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# Cyclovirus CyCV-VN species distribution is not limited to Vietnam and extends to Africa

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**Cycloviruses, small ssDNA viruses of the *Circoviridae* family, have been identified in the cerebrospinal fluid from symptomatic human patients. One of these species, cyclovirus-Vietnam (CyCV-VN), was shown to be restricted to central and southern Vietnam. Here we report the detection of CyCV-VN species in stool samples from pigs and humans from Africa, far beyond their supposed limited geographic distribution.**

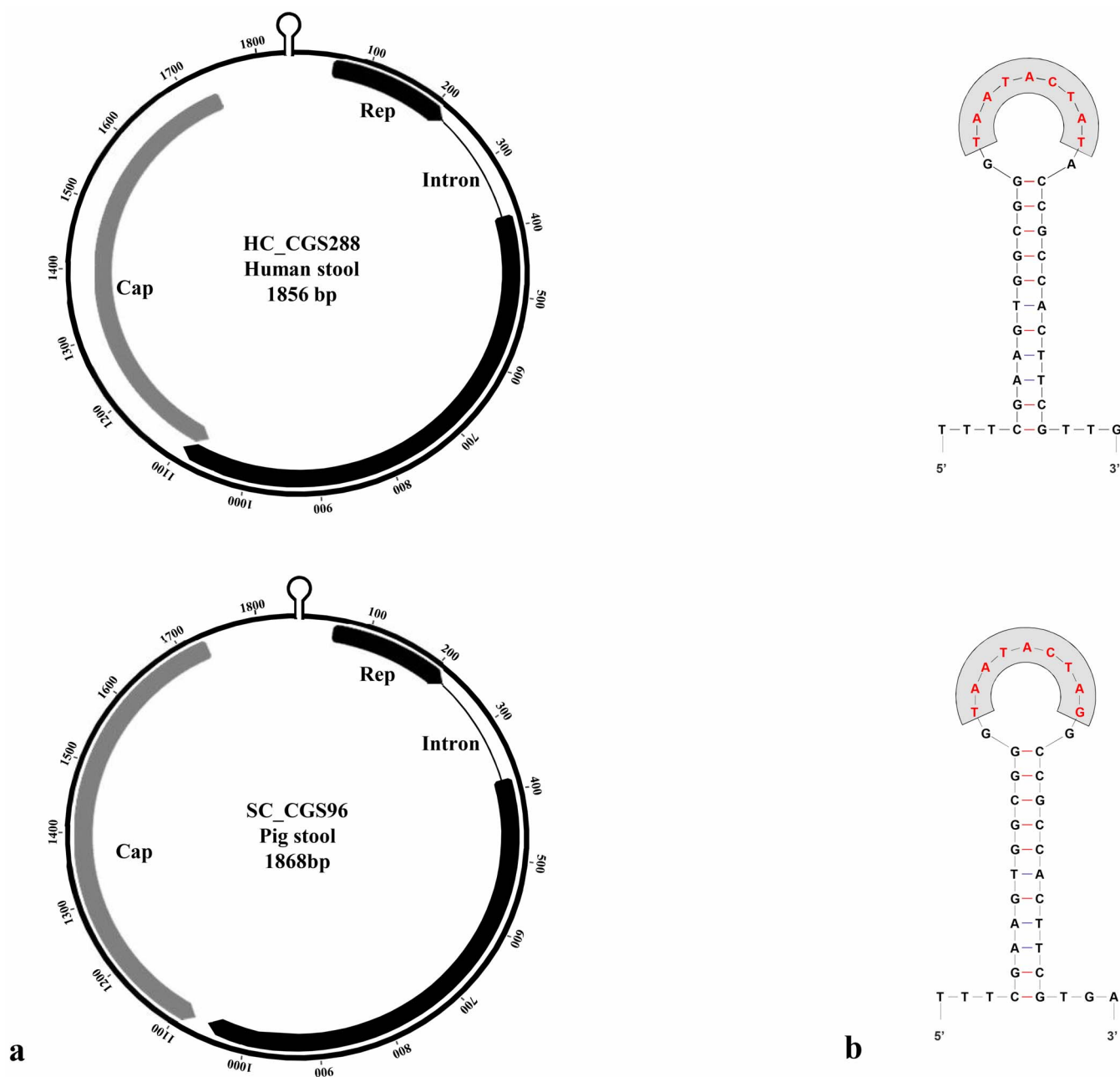
Cycloviruses (CyCVs), a recently proposed genus within the *Circoviridae* family, are non-enveloped single-stranded DNA (ssDNA) viruses with a  $\approx 2$  kb circular genome containing two major inversely arranged open reading frames (ORFs) encoding a putative replication-associated protein (Rep) and a capsid (Cap) protein<sup>1</sup>. An ever growing number of species of CyCVs have been described in mammals (human, chimpanzee, goat, cattle, bat, pig, sheep, and camel), birds (chicken and duck) and insects (dragonflies and wood cockroach)<sup>1–6</sup>. In spite of a high genetic diversity<sup>1,5</sup>, cross-species transmission seems likely, at least in mammals<sup>3,7</sup>.

