**Longitudinal Survey of Astrovirus infection in different bat species in Zimbabwe**: **Evidence of high genetic** Astrovirus **diversity**

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**Abstract**

Astroviruses (AstVs) have been discovered in over 80 animal species including diverse bat species and avian species. A study on Astrovirus circulation and diversity in different insectivorous and frugivorous chiropteran species roosting in trees, caves and building basements was carried out at 11 different sites across Zimbabwe. Pooled and individual faecal sampling methods were used for this study, with collection dates ranging from June 2016 to July 2021. In two sites, Magweto and Chirundu, sampling was carried out at monthly intervals from August 2020 to July 2021. Astroviruses and bat mitochondrial genes were amplified using pan-AstVs and CytB /12S RNA PCR systems respectively. Phylogenetic analysis of the *RdRp* Astrovirus sequences revealed a high genetic diversity of astroviruses. All the bat astroviruses tested in this study clustered with the *Mamastrovirus* genus. Two distinct groups of amplified sequences were identified. One group was comprised of sequences isolated from *Hipposideros, Rhinolophus* and *Eidolon helvum* spp*.* clustered with Human Astrovirus strains: *HuAstV* types1-6, *HuAstV*-MLB1-3 and *HuAstV*-VA-1. The second group comprising of the majority of the sequences clustered with different strains of Bat AstVs. Results from the longitudinal study at Magweto and Chirundu showed an overall AstV prevalence of 13.7% and 10.4% respectively. Peaks of AstV infection at Chirundu coincided with the period when juveniles are 4-6 months old. By combining data from our previous work on Coronaviruses, we also anaylzed co-infection of bats with Coronaviruses and Astroviruses at Magweto and Chirundu sites for which the prevalence of co-infection was 2.6% and 3.5% respectively.

**Introduction**

Astrovirus (AstVs) are non-enveloped single-stranded positive sense RNA viruses with a genome length of approximately 6.2 to 7.7 Kb [1]. They have been described in over 80 non-human species and are classified into two genera: *Mamastrovirus* and *Avastrovirus,* respectively infecting mammals and avian species [2]⁠. According to the International Committee on Taxonomy of Viruses (https://talk.ictvonline.org), *Mamastrovirus* are classified into 19 recognized species, *MAstV*-1 to -19, and two genogroups GI and GII [3]. All classic and novel Human Astrovirus (*HuAstVs*) belong to four different species, *MAstV*-1, *MAstV*-6, *MAstV*-8 and *MastV*-9 [2,4,5]. Recently, studies have reported AstVs infections in fish and insects [6,7], however bats and wild birds are considered to be natural reservoirs of the *Astroviridae* family [8,9]. Generally *HuAstVs* have been identified as causal agents of acute viral gastrointestinal illness worldwide particularly in children, immunocompromised people and the elderly [6–8]⁠. Of note, in humans, the majority of AstV-associated encephalitis or meningitis illnesses were reported in immunocompromised people [10–12]. Furthermore, beyond this well-known clinical manifestation of AstVs infections, neurovirulent AstVs infections have also been reported in both humans and domestic animals [13,14]. Moreover studies have reported numerous cross-species transmissions of AstVs [7], where some AstVs detected in humans closely clustered with strains identified in rodents, minks, feline species and ovine [2,5,7]. Bat-borne viruses represent an extensive research field owing to the plethora of viruses hosted by Chiropterans [15]. Furthermore, this mammalian order is known to be persistently infected by astroviruses [16] and this is inclusive of many species of insectivorous bats [17]. To date, a high diversity of bat AstVs from different bat families has been reported worldwide [11,18-19]. Bat AstVs belong to *MAstV*-12, and *MastV*-14 to -19 species [1]. In Europe a variety of bat species belonging to the Yangochiroptera sub-order have been discovered to harbor several astroviruses [20]⁠. In Africa, bat astroviruses were reported in Egypt, Gabon, Madagascar, Kenya and Mozambique [18, 21–23]⁠.

AstVs are known to occur in bat species with high prevalence and exceedingly high genetic diversity whereby infections in these hosts are usually non-pathogenic [24,25]. Despite all these studies, large gaps exist on astroviruses prevalence and infection dynamics in bats [9]. Astrovirus prevalence and detection in bat colonies seemed rather correlated to abiotic factors such as seasons and year of sampling and biotic factors such as sex and reproductive status and also viral co-infection [26]⁠⁠⁠. Viral shedding in bats occurs in spatial and temporal pulses that can drive spillover to other animals or humans [27-28].⁠ The varying prevalence of viruses in bats is determined by shedding pulses, where rare or very low prevalences are detected in the absence of shedding pulses and higher prevalence is detected when shedding pulses occur [27,29].

