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1 ***In vivo* cellular tropism of gorilla simian foamy virus in blood of infected**
2 **humans**

3

4 **Running title: Tropism of simian foamy virus in humans**

5

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21

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25 **Abstract**

26 Simian foamy viruses (SFV) are retroviruses that are widespread among non-human
27 primates. SFV can be transmitted to humans, giving rise to a persistent infection.
28 Only few data are available concerning the distribution of SFV in human blood cells.
29 Here we purified blood mononuclear cell subsets from 11 individuals infected with a
30 *Gorilla gorilla* SFV strain and quantified SFV DNA levels by quantitative PCR. SFV
31 DNA was detected in the majority of the CD8+, CD4+ and CD19+ lymphocyte
32 samples, and in rare CD14+ monocyte and CD56+ NK lymphocyte samples. The
33 median (Interquartile range, IQR) SFV DNA were 16.0 (11.0-49.8), 11.3 (5.9-28.3)
34 and 17.2 (2.0-25.2) copies/ 10^5 cells in CD8+ T lymphocytes, CD4+ T lymphocytes
35 and CD19+ B lymphocytes, respectively. Looking at the CD4 compartment, SFV
36 DNA was detected in both memory and naive CD4+ T lymphocytes. SFV DNA levels
37 in CD4+ T cells were positively correlated with the duration of the infection.
38 Our study shows with a quantitative method that CD8+, CD4+ and B lymphocytes are
39 major cellular targets of SFV in the blood of infected humans.

40

41 **Importance**

42 Investigation of simian foamy virus (SFV) infections in humans is important due to the
43 origin of human immunodeficiency viruses (HIV) and human T-cell lymphotropic
44 viruses (HTLV) from cross-species transmission of their simian counterpart to
45 humans. Surprisingly little is known about many aspects of the biology of SFV in
46 infected humans, including quantitative data concerning the cellular targets of SFV *in*
47 *vivo*. Here we show that the distribution of SFV DNA among the different leukocyte
48 populations is not homogeneous, and that viral load in CD4+ T lymphocytes is
49 correlated with the duration of infection. These new data will help in understanding

50 the biology of retroviral infections in humans and can be useful in the growing field of
51 SFV-based gene therapy.

52

53 **Introduction**

54 Like simian immunodeficiency virus (SIV) and simian T cell lymphotropic virus
55 (STLV), simian foamy viruses (SFV) are complex retroviruses that can infect humans
56 after cross-species transmission. No pathology has been reported so far in the
57 natural hosts or in humans, although large-scale dedicated studies have not been
58 performed yet (1,2). Humans are not the natural hosts of foamy viruses, but more
59 than 100 cases of SFV infection have been reported so far within individuals in close
60 contact with nonhuman primates (NHP). In such persons, SFV infection is persistent,
61 and SFV viral load ranges from 1 to >1000 SFV DNA copies/10⁵ blood cells (3-6).
62 While it is known that both retroviruses, HIV-1 and HTLV-1, can infect T cells (7,8),
63 little is known about the distribution of SFV in human blood cells (9). One pioneer
64 study in chimpanzees and African green monkeys revealed a preferential infection of
65 lymphocytes, especially CD8+ T cells, in 13 SFV-infected animals (10). In humans,
66 using semi-quantitative PCR assays, SFV DNA was found exclusively in CD8+ T
67 cells in two tested individuals (10). Interestingly, SFV DNA was detected in
68 monocytes and B lymphocytes, but not in CD4+ and CD8+ T cells, in one individual
69 who was infected with an African green monkey viral strain containing deleterious
70 mutations in the viral *bet* gene (11). However, partly due to the difficulty to obtain,
71 quickly process, and accurately quantify SFV DNA in samples from a series of SFV-
72 infected individuals whose history of infection is known or presumed, no further data
73 are available concerning SFV tropism in humans.

74 In this study, we aimed to quantify and analyze SFV viral load in the major
75 populations of peripheral blood mononuclear cells (PBMC) in a series of 11 hunters
76 from Central Africa who were chronically infected with a gorilla strain of SFV (4). In
77 contrast, the two previous studies were related to SFVs from African green monkeys
78 and chimpanzees and comprise more isolated reports compared to this more
79 systematic study (10,11). Moreover, for the first time, quantitative assays were
80 performed to better evaluate the levels of SFV DNA in the different cell populations.

81

82 **Materials and methods**

83 **Participants**

84 We studied 11 participants; all of them live in villages or settlements in the rain forest
85 of South Cameroon (Table 1). All participants received detailed information about the
86 study and gave their written consent. The study received administrative and ethical
87 clearance in Cameroon from the National Committee of Ethics, and in France, from
88 the 'Comité de protection des Personnes' and the 'Commission Nationale de
89 l'Informatique et des Liberté'.

90

91 **SFV status of enrolled individuals**

92 We enrolled in the present study 11 participants who had been shown to be infected
93 with a gorilla SFV strain in our previous work (4) (Table 1). Briefly, the 11 participants
94 were determined infected with SFV based on both serological results (Western-Blot
95 positivity) and molecular studies on PBMCs (SFV detection by nested- polymerase
96 chain reaction PCR on the pol-in fragment (integrase coding sequence of the
97 polymerase gene) and/or the LTR fragment). All of them have been bitten by a
98 gorilla, which is very probably the cause of SFV infection, as suggested previously

99 (4). Of note, the strain of SFV infecting each individual always matched with the NHP
100 species they were bitten by, as declared in the interview (herein, gorillas).

