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Novel Human Herpesvirus 8 Subtype D Strains in Vanuatu, Melanesia

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We show human herpesvirus 8 with diverse molecular subtype D variants to be highly endemic among the Ni-Vanuatu population. Most K1 genes were nearly identical to Polynesian strains, although a few clustered with Australian or Taiwanese strains. These results suggest diverse origins of the Ni-Vanuatu population and raise questions about the ancient human population movements in Melanesia.

Human herpesvirus 8 (HHV-8), or Kaposi Sarcoma-Associated Herpesvirus (KSHV), is the etiologic agent of Kaposi sarcoma (KS). HHV-8 is not a widespread ubiquitous virus; its presence is mainly restricted to areas where classic or endemic KS is highly prevalent, i.e., estimates of HHV-8 seroprevalence in the general adult population range from 5% to >50% (1).

Exploiting the highly genetic variability of the HHV-8 K1 gene, molecular epidemiology led to the identification of 5 major K1 subtypes (A–E), some of which appear to be strongly linked to the geographic origin of the samples. Thus, the few known subtype D strains have been reported only in inhabitants from the Western Pacific region (2).

For people of Oceanian ancestry (including Melanesian, Polynesian, and Micronesian), very little data are available on the clinical and molecular epidemiology of HHV-8 and its associated diseases (3–9). Thus, we studied HHV-8 in the Vanuatu, an archipelago in the Southwest Pacific region, formerly named New Hebrides, which contains >80 islands (6 provinces) (Figure 1). Indigenous Melanesians, also called Ni-Vanuatu, constitute 98% of the current population of ≈210,000. A recent study suggested that HHV-8 was rare in the Ni-Vanuatu population (10).

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Figure 1. A) Age-dependent herpesvirus 8 (HHV-8) seroprevalence rates in 376 Ni-Vanuatu persons >65 years and living in 18 islands representative of the 6 provinces of the Vanuatu Archipelago. Seropositivity was based on strict criteria, and only samples clearly reactive at a dilution >1:160 were considered HHV-8 positive. B) Age-dependent HHV-8 seroprevalence rate in 283 Ni-Vanuatu persons from 13 families originating from 4 islands (3 from Loh, 2 from Tanna, 4 from Ambae, and 4 from Espiritu Santo) of the Vanuatu archipelago. Error bars indicate 95% confidence intervals. C) Pedigrees of 2 families from Loh Island in which the presence of HHV-8 was examined in members of 3 generations. Gray circles and squares denote infected women and men, respectively. Black circles denote infected women for whom sequence of K1 gene fragment was obtained. Numbers within circles and squares indicate ages of the patients; NT, not tested.
Our goal for this cross-sectional study was to evaluate the prevalence of HHV-8 in the Vanuatu archipelago by using stringent serologic criteria and to characterize its genetic diversity.

The Study

Our work was performed on a large collection of ≈4,500 plasma and peripheral blood buffy coat (PBBC) samples from different islands of the archipelago, obtained in the framework of our previous studies on human T-cell lymphotropic virus (HTLV-1) (11,12). The field survey, carried out from April 2003 through August 2005, has been extensively described (11).

To detect plasma HHV-8 antibodies, an inhouse immunofluorescence assay (IFA) using BC-3 cells expressing only latent-associated nuclear antigens encoded by ORF73, was performed to detect plasma HHV-8 antibodies (13). Because HHV-8 seroprevalence increases with age in a virus-endemic population, we first tested a series of 376 samples, from persons >65 years of age (mean 72, median 70, range 65–96 years; 182 men and 194 women) originating from the 6 provinces of the archipelago (online Appendix Figure, available from www.cdc.gov/EID/content/13/11/1745-appG.htm). Among these 376 plasma samples, 170 (45.2%) were IFA positive at a 1:160 dilution, showing a clear typical nuclear spotted seroreactivity. The HHV-8 seroprevalence was similar between men (45.6%) and women (44.8%). The prevalence of HHV-8 increased with age, rising from 29.6% (65–69 years) to 57.1% (>80 years) (Figure 1, panel A) (p = 0.0005 trend χ² test). This high level of HHV-8 seroprevalence was present in all 6 provinces (online Appendix Figure).