Until now, no data was available on genetic diversity and prevalence of circulating AstVs in insectivorous bats in Zimbabwe. In this study we enlarge the spectrum of bat AstVs knowledge in Africa by identifying and analyzing bat-AstVs, their diversity and prevalence from different bat species colonies according to reproductive season phenology in Zimbabwe. Furthermore, we will also show and highlight the presence of co-infection of bats by astroviruses and coronaviruses and how it varies according to reproductive phenology.

**Material and Methods**

**Sampling approaches and sites**

Two different approaches were followed in this study: bat community sampling and individual bat sampling.

***Bat community pooled sampling***

Between February 2016 and December 2020, faecal samples from both insectivorous and frugivorous bat species were respectively collected in different sites including caves, an ancient mine and trees in Zimbabwe (Figure 1, Table 1).



**Figure 1:** The study sites highlighted on the map show the sampling areas for insectivorous and frugivorous bats colonies across Zimbabwe. In Honde valley, the *Eidolon helvum* colony was established in at least four different sites represented by the rectangle at the right of the figure. Red circles represent community approach study sites; yellow circles represent longitudinal survey sites (individual approach).

**Table 1:** Number of Astrovirus positives per site/sampling year in pooled faecal samples from frugivorous and insectivorous bats for the selected 14 sites.



All these sites were chosen according to the presence of bat colonies and the existence of anthropic activities. For instance, the selected caves and ancient mine are regularly visited by local people to collect bat guano, which is used as fertilizer, and/or to hunt bats for consumption. In Honde valley, frugivorous bats roosting sites in trees were close to maize crops or/and orchards.

All sites, except for three, were visited at different times during the sampling period (Table 1). The same sampling method was used at all sites and in every session as previously described by Bourgarel et al, [30]. Briefly, two square meters of plastic sheets were laid down at each site/per sampling session, underneath the bat colonies and left overnight. Six grams of fresh faeces from different bat individuals were collected from each plastic sheet in a 15 ml tube with 6 ml of RNA stabilization solution (*https://protocol-online.org/*). Back in the laboratory, samples were stored at -80°C for subsequent analysis. Of note, for all sampling sessions, personnel protection equipment (gloves, FFP3 mask, Tyvek coverall, etc) was used in order to respect the biosafety procedures and to protect the sampling teams. Good biosafety practices training are regularly dispensed for our sampling teams.

***Bat individual sampling***

Individual bat samples which had already been collected from two study sites for a study on coronaviruses by Chidoti et *al*. were used in this study (Figure 1) [31]. These two sites, one cave (Magweto) and one building basement (Chirundu Farm) had been visited from August 2021 to July 2022 on a monthly basis (Figure 1, Table 2) [31]⁠. Unfortunately, this study was conducted during the COVID 19 crisis and we were not able to access the study sites every month as planned owing to the imposed lock downs. Faecal samples had been collected by placing two square meters plastic sheets underneath the bat colonies. Only one faecal dropping (not contaminated by other faeces or urine) per 20 cm2 was collected, assuming it represented one individual. Faeces were conserved individually in a 1.5 ml tube filled up with 0.5 ml of home-made RNA stabilization solution ([*https://protocol-online.org/*](https://protocol-online.org/)) and stored at -80°C before further laboratory analyses.

**Table 2:** Prevalence of Astroviruses and co-infection with Coronaviruses and confidence intervals (CIs) per month at both Magweto and Chirundu sites in faecal samples from individual sampling of insectivorous bat communities per reproduction cycle stage.



Periods of gestation (pregnancy), parturition, lactation, weaning and presence of 4-6 months old juveniles had been determined based on observations (from captures and observation of the roosting bats) combined with literature, as already reported in Chidoti et *al.* [31]. The reproductive season at Chirundu and Magweto site had been observed to begin in September to February for the predominant bat species. The different insectivorous bat families observed at both sites had been found to be synchronous regarding their reproductive cycles, and consensus reproduction periods had been determined based on the literature and observations [31,32]⁠⁠.