101

102 **Purification of PBMC subpopulations**

103 Fresh blood samples were collected in EDTA tubes and shipped at room temperature
104 from the Pasteur Institute of Yaoundé to the Pasteur Institute of Paris for processing.
105 Within 20 hours following sampling, fresh PBMC were isolated by a Ficoll-Hypaque
106 gradient. PBMC subpopulations were purified by sequential separation using
107 magnetic beads: CD19+ B lymphocytes, CD14+ monocytes, CD4+ T cells, CD56+
108 NK cells and CD8+ T cells were isolated using the corresponding positive selection
109 kits (Miltenyi Biotec, Paris). For the 'memory study', CD4+ T cells were first selected
110 by the T cell isolation kit followed by positive selection of CD45RO+ memory cells
111 (Miltenyi Biotec, Paris). The flow-through contained the naive CD4+ T cell-enriched
112 fraction. The purity of the separated cells was checked by flow cytometry, using
113 BDFacsCanto for data acquisition and FlowJo V8.7.1 software for data analysis, after
114 labeling with a composition of the following antibodies: anti-CD3-FITC, anti-CD45RA-
115 V450 (BD Biosciences, Rungis, France), anti-CD8-APC, anti-CD14-PE (Beckman
116 Coulter, Villepinte, France), anti-CD16-V450, anti-CD56-PC7, anti-CD20-PC5 (BD
117 Pharmingen, Rungis, France), anti-CD4-APC-H7 (eBioscience, Paris, France).
118 LIVE/DEAD® Fixable Dead Cell Stain Kit (Life Technologies, Saint Aubin, France)
119 was used to identify viable cells. Gating strategy among live cells identified CD3+
120 CD4+ cells as CD4+ T lymphocytes, CD3+ CD8+ cells as CD8+ T lymphocytes,
121 CD3- CD19+ cells as B lymphocytes, CD3- CD14+ as monocytes and both CD3-
122 CD56+ and CD3- CD16+ as NK lymphocytes. The purity of CD4+, CD8+ and CD19+
123 lymphocytes, CD14+ monocytes, and CD56+ NK cells was $94.0\pm 3.1\%$, $87.9\pm 8.1\%$,

124 79.9±7.5%, 73.2±15.2%, and 61.4±16.4%, respectively (Supplementary Table S1).

125 The purity of the CD4⁺ naive cells was 77.1±18.5% and the purity of the CD4⁺

126 memory cells was 86.5±4.1% (data not shown).

127

128 **Quantification of SFV DNA viral loads**

129 Quantitative polymerase chain reaction assays (qPCR) targeting SFV polymerase

130 gene (*pol*) were performed mainly as previously described (5). Briefly, the primers

131 used (GF5qpcr-TAGACCTGAAGGAACCAAAATAATTCC, and GR5qpcr-

132 TCCTTCCTCATATTAGGCCACC) are designed to detect a 144 bp nucleic acid

133 region of the gorilla SFV polymerase gene. We used the Eppendorf realplex master

134 gradient detection system and SYBR Green Quantitect (Qiagen) in a 20 µl volume

135 reaction containing 10 µl of SYBR Green buffer, 150 nM of each primer and 500 ng

136 DNA sample (approximately 75 000 cells equivalent) quantified by ND 8000

137 (Nanodrop Technologies). The optimized qPCR conditions used were as follows:

138 95°C for 15min, 40 cycles of: 95°C for 15s, 60°C for 30s and 72°C for 30s. Samples

139 were tested in triplicate in two independent assays, and the mean and standard

140 deviation were calculated.

141 To standardize quantitative PCR (qPCR), a 465 bp region that included the PCR

142 target sequence from one primary isolate (BAK74, gorilla strain of SFV) was cloned

143 into a PCR cloning plasmid, TOPO TA cloning kit (Invitrogen), and diluted in 500ng of

144 human genomic DNA (Promega). All SFV qPCR assays included a standard curve

145 (from 3 to 3⁷ =2187 SFV DNA copies, 3-fold serial dilutions), performed in triplicate.

146 In each assay, SFV DNA copies from 3 to 3⁷ were in the range of linear detection,

147 and coefficient correlation of SFV standard curves was >0.95. The limit of detection

148 (LOD) was determined as the amount of SFV DNA copies corresponding to the

149 threshold cycle at which at most 5% of true-positive samples scored arbitrarily
150 negative. Using this definition, the LOD was determined as 3 SFV DNA copies/75
151 000 cells equivalent (4 SFV DNA copies/10⁵ cells equivalent). qPCR below the LOD
152 were arbitrarily set up at half the LOD (2 SFV DNA copies/10⁵ cells equivalent).

153 A 141bp-cellular albumin qPCR was performed on a 250x dilution of the samples
154 (2ng) to normalize with cellular DNA content (albF-
155 AAACTCATGGGAGCTGCTGGTT, albR- GCTGTCATCTCTTGTGGGCTGT), using
156 the same PCR conditions as for SFV quantification. All albumin qPCR assays
157 included an albumin standard curve generated by adding 5-fold dilutions of human
158 genomic DNA (Promega) to the PCR reagents, ranging from 0.08ng DNA to 50ng
159 DNA. Coefficient correlation of albumin standard curves was >0.95.

160 We checked in every individual assay the specificity of the primers by using a melting
161 curve.

162

163 **SFV *bet* DNA coding sequences**

164 We performed strain-specific nested-PCR to sequence the entire SFV *bet* coding
165 DNA sequence (CDS) from genomic DNA of PBMC of SFV-infected individuals. We
166 used nested-PCR to generate three PCR products, A (572 bp, *bet* open reading
167 frame orf-1), B (761 bp, 5'-fragment of *bet* orf-2), and C (861bp, 3'-fragment of *bet*
168 orf-2, overlapping with B).

