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A Thesis Submitted to the Graduate School in Partial Fulfilment of the Requirements for the Master of Science Degree in Biochemistry of Egerton University

EGERTON UNIVERSITY

## DECLARATION AND RECOMMENDATION

## Declaration

This thesis is my original work and has not been presented in this university or any other for the award of a degree to my knowledge.

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## DEDICATION

To the family and friends for whose support contributed to completion of this work.

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#### Abstract

Africa green monkeys (AGMs, Chlorocebus aethiops) (Gray, 1870) and olive baboons (Papio anubis anubis) (Lesson, 1827) are common non-human primates (NHPs) found within major urban centres in Kenya. The widely distributed AGMs are naturally infected with simian immunodeficiency virus (SIV) of the genus Lentivirus. While a baboon specific SIV has not been reported so far, studies have demonstrated that baboons are non-receptive to SIV infection leading to low prevalence in these species. Due to enhanced human-wildlife interactions, SIV can potentially infect humans during aggressive encounters resulting in injuries from bites and scratches and potential exposure to infected blood and other body fluids making SIV a virus of medical importance. Despite its zoonotic potential, a comprehensive investigation of SIVs in free-ranging Kenyan monkeys has not been undertaken. Therefore, this study sought to investigate the diversity of SIV strains infecting AGMs and olive baboons found within selected Kenyan urban centres using molecular approaches. Free-ranging NHPs (124 AGMs and 65 olive baboons) from within Mombasa, Kisumu, and Naivasha towns were trapped, and SIV identified by PCR targeting a partial pol and env gene fragments. Polymerase chain reaction-high resolution melt (PCR-HRM) analysis of pol amplicons revealed distinct melt profiles illustrating diverse virus strains. Detected SIV genetic fragments were further characterised by sequencing and phylogenetic analysis. We detected SIV in 32\% (39/124) and $3 \% ~(2 / 65)$ of AGMs and baboons, respectively. Phylogenetic analysis of the pol and env gene sequences demonstrated that diverse host species-specific SIV (SIVagm) strains infect AGMs populations without definite phylogeographical groupings. Moreover, analysis of the evolutionary selection demonstrated signatures of episodic and pervasive diversification on the env gene suggesting continuous SIV evolution within the natural host which is crucial for a virus to be able to cross the species barrier and infect a new host. Notably, for the first time, this study partially characterised a strain of SIVagm infecting olive baboons indicating putative cross-infection among sympatric NHP species. Additional elaborate studies are required to conclusively decipher the prevalence, pathogenesis and immunological response associated with SIVagm infection of baboons. Better understanding of prevalence and diversity of potentially zoonotic SIV strains circulating in NHP hosts is crucial in controlling emergence of infections in human.


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## LIST OF ABBREVIATIONS AND ACRONYMS

| AIDS | Acquired Immunodeficiency Syndrome |
| :---: | :---: |
| AGM | African Green Monkey |
| BLAST | Basic Local Alignment Search Tool |
| CA | California |
| cDNA | Complementary Deoxyribonucleic Acid |
| CRFs | Circulating Recombinant Forms |
| DNA | Deoxyribonucleic Acid |
| dNTPs | Deoxynucleotide Triphosphates |
| EI | Emerging Infection |
| GAC | Government Administration Compound |
| GC | Golf Course |
| HIV | Human Immunodeficiency Virus |
| icipe | International centre of insect physiology and ecology |
| ISERC | Institutional Scientific Review Committee |
| IPR | Institute of Primate Research |
| KAJ | Kajulu |
| KWS | Kenya Wildlife Services |
| MBK | Mbaraki |
| MERS | Middle East Respiratory Syndrome |
| NEB | New England Biolabs |
| NCBI | National Centre for Biotechnology |
| NHP | Non-Human Primate |
| PCR-HRM | Polymerase Chain Reaction- High Resolution Melting |
| RNA | Ribonucleic acid |
| SARS | Severe Acute Respiratory Syndrome |
| SIV | Simian Immunodeficiency Virus |
| TB | Tuberculosis |
| TRIM5 $\alpha$ | Tripartite Motif-containing protein 5 |
| TWS | Tudor water sport |
| UNAIDS | United Nations Programme on HIV/AIDS |
| VPU | Viral Protein U |
| WFS | World Federation of Scientists |
| WHO | World Health Organization |

## CHAPTER ONE

## INTRODUCTION

### 1.1 Background information

Simian immunodeficiency virus (SIV) belongs to the Lentivirus genus Retroviridae family (Desrosiers, 1990) and naturally infects over 30 species of African anthropoid nonhuman primates (NHPs) including chimpanzees and gorilla (Apes), and simian primates (monkeys) (Peeters et al., 2002). It has been shown that SIVcpz strain originating from chimpanzees (Pan troglodytes) and SIVsmm strain from sooty mangabeys (Cercocebus atys) represents zoonotic cross-species infections that produced the human immunodeficiency virus type 1 (HIV-1) and HIV-2 respectively (Chen et al., 1996; Sharp et al., 2005). HIV is one of the most devastating infectious disease in humans responsible for 1.1 million deaths in 2018 while an approximately annual budget of 26.2 billion dollars is spent on acquired immunodeficiency syndrome (AIDS) response (UNAIDS, 2019). The HIV virus first discovered in 1985 by Gallo \& Montagnier (2003), is thought to have emerged from zoonotic cross-species transmission from NHPs (Sharp \& Hahn, 2011) and was considered an emerging infection (EI) in the twentieth century. This demonstrates the human risk of contracting deadly infectious pathogens that originate from wild and domestic animals.

More than $60.3 \%$ of emerging infections (EIs) are caused by pathogens that infect animals and can be transmitted into humans i.e. zoonotic and $71 \%$ are of wildlife origin (Jones et al., 2008; Petersen et al., 2018). They are responsible for more than $25 \%$ of 57 million annual deaths (Wang et al., 2016) and approximately $30 \%$ of disability adjusted life years globally (Chua \& Gubler, 2013; James et al., 2018). In addition, economic losses of approximately 120 billion dollars in the past decade has been incurred due to EIs alone (Smith et al., 2019). Due to the nature of EIs, their direct and indirect impact on socio-economic activities will increase. For example, most of these diseases are zoonotic and with increasing human population necessitating change of land use, increased interactions between people and animals is inevitable, resulting into acquisition of new infections that can be rapidly translocated through international travel and trade (Rhyan \& Spraker, 2010). Further, the poor state of public health infrastructure in some regions does not allow efficient control and management of existing and new infections. An excellent example is the recent (2019) Ebola outbreak in Democratic Republic of Congo (DRC). This therefore calls for improved knowledge to inform on appropriate management and control of known EIs and/or emergent ones. In part, it is important to understand potential zoonotic pathogens in their natural non-human hosts to aid in predicting the potential risks of cross-species transmission (Rhyan \& Spraker, 2010).

A potentially high risk virus is SIV, a single stranded RNA virus is known to naturally infect 45 different species of NHPs including apes and monkeys in Africa (Peeters et al., 2012). Several studies have demonstrated that SIV has crossed and infected various NHPs such as chimpanzee and sooty mangabeys on multiple occasions (Gao et al., 1999; Sharp \& Hahn, 2011; D'arc et al., 2015), and possibly evolved into HIV that caused the AIDS pandemic (Sharp \& Hahn, 2010). This strongly suggests that active transmission of retroviruses from NHP to humans (Wolfe et al., 2004; Locatelli \& Peeters, 2012; Peeters \& Delaporte, 2012) is possible, and is a potential source of novel EIs. Therefore, there is need to identify and understand microbes of NHPs, specifically viruses that can evolve into highly infectious human pathogens.

SIV poses a threat of transmission to humans because of hunting and consumption of NHPs as bushmeat (Friant et al., 2015; Steve et al., 2017). Handling and processing of NHP bushmeat exposes humans to infected blood and other body fluids thereby serving as major route of human infection with SIV. Phylogenetic analysis of SIV in African NHPs has demonstrated species specificity where viruses cluster according to their respective hostspecies (Locatelli \& Peeters, 2012). This has been documented in viruses infecting Chlorocebus genus that includes AGMs (Wertheim \& Worobey, 2007) and the arboreal Cercopithecus genus (Bibollet-Ruche et al., 2004). Further, epidemiological studies have revealed high SIV prevalence in wild AGM populations in West and Southern Africa (Ma et al., 2013). High prevalence of SIV and diversity in NHPs sharing a habitat (sympatric) has been associated with cross-species transmission of the virus. Examples include cross-species transmission between sympatric AGMs and chacma baboons (Papio ursinus) in Southern Africa (van Rensburg et al., 1998), yellow baboons (Papio cynocephalus) in Tanzania (Jin et al., 1994) and patas monkey (Cercopithecus aethiops sabaeus) in West Africa (Bibollet-Ruche et al., 1996). Despite the high prevalence of natural SIV infections in different NHPs species, comprehensive surveys and molecular characterisation of this lentivirus, especially from East African monkeys remain limited.

AGM/vervet monkey (Chlorocebus aethiops), Sykes' monkey (Cercopithecus mitis) and olive baboon (Papio anubis anubis) are common NHPs found within most urban and periurban environments in East Africa (Newman et al., 2004; de Jong \& Butynski, 2010). These are the most widely distributed NHPs in Kenya and regarded as pests in most towns and villages (Jeneby et al., 2011; Kagira et al., 2017; Mutemi et al., 2017). As primates of diverse habitats, synanthropic or urban-restricted free ranging monkeys are potential sources of zoonotic viral diseases of human health importance such as chikungunya (Tsetsarkin et al., 2016), dengue, yellow fever, and Zika viruses (Weaver et al., 2018). Despite their importance as reservoir
hosts of zoonotic viral diseases, surveys for SIV infections in Kenyan monkeys have been opportunistic, and mostly on captive AGMs (Otsyula et al., 1996) and Sykes' monkeys (Ellis et al., 2004). In addition, comprehensive molecular epidemiology studies of SIV strains circulating in wild monkey populations in Kenya have not been performed to date. Opportunistic sampling of captive animals, difficulties in collecting and storing field samples have largely contributed to the understudy of SIV epidemiology in NHPs found within Kenya and East Africa at large.

Natural SIV infections of AGMs and Sykes' monkeys generally do not progress to AIDS like symptoms, contrary to SIV experimental infection of Asian macaque monkeys (Macaca mulatta) (Silvestri et al., 2007) and HIV infection in humans (Hirsch et al., 1995). These observations suggest the possibility of African NHPs being natural hosts to species-specific SIV lineages as demonstrated by phylogenetic clustering of known strains (Peeters et al., 2002; Ma et al., 2013; Steve et al., 2017). The AGMs are a super-species comprising of several phenotypically and geographically distinct subspecies of monkeys (Groves, 2016). These subspecies harbour distinct, but related SIV strains considered as subtypes that cluster according to their respective hosts within the SIVagm lineage (Johnson et al., 1989; Johnson et al., 1990; Hirsch et al., 1993; Carr et al., 2017). The AGMs are of specific interest in SIV epidemiology given their status as the largest known reservoirs of the virus with reported prevalence of up to $60 \%$ (Hirsch et al., 1995; Vanderford et al., 2006). Unlike AGMs, baboonspecific SIV strain has not been documented. However, previous seroprevalence survey of yellow baboons in Tanzania showed exposure to SIV (Jin et al., 1994). Follow-up PCR amplification and sequence analyses of these specimens reported infection by SIVagm originating from AGMs (Jin et al., 1994). Another study isolated and sequenced SIVagm strain from a healthy chacma baboon (Papio ursinus) sampled in South Africa (van Rensburg et al., 1998). Both studies suggest that cross-species transmission of SIV from AGMs to sympatric populations of baboons is occurring due to exposure to infected AGMs during predation. However, SIV status in olive baboons (Papio anubis) remains unknown despite reports of SIVagm exposure in its counterparts, yellow and chacma baboons.

Transmission of SIV from NHPs to humans can be influenced by the frequency of contacts and prevalence of infection in the animal host among other factors (Steve et al., 2017). In Kenya, there has been increased frequency of contacts between humans and NHPs mainly through increased illegal bush-meat trade, which involves diverse monkey species (Kimwele et al., 2012). However, investigations on SIVs distribution in NHPs found near large urban populations has been scarce. Most studies on SIVs are focused on surveillance of the virus in

African NHPs inhabiting rural and forest habitats (Ma et al., 2013; Steve et al., 2017). Nonetheless, urban-restricted NHPs can also harbour these pathogens, with the potential of cross-species transmission into naïve human population.

Therefore, this study sought to determine the prevalence and diversity of SIV strains recovered from common free-ranging NHPs in Kenya. The targeted NHPs included AGMs and olive baboons within urban and peri-urban centres in Kenya, where interaction with human is potentially high. Molecular techniques including polymerase chain reactions (PCR) coupled with high-resolution melting (HRM) analysis, gel electrophoresis and sequence analysis were used to screen blood samples from the targeted NHP species for SIV.

### 1.2 Statement of the problem

Molecular characteristics of pathogens such as SIV from free-ranging NHPs in Kenya remain poorly understood despite their immense risk of zoonotic transmission due to enhanced human-NHP interactions. This interaction usually occur during the process of butchering and preparation of bushmeat thereby increasing the risk of human infection. This phenomenon necessitates molecular detection, characterisation and phylogenetic analysis of this potentially zoonotic virus in order to determine their prevalence, diversity and its evolutionary changes. Currently, SIV prevalence has largely been determined from serological surveys on a few opportunistic samples from wild caught NHPs and those from captive animals. Comprehensive data on molecular prevalence and diversity of SIV variants circulating in common and widely distributed monkeys in East Africa is also scarce.

### 1.3 Objectives

### 1.3.1 General objective

This study investigated the prevalence and diversity of SIV strains infecting free-ranging Chlorocebus aethiops (Africa green monkeys - AGMs) and Papio anubis anubis (olive baboons) from selected urban centres in Kenya.

### 1.3.2 Specific objectives

i. To determine the prevalence of SIV in free-ranging AGMs and olive baboons inhabiting Mombasa, Naivasha and Kisumu urban environs.
ii. To determine genetic diversity of SIV in free-ranging AGMs and olive baboons in Mombasa, Naivasha and Kisumu urban environs.
iii. To identify SIV evolutionary patterns associated with host-species.

### 1.4 Hypotheses

The following were the null hypothesis for this study
i. There is no difference in SIV prevalence and genetic diversity among non-human primates found in Mombasa, Naivasha and Kisumu geographical regions.
ii. There is no genetic diversity, cross-species transmission and recombination of SIVs among sympatric primate species from urban ecological zones in Kenya.
iii. There is no SIV evolutionary patterns associated with the host-species in Kenya.

### 1.5 Justification of the study

This study proposes to bridge the knowledge gap on comprehensive molecular survey of SIV diversity in common East African monkeys, Chlorocebus aethiops (Africa green monkeys - AGMs) and Papio anubis anubis (olive baboons) found in densely populated urban centres in Kenya. Free-ranging populations of NHPs are regarded as potential sources of pathogens that can cause infectious diseases in humans. One such potential pathogen is SIV discovered in AGMs and considered as a human zoonotic virus similar to HIV (Peeters et al., 2014). Molecular analysis of SIV strains in free-ranging NHPs and their evolutionary trends are important epidemiological factors. First, frequent human-NHP interactions increase the risk of SIV transmission to humans and probably initiate new HIV or HIV-like pandemic. Secondly, extensive SIV transmission and strains co-circulation in urban-restricted NHPs may lead to virus recombination resulting into new strains capable of infecting other hosts species including humans. This potential risk can only be demonstrated if our knowledge on SIV prevalence in monkeys found near human settlements is well understood with detailed analysis of the virus genetic diversity and its evolution. Lastly, knowledge on NHP retroviral reservoir will enable us to update, adapt and improve the current diagnostic tools and thus be able to identify a wide diversity of simian retroviruses at interfaces where the emergence of new zoonotic diseases is likely to occur.

## CHAPTER TWO

## LITERATURE REVIEW

### 2.1 Emerging infections

Emerging infections (EIs) are newly recognised infections or those that previously existed, but whose incidence is rapidly increasing or expanding in geographic boundaries and host range (Morse, 2001). EIs are grouped into two, namely, emerging infectious disease (EIDs) and re-emerging infectious diseases (re-EIDs). EIDs are defined as infectious diseases whose incidence is increasing following its first introduction into a new host population or whose incidence is increasing in an existing host population (Engering et al., 2013). Examples include HIV/AIDS, Ebola haemorrhagic fever and severe acquired respiratory syndrome (SARS), Middle East respiratory syndrome (MERS) and H1N1 influenza (Swine flu) amongst others. Re-EIDs are those that had been considered eradicated or controlled, but their incidences are threatening to increase or reappear in new geographical locations or in drug resistant forms due to mutations (Morens \& Fauci, 2013). Examples of re-EIDs include multidrug resistant tuberculosis, cholera, dengue fever and polio. EIs are of global health concern, with the 2014-2016 Ebola outbreak in West Africa, and present cases in DRC demonstrating their global impact. Therefore, their control and management is of great importance.