**Nucleic acid extraction and RT-PCR**

***Community samples***.

Nucleic acids extraction was done from all faecal samples as previously described [30]⁠. Briefly, biological material (Faeces) of three or four sample tubes collected from the same plastic sheet were pooled and transferred in a 50 ml tube with 20 ml of PBS 1X then vigorously mixed. Tubes were centrifuged at 4500 rpm for 10 min. The supernatant was first filtered using gauze in order to eliminate faecal matter and transferred in fresh tubes then re-centrifuged at 4500 rpm for 10 min. The supernatant was filtered through a 0.45 µm filter to remove eukaryotic and bacterial sized particles. Seven millilitres of filtered samples were centrifuged at 250,000 g for 2.5 h at 4°C. The pellets were re-suspended in 600 µl H20 molecular grade and 150 µl were used to extract RNA and DNA using NucleoSpin® RNA Kit (Macherey-Nagel, Hoerdt, France) according to the manufacturer’s protocol.

***Individual samples***

Nucleic acids were extracted from 200 μl of faecal samples preserved in 0.5 ml RNA stabilization solution using 5X MagMax Pathogen RNA/DNA Kit (ThermoFisher Scientific, Illkirch-Graffenstaden, France), as already described in [30]. The faeces were vortexed vigorously(30Hz) using Retsch MM400 Tissuelyser for5 min to fully homogenise and mix the faecal particles, followed by centrifugation at 16000 g for 3 min to fully separate the supernatant from the faecal debris. A volume of 130μl of the supernatant was used for the isolation and purification stage of the nucleic acids using Mag Max extraction kit with the automatic KingFisher Duo Prime Purification System extractor (ThermoFisher Scientific, Illkirch-Graffenstaden, France). A final volume of 80μl of eluted RNA/DNA was stored at -80◦C.

***Bat species identification***

For all bat samples that tested positive forAstVs, bat species identification was carried out by the amplification and sequencing of mitochondrial C*ytochrome b* gene as previously described by Kocher et *al.* [33]⁠. *S*equences were then compared to available bat sequences in the GenBank database using *Basic Local Alignment Search Tool* (BLAST).

***Astrovirus detection***

Reverse Transcription (RT) using random hexamers were done on 5μl of RNA sample template using 1μl random hexamers, 0.5μl Oligo dT primer, 0.4μl of dNTPs (10mM) (ThermoFisher Scientific, Illkirch-Graffenstaden, France) and 5.5μl molecular grade water incubated at 65°C for 5 min. This was followed by addition of 4μl of Buffer 5X, 2μl of 0.1M DTT (M-MLV Reverse Transcriptase, Invitrogen, ThermoFisher Scientific) and 1μl of RNAse OUT, incubated at 37°C for 2 min. A volume of 1μl of M-MLV (M-MLV Reverse Transcriptase, Invitrogen, ThermoFisher Scientific, Illkirch-Graffenstaden, France) reverse transcriptase was added to the mixture followed by incubation at 25ºC for 10 min, 37ºC for 50 min and 70ºC for 15 min. The cDNA obtained was then used to partially amplify the *Astrovirus RNA-dependent-RNA polymerase* gene (*RdRp*) by using a semi-nested Pan-Astrovirus PCR system developed by Chu et *al* [16]⁠. Visualization of positive PCR products was performed with agarose gel electrophoresis using SYBR green on a 1% gel. Positive PCR products (422 bp) were gel-agarose or directly purified (Geneclean Turbo Kit, MP Biomedicals, Illkirch-Graffenstaden, France) and then sequenced in both 5’ and 3’ directions (LGC, Berlin, Germany) by using Sanger method.

For the community approach, purified PCR products were cloned by using Topo PCR Cloning kit according to the manufacturer’s protocol (ThermoFisher Scientific, Illkirch-Graffenstaden, France). Ten clones per PCR product were sequenced in both 5' and 3' direction using the Sanger method (Eurofins, Germany).

**Phylogenetic analyses**

Bat AstVs overlapping sequences were assembled into contiguous sequences using Geneious software package V. 2021.2.2 (Biomatters Ltd, Auckland, New Zealand). Partial non-concatenated nucleic acid sequences of the new AstVs were aligned using Clustal W [34] implemented in MEGA 7 [35], with minor manual adjustments. Ambiguously aligned and divergent regions were excluded from subsequent analyses. Phylogenies were inferred using Maximum Likelihood (ML) method implemented in PhyML V. 3.0 [36]⁠⁠. The suited evolution model was selected by Akaike’s Information criterion (AIC) using Topali software [37]⁠.⁠ The reliability of branching orders was tested using the bootstrap approach (1000 replicates) [38]⁠ and the GTR + F+ I substitution model was determined as the best suited evolution model.