However, in contrast with circoviruses, the pathological significance of CyCV infection remains unclear. CyCVs have been suggested to cause respiratory and neurological infections in humans<sup>1,7–9</sup>. The presence of the virus was demonstrated in serum, stool or cerebrospinal fluid (CSF) of patients with acute flaccid paralysis, paraplegia or suspected central nervous system infections from Pakistan, Tunisia, Malawi and southern and central Vietnam<sup>1,7,8</sup>. Of particular interest is the description of a high detection rate of the CyCV-Vietnam (CyCV-VN) species in the feces of pigs and chicken from the same area as infected humans<sup>7</sup>. This observation, together with the high level of conservation between human, swine and chicken CyCV-VN strains suggest that pigs and chicken might act as a reservoir for human infections<sup>7</sup>. Subsequent investigations on the circulation of CyCV-VN in human CSF specimens from northern Vietnam, Cambodia, Nepal and the Netherlands were negative, suggesting a limited geographic distribution of this CyCV species<sup>10</sup>.

In this study, the presence of CyCV-VN and CyCV-VN-like strains was demonstrated in human stool samples from Madagascar and swine feces from Cameroon, showing that the geographic distribution extends far beyond southern and central Vietnam as originally thought.

## Results

To assess the circulation of CyCV-VN in Africa, we tested, using a CyCV species-specific PCR, 410 human stool samples from Madagascar, 1337 human stool samples from Ghana and 520 pig stool samples from Cameroon.



**Figure 1** | Genomic organization (a.) and the stem-loop structures (b.) of the cycloviruses discovered in human and animal stool samples from Cameroon and Madagascar. The locations of two major ORFs, encoding the putative Rep and Cap protein are indicated by arrows. The nonamer sequences (marked in red) within stem-loop structures are also shown.

Out of the 410 human samples from Madagascar, three were positive for CyCV-VN-like species and the full genome was amplified, sequenced and characterized (Genbank acc. no. KM392287–KM392289). Ten swine samples from Cameroon were positive (1 from Bamenda, 9 from Douala). The full genome of three positive samples from Douala was obtained (Genbank acc. no. KM392284–KM392286). All human samples from (cases and controls) Ghana remained negative for CyCV-VN.

The genomic sequences of human CyCV strains from Madagascar were highly similar (98% identity) to that of human CyCV-VN species<sup>7</sup>. The genomic organization was identical with a Rep ORF interrupted by a 166 nt long intron with a typical splice donor (GT) and acceptor site (AG) and Cap ORF of the same length and positions (Figure 1). Rep sequence (292 aa) was  $\approx$ 98% identical to that of CyCV-VN species<sup>7</sup> and showed  $\approx$ 77% and  $\approx$ 73% identity to the

next closely related CyCV species, i.e. Malawian human CSF CyCV (VS5700009) and Tunisian human stool CyCV (TN25), respectively (Table 1)<sup>1,8</sup>. Cap protein of Madagascar CyCV was 222 aa long like typical CyCV-VN and equally conserved ( $\approx$ 98–99% identity when compared to CyCV-VN) but more divergent compared to Malawian VS5700009 ( $\approx$ 35% identity) and Tunisian TN25 ( $\approx$ 48%) CyCV species (Table 2). In contrast, the swine-derived CyCV strains from Cameroon exhibited a less conserved genome (86% identity to CyCV-VN)<sup>7</sup>. Although the genome organization was similar (Figure 1), the Rep gene was shorter (earlier STOP codon, at position 1055–57), leading to a Rep protein of 277 residues instead of 292<sup>7</sup>. As a consequence, Rep and Cap genes did not overlap like in the Madagascar and CyCV-VN strains<sup>7</sup>. In spite of this, the Rep protein sequence of Cameroonian swine CyCV-VN-like strains was highly conserved ( $\approx$ 94% identity compared to Madagascar and



