Full Genome Sequencing of Corriparta Virus, Identifies California Mosquito Pool Virus as a Member of the *Corriparta virus* Species

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Abstract

The species *Corriparta virus* (CORV), within the genus *Orbivirus*, family *Reoviridae*, currently contains six virus strains: corriparta virus MRM1 (CORV-MRM1); CS0109; V654; V370; Acado virus and Jacareacanga virus. However, lack of neutralization assays, or reference genome sequence data has prevented further analysis of their intra-serogroup/species relationships and identification of individual serotypes. We report whole-genome sequence data for CORV-MRM1, which was isolated in 1960 in Australia. Comparisons of the conserved, polymerase (VP1), sub-core-shell 'T2' and core-surface 'T13' proteins encoded by genome segments 1, 2 and 8 (Seg-1, Seg-2 and Seg-8) respectively, show that this virus groups with the other mosquito borne orbiviruses. However, highest levels of nt/aa sequence identity (75.9%/91.6% in Seg-2/T2: 77.6%/ 91.7% in Seg-8/T13, respectively) were detected between CORV-MRM1 and California mosquito pool virus (CMPV), an orbivirus isolated in the USA in 1974, showing that they belong to the same virus species. The data presented here identify CMPV as a member of the *Corriparta virus* species and will facilitate identification of additional CORV isolates, diagnostic assay design and epidemiological studies.

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Introduction

Corriparta viruses are mosquito-borne arboviruses, classified within one of the 22 virus species currently recognised within the genus *Orbivirus*, family *Reoviridae*. Currently there are also 15 'unclassified' orbiviruses in the genus, which may represent additional species [1–6]. The orbivirus genome is composed of 10 segments of linear dsRNA, packaged as one copy of each segment within each of the non-enveloped icosahedral virus particles. The intact virion is composed of three concentric protein shells (the 'outer-capsid', 'core-surface layer and the 'subcore-shell'). Orbiviruses are transmitted by ticks or hematophagus-insect vectors (including *Culicoides*, mosquitoes or sand flies) and collectively have a wide host-range that includes both domesticated and wild ruminants, equids, camelids, marsupials, sloths, bats, birds, large canine and feline carnivores, and humans [2,7–9].

The species *Corriparta virus* currently contains six distinct viruses, that are identified as: corriparta virus MRM1 (CORV-MRM1); CS0109; V654; V370; Acado virus; and Jacareacanga virus [2]. The structural and chemical properties of the corriparta viruses are similar to those of other orbiviruses [2]. They are sensitive to

low pH and heat, and can be modified by treatment with trypsin or chymotrypsin [10]. They have also been shown to multiply in mosquitoes after intra-thoracic inoculation [10].

Members of the *Corriparta virus* species/serogroup have been detected in Australia, Africa and South America [11]). They have been isolated from wild birds, and neutralizing antibodies were found in wild and domestic birds, cattle, marsupials, horses and man [12–15]. Corriparta virus MRM1 was isolated in 1960, from *Culex* mosquitoes, as well as from *Aedeomyia catasticta*, a rare mosquito species collected near Mitchell River in North Queensland, Australia. Subsequently, strains CS0109, V654 and V370 were also isolated in Australia [2,8,15,16]. Acado virus and Jacareacanga virus were isolated from pools of *Culex* mosquitoes collected in Ethiopia and Brazil during 1963 and 1975 respectively [8,11].

The International Committee on Taxonomy of Viruses (ICTV) has agreed 'polythetic' definitions for individual virus species [17]. The ability to exchange genome segments with other viruses belonging to the same virus species by 'reassortment' is recognised as the primary determinant of *Orbivirus* species [2,7]. However, in the absence of data concerning their compatibility for reassort-

ment, the members of individual species can be identified by other 'polythetic' parameters that include similarities in RNA and protein sequences, their RNA-segment size distribution (reflected by their migration patterns - electropherotype) during agarose gel electrophoresis [AGE], host and/or vector range, the clinical signs of infection, and serological relationships [2,7,18–20].

The members of the different *Orbivirus* species were originally identified as belonging to distinct 'serogroups', based on their cross-reactivity in 'group-specific' serological assays that include complement fixation (CF) tests, group-specific ELISA, or agar-gelimmuno-diffusion (AGID) tests, most of which target outer-core protein VP7(T13) [2,7,21]. The corriparta viruses were initially grouped primarily on the basis of CF tests [8,22]. However, a lack of neutralization assays has prevented further analysis of their intra-serogroup serological-relationships and the identification of distinct serotypes.

Recently, full genome sequencing and phylogenetic analyses have been used to determine the genetic relatedness and taxonomic status of individual isolates belonging to different Orbivirus species, including Bluetongue virus (BTV); African horsesickness virus (AHSV) and Epizootic hemorrhagic disease virus (EHDV) [1,23-26]. These sequence data have supported development of faster and more reliable, virus-species/serogroup, and virus-serotype specific diagnostic assays, using both conventional and real-time RT-PCRs [27-31]. Comparisons of nucleotide (nt) and amino acid (aa) sequence data also provide a basis for grouping of orbivirus isolates into topotypes and for molecular epidemiology studies [23,25,32,33]. However, full-genome sequence data are currently available for representatives of only 11 of the recognized Orbivirus species [24] (accession numbers given in Table S1). Additional partial-sequences are available for the highly conserved genome-segment encoding the subcore 'T2' protein (VP3 of BTV) of some Orbivirus species, including Warrego virus (WARV), Wallal virus (WALV), Wongorr virus (WGRV) and CORV [32] (Table S1).