Globally, emerging infections directly account for 15 million deaths (Wang et al., 2016), and $30 \%$ of disability adjusted life years annually (Chua \& Gubler, 2013; James et al., 2018) with up to 120 billion dollars in economic loses (Smith et al., 2019). Developing countries have been majorly impacted with the burden of morbidity and mortality, where more than 3 million children succumb annually due to EIs alone (WHO, 2016) despite the risk of EIDs/re-EIDs being globally distributed (Figure 1).

Reasons for increased risk from EIs include increasing human population (Peeters et al., 2002; Rhyan \& Spraker, 2010) that has necessitated changes in land use thus increasing humanwildlife interactions and chances of zoonotic transmission of pathogens (Pigott et al., 2015). Trade globalisation on the other hand propagates spread of the diseases (Morens et al., 2004).


Figure 1: A global map of emerging infectious diseases. The global distribution of emerging, re-emerging and deliberately emerging pathogens of viral, bacterial, fungal and protozoan origins are shown. The figures is adapted from Paules et al., (2017).

Additionally, high pathogen prevalence in animals that are in frequent interaction with humans have also been implicated as a factor enhancing zoonotic transfer of pathogens into human population (Keesing et al., 2010). Poor public health infrastructures in developing countries hamper the efforts to efficiently control and manage present and new infections. Some pathogens that have crossed species barrier and caused diseases in humans include bacteria, viruses, protozoan parasites and helminths (Bengis et al., 2004). Pathogens with high mutational rates such as RNA viruses and those capable of acquiring foreign genetic materials such as bacteria and pathogens infecting multiple hosts are more likely to turn into an emerging disease agent (Engering et al., 2013).

### 2.2 Viral emerging infectious diseases

Viruses accounts for over $50 \%$ of zoonotic pathogens with more than a third of EIs emanating from viruses with RNA genomes (Howard \& Fletcher, 2012). Ebola haemorrhagic fever, H1N1 influenza, SARS, MERS, Zika virus and HIV/AIDS are examples of viral EIs of significant public health concern and a considerable global socioeconomic impact. On average, at least two new human viruses are reported annually further increasing the existing disease burden (Woolhouse et al., 2008; 2012). Examples of newly reported viruses include human bocavirus, parvovirus 4 (Woolhouse et al., 2012), human hepegivirus 1 (HHpgV-1) and Merkel

Cell Polyomavirus (Kapoor et al., 2015), human Usutu virus (Simonin et al., 2018), porcine enteric alphacoronavirus (Fu et al., 2018) and Matryoshka RNA virus 1 (Charon et al., 2019). Emergence of pathogenic human viruses correlate to factors such as broad host range and a conserved host receptor across diverse potential hosts that permit invasion (Woolhouse et al., 2012). Bats, rodents and NHPs are important sources of zoonotic viruses (Peeters \& Delaporte, 2012; Friant et al., 2015) thus making increased human interactions with these animals a potential source of human infections. Consequently, upon zoonotic acquisition of the virus, urbanisation and increased transnational travels have enhanced human-human transmission on a global scale (Engering et al., 2013).

### 2.3 Origin of Human Immunodeficiency Virus (HIV)

Acquired immunodeficiency syndrome (AIDS) is a worldwide pandemic in humans caused by two strains of HIV virus (HIV-1 and HIV-2) suspected to have emerged from crossspecies infection of humans with SIV (Sharp \& Hahn, 2011). AIDS is one of the most devastating infectious disease in the modern world associated with 1.2 million deaths and annual losses of USD 21.7 billion (UNAIDS, 2019). Developing countries have experienced the greatest HIV/AIDS morbidity and mortality, with the highest prevalence rates recorded in young adults in sub-Saharan Africa (Dwyer-Lindgren et al., 2019). Although AIDS related deaths have been reduced by antiviral administration (Barouch, 2008; Richman et al., 2009), the realisation of curative therapy for HIV eradication is still elusive. Both HIV-1 and HIV-2 comprise several subgroups that probably emerged from independent cross-species transmission events of SIV from NHPs into human (Van Heuverswyn et al., 2007).

Despite infecting different hosts and existence of differences in some of their regulatory genes, both SIV and HIV commonly share structural genes also found in other lentiviruses thus suggesting their relatedness (Coffin, 1992). Phylogenetic analyses of various HIV subgroups have consistently shown the virus interspersing between respective SIV lineages thus suggesting the origin of HIV to be cross-species transmission of SIV from NHPs (Sharp \& Hahn, 2011). Studies have shown that over 45 species of old world monkeys are infected with SIV without pathogenic outcomes leading to conclusion that these species are natural hosts of SIV (Peeters et al., 2014). Transmission of SIV to human must have occurred through exposure to infected ape tissues and body fluids, which are common in bush-meat hunting and human wildlife conflicts (Peeters et al., 2002). Monkey bushmeat hunting and consumption has been reported in Kenya (Kimwele et al., 2012; Mwangi et al., 2016), thus implying that the risk of SIV transmission into humans is high. Recently, repeated SIV cross-species transmission into
humans leading to emergence of a divergent HIV-2 strain was documented in Central Africa (Ayouba et al., 2013). The AIDS pandemic can be managed by preventing transmission of the existing strains, while also actively monitoring SIV to avert potential risk of cross-species transmission of the virus into humans.

### 2.4 Simian immunodeficiency virus (SIV)

Simian immunodeficiency virus (SIV) is a single stranded RNA virus that naturally infects over 36 different species of non-human primates (NHPs) including apes and monkeys in Africa. SIV and HIV are closely related species of viruses with taxonomy classifying them within lentivirus genus of Orthroretrovirinae subfamily and Retroviridae family (Petropoulos, 1997). It is estimated that SIV has been infecting NHPs for over 32,000 years before being recently introduced to humans as HIV via cross-species transmission (Klatt et al., 2012).

### 2.4.1 Lineages of primate lentiviruses

Based on comparative nucleotide sequences, primate lentiviruses are categorised into different lineages (Table 1) though continued characterisation of viruses from more NHPs has resulted in an increase of these lineages.

Table 1: The major SIV lineages that have been characterised in Africa.

| Primate Species | Lineage | Reference |
| :--- | :--- | :--- |
| Chimpanzees (Pan troglodytes troglodytes) | SIVcpz | Gao et al., (1999) |
| Sooty mangabeys (Cercocebus atys) | SIVsmm | Hirsch et al., (1995) |
| African green monkeys (Chlorocebus genus) | SIVagm | Jin et al., (1994) |
| Sykes' monkeys (Cercopithecus albogularis) | SIVsyk | Locateli (2008) |
| L’Hoest monkeys (Cercopithecus lhoesti), | SIVlhoest/sun | Hirsch et al., (1995) |
| sun-tailed monkeys (Cercopithecus solatus) |  |  |
| Colobus monkey (Colobus guereza) | SIVcol | Locatelli et al., (2008) |
| Greater spot-nosed monkey | SIVgsn | Courgnaud et al., (2002) |
| De'Brazza monkey | SIVdeb | Bibollet-Ruche et al., |
|  |  | (2004) |
| Dent's Mona monkey (Cercopithecus denti) | SIVmon | Dazza et al., (2005) |
| Mustached monkey (Cercopithecus cephus) | SIVmus | Aghokeng et al., (2007) |
| Black and white colobus monkey (Colobus | SIVcol | Lauck et al., (2013) |
| guereza) |  |  |

Due to co-evolution and co-speciation with the host, SIV lineages have been observed to be host specific where closely related host species might also share related SIV lineages. Studies have demonstrated that these various lineages are approximately equidistant from each other, sharing between 40-50\% identity in the most conserved gag and pol genes, while also exhibiting a common genomic organisation encoding structural and enzymatic proteins (Klatt et al., 2012). However, these viruses also might selectively contain other unique genes such as $v p u$ and $v p x$ that are not common to all primate lentiviruses. Despite these viruses being categorised under different lineages, consistent clustering together upon phylogenetic analysis (Bibollet-Ruche et al., 2004), thus suggesting existence of an ancient SIV lineage in the distant past has implied a common evolutionary history. Existence of more than 60 species of old world monkeys necessitates continued SIV characterisation in order to understand their zoonotic potential.

### 2.4.2 Pathogenesis and transmission

Non-human primates that naturally harbour SIV do not exhibit disease conditions (Chakrabarti et al., 1991; Ma et al., 2014), while AIDS-like symptoms are observed in NHPs that do not naturally harbour the virus such as the Asian macaques (Pandrea et al., 2009). This phenomenon has been attributed to prolonged association between the virus and the host leading to evolution of a weaker immune response protecting the host against immunopathological damage (Seki \& Matano, 2012). Down-regulation of immune response during acute infection, which is readily reversed by the end of the acute phase, has been observed in NHPs (Liovat et al., 2009). Despite an overall non-pathogenicity in their natural host species, studies have revealed that SIVs can potentially cause AIDS upon accidental or experimental introduction into unnatural hosts such rhesus macaques (Apetrei et al., 2005). Pathogenicity of SIV in susceptible hosts has also been attributed to allelic variations in virus restriction factors such as Trim5 $\alpha$ in rhesus macaques (Kirmaier et al., 2010).

Within species, there are various routes for SIV transmission, with sexual route being the main mode of transmission amongst feral NHPs (Ma et al., 2013; 2014); few cases of mother to child transmission can also occur (Chahroudi et al., 2011; 2014). However, low levels of CCR5 receptors expressed in CD4+ cells confer SIV resistance to breastfeeding infants in AGMs and Mandrills (Sodora et al., 2009). Secondly, oral viral transmission is also possible in the course of cleaning wounds sustained from injuries inflicted from biting or fighting (Ma et al., 2013). Injuries sustained during aggressive confrontations amongst sympatric NHPs have also resulted in cases of cross-species transmission (Jin et al., 1994). Some apes and
monkeys harbour unique SIVs capable of replicating in human cells in vitro (Wolfe et al., 2004) suggesting potential of infecting humans. Thus, enhanced human-NHP interaction is a possible source of new human infections. Moreover, high pathogen prevalence in NHPs in addition to viral and host factors that are important in invasion, establishment of infection and consequently disease (Aghokeng et al., 2010) enhances the zoonotic risk.

### 2.4.3 Lentiviruses structure and replication mechanism

SIV is a spherical retrovirus whose structure comprises a viral envelope enclosing a dense core composed of two identical single-stranded RNA molecules, viral enzymes and structural proteins (Coffin, 1992). The viral gene is flanked by LTR (long terminal repeat) region at both 3'and 5'ends (Shah et al., 2014). The 5' LTR regions is critical for transcription of viral genes since it codes for the transcriptional promotor. The viral genetic material codes for nine genes encoding 19 proteins. Three of these genes encode for structural proteins including the groupspecific antigen (gag), envelope polyprotein (env) and the polymerase (pol) that are important for virus replication (Coffin, 1992). Gag forms the viral core protein that include the capsid protein (CA, P24), nucleocapsid (NC, P7) and outer core membrane protein (MA, P17) (Coffin, 1992). The pol polyprotein is cleaved into reverse transcriptase, integrase and protease enzymes.

Six of the remaining genes code for proteins that play regulatory roles and hence control virus tropism and replication (Faust et al., 2017). These include tat that initiates viral genes transcription, rev that facilitates mRNA translocation from the nucleus into the cytoplasm during replication the nef protein that regulates expression of surface receptors on virusinfected. The other include vif protein that enhances viral infectivity, the $v p r$ that facilitates translocation of the viral core from the cytoplasm into the nucleus and $v p u$ that enhances virion release from the host-cell (Faust et al., 2017). Other proteins include the reverse transcriptase enzyme that converts viral RNA into viral cDNA that is subsequently integrated into the host genome by integrase enzyme. This enables the virus to utilise the host's replication machinery (Luciw \& Leung, 1992). The extreme error-rate exhibited by reverse transcriptase due to lack of proof-reading activity alongside the high chances of recombination during replication of viral RNA results in high genetic diversity of retroviruses. The envelope glycoprotein is the main viral structural protein that facilitate attachment and host-cell entry.

SIV entry into host-cells is initiated by the virus binding to cluster differentiation antigen 4 (CD4) receptors on the T-cells using viral receptors on the env glycoprotein (gp120) (Petrovas et al., 2012). CCR5 is the main chemokine coreceptors that enhance SIV binding to

CD4 receptors (Edinger et al., 1999). Following attachment, the SIV env glycoprotein undergo conformational changes leading to exposure of the variable domains (V3 loop) that activates proteolytic enzymes necessary for virus and host-cell membrane fusion (Prabakaran et al., 2007). After fusion, the capsid (which contains the RNA, reverse transcriptase, proteases, ribonuclease, and integrase) is injected into the host cells’ cytoplasm (Engelman \& Cherepanov, 2012). Inside the host-cell, P24 facilitates viral RNA dissociation from the capsid which is then reverse transcribed into a double stranded cDNA molecule. The generated cDNA is translocated into the nucleus where it is integrated into the host's genome by integrase and awaits transcription (Asante-Appiah \& Skalka, 1997).

As the infected host cell transcribes its genetic material, it also transcribes the integrated viral genetic materials into mRNA (Luciw \& Leung, 1992). Earlier mRNAs to be transcribed are major viral regulatory genes that include tat, rev and nef (Faust et al., 2017). The mRNA transcripts encoding viral structural proteins such as envelope polyprotein (gp160) and the gagpol polyprotein are translated into respective proteins using the host-cell machinery. The gp160 is then translocated through the endoplasmic reticulum into the Golgi complex, where it is cleaved by cellular proteases into gp41and gp120 proteins that are subsequently glycosylated (Luciw \& Leung, 1992). As the virion matures, the gag-pol polyprotein is cleaved into individual SIV core proteins P24, P4 and P7 by viral proteases. It is during this cleavage process that reverse transcriptase, integrase and protease enzymes are also formed. Assembly of viral proteins alongside two genomic viral RNA molecules is catalysed by $v p u$ and vif on the hostcell's plasma membrane after which the mature SIV virion buds off and can infect other cells (Luciw \& Leung, 1992).

### 2.4.4 Recombination

The rates of recombination in RNA viruses such as SIV are known to be high due to extensive cross species transmission and co-infection with related strains in each species. Recombination plays a role in generation of new SIV variants in NHPs including strains with zoonotic potential (Chen et al., 1996). Divergence of recombinant strains from the representative SIV lineage strains has hampered the efforts of establishing their parental strains upon comprehensive sequence analysis (Klatt et al., 2012). The most notable examples of recombinant lentivirus strains include HIV-1 in humans and its intermediate ancestor SIVcpz found in chimpanzee (Sharp \& Hahn, 2011). Recombination of viruses from Cercopithecinae family such as SIVrcm, and SIVmus (Figure 2) is thought to have generated SIVcpz infecting chimpanzees (Bailes et al., 2003) which further evolved into HIV-1 (Sharp \& Hahn, 2010). To
successfully infect human host, SIV underwent evolutionary changes such as loss of viral protein X (VPX) followed by generation a unique viral protein R (VPR) (Figure 2) that was crucial in overcoming human restriction factors such as tetherin and TRIM5 $\alpha$ proteins (Neil et al., 2008; Roesch, 2018). Therefore, characterisation of SIV infecting NHPs from Cercopithecinae family is therefore particularly important due to their recombination potential and the capability of crossing species barriers. It has also been reported that sequential or co-infection with different HIV-1 strains in human results in recombinant viruses known as circulating recombinant forms (CRFs), which result in enhanced viral genetic diversity (Peeters et al., 2013). Enhanced genetic diversity has been implicated as the main cause of antiretroviral drug resistance whose complexity is increasing (Santoro \& Perno, 2013). The complexity of enhanced genetic diversity in SIV could also be increasing owing to recombination between SIV CRFs as seen in HIV (Peeters et al., 2013). This therefore necessitates comprehensive investigation into the genetic diversity of SIVs circulating in NHPs.


Figure 2: Proposed origin of human immunodeficiency virus (HIV). Cross-species transmission of SIVrcm from red-capped mangabeys and SIVmus from mustached monkey resulted in co-infection of the two strains in a new chimpanzee host. Both SIV from red-capped mangabeys and mustached monkey encode for their species-specific a viral infectivity protein (VPX) highlighted in red and brown respectively. Co-infection in a chimpanzee led to recombination of the two strains during replication. The VPX segment from the SIVrcm was
lost during the recombination process. Through genetic recombination, a unique viral infectivity gene (VPR) was generated in the chimpanzee host. This unique VPR gene was likely critical for viral adaptation to replicate in chimpanzees and humans by overcoming viral restriction factors such a TRIM5 $\alpha$. This figure was modified from Roesch, (2018).

