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2 HIV-1 provirus transcription and translation in macrophages differs from pre-  
3 integrated cDNA complexes and requires E2F transcriptional programs  
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5 Running Title: Active HIV-1 replication favors E2F transcriptional programs  
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22

## 23 Abstract

24 HIV-1 cDNA pre-integration complexes have been shown to persist for weeks in  
25 macrophages and to be transcriptionally active. Early and late gene transcripts are produced,  
26 along with some viral proteins, yet whole virus is not. While previous work has focused on the  
27 transcription and translation of HIV-1 genes; our understanding of cellular milieu that  
28 accompanies viral production is incomplete. We have used an *in vitro* system to model HIV-1  
29 infection of macrophages, and single cell RNA sequencing to compare the transcriptomes of  
30 uninfected cells, cells harboring pre-integration HIV-1 complexes (PIC) and those containing  
31 integrated provirus and actively making late HIV proteins. These are also compared to control  
32 cells, not exposed to virus.

33 Several observations provide new perspective on the effects of HIV-1 transcription from  
34 pre-integrated cDNA versus from integrated provirus. First, HIV-1 transcript levels do not  
35 necessarily correlate with virus production, cells harboring PIC cDNA have transcript loads  
36 comparable to cells transcribing from provirus and making p24, mCherry, and vpu proteins.  
37 Second, all HIV-1 transcripts are easily detectable in abundance from PIC cDNA transcription,  
38 as is the case with cells transcribing from provirus, although the frequency of PIC cells with  
39 detectable gag-pol, tat, env, and nef transcripts is higher than the corresponding frequencies  
40 observed for “Provirus cells”. Third, the background transcriptomes of cells harboring pre-  
41 integrated HIV-1 cDNA are not otherwise detectably altered from cells not containing any HIV-  
42 1 transcript. Fourth, integration and production of p24, mCherry, and Vpu proteins is  
43 accompanied by a switch from transcriptomes characterized by NFkB and AP-1 promoted  
44 transcription to a transcriptome characterized by E2F family transcription products. While some  
45 of these observations may seem heretical, single cell analysis provides a more nuanced

46 understanding of PIC cDNA transcription and the transcriptomic changes that support HIV-1  
47 protein production from integrated provirus.

48

49 **Author Summary**

50 Single cell analysis is able to distinguish between HIV-1 infected macrophage cells that  
51 are transcribing pre-integrated HIV-1 cDNA and those transcribing HIV-1 provirus. Only cells  
52 transcribing HIV-1 provirus are making p24, marker mCherry and Vpu proteins, which  
53 corresponds with a change in the host cell's background transcriptome from one expressing viral  
54 restriction and immunological response genes to one that is expressing genes associated with cell  
55 replication and oxidative phosphorylation.

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58

## 59      **Introduction**

60              The major barrier to curing HIV-1 in patients is a small reservoir of cells that are latently  
61   infected and impervious to immune recognition and clearance [1-3]. The study of HIV-1 latency  
62   is complicated by the fact that latently infected cells *in vivo* are extremely rare. It is a drawback  
63   that many studies of latency have relied on bulk sequencing endpoints. Under these conditions,  
64   the specific parameters defining the latently infected cell are diluted in the context of a vast  
65   heterogeneous population. In addition, multiple mechanisms can result in latency reversal and  
66   therefore one latently infected cell may differ from the next (reviewed in ref. [4]). Thus,  
67   averaging data from a heterogeneous cell population, such as data obtained by bulk sequencing  
68   studies, leads to confusion rather than clarification.

69              Following infection and reverse transcription, the pre-integration cDNA complex (PIC),  
70   in which the HIV-1 genome may take linear or circular forms, serves as a template for  
71   transcription [5-7]. In dividing T cells, the PIC is short-lived, and the transient transcription of  
72   genes from the PIC is considered irrelevant [6]. However, in quiescent T cells the PIC has been  
73   shown to be longer lived and even results in sufficient transcription for virion production in  
74   response to latency reversal agents [8]. In macrophages the dynamics of PIC integration are  
75   different [9-12]. Macrophages have been shown to harbor transcriptionally active pre-integrated  
76   HIV-1 cDNA for months [10, 11], however this PIC mRNA is not thought to result in the  
77   production of virus [12].