'California mosquito pool virus' (CMPV) was isolated in 1974 from pooled *Culex tarsalis* mosquitoes collected as part of an infectious agent surveillance program conducted by The California Department of Public Health [5]. Partial sequences for genome segments 2, 4, 6, 7 and 9 from CMPV (accession numbers EU789391 to EU789395) were compared to available data for other orbiviruses, suggesting that CMPV might represent a novel virus species [5]. However, the lack of reference sequences for representatives of all *Orbivirus* species, made it impossible to confirm the taxonomic status and species identity of CMPV at that time.

We report the full genome sequence of CORV–MRM1 (AUS1960/01). Comparisons of nucleotide (nt) and deduced amino acid (aa) sequences for the conserved polymerase 'VP1(Pol)', subcore-shell 'T2-protein', and outer-core 'T13-protein', to data published for other orbiviruses, indicate that CORV and CMPV belong to the same species - *Corriparta virus*.

Results

Virus propagation and dsRNA 'electropherotype'

CORV–MRM1 induced severe cytopathic effect (CPE) in BHK cell monolayers, by 48–72 hours post infection (pi). The viral dsRNA was purified from infected cell-cultures and analysed by 1% agarose gel electrophoresis (AGE) (Figure 1). CORV Seg-3 (encoding the larger outer capsid protein VP3(OC1)) migrates close to the medium sized genome-segments (Seg-4, Seg-5 and Seg-6) giving a 2-4-4 (2-4-3-1) migration pattern (Figure 1). This contrasts with the 3-3-4 (3-3-3-1) pattern that is more typical of the



Figure 1. Agarose gel (1%) electrophoresis for dsRNAs of CORV, BTV and EHDV. The purified genomic RNA of LIB2007/07 (a western-topotype isolate of BTV serotype 1 (BTV-1w)) - lane 1: AUS1982/05 (an eastern-topotype isolate of EHDV serotype 8 (EHDV-8e)) - lane 2: and CORV-MRM1 (AUS1960/01) - lane 3, were analysed by 1% agarose gel electrophoresis (AGE), then stained with ethidium bromide, visualised and photographed under UV illumination. The genome segment numbers (encoded proteins) are indicated at the side of the gel.

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Culicoides borne orbiviruses (as illustrated for BTV-1w [LIB2007/07] and EHDV-8e [AUS1982/05]).

Characterisation and coding assignments of CORV genome segments

Sequences for Seg-1 to Seg-10 CORV-MRM1 (AUS1960/ 01) have been deposited in the GenBank with accession numbers KC853042 to KC853051, respectively. They range from 3,925 bp to 790 bp (encoding proteins of 1,290 aa to 108 aa) with a total length of 19,093 bp, (Table 1). The different genome segments of CORV-MRM1 all share six fully conserved nucleotides at their 5' ends, and 10 at their 3' ends (+ve: 5'-GUAUAG......CAAAGGAUAC-3'). Two terminal nucleotides at the 5' end and three nucleotides at the 3' end (5'-GU.....UAC-3') are characteristic of the genus *Orbivirus* and the first and last two nucleotides represent inverted complements (http://www.reoviridae.org/dsRNA_virus_proteins/ CPV-RNA-Termin.htm).

Collectively the terminal non-coding regions (NCR) represent 4.99% of the CORV-MRM1 genome (Table 2). Comparisons with data from GenBank for other mosquito-borne-orbiviruses (MBOs) showed similar or higher percentages, while the tick-borne orbiviruses (TBOs) and *Culicoides*-borne orbiviruses (CBOs) show similar or lower values (Table 2).

Like other orbiviruses, most genome segments of CORV-MRM1 (AUS1960/01), have shorter 5' than 3' NCRs, except for Seg-6 (encoding the smaller outer capsid protein VP5(OC2)) which has a longer 5' NCR and Seg-9 (encoding VP6) which has 3' and 5' NCRs of equal length (Table 1). Exceptions also occur in Umatilla virus (UMAV) and Great Island virus (GIV) although the significance of these variations is unclear.

Seg-5 (1,936 bp) of CORV-MRM1 which encodes the 'tubule' protein NS1(TuP) (one of the most abundantly expressed orbivirus

