Limited geographic distribution of the novel cyclovirus CyCV-VN

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A novel cyclovirus, CyCV-VN, was recently identified in cerebrospinal fluid (CSF) from patients with central nervous system (CNS) infections in central and southern Vietnam. To explore the geographic distribution of this novel virus, more than 600 CSF specimens from patients with suspected CNS infections in northern Vietnam, Cambodia, Nepal and The Netherlands were screened for the presence of CyCV-VN but all were negative. Sequence comparison and phylogenetic analysis between CyCV-VN and another novel cyclovirus recently identified in CSF from Malawian patients indicated that these represent distinct cycloviral species, albeit phylogenetically closely related. The data suggest that CyCV-VN has a limited geographic distribution within southern and central Vietnam. Further research is needed to determine the global distribution and diversity of cycloviruses and importantly their possible association with human disease.

Cycloviruses (CyCVs) belong to the Circoviridae family and have recently been found in different sample types from different hosts, including mammals and insects1–3. Recently, we reported a new cyclovirus species, tentatively named cyclovirus-Vietnam (CyCV-VN), in cerebrospinal fluid (CSF) of two Vietnamese patients4. The virus was subsequently detected in 4% of 642 CSF samples of patients with central nervous system (CNS) infections in northern Vietnam, Cambodia, Nepal and The Netherlands were screened for the presence of CyCV-VN but all were negative. Sequence comparison and phylogenetic analysis between CyCV-VN and another novel cyclovirus recently identified in CSF from Malawian patients indicated that these represent distinct cycloviral species, albeit phylogenetically closely related. The data suggest that CyCV-VN has a limited geographic distribution within southern and central Vietnam. Further research is needed to determine the global distribution and diversity of cycloviruses and importantly their possible association with human disease.

Results

To explore whether CyCV-VN is circulating beyond central and southern Vietnam, we screened a total of 615 CSF specimens from patients with acute CNS infections from northern Vietnam (n = 233), Cambodia (n = 123), Nepal (n = 80) and The Netherlands (n = 179) for the presence of CyCV-VN DNA (Table 1).

CyCV-VN DNA, however, was absent from all 615 CSF specimens tested, suggesting a confined geographical distribution of this virus.

To determine whether CyCV-VN represents a cycloviral species distinct from CyCV-VS5700009, the novel cyclovirus recently detected in CSF from Malawian patients5, we conducted sequence comparisons and phylogenetic analyses to examine the genetic relationship between CyCV-VN and CyCV-VS5700009. Pairwise comparisons showed that the degree of sequence similarity between CyCV-VN and CyCV-VS5700009 was 60% at the nucleotide level of the complete genome sequence and 36% and 76% at the amino acid level of the capsid protein.
Table 1 | Patient cohorts from which samples were used in this study to detect CyCV-VN

| Number of samples | Population | City* | Country   | References | Time frame       | P value* |
|-------------------|------------|-------|-----------|------------|------------------|----------|
| total             | 642        | 26    | southern and central Vietnam | 6          | 1999–2009        |          |
| positive          | 233        | adults | Ha Noi northern Vietnam | 11         | 2007–2008        | 0.0004   |
|                   | 123        | children | Siem Reap Cambodia | 12         | 2009–2010        | 0.01     |
|                   | 80         | all ages | Kathmandu Nepal | 13         | 2009–2011        | 0.1      |
|                   | 179        | all ages | nationwide | The Netherlands | unpublished | 2010–2011 | 0.003 |
| All               | 615        | 0     |           |            |                  | <0.0001  |

*Name of the city indicates the location of the hospital where the patients were admitted.
#Fischer exact; compared with results from the study in reference #6.

Figure 1 | Reconstructed phylogeny tree of complete genome sequences of cycloviruses, CyCV-VN (red) and CyCV-VS5700009 (blue); circoviruses and a gyrovirus were used as outliers. CV: circovirus, Sequence accession number are in square brackets.
and replication association proteins, respectively. These values are all well below the proposed demarcation criteria to distinguish different CyCV species (75%, 70% and 85% identity, respectively)\(^1\). The genetic distance between the two viruses was confirmed by phylogenetic analysis based on complete genome sequences (Figure 1). These analyses indicate that CyCV-VN and CyCV-VS5700009 represent two distinct cycloviral species.

**Discussion**

In a previous study we showed that the newly discovered CyCV-VN was widely distributed in central and southern Vietnamese provinces\(^6\). In this study we further explored the geographic distribution of CyCV-VN by testing 615 CSF samples from patients with suspected CNS infections from northern Vietnam, Cambodia, Nepal and the Netherlands. CyCV-VN was absent from all 615 tested CSF samples from these locations, suggesting that the geographic distribution of CyCV-VN may be limited to southern and central Vietnam.

