Vironome of Kaposi sarcoma associated herpesvirus-inflammatory cytokine syndrome in an AIDS patient reveals co-infection of human herpesvirus 8 and human herpesvirus 6A

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ABSTRACT

KSHV inflammatory cytokine syndrome (KICS) is a newly described condition characterized by systemic illness as a result of systemic, lytic KSHV infection. We used Illumina sequencing to establish the DNA vironome of blood from such a patient. It identified concurrent high-level viremia of human herpesvirus (HHV) 8 and 6a. The HHV8 plasma viral load was 5,300,000 copies/ml, which is the highest reported to date; this despite less than five skin lesions and no HHV8 associated lymphoma. This is the first report of systemic HHV6a/KSHV co-infection in a patient. It is the first whole genome KSHV sequence to be determined directly from patient plasma rather than cultured or biopsied tumor material. This case supports KICS as a new clinical entity associated with KSHV.

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Introduction

Kaposi sarcoma (KS) and the plasmablastic variant of multicentric Castleman disease (MCD) are associated with Kaposi sarcoma-associated herpesvirus (KSHV) infection and occur with increased prevalence in AIDS patients. KS remains among the leading causes of death for people living with HIV/AIDS today. It is the most frequent cancer overall in many sub Saharan countries (e.g. Uganda) (Franceschi et al., 2008). Since skin KS lesions are relatively easy to diagnose clinically, KSHV viral load assays are seldom conducted. KSHV gene transcription in KS is variable, but predominantly restricted to the latent viral genes (Dittmer, 2003, 2011). Compared to diseases caused by other herpesviruses (HSV-1, HCMV, EBV) systemic viremia in KS is an order of magnitude less (Lin et al., 2009). As a consequence, infectious virus has not been efficiently recovered from patients, perhaps due to comparatively low systemic viremia vis-à-vis other herpesviruses. There exists a gap in our knowledge as all known sequence information for KSHV was derived from cell line propagated or tumor derived clones, but not from free virus systemically replicating in the natural host.

MCD is a hyperproliferative B cell disorder. MCD patients express high levels of viral IL6 (vIL6) and human IL6, which are thought to be responsible for the systemic illness and inflammatory symptoms (Brandt et al., 1990; Oksenhendler et al., 2000). Clinical symptoms of MCD include edema, weight loss, pleural effusion (mostly cell free, as opposed to PEL), anemia, thrombocytopenia, lymphadenopathy, and splenomegaly (reviewed in
Polizzotto et al., 2012). An alternative clinical diagnosis is that of
systemic inflammatory response syndrome (SIRS), which can
cause rapid death in AIDS patients. It is thought of as an extreme,
uncontrolled, cytokine-mediated response to infection in CD4
deficient patients. The underlying trigger of this response is
typically unknown. Thus, symptomatic treatment remains the
only option.

KSHV inflammatory cytokine syndrome (KICS) has been
recently been proposed as a new KSHV associated disease
(Uldrick et al., 2010). The six patients originally described had
no evidence of MCD despite thorough examination, and no
development of MCD upon follow up. However, they presented
with clinical inflammatory syndromes as seen in MCD, high
human IL6 levels, and high systemic KSHV viral load. The patient
described here had the highest systemic KSHV viral load on
record to date, with no identifiable MCD. He fits the novel clinical
description for KICS and we were able to establish the complete
DNA vironome from blood.

Systemic KSHV–HHV6a co-infection was uncovered by next
generation sequencing of peripheral blood mononuclear cells and
plasma. We obtained a complete KSHV genome sequence from
high titer, replicating virus rather than tumor associated or tissue
culture propagated variants (Depledge et al., 2011; Lin et al.,
2012), which in the case of KSHV tend to replicate extremely
poorly and may have accumulated mutations.