Seg No.	Seg size (bp)	G+C content (%)	Protein encoded (structure/putative function)	ORFs bp (including stop codon)	5'NCR	3'NCR	No. of amino acids	Predicted protein molecular mass (<i>kDa</i>)	5' Termini	3′ Termini	Accession numbers
-	3,925	41.66	VP1 (Pol)	12-3884	11	44	1290	147.07	5'-GUAUAGG	CCAAAGGAUAC-3	KC853042
7	2,952	43.43	VP2 (T2)	33-2894	32	61	953	108.75	5'-GUAUAGG	CCAAAGGAUAC-3'	KC853043
m	2,284	42.43	VP3 (OC1)	20-2236	19	51	738	84.81	5'-GUAUAGG	CCAAAGGAUAC-3	KC853044
4	2,032	43.11	VP4 (Cap)	12-1943	11	92	643	74.83	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853045
5	1,936	46.33	NS1 (TuP)	20-1789	19	150	589	68.61	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853046
9	1,683	46.11	VP5(OC2)	56-1639	55	47	527	57.7	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853047
	1,187	46.5	NS2 (ViP)	26-1141	25	49	371	41.86	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853048
8	1,176	47.62	VP7 (T13)	17-1081	16	98	354	38.96	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853049
6	1,128	47.16	VP6 (Hel)	49-1083	48	48	344	36.85	5'-GUAUAGA	UCAAAGGAUAC-3'	KC853050
		I	NS4	143-601	142	530	152	18.37			
10	790	47.22	NS3	17-733	16	60	238	25.92	5'-GUAUAGG	CCAAAGGAUAC-3'	KC853051
			NS3a	407-733	406	60	108	11.848			

The size of the highly conserved T13 core-surface protein (VP7 of BTV) is also relatively consistent in most orbiviruses, while the viral inclusion-body matrix-protein, (non-structural protein 2 [NS2(ViP)]) is more variable in size (Table 2). As a result, NS2 of CORV-MRM1 (AUS1960/01) is encoded by Seg-7, while Seg-8 encodes the core-surface protein VP7(T13) (Table 1), in a manner similar to the other MBOs (Table 2). However, this coding-assignment is again reversed in BTV and in some CBOs (EUBV and CHUV), due to variability in the size of NS2 between different viruses [34,35]. The capping enzyme 'VP4(CaP)' of CORV-MRM1 (at 643aa) is smaller than that of the other MBOs and some of the CBOs (Table 2). However, it is encoded by the largest Seg-4 (at 2,032 bp) so far identified in any orbivirus. Most of the genome segments of CORV-MRM1 (AUS1960/ 01) except Seg-9 and Seg-10 are monocistronic, encoding a protein from a single large ORF, starting from an initiation codon with a strong Kozak sequence (RNNAUGG) [36]. However, like BTV, CORV Seg-10 has two in-frame AUG initiation sites encoding the NS3 and NS3a proteins of 238 aa and 108 aa (starting at 17 bp and 407 bp) respectively (Table 1). The first of these (coding for NS3) has a 'weak Kozak context' (GUAAUGU), possibly enhancing read-through and initiation of translation from the second 'in frame' initiation site (at 407 bp). This has a strong Kozak context (GUUAUGG), but would express the smallest NS3a in any of the orbiviruses characterized to date.

The first start codon of CORV-MRM1 (AUS1960/01) Seg-9 also has a moderate Kozak sequence (UUGAUGA) and a second down-stream ORF. However, this is in the +2 reading-frame (at 143-598 bp), encoding the 152 aa NS4 protein. NS4 has previously been identified in several other orbiviruses and has been characterised in BTV and GIV [24,37,38]. The downstream Seg-9 ORF of CORV has a strong Kozak context (AGGAUGG) and is expected to produce a protein in infected cells. Weak or moderate Kozak sequences have also been observed in several of the genome segments of other orbiviruses, but they still appear to be translated effectively [24,39].

The G+C content of the CORV genome is 45.16%, which is considerably higher than that of other MBOs [Peruvian horse sickness virus (PHSV) and YUOV with 36.66% and 41.59% respectively] but within the overall G+C range of the insect-borne orbiviruses [36.66% in PHSV (mosquito), to 45.86% in Equine encephalosis virus (EEV) (Culicoides)]. However, it is lower than that of the tick-borne or tick-associated orbiviruses [57.29% in GIV and 51.93% in St Croix River virus (SCRV)].

Phylogenetic comparisons of subcore-shell 'T2' proteins

Comparison of homologous orbivirus proteins using BlastX analysis identified VP2, encoded by Seg-2 of CORV, as the subcore shell 'T2' protein (equivalent to VP3(T2) of BTV and VP2(T2) of YUOV and SCRV). An unrooted neighbour-joining (NJ) phylogenetic tree (Figure 2) constructed for different orbivirus

proteins) is longer than the homologous Seg-5 of the CBOs or TBOs (Table 2) with a long (150 bp) 3' NCR, but is smaller than that of some other MBOs [1]. Long 3' NCRs were also observed in the NS1 genome segment of some other MBOs (e.g. UMAV-278 bp; Yunnan orbivirus [YUOV]-205 bp), but not in all other orbiviruses (<118 bp).

The 'highly conserved' subcore-shell VP2(T2) protein and the 'highly variable' outer-capsid/cell-attachment protein VP3(OC1) of CORV-MRM1, are encoded by Seg-2 and Seg-3, respectively. A similar coding pattern is seen in other MBOs, but the presence of a larger OC1 in the CBOs (including bluetongue virus, the orbivirus 'type' species) results in a reversed coding assignment for these two genome-segments (Figure 1 and Table 2).