169 The primers used to generate PCR-product A were primers A1 (5'-
170 AACTGGAGAGAGCTAAAGCAG-3') and A2 (5'-CCAGACACCAACATCAGCA-3') for
171 the outer PCR and primers A3 (5'- ATTGGTAACTTCTTAACTGG -3') and A4 (5'-
172 GTAACAGGCATAGGTTTCATG -3') for the inner PCR. The reaction conditions were

173 95°C for 15min, 40 cycles of: 95°C for 15s, 50°C for 1min and 72°C for 1min,
174 followed by 72° for 7 min.

175 The primers used to generate PCR-product B were primers B1 (5'-
176 CATGAACCTATGCCTGTTAC -3') and B2 (5'- CTGATCTGAAAGATTTGCAGC -3')
177 for the outer PCR and primers B3 (5'- TGCTGATGTTGGTGTCTGG -3') and B4 (5'-
178 CTTGATAAGCATATTGGAGCC -3') for the inner PCR. The reaction conditions were
179 95°C for 15min, 40 cycles of: 95°C for 15s, 65°C for 1min, and 72°C for 1min,
180 followed by 72° for 7 min.

181 The primers used to generate PCR-product C were primers C1 (5'-
182 CAGAACCTGATGTATGGTG -3') and C2 (5'- CCTCTCTAGGGATTATACAG -3') for
183 the outer PCR and primers C3 (5'- GGTGTAGTGCAGCACTTTG -3') and C4 (5'-
184 CTCTTCTTACTACTTCC -3') for the inner PCR. The reaction conditions were
185 95°C for 15min, 40 cycles of: 95°C for 15s, 50°C for 1min and 72°C for 1min,
186 followed by 72° for 7 min.

187 The inner primers A3 and A4, B3 and B4, or C3 and C4 were used to sequence the
188 PCR products A, B, or C, respectively. For each donor, one of each PCR product A,
189 B and C was sequenced by direct sequencing of the PCR product (this implies
190 sequencing of the major variants). Each base was sequenced twice, using forward
191 and reverse primers. Sequences were aligned using the software CLC Genomics
192 workbench (Gap open cost: 10.0, Gap extension cost: 1.0). GenBank accession
193 numbers are: KF982250-KF98226 (*bet* orf-1) and KF982262-KF982272 (*bet* orf-2).

194

195 **Statistical analysis**

196 Concerning qualitative analysis, SFV DNA detection rates across the 5 cell subsets
197 were compared using the Cochran's Q test, and two-by-two comparisons were

198 performed by the McNemar's test. Concerning quantitative analysis, the Friedman
199 test was used to compare SFV DNA levels across T and B lymphocytes. Spearman's
200 rank test was used to assess correlations between SFV DNA levels and variables
201 related to SFV infection. Graphpad Prism software was used for statistical analysis.

202

203 **Results**

204 SFV DNA was quantified in selected subsets of peripheral blood cells from 11
205 individuals (Figure 1). SFV DNA was detected in CD8+ T lymphocytes for 10 out of
206 11 (91%) donors, in CD4+ T lymphocytes for 9 (82%) of them and in CD19+ B
207 lymphocytes from 7 (64%) of them. In contrast, SFV DNA levels were above the limit
208 of detection in only two (18%) CD14+ monocyte samples and one (9%) CD56+ NK
209 lymphocyte samples. The frequency of SFV DNA detection across the 5
210 mononuclear cell subsets was significantly different (Cochran's Q test $P < 0.001$). The
211 frequency of SFV DNA detection was significantly higher in CD8+, CD4+ and CD19+
212 B lymphocytes than in CD56+ NK lymphocytes, and in CD8+ and CD4+ T
213 lymphocytes than in CD14+ monocytes (McNemar's test $P < 0.05$ for each two-by-two
214 comparisons). There was a trend for a higher frequency of SFV DNA detection in B
215 lymphocytes than in monocytes, although not reaching statistical significance
216 (McNemar's test $P = 0.07$).

217 We then performed quantitative analysis on CD8+, CD4+ and CD19+ lymphocytes
218 as SFV DNA was generally above the limit of detection in these subsets. The mean
219 (\pm standard deviation, SD) SFV DNA loads were 49.2 ± 78.1 , 19.3 ± 18.9 , and
220 21.1 ± 25.9 copies/ 10^5 cells in CD8+ T lymphocytes, CD4+ T lymphocytes and CD19+
221 B lymphocytes, respectively. The median (Interquartile, IQR) SFV DNA values were
222 16.0 (11.0-49.8), 11.3 (IQR, 5.9-28.3) and 17.2 (IQR, 2.0-25.2) copies/ 10^5 cells in

223 CD8+ T lymphocytes, CD4+ T lymphocytes and B lymphocytes, respectively. SFV
224 DNA levels in CD4+ T lymphocytes, CD8+ T lymphocytes and B lymphocytes were
225 not statistically different (Friedman's test $P=0.12$). SFV DNA load in CD4+ T
226 lymphocytes and B lymphocytes were correlated (Spearman's $\rho=0.646$, $P=0.04$) but
227 neither correlated with SFV DNA load in CD8+ T lymphocytes (CD4+ T lymphocytes:
228 Spearman's $\rho=0.469$, $P=0.15$; B lymphocytes: Spearman's $\rho=0.312$, $P=0.35$).