### 2.4.5 Epidemiology and prevalence of SIV

Serological and molecular surveys of bush meat, wild caught primates as well as applying non-invasive sampling techniques in Africa has shown that SIV infection is widespread and prevalence may vary depending on species and geographic origin (Aghokeng et al., 2010). Wide distribution of NHPs that harbour a plethora of potentially zoonotic pathogens enhances the risk of cross-species transmission and super-infection with multiple virus strains with possibilities of virus recombination generating new strains that can infect human (Peeters et al., 2002). SIV infection has been reported to be uniformly distributed amongst feral NHPs such as AGMs, mandrills and sooty mangabeys due to high-steady state viral replication facilitating sexual transmission (Ma et al., 2014). SIV prevalence ranging between 46-50\% has been reported, and is lower in males than females (Ma et al., 2013). This is probably because most male NHPs are expected to have significantly lower exposure, as only a few attain the dominance status necessary for sexual access to breeding females (Galat-Luong et al., 1994). Studies have shown that SIV prevalence increases with age, but is uniformly low in juveniles and infants (Ma et al., 2013), thus further corroborating suggestions that sexual routes as well as aggression could play important role in SIV transmission. For instance, uneven prevalence across different age groups with $28 \%$ infants and juveniles being infected compared to $67 \%$ infection in sexually mature adults has been reported (Ma et al., 2014). High SIV prevalence could be an important factor that drives cross-species transmission, and therefore necessitates evaluation of viral prevalence in monkeys from Kenya to estimate zoonotic risk more so in regions experiencing regular human-NHP interactions.

### 2.4.6 Genetic diversity

Phylogenetic analysis has revealed high SIV genetic diversity from geographically isolated NHPs of the same species such as chimpanzees and AGMs. For instance, both West African chimpanzees (Pan troglodytes troglodytes) and East African chimpanzees (Pan troglodytes schweinfurthii) are infected with viruses belonging to the same SIV lineage (Santiago et al., 2003) though they exhibited $30-50 \%$ sequence divergence (Haesevelde et al., 1996). This divergence could have occurred due to viruses infecting the two species having evolved in isolation over time. Phylogeographic diversity of up to $19 \%$ (Bibollet-Ruche et al.,
2004) has also been reported for SIVdeb despite these strains clustering together phylogenetically, hence suggesting existence of intra-clade divergence. Analysis of the $v p u$ gene in SIVden virus infecting Dent's Mona monkey revealed a significant divergence from other virus lineages known to harbour this gene where SIVdens' $v p u$ was found to be truncated at its C-terminal end (Dazza et al., 2005). Moreover, the env genes in SIVden were also shown to overlap as opposed to the genes being separate in HIV-1/SIVcpz lineage (Dazza et al., 2005). Mustached monkeys were also shown to be infected by multiple genetically distinct SIVmus strains, one being a recombinant between SIVmus and SIVgsn from cohabiting greater spot nosed monkeys while the second strain did not cluster with previously reported SIVmus strains (Aghokeng et al., 2010).

Wide genetic diversity of SIV and functional motifs variations suggests a complex evolutionary history and strong indication that co-infections are common. Super-infection of NHPs with genetically diverse virus strains is a potential avenue for recombination events that can lead to emergence of new viruses that could infect humans (Smith et al., 2005). Newly uncharacterised viruses are of particular interest as their zoonotic risks are unknown. Recent discovery of HIV-1 subtype P isolated from Cameroonian patients in 2009 (Peeters et al., 2013) and the possible new groups of HIV-2 reported in 2013 (Ayouba et al., 2013) clearly reveals that our knowledge on lentivirus genetic diversity and cross-species transmission is still incomplete. Hence, there is need for wider geographical sampling of NHPs and determination of prevalence and molecular characteristics of the viruses they harbour.

### 2.5 Non-human primates as a source of emerging infections

Close synanthropic interaction between humans and animals regarded as potential sources of zoonotic pathogens increases the risk of human retroviruses emergence (Wolfe et al., 2004; Peeters \& Delaporte, 2012). Humans, being closely related in physiologic and genetic characteristics to NHPs, (Wolfe et al., 1998) can therefore be equally susceptible to many pathogens found in NHPs. Pathogens infecting NHPs such as bacteria, haemoprotozoan parasites, ectoparasites and viruses are known to be closely related to human pathogens (Wolfe et al., 1998). Significantly high prevalence of zoonotic pathogens such as African trypanosomes, piroplasms and even leishmania have previously been reported in wild caught NHPs including baboons and AGMs from Kenya (Jeneby et al., 2002; 2011). On the other hand, studies have revealed that apes are infected with a plethora of diverse Plasmodium species closely related to malaria-causing Plasmodium falciparum (Krief et al., 2010).

Phylogenetic analysis of these strains seemed to suggest that Plasmodium infecting humans could be having a simian origin (Liu et al., 2010).

In general, over half of emerging pathogens in humans are viruses and a large number of them have been isolated from NHPs (Jones et al., 2008). Over the last decade, viral diseases associated with African NHPs as reservoirs or incidental hosts have emerged as major disease outbreaks in humans (Wolfe et al., 2007). These include a series of Ebola virus outbreaks in West and Central Africa (Saéz et al., 2015) and a case of Marburg zoonotic infection (Nyakarahuka et al., 2017) in Uganda, East Africa. High prevalence of viruses such as SIV and simian T-cell lymphotropic virus (STLV) has been documented in many species of old world monkeys (Aghokeng et al., 2010; Liégeois et al., 2012) thus increasing the risk of zoonotic transmission into humans. Zoonotic transfer of these pathogens occurs through exposure to infected tissues and fluids (Peeters et al., 2002). This therefore makes NHPs important species for sentinel surveillance for emergence of pathogens that can potentially cross the species barrier into humans. Additionally, bats that also share habitats with free-ranging NHPs have been identified as the potential source of Ebola virus and Nipah virus (Friant et al., 2015; Saéz et al., 2015). Therefore, it is important to determine the molecular characteristics of these zoonotic pathogens in their natural host to enable prediction of cross-species transmission. Finally, failure of standard assays to detect newly emerged pathogens in humans (Peeters et al., 2002; Ndembi et al., 2009) calls for continued characterisation of zoonotic pathogens to facilitate development of specific assays capable of detecting potentially new infections emerging from zoonotic events before they can progress to the epidemic magnitudes (WHO, 2004).

Therefore, this study determined prevalence, and molecular characteristics of SIV in freeranging NHPs that are in close proximity to large human populations in Mombasa, Naivasha and Kisumu urban centres with the aim of estimating the magnitude of zoonotic risk due to human exposure to these NHPs.

## CHAPTER THREE

## MATERIALS AND METHODS

### 3.1 Ethics statement and permits

The Institutional Scientific Review Committee (ISERC) at the Institute of Primate Research (IPR), Kenya, reviewed the experimental design and methods used in this study. Authorisation permit (ISERC/04/18) for in situ sampling of wildlife species was acquired from Kenya Wildlife Service (KWS). See appendix for the permit details.

### 3.2 Summary of experimental design

This was a cross-sectional study. NHPs were captured with care to ensure minimal bruises or injuries and sampled once before they were released back into their natural habitats. The experimental procedures carried out in this study are summarised in Figure 4.


Figure 3: Summary of the experimental pipeline for the study. A workflow showing the sequence of various tasks and activities undertaken in the study

### 3.3 Sampling sites

The urban and peri-urban centres targeted for sampling were Mombasa, Kisumu and Naivasha in Kenya (Figure 3) where urban-dwelling NHPs can be found. The sampling sites in Mombasa Island were the Tudor water sports (TWS) facility located northwest of the Island, Mbaraki (MBK) in the south of the island, and government administration compounds (GAC) along the eastern parts of the Island. Also targeted was Mazeras ( $3^{\circ} 58^{\prime} \mathrm{S}, 39^{\circ} 33^{\prime} \mathrm{E}$ ) on the mainland and within the outskirts of Mombasa town. In Kisumu ( $0^{\circ} 6^{\prime} \mathrm{S}, 34^{\circ} 45^{\prime} \mathrm{E}$ ), a city located along the shores of Lake Victoria in Western Kenya three sites were targeted. These are the Impala park zoo within the town centre, residential golf course (GC) area near the Kisumu international airport and Kajulu (KAJ), a peri-urban centre located 5 km from the city. Kimana area, an eco-tourism destination at the shores of Lake Naivasha was the only sampling site considered in Naivasha $\left(0^{\circ} 43^{\prime} \mathrm{S}, 36^{\circ} 26^{\prime} \mathrm{E}\right)$, located in the Great Rift Valley.


Figure 4: Urban and peri-urban sampling sites. Map of Kenya (center) showing the geographical locations of the three study urban and peri-urban centres (red circles) namely Kisumu, Mombasa and Naivasha where the non-human primates were trapped. The three sampling centres are shown offset with sampling sites represented by the green circles.

### 3.4 Sampling of NHPs

Free ranging NHPs were baited with fresh carrots, bananas and maize cobs and captured using wire mesh cages fitted with self-triggering slide doors as previously described (Jeneby et al., 2011). Individual animals were anaesthetised by administration of xylazine (2\%): ketamine hydrochloride (10\%) (Sigma Aldrich, Darmstadt, Germany) in the ratio of 1:3 and a dosage of $0.1 \mathrm{ml} / \mathrm{kg}$ body weight. Age was determined using dental eruption (Akinyi et al., 2013) before blood was collected once in situ. Whole blood ( 10 ml ) was drawn from the femoral vein by venipuncture using 21 gauge needle and 10 ml syringe by an authorised wildlife veterinarian from KWS. The 21 gauge needle was chosen according to the body size of the NHPs captured. A bigger sized needle ( $<21$ gauge) was avoided to minimize injury to the animal while drawing blood, whereas a smaller needle (>21 gauge) was avoided to minimize hemolysis. An aliquot ( 5 ml ) of each blood sample was preserved in sterile vacutainer tubes with EDTA as anticoagulant ( 10 ml BD vacutainer ${ }^{\ominus}$ K2E (EDTA, BD-Plymouth UK), while the other 5 ml was preserved in serum tubes ( 6.0 ml BD vacutainer ${ }^{\circ}$, BD-Plymouth UK). The blood samples were transported in dry ice to the laboratory and stored at $-80^{\circ} \mathrm{C}$. Two drops of blood were spotted onto Whatman FTA cards (Sigma Aldrich, Darmstadt, Germany) (Strøm et al., 2014), air dried and stored at room temperature. For infants, drops of blood from sterilised fingertips was used to alleviate stressful venipuncture sampling. These drops of blood were spotted onto Whatman FTA cards, air dried, and stored at room temperature. Sampled animals were monitored until they regained full consciousness before release.

### 3.5 DNA extraction

Total viral genomic material (viral RNA and proviral DNA) was extracted from $200 \mu \mathrm{~L}$ of each EDTA-blood sample using MagNA 96 Pure DNA and Viral NA Small Volume Kit (Roche Applied Science, Penzberg, Germany) in a MagNA Pure 96 (Roche Applied Science, Germany) automated extractor as previously described (Villinger et al., 2017). Briefly, $200 \mu \mathrm{~L}$ of each EDTA blood samples was transferred into MagNA pure 96 processing cartridge (Cat No. 06241603 001). The cartridge was then loaded into the MagNA Pure 96 automated extractor and programmed to elute the recovered genomic materials in $50 \mu \mathrm{~L}$ elution buffer.

### 3.6 Molecular identification of SIVs

Screening for SIV was done by nested PCR-HRM analysis targeting the proviral DNA (Ayouba et al., 2015) to detect and differentiate primate lentiviruses naturally infecting NHPs. The primary amplification was done using NDR1-POLOR1 primer set (Figure 4) in a Veriti ${ }^{\mathrm{TM}}$ thermocycler (Life technologies holdings PTE LTD, Singapore). Each $25 \mu \mathrm{~L}$ reaction
contained $2.5 \mu \mathrm{~L}$ of 10x standard buffer (NEB Inc., Massachusetts, USA), $0.5 \mu \mathrm{~L} 10 \mathrm{mM}$ dNTPs, $0.5 \mu \mathrm{~L}$ of $10 \mu \mathrm{M}$ forward primer and reverse primer each and $0.125 \mu \mathrm{~L}$ of 5 U Taq polymerase (New England Biolabs Inc., Ipswich, Massachusetts, United States), $1 \mu \mathrm{~L}$ of template, and topped up with nuclease free PCR water. Amplification conditions included an initial denaturation at $95^{\circ} \mathrm{C}$ for 3 min followed by 35 cycles at $95^{\circ} \mathrm{C}$ for 30 secs, $45^{\circ} \mathrm{C}$ for 30 secs, and $72^{\circ} \mathrm{C}$ for 2 min and a final extension at $72^{\circ} \mathrm{C}$ for 10 min . Secondary PCRs were performed in a Rotor-Gene Q thermocycler (QIAGEN, Germany) using Polis4/Uni2 primers targeting a 650-nucleotide (nt) fragment pol gene broadly conserved in lentiviruses. The $10 \mu \mathrm{~L}$ reaction volumes contained $2 \mu \mathrm{~L}$ of 5X HRM EvaGgreen Mastermix (Solis Biotype, Riia Estonia), $0.5 \mu \mathrm{~L}$ of each $10 \mu \mathrm{M}$ primer and $1 \mu \mathrm{~L}$ DNA template as described by (Villinger et al., 2017). Amplification conditions included an initial denaturation at $95^{\circ} \mathrm{C}$ for 15 min followed by 35 cycles at $95^{\circ} \mathrm{C}$ for 30 secs, $55^{\circ} \mathrm{C}$ for 30 secs, and $72^{\circ} \mathrm{C}$ for 30 secs and a final extension at $72^{\circ} \mathrm{C}$ for 5 min . Subsequent HRM analysis was conducted as previously described (Villinger et al., 2017). Briefly, PCR products were subjected to further denaturisation at 95 ${ }^{\circ} \mathrm{C}$ for 1 min , annealing at $40^{\circ} \mathrm{C}$ for 1 min and equilibrating at $75{ }^{\circ} \mathrm{C}$ for 90 secs, and then increasing the temperature in $0.1^{\circ} \mathrm{C}$ increments up to $90^{\circ} \mathrm{C}$, with fluorescence acquisition after 2 secs incremental holding periods. Distinct HRM profiles, normalized in the range of $80-90^{\circ} \mathrm{C}$, were visually determined for each reaction after completion of HRM data acquisition.


Degenerate nucleotides: $\mathrm{R}=\mathrm{A}$ or $\mathrm{G} ; \mathrm{Y}=\mathrm{C}$ or $\mathrm{T} ; \mathrm{B}=\mathrm{G}$ or $\mathrm{T} ; \mathrm{W}=\mathrm{A}$ or $\mathrm{T} ; \mathrm{H}=\mathrm{A}, \mathrm{C}$, or T and $\mathrm{N}=\mathrm{A}, \mathrm{C}, \mathrm{G}$ or T

Figure 5: SIV genome architecture and the PCR amplification strategy. A. A cartoon representing SIV genome architecture showing respective genes. The genes targeted for amplification are shown by the horizontal bars and include pol, RT-integrase and two regions of gp120. The primers flanking the target gene fragments are represented by the alphabets (AH) at the terminal ends of each fragment. B. The primers used for amplification include A (NDR1) and B (POLOR1) targeting a 2.7 kb of the pol gene fragment, primer sets C (Polis4) and D (Uni2) used in nested PCR to amplify the 650 bp fragment of integrase gene within the pol gene. The primer sets $\mathrm{E}(\mathrm{ENVA})$ and F (ENVB) targeting a 1 kb fragment of the env gene while nested PCR using primer sets G (ENVC) and H (ENVD) amplified a 900bp partial env gene. (LTR - Long terminal repeat, gag - gene specific antigen, PR - protease, RT- reverse transcriptase, INT - integrase, env - envelope).