A second serologic survey that used 237 plasma samples taken from 13 families with genealogic trees was performed (Figure 1, panels B and C). Among these 237 samples, 12 originated from elderly persons included in the 376 samples tested before. The HHV-8 seroprevalence was clearly age-dependent, rising from 6.9% among children 1–9 years of age to 28.2% in adults 50 years of age, followed by a new increase in persons ≥51 years of age (55.9%) (p<10⁻⁴ trend χ² test). These results demonstrate that HHV-8 infection is endemic, widespread, and circulates in the Ni-Vanuatu population.

We then characterized these HHV-8 strains molecularly. All DNA samples (1 μg), extracted from the PBBC, were first amplified by PCR for human β-globin sequences to control amplifiability. HHV-8 infection was determined by a nested PCR to obtain a 737-bp fragment of the open reading frame of the K1 gene (ORFK1). The first PCR was performed with the primer set K1AG75S/K1AG1200AS.
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(14) and followed by a nested PCR with a second set of primers VR1S/VR2AS1 (15). All PCR products were purified from gel, cloned, and sequenced. Sequences were verified on both DNA strands. ORFK1 amplification was obtained from 32 (21.6%) of the 148 HHV-8–seropositive samples tested but in none of the 26 HHV-8–seronegative samples. Sequences were obtained for only 30 of the 32 ORFK1-positive PCRs (Table).

Comparative sequence analysis indicates that the 30 new sequences differed from each other. Furthermore, among them, 3 groups can be clearly identified. The first group comprises most strains (23/30) and corresponds to the sequences found in persons from the south central islands of the archipelago (Mallicolo, Ambrym, Epi, Tongoa, Emae, and Tanna); the second group comprises 4 sequences from persons living in the northern islands of Loh (LO13, LO17, LO48) and Santo (CESW32); and the third group involves only 3 sequences (ML10, ML36, ML46) from persons living in the northern island of Motalava.

Phylogenetic analyses were performed on the 30 novel sequences obtained in this study, on all subtype D available K1 sequences, and on representatives of the different HHV-8 subtypes/subgroups, as described (4). The phylogenetic analyses were performed with all of the sequences available. These sequences include 3 strains from Japan (J24, J25, and J26), 1 from Australia (3Au1), 1 from Taiwan (TKS10), 1 from New Zealand (ZKS3), and 1 from Wallis (WalKS1) (2, 4, 6–8). Our results demonstrate that the Ni-Vanuatu HHV-8 clustered in 3 different genotype D subclades, which are highly supported phylogenetically with high bootstrap values of 99% or 100% (Figure 2). The first one comprising most strains corresponds to sequences closely related to each other and to the 2 Polynesian strains WalKS1 and ZKS3 (4, 8). The second group (Loh/Santo), with only 4 sequences, was closely related to the Taiwanese strain TKS10 (8). The last 3 sequences from Motalava were nearly identical to the only strain from Australia (6).

### Table. Demographic, geographic, and serologic data of HHV-8 seropositive persons from the Vanuatu Archipelago, confirmed by molecular analysis