**Temporal variations of Astrovirus prevalence and bat reproductive phenology**

In the same way as in Chidoti et *al*, the prevalence of astrovirus infection was calculated at the community level for small insectivorous bat species from the two longitudinal study sites, Magweto and Chirundu farm [31]. The proportion of RNA AstV-positive samples were estimated per month and site with 95% confidence intervals (CI) using Wilson score test*(https://epitools.ausvet.com.au/ciproportion)*[39].⁠

The influence of the different phases of the bat reproductive cycle stage (periods of pregnancy, parturition/lactation, weaning, weaned juveniles of 4 to 6 months old) and of *Coronavirus* infection/shedding on the prevalence of astroviruses was tested by running a generalized linear mixed model (GLMM) for each site (Magweto and Chirundu) as described in Chidoti et *al* [31]. Parturition and lactation were analysed as one season because the observations did not allow the separation of the two periods as they were interlinked. The response variable with a binomial distribution was the AstVs PCR result of the samples, and the explanatory variables with fixed effects were the different phases of the reproduction cycle. Given CoronavirusPCR results on the same samples were available through our previous study (see below), Coronavirus status was also integrated into the model as fixed variable to investigate any effect of co-infection on AstVs infection/shedding. The reproductive phases were coded as 1 if it was during the corresponding reproduction phase and 0 if it was not. A session identification code was included as a random effect to control for repeated measures from the same sampling session to account for clustered samples collected.

***Astrovirus and Coronavirus co-infection***

Coronavirus data, produced by Chidoti et *al* [31], on the same sample sets were used in this study to assess the AstVs and *Coronavirus* co-infection in bat communities from Magweto and Chirundu sites and a descriptive analysis of prevalences was carried out (**Table 2**).

**Results**

Astrovirus ***detection in pooled sampling sites***

The sampling was carried out from June 2016 to December 2020 (Table 1) at the selected sites. At Magweto cave site, 24 samples were collected. In Bashongwe collection was carried out in March, July and December 2018, and 251 samples were collected. For Mabura cave site, a total of 58 samples were collected, in 2016 and 2017, with sampling only carried out once in both years. In Mugunza cave site, a once off sampling session was done in 2017 and a total of 50 samples were collected. In the Honde Valley site, 336 samples were collected under nine *Eidolon helvum* roost trees (Table 1). In Magweto, five of the eight pooled samples (62.5%) tested positive for AstV. In Bashongwe cave, all the 72 samples collected tested negative for AstV (Table 1). In Mabura, Six (31.6%) out of 19 pooled samples tested positive for AstVs and all of the 13 samples from Mugunza tested for AstVs were negative (Table 1). In Honde Valley, 173 pooled samples were tested for AstVs and only four (2.3%) were positive (Table 1). In Chirundu Baobab tree only one (4.8%) out of the 21 samples collected site was positive for AstV.

***Prevalence and seasonality of Astroviruses at Chirundu and Magweto site***

In total, we analyzed 1559 samples from Magweto cave site, and 1728 samples from Chirundu farm. (Table 2).The overall prevalence of AstVs at bat community level at Chirundu was 10.4% [95% CI: 9.0-11.8] (179 positives out of the 1728 samples). The highest prevalence of 21.3% [95% CI: 16.5-26.8] was observed in March 2021 and corresponded to the 4-6 months old juvenile period (Table 2). The prevalence of *AstV* increased from December 2020 to March 2021, which corresponded to the lactation period, followed by the weaning of four to six months old juveniles. The lowest prevalence was observed in August (0%), October (1% [95% CI: 0.3-2.9]) and November (1.2% [95% CI: 0.4-3.6]), which correspond to the transition between non-gestation and gestation period, as well as the beginning of the parturition period (Table 1, Figure 2a).

**Figure 2:**

Bat communityAstVs prevalence per month at Magweto site. **Fig 2a & 2b** show estimation of the astrovirus prevalence (with CI 95%) at Chirundu (a) and Magweto (b) sites respectively for the months and year sampled. The graphs were plotted with y-axes as the percentage of AstVs prevalence and x-axis as the sampling collection date and the corresponding reproductive cycle stage, each colour code representing a different season.



a/



**2020**

**2021**

**Non-gestation**

**Pregnancy**

**Parturition**

**Parturition & Lactation**

**No sampling done**

**Weaning**

**4-6\_months Juveniles**

b/

In Magweto site, the overall prevalence of Astroviruses for the insectivorous bat community was 13.7% [95% CI:12.1-15.5] (214 positives out of 1559 samples). There was no clear pattern for the prevalence of AstVs according to reproductive cycle stages in this site (Table 2, Figure 2b). The highest prevalence of 19.4% [95% CI: 14.9-24.9] was observed in June 2021 during the non-gestation period , and very close prevalences observed during the pregnancy (17.9% [95% CI: 13.7-23]) and 4-6 months old juvenile (19% [95% CI: 14.5-24.4]) periods (Table 2). Lower prevalence was observed in September at the end of the non-gestation period (5.2% [95% CI: 3.3-8]).