SIV strains were characterised by amplifying a 650 nt pol gene fragment (spanning reverse transcriptase and integrase regions) and a 900 nt envelope (env) gp120 fragment encompassing V3-V5 regions. The pol gene (Figure 4) was amplified using NDR1-POLOR1 primers for primary amplification, followed by nested PCR using the Polis4/Uni2 primers (Figure 4) in a Veriti ${ }^{\mathrm{TM}}$ thermo cycler (Life technologies holdings PTE LTD, Singapore). The reaction composition and amplification conditions for the primary reaction are as described in section 3.2. Each $25 \mu \mathrm{~L}$ secondary reaction contained $2.5 \mu \mathrm{~L}$ of 10 x standard buffer (NEB Inc., Massachusetts, USA), $0.5 \mu \mathrm{~L} 10 \mathrm{mM}$ dNTPs, $0.5 \mu \mathrm{~L}$ of $10 \mu \mathrm{M}$ forward primer and reverse primer each and $0.125 \mu \mathrm{~L}$ of 5U Taq polymerase (NEB Inc., Massachusetts, USA). The primary PCR products ( $1 \mu \mathrm{~L}$ ) was used as the template for secondary PCR amplification. The, cycling conditions applied were denaturation at $95^{\circ} \mathrm{C}$ for 3 min , followed by 35 cycles at $95^{\circ} \mathrm{C}$ for 30 secs, annealing was done at $55^{\circ} \mathrm{C}$ for 30 secs, extension at $72^{\circ} \mathrm{C}$ for 2 min and a final extension was done at $68^{\circ} \mathrm{C}$ for 10 min .

For further characterisation of the env gp120 gene, we amplified longer fragments by nested PCR (Ma et al., 2013) using the ENVA/ENVB (E-F) primers for the primary amplification and ENVC/ENVD (G-H) primers for the second nested amplification (Figure 4). Each $20 \mu \mathrm{~L}$ PCR mixture for primary amplification contained; $10 \mu \mathrm{~L} 2 \mathrm{x}$ Multiplex PCR Plus Kit Mastermix (QIAGEN, Hielden, Germany), $2 \mu \mathrm{~L}$ Q-Solution and $0.5 \mu \mathrm{~L}$ of $10 \mu \mathrm{M}$ each forward and reverse primers, $5 \mu \mathrm{~L}$ of nuclease free PCR water, and $2 \mu \mathrm{~L}$ of template DNA. Nested amplifications also contained the same PCR components as described above but $1 \mu \mathrm{~L}$ of the primary amplicon was used as template. Amplification reactions were done in a SimpliAmp ${ }^{\text {TM }}$ thermocycler (Applied Biosystems, Branchburg, NJ). The cycling conditions included an initial incubation at $95^{\circ} \mathrm{C}$ for $5 \mathrm{~min}, 35$ cycles of extension at $95^{\circ} \mathrm{C}$ for 30 secs,
primer annealing at $55^{\circ} \mathrm{C}$ for 30 secs, extension at $72^{\circ} \mathrm{C}$ for 1 min and final extension at $68^{\circ} \mathrm{C}$ for 10 min for both primer sets. Nested PCR products were visualised under UV after electrophoresis on a $1.5 \%$ agarose gel stained with $0.5 \mu \mathrm{M} / \mu \mathrm{L}$ ethidium. Amplicons with the expected fragment size of 650 nt and 900 nt were purified using ExoSAP-IT protocol (USB® Products AffyInc., Ohio, USA) as described below.

### 3.7 Purification of PCR amplicons

Representative samples that generated the expected fragment size of 650bp and 900bp were purified using ExoSAP-IT protocol (USB® Products AffyInc., Ohio, USA) as described by the manufacturer. Briefly, $5 \mu \mathrm{~L}$ of post-PCR reaction products were mixed with $2 \mu \mathrm{~L}$ of ExoSAP-IT ${ }^{\text {TM }}$ reagent in sterile 0.2 mL PCR tubes. The mixture was vortexed briefly and centrifuged before incubation at $37^{\circ} \mathrm{C}$ for 15 min . The reaction was terminated by incubation at $85^{\circ} \mathrm{C}$ for 15 min , and the resulting purified samples submitted for sequencing. This service was outsourced at Macrogen ${ }^{\text {TM }}$ (Seoul, South Korea).

### 3.8 Phylogenetic and evolutionary selection analyses

### 3.8.1 Phylogenetic analyses

Consensus sequences generated using Geneious v10 (Kearse et al., 2012) were aligned using MUSCLE (Edgar, 2004). Phylogenetic reconstruction was performed by the maximum likelihood approach of PhyML (Guindon \& Gascuel, 2003). The robustness of phylogenetic trees' topology was ascertained by bootstrapping using 1000 replicates. Trees were visualized using iTOL v3 (Letunic \& Bork, 2016).

### 3.8.2 Analyses of evolutionary selection

Analysis of the evolutionary selection pressure on the SIV env gene was carried out using various statistical methods implemented on the adaptive evolution server (www.datamonkey.org) (Delport et al., 2010). These methods included single likelihood ancestor counting (SLAC), Fixed Effects Likelihood (FEL) (Pond \& Frost, 2005), Fast Unconstrained Bayesian Approximation (FUBAR) (Murrell et al., 2013), mixed effects model of evolution (MEME) (Murrell et al., 2012), and Internal Fixed Effects Likelihood (IFEL) (Pond et al., 2006). FEL and SLAC methods combine maximum-likelihood (ML) and counting approaches to infer nonsynonymous (dN) and synonymous (dS) substitution rates on a per-site basis for a given coding alignment and corresponding phylogeny while assuming selection pressure for each site is constant along the entire phylogeny. However, FEL uses Likelihood Ratio Test, to ascertain if dN is significantly greater than dS while SLAC ascertains the
significance at each site using an extended binomial distribution approach. Under MEME, the hypothesis that individual sites are subject to episodic positive or diversifying selection is tested using a mixed-effects maximum likelihood approach. On the other hand, FUBAR infers the substitution rates per site of the provided codon alignment using the Bayesian approach with the assumption of constant selection pressure on the entire phylogeny. For SLAC, MEME, IFEL and FEL, results obtained were considered significant at 0.05 level of significance while for FUBAR, a posterior probability > 0.9 was deemed significant.

## CHAPTER FOUR

## RESULTS

### 4.1 Species of monkeys and their distribution

One hundred and twenty-four urban restricted AGMs (53 females, 71 males) and 65 (21 females, 44 males) peri-urban olive baboons of different age groups were trapped with care to minimise bruises (Table 2) and sampled under anaesthesia. Troops of AGMs were found within urban peri-urban areas while baboons were exclusively located in peri-urban sites. The ages of the captured NHPs were estimated based on external features such as body carriage, dental eruption and pelage condition by an experienced wildlife veterinarian (Akinyi et al., 2013). The sampled AGMs included 72 adults, 36 subadults and 16 infants while olive baboons included 48 adults, 6 subadults and 13 infants.

Table 2: The distribution and demographic of non-human primates (NHPs) sampled from the various sites of urban and peri-urban centre namely Mombasa, Naivasha and Kisumu in Kenya.

| Sampling <br> Towns | AGMs |  |  |  | Olive baboons |  |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Sampling Sites | Infants | Sub-adults | Adults | Infants | $\begin{aligned} & \text { Sub- } \\ & \text { adults } \end{aligned}$ | Adults |
| Kisumu |  | T[M/F] | T[M/F] | T[M/F] | T[M/F] | T[M/F] | T[M/F] |
|  | Kajulu | - | 10[4/6] | 11[7/4] | - | 2[2/0] | - |
|  | Golf Club | - | 1[1/0] | 4[3/1] | - | - | - |
|  | Impala Park | 1[1/0] | 10[7/3] | 14[8/6] | - | - | - |
|  | Subtotal | 1[1/0] | 21[12/9] | 29[18/11] |  | 2[2/0] | 8[6/2] |
| Naivasha | Kimana | - | 4[2/2] | 18[11/7] | - | - | - |
| Mombasa | Mbaraki | 6[3/3] | 8[1/7] | 9[5/4] | - | - | - |
|  | Tudor | 3[2/1] | 2[1/1] | 7[5/2] | - | - | - |
|  | GAC | 6[3/3] | 1[1/0] | $9[6 / 3]$ | - | - | - |
|  | Mazeras | - | - | - | 13[4/9] | 4[3/1] | 38[24/14] |
|  | Subtotal | 15[8/7] | 11[3/8] | 25[16/9] | 13[4/9] | 4[3/1] | 38[24/14] |
|  | Grand total | 16[9/7] | 36[17/19] | 72[45/27] | 13[4/9] | 6[5/1] | 46[30/16] |

Key: $\mathbf{T}=$ Total, $\mathbf{M}=$ Male, $\mathbf{F}=$ Female

### 4.2 Prevalence of SIV in AGMs and baboons

SIV infection status was determined using a combination of PCR-HRM of the partial pol gene and nested PCR amplification of partial env gene. The fragment sizes of amplified PCR products were confirmed by agarose gel electrophoresis (Figure 6) and sequencing.



Figure 6: Detection of SIV in NHPs by a combination of PCR-HRM and gel electrophoresis. A. Representative melt peaks (left panel) and HRM melt profiles (right panel) generated by PCR-HRM of the partial pol gene ( 650 bp ) from SIV positive samples. B. The amplified and electrophoresis size separated SIV partial pol gene (left panel) and env gene (right panel) fragments of 650 bp and 900 bp respectively. L represents the GelPilot 100 bp Plus molecular weight ladder (QIAGEN, Hielden, Germany) ; + and - represent SIV positive and negative controls respectively.

From these amplifications, SIV prevalence was $32 \%$ (39/124) in AGMs and 3\% (2/65) in baboons (Table 3). Infection rates in male AGMs was $34 \%$ ( $24 / 71$ ) and females $28 \%$ ( $15 / 53$ ). Two male baboons, one from Kajulu in Kisumu and the other from Mazeras, in Mombasa were positive for SIV. Prevalence of SIV among age groups of NHPs was higher in adults ( 27 out of $72,37 \%$ ), followed by sub adults ( 9 out $36,25 \%$ ) and infants ( 3 out of $16,19 \%$ ) (Table 3 ). The distribution of SIV infections in different sampling locations within urban and peri-urban location was uneven and ranged from $8 \%$ to $48 \%$ in AGMs (Table 3). Infections by sampling sites was highest in Mbaraki, Mombasa (10/22, 48\%) followed by Kimana in Naivasha (17/51, $33 \%$ ) and Impala park in Kisumu (7/25, 28\%). The AGMs sampled in Tudor, Mombasa has the lowest SIV prevalence of $8 \%(1 / 12)$ (Table S1. Appendix).

Table 3: Prevalence of SIV infection in NHPs (AGMs and olive baboons) sampled from Kisumu, Naivasha and Mombasa towns in Kenya. Prevalence of SIV infection in NHPs (AGMs and olive baboons) sampled from Kisumu, Naivasha and Mombasa towns in Kenya.

|  | SIV positive /Total sampled (\% Infected) |  |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: | :---: | :---: |
|  | Male | Female | Total | Male | Female | Total |
| Kisumu | $10 / 31(32)$ | $2 / 20(10)$ | $12 / 51(24)$ | $1 / 8(13)$ | $0 / 2(0)$ | $1 / 10(10)$ |
| Mombasa | $8 / 27(30)$ | $9 / 24(38)$ | $17 / 51(33)$ | $1 / 36(3)$ | $0 / 19(0)$ | $1 / 55(1)$ |
| Naivasha | $6 / 13(46)$ | $4 / 9(44)$ | $10 / 22(45)$ | - | - | - |
| Total | $\mathbf{2 4 / 7 1 ( 3 4 )}$ | $\mathbf{1 5 / 5 3 ( 2 8 )}$ | $\mathbf{3 9 / 1 2 4}(\mathbf{3 1 )}$ | $\mathbf{2 / 4 4}(\mathbf{5})$ | $\mathbf{0 / 2 1}(\mathbf{0})$ | $\mathbf{2 / 6 5 ( 3 )}$ |

### 4.3 High resolution melting profiles

HRM analysis of the PCR amplified SIV pol gene demonstrated distinct melt profiles (Figure 6A) illustrating either diversity of detected viruses or unknown product of the predicted sizes. These results most likely illustrate virus diversity but must be supported by empirical evidence from amplicon sequencing. The normalization regions used for the analysis of the amplified pol gene ranged from $82.0^{\circ} \mathrm{C}$ to $82.14^{\circ} \mathrm{C}$ in the leading range and $85.86^{\circ} \mathrm{C}$ to $86.0^{\circ} \mathrm{C}$ in the trailing range. Positive virus amplification from specimens MG590127-AGM675 and MG590131-AGM699from NHPs sampled in Naivasha resulted in two distinct melt profiles (Figure 6A) suggesting existence of virus divergence. In contrast, SIV isolates from MG590129-AGM682 and MG590127-AGM675 from Naivasha generated similar melt peaks but distinct melt profiles as well further illustrating existence of diversity. Amplifications of DNA specimens from MG590133-AGMKis18 from Kisumu, MG757169 AGMGAC6 from Mombasa, and MG590135 AGMMbk11 from Mombasa had distinct melt profiles from each other. The plausible divergence was confirmed by sequencing the amplicons and querying them against GenBank database (Clark et al., 2016). This generated hits of $82 \%-84 \%$ identity to the already characterised SIVagm strains (Figure 7). Generally, there was nucleotide sequence plasticity in this gene fragment upon sequence alignment also demonstrated by the percent identities of the gene products i.e. protein sequences (Table S2).


Figure 7. Multiple sequence alignment for the $\mathbf{6 5 0} \mathbf{~ b p}$ partial pol gene amplified by PCR-HRM showing the distribution of conserved nucleotides. Amplified and sequenced gene fragments of SIV pol gene from AGMs were aligned, showing a range of between 75 to $83 \%$ identity to our reference SIVagm sequences previously characterised from Kenya. The conserved nucleotides are highlighted with a black background and an asterisk (*) at the top of the alignment. Characterised SIVagm sequences from South Africa, West Africa and previously characterised sequences from Kenya were included in the alignment. The sequence identities including their accession numbers and the sampling region are shown on the left side of the alignment. Sequences were aligned using MUSCLE and the alignments visualised and exported for printing on ESPript (Robert \& Gouet, 2014) and modified on Adobe Illustrator.

### 4.4 SIV genetic diversity

### 4.4.1 Identification of SIV sequence lineages

A stringent basic local alignment search (BLAST) (Altschul et al., 1990) at an e- value of 0.0001 was used at NCBI's GenBank database (www.ncbi.nlm.nih.gov) to identify the strains of SIV sequences generated in this study. BLAST search using partial pol nucleotide sequences revealed ' $81-93 \%$ ' sequence identity to the published SIVagm pol gene from AGMs isolated in Africa while the partial env nucleotide sequences showed identity of ' $79-85 \%$ ' to "simian immunodeficiency virus isolate 9649env05" (GenBank Accession U93806) also isolated from an AGM. Further, these identities were confirmed by performing sequence alignments alongside reference sequences from GenBank.

Generally, taxonomic classification of SIV into strains/lineage correlate with their respective primate host species (Foley et al., 2016). In addition, significant diversity has been documented in SIVs from the same NHP host from different geographic locations (Ma et al., 2013; 2014). Therefore, this diversity pose a limitation to the criterion of grouping SIV according to the host species. This is further complicated when cross-species transmission or co-infection with multiple strains occur. To interrogate the lineage identification further, we first determined the SIV sequence identity ranges for viruses from GenBank isolated from various NHPs species in different geographic locations namely West, Central, Eastern and Southern Africa.

These sequences included SIV isolated from 18 NHP species including chimpanzees (Pan troglodytes), sooty mangabeys (Cercocebus atys), mandrills (Mandrillus sphinx), colobus monkey (Colobus guereza), monkeys (Cercopithecus albogularis), De Brazza's monkeys (Cercopithecus neglectus), greater spot nosed monkeys (Cercopithecus nictitans), mona monkeys (Cercopithecus mona), moustached monkey (Cercopithecus cephus) and Africa green monkeys (Chlorocebus aethiops). The separate multiple sequence alignments for partial pol and env gene fragments were used to generate percent identity matrices for each species from different regions. Resultant matrices were used to derive SIV identities range per strain per region (Figure 8). The SIVs derived from AGMs (SIVagm) are widely distributed in Africa. Their sequence identities ranged from 81-98\% in West Africa, $81 \%$ - $100 \%$ in Eastern Africa and $82-90 \%$ in Southern Africa. Only one SIVagm sequence was available from monkeys in Central Africa region, explaining the $100 \%$ identity observed. The comparison of SIVagm sequences from different regions in Africa showed the identities ranging from 71\%-83\%.

These results supported the observation of nucleotide conservation plasticity upon alignment of SIV sequences amplified in NHPs from Kenya in this study (Figure 7). As illustrated in Figure 8, colobus monkeys had the lowest range of SIV pol identities ( $52 \%-60 \%$ ). In general, the identities of SIV sequences within an NHP species ranged between $70 \%$ and $85 \%$. Amongst the 18 NHP species from which the SIV sequences were analysed, only the AGMs had representatives from all the four regions.

