Anti-HIV-1 Activity of *Crocodylus mindorensis* (Philippine Crocodile) Serum in Cell-free and Cell-associated Virus Interactions to Human Peripheral Blood Mononuclear Cells

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Abstract

Highly-Active Antiretroviral Therapy (HAART) is the recommended treatment and management strategy for HIV infection. Although the existing antiretroviral drugs are indispensably significant in improving the quality and extending the lives of HIV/AIDS individuals, the drugs still have many limitations including development of resistance, production of toxicity, and their limited availability. These limitations continue to open new opportunities in the use of ethno-medicine for the management of HIV/AIDS. With this, few researchers have made effort to test the inhibitory activity of crocodile serum as it has a unique and diverse molecular activity in preventing HIV-1 replication. In this study, a cell culture-based assay was utilized using peripheral blood mononuclear cells coupled with colorimetric enzyme immunoassay to determine the HIV-1 reverse transcriptase activity of the treated cell-culture system. One HIV-1 seropositive serum was processed for Peripheral Blood Mononuclear Cells (PBMC) co-culture from which HIV-1 isolates were obtained. The HIV-1 reverse transcriptase activity after 21 days was 0.5928 pg/well. Moreover, a baseline Philippine crocodile serum concentration of 0.5% vol/vol was used based on the previous study conducted by Hinay and Sarol in 2018, and the cell viability results showed no cell reduction of mononuclear cells after 72 hours incubation. The inhibitory activity of the Philippine crocodile serum at 0.5% and 0.25% vol/vol concentrations inhibited 65.68 ± 2.93% and 69.92 ± 0.45% respectively in post-infection interactions. In addition, the Philippine crocodile serum in pre-infection interaction at 0.5% and 0.25% vol/vol concentrations inhibited 68.61 ± 1.67% and 69.95 ± 2.24% respectively. As has been noted, the inhibitory actions of the Philippine crocodile serum effectively regulate the HIV-1 replication in both pre- and post-infection interactions. With this, the Philippine crocodile serum could be a novel source of HIV-1 replication inhibitors.

Keywords: Philippine freshwater crocodile, HIV-1 co-culture, HIV-1 inhibition, HIV-1 reverse transcriptase.

Introduction

Reptiles such as alligators exhibit remarkable ability to heal rapidly considering the septic environment in which they live. Merchant and colleagues provided observation on the broad range of antiviral activity of alligator serum [1,2]. Merchant’s recent studies also showed that these activities are due to a potent and broad-acting serum complement system [2]. In the Philippines, there are two species of crocodile-the *Crocodylus porosus* locally known as Philippine saltwater crocodile and the *Crocodylus mindorensis*, the endemic Mindoro crocodile. The relatedness of *Alligator mississippiensis* (American alligator) described by Merchant and colleagues and the Philippine crocodile *Crocodylus mindorensis* based on phylogeny shows that they belong to the same phylum (Crocodilia) but from different families [3].

Crocodiles may acquire serious injuries when they hunt food and may encounter higher order animals.
The antiviral activity of Alligator mississippiensis (American alligator) serum described by Merchant [7], conducted using cell culture-based method provided the evidence that it contains anti-HIV-1 activity with half maximal inhibitory activity of 0.9% [1,2]. In the Philippines, particularly in Davao, crocodiles which are characteristically similar to alligators are not only abundant but are scientifically cultured and propagated. With the availability of Philippine crocodile, a scientific research that will be a foothold and basis that Crocodylus mindorensis serum could be used as a potential inhibitor of HIV/AIDS infection was investigated.

**Subjects and Methods**

**Specimen collection**

A minimum of 2 mL serum sample from one purposively selected adult- male Crocodylus mindorensis was collected at the Animal Clinic of the Davao Crocodile Park facility. The freshly collected sera were separated from whole blood samples collected by a veterinarian as part of the health check of the crocodile. Specifically, 5 mL of whole blood were allowed to clot at room temperature for approximately three hours. The serum was separated by centrifugation at 3,000 xg for 15 minutes and processed immediately.

**Ethical considerations**

**Blood Sampling Protocol:** Peripheral blood mononuclear cells were obtained from two different individual sources. First from a healthy non-HIV-1 reactive individual and second from a HIV-1 reactive individual. An inform consent were given to both participants before blood collection.

A volunteer HIV-1 reactive individual with a signed inform consent was the source for HIV-1 co-culture method. A three (3) 5 mL EDTA tube was used to collect the blood using venepuncture technique with strict compliance to Occupational Safety and Health Administration 1910.1030 Blood-borne Pathogens guidelines (OSHA 1910.1030). After collection, the sample was processed immediately. The same protocol was followed for healthy non-HIV-1 reactive volunteer.