**Table 1** | Sequence identity matrix for the replication-associated protein sequence (%). Cameroonian swine (KM392284–KM392286) and Madagascan human cycloviruses (KM392287–KM392289) described in this study are bolded. Vietnamese VN strains are colored in grey, Malawian VS5700009 in light yellow and Tunisian TN25 and TN18 appear in orange<sup>1,7,8</sup>

	KM392285	KM392284	KM392286	KM392289	KM392288	KM392287
<b>KM392285</b>		100	100	93,5	93,5	93,5
<b>KM392284</b>	100		100	93,5	93,5	93,5
<b>KM392286</b>	100	100		93,5	93,5	93,5
<b>KM392289</b>	93,5	93,5	93,5		100	99,3
<b>KM392288</b>	93,5	93,5	93,5	100		99,3
<b>KM392287</b>	93,5	93,5	93,5	99,3	99,3	
KF031465	93,5	93,5	93,5	97,6	97,6	97,6
KF031467	93,5	93,5	93,5	97,6	97,6	97,6
KF031468	93,5	93,5	93,5	97,6	97,6	97,6
KF031469	93,5	93,5	93,5	97,6	97,6	97,6
NC_021707	93,5	93,5	93,5	97,6	97,6	97,6
KF031470	93,5	93,5	93,5	98,3	98,3	98,3
KF031471	93,5	93,5	93,5	98,6	98,6	98,6
KF031466	93,1	93,1	93,1	97,9	97,9	97,9
KC771281	77,4	77,4	77,4	77,1	77,1	76,3
NC_021568	77,4	77,4	77,4	77,1	77,1	76,3
GQ404857	73,6	73,6	73,6	72,7	72,7	72,7
GQ404858	74,3	74,3	74,3	73,4	73,4	73,1
HQ738635	71,7	71,7	71,7	70,5	70,5	70,5
NC_014927	71,7	71,7	71,7	70,5	70,5	70,5
HQ738634	71,3	71,3	71,3	70,1	70,1	70,1
HQ738636	50,5	50,5	50,5	48,8	48,8	49,1
NC_014928	50,5	50,5	50,5	48,8	48,8	49,1
GQ404844	49,5	49,5	49,5	47,3	47,3	47,7
GQ404846	50,7	50,7	50,7	48,6	48,6	48,9
GQ404848	50,7	50,7	50,7	48,6	48,6	48,9
GQ404849	48,2	48,2	48,2	45,4	45,4	45,8
GQ404850	47,9	47,9	47,9	45,1	45,1	45,4
HQ738644	49,3	49,3	49,3	48,6	48,6	49,3
NC_014930	49,3	49,3	49,3	48,6	48,6	49,3
HQ738643	49,6	49,6	49,6	48,9	48,9	49,6
KC512919	50,7	50,7	50,7	48,8	48,8	48,8
GQ404847	52,7	52,7	52,7	50,2	50,2	50,5
KC512916	50,4	50,4	50,4	48,6	48,6	48,9
HQ638063	50,7	50,7	50,7	50,2	50,2	49,8
HQ638066	50,7	50,7	50,7	49,8	49,8	49,5
HQ638060	50,7	50,7	50,7	50,5	50,5	50,2
GQ404845	52	52	52	49,8	49,8	50,2
HQ738637	51,8	51,8	51,8	51,8	51,8	51,4
NC_014929	51,8	51,8	51,8	51,8	51,8	51,4
GQ404854	51,8	51,8	51,8	50,2	50,2	50,9
JF938081	49,8	49,8	49,8	48,8	48,8	49,1
HM228874	50,7	50,7	50,7	49,8	49,8	49,5
JF938079	50	50	50	48,6	48,6	48,9
JX185426	49,8	49,8	49,8	49,1	49,1	49,5
NC_023866	49,8	49,8	49,8	49,1	49,1	49,5
JX569794	44,8	44,8	44,8	43,7	43,7	44
NC_020206	44,8	44,8	44,8	43,7	43,7	44
GQ404855	52,5	52,5	52,5	50,4	50,4	50,7
KF726986	50	50	50	49,6	49,6	50
KF726984	50	50	50	49,6	49,6	50
NC_023874	50	50	50	49,6	49,6	50
KF726987	50	50	50	49,6	49,6	50
KF726985	50	50	50	49,6	49,6	50
NC_023869	50,7	50,7	50,7	50	50	50
KM017740	40,1	40,1	40,1	39,4	39,4	39,4
JX260426	11	11	11	11	11	10,7