B


| Suntailed |  |  |  |  |
| :--- | :---: | :---: | :---: | :---: |
|  | West | Central | East | South |
| West | - | - | - | - |
| Central | - | 100 | - | - |
| East | - | - | - | - |
| South | - | - | - | - |

Figure 8: The SIV percentage sequence identity matrices generated from sequence comparison. A. Range of percentage identity of SIV partial pol gene and B. env gene fragments from different NHP species from different geographical regions (West, Central, East and Southern) in Africa. The SIV sequences were grouped according to the NHP host species and the geographical region of isolation. The sequence alignments were first compared within the NHP species within the same region and then across different regions. The percent identities
are recorded in the matrices as a range from the lowest to the highest identity observed for every sequence alignment done.

The results for the partial env gene were comparable to the partial pol gene. The identities for AGMs ranged from $71 \%-79 \%$ in West African region, $67 \%-99 \%$ in Eastern Africa and 79\% -97\% in Southern Africa. Likewise, only one sample was available from Central Africa thus accounting for $100 \%$ identity observed in this region. For Central African suntailed monkeys and drill monkeys, single env gene sequence was available for each NHP species, representing the observed $100 \%$ identity. There was a significant difference in the identity ranges for the mona monkey ( $52 \%-72 \%$ in the env gene and $67 \%-100 \%$ in the pol gene) from Central Africa suggesting a higher diversity.

Once the identities range was determined (Figure 8), representative SIV sequences from different NHP species were aligned with the SIV sequences recovered from AGMs in Kenya and used to generate percent identity matrices for partial pol and env gene fragments. From the percentage identities matrices, separate heat maps were generated for the partial gene fragments (Figures 9 and 10). Generated heat maps enabled placement of SIV sequences from this study into SIVagm lineage as shown in heat maps. The highest identities are highlighted in green thus, the sequences from this study were very close to AGM derived viruses than the mandrills and chimpanzees whose identities to AGM derived viruses was low ( $41-42 \%$ ) as shown by the red highlights of the heat map.

From the study, the identities for SIVs to other characterised AGMs' isolates ranged from $75 \%$ to $83 \%$ for the partial pol gene and $62 \%$ to $80 \%$ for the partial env gene. Comparing SIV isolated from AGMs and Sykes in Kenya, whose geographic source is the same, the identities ranged from $63 \%$ to $70 \%$. Moreover, in this study, the SIV sequenced from an archived sample of a Sykes monkey from IPR (MG590108.1_SIVsyk) had 64\% to 65\% identity to viruses from AGMs while the identity was $77 \%$ to $86 \%$ when compared with viruses from Sykes monkeys in GenBank (Figure 10). Comparing sequences derived from AGMs with other NHP species, the identities ranged from $63 \%-72 \%$ against Chimpanzees, $63 \%-68 \%$ against gorillas, $66 \%$ - $70 \%$ against greater spot-nosed monkey, $58 \%-63 \%$ against colobus monkey, $65 \%-70 \%$ against mona monkey, $65 \%-71 \%$ against sooty mangabeys and $68 \%-73 \%$ against the redcapped mangabeys. The SIV sequence identities remained marginally low amongst the different NHP species with overlapping geographic ranges. For instance, between the redcapped mangabeys and the mustached monkeys from Central Africa, the identities ranged
between $66 \%-77 \%$, colobus monkeys and the debrazza's monkeys had an identity range of $60 \%-66 \%$ while the chimpanzee and colobus monkeys had an identity range of $55 \%-62 \%$. SIV sequences from evolutionarily related chimpanzee and gorilla had a range of $76 \%-80 \%$. The plausible explanation for this high identity could be that SIVs infecting gorillas originated from chimpanzees (Takehisa et al., 2009). Collectively, these findings reveal highest identity of the sequences from this study to AGM derived SIVs. This observation indicates that the SIV amplified from AGMs qualify for classification within the SIVagm lineage. However, the interpretation of these identities needs careful consideration in cases of cross-species transmission and/or recombination. For instance, SIVs from red-capped mangabeys and chimpanzee shared $76 \%-81 \%$ nucleotide identities for the partial pol gene. Notably, the env fragments for red-capped mangabeys were unavailable for comparison. These pol identities could probably be emanating from the possibility of SIV in chimpanzee being a recombinant involving the viruses from red-capped mangabeys (Bailes et al., 2003). This therefore necessitated phylogenetic analysis to determine the evolutionary relationship of SIVs within the AGM lineage.


Figure 9. Percentage identity matrix of SIV partial pol sequence alignments for SIV lineage identification. Sequences of pol gene from SIV identified in the study were aligned against SIV sequences retrieved from GenBank database originating from different non-human primate species from Africa. The alignment was done using MAFFT run in Geneious software v10 (Kearse et al., 2012). The percent identities are shown in the table with the heat map indicating the identities of SIV sequences in a descending order with green representing most identical sequences and red showing the least identical sequences.


Figure 10. Percentage identity matrix of SIV partial env sequence alignments for SIV lineage identification. Sequences of env gene from SIV identified in the study were aligned against SIV sequences retrieved from GenBank database originating from different non-human primate species from Africa. The alignment was done using MAFFT run in Geneious software v10 (Kearse et al., 2012). The percent identities are shown in the table with the heat map indicating the identities of SIV sequences in a descending order with green representing most identical sequences and red showing the least identical sequences.

### 4.4.2 Phylogenetic analyses of SIVagm infecting AGMs and baboons in Kenya

All samples determined to be SIV positive by PCR-HRM were amplified using primers targeting pol ( 19 samples) or env ( 22 samples) genes and sequenced. Out of these, 14 samples generated unambiguous sequences for pol and 17 for $e n v$ gene fragments respectively. These sequences have been deposited in GenBank under accession numbers MG590108 MG590139, MG757169-MG757171(Table S5). Phylogenetic analysis revealed that the partial pol sequences obtained in this study clustered in a monophyletic clade with those from SIVagm isolates previously obtained from Kenya (Figure 11). No clear segregation of the

Kenyan isolates of SIVagm according to the respective sampling towns was detected. Within the Kenyan clades, sub clusters had robust bootstraps support values greater than $90 \%$. One isolate from an AGM, MG590133-AGMKis18, sampled from Kisumu is closely related to SIVagmTAN-1 (Accession number U58991) isolated from an AGM in Eastern Uganda (Figure S1).


Figure 11. Maximum likelihood tree of the partial pol gene ( 650 bp ) illustrating genetic diversity of SIV from synanthropic (urban dwelling) AGMs from three urban centres in Kenya. Sequences from Kenya, South Africa and West Africa are highlighted in red, blue and green respectively. Sequences obtained from GenBank with the host species originating from Kenya are shown in black. The sequence obtained from tantalus subspecies of AGMs from Uganda is highlighted in lightblue. Solid brown circles represent bootstrap support values of $>70$ while the bar represents the number of nucleotide substitutions (0.1) per site. Naming of sequences generated in this study is according to the monkey species (AGM), the animal number and sampling site.

Phylogenetic results obtained for the env and pol genes show similarity in the geographical clustering of the sequences according to the country of origin (Figure 12). No clear segregation of SIV variants according to the geographical locations of host sampling points was evident within Kenya. SIV sequence isolated from an olive baboon sampled in Kisumu clustered with sequences from sympatric AGMs with high bootstrap support. Based on the branching pattern of the phylogenetic trees, SIV from animals sampled in Mombasa and Kisumu appeared to be ancestral to those infecting animals sampled in Naivasha. Epidemiologically linked SIVs (AGM682 and AGM-NV4, AGM - NV681 and AGM - NV2, AGM - GC5 and baboon - Kaj5) were identified in Naivasha and Kisumu (Figure 12).


Figure 12: Maximum likelihood tree of the partial env gene ( 900 bp ) illustrating genetic diversity of SIV infecting synanthropic AGMs from several urban centres in Kenya. Sequences from Kenya, South Africa and West Africa are highlighted in red, blue and green respectively. Sequences obtained from GenBank with the host species originating from Kenya are shown in black. Solid brown circles represent bootstrap support values of $>70$ while the bar represents the number of nucleotide substitutions (0.1) per site. Naming of sequences generated in this study is according to the monkey species (AGM), the animal number and sampling site.

Within the phylogenetic cluster of SIVs isolated from Kenya, there was no clear segregation according to the sampling towns amongst the pol and env sequences from this study (Figure 13). This topology indicates existence of SIVagm sequence divergence despite falling within the same Kenyan cluster. SIV sequences from the same urban centre for instance

Naivasha (Figure 13) clustered in phylogenetic branches with low bootstrap support values (<70), demonstrating that AGMs sharing a habitat are infected with highly divergent SIV strains. This observation is well supported by the results from HRM analysis which showed unique melt profiles and thus limitation in identifying the SIV strains by HRM alone. On the other hand, sequences with high bootstrap support values (100) illustrates existence of a common evolutionary history and possibilities of sharing a common ancestor in the recent past.

The phylogenetic topology exhibited by nucleotide sequences was also observed for amino acids where there was no clustering according to the location of the hosts in Kenya. Whereas, the env protein sequences from Kenya interspersed with sequences from Africa (Figure S2), the pol protein maintained the tree topology (Figure S3) observed in the nucleotide sequences (Figure 11). The diversity results from the protein sequences are in agreement with HRM and phylogenetic results where samples from Naivasha, (MG590129-AGM682 and MG590127-AGM675) with similar melt profiles had 99\% identity. SIVs infecting MG590135AGMMbk11 from Mombasa and MG590131-AGM699 from Naivasha had 90\% identity, a trend illustrated by their melt profile differences. Whereas, SIV variants from MG590131AGM699and MG590127-AGM675 having different melt profiles, had 97\% in their pol amino acids sequences. On the other hand, the amino acids conservation was lower in the env protein compared to the pol protein as shown from the amino acid alignments (Figure S3). The sequences from Naivasha had $77 \%-98 \%$ amino acid identity in the env protein. These sequences from Naivasha exhibited $77 \%-89 \%$ identity to sequences from Mombasa and $80 \%$ - $91 \%$ identity to sequences from Kisumu. The env amino acid sequence from a from an olive baboon (MG590126 Baboon5) sampled in Kisumu had 96\% identity with the amino acid sequence from an AGM (MG590111 AGM5) also from Kisumu despite maintaining 80\% $89 \%$ identity to the rest of the sequences. This further supports epidemiological linkage of these two isolates as implied by phylogenetic analysis (Figure 12).

A

B


Figure 13: Maximum likelihood phylogenetic tree generated from partial pol gene (A) and env gene (B) illustrating genetic diversity of SIV from synanthropic (urban dwelling) AGMs from three urban centres in Kenya alongside previously characterized sequences from Kenya. Sequences from this study are highlighted in red while previously characterised Kenyan sequences retrieved from GenBank are shown in black. The percent bootstrap support values are shown at the nodes of the phylogenetic tree. The branch support evaluation was done using 1000 bootstrap replicates.

### 4.4.3 SIV pol and env protein diversity

Overall, protein sequence conservation was higher in the pol protein sequences compared to the env protein sequences (Figure S1). The identity of the amino acids for the partial pol protein ranged between $91-99 \%$ for the samples obtained from Naivasha. Samples obtained from Mombasa had an amino acid identity of $88 \%$. Amino acid sequences from AGM Kis18 sampled in Kisumu, had $79 \%$ - $84 \%$ identity with other sequences from Kenya further supporting the genetic distant demonstrated by phylogenetic analysis (Figure 11). Similar to the pattern observed for nucleotide sequences, the sequences from Naivasha had 91-94\% identity to sequences from Mombasa while also having $88-92 \%$ identity to sequences from Kisumu supporting lack of phylogeographical clustering. The overall amino acid identity observed for the pol protein suggests that codon usage was maintained despite the high nucleotide sequences diversity. Despite phylogenetic and amino acid diversity, the conserved domains and hypervariable regions (V3-V5) were conserved as expected in the env protein sequence (Figure 13 B ). The identified V3 domain had 34 amino acids as reported previously while maintaining the well-conserved MAG motif that largely determines cellular tropism in SIV (Figure 13. C) (Carr et al., 2017). Eight of 12-conserved cysteine in the env region (Müller et al., 1993; Ma et al., 2013), were present. Conservation of major env functional domains including the CD4 binding sites, N -linked glycosylation sites and fusion peptide suggests protein function was maintained.

A


B


C
M29975.1 SIVagm KE AGM682 SIVagm_KE
JX462429 SIVaḡm_SA KR862356 SIVagm WA LC114462.1 SIVagm SA MG590111_AGMGC5 SIV̄agm_KE MG590126_PAN5KAJ SIVagm_KE MG590109-AGM17KAJ SIVagm_KE U03997.1-SIVagm WA AB253736.1 SIVagm KE MG590123_AGMGAC SIV̄agm_KE MG590122_AGM699 SIVagm_KE MG590114 ${ }^{-}$AGM680 SIVagm ${ }^{-}$KE MG590115_AGM681 SIVagm KE MG590118_AGM685 SIVagm_KE MG590113-AGM675 SIVagm_KE
MG590124_AGM7MBK SIVagm_KE


Figure 14. SIV genome architecture and partial env protein domains. A. A cartoon representing SIV genome architecture showing respective genes and offset is a representation of the V3-V5 and the fusion protein (FP) domain) domains of the partial env glycoprotein. B. Env protein amino acids alignment showing the V3 to V5 domains, the fusion protein, the potential N -linked glycosylation sites highlighted with asterisks ( ${ }^{* * *)}$. The conserved cysteine residues are highlighted with black dots while potential CD4 binding sites are indicated as black squares, the MAG motif is highlighted hash signs (\#\#\#). C. A trimmed alignment of V3 hypervariable domain comprising 34 residues and showing the MAG motif as reported from previous studies.

### 4.5 Evolutionary selection

Analysis of evolutionary selection pressure demonstrated signatures of episodic and pervasive diversification on the env gene. IFEL and FEL approaches revealed codons 18 and 58 (Table 4) to be under purifying/negative selection ( 0.05 significance level). FUBAR revealed 14 sites under pervasive diversifying selection at posterior probability of $>0.9$ while MEME revealed 21 sites to be under episodic diversifying selection at a 0.05 level of significance. Overall, eight sites were identified as being under diversifying selection by FUBAR and MEME (Table 4).

Table 4: SIV env gene codons under evolutionary selection pressure. Analysis was undertaken using methods implemented on the Datamonkey web server (Delport et al., 2010).

| Codon | Diversifying selection | Negative selection |  |  |
| :--- | :--- | :--- | :--- | :--- |
|  | FUBAR $(\boldsymbol{\beta}>\boldsymbol{\alpha})$ | $\mathbf{0 . 9})$ | MEME $(\mathbf{P}<\mathbf{0 . 0 5})$ | IFEL $(\mathbf{P}<\mathbf{0 . 0 5})$ |
|  | FEL $(\mathbf{P}<\mathbf{0 . 0 5})$ |  |  |  |
|  | Pervasive | Episodic | Negative | Negative |
| $\mathbf{2}$ | 0.930754 | 0.006223 | - | - |
| $\mathbf{1 8}$ | - | - | 0.010375 | 0.03724 |
| $\mathbf{5 8}$ | - | - | 0.020702 | 0.046966 |
| $\mathbf{7 0}$ | 0.926138 | 0.001725 | - | - |
| $\mathbf{1 2 4}$ | 0.939274 | 0.002007 | - | - |
| $\mathbf{1 6 1}$ | 0.926871 | 0.019448 | - | - |
| $\mathbf{1 7 0}$ | 0.965838 | 0.01065 | - | - |
| $\mathbf{2 9 6}$ | 0.927782 | 0.01812 | - | - |
| $\mathbf{2 9 7}$ | 0.960401 | 0.00871 | - | - |
| $\mathbf{3 0 1}$ | 0.927481 | 0.012432 | - | - |

### 4.6 Summary

In summary, this study sought to determine the prevalence and genetic diversity of simian immunodeficiency virus (SIV) in common free-ranging urban restricted non-human primates. The target NHP species included, AGMs and olive baboons captured in Mombasa, Naivasha and Kisumu urban centres. Molecular techniques by PCR-HRM, gel-electrophoresis and sequencing were used to screen DNA samples extracted from the sampled NHPs for SIV. The overall SIV prevalence was $32 \%$ (39/124) in AGMs and 3\% (2/65). More adults (27 out of 72, $37.5 \%$ ), than subadults ( 9 out $36,25 \%$ ) and infants (3 out of $16,18.8 \%$ ) were positive for SIV. High-resolution melt analysis of the PCR amplicons revealed high diversity of SIV in the sampled animals. Despite demonstrating extensive diversity, HRM did not distinguish between different strains circulating in the sampled NHPs that led to sequencing to assist in characterisation of the detected SIVs. The nucleotide sequence plasticity observed from sequence comparison could be the reason why HRM did not resolve different SIV strains detected. Partial SIV pol and env genes were sequenced from this study. Comparison with characterised sequences from GenBank placed these sequences within the SIVagm lineage including the one isolated from an olive baboon. This also revealed nucleotide sequence plasticity upon sequence alignment. Phylogenetic analysis confirmed the extensive diversity that was demonstrated by HRM. There was no phylogeographic clustering according to the NHPs sampling location but rather all the sequences were homogenously distributed within the Kenyan clade of SIVs. Analysis of evolutionary selection pressure revealed episodic and pervasive diversifying selection within the env gene of SIVs indication continuous SIV evolution within its host.