	CORV		Ň	AV		PHSV		۲.	Ō		BTV-8	3	A	HSV-1		EHD	V-1e	Ш	>		EUB			NUH		٩N		Ň	CRV	
Protein	qd ni əziz tnəmpəz	protein size in aa	dq ui əzis	Jnemes	ss ni sziz nistorq	dq ui əzis	Segment protein size in aa	dq ui əzis	JnemteS	ss ni sziz nistorq	dq ui əzis	Segment	ad ni sziz	da mengel	es ni sziz nistord	dq ui əzis	se ni szis nistorg	dq ui əzis	tnempez	ee ni sziz nistorq	dq ui əzis	2egment	es ni sziz nisjord	Seament	ee ni sziz nistord	dq ui əzis	tnempez	dq ui əzis	tnempe2	protein size in aa
Pol	3925 S1	1 129	10 393.	3 S1	1299	3987	S1 15	311 395	33 S1	1315	3944	S1 1	302 35	966 S1	1 1305	3942	S1 1:	302 39	53 S1	1307	3962	S1	307	930 S	1 1295	3897	S1 1	285 40	89 S1	1345
T2	2284 S2	2 755	279	1 S2	905	2856	S2 92	25 290	00 S2	940	2772	S3 9	01 2;	792 SE	3 905	2768	S3 8:	31:	78 S3	3 902	2773	23 23	006	774 S	3 904	2794	S2 9	08 27	'47 S2	890
0C1	2952 S3	3 953	3 241	1 S3	797	2747	S3 8{	81 266	38 S3	873	2939	S2 9	61 3.	218 Sž	2 1056	2968	S2 9;	71 27.	75 S2	2 1042	2958	S2	968	055 S.	2 1002	1722	S4 5	51 20	24 S4	654
Cap	2032 S4	4 643	1 202	1 S5	648	1996	S4 64	46 199	93 S4	645	1981	S4 6	44 19	978 S4	4 642	1983	S4 64	198	36 S4	t 646	1984	S4	144	967 S.	4 640	1936	S3 6	35 20	17 S3	643
Tup	1936 S5	5 589	204	7 S4	580	1784	S5 55	54 195	57 S5	574	1776	S5 5	52 1.	748 S5	5 548	1806	S5 55	51 174	49 S5	548	1763	S5	56 1	764 S	5 545	1731	S5 5	31 16	57 S5	517
0C2	1683 S6	6 527	168	1 S6	529	1695	S6 52	29 168	33 S6	535	1637	S6 5	26 1	564 St	5 505	1640	S6 5.	27 16(07 S6	515	1658	S6	529 1	610 S	5 521	1666	S6 5	37 16	64 S6	517
T13	1176 SE	8 354	t 117	1 S8	351	1180	S8 35	53 115	91 S8	355	1156	S7 3	49 1	167 S£	3 349	1162	S8 34	11.	75 S8	350	1175	S7	350 1	151 S	7 348	1181	S7 3	57 12	56 S7	379
ViP	1187 S7	7 371	135	7 S7	407	1613	S7 43	35 150	04 S7	435	1125	S8 3	54 1	166 SS	365	1186	S7 3.	73 118	31 S7	7 362	1128	S8	351 1	059 S	3 333	1172	S8 3	59 14	63 S8	462
Hel	1128 SS	9 344	110	3 S9	341	1071	S9 35	34 106	32 S9	338	1049	S9 3	29 1	169 S7	7 369	1143	S9 3(50 108	30 SG	341	1047	S9	328 8	77 S	9 272	1056	S9 3	21 76	4 S9	232
NS4	- S5	9 152		S9	149		S9 1	11	S9	113		S9 7		- S	7 117		S9 7(SS	9 145	I	S9	34	ŭ.	9 83	I	S9 1		S9	93
NS3	790 S1	10 238	887	S10	264	819	S10 25	55 825	5 S1C) 253	822	S10 2	29 7(63 S1	10 218	810	S10 2.	28 755	9 S1	0 240	856	S10.2	242	28 S	10 211	703	S10 1	71 76	4 S10	224
Total Genome (bp)	19093		194	02		19748		196	316		19201		-	9531		1940	œ	19	153		1930	4	-	8915		17858		31	445	
%NCR	4.99		5.71			5.46		5.1	8		3.96		т.	.79		4.09		3.5	7		4.04		,	.71		5		4.	4	
G+C (%)	45.16		41.5	5		36.66		41.	59		44.98		4	3.46		43.02		45.	86		45		,	9.89		58.13		51	.92	
Vectors	Mosquit	itoes									Culico	ides s	sdd.													Tick o	or Tid	assoc	iated	
Pol = Poly T13 = Out doi:10.137	/merase, O ter core pr '1/journal.p)C1 =(rotein (pone.0	Outer c (T13 sy. 1070775	apsid p mmetry .t002	y), ViP =	1 (VP2 (= Viral i	of BTV nclusic), T2 = Ir on body	nner co proteir	n (NS2)	tein (T2 , Hel =}	symm Helica:	etry), C se prote	i ap = Ci ein. NCI	apping - R and G	enzym€ i+C con	, Tup = tent we	Tubule re calcu	formir lated	for full	ein or T genom	ubular e sequ	proteir ences.	(NS1),	0C2 = (Duter ca	psid p	otein 2	(VP5 of	BTV),

Table 2. Full genome sequence database available for recognised species of genus Orbivirus and their genome coding assignments.