Results from the GLMM didn’t show any effect of bats seasonal reproduction periods on detection of AstV RNA in samples collected at Magweto site (Table 3a). For Chirundu site, the GLMM showed significantly higher probability of being positive to astrovirus during the period of weaned 4-6 months old juveniles (odds ratio (OR) = 5.088, 95% CI = 1.33- 36.32, p= 0.016), and significantly lower probability of astrovirus positivity during the pregnancy period (odds ratio (OR) = 0.136, 95% CI = 0.03- 0.81, p=0.006) (Table 3b).

**Table 3*:*** Results of the GLMM with the following explanatory variables for both site (Chirundu and Magweto): PCR\_Pan-*Coronavirus*, Pregnancy, Weaning, Weaned juveniles 4 to 6 months. With references OR odds ratio, CI95 95% confidence interval.

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***Prevalence and seasonality of coinfection with Astroviruses and Coronavirus at Chirundu and Magweto site***

The co-infection prevalence in bat communities from Chirundu and Magweto sites was relatively low. The co-infection prevalence was observed to follow a similar trend as that of CoVs and AstVs prevalences, where it showed peaks that correlated with peaks of the individual viral families (Figure 3 a & b. The overall prevalence of co-infection at Chirundu site was 3.6% [95% CI: 3.8-5.8] (Table 2) with the highest co-infection prevalence of 10.6% [95% CI: 6.8-16.1] observed in February 2021 during the weaning period. The lowest prevalences of 0% were observed in August [95% CI: 0.0-0.2] and October [95% CI: 0.0-1.3] during the end of the non-gestation and the pregnancy period, respectively (Table 2, Figure 3a).

The overall prevalence of co-infection at Magweto site was 2.6% [95% CI: 1.8-3.5] (Table 2, Figure 3b). Co-infection in this site was recorded from September 2020 to December 2020 and from March 2021 to June 2021 (Table 2, Figure 3b). The highest prevalence of co-infection was 8.3% [95% CI: 5.4-12.4] in March during the 4-6 months old juvenile period; while the lowest 0.3% [95% CI: 0.05-1.6] was observed in September 2020 during the non-gestation period (Table 2, Figure 3b).

**Figure 3*:*** Prevalence of *Astrovirus*, *Coronavirus* andco-infections involving both viruses per month: co-infections are represented in green, coronaviruses in blue and astroviruses in red for a/ Chirundu site and b/ Magweto site according to reproduction cycle stages.

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a/

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b/

***Astroviruses genetic diversity***

All AstV positive samples (N=423) were sequenced, representing 179 samples from Chirundu, 229 from Magweto, seven from Mabura, seven from Honde valley, and one from Baobab tree. In this study we sequenced all AstVs positive samples from 5 sites: (Chirundu farm N=179; Magweto cave N=229; Mabura cave N=7; Honde Valley N=7 and Baobab tree N=1). Of the 423 samples, 214 (50.6%) were amplified from *Hipposideros* spp., 63 (14.9%) from *Rhinolophus* spp., 53 (12.5%) from *Miniopterus* spp., 10 (2.4%) from *Nycteris* spp., seven (1.7%) from *Eidolon* spp. and for the remaining 76 samples known to originate from bats for which we couldn’t determine the genus/species.

A phylogenetic analysis was conducted with the 155 partial sequences generated during this study. All clustered within the *Mamastrovirus* genus (Figure 4).

Two sequences isolated from Magweto cave, one from *Rhinolophus* spp*.* (MAG-573) and 1 from *Hipposideros* spp. (MAG-292) clustered with a group of Human astroviruses (*HuAstVs*) (Figure 4).

This cluster was well supported (Bootstrap >90%) and the nucleotidic acid identities between Bat Astroviruses MAG-573, MAG-292 and HuAstV-2 were 96 and 93% respectively (Data not shown).