## CHAPTER FIVE

## DISCUSSION

This study reports on comprehensive molecular survey of SIV variants circulating in common monkeys, the AGMs and olive baboons inhabiting urban and peri-urban centres in Kenya. Apart from serologic survey (Ellis et al., 2004) and opportunistic sampling for SIV in captive animals (Otsyula et al., 1996), comprehensive molecular epidemiology studies of SIVs in free ranging monkey populations in Kenya have not been previously conducted to the best of our knowledge. We report an overall SIV prevalence of $32 \%$ amongst free-ranging urbanrestricted AGMs and low SIV infection (3\%) in peri-urban dwelling olive baboons in Kenya. Data from this study reveals higher overall SIV prevalence infecting AGMs in Kenya compared to previously documented $26 \%-28 \%$ seroprevalence in captive and semi-captive AGMs (Ohta et al., 1988; Otsyula et al., 1996). We also report for the first time detection of SIVagm strain in olive baboons from Kenya. Previous studies have reported SIVagm strains from yellow baboons and chacma baboons from Tanzania and South Africa respectively (Kodama et al., 1989; Jin et al., 1994).

Although the overall SIV infection rate in AGMs from the urban centres was $33 \%$ in Mombasa, $24 \%$ in Kisumu and $45 \%$ in Naivasha, some sampling sites such as Mbaraki (Mombasa) and Kimana (Naivasha) had much higher infection rates of $48 \%$ and $46 \%$, respectively (Table S1). The findings from these sites are comparable to previous molecular studies that reported $44 \%$ and $45 \%$ overall SIVagm prevalence in wild caught AGMs from Gambia and South Africa, respectively (Ma et al., 2013; 2014). Likewise, the AGMs sampled in Tudor (Mombasa) and Kajulu (Kisumu) had lower infection rates of $8 \%$ and $19 \%$, respectively (Table S1). This was comparable to a study in Zambia that reported $19 \%$ SIVagm prevalence in AGMs (Carr et al., 2017). Generally, the prevalence according to sampling sites ranged from $8 \%-48 \%$ in our study. This was similar to SIV incidence variations reported by Baileys and co-authors (2016) from AGMs from various sampling locations in South Africa. The variations in the prevalence exhibited from our study could be attributed to a smaller sample size as compared to the study from Ma and co-authors (2013). The study from Baileys et al. (2016) also had a smaller sample size per sampling site (ranging from 12 to 40 individuals) thus implying a higher prevalence when all the NHPs captured per site individuals turned out to be infected with SIV. Our findings provide further molecular evidence of high SIVagm prevalence in Kenya thus corroborating the consideration of AGMs as natural hosts of SIV in Africa. Despite not showing uniformity of SIV prevalence as suggested from other
studies, our findings corroborate the observation that SIVagm infections are high and common in AGM species across Africa.

Prevalence of SIV amongst different age groups was non-uniform with more adult than sub-adult and infant AGMs being SIV positive. The plausible reason for low infection rate amongst infants is the rare vertical transmission of SIV in AGMs as previously reported (Otsyula et al., 1996; Chahroudi et al., 2012). Ma and co-authors in 2014 demonstrated immunological suppression of SIV replication in infants thereby limiting infection. In addition, immunological adaptations in infants and juveniles has been shown to confer them with resistance to maternal virus transmission thus lowering SIV prevalence amongst this age-group (Pandrea et al., 2012). In contrast, sexual transmission is the main route of SIV transmission in adults and therefore enhanced exposure during mating leads to high prevalence among adults (Phillips-Conroy et al., 1994). The prevalence data from this study is comparable to previously documented seroprevalence exceeding $30 \%$ amongst extensively studied species including wild and captive AGMs in Africa (Santiago et al., 2005; Aghokeng et al., 2006; Ma et al., 2013).

Detection of SIVagm in an olive baboon from Kenya corroborates epidemiological findings from earlier studies from chacma baboons in Southern Africa and yellow baboons in Tanzania (Kodama et al., 1989; Jin et al., 1994; Carr et al., 2017). Elsewhere, wild caught chacma baboons also tested positive for SIV serologically, while sequencing confirmed the virus to be of SIVagm strain (van Rensburg et al., 1998) suggesting natural infection from sympatric AGMs. Our findings confirm the possibility of SIVagm establishing in new hosts. Given that a baboon-specific SIV has not been identified this far, it would be necessary to determine the extent of SIV transmission from other species into baboons. Characterisation of these viruses will not only be important in elucidating the determinants of cross-species transmission but also host adaptation to the virus. This complicates the lineage classification based on host. The infectious outcomes of SIVagm in baboons could offer insights into development of a baboon model for HIV therapeutic research.

To evaluate the diversity and attempt characterisation of SIV strains detected in these free ranging NHPs, PCR amplicons for SIV were subjected to HRM analysis. Despite its robustness, this technique could not explicitly distinguish SIV strains in this study based on their melt profiles. However, variation in melting profiles showed extensive diversity of SIV strains from the same sampling site (Figure 6). In studies where single nucleotide polymorphisms are critical, the robustness of HRM in analysing miniature differences between PCR generated DNA fragments has proved to be reliable (Hinsberger et al., 2019). To further
characterise SIV sequences amplified by PCR-HRM and sequenced, we conducted a BLAST search on NCBI's GenBank database. This gave SIVagm as topmost hits for all the sequences from this study. The identities were further confirmed by conducting sequence alignments and generation of identity matrices that placed our sequences within the SIVagm lineage (Figures 8 and 9).

Therefore, variation in HRM melt profiles suggests existence of SIV intra-lineage diversity. This was supported by the results from multiple sequence alignments and generation of percent identity matrices (Figure 8). From the alignments, SIV sequences showed nucleotide plasticity, which could probably contribute to the observed varying melt profiles for the same SIVagm lineage. Likewise, the percent identities from the matrices ranged from $70 \%$ to $97 \%$ (Figure 9). This shows that despite the sequences falling within the SIVagm lineage, there were imminent differences in their sequences, thus giving rise to varying melt profiles. Therefore, this observation presents a limitation in HRM characterisation of SIV that exhibits low nucleotide conservation on the sequence alignments. As reviewed by Tong \& Giffard (2012), an increase in length of the amplicon size beyond 100bp becomes difficult to resolve by HRM for sequences with nucleotide plasticity such as SIV and HIV. Despite the HRM's shortfall, the technique stood out as a robust method in longitudinal evaluation of HIV diversity within an infected individual (Towler et al., 2010). In Towler's study, HIV diversity was documented in terms of variation in the melt curve width and not the different shapes/profiles of the melt curves. Progression of HIV infection correlated with virus diversification from founder strains, which correlated with increasing melt curve width. However, PCR-HRM is a cost-effective technique for monitoring prevalence of primate lentiviruses in urban-restricted NHPs rather than independent evaluation of SIV diversity/characterisation especially in virus isolates from different hosts. Despite its shortfalls in SIV characterisation, HRM is emerging as a sensitive and robust probe free technique for detecting infectious pathogens and differentiate genetic variants of different viruses (Lin et al., 2008; Steer et al., 2009; Yu et al., 2015; Villinger et al., 2017) including HIV (Towler et al., 2010).

Upon phylogenetic analysis, both env and pol sequences from this study clustered with previous isolates infecting captive AGMs from Kenya (Figure 11 and 12). However, these Kenyan isolates were distinct from those from infected monkeys in South Africa and West Africa. One sequence MG590133-AGMKis18 from Kisumu clustered with SIVagm Tan-1 isolated from Eastern Uganda (Soares et al., 1997), suggesting that the two had a common evolutionary history (Figure 11). This could be attributed to geographic proximities of their hosts' ranges as previously reported for simian T-lymphotrophic virus (STLV) infecting
mustached monkeys in Gabon (Liégeois et al., 2012). This observation agrees with the west-to-east SIV transmission model across existing AGM geographic ranges given the location of Kisumu relative to Uganda boarder (Wertheim \& Worobey, 2007). However, there is need to have a more detailed genetic analysis including full length genome sequencing of the isolate to exclude the possibilities of it being a recombinant/mosaic strain. Based on the overall phylogenetic tree topologies, both pol and env trees, SIV isolates from West Africa appeared to be ancestral to the SIV strains from East Africa and South Africa. Despite extensive diversity demonstrated by PCR-HRM, phylogenetic analysis did not show phylogeographical clustering of SIV according to different locations in Kenya from which monkey populations were sampled. However, based on identification of closely related phylogenetic clusters with strong node bootstrap supports, epidemiologically linked SIV sequences could be inferred from NHPs sampled in the same population such as Naivasha suggesting high levels of local virus transmission (Figures 13). Bootstrap support values (100\%) amongst SIV isolates from Naivasha imply continuous local virus transmission. In contrast, the interspersing of the isolates from Kisumu and Mombasa indicate that divergent strains can also occur in one location as previously reported in mustached monkeys (Liégeois et al., 2012) and chimpanzees (Van Heuverswyn et al., 2007). Within the cluster of SIV isolated from Kenya, there was no clear segregation according to the sampling towns amongst the newly sequenced pol and env (Figure 13) SIV gene fragments. This suggests and overall relatedness of these sequences as demonstrated by robust bootstrap values for the clusters on both phylogenetic trees.

For the first time, using molecular tools, we confirmed SIV infection in two olive baboons sampled in Kenya. The SIV sequence recovered from a baboon shared $97 \%$ similarity with an isolate from an AGM (MG590111-AGM5) from Kisumu (Figure 10). Phylogenetic analysis placed the baboon-derived sequence in a strongly supported cluster ( $100 \%$ bootstrap) with MG590111-AGM5 sampled in Kisumu (Figure 12). Upon translation and alignment of the nucleotide sequences, these two isolates had $98 \%$ amino acid identity (Table S3). Based on these results, the two SIV isolates were closely related and could be epidemiologically linked to a common ancestor within the SIVagm lineage. Detection of SIVagm in an olive baboon implies cross-species SIVagm transmission and diversification, increasing probabilities of establishing in this new baboon host. The most plausible explanation for cross-species transmission could be attributed to the aggressive behaviour of baboons (Jin et al., 1994; Carr et al., 2017), which strongly imply baboons could be infected from preying on AGMs (Kodama et al., 1989). Further, cross-species acquisition of SIVagm in Patas monkey resulted from physical contact involving aggression and bites amongst AGMs and Patas monkeys sharing a
habitat in Senegal (Bibollet-Ruche et al., 1996). Likewise, the behavior of hunting and preying on small monkeys resulted in SIVgsn and SIVrcm co-infection in chimpanzees whose outcome was recombination and emergence of SIVcpz in chimpanzees (Bailes et al., 2003). Elsewhere, physical aggression resulting in bites from NHPs has been reported as a cause of human zoonotic infection with simian foamy virus (SFV) in Gabon (Mouinga-Ondémé \& Kazanji, 2013). All these scenarios provide the chance of exposure to body fluids, one of the modes of retroviruses transmission. This further substantiates the evidence of the complex evolution and cross-species transmission of primate lentiviruses in their natural habitat. The potential of freeranging urban-restricted AGMs and baboons serving as sources of emerging retroviruses in human populations cannot be understated. Notably, SIVs from AGMs have successfully been cultured on immortalized human CD4 cells (Carr et al., 2017), suggesting the possibility of replicating in human cells.

There was an overall amino acid conservation in terms of percentage identity and sequence alignments within the pol protein (Table S2 and Figure S 1 ) compared to the env protein (Table S3 and Figure S1) as previously reported in northern talapoins (Miopithecus ogouensis) (Liegeois et al., 2006). However, the amino acid's percent identity ( $81 \%-99 \%$ ) was higher than the nucleotide sequences ( $66 \%-99 \%$ ) for pol gene and ( $77 \%-98 \%$ ) amino acid identity compared to ( $62 \%-99 \%$ ) in nucleotide sequence similarity for env gene. Equally, percentage amino acid similarities were higher suggesting prevalence of silent mutation. Despite the reduction of the amino acid identity/similarity range compared to the nucleotide sequences suggesting its utility to improve lineage classification, its reliability is still complex. For instance, up to $31 \%$ amino acid differences within SIVagm sequences have been reported suggesting high variability in sequences from within the AGM species (van Rensburg et al., 1998). In contrast, $84 \%-94 \%$ amino acid similarity between SIV sequences from red-capped mangabeys (SIVrcm) and agile mangabeys (SIVagi) were reported suggesting close relationship of these two sequences from two different host species (Ahuka-Mundeke et al., 2010). Elsewhere, genetic distance assessment of the entire SIVmus proteome revealed identity differences for the isolates from this lineage within the env proteome (Aghokeng et al., 2007). These findings suggest that classification of SIV into lineages is complex and requires holistic combination of various factors including identity of the host species this gets complex in case of cross-species transmission and recombination. Generally, the pol protein conservation enables accurate assignment of newly sequenced SIVs into their respective lineages. Whereas diversity in the env domains are associated with subtle but important structural and hence functional changes (Leonard et al., 1990).

Despite env amino acid diversity, V3-V5 hypervariable domains revealed a similar structure to a reference SIVagmVER-2 (GenBank accession U04004.1) as previously described (Ma et al., 2013), suggesting that these sequences could be translated into functional proteins. Further, the expected N -linked glycosylation sites, the cysteine residues and the potential CD4 binding sites were conserved in the env proteins suggesting maintained biological functions despite observed nucleotide sequence diversity (Figure 13). These results suggests existence of synonymous and non-synonymous nucleotide substitutions. Analysis of the synonymous and non-synonymous substitution rates ( dS and dN ) for the env protein revealed average $\mathrm{dS}=0.0309, \mathrm{dN}=0.0957, \mathrm{dS} / \mathrm{dN}=0.3149$ and $\mathrm{pS} / \mathrm{pN}=0.3286$ (Table S4). The observed $\mathrm{dS} / \mathrm{dN}$ ratio ( 0.3149 , which is $<1$ ) illustrates the presence of purifying selection against deleterious mutations and hence maintain a viable virus population. Non-synonymous mutations induced by selection pressure affects protein function thereby influence virus fitness. The SIVs from the targeted animals in this study exhibited extensive genetic diversity, indicating that they thrive in a suitable niche for viral evolutionary changes within natural hosts (Simmonds et al., 2018). When selection favours mutations within amino acids critical for hostvirus interactions, the virus fitness in a new host is favoured (Shackelton et al., 2005). Therefore, the diversity and selection patterns of SIV in AGMs could be a recipe for emergence of novel strains that could cross species barriers. The risk of SIV transmission into different host species including human cannot be ignored in the wake of high SIV prevalence in freeranging urban restricted AGMs with evidence of SIVagm infection in an olive baboon because of cross-species transmission.

The codon-oriented FUBAR and MEME methods revealed evidence of diversifying (positive) selection within the SIV env genes sequenced in this study. These signatures of evolutionary selection pressure implies ongoing viral molecular adaptation. The sequenced gene fragment corresponded to the surface (SU) fragment that is accessible to the immune system thus playing a role in host-virus interaction. This interaction with the host elicits immunological pressure on the env gene resulting in enhanced evolution, sequence diversity and increased fitness (Lynch et al., 2009). Under diversifying or positive selection, more fit variants from a virus quasispecies (a virus population comprising related sequences arising from high mutation rates during viral RNA replication) are selected over variants with deleterious mutations thereby ensuring viable virus population are able to infect new cells. Despite resulting in non-viable individuals, existence of virus quasispecies creates a pool of beneficial mutations that enhance adaptation to new environments and host immunological pressure during infection. This sequence diversification and evolutionary selection in a natural
host is a typical characteristic of potentially zoonotic viruses as opposed to viruses that diversify in a dead-end host thus limiting onward transmission (Vanderford et al., 2006). Located within the SIV env glycoprotein is a V3 domain, which is the most conserved of all the five hypervariable regions. V3 is crucial to viral entry by being a critical determinant of the chemokine receptor's specificity. Therefore, amino acid sequence changes that affect overall charge will have an impact on co-receptor usage during the host cell invasion, thereby influencing adaptation to a new host (Demma et al., 2005; Van Heuverswyn et al., 2007). This phenomenon underscores the diversification of SIV as a potential precursor for pathogen spill over.