Figure 2. Unrooted neighbour-joining tree for orbivirus subcore-shell T2 proteins. A phylogenetic tree was constructed using distance matrices, generated using the p-distance determination algorithm and pairwise deletion parameters in MEGA 5 (1000 bootstrap replicates) [65]. Since many of the available sequences are incomplete, the analysis is based on partial sequences (aa 393 to 548, numbered with reference to the aa sequence of BTV-VP3(T2)). The numbers at nodes indicate bootstrap confidence values after 1000 replications. The tree shown in Figures 3 and 4 were drawn using the same parameters. The CORV and CMPV isolates are shown in red font in the amber coloured circle. Full names of virus isolates and accession numbers of T2 protein sequences used for comparative analysis are listed in Table S1 (supplementary data). 'e' and 'w' after serotype number indicate eastern and western topotype strains, respectively. doi:10.1371/journal.pone.0070779.q002

T2 protein sequences (listed in Table S1 supplementary data) identified two major clusters (Figure 2). The larger group in which VP3(T2) is encoded by Seg-3, comprises the Culicoides transmitted orbiviruses (including BTV, AHSV, EHDV, WALV, Eubenangee virus [EUBV], WARV and Palyam virus [PALV]). However, the second group in which VP2(T2) is encoded by Seg-2, includes two sub-groups transmitted by ticks (the Great Island viruses), or by mosquitoes (WGRV, UMAV, PHSV and YUOV) respectively. SCRV, which has previously been suggested as a tick-orbivirus rather than a tick-borne-orbivirus [40], branches separately from these groups and appears to be more distantly related to other orbiviruses. VP2(T2) of CORV-MRM1 (AUS1960/01) clusters very closely with partial sequence data available for CMPV(98.78%/98.99% aa/nt identity Table 3 and 4), within the mosquito-borne group, indicating that they belong to the same virus species. These two viruses show lower levels of identity with the other Orbivirus species analysed (<58.53%/57.98% aa/nt identity - Table 3).

Phylogenetic comparisons of VP1(Pol) proteins

Seg-1, which encodes the viral RNA polymerase VP1(Pol), is highly conserved across the genus *Orbivirus*, and provides a target for sequencing and RT-PCR assays to identify virus isolates at both species and genus levels [1,24,31,40]. The aa sequence of VP1(Pol) from CORV-MRM1 (AUS1960/01) was compared with representatives of other *Orbivirus* species to construct an unrooted NJ phylogenetic tree (Figure 3). Three major clusters were identified, corresponding to the different vector-groups: CBOs, MBOs and TBOs. CORV-MRM1 (AUS1960/01) again clusters as a distinct species, close to other MBOs (>51.44/54.9% aa/nt identity - Table 3), with more distant relationships to VP1 of the *Culicoides* and tick-borne groups (<50.51%/53.99% aa/nt identity - Table 3).

Phylogenetic comparisons of outer-core 'T13' proteins

The orbivirus core-surface protein VP7(T13) is immunodominant and represents a major serogroup-specific antigen [21]. An unrooted phylogenetic tree for VP7(T13) (encoded by Seg-8 of CORV-MRM1) (Figure 4) shows a similar topology to the VP2(T2) and VP1(Pol) trees (Figures 2 and 3), grouping viruses according to their vectors. CORV-MRM1 clusters with CMPV (sharing 91.7%/77.6% aa/nt identity) as members of a distinct species (Table 4), showing lower identities with the other MBOs (<42.57%/47.62% aa/nt identities (with UMAV)), or the CBOs and TBOs (<32.95%/41.57% aa/nt identities (with GIV)) (Table 3).

Relationships between CORV and CMPV

The sequences of CORV-MRM1 Seg-4/VP4(Cap), Seg-6/ VP5(OC2), and Seg-9/VP6(Hel)/NS4 were also compared with

Table 3. Percent nucleotide (nt) and amino acid (aa) identities of Corriparta virus MRM1(AUS1960/01) with other orbiviruses.

Sl no.	Orbivirus species	Abbreviation	T2		VP1		T13	
			aa	nt	aa	nt	aa	nt
1	Bluetongue virus	BTV	37.11	47.33	47.40	52.41	25.07	38.71
		BTV25	36.22	47.00	46.84	52.12	23.63	38.14
		BTV26	37.11	46.56	46.84	51.68	24.21	37.85
2	African horse sickness virus	AHSV	37.06	47.64	47.90	53.03	23.85	39.85
3	Epizootic haemorrhagic disease virus	EHDV	37.19	46.88	47.40	53.06	24.50	39.67
4	Eubenangee virus	EUBV	36.60	47.53	47.60	52.88	27.30	42.53
5	Palyam virus	PALV	35.77	47.14	50.51	53.99	27.67	41.02
5	Equine encephalosis virus	EEV	33.96	45.36	46.61	51.26	24.71	39.18
7	Warrego virus	WARV	44.72	51.78	—	_	—	—
B	Wlallal virus	WALV	44.64	50.83	—	—	—	—
Ð	Umatilla virus	UMAV	50.17	55.21	59.66	58.12	42.57	47.62
		SLOV	50.39	54.29	57.94	57.23		
10	Peruvian horse sickness virus	PHSV	46.37	53.45	52.06	55.38	33.52	43.94
11	Yunnan orbivirus	YUOV	44.78	53.16	51.44	54.90	34.09	45.36
12	Corriparta virus*	CORV	98.78	98.99	_	_	_	_
13	Wongorr virus	WGRV	58.53	57.98	_	_	_	_
14	Great Island virus	GIV	45.20	49.87	50.51	52.13	32.95	41.57
15	St Croix river virus	SCRV	24.15	38.46	39.01	45.23	22.03	35.94
16	Pata virus	PATAV	36.19	46.84	47.23	52.28	26.59	40.75

*data previously published for corriparta virus MRM1 (Ac No. AF530086).

