Method for Discovering Novel DNA Viruses in Blood Using Viral Particle Selection and Shotgun Sequencing

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Method for discovering novel DNA viruses in blood using viral particle selection and shotgun sequencing

Mya Breitbart and Forest Rohwer

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Rapid identification of viruses is needed to monitor the blood supply for emerging threats. Here we present a method that meets these criteria and allows for the shotgun sequencing of novel, uncultured DNA viruses directly from human blood. This method employs selection based on the physical properties of viruses combined with sequence-independent amplification and cloning. We show that both single- and double-stranded DNA viruses can be recovered from blood samples using this approach. In addition, we report the discovery of novel anellovirus sequences in the blood of healthy donors. PCR primers designed to amplify these novel anellovirus sequences were then used to verify the presence of these viruses in the general donor population.

INTRODUCTION

New viral infections in humans may be emerging at an increasing rate (1). The safety of the blood supply is currently maintained by selective screening of donors (2), pathogen reduction (3), and nucleic acid and antibody testing for known viruses, including human immunodeficiency virus (HIV), hepatitis B virus (HBV), hepatitis C virus (HCV), human T cell lymphotrophic virus (HTLV), and West Nile virus (WNV). These practices have made the blood supply extremely safe from the transmission of known viruses (the risk of contracting HIV or HCV through a blood transfusion is less than 1 in 2,000,000) (4).

The largest potential threats to the blood supply are posed by novel pathogens, which would escape current screening techniques. In order to identify emerging viruses, an active surveillance system capable of continually monitoring the blood supply is needed. Hybridization to DNA microarrays containing all known viral sequences and low-stringency PCR using degenerate primers to highly conserved viral sequences have successfully been used to identify and characterize novel viruses (5,6). These methods rely on conserved sequences being present within viral genomes, which makes them inadequate for the discovery of completely novel viral types in human blood samples.

The use of representative differential amplification (RDA) (7) has led to the discovery of several novel viruses in humans, including HCV (8), hepatitis GB virus (9), herpesvirus 8 (10), and Torque teno virus (TTV) (11). RDA was originally intended for comparing two cellular genomes, and applying this method to viral discovery in human blood requires that the composition of the background host DNA to be subtracted out is similar in both the tester and driver samples. In addition, RDA typically involves large quantities of DNA, which would require a high level of viremia before the virus could be recovered. While RDA is appropriate if samples from the same individual are being compared before and after viral infection, it is not clear that this approach will be applicable to blood pooled from multiple individuals.

Two recent studies have presented viral discovery methods that utilize selection for viral particles (based on size, density, and nuclease resistance), followed by sequence-independent amplification (12,13). These approaches have been used to identify new bovine parvoviruses present in commercial bovine serum (12), sequence novel viruses from the plasma of patients with symptoms of acute viral infection (14), and diagnose herpes simplex virus type 1 (HSV-1) in the supernatant of infected cells isolated from the mouth-washing material of a chronic fatigue syndrome patient (13).

Several viruses, such as TTV (15–17), TTV-like mini virus (TLMV) (18),
SEN virus (SENV) (19), Epstein-Barr virus (20), and B19 (21) are found in the blood of healthy people. The process of discovering novel blood viruses is hindered by methodological limitations such as low viral DNA content, abundance of host and mitochondrial DNA, and the absence of genes that are conserved among all viruses. Here we present a method to shotgun-sequence novel, uncultured DNA viruses directly from human blood. This method employs selection based on the physical properties of viruses combined with sequence-independent amplification and cloning. Using this method, we discovered novel anellovirus sequences in the blood of healthy donors.

**MATERIALS AND METHODS**

**Blood Collection**

All blood collection was performed by technicians at the General Clinical Research Center at the Scripps Research Institute (San Diego, CA, USA). Ten milliliters of blood were collected from healthy blood donors into sodium heparin vacutainers (BD Biosciences, Franklin Lakes, NJ, USA) and stored at 4°C until processing (within 1 week).

**Sample Descriptions**

Library 1 was constructed from blood pooled from 20 healthy donors collected in November 2004. Libraries 2 and 3 were constructed from blood pooled from 15 healthy donors collected in February 2005. The pooled plasma samples for Libraries 1 and 2 were spiked with 10^7 bacteriophage M13 particles per milliliter as a tracer to ensure the recovery of single-stranded DNA (ssDNA) viruses. Library 3 was not spiked with any tracers. For screening with specific primers, plasma was pooled from approximately 100 additional healthy donors (collected between September 2004 and February 2005).