Vietnamese strains). Since CyCVs in the same species are defined as having >85% identity in the Rep protein sequence, these three Cameroonian strains clearly belong to the CyCV-VN group/species (Figure 2). Their Rep sequence identity to Malawian VS5700009 and Tunisian TN25 CyCVs was ≈77% and ≈74%, respectively (Table 1). The Cap gene was 3 nucleotides shorter (deletion of 3 nt after position 1634) and consequently the Cap protein was only 221 residue

long. Cap protein was less conserved compared to that of Madagascan and Vietnamese CyCV strains (≈84–85% identity), but was much closer to CyCV-VN Cap proteins than to that of Malawian VS5700009 and Tunisian TN25 CyCV species (≈34% and ≈48%, respectively) (Table 2). Overall, Madagascan human-derived CyCV strains appeared as typical CyCV-VN strains, highly conserved, while Cameroonian swine strains belonged to the same



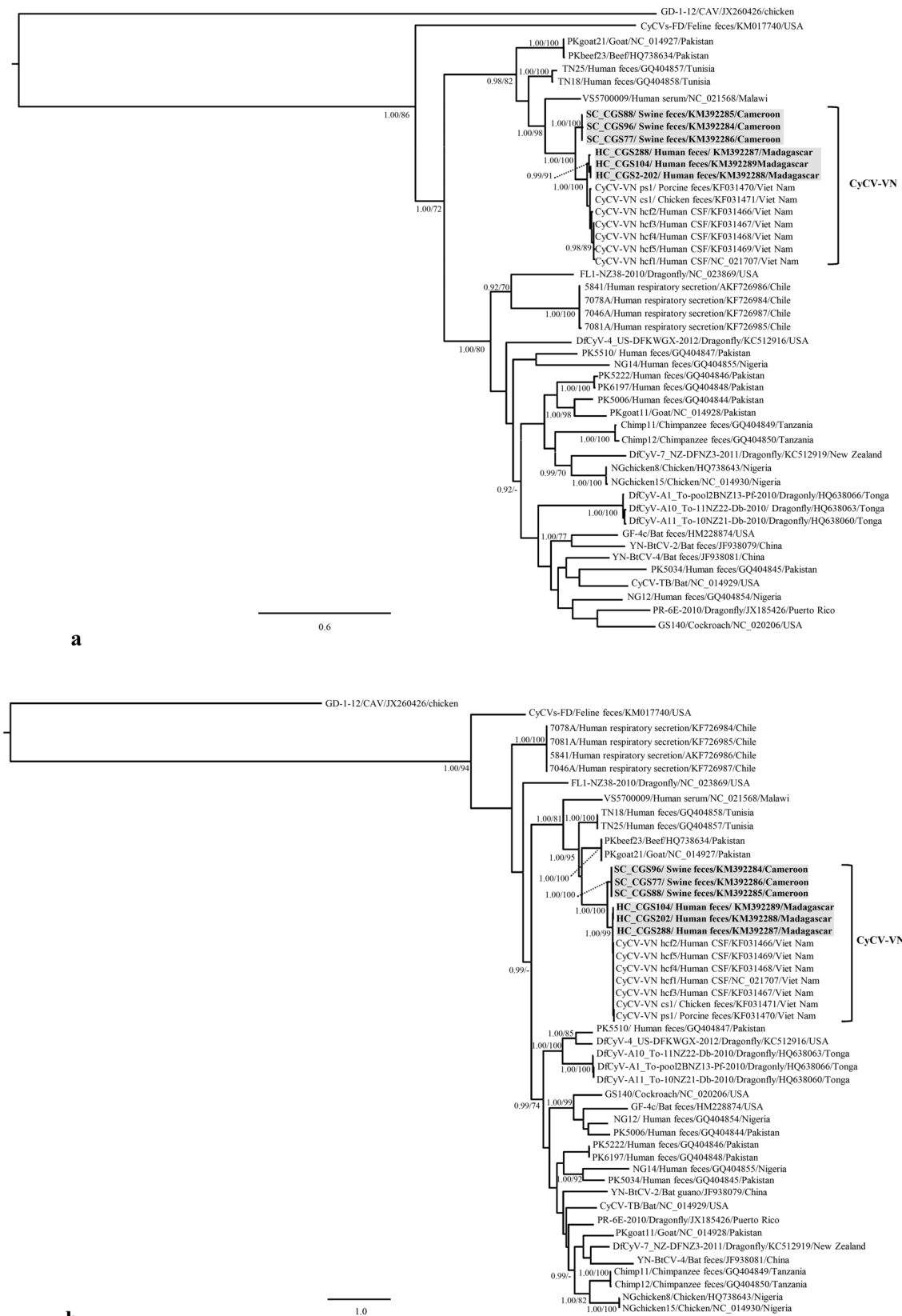
**Table 2 |** Sequence identity matrix for the Capsid protein sequence (%). Cameroonian swine (KM392284–KM392286) and Madagascan human cycloviruses (KM392287–KM392289) described in this study are bolded. Vietnamese VN strains are colored in grey, Malawian VS5700009 in light yellow and Tunisian TN25 and TN18 appear in orange<sup>1,7,8</sup>