78              We have recently used single cell RNA sequencing (sc RNA-seq) techniques [13, 14] to  
79   show that in an activated monocytic cell line, most of the cells in a culture infected with HIV-1  
80   are in fact producing HIV transcripts, while only a minority are producing Gag/p24 and other  
81   proteins. Levels of HIV-1 transcript do not correlate with virus production, since many of the

82 cells harboring PIC complexes have transcript loads comparable to cells making p24. This means  
83 that a high load of viral transcript is not a sufficient switch to reverse latency. Overall, within the  
84 limitations of 10X Genomics technology [15], transcripts for gag-pol, tat, env, and nef are found  
85 in higher numbers of PIC cells, compared to cells transcribing from provirus. However, the loads  
86 of these transcripts detected in PIC and provirus cells is similar. Transcripts for gag, vif, vpr, rev,  
87 vpu, and the marker gene mCherry are detectable in relatively equivalent percentages of cells  
88 transcribing from provirus or PIC, and at similar levels per cell. In no case did “Provirus” cells  
89 (those transcribing from integrated HIV-1 cDNA and producing gag proteins) appear to be  
90 producing any of the detected HIV-1 transcripts at a higher prevalence, or at higher  
91 concentrations per cell.

92 Quite notably, the background transcriptomes of cells harboring HIV-1 PIC are not  
93 detectably altered by PIC transcription. Unsupervised clustering shows cells containing PIC  
94 transcripts to be distributed equally throughout the PIC/Bystander cell cluster. “Bystander” cells  
95 are defined as those cells not containing any detectable HIV-1 transcript. Thus, cells harboring  
96 HIV-1 PIC appear oblivious to the presence of HIV-1 gene transcription, even though some have  
97 been reported to be producing detectable levels of Nef, other HIV-1 proteins and chemokines [9,  
98 10, 15-19].

99 Transcription Factor Targeting analysis [20-22] shows that NFkB and AP-1 transcription  
100 products are predominant in the transcriptomes of PIC and Bystander cells. In contrast, Provirus  
101 cluster cells, on average, contain higher total amounts of viral transcripts and their  
102 transcriptomes predominantly exhibit E2F promoter family transcription products. These cells  
103 make detectable amounts of p24, mCherry and Vpu proteins. This seems counter-intuitive  
104 because NFkB and AP-1 are transcription sites in the HIV-1 LTR that promote transcription

105 from the provirus [23, 24]. In addition, E2F has been reported to suppress HIV-1 transcription  
106 [25, 26]. Nevertheless, in our model it is clear that when cells are making late HIV-1 gene  
107 proteins, the transcriptomes exhibit activation of E2F regulated transcripts, while NFkB and AP-  
108 1 regulated genes are relatively suppressed. Western blotting data agrees with the transcription  
109 factor analysis in that Rb phosphorylation is increased in the Proivirus cluster cells, which  
110 correlates with increased E2F activation.

111 Western blotting also shows that cells transcribing from integrated provirus are making  
112 other HIV-1 proteins. Synthesis of p24 and Gag protein was detected in these cell preparations,  
113 as was mCherry, in positive correlation with flow cytometry results. Vpu protein was also  
114 detected only in protein preparations containing Proivirus cluster cells.

115 E2F domination of the transcriptome is accompanied by activation of regulomes  
116 associated with cell division and RNA processing [27, 28]. The idea that the general cell  
117 transcriptome must transition to support virus production, indicated by E2F promoter family  
118 signaling, is supported by an experiment that shows that cells already making virus are 2 to 5  
119 times more likely to make virus on a second round of infection than PIC or Bystander cells in the  
120 same culture. This was also observed in primary macrophage and T cell cultures, suggesting that  
121 a cell's transcriptional background, in general, is a key factor in determining if virus will be  
122 produced or not.

123

124 **Results**

125 **DHIV3 infection of activated THP-1 macrphages yields cells with**  
126 **transcriptomes containing DHIV3 mRNA but otherwise identical to**  
127 **uninfected cells**

128 We used a VSV-G-pseudotyped DHIV3 virus that expressed mCherry to promote  
129 consistent levels of viral entry in PMA-activated THP-1 cells and to allow for easier  
130 interpretation of post-cell entry events [29, 30] (Fig. S-1). Following infection, flow cytometry  
131 data clearly revealed two clusters of THP-1 cells, mCherry positive cells (usually from 2 to 20  
132 percent of the total population, depending on DHIV3 titer) and mCherry negative cells (Fig. 1).

