

## Abstract

**Background:** A novel, isothermal nucleic acid amplification method, RT-LAMP, presents potential for nucleic acid amplification-based diagnostics in resource-limited settings. Low-cost HIV-1 viral load monitoring will improve access to ART for HIV-1-infected individuals present in settings where on-site viral load testing is unavailable.

**Aim:** The aim of this dissertation was to develop an RT-LAMP HIV-1 viral load assay by combining the RT-LAMP reaction with colorimetric amplification detection by hydroxynaphthol blue dye.

**Methods:** Different approaches for HIV RNA extraction from patient plasma and culture supernatant were studied to obtain template for RT-LAMP. Reaction products for 4 different RT-LAMP primer sets were analysed using agarose gel electrophoresis and restriction digestion.

**Results:** The first 3 primers sets produced persistent off-target amplification. The fourth primer set, designed against culture supernatant DU179, produced a target-specific colour change from violet to blue after 1 hour, following optimisation of amounts of  $Mg_2SO_4$  and AMV RT. Further studies showed HNB detection sensitivity to template copy number.

**Conclusions:** Initial reaction conditions pertaining to an RT-LAMP based, colorimetric HIV-1 viral load assay were established. Further work is required to determine the reaction duration at which the colour change represents a viral load of  $\geq 1000$  copies HIV RNA per ml plasma.