**Isolation of Viral DNA**

The methods for obtaining viral DNA from blood samples are outlined in Protocol 1. Whole blood samples were centrifuged at 3000x g at 4°C for 10 min to pellet the blood cells. Eight and a half milliliters of cell-free plasma were loaded onto a cesium chloride (CsCl) step gradient consisting of 1 mL each of 1.7, 1.5, and 1.2 g/mL in 1× phosphate-buffered saline (1× PBS; 137 mM NaCl, 2.7 mM KCl, 4.3 mM Na_2HPO_4, 1.4 mM KH_2PO_4, pH 7.3). The gradient was designed to capture all known groups of DNA viruses (22). The gradients were ultracentrifuged at 55,000x g at 4°C for 2 h, and the 1.2–1.5 g/mL fraction was collected. This fraction was stained with SYBR Gold (Molecular Probes, Eugene, OR, USA) and viewed under epifluorescence and electron microscopy to ensure that it contained virus-like particles (23,24). The CsCl-purified viral particles were then incubated with 0.2 volumes of chloroform for 10 min at room temperature. The chloroform was pelleted, and the supernatant was removed and incubated with 2.5 U DNase I (Sigma-Aldrich, St. Louis, MO, USA) per microliter of sample for 2 h at 37°C. This sample was then concentrated and washed twice with 1× PBS on a Microcon® 30 (Millipore, Billerica, MA, USA). EDTA was added to a final concentration of 20 mM to inactivate the DNase I, and DNA was then extracted using the QIAamp® MinElute™ Virus Spin Kit (Qiagen, Valencia, CA, USA) according to the manufacturer’s instructions. Total DNA was also extracted from unprocessed cell-free plasma from the healthy donors using the QIAamp Blood Mini Kit (Qiagen). This was used for the comparison of human, mitochondrial, and TTV DNA found in unprocessed plasma versus viral DNA purified using the method outlined in Protocol 1.

**PCR to Check DNA Quality**

Viral DNA extracted from the blood samples was screened by PCR for the presence of (i) human 18S rDNA, (ii) human mitochondrial DNA, and (iii) TTV. All primer sequences are listed in Table 1. PCR products positive for TTV in the nested reaction were sequenced for verification. The PCR mixture (50 μL total volume) for the specific primer

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**Figure 1. Transmission electron micrographs of some viruses found in the blood of healthy donors.**

<table>
<thead>
<tr>
<th>Protocol 1. Overview of the Methodology for Obtaining Purified Viral DNA from Whole Blood Samples</th>
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<tr>
<td>1. Removal of blood cells by centrifugation</td>
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<td>2. Purification of viral particles by CsCl density-dependent centrifugation (results may be visualized by electron microscopy; see Figure 1)</td>
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<td>3. Chloroform treatment</td>
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<td>12. Bioinformatic analyses</td>
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<td>13. Confirmation by specific PCR (see Figure 3)</td>
</tr>
</tbody>
</table>

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In order to recover ssDNA viruses, the pooled plasma used to construct Libraries 1 and 2 were spiked with 10^7 particles of bacteriophage M13 (22) per milliliter before processing the samples as outlined in Protocol 1. The presence of spiked M13 in the purified viral samples was not monitored by PCR, but was determined based on TBLASTX similarities of the shotgun library sequences.

**Construction of Viral Shotgun Libraries**

**Linker-amplified shotgun libraries—Libraries 1 and 2.** Once viral DNA samples were determined to be free of mitochondrial and host DNA, linker-amplified shotgun libraries (LASLs) were constructed by Lucigen (Middleton, WI, USA) using a modification of the protocol described by Breitbart et al. (www.sci.sdsu.edu/PHAGE/LASL) (26). Total DNA was initially amplified using the strand-displacement enzyme Bst DNA polymerase (New England BioLabs, Beverly, MA, USA). The amplified DNA was then sheared into 1–2 kb fragments using a HydroShear® (GenMachines, San Carlos, CA, USA), the fragments were end-repaired, and short double-stranded DNA (dsDNA) linkers were ligated to the ends. These fragments were then PCR-amplified using primers to the linkers and cloned into the pSMART vector as previously described (26). Transformants were sequenced by the San Diego State University Microchemical Core Facility (San Diego, CA, USA) using the M13F primer.