Results

The patient is a 42-year old Caucasian male who was diag-
nosed with human immunodeficiency virus (HIV) infection in
1993 and treated with zidovudine/lamivudine and nevirapine. He
discontinued medication in 2003, when his HIV-1 viral
load was 600,000 copies/ml and a CD4 count of 27 cells/
undetectable. He presented in December 2008 with a HIV-1 viral
load of 600,000 copies/ml and a CD4 count of 27 cells/
undetectable. He was diagnosed with Castleman disease and treated
with rituximab and tocilizumab, but the disease progressed and
he developed Kaposin, and orf69 (Fig. 1, panel E). We chose multiple genes to
increase the specificity of our assay. These primers were pre-
viously validated on clinical and experimental samples (Dittmer,
2011, 2003). As a positive control for DNA isolation from clinical
specimens we used gapdh primers. The positive control was total
DNA from BCBL-1 cells and the negative control sample was
water. All primers amplified the predicted size fragment from the
patient sample and positive control, but not the negative control
(the very faint bands in lane 1 and 3 represent primer dimers,
which were smaller than the predicted size). No band was
observed in the ‘no primer control’ lane (Fig. 1, panel E, lane “-”). As this assay is a conventional endpoint PCR assay, it is not
quantitative. The brightness of the individual bands is dependent
on their size as well as accumulation. The three longest PCR
products correspond to KSHV genes LANA, vGPCR and gapdh
(Fig. 1, panel E, lanes 1, 4, “+”). The four very short PCR products
yielded shorter and thus less bright products (Fig. 1, panel E, lanes
2, 3, 5, 6). This demonstrates that KSHV was present in the plasma
of this patient.

To detect free virus, we determined the plasma viral load of
KSHV. The patient had a plasma KSHV viral load of 5,300,000
copies/ml (Viracor-IBT Laboratories, MO). This represents the
highest KSHV viral load on record (Table 1). By comparison KS
patients (Polstra et al., 2003; Sayer et al., 2011) typically have a
mean KSHV viral load of 1000 copies/ml (range 151–26,915), PEL
patients (Simonelli et al., 2009) of 10,284 copies/ml (range 2,558–
36,300) and MCD (Simonelli et al., 2009) of 4,400 copies/ml
(range 600–1,678,000). KS malignancies are AIDS defining, but
rarely associated with high-level virus replication. Average KSHV
plasma viral loads are 1–2 log10 units lower than EBV or HCMV
viral loads even in symptomatic patients (Laney et al., 2007;
Whitby et al., 1995). Hence, for this exceptional case we con-
sidered that the patient succumbed to acute KSHV viremia.

We turned to Illumina-based sequencing to determine the
complete vironome and complete genome sequence of this KSHV
isolate. Total DNA was isolated from white blood cells. No
attempts were made to enrich for viral sequences. Illumina™
sequencing was performed by sonication of DNA, primer ligation
and tethered amplification, followed by DNA synthesis to yield
> 35 base reads. To determine the vironome, sequencing reads
were aligned to the reference genomes of all known viruses.
Combining five individual sequencing runs we obtained a total of
145,179,479 reads of 36 bp (paired), 50 bp (single) and 75 bp
(paired and single) from samples of this patient. A discovery score
(D score) based on the total number of aligned reads and coverage
for each viral genome was computed and subjected to hierarch-
cal clustering. Sequencing reads from samples of known herpes-
virus genome status (two copy EBV – integrated cell line
Nalmalwa, PEL cell line BCBL-1) were included to calibrate the
virus discovery algorithm. In four of the sequencing runs for this
patient, purified Guinea Pig CMV DNA was spiked in as an internal
control, and was recovered in sequence alignment for these
samples (Genbank ID: AB592928).

The clustering results are displayed as heatmap and dendro-
grams (Fig. 2, panel A). Positivity is displayed as color (white,
orange, red), with deeper color indicating a higher D score.
D score is a weighted combination of coverage, genome-size and number of hits. For the 24 human chromosomes we calculated at 95% CI of \( \text{95CI} \) of \( 0.03 \) to \( 0.07 \log_{10} \) copies/cell, i.e. two copies per cell.

Our initial screen identified sequences for 648 genbank entries, which were classified as either fungal, viral, protozoal or bacterial. However, most were present at \( \text{r} / C_{0} = 2 \log_{10} \) genome copies/cell, i.e. fewer than 1 copy per cell. Several bacterial pathogens, including toxoplasma, were identified for but could not be confirmed by standard tests. Mitochondrial genomes were present at \( 1.52 \log_{10} \) copies/cell, and KSHV was present at \( 0.23 \log_{10} \) copies/cell, i.e. at \( > 1 \) copy per cell.

In total, 22,455 reads aligned to KSHV, resulting in a 9.23 fold median genome coverage (Fig. 2, panel B). Since coverage is a function of copy number and genome size this demonstrates for the first time HHV8 and HHV6a co-replication in a patient. Both KSHV and HHV6a DNA sequences were found in roughly equal proportion in cell-free plasma, where the KSHV plasma viral load was \( > 10^{6} \). This would argue that the HHV6a signal, too, stemmed from replicating virus rather than being horizontally transmitted.