133 Quantifying DHIV3 infection by scRNA-seq closely agreed with the flow cytometry data  
134 in that two clusters of cells were clearly identified, based on an individual cell's background  
135 transcriptome, in percentages that agreed with flow cytometry of replicate cultures (Fig. 2). Data  
136 analysis is described in detail in the Methods. Briefly, raw fastq files were generated, aligned to a  
137 custom reference genome (GRCh38 augmented with mCherry and HIV genes) and per cell gene  
138 counts generated with the 10X Cell Ranger pipeline. Following basic QC and filtering as  
139 suggested by Seurat, we generated both UMAP and t-SNE clustering projections [15]. Library  
140 construction targeted 5,000 cells and routinely yielded greater than 4,000 cells following Seurat  
141 analysis and quality control. Library construction and sequencing experiments were performed  
142 with technical repeats. The technical repeats were found to be statistically identical and were  
143 therefore combined (Fig. S-2).

144 Following Seurat analysis the number of mean reads per cell was approximately 35,000,  
145 with a median of more than 2,500 genes detected per transcriptome, greater than 15,000 different

146 gene transcripts detected in the overall library. Control cultures yielded slightly larger numbers  
147 of cells captured, with greater than 6,000 cells each and with more than 3,000 gene detected per  
148 cell. All experiments were conducted with parallel cultures that were analyzed by flow  
149 cytometry. In the HIVreplicate1 experiment, flow cytometry indicated 8.6 percent mCherry  
150 positive cells (Fig.1).

151 Figure 2 shows the scRNA-seq UMAP analyses of HIVreplicate1 (t-Sne projection  
152 shown in Fig. S-3). When we quantified HIV-1 transcripts in individual cells, we found that cells  
153 in the smaller, “Provirus” cluster (defined as what we now understand to be cells transcribing  
154 provirus template) were characterized by transcriptomes generally containing higher minimum  
155 levels of HIV-1 transcripts. This “Provirus” cluster accounted for approximately 8.1 % of the  
156 total cell population, and closely matched the percentage of cells identified as mCherry positive  
157 by flow cytometry in the duplicate culture, shown above. In the second “PIC/Bystander” cluster  
158 of mCherry negative cells, we found that many of the macrophages (more than 50%) were  
159 actually producing HIV-1 transcript (Fig. S-3 plots cells with any detected HIV-1 transcript). We  
160 now understand that these cells are “PIC” cells, defined as cells containing HIV-1 mRNA  
161 transcribed from pre-integrated cDNA complexes. Some PIC cell transcriptomes appeared to  
162 contain HIV-1 transcript loads as high as many cells detected in the Provirus cluster, but most  
163 expressed lower amounts (Fig. 2; S-3). Remarkably, these PIC cells had no other statistically  
164 significant changes to their transcriptomes that would differentiate them from the truly  
165 uninfected “Bystander” cells (defined as cells in the PIC/Bystander cluster not containing  
166 detectable HIV-1 mRNA). Thus, PIC and Bystander cells made up the “PIC/Bystander” cluster.

167 We used a Feature map to plot the influence of cell cycle, number of genes detected or  
168 mitochondrial transcript number on the distribution of cells containing HIV-1 transcripts (Fig 3

169 B-D). None of these factors had any significant influence of the distribution of PIC cells  
170 throughout the PIC/Bystander cluster. We then used unsupervised clustering and violin plots of  
171 HIV transcript load to examine the distribution of cells containing HIV-1 transcripts (Fig. 4)  
172 throughout the Proivirus and PIC/Bystander clusters. Using K-nearest neighbors clustering, at  
173 K10, 2 clusters of cells (clusters 6 and 8) were identified that accounted for most of the Proivirus  
174 cluster cells (372 of the 380 cells), as determined by semi supervised clustering shown above  
175 (Fig. 2). Therefore, the combined transcriptomes of cells in clusters 6 and 8 were compared to  
176 the combined transcriptomes of cells in clusters 1-5, 7 and 9-10, and used to generate the DGE  
177 analyses shown below. Unsupervised clustering using a range of designated K values from 10 to  
178 130 is shown in Figure S-4.