229 We repeated peripheral blood sampling, cell subset separation and SFV DNA
230 quantification 8-14 months later for three donors (BAD348, BAD468 and BAK74)
231 (Figure 2). For these three individuals, the CD8+ T lymphocytes contained the
232 highest SFV DNA levels at both time points. In CD4+ T cells from BAD468, and in
233 both CD4+ and CD19+ cells from BAK74, SFV DNA was detected at both time
234 points. Finally, samples with undetectable level of SFV DNA at one time point
235 (CD14+ monocytes and CD56+ NK cells from the 3 donors, as well as CD4+ and
236 CD19+ cells from BAD348 and CD19+ cells from BAD468) displayed undetectable or
237 low levels (< 10 copies/ 10^5 cells) of SFV DNA at the other time point. In conclusion,
238 distribution of SFV DNA across blood cell subsets appears mainly stable over a 8-14
239 months period.

240 We quantified SFV DNA load in naive and memory CD4+ T cells from five individuals
241 (Figure 3). SFV DNA load was above the limit of detection in memory CD4+ T cells
242 from all donors and in naive CD4+ T cells from 3 donors. The mean (\pm SD) SFV DNA
243 load was 21.8 ± 18.7 copies/ 10^5 cells in memory CD4+ T cells and 10.1 ± 11.3
244 copies/ 10^5 in naive CD4+ T cells; medians (IQR) were 15.0 (6.2-40.9) and 2.0 (IQR,
245 2.0-18.7) SFV DNA copies/ 10^5 cells in memory and naive CD4+ T cells, respectively.
246 Thus, SFV targets both CD4+ naive and CD4+ memory cells. Due to limitations in the

247 biological material available, analysis of SFV DNA load in naive and memory CD8 T
248 lymphocytes could not be performed.

249 We then studied the association between SFV DNA load and characteristics of
250 infection. SFV DNA load in PBMC tended to be positively correlated with the duration
251 of infection without reaching statistical significance (Spearman's $\rho=0.563$, $P=0.07$,
252 Figure 4A). SFV DNA load in CD4+ T cells was positively correlated with the duration
253 of the infection (Spearman's $\rho=0.632$, $P=0.04$, Figure 4B). In contrast, such
254 correlation was not found for CD8+ and CD19+ lymphocytes (Spearman's $\rho=0.032$,
255 $P=0.93$ for CD8+ lymphocytes, and Spearman's $\rho=0.425$, $P=0.19$ for CD19+
256 lymphocytes, Figure 4C and 4D respectively). These data suggest a selective
257 increase in viral load in CD4+ T lymphocytes over time, when considering long
258 periods of time (over decades). The number of SFV DNA copies in the different cell
259 subsets examined did not correlate with age at infection or age at sampling (data not
260 shown).

261 Interestingly, premature stop codons in the viral accessory gene *bet* were reported
262 for one SFV-infected individual (11). In this individual, SFV DNA was found in CD19+
263 B lymphocytes and CD14+ monocytes cells but it was not detected in CD4+ T cells
264 and CD8+ T cells and the authors suggested that this may be linked to SFV *bet*
265 mutations. Therefore, we sequenced the entire SFV *bet* CDS (1443 bp) from the 11
266 individuals. We did not find deletion events nor stop codons in the 11 *bet* gene
267 sequences (Supplementary Figures S1 and S2).

268

269 **Discussion**

270 Here, we quantified SFV DNA load in the major PBMC subpopulations of 11
271 individuals infected with a *Gorilla gorilla* SFV strain. Previous reports on three

272 infected persons showed that B and T lymphocytes as well as monocytes harbor
273 SFV DNA (10,11). Semi-quantitative analysis of SFV DNA loads in the different
274 PBMC populations of 2 individuals revealed that 10^4 - 10^5 CD8 T cells were sufficient
275 to amplify SFV DNA, indicating viral loads of 1-10 SFV DNA copies/ 10^5 cells, which
276 is slightly lower than our results, and might be related to PCR sensitivity, infecting
277 strain of SFV and/or samples size. Our study is the first that accurately quantifies
278 SFV DNA loads in PBMC populations in a series of SFV-infected individuals. We
279 observed that the frequency of SFV detection was higher in T and B lymphocytes
280 than in NK lymphocytes and monocytes. No difference in SFV DNA levels were
281 observed between T and B lymphocytes. Among CD4+ T cells, SFV DNA was found
282 in both naive and memory CD4+ T lymphocytes. Interestingly, HIV-1 (12) and HTLV-
283 1 (8) preferentially infect memory CD4+ lymphocytes. More data would have been
284 necessary to conclude for a preferential infection of memory CD4+ T lymphocytes if
285 any. Definition of naive and memory CD8+ T lymphocytes and B lymphocytes relies
286 on coexpression of two molecules and their purification requires a cell-sorter in
287 biosafety level 2 laboratory, that was not available at the time of the study. Further
288 experiments will be needed to understand if the observed cellular targets of SFV in
289 humans are linked to biological properties of SFV, such as replication in the different
290 cell subsets, as previously suggested *in vitro* (13). It would also be interesting to
291 know whether infection of the T and B cell progenitor, clonal expansion of
292 lymphocytes and/or homeostatic CD8+ proliferation account for SFV DNA levels.
293 We observed that the distribution of SFV DNA in peripheral blood cell subsets was
294 stable over several months. However, our cross-sectional study suggests that SFV
295 cellular targets may evolve over several decades. We found that SFV DNA load in
296 CD4+ T cells was positively correlated with the duration of the infection whereas this

297 correlation was not observed for CD19+ B cells or CD8+ T cells. Further longitudinal
298 studies will be needed to better understand the evolution of SFV DNA load and
299 cellular target in the course of infection (5). Finally, it would be interesting to have
300 quantitative data of the natural host of SFV infections, gorillas in our case, but blood
301 sampling of a large series of SFV-infected gorillas in this area is extremely
302 challenging.