	KM392288	KM392289	KM392287	KM392285	KM392286	KM392284
NC_021568	35,3	35,3	35,3	34,1	34,1	34,1
KC771281	35,3	35,3	35,3	34,1	34,1	34,1
KF031470	97,7	97,7	97,3	83,8	83,8	83,8
KF031471	97,7	97,7	97,3	83,8	83,8	83,8
KF031469	98,6	98,6	98,2	84,2	84,2	84,2
KF031468	98,6	98,6	98,2	84,2	84,2	84,2
KF031467	98,6	98,6	98,2	84,2	84,2	84,2
NC_021707	98,6	98,6	98,2	84,2	84,2	84,2
KF031465	98,6	98,6	98,2	84,2	84,2	84,2
KF031466	98,2	98,2	98,6	84,2	84,2	84,2
<b>KM392288</b>		100	98,6	84,7	84,7	84,7
<b>KM392289</b>	100		98,6	84,7	84,7	84,7
<b>KM392287</b>	98,6	98,6		84,2	84,2	84,2
<b>KM392285</b>	84,7	84,7	84,2		100	100
<b>KM392286</b>	84,7	84,7	84,2	100		100
<b>KM392284</b>	84,7	84,7	84,2	100	100	
GQ404857	48	48	47,5	48	48	48
GQ404858	47,5	47,5	47,1	47,5	47,5	47,5
HQ738635	49,3	49,3	49,8	47,9	47,9	47,9
NC_014927	49,3	49,3	49,8	47,9	47,9	47,9
HQ738634	49,3	49,3	49,8	47,9	47,9	47,9
KF726987	22,8	22,8	22,4	23,8	23,8	23,8
KF726986	22,8	22,8	22,4	23,8	23,8	23,8
KF726984	22,8	22,8	22,4	23,8	23,8	23,8
NC_023874	22,8	22,8	22,4	23,8	23,8	23,8
KF726985	22,8	22,8	22,4	23,8	23,8	23,8
HQ638060	24,9	24,9	24,9	24,5	24,5	24,5
HQ638066	24,5	24,5	24,5	24	24	24
HQ638063	25,3	25,3	25,3	25,8	25,8	25,8
KC512916	24,1	24,1	24,6	24,6	24,6	24,6
GQ404847	25,4	25,4	25,4	24,1	24,1	24,1
NC_020206	19,7	19,7	19,2	19,7	19,7	19,7
JX569794	19,7	19,7	19,2	19,7	19,7	19,7
GQ404844	19	19	19	19,5	19,5	19,5
GQ404854	22,2	22,2	22,2	22,2	22,2	22,2
HM228874	17,3	17,3	17,3	16,5	16,5	16,5
GQ404845	18,7	18,7	18,7	19,2	19,2	19,2
GQ404855	17,2	17,2	17,2	16,8	16,8	16,8
HQ738636	22,6	22,6	22,2	23,6	23,6	23,6
NC_014928	22,6	22,6	22,2	23,6	23,6	23,6
KC512919	23,8	23,8	23,4	22,6	22,6	22,6
JF938081	23,2	23,2	23,2	23,3	23,3	23,3
HQ738644	20,4	20,4	20,4	22,6	22,6	22,6
NC_014930	20,4	20,4	20,4	22,6	22,6	22,6
HQ738643	20,4	20,4	20,4	22,6	22,6	22,6
GQ404850	21,3	21,3	21,3	21,3	21,3	21,3
GQ404849	21,3	21,3	21,3	21,3	21,3	21,3
JF938079	23,5	23,5	23	22,6	22,6	22,6
NC_023866	21,7	21,7	20,9	22,1	22,1	22,1
JX185426	21,7	21,7	20,9	22,1	22,1	22,1
HQ738637	20,9	20,9	20,9	20,9	20,9	20,9
NC_014929	20,9	20,9	20,9	20,9	20,9	20,9
GQ404848	23,8	23,8	23,4	22,1	22,1	22,1
GQ404846	23,4	23,4	22,9	21,6	21,6	21,6
NC_023869	24,2	24,2	23,7	24,7	24,7	24,7
KM017740	18,7	18,7	18,7	17,4	17,4	17,4
JX260426	6,6	6,6	6,6	7,5	7,5	7,5