179 Clusters 6 and 8 (Fig. 4B) were characterized as containing few cells with low HIV  
180 transcript load. Thus, cells in what we define as the Proivirus cluster differ significantly from PIC  
181 cells in terms of average minimal DHIV3 transcript load. They also differ significantly in  
182 background transcriptome, as shown below. The multiple clusters generated in this unsupervised  
183 clustering are likely stochastic. Most importantly, they were not influenced by the presence or  
184 absence of PIC cells (Fig. 4B). PIC cells are distributed throughout all the PIC/Bystander  
185 clusters, regardless of the specified K value. So, for our purposes, the two cluster model shown  
186 in Figure 2 was considered to accurately represent the interpretation gained from flow cytometry  
187 (Fig. 1), namely: either the cells were making mCherry protein, or they were not.

188 Inferring from our flow cytometry percentages, we hypothesized that only cells in the  
189 Proivirus cluster were making fluorescing mCherry protein. To confirm that cells making  
190 mCherry were also making late viral proteins, the production of p24 capsid protein was  
191 quantified using flow cytometry and monoclonal mouse IgG-AG3.0 anti-Gag p24 antibody.

192 Cells were fixed in formaldehyde, and secondary anti-mouse Alexa Fluor antibody allowed  
193 detection during flow cytometry. When cells were analyzed using FACS Canto, only cells  
194 making mCherry were found to stain for p24 (Fig. 5). This suggests that the presumed PIC cells  
195 are not making late viral proteins in detectable amounts, while only Proivirus cells appear to be  
196 making late viral proteins.

197

## 198 **The DHIV3 mRNA in “PIC” cells is due to PIC transcription**

199 Integrase inhibitors have proven to be useful tools in the study of PIC transcription [16].  
200 We used 25nM MK-2048 [31] for our experiments to confirm the “Proivirus” versus “PIC” status  
201 of our cell clusters. When integrase inhibitor treated cultures were analyzed by flow cytometry,  
202 mCherry producing cells were routinely reduced from an average of about 12% to less than 2%  
203 of the total (Fig. 6). The use of the viability stain assured that this was not due to death of  
204 infected and drug-treated cells. We then analyzed protein preparations from parallel cultures of  
205 these cells by Western blotting. For the Western analysis, we initially purified populations of  
206 mCherry positive and mCherry negative cells using FACS analysis, collecting cells in the viable  
207 windows shown in Figure 6. However, because of the laborious difficulty in obtaining sufficient  
208 quantities of protein from the sorted cells, we resorted to comparing whole cultures of cells  
209 obtained either in the presence or in the absence of integrase inhibitor. We achieved >80%  
210 inhibition of mCherry production by integrase inhibitor treatment in these experiments.  
211 However, this means that in the integrase inhibitor treated/DHIV3 infected cell protein  
212 preparations there probably remained proteins from some mCherry positive cells, albeit in  
213 amounts greatly reduced from amounts present in the protein preparations from the  
214 corresponding DHIV3 infected cultures without integrase inhibitor treatment. Furthermore, in the

215 DHIV3 infected cell preparations in the absence of integrase inhibitor, the majority of protein  
216 (>80%) still came from the mCherry negative cells. Both these protein preparations were  
217 compared to an equivalent amount of protein from Control cells. Control cell protein  
218 preparations were from PMA activated THP-1 cells not exposed to DHIV3.

219 We found protein preparations from DHIV3-mCherry cultures in the absence of MK-  
220 2048 to contain mCherry (Fig. 7 A). In addition, these same protein preparations contained p24  
221 and p24 precursor proteins (Fig. 7 B). As expected, p24 protein was also detectable in protein  
222 from the integrase inhibitor treated culture. In the literature, this observation has been attributed  
223 to residual p24 protein lingering from the initial infection [10]. Fortunately, the p24 antibody we  
224 used is known to bind both p24 and Gag precursor proteins [10, 32]. We found both p24 and  
225 precursor p24 proteins only in the protein samples from cells not treated with integrase inhibitor.  
226 We obtained additional support for this interpretation by taking a 48 hr time point (Fig. 7D). The  
227 extended time point showed p24 precursor proteins only appearing in the integrase inhibitor  
228 cultures 48 hours after treatment. In contrast, p24 and Gag precursor proteins were increasingly  
229 detectable in cell proteins from integrase competent infections. Thus, p24 detected in DHIV3  
230 infected cells was likely due to residual p24 from the original infection.