303 In humans, SFV DNA was found exclusively in CD8+ T cells in two tested individuals
304 (10), and in monocytes and B lymphocytes in one individual who was infected with a
305 viral strain containing deleterious mutations in the SFV *bet* accessory gene (11). Of
306 note, we have also found a premature stop codon in the *bet* gene of the *Pan*
307 *troglydytes troglydytes* strain of SFV (in 3 chimpanzees and 4 humans infected with
308 SFV), resulting in a predicted Bet protein that is shortened by 2% (14). These
309 observations led us to sequence the *bet* coding sequence from the 11 individuals. No
310 sequences harbored stop codons or major deletions, suggesting that at least in our
311 series of individuals infected with a gorilla foamy-virus, deleterious mutations in *bet*
312 are not required for infection of monocytes and B lymphocytes. Of note, 2/11 *bet*
313 sequences were obtained after virus isolation in our previous study (14) and showed
314 >99.5% similarity with the ones presented here. The number of cell fractions with
315 detectable SFV DNA varied between one and five and we did not observe a bimodal
316 distribution of the virus across the mononuclear cell subsets. The variations across
317 the two previous studies and our results may be related to the small size of study
318 populations, the differences in assay sensitivity, the inter-individual variations, or to
319 distinct specificities of SFV from different clades. Indeed, we have studied individuals
320 infected with SFV from gorillas, whereas studies reported by Callahan et al. and van

321 Laer et al. focused on individuals infected with SFV from African green monkey or
322 chimpanzee (10,11).

323 Sample size is small but this comes as a result of the relative rarity of SFV-infected
324 individuals, and the difficulty to obtain and preserve high-quality samples in the field.

325 In most donors, SFV DNA load in purified CD8, CD4 and CD19 subsets were similar
326 or higher than in whole PBMC, in line with the preferential infection of these
327 lymphocyte subsets. For some samples, a fraction of PBMC has partially lost the
328 expression of CD markers after overnight shipment, or was undergoing apoptosis. As
329 a consequence, SFV DNA load could be higher in enriched viable subsets than in
330 whole PBMC (as for donor AKO394). Importantly, SFV DNA loads in different PBMC
331 and/or buffy coat samples from these same individuals were reported by our team in
332 previous studies, and they were in the same range (4,5). Moreover, the lower purity
333 of certain cell subsets (especially NK cells) is another limitation of our study, although
334 SFV DNA in NK cell subsets is generally undetectable. These technical issues do not
335 change our major conclusion of the preferential infection of CD8+ and CD4+ T
336 lymphocytes and B lymphocytes in individuals infected with a gorilla SFV strain.

337 Our findings raise concerns about the consequences of *in vivo* retroviral co-
338 infections, because we find that SFV, as HIV-1 and HTLV-1, show long-term
339 persistence in leukocytes including T lymphocytes. Furthermore, SFV can increase
340 HIV binding at the cell surface *via* heparan sulfate glycosaminoglycans, and can up-
341 regulate HIV-1 LTR transactivation (15,16). Cases of HIV-1/SFV (17) or HTLV-1/SFV
342 (unpublished data) co-infections have been found in humans. In macaques,
343 experimental SIV infection in either non SFV-infected or naturally SFV-infected
344 animals revealed that SFV infection increases the pathology of SIV infection (18).
345 Given the ongoing risk of exposure to these three primate retroviruses, especially in

346 Central Africa, microbiological surveillance is required to better evaluate the potential
347 clinical and physiopathological impact of such retroviral interactions. Furthermore,
348 our findings indicate the cellular distribution of SFV *in vivo*. This could have important
349 implications for the development of SFV-based viral vectors for use in human gene
350 therapy. For instance, recent approaches for treating blood-related diseases consist
351 of direct intravenous administration of SFV viral vectors. Identifying the cellular
352 targets of SFV is critical to better estimate the efficiency and safety of such vectors
353 (19,20).

354

355 **Authorship Contributions**

356 R.R., E.B., T.M. and A.G. performed experiments; R.R. and F.B. analyzed data; R.R.,
357 F.B. and A.G. wrote the manuscript. A.G. supervised the research project.

358

359 **Conflict of interest disclosure**

360 The authors declare no competing financial interests.

361

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Table 1: Epidemiological features of the 11 simian foamy virus (SFV)-infected humans.

Individual ^a	Ethnicity	Age at contact ^b (years)	Age at sampling (years)	Estimated duration of infection (years)	Wound location	Severity ^c	Sampling date and study ^d	Complete SFV genome available ^e
AKO394	Bantu	53	58	5	Thigh	1	18/09/2012 (P)	No
BAD348	Bantu	19	32	13	Leg	3	14/03/2013 (P); 19/11/2013 (P+M)	No
BAD447	Bantu	40	60	20	Hand	2	10/03/2013 (P)	No
BAD456	Bantu	24	34	10	Leg	2	10/03/2013 (P)	No
BAD463	Bantu	37	47	10	Leg	3	19/11/2013 (P+M)	No
BAD468	Bantu	25	38	13	Several	3	04/09/2012 (P); 10/03/2012 (M); 15/11/2013 (P)	Yes (JQ867465.1)
BAK177	Pygmy	26	40	14	Leg	3	15/11/2013 (P)	No
BAK232	Pygmy	40	63	23	Hand	2	19/11/2013 (P+M)	No
BAK55	Pygmy	30	70	40	Arm	2	15/11/2013 (P)	No
BAK74	Pygmy	26	51	25	Foot	2	04/09/2012 (P); 14/03/2012 (M); 15/11/2013 (P)	Yes (JQ867464.1)
LOBAK2	Pygmy	37	64	27	Thigh	3	10/03/2013 (P)	No

^a All individuals lived in South Cameroon rainforest villages/settlements.