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the incomplete sequence data available for CMPV (Table 4). CMPV again grouped closely with CORV-MRM1 (AUS1960/01), sharing aa/nt identities of 75/67.4% and 86.2/74.3% in VP4(Cap) and VP5(OC2) respectively, further supporting the identification of CMPV as a member of *Corriparta virus* species. Seg-9 of CORV-MRM1 (coding for NS4 and VP6(Hel)) shares only 42.6/61.6% aa/nt identities with CMPV. However, no closer matches were found in either of these proteins with members of other *Orbivirus* species. Further analysis of CMPV Seg-9 confirmed the presence of an alternate ORF, encoding a 153 aa NS4 protein, which shares 55.9%/63.2% aa/nt identity with NS4 of CORV-MRM1.

Discussion

Collectively the orbiviruses infect a wide range of hosts and are transmitted by a diverse group of vectors, including Culicoides, mosquitoes, sand flies and ticks [7]. Initially, the different 'serogroups' of orbiviruses, which are now recognised as distinct virus species, were identified by CF, AGID, immunofluourescence (IF) tests and/or enzyme-linked immunosorbent assays (ELISA). However, low level serological crossreactions have been detected between some of the more closely related Orbivirus species (for example between isolates of BTV (the Orbivirus type species) and EHDV [41] making it difficult to conclusively identify the species of new isolates by these techniques alone. The high quality reference-strains and antisera needed for these assays are not widely available for all of the existing serogroups/species, and may themselves represent a biosecurity risk. In addition, serological methods do not generate absolute quantitative values for the relatedness of individual virus strains.

In contrast, nucleotide sequence data for reference orbivirus strains and novel isolates can be compared and transmitted easily between laboratories, without risk, providing highly reproducible and fully quantitative numerical values for the relatedness of each genome segment/protein. These data can also be used to unambiguously identify different genome segments, proteins, virus species, topotypes and serotypes [1,25,33,39,42].

Due to their economic significance, full genomes of multiple isolates (including reference strains) of BTV, EHDV and AHSV have already been characterized [23,43–49]. These data have supported development of rapid and reliable molecular methods and diagnostic tools (RT-PCR assays) for identification of virus serogroup/species and serotype [27–31,49–51].

Sequence variations in the outermost orbivirus capsid and cellattachment protein, VP2(OC1) of BTV, correlate with both the geographic origin of the virus (topotype) and with its serotype [42,47]. In contrast sequence variation in the core proteins VP1(Pol) and T2 (VP3 of BTV) correlate only with virus genus, species and topotype [1,32,33,39,40]. However, a lack of fullgenome sequence data for all 22 recognized *Orbivirus* species has hindered identification of isolates belonging to novel *Orbivirus* species and development of nucleic acid based diagnostic tests.

The full-genome sequence data presented here for CORV-MRM1 (AUS1960/01) provides a primary reference for identification of other (novel) members of the species *Corriparta virus* (CORV). Conserved nucleotide sequences are present at both the upstream and downstream termini of the genome segments of different *Orbivirus* species [2,52]. All of the genome segments of CORV-MRM1 (AUS1960/01) contain the terminal sequences (5'-GUAUAG.....CAAAGGAUAC-3') showing several differences from those of BTV (e.g. 5'-GUUAAA......ACUUAC-3'). The significance of a longer conserved 3' terminal region in CORV-

Segment (Protein)	Seg-1/VP1 (Pol)	Seg-2/VP2 (T2)	Seg-3/VP3 (OC1)	Seg-4/VP4 (CaP)	Seg-5/NS1 (TuP)	Seg-6/VP5 (OC2)	Seg-7/NS2 (ViP)	Seg-8/VP7 (T13)	Seg-9#/VP6 (Hel) & NS4	Seg-10/(NS3)
Genome segment of CORV (bp)	3,925	2,952	2,284	2,032	1,936	1,683	1,187	1,176	1,128	062
Partial sequence of CMPV (bp)*		358	I	288	I	646	I	737	596	I
% aa identity (CORV vs CMPV)		91.6	I	75		86.2	I	91.7	42.1/55.9 [§]	I
% nt identity (CORV vs CMPV)		75.9		67.4	I	74.3		77.6	61.6/63.2 [§]	
*Only partial CMPV sequences are av #Identities calculated using reverse c	ailable in GenBank omplement sequei	c nces of CMPV.								

^sIndicates identities of NS4.

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MRM1 is unknown, although it has been suggested that these regions may play a role in initiation of transcription or translation of the RNA or its packaging during virus replication [53,54].