**Bioinformatic Analysis of Shotgun Library Sequences**

Quality scores were assigned to the sequences using Phred (27,28), and the sequences were trimmed using Sequerencer™ 4.0 (Gene Codes, Ann Arbor, MI, USA). Sequences were trimmed until there were not more than 3 bases with confidence scores below 25 within 25 bases of either end. Sequences were compared against GenBank using TBLASTX (29,30). For phylogenetic analyses of the novel anellovirus sequences (SDSU1 and SDSU2), sequences showing significant TBLASTX hits were downloaded from GenBank. The sequences were then aligned using CLUSTALX (31) and neighbor-joining trees were constructed, bootstrapped 1000 times, and analyzed using Treeview (see Figure 3) (32).

**PCR for Specific Sequences Identified by Shotgun Library Sequencing**

Primer 3 (33) was used to design primers to amplify three of the viral sequences identified through sequencing of the shotgun libraries.
The primers were designed to amplify sequences that had top TBLASTX hits to an uncultured anellovirus (DQ092578), SENV (DQ092579), and TLMV (DQ092580).

The PCR mixture (50 μL total volume) for the specific primer sets contained 5 μL of target DNA, 1× FideliTaq PCR Master Mix, and 1 μM each primer. The thermal cycler conditions were 5 min at 94°C; 35 cycles of 1 min at 94°C, 1 min at 60°C–0.5°C per cycle, 1 min at 72°C; and 10 min at 72°C. The identity of PCR positives was confirmed by sequencing.

RESULTS

Electron Microscopy of Purified Blood Viruses

Electron microscopy (Figure 1) verified the presence of viral particles in the plasma of healthy blood donors. The majority of the viral particles observed were approximately 100 nm in diameter. In all cases, the viral particles were concentrated in the band isolated from the CsCl gradient, and no viruses were observed in the other fractions after centrifugation (data not shown).

Removal of Human and Mitochondrial DNA

Because high host and mitochondrial DNA concentrations overwhelmed the viral DNA signal (data not shown), CsCl density-dependent centrifugation, chloroform, and DNase I treatments were used to remove human and mitochondrial DNA. Free human DNA was removed beyond the detection limit of the PCR assay by the CsCl centrifugation and DNase I treatment (Figure 2). The addition of chloroform was necessary to disrupt the mitochondrial membranes, making them susceptible to DNase I digestion. Viral nucleic acids were protected from DNase I degradation by the viral protein capsids, although some enveloped viruses may have been lost. PCR assays showed that this method significantly reduced the concentrations of host and mitochondrial DNA compared with whole plasma, without resulting in observable losses of viral DNA (Figure 2 and data not shown).

Recovery of Viral DNA

PCR-based control experiments showed that both naturally present TTV and artificially spiked adenovirus 7 tracers were recovered from human plasma using this method (Figure 2). Because concentrations of adenovirus can reach up to $10^{10}$ viruses per milliliter in the blood of infected patients (34), an addition of $10^7$ viruses per milliliter is physiologically relevant.

TBLASTX analysis of the sequences from the viral shotgun libraries showed that this method recovered artificially spiked bacteriophage M13 as well as naturally present anelloviruses. For the libraries that were artificially spiked with $10^7$ bacteriophage M13 particles per milliliter, 20%–40% of the shotgun library sequences originated from bacteriophage M13 (as determined by TBLASTX similarity). These sequences could be assembled into larger contigs, demonstrating that the genome of a virus present at a concentration of $10^7$ virions...
This study

Novel TLMV
Novel SENV

This study

Sequence

SD

This study

small anellovirus 2 (AY787831).

41% identical on the nucleotide level

Library 2 (A 286-bp fragment recovered from to previously sequenced anelloviruses.

relationships of these novel sequences

Figure 3 shows the phylogenetic

further analyzed two of these sequences.

the blood of the general population, we

numerous recent studies have demon

known viruses were observed in each

complete genomes could be assembled

per milliliter would be recovered at

high coverage, and it is possible that

can be assembled using this method. Several naturally occurring sequences with similarity to known viruses were observed in each library (Table 2). In addition, approximately 10% of the shotgun sequences had no similarities in GenBank with an E-value <0.001. These sequences may represent undescribed groups of viruses.

Identification of Novel Viral Sequences

Sequences from the healthy donor plasma shotgun libraries with significant TBLASTX similarities to known viruses are listed in Table 2. Because numerous recent studies have demonstrated the presence of anelloviruses in the blood of the general population, we further analyzed two of these sequences. Figure 3 shows the phylogenetic relationships of these novel sequences to previously sequenced anelloviruses. A 286-bp fragment recovered from Library 2 (SDSU1; DQ092578) was 41% identical on the nucleotide level and 35% on the amino acid level to small anellovirus 2 (AY787831).