^b All individuals had been bitten by a gorilla during hunting activities and are infected with a gorilla SFV (3). Age at contact was assumed, based on interview.

^c Wound bite severity (based on interview and physical injury): 1 = low, 2 = medium, 3 = high.

^d (P) corresponds to sampling for study of the general tropism of SFV in PBMC and (M) corresponds to sampling for study of the tropism of SFV in naive and memory cells.

^e Complete sequence of isolated viral strains from the indicated individuals are available in our previous study, and their GenBank accession numbers is provided (14).

Figure legends

Figure 1: SFV DNA load in PBMC populations of 11 hunters infected with a *Gorilla gorilla* SFV strain

PBMC populations from 11 SFV-infected individuals were isolated and SFV DNA load was quantified in each PBMC population. Each quantification was performed twice in triplicate (e.g. n=6). Means \pm Standard deviations are shown. The values below the limit of detection (LOD) were arbitrary set as half the LOD, 2 SFV DNA copies/ 10^5 cells.

Figure 2: SFV DNA load in PBMC populations of 3 hunters infected with a *Gorilla gorilla* SFV strain at two time points

PBMC populations from 3 SFV-infected individuals were isolated and SFV DNA load was quantified in each PBMC population at two different time points. Each quantification was performed twice in triplicate (e.g. n=6). Means \pm Standard

deviations are shown. NA: not available. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/ 10^5 cells.

Figure 3: SFV DNA load in memory and naive CD4+ T cells of 5 hunters infected with a *Gorilla gorilla* SFV strain

CD4+ cells were separated into naive and memory cells before quantification of SFV DNA load in 5 SFV-infected individuals. Each quantification was performed twice in triplicate (e.g. n=6). Means \pm Standard deviations are shown. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/ 10^5 cells.

Figure 4: SFV DNA load in PBMC, CD4+ T cells, CD8+ T cells and CD19+ B cells as a function of duration of infection

PBMC populations from 11 individuals infected with a *Gorilla gorilla* SFV strain were isolated and SFV DNA load was quantified in each PBMC population. Each quantification was performed twice in triplicate (e.g. n=6) and plotted as a function of duration of infection. Time of infection was based on interviews as described previously (4). Means are shown, as well as Spearman's test results (bold characters). Of note, Spearman's test is non-parametric and thus appropriate for our set of data (no assumptions about the probability distributions of the variables, here viral loads). The linear regression (parametric) is provided for visualization but has no statistical relevance, as the correlation existing between viral loads and duration of the infection is not linear. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/ 10^5 cells.

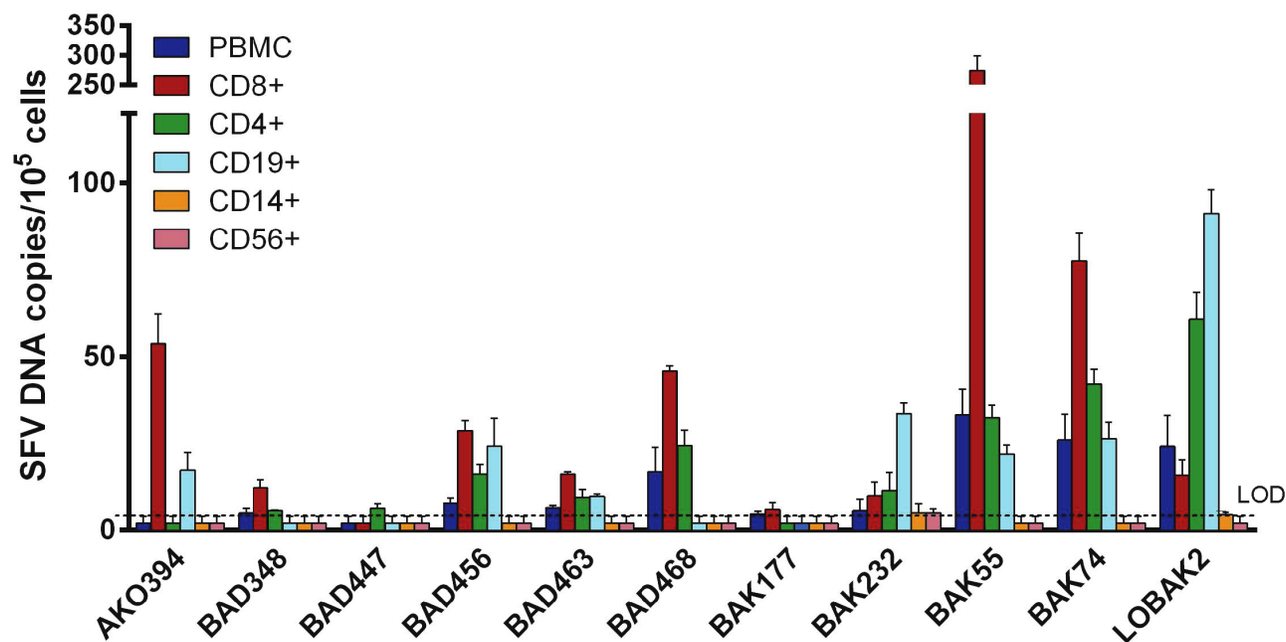


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PBMC populations from 11 SFV-infected individuals were isolated and SFV DNA load was quantified in each PBMC population. Each quantification was performed twice in triplicate (e.g. n=6). Means \pm Standard deviations are shown. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/ 10^5 cells.