231 Further confirmation of the PIC versus Proivirus status of our semi supervised cluster cell  
232 populations was obtained using real-time polymerase chain reaction (real-time PCR) analysis  
233 [33, 34]. DNA from the respective cell treatment groups described above (Control, DHIV3  
234 infected, DHIV3 infected plus integrase inhibitor) was isolated and analyzed by PCR using  
235 multiple sets of primers. For detection of integrated proviral DNA a set of primers [34] was used  
236 to amplify from random nascent human genomic Alu sequences to an internal HIV LTR  
237 sequence. This initial amplification step was followed by a second PCR amplification step using

238 nested primers, which would only amplify discrete DNA products that contain integrated  
239 provirus [34]. Evidence for integrated provirus was detected in much higher abundance from  
240 DNA of DHIV3 infected cells in the absence of integrase inhibitor (Fig S-5), although small  
241 amounts of integrated provirus DNA was detected in DHIV3-mCherry infected and integrase  
242 inhibitor treated cell culture DNA, when using higher amounts (200 ng) of starting DNA. This is  
243 in agreement with Flow cytometry results (Fig. 6) and Western analysis (Fig. 7) indicating small  
244 amounts of proviral DHIV3-mCherry DNA in our integrase inhibitor treated cultures.

245 To demonstrate unequivocally that our PIC cluster cells do indeed contain PIC HIV  
246 cDNA, we used the primers of Brussels and Sonigo [33], which are internal to the HIV-1 LTR  
247 sequence. These primers were oriented in a way so as to detect only circular 2-LTR PIC HIV-1  
248 DNA by bridging the 2-LTR junction [33, 34]. We found that we required 2 rounds of PCR  
249 amplification to obtain the predicted 2-LTR amplicon, suggesting that this particular PIC species  
250 is in low abundance in our model cells. We tested this conclusion using bracketing primers (see  
251 Methods) to generate a product to contain the predicted amplicon of Brussels and Sonigo [33],  
252 and then followed with a round of amplification using the previous primers to generate a nested  
253 product. This approach also generated the predicted amplicon product from DHIV3-mCherry  
254 infected culture DNA, whether in the presence of integrase inhibitor or not. The predicted 2-LTR  
255 PIC cDNA PCR product was not detected in Control cell DNA.

256 To confirm that total PIC cDNA is in relatively high abundance in our DHIV3-mCherry  
257 infected cultures, we adapted previous primers to amplify total DHIV3 LTR DNA. With these  
258 primers, similarly high levels of total PIC cDNA was detectable in DHIV3 infected cells,  
259 whether in the presence of integrase inhibitor or not (Fig S-5).

260                    Taken together, the Western blot and real-time PCR data confirm our flow cytometry  
261                    observations that mCherry producing cells were also producing p24. The fact that the mCherry  
262                    cells are selectively suppressed by integrase inhibitor treatment leads to the conclusion that only  
263                    Provirus cluster cells are making p24 or mCherry proteins from the DHIV3 transcripts.  
264                    Conversely, transcripts detected in PIC cells, due to PIC transcription, and do not lead to  
265                    detectable mCherry or p24 synthesis.

266                    We tried antibodies for all the other major HIV-1 proteins to test the correlation of  
267                    protein expression with the detection of transcript. One antibody that yielded a clear result was  
268                    the polyclonal antibody against Vpu. In this case, we see a result very similar to that obtained  
269                    with mCherry and p24, in that protein was clearly detected in samples from infected cell cultures  
270                    that were not treated with integrase inhibitor (Fig. 7). Vpu was not detectable in protein from the  
271                    integrase inhibitor treated cells. It is an interesting side-note that Vpu has been associated with  
272                    inhibition of NFkB promoted transcription [35, 36].