A 284-bp sequence recovered from Library 3 (SDSU2; DQ092579) was 67% identical on the nucleotide level and 63% identical on the amino acid level to SENV (AY206683) (Figure 3, C and D). These divergent anellovirus sequences constitute novel viruses identified using the methods presented here and contribute to the wide genomic heterogeneity of this group.

Specific PCR to Confirm the Presence of Novel Anelloviruses

PCR primers designed to amplify three of the novel anellovirus sequences identified in this study confirmed their presence in blood pooled from 100 different healthy donors (Figure 3A). PCR positives were sequenced for confirmation, and all amplified sequences were identical to those initially recovered from the shotgun libraries.

DISCUSSION

The method described here provides a new technique for sequencing novel DNA viruses from plasma samples. There is a high level of selection for viruses in this protocol through the removal of blood cells by centrifugation, the purification of viruses by CsCl density-dependent centrifugation, chloroform addition to lyse mitochondria, and DNase I treatment to remove nonviral DNA. This selection eliminates the need for the background human DNA signal subtraction that is performed in RDA analysis (8). It is important to note that the method outlined here employs a CsCl gradient for purifying viruses instead of filtration-based techniques. Filtering samples through a 0.2-μm filter (12,13) to remove cells and mitochondria may result in the loss of large viral particles (e.g., some herpesviruses are 0.2 μm in diameter) (22).

Depletion of Human and Mitochondrial DNA

Removal of free human host DNA was a critical step in this method because free human DNA may be present at significantly higher concentrations than viral DNA in plasma. Free human DNA was removed beyond the detection limit of our PCR conditions for human 18S rDNA by the CsCl density-dependent centrifugation and DNase I treatment (Figure 2).

In addition to removing free DNA, it was also necessary to remove mitochondria from the plasma sample. Chiu et al. (35) have shown that mitochondrial DNA is present in human plasma in both particle-associated and free forms. Preliminary experiments showed that if the mitochondria were not specifically removed from the plasma sample (by chloroform and then DNase I treatment), all the sequences recovered from the shotgun libraries were mitochondrial in origin (data not shown). The depletion protocols described here efficiently reduced the concentrations of mitochondrial DNA.
Table 2. Sequences from the Healthy Blood Donor Shotgun Libraries with Significant TBLASTX Similarities to Known Viruses

<table>
<thead>
<tr>
<th>Library (No.)</th>
<th>Top Known Virus Hit</th>
<th>TBLASTX E-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>TTV-like mini virus</td>
<td>9e-04</td>
</tr>
<tr>
<td>1</td>
<td>TTV-like mini virus</td>
<td>3e-25</td>
</tr>
<tr>
<td>2</td>
<td>TTV-like mini virus</td>
<td>2e-15</td>
</tr>
<tr>
<td>1</td>
<td>Heliothis zeae virus 1</td>
<td>1e-15</td>
</tr>
<tr>
<td>1</td>
<td>Heliothis zeae virus 1</td>
<td>2e-16</td>
</tr>
<tr>
<td>1</td>
<td>Heliothis zeae virus 1</td>
<td>9e-17</td>
</tr>
<tr>
<td>1</td>
<td>Heliothis zeae virus 1</td>
<td>4e-11</td>
</tr>
<tr>
<td>1</td>
<td>Pseudorabies virus</td>
<td>7e-05</td>
</tr>
<tr>
<td>1</td>
<td>Bacteriophage φ3626</td>
<td>2e-27</td>
</tr>
<tr>
<td>1</td>
<td>Cowpox virus</td>
<td>1e-19</td>
</tr>
<tr>
<td>1</td>
<td>Cowpox virus</td>
<td>1e-19</td>
</tr>
<tr>
<td>1</td>
<td>Human endogenous retrovirus H (HERV-H)</td>
<td>1e-25</td>
</tr>
<tr>
<td>1</td>
<td>Human endogenous retrovirus H (HERV-H)</td>
<td>5e-25</td>
</tr>
<tr>
<td>1</td>
<td>Bacteriophage EJ-1</td>
<td>2e-05</td>
</tr>
<tr>
<td>1</td>
<td>Human herpesvirus 3</td>
<td>5e-05</td>
</tr>
<tr>
<td>1</td>
<td>TTV</td>
<td>2e-24</td>
</tr>
<tr>
<td>1</td>
<td>TTV</td>
<td>1e-04</td>
</tr>
<tr>
<td>2</td>
<td>Small anellovirus 2a</td>
<td>3e-06</td>
</tr>
<tr>
<td>1</td>
<td>Methanobacterium phage psiM2</td>
<td>1e-04</td>
</tr>
<tr>
<td>1</td>
<td>Spodoptera litura nucleopolyhedrovirus</td>
<td>7e-08</td>
</tr>
<tr>
<td>1</td>
<td>Chlamydia phage φCPAR39</td>
<td>1e-09</td>
</tr>
<tr>
<td>3</td>
<td>SEN virusa</td>
<td>8e-38</td>
</tr>
</tbody>
</table>