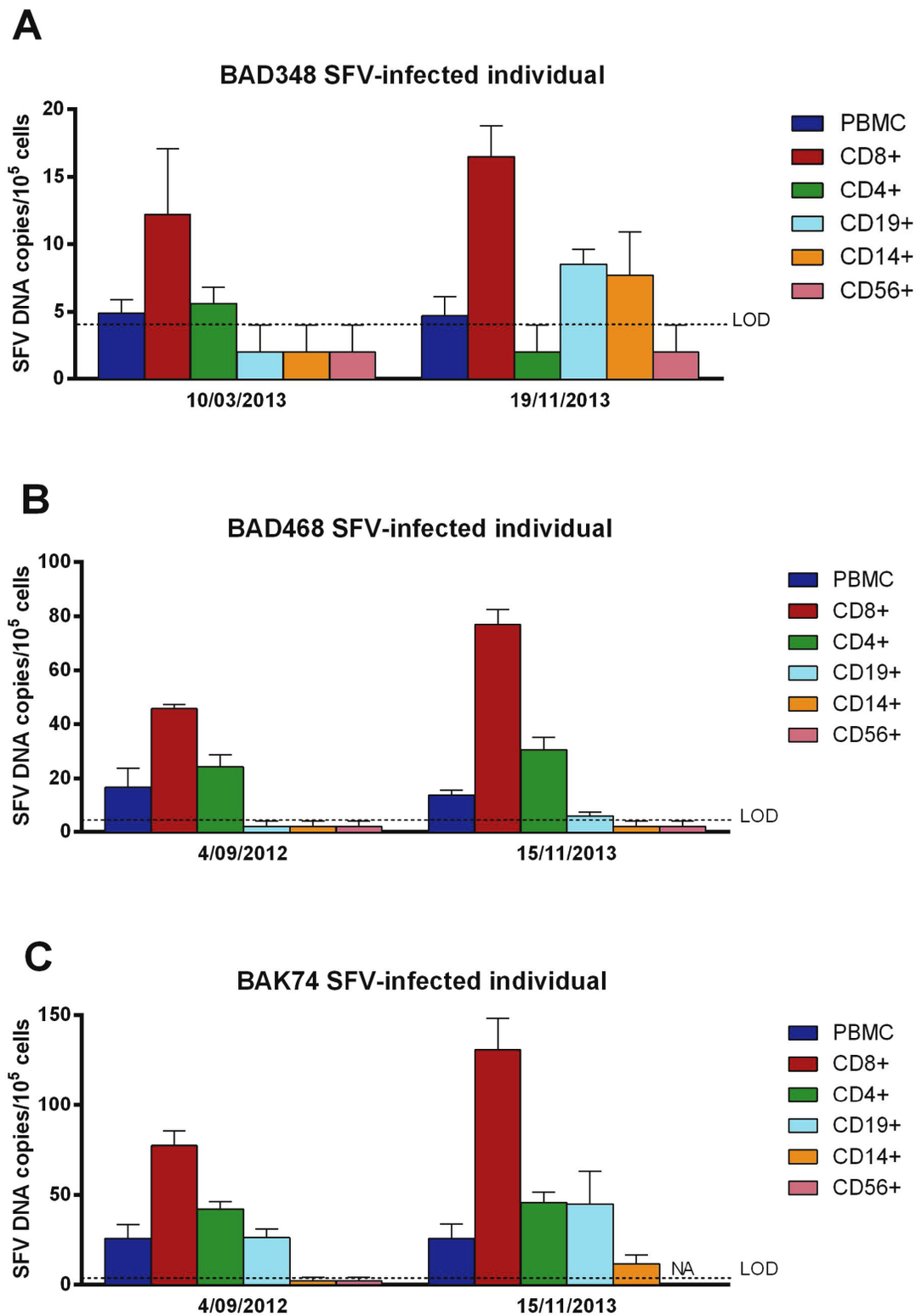


Figure 2: SFV DNA load in PBMC populations of 3 hunters infected with a *Gorilla gorilla* SFV strain, at two time points

PBMC populations from 3 SFV-infected individuals were isolated and SFV DNA load was quantified in each PBMC population at two different time points. Each quantification was performed twice in triplicate (e.g. n=6). Means +/- Standard deviations are shown. NA: not available. The values below the limit of detection (LOD) were arbitrarily set as half the LOD. 2 SFV DNA copies/10⁵ cells.