273

274                    **scRNA-seq analysis of integrase inhibitor treated DHIV3 infected  
275                    cultures.**

276                    DHIV3 infected THP-1 cells were treated with 25nM MK-2048 at the time of infection,  
277                    using our established protocol, and analyzed by scRNA-seq. In this experiment, the integrase  
278                    inhibitor blocked ~87.5% of mCherry production by flow cytometry analysis of a parallel culture  
279                    (Fig. 6). The UMAP image of DHIV3 transcript features is shown in Figure 8A. Transcriptome  
280                    clustering using varying nearest neighbor K-values (K10, K50, K90, and K130) yielded 3 to 7  
281                    clusters, depending on the K-value applied (S-6). Regardless of the specified K-value, we did not  
282                    detect a Provirus cluster as was observed with integrase competent infections (Fig. 8C). Again,

283 the distribution of HIV-1 transcript containing cells throughout the semi supervised integrase  
284 inhibitor single cell cluster was not effected by cell cycle, percent mitochondrial gene expression  
285 or numbers of genes detected per cell (Fig. S-7). This result confirms that integrase inhibitor-  
286 treatment selectively suppresses cells in the Proivirus cluster, agreeing with results obtained by  
287 Western blot and PCR analysis, and confirms that the DHIV3 mRNA detected in the PIC cluster  
288 cells is due to transcription of PIC complexes.

289

290 **Hallmark and REACTOME analyses indicate mitogenic associated**  
291 **pathways are up-regulated in Proivirus cluster cells whereas viral**  
292 **restriction and interferon associated pathways are upregulated in**  
293 **PIC cluster cells.**

294 Differential gene expression (DGE) in Proivirus versus PIC/Bystander cluster  
295 transcriptomes was analyzed using GSEA with Hallmark or REACTOME gene lists (Appendix  
296 I), the pairwise TTests function from Scran was used to determine the statistically significant  
297 differentially expressed genes between groups. Table I presents the statistically significant  
298 Hallmark results. By comparing the DGE between the Proivirus and PIC/Bystander cell clusters  
299 using Hallmark and REACTOME tools, it was clear that Proivirus cluster cells were  
300 differentiated from PIC/Bystander cluster cells in several key ways. In general, the  
301 transcriptomes of cells in the Proivirus cluster were characterized by gene transcripts associated  
302 with cell replication, whereas the transcriptomes of cells in the PIC/Bystander cluster were  
303 characterized by pathways associated with immune-response and interferon signaling. The  
304 analyses clearly showed upregulation of cell replication, oxidative phosphorylation, protein

305 synthesis and E2F family targeted pathways in the Proivirus cluster cells. On the other hand,  
306 NFkB, AP1, interferon responsive and immune response pathways relatively upregulated in the  
307 PIC/Bystander cluster cell transcriptomes. Intuitively, this makes sense, but it runs contrary to  
308 established literature that associates E2F transcription factors with decreased viral production  
309 [24, 25]. This finding would not be obvious without the use of single cell analysis. In the absence  
310 of single cell analysis the Proivirus cluster's differentially expressed gene transcripts would have  
311 been swamped out by the 90% of mRNA obtained from PIC and Bystander cells.

312 Using the UMAP Feature plots shown in Figure 9, we visualized the distribution of cells  
313 containing some of the most statistically significant differentially expressed transcripts in our  
314 Proivirus cell transcriptomes versus the PIC/Bystander cell transcriptomes. The GSEA lists of the  
315 differentially expressed genes are provided in supplementary Appendix I. This leads us to  
316 hypothesize that the cell's transcriptome background is what determines if virus transcript will  
317 lead to virus protein production, not the differential transcription of HIV-1 genes. This  
318 hypothesis was supported by Transcription Factor Targeting and Western blot analyses (below).

319

320 **PIC and Proivirus Cells express all DHIV3 genes at equivalent levels,**  
321 **although higher numbers of PIC cells detectably express some**  
322 **transcripts.**

323 We were interested to know if early HIV-1 gene transcripts (tat, nef, or rev)  
324 predominated in PIC cell transcriptomes, versus later transcripts in the Proivirus cells. We used  
325 Feature plots to determine the distribution of selected HIV gene transcripts expressed in

326 individual cells. It was found that cells expressing the respective early or later gene transcripts  
327 were distributed throughout the image (Fig. 10 A).