Our results demonstrate extensive SIVagm diversity in free-ranging AGMs from Kenya, besides complementing the high prevalence findings from other African countries. This study also documents, for the first time in Kenya, SIVagm infection in olive baboons, suggesting cross-species SIVagm transmission. The SIVagm virus is known to be species-specific to the genus Chlorocebus (Allan et al., 1991). Further, the occurrence of highly related viruses infecting AGMs from Naivasha indicates identifiable local transmission within the population. Immense immunological pressure is being exerted on the env protein leading to concentration of episodic diversifying selection in this region and probably increasing chances of establishing in this new host, the baboon. Further sampling of AGMs and baboons is necessary to reveal the historical connection of SIVs infecting monkeys in East Africa and identify the extent of cross-species transmission into baboons in the wild and the associated outcomes of the infection in the new primate host. This could be important for the establishment of a baboon model for HIV vaccine research. With reports of continued transmission of simian retroviruses, including SIV, STLV, and SFV to humans, our findings underscore the importance of characterising diverse SIV strains in East African primate populations to identify those that can potentially cross species barriers and could establish in humans. Finally, our findings demonstrate the urgency of developing local surveillance programmes to identify the extent of retrovirus genetic diversity, including possible cross-species transmission events across major human-NHP interfaces with high transmission risks, and adequate SIV control strategies at human-wildlife interfaces.

## CHAPTER SIX

## CONCLUSIONS AND RECOMMENDATIONS

### 6.1 Conclusions

i. This study demonstrates SIV prevalence above $30 \%$ in AGMs which is comparable to previous studies that have shown prevalence greater than $40 \%$ in wild AGMs in Africa, implying that urban-restricted AGMs also exhibit high levels of SIV infection. Further, this study detected SIVagm strain in olive baboons that commonly shared a habitat with AGMs in Mombasa and Kisumu suggesting possibility of cross-species transmission.
ii. There is extensive genetic diversity of SIV infecting free-ranging NHPs found in urban and peri-urban centres in Kenya as illustrated by both phylogenetics and by amino acid sequence similarity.
iii. Phylogenetic analysis of SIV sequences from this study did not show evidence of phylogeographical clustering according to the origin of the host NHPs. This implies the strains from Kenya could be having a common ancestor including the strain detected in olive baboons. Signatures of diversification detected on the SIV partial env sequences implies continuous virus evolution in their hosts that probably generated the strain with the ability to establish in a new host.

### 6.2 Recommendations

The following are the recommendations drawn from this study:
i. There is potential of cross-species transmission of SIV amongst sympatric NHP species. Cross-species transmission increases the risk of co-infection with different strains, which provides a probable chance for virus recombination which can generate unique SIV strains with unknown zoonotic potential.
ii. With increased frequency of human-wildlife contacts across different human-wildlife interfaces in Kenya, the potential of emergence of novel zoonotic human retroviruses with a significant impact on human health as evidenced by HIV/AIDS pandemic cannot be overlooked. This calls for elucidation of the zoonotic risks posed by SIV circulating in NHP species across various human-NHP interfaces in terms of infection outcomes in a non-natural hosts such as olive baboons. Such studies will help describe the overall picture of the emergence of zoonotic pathogens with a primate origin such as HIV/SIV and inform strategies for mitigating unforeseen pandemics.
iii. There is also a need for community sensitisation and engagement with policy makers about the zoonosis risks involved during aggressive human encounters with NHPs.

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## APPENDICES

## Appendix 1: Supplementary tables

Table S1: Prevalence of SIV infection in AGMs and olive baboons according to species and sampling sites from Mombasa, Naivasha and Kisumu as detected by PCR.

| Sampling <br> Town | Sampling Sites | Infected (\%) SIV positive/Total sampled NHPs |  |
| :--- | :--- | :--- | :---: |
|  | AGMs | Baboons |  |

Mombasa

| Mbaraki | $11 / 23(48 \%)$ |  |
| :--- | :--- | :--- |
| Tudor | $1 / 12(8 \%)$ | - |
| GAC | $5 / 16(31 \%)$ | - |
| Mazeras |  | $1 / 55(1 \%)$ |
| Subtotal | $\mathbf{1 7 / 5 1}(\mathbf{3 3 . 3 \%})$ | $1 / 55(1 \%)$ |

## Kisumu

| Kajulu | $4 / 21(19 \%)$ | $1 / 10(10 \%)$ |
| :--- | :--- | :--- |
| Golf Course | $1 / 5(20 \%)$ | - |
| Impala | $7 / 25(28 \%)$ | - |
| Subtotal | $\mathbf{1 2} / 51(\mathbf{2 3 . 5 \%})$ | $1 / 10(10 \%)$ |

## Naivasha

|  | Kimana | $10 / 22(46 \%)$ |
| :--- | :--- | :--- |
| Total | $\mathbf{3 9 / 1 2 4 ( 3 2 \% )}$ | $\mathbf{2 / 6 5 ( 3 . 1 \% )}$ |

Table S2. Percent SIV pol amino acid identity matrix generated by multiple sequence alignment using clustal O implemented on the webserver at the European Bioinformatics Institute (EMBL-EBI) (Sievers et al., 2011).

|  |  |  |  |  | AB253736-SIVagmTYO-1 | AF395567_SIVagmTYO1 | X07805_SIV-AGM | $\begin{aligned} & \overline{2} \\ & \stackrel{+}{n} \\ & \underset{2}{2} \end{aligned}$ |  | MG590128_AGM681 | N |  |  |  |  |  |  |  |  | MG590134 AGMGAC16 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| AF395565_SIVagm | 100 | 95 | 95 | 88 | 91 | 90 | 90 | 81 | 89 | 90 | 90 | 90 | 90 | 90 | 91 | 97 | 85 | 85 | 91 | 90 |
| M29975.1_SIVagm | 95 | 100 | 95 | 91 | 92 | 91 | 92 | 82 | 91 | 93 | 92 | 92 | 92 | 92 | 94 | 97 | 83 | 88 | 93 | 92 |
| AF395564_SIVagm | 95 | 95 | 100 | 92 | 94 | 93 | 93 | 82 | 92 | 94 | 94 | 94 | 94 | 94 | 95 | 94 | 82 | 89 | 91 | 93 |
| L40990_SIVagm | 88 | 91 | 92 | 100 | 92 | 91 | 92 | 79 | 90 | 92 | 91 | 92 | 92 | 92 | 93 | 90 | 81 | 88 | 88 | 93 |
| AB253736-SIVagmTYO-1 | 91 | 92 | 94 | 92 | 100 | 99 | 99 | 82 | 89 | 91 | 90 | 90 | 90 | 90 | 92 | 93 | 80 | 91 | 91 | 94 |
| AF395567_SIVagmTYO1 | 90 | 91 | 93 | 91 | 99 | 100 | 98 | 81 | 88 | 90 | 88 | 89 | 89 | 89 | 90 | 92 | 79 | 90 | 90 | 92 |
| X07805_SIV-AGM | 90 | 92 | 93 | 92 | 99 | 98 | 100 | 81 | 88 | 90 | 89 | 90 | 90 | 90 | 91 | 92 | 79 | 90 | 90 | 93 |
| U58991 | 81 | 82 | 82 | 79 | 82 | 81 | 81 | 100 | 81 | 82 | 83 | 82 | 82 | 82 | 81 | 82 | 93 | 82 | 83 | 81 |
| MG590131_AGM699 | 89 | 91 | 92 | 90 | 89 | 88 | 88 | 81 | 100 | 95 | 95 | 97 | 97 | 97 | 96 | 90 | 82 | 88 | 90 | 91 |
| MG590128_AGM681 | 90 | 93 | 94 | 92 | 91 | 90 | 90 | 82 | 95 | 100 | 99 | 97 | 97 | 98 | 98 | 93 | 82 | 88 | 90 | 94 |
| MG590139_AGMNV21 | 90 | 92 | 94 | 91 | 90 | 88 | 89 | 83 | 95 | 99 | 100 | 97 | 97 | 98 | 97 | 92 | 82 | 88 | 88 | 92 |
| MG590129_AGM682 | 90 | 92 | 94 | 92 | 90 | 89 | 90 | 82 | 97 | 97 | 97 | 100 | 100 | 99 | 97 | 91 | 83 | 89 | 89 | 92 |
| MG590137_AGMNV4 | 90 | 92 | 94 | 92 | 90 | 89 | 90 | 82 | 97 | 97 | 97 | 100 | 100 | 99 | 97 | 91 | 83 | 89 | 89 | 92 |
| MG590127_AGM675 | 90 | 92 | 94 | 92 | 90 | 89 | 90 | 82 | 97 | 98 | 98 | 99 | 99 | 100 | 97 | 92 | 82 | 90 | 89 | 93 |
| MG590130_AGM684 | 91 | 94 | 95 | 93 | 92 | 90 | 91 | 81 | 96 | 98 | 97 | 97 | 97 | 97 | 100 | 93 | 82 | 87 | 90 | 94 |
| MG590136_AGMNV22 | 97 | 97 | 94 | 90 | 93 | 92 | 92 | 82 | 90 | 93 | 92 | 91 | 91 | 92 | 93 | 100 | 84 | 88 | 92 | 93 |
| MG590133_AGMKis18 | 85 | 83 | 82 | 81 | 80 | 79 | 79 | 93 | 82 | 82 | 82 | 83 | 83 | 82 | 82 | 84 | 100 | 79 | 82 | 80 |
| MG590132_AGMKis17 | 85 | 88 | 89 | 88 | 91 | 90 | 90 | 82 | 88 | 88 | 88 | 89 | 89 | 90 | 87 | 88 | 79 | 100 | 88 | 88 |
| MG590135_AGMMbk11 | 91 | 93 | 91 | 88 | 91 | 90 | 90 | 83 | 90 | 90 | 88 | 89 | 89 | 89 | 90 | 92 | 82 | 88 | 100 | 88 |
| MG590134_AGMGAC16 | 90 | 92 | 93 | 93 | 94 | 92 | 93 | 81 | 91 | 94 | 92 | 92 | 92 | 93 | 94 | 93 | 80 | 88 | 88 | 100 |

Table S3. Percent SIV env amino acid identity matrix generated by multiple sequence alignment using clustal O implemented on the webserver at the European Bioinformatics Institute (EMBL-EBI) (Sievers et al ., 2011).

|  | AB253736_SIVagmTYO-1 | $\begin{aligned} & \text { à } \\ & \text { 会 } \\ & 0 \\ & 3 \end{aligned}$ | U93746_babenv05 | U93744_babenv03 | X07805_SIVagm | MG590109_AGM17Kis | MG590110_AGM4Kis | MG590126_AGM5Kis | MG590112_AGM672 | MG590113_AGM675 | 089W9V $\dagger$-II06SDN | MG590115_AGM681 | MG590116_AGM682 | MG590117_AGM684 | MG590118_AGM685 | MG590119_AGM686 | 069W9V 0ZL06SON | 869WDV ${ }^{-}$IZI06SDN | MG590122_AGM699 | U | MG590124_AGM7Mbk | MG590126_Baboon5 |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| AB253736_SIVagmTYO-1 | 100 | 84 | 78 | 84 | 99 | 86 | 88 | 86 | 81 | 83 | 88 | 89 | 88 | 86 | 89 | 88 | 88 | 82 | 89 | 86 | 82 | 86 |
| AF015909 | 84 | 100 | 73 | 79 | 82 | 81 | 83 | 82 | 80 | 80 | 81 | 81 | 84 | 84 | 82 | 84 | 84 | 76 | 85 | 81 | 79 | 83 |
| U93746_babenv05 | 78 | 73 | 100 | 90 | 77 | 78 | 79 | 77 | 73 | 75 | 80 | 79 | 80 | 78 | 80 | 79 | 80 | 74 | 79 | 77 | 75 | 78 |
| U93744_babenv03 | 84 | 79 | 90 | 100 | 83 | 83 | 84 | 83 | 78 | 81 | 85 | 85 | 85 | 84 | 86 | 84 | 85 | 80 | 84 | 83 | 82 | 84 |
| X07805_SIVagm | 99 | 82 | 77 | 83 | 100 | 85 | 87 | 85 | 79 | 82 | 87 | 88 | 87 | 85 | 88 | 87 | 87 | 81 | 88 | 85 | 81 | 85 |
| MG590109_AGM17Kis | 86 | 81 | 78 | 83 | 85 | 100 | 89 | 88 | 81 | 82 | 84 | 85 | 87 | 86 | 84 | 84 | 87 | 81 | 86 | 84 | 82 | 89 |
| MG590110_AGM4Kis | 88 | 83 | 79 | 84 | 87 | 89 | 100 | 88 | 82 | 85 | 88 | 89 | 89 | 87 | 89 | 90 | 94 | 84 | 91 | 86 | 83 | 87 |
| MG590126_AGM5Kis | 86 | 82 | 77 | 83 | 85 | 88 | 88 | 100 | 80 | 84 | 86 | 87 | 89 | 87 | 87 | 87 | 88 | 80 | 87 | 84 | 82 | 96 |
| MG590112_AGM672 | 81 | 80 | 73 | 78 | 79 | 81 | 82 | 80 | 100 | 80 | 82 | 82 | 86 | 87 | 82 | 80 | 81 | 77 | 82 | 80 | 77 | 82 |
| MG590113_AGM675 | 83 | 80 | 75 | 81 | 82 | 82 | 85 | 84 | 80 | 100 | 87 | 87 | 87 | 87 | 88 | 86 | 87 | 80 | 88 | 85 | 83 | 85 |
| MG590114_AGM680 | 88 | 81 | 80 | 85 | 87 | 84 | 88 | 86 | 82 | 87 | 100 | 98 | 90 | 89 | 98 | 89 | 92 | 83 | 90 | 88 | 85 | 87 |
| MG590115_AGM681 | 89 | 81 | 79 | 85 | 88 | 85 | 89 | 87 | 82 | 87 | 98 | 100 | 90 | 89 | 98 | 89 | 92 | 83 | 90 | 89 | 85 | 87 |
| MG590116_AGM682 | 88 | 84 | 80 | 85 | 87 | 87 | 89 | 89 | 86 | 87 | 90 | 90 | 100 | 97 | 91 | 89 | 90 | 83 | 91 | 88 | 86 | 91 |
| MG590117_AGM684 | 86 | 84 | 78 | 84 | 85 | 86 | 87 | 87 | 87 | 87 | 89 | 89 | 97 | 100 | 90 | 88 | 89 | 83 | 89 | 87 | 85 | 90 |
| MG590118_AGM685 | 89 | 82 | 80 | 86 | 88 | 84 | 89 | 87 | 82 | 88 | 98 | 98 | 91 | 90 | 100 | 90 | 93 | 84 | 91 | 88 | 85 | 87 |
| MG590119_AGM686 | 88 | 84 | 79 | 84 | 87 | 84 | 90 | 87 | 80 | 86 | 89 | 89 | 89 | 88 | 90 | 100 | 93 | 85 | 92 | 87 | 84 | 87 |
| MG590120_AGM690 | 88 | 84 | 80 | 85 | 87 | 87 | 94 | 88 | 81 | 87 | 92 | 92 | 90 | 89 | 93 | 93 | 100 | 85 | 92 | 87 | 86 | 87 |
| MG590121_AGM698 | 82 | 76 | 74 | 80 | 81 | 81 | 84 | 80 | 77 | 80 | 83 | 83 | 83 | 83 | 84 | 85 | 85 | 100 | 86 | 81 | 80 | 80 |
| MG590122_AGM699 | 89 | 85 | 79 | 84 | 88 | 86 | 91 | 87 | 82 | 88 | 90 | 90 | 91 | 89 | 91 | 92 | 92 | 86 | 100 | 88 | 86 | 86 |
| MG590123_AGM7GAC | 86 | 81 | 77 | 83 | 85 | 84 | 86 | 84 | 80 | 85 | 88 | 89 | 88 | 87 | 88 | 87 | 87 | 81 | 88 | 100 | 83 | 84 |
| MG590124_AGM7Mbk | 82 | 79 | 75 | 82 | 81 | 82 | 83 | 82 | 77 | 83 | 85 | 85 | 86 | 85 | 85 | 84 | 86 | 80 | 86 | 83 | 100 | 83 |
| MG590126_Baboon5 | 86 | 83 | 78 | 84 | 85 | 89 | 87 | 96 | 82 | 85 | 87 | 87 | 91 | 90 | 87 | 87 | 87 | 80 | 86 | 84 | 83 | 100 |

Table S4. Analysis of the synonymous and non-synonymous substitution rates (dS and dN) for the env protein. Analysis was done on codon alignments of the env sequences and the run executed on hyphy platform via data monkey webserver (Delport et al., 2010).