The other specific cluster of bat astrovirus showing a close relation to *HuAstVs* is a sequence isolated from *Hipposideros* spp. (MAG-1236) in Magweto, it clustered with *HuAstV-MLB-1, -2* and *3* (Figure 4). Six sequences isolated from the frugivorous bats *E.helvum* found in Honde Valley and one from *Rhinolophus* spp. collected in a Baobab tree formed a sister clade with another group of Human astroviruses *HuAstV-VA-1;-2;-3;-4* and *HuAstV-HMO-A* (Figure 4). In this study, the majority of the sequences were closely related to astroviruses identified in insectivorous bats (Figure 4). These were mainly amplified from *Hipposideros* spp.(HC-1 to -9) and *Rhinolophus* spp.(Rhi-1 to -4)bat species. One cluster comprised specifically of sequences isolated from *Rhinolophus* spp.. Most of these sequences were isolated from Chirundu and a few of them from Magweto, and clustered together with *AstV* strains from *Nyctalus* spp.and *Verpitillio* spp. bat species isolated in Czech Republic (references Y & Z in the phylogenetic tree) (Figure 4). Another cluster comprising of the majority of sequences from Chirundu and some from Magweto site, were characterized from *Miniopterus* spp. and *Rhinolopus* spp. All together, they formed a phylogenetic cluster with strains of bat astroviruses isolated from *Miniopterus* spp. from Madagascar and China (references N, O, R, U, T), *Rhinolophus* spp. from Korea (references P, Q) as well as *Rousettus* spp. and *Paratrianops* spp. *BAstV* from Madagascar (references V, W) (Figure 4)

Two major clades from Chirundu and one from Magweto, isolated from *Miniopterus, Rhinolophus* and *Hipposideros* spp. of batsalso phylogenetically clustered with a strain from Mozambique isolated from *Nycteris thebaica* (reference M)*.* One sequence from Magweto isolated from *Rhinolophus* spp. and two from Mabura cave isolated from *Hipposideros* spp.*,* clustered with a strain of bat astrovirus isolated from *Myotis* spp. in Madagascar (references K, L) (Figure 4). A cluster, comprising two clades from Magweto and one sequence from Chirundu and three sequences from Mabura cave isolated from *Rhinolophus* and *Hipposideros,* were closely related to bat astrovirus strains from *Hipposideros* spp. of Mozambique and Laos (references H, I) (Figure 4). The last small group comprising one clade from Chirundu, four sequences from Magweto, all isolated from *Rhinolophus* spp.*,* and one sequence from Mabura isolated from *Hipposideros* spp. clustered together with bat astrovirus strains from *Rhinolophus* spp. of Laos and Korea (references D, E, F) (Figure 4).

One of the large clusters shows the phylogenetic relationship amongst sequences derived from, *Hipposideros* spp.(Clades HC-5 to -8), these making up the majority of the sequences in this clade and one isolated from *Miniopterus* spp., one from *E. helvum* and one from a *Nycteris* spp.. The bat astroviruses in this clade, all showed a close relation to a bat astrovirus strain isolated from *Hipposideros* *spp.* of Mozambique (reference C). The largest cluster constituted mainly by Magweto site sequences and some Chirundu sequences, from *Hipposideros* spp. (clades HC-1 to 4) and a few *Rhinolophus* and *Nycteris* spp., clustered with a strain from *Hipposideros* spp. of Mozambique (references A, B) (Figure 4).



**Figure 4*:* Phylogenetic tree of *Astroviruses* partial *RdRp* gene**. The sequences detected at Chirundu site are represented by red branches, at Magweto site by blue branches, at Honde Valley by green branches, at Mabura cave by brown branches and at Baobab tree site by an orange branch. Human *Astrovirus* are highlighted in red rectangles and bold letters represent the different bat AstV references used to build this tree (See Supplementary Data: Figure 4). The tree was built using the maximum likelihood method based on the GTR + G4 + I model. The robustness of nodes was assessed with 1000 bootstrap replicates. Bootstrap values >70 are in asterisk and those <70 are not shown. HC=*Hipposideros caffer*; RS=*Rhinolophus simulator*; NT=*Nycteris thebaica*; EH= *Eidolon helvum*; Min=*Miniopterus* spp.; Rhi=*Rhinolophus* spp.. MAG=Magweto site; CHI=Chirundu site; BAOB= Baobab; MAB= Mabura site; HV=Honde valley site. The scale bare represented the number of substitutions per site.