species but were less conserved, especially in the Cap protein sequence.

The typical nonamer sequence of the stem loop structure (TAATACTAT) was conserved in Madagascan human CyCV strain genomes and with one nucleotide substitution (TAATACTAG) in Cameroonian swine-derived CyCVs compared to CyCV-VN strains (Figure 1). The sequence of the loop itself was 11 nt long and

identical in all these strains (Figure 1). Highly conserved circovirus/cyclovirus Rep motifs WWDGY, DDFYGW and DRYP were present in both Madagascan human and Cameroonian swine strains (1). The dNTP binding site (G-GSK) was intact and the motifs associated with rolling-circle amplification (FTLNN, TPHLQG, CSK) were also identified, with the same alterations (WTLNN and TKHLQG) as in CyCV-VN strains<sup>1</sup>.



**Figure 2 |** Bayesian phylogenetic trees based on complete amino acid sequences of the Rep (a.) and Cap (b.) protein of the cycloviruses. Statistical support of grouping from Bayesian posterior probabilities (clade credibilities  $\geq 90\%$ ) and 1,000 parallel Maximum Likelihood bootstrap replicates ( $\geq 70\%$ ) is indicated at the nodes. A Gyrovirus (Chicken anemia virus, GenBank accession number JX260426) was chosen as an outgroup. Taxon information includes the name, source of isolation, GenBank accession number, and country of origin. The cyclovirus strains generated during this study are bolded. The scale bar represents amino acid substitutions per site.



Phylogenetic analysis based on the complete Rep and Cap protein sequences showed that the CyCVs from Cameroon, Madagascar and Vietnam clustered together forming a distinct and highly supported clade/group (CyCV-VN) within the cyclovirus phylogeny (Figure 2). Although, the Cameroonian swine-derived CyCV strains shared a common ancestor with the human-derived CyCVs, they form a distinct lineage within the CyCV-VN clade, suggesting separate evolutionary events.

## Discussion

Cycloviruses are recently described members of the *Circoviridae* family<sup>1</sup>. CyCVs species have been described in mammals, including humans, birds and insects<sup>1–6</sup>. Recently, two studies reported the detection of CyCVs in the cerebrospinal fluid of symptomatic human patients in Vietnam and Malawi<sup>7,8</sup>. An extensive specific screening failed to identify the Vietnam CyCV-VN strain in neighboring Asian countries (Cambodia, Nepal) and in the Netherlands, suggesting a very limited distribution to central and southern Vietnam, where it had been detected in human, pig and chicken<sup>10</sup>.

In order to assess the presence of CyCV-VN on the African continent, we screened a collection of human stool samples from Madagascar and Ghana and of pig stool samples from Cameroon. A total of three human samples and ten swine samples were PCR-positive and confirmed by sequencing. The full genome of the three human CyCV-VN and of three of the swine strains were obtained by primer walking methods.

Phylogenetic analysis showed that both Madagascan and Cameroonian CyCV strains clustered with Vietnam CyCVs forming two distinct sister lineages within CyCV-VN clade and showed distant relationship to previous African cyclovirus strains from Malawi, Tunisia or Nigeria<sup>1,3,7,8</sup>. It is likely that CyCV-VN can be found in other parts of Africa and Asia and possibly in other continents. Our data in pigs suggest a heterogeneous distribution in Cameroon since most positive cases were concentrated in the Douala region. Further, the 1337 human samples from Ghana were tested negative, suggesting an absence of this viral strain from this area. This apparent unequal circulation might be an explanation for the lack of CyCV-VN detection outside Vietnam in previous studies<sup>10</sup>.