| Virus strain | Age, y | Sex | Island of origin | Province | IFA titers (LANA) | PCR K1 | GenBank accession no. |
|--------------|--------|-----|-----------------|----------|------------------|--------|----------------------|
| LO 17        | 70     | F   | Loh             | TORBA    | 10.240           | +      | EF589758             |
| LO 13        | 23     | F   | Loh             | TORBA    | 160              | +      | EF589757             |
| LO 48        | 64     | F   | Loh             | TORBA    | 640              | +      | EF589759             |
| ML 10        | 60     | F   | Mota Lava       | TORBA    | 640              | +      | EF589763             |
| ML 36        | 60     | M   | Mota Lava       | TORBA    | 640              | +      | EF589764             |
| ML 46        | 61     | M   | Mota Lava       | TORBA    | 1.280            | +      | EF589765             |
| ES 65        | 82     | F   | Santo           | SANMA    | 160              | +      | EF589756             |
| CES W32      | 60     | F   | Santo           | SANMA    | 320              | +      | EF589746             |
| MA 55        | 60     | M   | Mallicolo       | MALAMPA  | 2,560            | +      | EF589760             |
| MAL 4        | 74     | M   | Mallicolo       | MALAMPA  | 320              | +      | EF589761             |
| MAL 24       | 70     | M   | Mallicolo       | MALAMPA  | 1,280            | +      | EF589762             |
| RYM 31       | 80     | F   | Ambrym          | MALAMPA  | 40               | +      | EF589766             |
| RYM 42       | 74     | F   | Ambrym          | MALAMPA  | 80               | +      | EF589767             |
| EP 43        | 80     | F   | Epi             | SHEFA    | 20.480           | +      | EF589753             |
| EP 58        | 79     | F   | Epi             | SHEFA    | 20.480           | +      | EF589754             |
| EP 111       | 74     | F   | Epi             | SHEFA    | 1.280            | +      | EF589755             |
| TON 72       | 75     | M   | Tongoa          | SHEFA    | 5.120            | +      | EF589772             |
| CTON H54     | 71     | M   | Tongoa          | SHEFA    | 2.560            | +      | EF589747             |
| EM 1         | 70     | M   | Emae            | SHEFA    | 640              | +†     | NA                   |
| EM 4         | 75     | F   | Emae            | SHEFA    | 2.560            | +      | EF589752             |
| CEM H2       | 73     | M   | Emae            | SHEFA    | 1.280            | +      | EF589744             |
| EF 43        | 73     | F   | Efate           | SHEFA    | 640              | +      | EF589748             |
| EF 52        | 74     | M   | Efate           | SHEFA    | 1.280            | +      | EF589749             |
| EF 55        | 81     | M   | Efate           | SHEFA    | 640              | +      | EF589750             |
| EF 56        | 75     | F   | Efate           | SHEFA    | 5.120            | +†     | NA                   |
| EF 60        | 73     | F   | Efate           | SHEFA    | 640              | +      | EF589751             |
| CEF H56      | 72     | M   | Efate           | SHEFA    | 80               | +      | EF589742             |
| CEF H67      | 70     | M   | Efate           | SHEFA    | 640              | +      | EF589743             |
| TA 115       | 77     | M   | Tanna           | TAFAEA   | 1.280            | +      | EF589768             |
| TA 116       | 70     | F   | Tanna           | TAFAEA   | 320              | +      | EF589769             |
| TA 124       | 70     | F   | Tanna           | TAFAEA   | 5.120            | +      | EF589770             |
| TA 162       | 70     | M   | Tanna           | TAFAEA   | 5.120            | +      | EF589771             |

*IFA, immunofluorescence assay; LANA, HHV-8 specific antibody directed against latent nuclear antigen; PCR K1, PCR for amplification of a 737-bp fragment of the ORFK1 genomic region of HHV-8; NA, not available.
†Weak PCR signal.
Furthermore, phylogenetic analysis showed a star-like tree with a long branch for the Polynesian clade, which includes most Ni-Vanuatu strains. This finding strongly suggests a common origin or ancestor for these strains, with a possible founder effect (Figure 2). Based on stringent serologic and molecular analyses, our study demonstrates for the first time, to our knowledge, that HHV-8 infection is endemic in a Melanesian population.

Our serologic findings are consistent with those found in some remote villages of Papua New Guinea (5, 9). On the basis of these studies, it is tempting to suggest that intrafamilial HHV-8 transmission occurs in such populations, as previously demonstrated in highly HHV-8 endemic populations of African origin (13).

From the molecular point of view, finding such a high molecular diversity of HHV-8 subtype D with some Polynesian-, Taiwanese- and Australian-like strains was surprising. These heterogeneous findings contrast with the more homogenous situation found for HTLV-1 genotypes in the same population (11).

Conclusions

Our molecular findings suggest that HHV-8 has been introduced in the Ni-Vanuatu populations by different migrations of infected persons. This conclusion is strengthened by the clustering of the Australian- and Taiwanese-like strains in the northern islands of Loh and Motalava. A variety of scenarios have been proposed to explain the peopling of near and remote Oceania, and our data highlight the possible multiple origins of Ni-Vanuatu ancestors. Ongoing molecular studies on both viral and mitochondrial/nuclear DNA will contribute to this debate through analyses of the variations observed. Indeed, these variations are intimately linked with the dispersal of early human settlers; analyses of the genetic variability of HHV-8 can help us reconstruct the patterns of human dispersal into Oceania (11).

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Mr Cassar is a PhD candidate with primary research interest in the clinical and molecular epidemiology and physiopathology of dengue viruses. He is working on the epidemiology of HTLV-1 and HHV-8 in Melanesian populations.

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