**Discussion**

The overall AstVs prevalence was respectively 10.0% and 13.7% in Chirundu and Magweto sites, confirming their circulation in bat populations from Zimbabwe. However, compared to studies in other African countries, the observed prevalence was low: Hoarau et *al* and Lebarbenchon et *al* detected *AstV* in 20-22% of individual bats in Mozambique and Madagascar, respectively [21,23]. This difference in prevalence of AstV compared to other countries can be attributed to different study designs, longitudinal sampling in our case versus transversal studies in Mozambique and Madagascar [21,23]. Furthermore, these differences could also result from the studied bat species having varying immunological responses and susceptibilities to different viruses [40], and the different localities resulting in subtle changes that may impact the infection dynamics.

In Chirundu site increasing RNA-AstV prevalences were observed from the lactation to the 4-6 months old juvenile seasons with 21.2% of RNA-AstV detection during the latter. Astroviruses have been described to display seasonal variations in prevalence [23]. Drexler et al observed different peaks of astrovirus detection associated with different stages of the reproductive season with peaks correlating to maternal aggregations during breeding season and after parturition season due to establishment of a susceptible sub-population of weaned new-born bats who did not yet have their own adaptive immunity [24]. In our study the prevalence was high during some of the months compared to overall prevalence, which might be related to the effect of reproductive cycle stages on the shedding of viruses, as demonstrated for other viruses. In another study the reproductive season with juveniles and immature individuals showed a very high prevalence of CoV as compared to prevalences in the absence of juveniles and presence of sexually mature bats thus suggesting different stages of the reproduction having effect on viral shedding [31, 41–43]. In this study we observed a similar trend for AstVs for one of the study sites. Furthermore, Mendenhall et *al*, also identified the bat juvenile stage as exhibiting a greater AstVs viral burden than any other stage of the reproductive season [43]. The range of prevalence observed during the highest peaks coincides with the ≥20% overall AstV detection reported by Hoarau et *al* [21] during the time of their sampling in February and May*.* Here we observe a peak during the 4-6 months old juvenile season in Chirundu site which corresponds to a high influx of immunologically immature individuals in the bat population, similar to the what is observed in the Chidoti et *al*, (2022) for CoV infection [31].

Juveniles and immature individuals are known to contribute to shedding pulses of viruses as they develop productive infections during the acute phase [27], which coincides with the waning protection of maternal antibodies [31], thus increasing the susceptibility and rate of infection in the young⁠.

We could not observe a clear trend in Magweto site, which might be related to the absence of samples from December 2020 to March 2021, which correspond to the lactation and weaning periods⁠⁠.

In our study we also investigated co-infection of bats by coronaviruses and astroviruses. We compared the co-infection prevalence with the prevalence of astroviruses and coronaviruses described at each site during the same reproductive cycles. The co-infection prevalence showed a similar trend to that observed in the individual viruses, whereby during observed peaks of high AstVs and CoVs detections, the co-infection also peaked at similar phases of the reproductive cycle. Thus the co-infection of bats by both CoVs and AstVs is high when there is high infection of bats by either virus.

The overall prevalence of co-infection was 3.5% and 2.6 % in Chirundu and Magweto respectively while the bats from Madagascar and Mozambique respectively showed a AstV-CoV co-infection of 5% (±2.7%) and of 1.7% (±1,1%) [9,40]. Co-infection and recombination of viruses in bats have been reported on several occasions including co-infection of bats by coronaviruses and astroviruses [9,40,44]⁠. Bats infected with either CoV or AstV were shown to be more likely co-infected with the respective virus [26]. In our study, no effect of coronavirus infection/shedding on astrovirus infection/shedding was observed.