We established uniformity of genome coverage as a second criterion for virus identification. All reads mapped randomly across the entire length of the HHV8 and HHV6a genomes (Fig. 2, panel B and C). Reads that aligned to more than one position, i.e. to repeat region, were randomly placed at only a single position. In contrast to HHV6a, the few reads that mapped to the HHV6b genome, all clustered within the terminal repeat region (data not shown), indicating insufficient coverage to demonstrate biologically relevant infection.

We derived the whole genome sequence for this KSHV strain by reference-guided assembly (Genbank ID: JQ619843). The KSHV consensus sequence was compared to the KSHV reference sequence (Genbank ID: NC_009333). We did not identify novel DNA insertions or orfs. We identified small InDels in specific genes (the polymorphic regions of K1 and K15) and in repeat regions, which is a result of the short read length of the Illumina platform, but not deletions of entire coding regions or gross aberrations. Thus no one gene could be linked to the highly replicative phenotype. These are annotated in the sequence submission.

To demonstrate relatedness of this primary KSHV isolate to tumor cell line derived viral sequences, we used ClustalW (Felsenstein, 1989) using bootstrap analysis with Neighbor-Joining. The pairwise distances between all the aligned whole genome sequences were computed using a maximum likelihood model, the number of replications for bootstrap analysis was 1000. The result is displayed using PHYLIP and demonstrated high sequence conservation expected of double-stranded DNA viruses than HHV6a, which is rarely found in patients (Boutolleau et al., 2006).

A total of 23,300 reads mapped to HHV6a, yielding 7.65 fold median coverage (Fig. 2, panel C). Since coverage is a function of copy number and genome size this demonstrates for the first time HHV8 and HHV6a co-replication in a patient. Both KSHV and HHV6a DNA sequences were found in roughly equal proportion in cell-free plasma, where the KSHV plasma viral load was \( > 10^{6} \). This would argue that the HHV6a signal, too, stemmed from replicating virus rather than being horizontally transmitted.

We estimated uniformity of genome coverage as a second criterion for virus identification. All reads mapped randomly across the entire length of the HHV8 and HHV6a genomes (Fig. 2, panel B and C). Reads that aligned to more than one position, i.e. to repeat region, were randomly placed at only a single position. In contrast to HHV6a, the few reads that mapped to the HHV6b genome, all clustered within the terminal repeat region (data not shown), indicating insufficient coverage to demonstrate biologically relevant infection.

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It is difficult to derive genome-wide conclusions from a comparison of just six genomes. We did note extraordinary conservation in primary nucleotide and amino acid sequences among all isolates. For the unique regions of the genome, approximately one nucleotide change was present per kb and less than one amino acid per coding regions, as expected for DNA viruses. The most divergent sequence was the original BC-1, which was obtained by Saenger sequencing; and which may not have had the extensive genome coverage afforded by NextGen based sequencing. The Sau3AKS sequence has deletions in K15 compared to all other sequences, which may be the result of phage library prep and propagation in *Escherichia coli*. We observed variations and gaps in the repeat regions (LIR1, DR1-6, and TR). These could be the result of misalignment, as short read-based protocols are known to generate incomplete alignments in the repeat regions, or they could be the result of genome evolution. For instance, a 36 bp gap in K1 was conserved in this sequence, GK18, JSC1 and Sau3A; a 9 bp deletion in DR3 was conserved in JSC1, BCBL1 and Sau3A. We did not have enough coverage for a de-novo assembly to unambiguously identify small DIPs or SNPs for this particular isolate.

Rather than directly testing for each potential pathogen sequentially, NextGen sequencing technology allowed us to detect a broad range of viruses, bacteria, fungi, and parasites, resulting in the detection of additional factors that may be playing a role in the disease, in this case HHV6a.

**Discussion**

This report highlights NextGen sequencing technology on patient material to identify infectious agents in a case of a fatal inflammation reaction of unknown origin. Rather than testing for each potential pathogen individually, sequencing of total DNA in...
plasma and PBMC allowed us to screen for a broad range of viruses, as well as bacteria, fungi, and parasites (data not shown), thus resulting in the detection of additional factors that may be playing a role in patient illness, in this case, HHV6a viremia. This novel application of Illumina sequencing to assist in patient diagnosis proved to be a useful tool in determining the cause of illness and death in this patient.