The orbivirus proteins VP1(Pol), 'T2' and 'T13', which are highly conserved have taken priority in development of molecular diagnostic assays and in phylogenetic analyses [24,29,31,32,55]. Studies with large numbers of different BTV and EHDV isolates show >73%, >83% and >73% intra-species as identities in VP1, T2 and T13 respectively, providing useful markers for the identification and classification of existing and novel orbivirus isolates [1.40.44].

CORV-MRM1 shares less than 60% as identity (Table 3) in VP1, T2 and T13, with members of the other recognised Orbivirus species, confirming the classification of Corriparta virus as a distinct species. CORV-MRM1 is most closely related to other MBO species, particularly Umatilla virus (UMAV), with 59.66% and 50.17% aa identity in VP1 and T2 proteins respectively (Table 3). However, CORV-MRM1 shows 91.6%/75.9% aa/nt identity and 91.7%/77.6% aa/nt identity to CMPV in its T2 and T13 protein/gene sequences, indicating that they belong to the same virus species and therefore that CMPV does not represent a new species as previously suggested [5].

Phylogenetic analyses show that although the size of the orbivirus sub-core-shell T2 protein is relatively constant across the genus (between 890 aa in SCRV to 953 aa in CORV), the size of the larger of the two outer coat proteins (OC1) is much more variable (between 551 aa in GIV to 1056 aa in AHSV). This results in a change in relative size, order and numbering of the genome segments encoding the T2 and OC1 proteins, between the tick-associated/transmitted orbiviruses [VP2(T2) and VP4(OC1) - encoded by Seg-2 and Seg-4], the MBOs [VP2(T2) and OC1(VP3) - encoded by Seg-2 and Seg-3 respectively], and the CBOs [VP3(T2) and VP2(OC1) - encoded Seg-3 and Seg-2 respectively] [1,23,34,39,40,56].

Previous studies have suggested that the MBOs have evolved from tick-borne ancestors, with CBOs being last to evolve [39]. The concatermerisation of orbivirus genome segments and subsequent mutations may provide a mechanism that can progressively increase the size of individual genome segments [39,56]. It may therefore be significant that the size of OC1 increases in the order: TBOs (551 aa in GIV and 654 aa in SCRV), MBOs (755 aa in CORV to 881 aa in PHSV) and CBOs (961 aa in BTV to 1056 aa in AHSV-1), (Table 2).

Previous studies also indicate that the orbiviruses have evolved through a process of 'co-speciation' with their vectors [39]. Phylogenetic analyses of the conserved Pol, T2 and T13 proteins (presented here - Fig. 2, 3 and 4), show consistent grouping of the CBOs, MBOs and TBOs. In each case CORV-MRM1 groups with the other MBOs (WGRV, UMAV, PHSV and YUOV).

Corriparta viruses have been isolated in Australia, Africa and South America [11]. However, the data presented here clearly identify CMPV, which was isolated in North America [5], as a member of the species Corriparta virus. The occurrence of these closely related viruses in the Americas and Australia indicates that there has been spread of viruses between these regions, which could be due to movement of infected hosts or vectors. Similar movements are also suggested by the detection of other orbiviruses (UMAV and PHSV, and individual serotypes of BTV and EHDV) in more than one continent, e.g., in Australia, Africa and the Americas [1,28,57-59]. Additional strains of each serogroup/ species, from different locations/origins, need to be isolated and characterised/sequenced to better understand their geographical distribution and its significance.



Figure 3. Unrooted neighbour-joining for orbivirus polymerase VP1(Pol) proteins. An unrooted NJ phylogenetic tree for orbivirus VP1(Pol) proteins was constructed using a p-distance algorithm and pairwise deletion parameters, as indicated in Figure 1. The CORV-MRM1 isolate characterised in this study is indicated in red font in amber coloured circle. Full names of virus isolates and accession numbers of polymerase sequences used for comparative analysis are listed in Table S1 (supplementary data). 'e' and 'w' after serotype number indicate eastern and western topotype strains, respectively. doi:10.1371/journal.pone.0070779.g003



Figure 4. Unrooted neighbour-joining for orbivirus outer-core VP7(T13) proteins. An unrooted NJ phylogenetic tree for orbivirus VP7(T13) proteins was constructed using a p-distance algorithm and pairwise deletion parameters, as indicated in Figure 1. The CORV-MRM1 and CMPV isolates are shown in red font in amber coloured circle. Full names of virus isolates and accession numbers of T13 protein sequences used for comparative analysis are listed in Table S1 (supplementary data). 'e' and 'w' after serotype number indicate eastern and western strains, respectively. doi:10.1371/journal.pone.0070779.g004

The sequences and relative sizes of the VP1(Pol), T2 and OC1 proteins are important evolutionary markers that can help differentiate/group orbiviruses by species, serotypes and topotype. The sequence data generated in this study will facilitate the use of phylogenetic analyses to identify other novel isolates belonging to the *Corriparta virus* species, as well as helping to identify the arthropod vectors involved in their transmission. Further studies are still needed to define the different serotype and topotypes of CORV.