**Biosafety considerations**

Upon arrival of crocodile serum sample, proper protocol was observed including serum separation to anti-HIV-1 activity assessment. At the end of the experiment, the used materials such as microplate and washing basin containing 5.25% sodium hypochlorite were decontaminated at 12°C at 15 psi for 30 minutes including all crocodile serum received.

**HIV-1 isolation**

Isolation of HIV-1 from one sero-positive sample was done using a PBMC micro-co-culture assay [8]. Briefly, one healthy, HIV-sero-negative donor PBMCs were stimulated with the mitogen phytohemagglutinin-P (PHA-P; at a final concentration of 5.0 μg/mL), in the presence of human interleukin 2 for 24-72 hours before use to promote blast formation and replication of T-cells. The cells were counted and 1×10⁶ PHA-stimulated donor cells and 1×10⁶ PBMCs from sero-positive HIV-1 individual was added in duplicate wells of a 24-well tissue culture plate. The final volume was adjusted to 2.0 mL with growth media containing RPMI1640 with 20% FBS and 10.0 U/mL IL-2. The plate was incubated at 37°C with 5% CO₂ in a CO₂ incubator. On day 7 and 14, the cultures were replenished with fresh growth media containing 5×10⁵ PHA stimulated donor cells (feeder cells). The culture was continued until day 21 and then terminated. Supernatant fractions from duplicate wells from day 14 and 21 were saved separately, and stored at -70°C until analysis for HIV-1 RT by ELISA. Cultures were considered positive only when day 21 supernatants showed an increase in HIV-1 RT from day 14 supernatants.

**Cell viability assay**

Before using the Philippine crocodile serum sample for antiviral assay, it was necessary to assess their toxicity against Peripheral Blood Mononuclear Cells (PBMC). Briefly, a 100 μL PBMC was suspended into a 0.5 mL microcentrifuge tube and added 90 μL of Safranin stain. The PBMC were counted to 1×10⁶ cells. After determining the PBMC count, cell viability assay was performed. Briefly, a 100 μL 1×10⁶ PBMC were suspended into a...
0.5 mL microcentrifuge tube and added 100 μL of 0.5% and 0.25% vol/vol Philippine crocodile serum. The interactions were allowed up to 72 hours and cell viability was counted.

**Post-infection interaction (cell-associated HIV)**

The procedure was based on the study of Dahake et al. [8]. Briefly, HIV-1 virions from co-cultured were first allowed to infect PBMCs and the Philippine crocodile serum were added to the suspension after one hour. A volume of 100 μL of HIV-1 virions were added to 1 mL of PHA-stimulated PBMCs and incubated for 2 hours in a CO₂ incubator. After incubation, the cells were washed carefully, and supernatants were aspirated set to leave 50 μL/well. After washing, 100 μL of infected cells were plated into wells and 100 μL of Philippine crocodile serum were added. The plates were incubated for 72 hours at 37°C in the CO₂ incubator. The contents of each well were transferred to microfuge tubes, cells were pelleted at 13,000 RPM for 5 minutes and the supernatants used for determining the HIV-1 RT levels.

**Data management**

The results of pre-infection and post-infection assays were expressed as means ± SD in triplicates. The Percent (%) reduction of HIV-1 activity was computed using the following mathematic formula:

\[
\% \text{ inhibition} = \frac{Absorbance \text{ of Negative control} - Absorbance \text{ of test sample}}{Absorbance \text{ of Negative Control} \times 100}
\]

*Negative control - without crocodile serum
**Test sample - with Crocodile Serum

**Results and Discussion**

**HIV-1 co-culture and cell viability**

The HIV-1 RT activity after 21 days was 0.5928 ng/well. On the other hand, a baseline Philippine crocodile serum concentration of 0.5% vol/vol was used based on the study conducted by Hinay and Sarol [9] and the cell viability results showed no cell reduction of mononuclear cells after 72 hours incubation.

<table>
<thead>
<tr>
<th>Concentration % vol/vol</th>
<th>Post-infection Mean ± SD</th>
<th>Post-infection Mean ± SD</th>
</tr>
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<tbody>
<tr>
<td>0.50</td>
<td>65.68 ± 2.93</td>
<td>68.61 ± 1.67</td>
</tr>
<tr>
<td>0.25</td>
<td>69.32 ± 0.45</td>
<td>69.95 ± 2.24</td>
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**Table 1:** Inhibitory activity of the Philippine crocodile serum in post-infection interaction (cell-associated HIV).
Post-infection interaction (cell-free HIV)

Table 1 shows the inhibitory activity of the Philippine crocodile serum. HIV-1 replication inhibition at 0.5% and 0.25% vol/vol concentrations were 65.68 ± 2.93% and 69.92 ± 0.45% respectively. This is also supported with the HIV-1 Reverse Transcriptase activity of the peripheral blood mononuclear cells (Figure 1). The PBMCs interacted with crocodile serum concentration of 0.5% and 0.25% vol/vol shows a decreased HIV-1 reverse transcriptase activity of 0.1216 and 0.0954 ng/well respectively.