Our findings further confirm that pigs might play a central role in the epidemiology of cycloviruses in general and for CyCV-VN species in particular<sup>7</sup>. Besides, the exact pathological significance of CyCV-VN infection in humans is still unknown and further studies are clearly needed to enlighten the transmission route(s), the host tropism and the pathological consequences of these CyCVs.

Overall, the results presented in this study indicate that CyCV-VN and CyCV-like strains circulate in humans and pigs in Africa, far beyond their supposed limited geographic distribution to Vietnam.

## Methods

**Samples collection.** A total of 410 stool samples from healthy children were obtained from Ampasina village, belonging to the township of Andina in the Ambositra region, Madagascar, in 2012<sup>11</sup>. All details including sample preparation are described in ref 11. A total of 1373 human stool samples were retrieved from children up to 15 years of age with diarrhea or vomiting (n = 673) and pediatric controls (n = 664) from a rural area in the Ashanti Region, Ghana. Besides, feces from 520 healthy pigs were sampled in 2012 in the cities of Bamenda (n = 75), Ngaoundere (n = 86), Yaoundé (n = 199) and Douala (n = 160) in Cameroon<sup>12</sup>.

**Ethics Statement.** Experiments were conducted in accordance with the ethical principles set out in the Declaration of Helsinki and informed consent was obtained from all subjects. All experimental protocols were approved by the Ministry of Health of the Republic of Madagascar (no. 015 – MSANP/CE, 22 February 2012) and from the Committee on Human Research, Publications and Ethics, School of Medical Science, Kwame Nkrumah University of Science and Technology, Kumasi, Ghana.

**DNA extraction, CyCV-VN specific PCR and sequencing.** Samples were extracted using Invitex PSP Spin Stool DNA Kit (Stratec Biomedical AG, Birkenfeld, Germany) or QIAamp DNA Stool Mini Kit (Qiagen, Hilden, Germany), according to manufacturer's instructions. PCR screening of CyCV-VN was performed on extracted nucleic acid using the CyCV-VN-specific diagnostic primers designed based on available CyCV-VN sequences in databases (see Supplementary Table S1

online). In order to further characterize the virus strains and to compare them with previously published data, the full-length genome was successfully amplified by primer walking using specific primers (Table S1 online). PCR were performed using HotStarTaq® Plus Master Mix Kit (Qiagen, Hilden, Germany). The PCR was performed in a 25 µl volume containing 3 µl of sample DNA, 10 µl of HotStarTaq Master Mix, 2 µl ml of each primer (20 pmol), and ddH<sub>2</sub>O up to 25 µl. The positive and negative controls (extraction and water controls) were included in every PCR run. Amplification conditions were as follows: denaturation at 95°C for 2 min; 45 amplification cycles were performed at 94°C for 20 s, 55°C for 45 s, 72°C for 1 min. Final extension was at 72°C for 7 min. The fragments were sequenced at least twice in each direction by conventional Sangers technology (LGC, Berlin, Germany).

**Sequence analysis.** Sequence analysis, genomic organization and multiple alignments were performed using Geneious v7.1.7 (Biomatters, Auckland, New Zealand). Secondary DNA structure predictions were performed with the web-based version of mfold (<http://mfold.bioinfo.rpi.edu>). The trees were constructed by using the Bayesian Markov Chain Monte Carlo sampling method in MrBayes v3.2.2<sup>13</sup> and maximum likelihood in PhyML v3.1<sup>14</sup> and included representatives of all cycloviruses species and those six novel human- and swine-derived cycloviruses whose full genome could be determined. The Whelan and Goldman (WAG) amino acid substitution model with gamma-distributed rate variation among sites and a proportion of invariable sites were used.

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## Author contributions

R.M.H., H.F., J.M., A.P., H.J., J.B., N.S., D.D., R.M.C., V.S.P., N.G.S., S.P., H.A.M., A.M.D., M.T.K., N.R., E.T. and R.R. collected the samples. M.G., A.P. and D.C. performed laboratory experiments and analyzed the sequences. J.S.C., M.G. and D.C. drafted the manuscript. All authors were involved in reviewing and editing of the manuscript.

## Additional information

**Nucleotide sequence accession number** The Genbank accession numbers of the cycloviruses described in the study are KM392284–KM392289.

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