Materials and Methods

Virus propagation

CORV-MRM1 (AUS1960/01), obtained at passage level MB6/BHK2 from the Orbivirus Reference Collection at The Pirbright Institute, was propagated in BHK-21 cell monolayers [clone 13 obtained from European Collection of Animal cell Cultures (ECACC – 84100501)], in Dulbecco's minimum essential medium (DMEM) supplemented with antibiotics (100 units/ml penicillin and 100 μ g/ml streptomycin) and 2 mM glutamine. Infected cell-cultures were incubated at 37°C until they showed widespread (100%) cytopathic effects (CPE). Then viruses were harvested, aliquoted and used for the extraction of viral dsRNA. All virus isolates used in these studies were obtained from diagnostic samples of naturally infected animals and were taken as a part of normal diagnostic investigations by qualified veterinarians in the individual countries.

Extraction and purification of CORV dsRNA

Cell monolayers showing 100% CPE after infection with CORV-MRM1, were harvested and pelleted at 3000 g for 5 min. The viral dsRNA was released and purified using TRIzol® reagent (Invitrogen) as described by Attoui et al [60]. Briefly, the infected cell pellet was lysed in 1 ml of TRIzol®, then 0.2 volume of choloroform was added, vortexed and the mixture incubated on ice for 10 min. The aqueous phase, containing total RNA, was separated from the phenol-chloroform phase by centrifugation at 10,000 g for 10 min before 900 µl of isopropanol was added prior to incubation at -20° C for 2 hours. The RNA was pelleted at 10,000 g for 10 min, washed with 70% ethanol, air dried and dissolved in 100 µl of nuclease free water (NFW). Single stranded RNA (ssRNA) was removed by precipitation with 2M LiCl (Sigma) at 4° C overnight, followed by centrifugation at 10,000 g for 5 min. An equal volume of isopropanol, containing 750 mM ammonium acetate, was mixed with the supernatant. After precipitation at -20° C for 2 hours, the viral dsRNA was pelleted at 10,000 g for 10 min, washed with 70% ethanol, air dried and suspended in 50 µl of NFW. The RNA was either used immediately or stored at -20° C.

Reverse transcription and PCR amplification

CORV-MRM1 genome segments were reverse-transcribed into cDNA using the full-length amplification (FLAC) technique described by Maan *et al.* [61]. Briefly, a 35 base oligonucleotide 'anchor-primer', with a phosphorylated 5' terminus, was ligated to the 3' ends of the viral dsRNAs using the T4 RNA ligase overnight at 16°C. Then dsRNA segments were fractionated on 1% agarose gel and recovered from the gel using a 'silica binding' method (RNaid[®] kit, MP Biomedicals) as per the manufacturer's

References

instructions. The dsRNA eluted in NFW, was denatured at 99°C for 5 minutes, and then snap chilled on ice before synthesising first-strand cDNA using RT system (Promega). The resulting cDNAs were amplified using primers complementary to the anchor primer and high fidelity KOD polymerase enzyme (Novagen). PCR amplicons were analyzed by agarose gel electrophoresis.

Cloning and sequencing of cDNA segments

cDNA amplicons were purified and cloned into the 'pCR®-Blunt' vector supplied with the Zero Blunt® PCR Cloning Kit (Invitrogen). Recombinant plasmid-vectors containing CORV-MRM1 inserts were transformed into One Shot® TOP10 competent cells supplied with the cloning kit. Clones containing the desired inserts were identified by colony touch PCR using M13 universal primers. Plasmids were extracted from the clones identified, using the QIAprep Spin MiniPrep Kit (Qiagen). The plasmids and PCR products were sequenced using an automated ABI 3730 DNA sequencer (Applied Biosystems).

Sequence and phylogenetic analysis

'Raw' ABI sequence data was assembled into 'contigs' using the SeqManII sequence analysis package (DNAstar version 5.0). The ORFs of CORV-MRM1 genome segments were identified and translated to aa sequences for further analysis using EditSeq (DNAstar version 5.0). The putative function of each protein was identified by Blast X comparisons to homologous orbivirus (BTV) proteins in GenBank (http://blast.ncbi.nlm.nih.gov/Blast. cgi?CMD = Web&PAGE_TYPE = BlastHome). Multiple alignments of consensus sequences were performed using Clustal X (Version 2.0) [62], Clustal Omega (http://www.ebi.ac.uk/Tools/ msa/clustalo/) and MAFFT [63] to ensure proper alignment. Aligned protein sequences were back translated to nucleotide sequences using DAMBE [64]) or RevTrans 1.4 server available online (http://www.cbs.dtu.dk/services/RevTrans/) for further nucleotide analysis. Pairwise distance (aa and nt) calculations and phylogenetic trees constructions were done using MEGA 5 software [65] with the p-distance parameter and neighbourjoining method [66]. GenBank nucleotide accession numbers of polymerase (VP1), T2 and T13 protein sequences that were used in phylogenetic analyses are provided in Table S1 (supplementary data).

Supporting Information

Table S1Nucleotide accession numbers for sequencesused in phylogenetic analysis.(DOCX)

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Author Contributions

Conceived and designed the experiments: MB SM NSM HA PPCM. Performed the experiments: MB SM NSM KN MG. Analyzed the data: MB SM NSM HA. Contributed reagents/materials/analysis tools: MB SM RT JB PPCM. Wrote the paper: MB SM PPCM.

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