In the current study we detected bat Astroviruses from *Hipposideridae, Rhinolophidae*, *Pteropodidae, Nycterida*e and *Miniopterida*e families. Astroviruses are known to show no host restriction and are widespread within the Chiroptera order [16,24]. The majority of the astroviruses amplified were from *Hipposideros* and *Rhinolophus* species and these clustered with AstV sequences derived from Mozambique, Madagascar and China of the same families [21,23,24]. The observed trend is due to the two genera being the most dominant species of bats at both sites where sampling was done. Therefore, the probability of higher detection of AstV in these species is expected and evidently observed to be higher than other rare or less dominant species. Active transmission events of astroviruses amongst *Rhinolophidae* and *Hipposideridae* bat species are known to occur [24]. In this study, a high degree of species-specific tropism, especially in bat astrovirus related clusters was observed with astrovirus strains isolated from *Hipposideros* spp. clustering with each other and similar trends for *Rhinolophus* and *Miniopterus* spp.. However, the clusters were not site-specific as sequences from Magweto, Mabura and Chirundu sites formed clusters with each other. We also observed a large phylogenetic cluster which comprised specific *Hipposideros and Rhinolophus* AstV sub-clades (HC-1 to -8 and Rhi-1 to -4) as well as few AstV sequences from other bat species (Figure 4). Previous studies on astroviruses in bats have also described this family of viruses to be involved solely in tropism transmission which is species specific [45,46]. AstVs sequences isolated from different bat species of the same site clustered together. This indicates potential cross species transmission of astroviruses within each site, as reported by Xiao et *al* in bats sharing the same habitat [46]⁠. We also observed phylogenetic clustering of astroviruses from Chirundu and Magweto sites into same clades showing a great diversity of astroviruses in circulation within the studied bat communities. Novel astroviruses species have been reported in bats hosts [25]⁠⁠⁠, and in this study it is evident due to the diverse astroviruses described. The *AstV* isolated from *E. helvum*, frugivorous bats, clustered together and showed close phylogenetic relation to *BAstV* isolated from *Rousettus* spp*.* bats in China and *Taphozous* spp*.* bats in Laos. One strain isolated from *E. helvum* clustered with *Hipposideros* bat astroviruses sequences isolated in Chirundu site. This could mean that frugivorous bats and insectivorous bats can be infected by the same type of AstVs strain thus indicative of the multi-host spectrum for the Astroviridae family. Detection of AstVs in the frugivorous bats was low compared to insectivorous bats, and this can be attributed to the insectivorous bats ability to adapt and harbour more astroviruses than frugivorous bats [46]⁠⁠⁠.

Two strains, Mag-292 *H. Caffer*, Mag-573 *Rhinolophus* spp*.* clustered within the *HuAstv* type 1 to 6, that are known to cause infection and diarrhoea in children and infants [46]. Six of our strains isolated from *E. helvum* and one from *Rhinolophus* spp. formed sister clades to *HuAstV* HMO, associated with severe extra-intestinal illnesses in humans [5]⁠ and VA-1-4 strains. Another AstV strain detected in *Hipposideros* bat species clustered with *HuAstv* MLB-1, -2 and -3. That are all known to cause gastro-intestinal and neurological diseases with mild-severe symptoms in humans [47]. This phylogenetic clustering could suggest that *HuAstV* and *BAstV* may have a shared common ancestor. However, with limited sequence data the evolutionary history is merely a speculation that needs to be further investigated through full genome analysis. Another hypothesis for the observed relation is that, since bats are known to be important reservoir hosts of astroviruses with great diversity, they could be ancestors of the *HuAstV* due to the high prevalence of AstV detected in bats. *HuAstV* transmission occurs via the faecal-oral route and contaminated food or water [1]⁠⁠⁠, therefore it is unlikely that anthroponosis may have occurred. Third, this observed relation could be due to a rare event or mutation, or a new strain of bat astrovirus that has not been described yet. Further full genomic data as well as epidemiological data on these bat colonies are needed to reach a full conclusion.As novel strains and host species continue to be discovered, understanding AstV transmission and their zoonotic potential is essential. Dissemination of *HuAstV* and AstV from zoonotic reservoir hosts remain a significant threat to public health [7].⁠

**Conclusion**

This study offered important contributions to the understanding of virus diversity in an under-sampled region of the world, and the sequences contribute to a better picture of co-evolution and transmission among bat species. Multiple sampling points as in the study further give a more comprehensive picture of how viruses vary over space and time and host species. Furthermore, the differences in prevalence between the two main individual populations can be attributed but not limited to differences between locations, sex, population dynamics and bat species and also anthropogenic factors therefore there is need for a more comprehensive study to fully establish the factors contributing to the AstV viral dynamics among bat populations. Astroviruses have a high genetic diversity, multiple mechanism of generating additional diversity, and infect a wide range of host species therefore understanding their prevalence and infection dynamics in their wildlife hosts can help to predict or prevent the emergence of novel astrovirus strains into domestic animals and human populations

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**Data, scripts, code, and supplementary information availability**

***Sampling Data, Astrovirus sequences, Graphs and Figures***

**https://doi.org/10.5281/zenodo.7856230**

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***GenBank Accession Numbers***

The AstVs sequences have been deposited to the GenBank under the following numbers: OQ271049 - OQ271203

The Cytochrome B sequences have been deposited to the GenBank under the following numbers: OM487705-OM488020

***Statistical analysis***

The script used for the GLMM analysis is available at: https://doi.org/10.5281/zenodo.7847934

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