Unexpectedly, an inverse relationship was observed between crocodile serum concentration and the anti-HIV-1 Reverse Transcriptase activity. This means that the inhibitory activity decreases as the Philippine crocodile serum concentration increases. This is similar to the previous study of Hinay and Sarol [9], were the result could be explained by the presence of an “inhibitor” of the inhibitor” in the crocodile serum. The best candidate inhibitor present in the crocodile serum with putative inhibition activity to HIV-1 is the Antimicrobial Peptides (AMPs) [10]. This is also supported by the study of Hinay and Sarol [9] where Philippine crocodile serum concentration showed in vitro inhibition of HIV-1 Reverse transcriptase by high as 92.93 ± 0.72% at 0.5 vol/vol %. Moreover, another putative component of the Philippine crocodile serum as described by Okada and colleagues provided a good support that the complement fragment C5a, an anaphylotoxin and a small peptidase product during complement activation, has a 30% HIV-1 Reverse transcriptase inhibitory activity [11]. The inhibition of the HIV-1 virion interacted with the Philippine crocodile serum indicates that the inhibition occurs either direct interaction to viral enzymes reverse transcriptase and or protease which are key viral enzymes of HIV-1 replication.

Pre-infection interaction (cell-free HIV)

The potential of the Philippine crocodile serum as inhibitor against HIV-1 replication was determined and the findings demonstrated a significant inhibition of HIV-1 replication prior to interaction to PBMCs. Table 1 shows the inhibitory activity of the Philippine crocodile serum in pre-infection interaction. Reverse Transcriptase inhibition at 0.5% and 0.25% vol/vol concentrations inhibited 68.61 ± 1.67% and 69.95 ± 2.24% respectively. This is also supported with the HIV-1 Reverse transcriptase activity of the peripheral blood mononuclear cells (Figure 2). The PBMCs interacted with crocodile serum concentration of 0.5% and 0.25% vol/vol with an HIV-1 reverse transcriptase activity of 0.1005 and 0.10054 ng/well respectively. With this the Philippine crocodile serum have both direct virucidal effects to HIV-1 virion or the crocodile serum inhibits the viral entry subsequently results through inhibition of fusion of the viral envelope with the host cell membrane [12]. Possible crocodile serum component such as Alpha-1 antitrypsin (AAT) which is an acute phase reactant and the most abundant circulating serine protease inhibitor can be a putative HIV-1 inhibitor [13]. Shapiro et al. described Alpha-1 antitrypsin (AAT) in their study as a natural HIV-1 antagonist. In their study the inhibitory activity of the Alpha-1 antitrypsin was observed in the induction of latent virus from chronically infected cells, viral infection and replication in PBMC, attachment CCR-5 - cell assay and infection and replication in whole blood [13].

Merchant et al. described the effectiveness of alligator serum against the laboratory strain of Human immunodeficiency virus type-1 using a cell culture-based assay. The results exhibited potent anti-HIV activity at 20- and 64-fold dilutions with 100 and 89% reduction in cytopathic effects, respectively. The study also showed
that the higher concentration of the serum was difficult to assess due to inherent toxicity to the cell culture used in the study [7]. In addition, the results suggest that both the antiHIV-1 activity and cell toxicity were mediated by serum complement activity as heat treatment of the serum destroyed both effects. Like human complement activity, alligator serum complement is sensitive to heat inactivation [2].

These findings are quite significant since the Philippine crocodile serum concentration of 0.5% and 0.25% vol/vol seems to work in both pre- and post-infection interactions and hence suggest that the effect may be mediated by a direct action on the virus particle. Perhaps it blocks binding or uptake to cells or causes immediate particle lysis.

Conclusion

The results showed that the HIV-1 Reverse Transcriptase activity was inhibited by 65.68 ± 2.93% and 69.92 ± 0.45% in Philippine crocodile serum concentration of 0.5% and 0.25% vol/vol respectively in post-infection interaction and 68.61 ± 1.67% and 69.95 ± 2.24% in Philippine crocodile serum concentration of 0.5% and 0.25% vol/vol respectively in pre-infection interaction. As has been noted, the 0.5% and 0.25% vol/vol concentration of the Philippine crocodile serum effectively regulates the HIV-1 replication in both pre- and post-infection interactions. With this, the study recommends identifying and characterize the active compounds in the Philippine crocodile serum and develop a novel source of HIV-1 replication inhibitors.

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Conflict of Interest

No conflict of interests is declared by authors for the contents in this manuscript.

Author’s Contribution

Alfredo A. Hinay Jr., Nelyn Mae T. Cadotdot and Marilou V. Tablizo designed and carried out the experiment and prepared the draft manuscript.

References