TTV, Torque teno virus.
- Sequences that were further analyzed to identify phylogenetic relationships to known viruses (Figure 3, B–D).

Applications of this Protocol

Here we have shown that a method based on the selection for viral particles followed by sequence-independent amplification can recover DNA viral sequences from the blood of healthy people. Applying this technique to large donor pools will allow for the identification of viruses normally present in the blood supply. This will provide a baseline against which samples can be compared in order to identify emerging threats even before clinical symptoms are observed. In addition to its use for monitoring the blood supply, this method can also be applied to diagnose unknown viral infections in patients. For example, more than 50% of pneumonia cases and almost 60% of encephalitis cases requiring hospitalization in the U.S. have no known etiology (38,39). In the case of many viral infections, viremia phases are observed. In addition to the removal of the envelope, and if these viruses were rendered DNase-sensitive, then they would not be recovered using this approach. One possible alternative to the chloroform treatment is antibody-based depletion of the mitochondria. However, we found that this method did not consistently reduce mitochondrial numbers and had variable effects on the efficiency of viral recovery (data not shown).

Blood Sample Collection and Storage

This method was tested on blood samples that were collected using heparin as the anticoagulant and stored at either room temperature or 4°C. One needs to take caution when using blood preserved in EDTA, acid citrate dextrose, sodium citrate, and potassium or ammonium oxalate because these compounds chelate cations that are required for effective DNase I digestion and may compromise the integrity of viral particles. Further studies need to be performed to determine the ability of this method to recover viruses from frozen samples, serum, or blood fraction paste.

to the point that we could no longer detect mitochondrial DNA by PCR (Figure 2), and only a few sequences with TBLASTX similarities to known viruses were recovered from the shotgun libraries.

Identification of Novel Anellovirus Sequences

Among the shotgun sequences obtained from healthy donor plasma, several sequences with similarities to the anellovirus family were identified (Table 2). The anellovirus family (formerly part of the circovirus family) is a highly heterogeneous group of single-stranded, circular viruses, including TTV, TL MV, and SENV. Using the methodology presented here, we have detected several novel anellovirus sequences, which suggests that there is still undiscovered diversity within this family of viruses. Anelloviruses are present worldwide and are found in over 70% of the general population of Asia, Africa, and South America (17). These viruses appear to be less prevalent in the U.S. and Europe (16); however, TTV viremia occurs in 8%–42% of healthy blood donors in the U.S. (36,37) and 47%–73% of individuals in European countries (17). It has been suggested that TTV and SENV are associated with posttransfusion hepatitis; however, no definitive links between the presence of these viruses and clinical pathology have been demonstrated.

Possible Loss of Enveloped Viruses

The chloroform treatment that is required for lysing mitochondria will also disrupt the membrane of enveloped viruses. In control experiments with the enveloped virus HSV-1, the protein capsid remained intact and protected the viral DNA from degradation even after removal of the envelope (data not shown). Therefore, at least some enveloped viruses can be recovered using this method. However, the capsid of some viruses may not be stable after
about the identity of these viruses may yield insights into treatment options.

Because many viruses are cell-associated (e.g., Epstein-Barr virus and cytomegalovirus), one logical step for further development of this method is its application to human tissues. The method outlined here can also easily be adapted to identify RNA viruses through the addition of a reverse transcription step before the sequence-independent amplification (12,13). Finally, this method can also be applied to detect novel DNA viruses within animal populations, which is of potential importance for both farming and human health.

The protocol described here can be completed (sample collection to sequence data) in fewer than 2 days, which is faster than traditional protocols that require obtaining the virus in culture. Once sequence data are obtained from the newly identified virus of interest, PCR primers can be designed to rapidly screen a large number of samples in order to determine the prevalence, abundance, and geographical distribution of the novel virus. The safety of the blood supply can be monitored, and viral sequences can be compared in order to determine the origin of the virus and epidemiological links between patients (40).

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