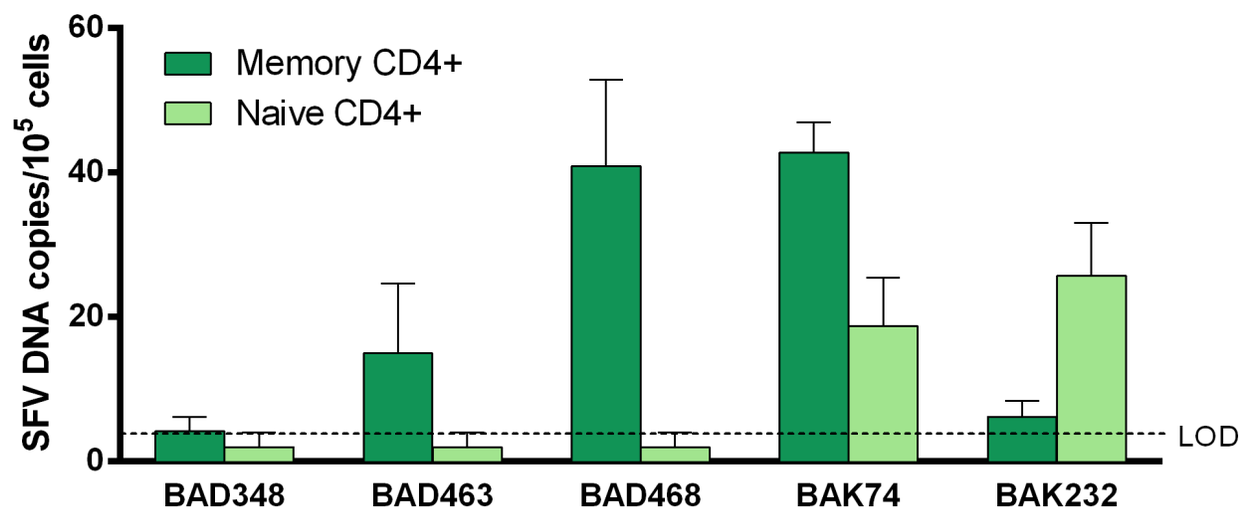


Figure 3: SFV DNA load in memory and naive CD4+ T cells of 5 hunters infected with a *Gorilla gorilla* SFV strain

CD4+ cells were separated into naive and memory cells before quantification of SFV DNA load in 5 SFV-infected individuals. Each quantification was performed twice in triplicate (e.g. n=6). Means +/- Standard deviations are shown. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/10⁵ cells.

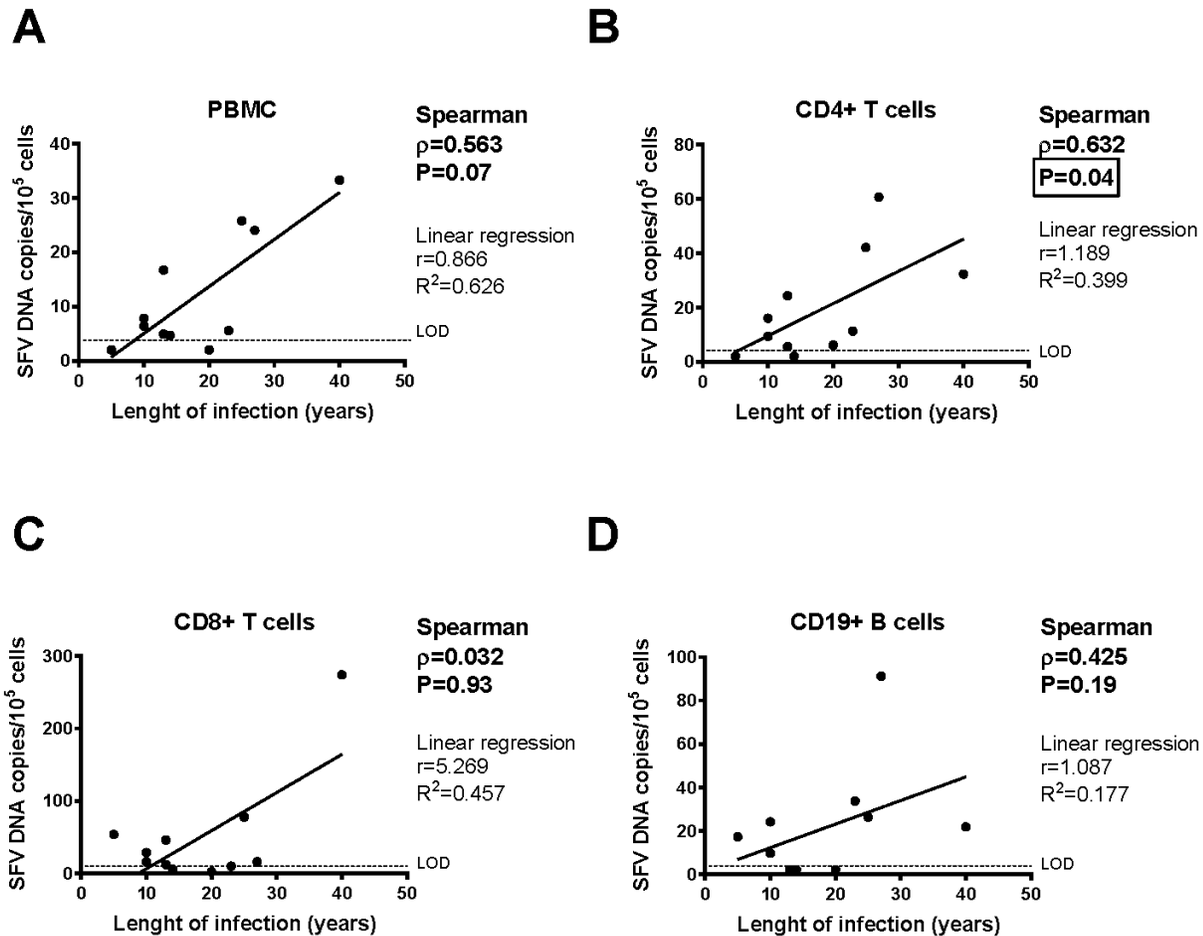


Figure 4: SFV DNA load in PBMC, CD4+ T cells, CD8+ T cells and CD19+ B cells as a function of duration of infection

PBMC populations from 11 individuals infected with a *Gorilla gorilla* SFV strain were isolated and SFV DNA load was quantified in each PBMC population. Each quantification was performed twice in triplicate (e.g. $n=6$) and plotted as a function of duration of infection. Time of infection was based on interviews as described previously (4). Means are shown, as well as Spearman's test results (bold characters). Of note, Spearman's test is non-parametric and thus appropriate for our set of data (no assumptions about the probability distributions of the variables, here viral loads). The linear regression (parametric) is provided for visualization but has no statistical relevance. The values below the limit of detection (LOD) were arbitrarily set as half the LOD, 2 SFV DNA copies/ 10^5 cells.