We previously reported a prevalence of 4% (28/644) among children and adults with confirmed or suspected CNS infections in overlapping time periods with the current study\(^6\), hence failure to detect CyCV-VN in this study is unlikely due to sample size restrictions or temporal bias. This is supported by observed statistically significant differences in prevalence (Table 1). However, it cannot be ruled out that related cycloviruses (including CyCV-VSS700009) may have gone undetected since our PCR method was designed for specific detection of CyCV-VN\(^6\). Indeed, sequence analysis indicates that the differences at primer and probe binding sites would render detection of the Malawian virus and other reported cycloviruses by CyCV-VN-specific PCR method highly unlikely (Figure 2\(^5\)). Testing for other (un)known cycloviruses in these 615 CSF samples is, therefore, desirable, but beyond the scope of the present study. Likewise, to fully explore the circulation of CyCV-VN in the regions from where 615 patients came it is necessary to test for the CyCV-VN in livestock species (e.g. pigs, chickens and ducks) that have been previously tested positive for the virus in more than 50%\(^6\).

In summary, our new data from clinical case detections suggest that circulation of CyCV-VN is restricted to southern and central Vietnam. Further research utilizing degenerate PCR primers is needed to examine to what extent diverse cycloviruses can be detected in CSF of patients with CNS infections worldwide. In addition, further investigations of cyclovirus prevalence in healthy subjects, clinical cases, and non-human species will be required to assess the epidemiological significance of these findings.

**Methods**

**Clinical materials.** CSF specimens analyzed in this study were derived from three prospective clinical studies designed to establish the etiology of CNS infections in northern Vietnam (n = 233), Cambodia (n = 123) and Nepal (n = 80). The Dutch CSF specimens (n = 179) represented a random sample of anonymized specimens from Dutch patients with suspected CNS infections submitted to the Netherlands Reference Laboratory for Bacterial Meningitis in Amsterdam.

**Nucleic acid extraction.** As part of standard operating procedures applied in our laboratory, CSF samples were spiked with an optical amount of phocid herpesvirus and equine arteritis virus as (reverse transcription) PCR internal controls\(^5\), and were then subjected to an automatic extraction procedure with use of the Viral NA Small Volume Kit (Roche applied sciences, UK) and a MagNA Pure 96 system (Roche applied sciences). The isolated nucleic acid were then stored at −80°C for subsequent PCR screening for CNS pathogens as well as CyCV-VN.

**Real time PCR.** PCR screening of CyCV-VN was performed on extracted nucleic acid samples that had been tested by PCRs for CNS pathogens and had expected crossing point values of internal controls between 33 and 35, and was carried out as previously described\(^5\). In brief, the PCR was done in a final 25 µl volume reaction containing 1.25 U of HotStarTaq DNA polymerase (QIAgen GmbH, Hilden, Germany) 800 µM of each primer, 100 µM of a FAM-labeled probe, 400 µM each dTTPs 5 mM of MgCl\(_2\) (provided with the polymerase), 1 × PCR buffer (provided with the polymerase), and 3 µl of extracted NAs. PCR reaction was performed on an LightCycler 480 Instrument II (Roche applied Sciences, UK) with an initial polymerase-activation step at 95°C for 14 min 30 sec followed by 45 cycles of 95°C for 30 sec, 55°C for 30 sec and 72°C for 30 sec. A positive control and several negative controls (including extraction controls and water controls) were included in every PCR run. All the PCR experiments were performed in molecular diagnostic facilities that consist of three physically separated laboratories for reagent preparation, extraction and amplification, and these were used a unidirectional workflow.

**Sequence analysis.** Pairwise comparisons and phylogenetic analyses were done using AligX (Vector NTI Advance 7; Invitrogen, Carlsbad (CA, USA)) and neighbor-joining in MEGA version 4\(^6\).

**Ethical statement.** All clinical studies from which the CSF samples were chosen for PCR screening in the present study were reviewed and approved by the Institutional Review Boards of collaborating hospitals and the Oxford Tropical Research Ethics Committee (OxTREC), University of Oxford, United Kingdom. Informed consent was obtained from patients or a parent or guardian of each enrolled patients.

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**Author contributions**
H.D.T.N., N.V.V.C., N.V.K., W.T., H.F.L.W., A.v.d.E., A.G., S.S. and T.T.H. collected CSF specimens. M.C., N.V.T., N.H.U., M.C., J.B. and N.T.T.C.B. conducted laboratory experiments. L.V.T. did sequence analyses. L.V.T., H.R.v.D., J.F., L.v.d.H., J.B. and M.D.d.J. designed the study. L.V.T., H.R.v.D. and M.D.d.J. drafted the manuscript. All authors were involved in subsequent reviewing and editing the manuscript.

**Additional information**
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