| Sequence | Compared against | SD | SN | S | N | pS | pN | dS | dN | dS/dN | pS/pN |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| MG590123 AGMGAC7 (Mombasa) | MG590111 AGM5 (Kisumu) | 3.67 | 60.33 | 181.83 | 715.17 | 0.02 | 0.08 | 0.02 | 0.09 | 0.23 | 0.24 |
| MG590123 AGMGAC7 (Mombasa) | MG590132 AGM17(Kisumu) | 7.17 | 71.83 | 180.50 | 716.50 | 0.04 | 0.10 | 0.04 | 0.11 | 0.38 | 0.40 |
| MG590123 AGMGAC7 (Mombasa) | MG590115 AGM681 (Naivasha) | 6.17 | 64.83 | 175.00 | 698.00 | 0.04 | 0.09 | 0.04 | 0.10 | 0.36 | 0.38 |
| MG590123 AGMGAC7 (Mombasa) | MG590121 AGM698 (Naivasha) | 5.17 | 71.83 | 181.00 | 716.00 | 0.03 | 0.10 | 0.03 | 0.11 | 0.27 | 0.28 |
| MG590123 AGMGAC7 (Mombasa) | MG590113 AGM675 (Naivasha) | 202.83 | 488.17 | 186.00 | 693.00 | 1.09 | 0.70 | nan | 2.10 | nan | 1.55 |
| MG590123 AGMGAC7 (Mombasa) | MG590114 AGM680 (Naivasha) | 7.67 | 79.33 | 180.67 | 716.33 | 0.04 | 0.11 | 0.04 | 0.12 | 0.36 | 0.38 |
| MG590123 AGMGAC7 (Mombasa) | MG590116 AGM682 (Naivasha) | 4.50 | 44.50 | 182.00 | 715.00 | 0.02 | 0.06 | 0.03 | 0.07 | 0.39 | 0.40 |
| MG590123 AGMGAC7 (Mombasa) | MG590117 AGM684 (Naivasha) | 11.50 | 103.50 | 169.83 | 685.17 | 0.07 | 0.15 | 0.07 | 0.17 | 0.42 | 0.45 |
| MG590123 AGMGAC7 (Mombasa) | MG590118 AGM685 (Naivasha) | 3.33 | 49.67 | 180.67 | 716.33 | 0.02 | 0.07 | 0.02 | 0.07 | 0.26 | 0.27 |
| MG590123 AGMGAC7 (Mombasa) | MG590119 AGM686 (Naivasha) | 3.83 | 54.17 | 181.33 | 715.67 | 0.02 | 0.08 | 0.02 | 0.08 | 0.27 | 0.28 |
| MG590123 AGMGAC7 (Mombasa) | MG590120 AGM690 (Naivasha) | 4.17 | 53.83 | 182.00 | 715.00 | 0.02 | 0.08 | 0.02 | 0.08 | 0.29 | 0.30 |
| MG590123 AGMGAC7 (Mombasa) | MG590122 AGM699 (Naivasha) | 3.67 | 43.33 | 181.67 | 715.33 | 0.02 | 0.06 | 0.02 | 0.06 | 0.32 | 0.33 |
| MG590123 AGMGAC7 (Mombasa) | MG590126 Baboon5 (Kisumu) | 3.67 | 61.33 | 175.67 | 685.33 | 0.02 | 0.09 | 0.02 | 0.10 | 0.22 | 0.23 |

Averages of the first sequence (MG590123 AGMGAC7, Mombasa) compared to others:
$\mathrm{dS}=0.0309 \mathrm{dN}=0.0957 \mathrm{dS} / \mathrm{dN}=0.3149 \mathrm{pS} / \mathrm{pN}=0.3286$

Table S5: Sample identification for NHPs that were positive for SIV. The accession numbers for respective sequences submitted to NCBI's GenBank database.

| Sample ID | GenBank <br> Accession | SIV gene | Sample ID | GenBank <br> Accession | SIV <br> gene |
| :--- | :--- | :--- | :--- | :--- | :--- |
| AGMKIS17KAJ | MG590108 | env | BABOON5 | MG590126 | env |
| AGMKIS4KAJ | MG590110 | env | AGMNVS675 | MG590127 | pol |
| AGMKIS5GC | MG590111 | env | AGMNVS681 | MG590128 | pol |
| AGMNVS672 | MG590112 | env | AGMNVS682 | MG590129 | pol |
| AGMNVS675 | MG590113 | env | AGMNVS684 | MG590130 | pol |
| AGMNVS680 | MG590114 | env | AGMNVS699 | MG590131 | pol |
| AGMNVS681 | MG590115 | env | AGMKIS17 | MG590132 | pol |
| AGMNVS682 | MG590116 | env | AGMKIS18 | MG590133 | pol |
| AGMNVS684 | MG590117 | env | AGM16GAC | MG590134 | pol |
| AGMNVS685 | MG590118 | env | AGM11MBK | MG590135 | pol |
| AGMNVS686 | MG590119 | env | AGMNVS22 | MG590136 | pol |
| AGMNVS690 | MG590120 | env | AGMNVS4 | MG590137 | pol |
| AGMNVS698 | MG590121 | env | AGMNVS5 | MG590138 | pol |
| AGMNVS699 | MG590122 | env | AGMNVS21 | MG590139 | pol |
| AGM17GAC | MG590123 | env | AGM6GAC | MG757169 | pol |
| AGM7MBK | MG590124 | env | AGM12NVS | MG757170 | pol |
| AGMNVS679 | MG590125 | env | AGM13NVS | MG757171 | pol |

## Appendix 2: Supplementary figures



Figure S1: Multiple sequence alignments for the pol protein amino acid residues (A) and env protein amino acid residues (B) generated from the partial pol and env gene fragments amplified in this study alongside SIVagm sequences from GenBank. The conserved residues are highlighted with a dark background and the column highlighted with an asterisks (*). The conservative amino acid replacements can be observed for the non-conserved residues indicating maintenance of the normal protein function.


Tree scale: $0.1 \longmapsto$
Figure S2: Maximum likelihood tree generated from partial pol protein amino acids illustrating genetic diversity of SIV from synanthropic (urban dwelling) AGMs from three urban centres in Kenya. Sequences from Kenya are highlighted in red, South Africa in blue and West Africa in green. Solid brown circles represent bootstrap support values of $>70$ while the bar represents the number of nucleotide substitutions (0.1) per site. Sequences generated in this study are named according to the monkey species (AGM), the animal number and sampling site.


Figure S3: Maximum likelihood tree generated from partial env protein amino acids illustrating genetic diversity of SIV from synanthropic (urban dwelling) AGMs from three urban centres in Kenya. Sequences from Kenya are highlighted in red, South Africa in blue and West Africa in green. Sequences in black and those highlighted in purple were deposited in GenBank from captive AGMs and infected baboons in the US that were positive for SIVagm. Solid brown circles represent bootstrap support values of $>70$ while the bar represents the number of nucleotide substitutions (0.1) per site. Sequences generated in this study are named according to the monkey species (AGM), the animal number and sampling site.

# Appendix 3: Publication 

# Broad diversity of simian immunodeficiency virus infecting Chlorocebus species (African green monkey) and evidence of cross-species infection in Papio anubis (olive baboon) in Kenya 

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#### Abstract

Background: Simian immunodeficiency virus (SIV) naturally infects African non-human primates (NHPs) and poses a threat of transmission to humans through hunting and consumption of monkeys as bushmeat. This study investigated the as of yet unknown molecular diversity of SIV in free-ranging Chlorocebus species (African green monkeys-AGMs) and Papio anubis (olive baboons) within Mombasa, Kisumu and Naivasha urban centres in Kenya. Methods: We collected blood samples from 124 AGMs and 65 olive baboons in situ, and detected SIV by high-resolution melting analysis and sequencing of PCR products. Results: Simian immunodeficiency virus prevalence was $32 \%$ in AGMs and $3 \%$ in baboons. High-resolution melting (HRM) analysis demonstrated distinct melt profiles illustrating virus diversity confirmed by phylogenetic analysis. Conclusions: There is persistent evolutionary diversification of SIVagm strains in its natural host, AGMs and cross-species infection to olive baboons is occurring. Further study is required to establish pathogenesis of the diverse SIVagm variants and baboon immunological responses.


## KEYWORDS

African green monkeys, evolution, high-resolution melting analysis, olive baboon, SIVagm, strain diversity

## 1 | INTRODUCTION

African non-human primates (NHPs) are reservoirs for diverse zoonotic retroviruses of human health importance including simian immunodeficiency virus (SIV), genus Lentivirus and simian foamy virus (SFV) ${ }^{1}$ besides being incidental hosts for Ebola ${ }^{2}$ and Marburg virus. ${ }^{3}$ Diverse strains of SIV infect over 40 species of African NHPs. ${ }^{4}$ Although not pathogenic to its natural host, the NHPs, the virus continuously poses a threat of transmission to humans because of
hunting and consuming NHPs as bushmeat. ${ }^{5,6}$ It has been shown that human immunodeficiency virus (HIV) originated from cross-species transmission of SIVcpz strain from chimpanzees (Pan troglodytes) and SIVsmm from sooty mangabeys (Cercocebus atys), resulting in zoonotic infections that respectively produced human immunodeficiency virus type 1 (HIV-1) and HIV-2 pandemic. ${ }^{7,8}$ Despite the high prevalence of natural SIV infections in African NHPs species, ${ }^{6,9,10}$ comprehensive surveys and molecular characterisation of lentiviruses infecting East African monkeys remain limited.
microbiologically documented without bacteremia (MDCD) $2-$ Clinically documented (CD) 3-Posible infection (PI): no MD no CD.

Results: 134 infectious episodes were diagnosed during 39 episodes of neutropenia in 33 patients, mean age 58( $\mathrm{DS} \pm 18$ ) years, female $78(58 \%)$, mean duration of neutropenia $30(\mathrm{DS} \pm 15)$ days, median ANC at diagnosis of infectious episode $41 \mathrm{cel} / \mathrm{mm}^{3}$ (range, $0-962$ ). We found 3,2( $\mathrm{DS} \pm 1,7$ ) infectious attacks by episode of neutropenia.

Among the 134, MD 58(43\%): BUO 9(7\%), BSI 30(22\%), MDCD 19(29\%); CD 58(43\%) and PI 18(13\%). Most common sites of infection: Lung $33(25 \%)$, intestinal tract $30(22 \%)$, skin and mucosa 20(15\%); 39 bacteremias ocurred with 43 isolations: gram negative (GN) 21 ( $49 \%$ ), gram positive (GP) $18(42 \%$ ), fungemia $3(7 \%)$, others $1(2 \%)$. K pneumoniae was the most common GN and coagulasenegative staphylococci between GP. The prevalence of multi drug resistance microorganisms (MDRM) in bacteremia was: $8(38 \%)$ in GN, and 6(33\%) in GP. Mortality rate was 12\%. Duration of neutropenia, median ANC, site of infection, and isolation of MDRM were not associated with mortality. On univariate and multivariate analyses the variables found to be significantly associated with mortality were the presence of bacteremia (OR 8: IC 2.4-28; $p=0,001$ ) and age $\geq 60$ years (OR 12: IC $2,56-61,7 ; p=0,002$ )

Conclusion: We found a high rate of bacteremia between infectious episodes, as well as high percentage of MDRM isolations. An age $\geq 60$ years and the presence of bacteremia were associated with more probability of death.
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Session: Moderated ePoster Presentations: HIV and Tuberculosis
Date: Friday, March 2, 2018
Time: 12:45-13:45
Room: San Telmo
Type: Electronic/Moderated Poster Presentation
Extensive diversity of SIVagm infecting
synanthropic African green monkeys and Olive Baboons in Kenya
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Background: Diverse strains of simian immunodeficiency virus (SIV) infect over 40 species of African nonhuman primates (NHP) and continuously pose potential threat of transmission to humans. However, strain diversity amongst common free-ranging Chlorocebus aethiops (African green monkey-AGMs) and Papio anubis (Olive baboon) in Kenya remains unclear. For the first time, we investigated epidemiology and diversity of SIV in AGMs and Baboon from major Kenyan urban centres.

Methods \& Materials: A total of 126 AGMs and 65 olive baboons from Mombasa, Kisumu and Naivasha were sampled once in situ and released. Blood samples were spotted on whatman FTA cards for DNA/RNA extraction. Total DNA from dried blood on FTA (DBFTA) was subjected to PCR followed by high resolution melting (HRM) analysis using consensus primers targeting a 650-bp fragment in partial pol gene and a 900-bp fragment of env gene encompassing hypervariable V3-V5 regions.

Results: PCR-HRM analysis illustrated amplification of SIV frag ments giving an overall infection rate of $30.95 \%$ (39/126) in AGMs and $3.07 \%$ (2/65) in Olive baboons. Subsequent sequence identification confirmed SIV infection in AGM and for the first time in olive baboon. Phylogenetic analysis of pol and env genes revealed extensive genetic diversity among newly generated SIVagm sequences within groups sympatric NHPs and within a geographical location Signatures of pervasive and episodic diversifying selection were also detected on the env gene indicating continuous SIV diversification.

Conclusion: This molecular evidence of SIVagm in olive baboons illustrates continuous simian-to-simian SIV transmission in the wild and can be linked to potential risks of transmission to humans through consumption of monkey bushmeat. In addition, this study shows that DB-FTA specimens and PCR-HRM analysis can be used as a cost-effective alternative sampling method for the surveillance and monitoring of SIV and other retrovirus of public health importance.
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Final Abstract Number: Fri_Station 02.2
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Type: Electronic/Moderated Poster Presentation
Functionality of CD8 ${ }^{+}$T-cells in subjects under cART: implications on cure strategies
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Background: Reaching HIV cure will largely depend on the capacity of HIV-specific memory CD8 ${ }^{+}$T-cells (CD8TC) to eliminate the viral reservoir. However, CD8TC response is limited in subjects on cART. Here, we aimed to investigate the phenotype and function of in vitro expanded CD8TCs in $\mathrm{HIV}^{+}$subjects and the impact of ART initiation timing on these parameters.

Methods \& Materials: PBMCs from $28 \mathrm{HIV}^{+}$subjects on CART for 1 year were obtained. Twelve initiated treatment cluring chronic infection (Delayed Treatment, DT) and 16 within four months post-infection (Early Treatment, ET). PBMCs were stimulated with peptides spanning Nef and Gag plus IL-2 during 14 days. ElisPOI (pre and post-expansion) and Flow Cytometry (FC, post-expansion) were performed to assess expanded CD8TC function (CD107a/b, IFN-g, IL-2, MIP-1b and TNF-a) and phenotype (CD45RO, CCR7 CD95 and PD1). Data was analyzed using non-parametric statistics

Results: Magnitude of ELISPOT responses increased after expansion by $10^{3}$ times ( $p<0.002$ ), in both groups, being this effect more pronounced in CD8TCs, compared to CD4TCs ( $\mathrm{p}<0.0001$ ) as confirmed by FC. Cells showed higher avidity after stimulation (evidenced by greater spot sizes, $\mathrm{p}<0.002$ ). DT subjects displayed a broader response to HIV than ET, after expansion ET group had a significantly higher proportion of monofunct.onal

## Our REF:KWS/VET/3003.24

$3^{\text {rd }}$ March 2020
The Director
Graduate School, Egerton University P.O Box 536-20115,

Egerton.
Dear Pruy Gitaka,

## REF: MSC PROJECT FOR RICHARD NYAMOTA STUDENT No. SM14/23502/14

Kenya Wildlife Service (KWS) is a state corporation mandated to conserve and manage wildlife in Kenya on behalf of the Kenyan people. One of the key responsibility of KWS is to maintain viable and healthy wildlife population. Through its veterinary department KWS regularly collects wildlife samples during scheduled veterinary interventions, translocations and routine medical health checks in collaboration with other government institutions including IPR under existing material transfer agreements (MTA).

Richard Nyamota in his study entitled Prevalence and genetic diversity of simian immune deficiency virus infecting free ranging non-human primates in Kenyan Urban centres was part of KWS routine medical checks and did not require any external permits. Further all molecular analysis related to this work were conducted at the Forensic and Genetics laboratory at Kenya Wildlife Services under my direct supervision in collaboration with experts from IPR and ICIPE.

Yours


Moses Y. Otiende PhD.
Manager Forensics and Genetics Laboratory, Kenya Wildlife Services.

KENYA WILDLIFE SERVICE
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## INSTITUTIONAL REVIEW COMMITTEE (IRC) <br> FINAL PROPOSAL APPROVAL FORM

## Our ref: ISERC/04/18

Dear Dr. Mercy Akinyi,

It is my pleasure to inform you that your proposal entitled "Dynamics of co-infection in wild primates" has been reviewed by the Institutional Review Committee (IRC) at a meeting of $17^{\text {th }}$ April 2018. The proposal was reviewed on the scientific merit and ethical considerations on the use of animals for research purposes.

The committee is guided by the Institutional guidelines as well as International regulations, including those of WHO, NIH, PVEN and Helsinki Convention on the humane treatment of animals for scientific purposes and GLP.

This proposal has been approved and you are bound by the IPR Intellectual Property Policy.