This represents the first documentation of systemic KSHV/HHV6 co-replication in a patient. Primary HHV6 infection is mostly subclinical and HHV6 has near 100% sero-prevalence in the US. HHV6a is primarily associated with exanthema subitum (Roseola). It has been detected in end-stage AIDS patients, but these instances are extremely rare. HHV6a DNA has been detected in a single archival case of a PEL biopsy before (Asou et al., 2000) but not as free, replicating virus. HHV6a is also found in forms of reactive lymphoid hyperplasia, such as Crohn’s disease, histiocytic necrotizing lymphadenitis or Kikuchi–Fujimoto disease in high prevalence areas, although the present studies are insufficient to support a conclusive association of HHV6a with these or other forms of rare reactive lymphoid hyperplasia (Bouvard et al., 2009). HHV6a can cause KSHV reactivation from PEL cell lines in culture, as can HCMV (Lu et al., 2005; Vieira et al., 2001).

Whether, in turn, KSHV can stimulate HHV6a replication and/or its pathogenesis is presently unknown. Both viruses encode as well as induce cytokines and chemokines, including human IL6 and vIL6, which can induce inflammatory symptoms. We surmise that systemic replication of both viruses, rather than KSHV-associated neoplasia, led to the clinical symptoms. The clinical description of SIRS as a cytokine storm is consistent with that of KICS, which was recently proposed as a new KSHV associated clinical entity and which is defined by high cytokine production, particularly of IL6 and the presence of KSHV, but a negative diagnosis for MCD. We propose that coinfection may have added to the intensity of KICS. Further studies are necessary to determine whether HHV6a coinfection is present in other KICS patients, or if it is necessary for the development of KICS.

Conclusions

We have described a fatal case of systemic KSHV and HHV6a co-viremia in an AIDS KICS patient. Using Next-Gen sequencing, we were able to generate the first KSHV sequence directly from patient sample, without implementing cloning techniques. This novel sequence has been deposited into Genbank.

Materials and methods

DNA extraction

Genomic DNA was isolated from white blood cells or plasma using Wizard SV™ kit (Promega Inc.). All PCR products were the expected size by gel electrophoresis. No amplification was observed without DNA extraction.

Illumina sequencing

Illumina™ sequencing was performed by sonication of DNA, primer ligation and tethered amplification, followed by DNA synthesis to yield 35 base reads. The high viral load allowed us to use total DNA without virus-specific enrichment (e.g. Kwok et al., 2012). To prepare the library, DNA was sonicated, then ends were repaired and adenylated. Paired end adapters were ligated to DNA fragments and purified via gel electrophoresis excision and QIAGEN gel extraction kit (QIAGEN, part #28704). Ligated products were amplified using PCR prior to sequencing run. Sequencing was performed using a variety of conditions: paired end 2 \times 76 cycles. Additional runs include 50 bp single reads, 36 bp paired reads, and 76 bp single reads. The reads were aligned to human, viral (Genbank ID: NC_003409 for KSHV, NC_001664 for HHV6a) and microbial genomes using CLCbio software (CLCbio Inc.) and EMBOSS (Rice et al., 2000).

By computing the number of sequence reads that aligned to a given pathogen genome against the genome size, we estimated the relative frequency of the pathogen. For human DNA, we obtained an average 0.005 reads/bp; more for mitochondrial genomes, since there are multiple mitochondria per cell genome. For KSHV and HHV6a we obtained almost equal coverage with 0.003 reads/bp. In contrast, HHV6b coverage was 10 fold lower (5 \times 10^{-4} reads/bp), as was EBV (3 \times 10^{-5} reads/bp) and (HCMV 8 \times 10^{-6} reads/bp) coverage. Our discovery score D takes a linear combination of the number of hits, the target genome and coverage. It was calculated for each genbank entry and each sample and subjected to unsupervised clustering using R v2.8.0 (Maingdonald and Braun, 2007).

Phylogenetic comparison

To compare our KSHV isolate to the existing whole genome data, we used clustalW and the following sequences: BCBL1 (Genbank ID: HQ404500), JSC1 (Genbank ID: GQ994935), KS biopsy (Genbank ID: U93872), BC1 (Genbank ID: KSU75698), and GK18 (Genbank ID: NC_009333). Our patient sequence is indicated as Consensus and submitted as Genbank ID: JQ619843.

Immunohistochemistry

A formalin fixed, paraffin-embedded lymph node was sectioned and dehydrated through a series of ethanol washes. Sections were then stained using a rat monoclonal anti-LANA antibody (Advanced Biotechnologies Inc.) diluted at 1:1000, and Vectastain™ (Vectorlabs Inc.) kit, counterstained with hematoxylin, and rehydrated through a series of xylene washes before being fixed with a xylene-based mounting solution. Images were collected on a LEICA DM LS microscope with DFC480 camera.

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