------------|-----------------------------------|--------------|
| Glycoprotein     | Kudu         | 34       | Asparagine          | Serine                            | none         |
| Glycoprotein     | Kudu         | 112      | Threonine           | Alanine                           | non-polar to polar |
| Glycoprotein     | Kudu         | 191      | Serine              | Asparagine                        | none         |
| Glycoprotein     | Kudu and Jackal | 1140    | Serine              | Glycine                           | non-polar to polar |
| Polymerase       | Kudu and Jackal |        |                     |                                   |              |

Sequence analyses revealed several aa variations unique to Namibian kudu RABV’s. The variations specifically seen in the RABV isolates from kudu may suggest that these changes arose due to the adaptation of the virus to the host, but this notion will remain speculative until proven. Alternatively, this may suggest the divergence of the virus due to a separate RABV cycle being maintained within the kudu population (geographical and temporal drift), as kudu in Namibia may be geographically isolated due to separation of these animals by barriers such as game fences. Variations observed in viruses from both kudu and jackal may be due to the geographical isolation (on a larger scale) of the viruses from other rabies viruses and RABV cycles in Namibia and southern Africa, and support the evolutionary link between kudu and jackal rabies. In contrast, a recent study showed that RABVs did not undergo significant mutational differences after the host shift from bats to skunks, in both the coding and non-coding regions of the genomes [9]. However, the epizootic and maintenance of rabies in skunks occurred in a brief time frame in comparison with the emergence of rabies in kudu, and thus the time for divergence is greater in the kudu RABV cycle.

Complete genome sequencing is an important and useful method utilized in this study. The need for more complete genome sequences is becoming more important, especially with the molecular data that can only be generated through this technique. This study has shown the need for more complete genome sequencing as the sequences from this study were the first complete genome sequences for RABV from sub-Saharan Africa. These sequences will be important for a future databank of complete genomes that can be used for improved phylogenetic resolution as well as other molecular studies. Further work will need to be performed in order to determine virus evolution, the effects of certain mutations on protein folding, as well as continued insight into host-virus interactions and pathogenicity [20].

In conclusion, phylogenetic and full genome sequence analyses showed several correlations between Namibian ‘street viruses’ to attenuated vaccine strains, which can be explained by the introduction of the cosmopolitan RABV strain from Europe into Africa during the mid-20th century. Most importantly, several unique mutations were observed in the RABV isolates from kudu, suggesting that these mutations may have occurred due to the adaptation of the virus to the host. Evidence from this study strongly argues for the maintenance of an independent RABV cycle in kudu, separate from the jackal cycles in Namibia. Future studies will be needed to determine the potential significance of the genomic regions identified as unique to kudu RABVs and to determine whether host specific regions occur in RABV and other lyssaviruses. The important implications of the phenomenon of the independent maintenance of RABV among kudu are that the control and eradication of a rabies cycle in this, and other species, will pose unique and uncharted challenges. From a continental perspective it is important to note that endemic canine rabies remains of primary concern and Namibia and other countries of the southern African region are no exception. Eventual effective control of rabies in kudu and jackal is likely to be subject to an effective control program for rabies in dogs.

Supporting Information

Table S1 Primers used for full RABV genome sequencing.

(IMPORTANT) Table S2 RABV full genomes used in phylogenetic and mutational analysis.

Author Contributions

Obtained samples and performed FAT tests: SK. Conceived and designed the experiments: WM BH TM LHN. Performed the experiments: TPS MF. Analyzed the data: TPS. Contributed reagents/materials/analysis tools: WM LHN BH TM DH CF. Wrote the paper: TPS LHN.

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