328 We then used violin plots to compare the load of individual DHIV3 gene transcripts in  
329 Provirus cluster cells to PIC cells (Fig. 10 B). The Provirus cluster (from Fig. 4) contained  
330 transcriptomes of 372 cells, the PIC/Bystander cluster contained 569 cells, thus the ratio of  
331 Provirus to PIC cells was ~0.65. This visualization was much more informative and yielded a  
332 more nuanced understanding (Fig. 10 B). Some transcripts, such as gag-pol, tat, env and nef were  
333 detected in far more cells in the PIC cluster, albeit often at the lower level of detectable loads per  
334 cell. In contrast, gag, vif, vpr, rev, vpu and mCherry, were clearly detectable, but in lower  
335 numbers of Provirus and PIC cells. Cells containing these transcripts were comparably prevalent  
336 in the two groups. All transcripts were easily detectable in both Provirus cells and PIC cells.  
337 Furthermore, there is a clear overlap in the levels of HIV-1 gene transcripts detected in Provirus  
338 and PIC cells.

339 During transcription of pro-virus, HIV-1 does not produce all transcripts in equal number  
340 or at the same time [37, 38]. The specific processed gene transcripts produced initially in viral  
341 replication differ from transcripts produced later on. Furthermore, 10X Genomics scRNA-seq  
342 library production is known for significant numbers of dropouts, and cDNA copying of various  
343 gene transcripts during library construction varies in efficiency [15, 39, 40]. Furthermore, in  
344 using poly-T primers in the cDNA library construction, the 10X process introduces a 3' bias  
345 toward the detection of given sequences in a transcript [15]. Thus, it is not possible to make  
346 quantitative comparisons between the different transcripts using this approach. Nevertheless, the  
347 overarching take-away from this single cell analysis is that cells making fully spliced transcripts  
348 such as nef, tat, and nev are equally prevalent with cells making gag-pol transcripts (Fig. 10).

349 Furthermore, there appears to be two patterns of transcription from PIC. One pattern, observed  
350 with gag-pol, tat, env, and nef, is characterized by gene transcripts being more frequently  
351 detectable in PIC cells than Provirus cells. The other pattern, observed with gag, vif, vpr, rev,  
352 vpu, and mCherry, suggests relative equal frequency of transcription in Provirus and PIC cells.

353

### 354 **Biological repeat experiments confirm observations.**

355 To confirm the conclusions obtained from the analyses presented above, an independent  
356 biological repeat experiment was conducted. The repeat experiments captured over 4,500 cells,  
357 with an average greater than 3,000 genes per transcriptome. The control cell cultures again  
358 yielded a slightly larger number of cells captured with more than 3,000 genes transcripts per cell.  
359 The biological repeat experiments, (HIVreplicate2 and WT2) were also conducted with parallel  
360 cultures that were analyzed by flow cytometry. Flow cytometry indicated 2.6 percent mCherry  
361 positive cells in the HIVreplicate2 experiment. Figure S-8 shows a UMAP analysis of the  
362 biological repeat experiments HIVreplicate1 and HIVreplicate2. Figure S-9 shows unsupervised  
363 clustering of HIVreplicate2 using K nearest neighbor values from 10 to 130. Figure S-10 shows  
364 distribution of respective DHIV3-mCherry transcripts throughout HIVreplicate2 UMAP.

365 Several comparisons were made to confirm that transcriptomes from Provirus and  
366 PIC/Bystander clusters in the repeat experiments were identical. In the first comparison,  
367 differentially expressed genes (positive or negative) were identified between the Provirus and  
368 PIC/Bystander clusters in the respective biological repeats. The log2 fold changes from these  
369 gene sets were then compared to test if the differences between the transcriptomes of Provirus  
370 and PIC/Bystander cells in the repeat experiments was consistent (Fig. S-11). Because there  
371 almost 4 times as many Provirus cluster cells in the HIVreplicate1 experiment, there were more

372 significantly differentially expressed genes in the HIVreplicate1 case, compared to the  
373 HIVreplicate2 case. Nevertheless, two the gene sets were positively correlated.  
374 Correspondingly, GSEA with Hallmark and REACTOME gene sets identified many of the same  
375 pathways as differentially regulated in the two biological repeats (Appendix I).

376 To obtain additional statistical certainty for our conclusion that results obtained for  
377 HIVreplicate1 were repeated in HIVreplicate2, we compared log2 fold change values from the  
378 Proivirus or PIC/Bystander clusters in HIV infected cells to the log2 fold change values from the  
379 Control (WT) PMA-treated THP-1 cultures. This 8-way comparison (shown in Fig. 11) provided  
380 statistical certainty that DGE sets from the Proivirus cluster and PIC/Bystander cluster gene sets  
381 from the biological replicates were not different. The replicate Proivirus and PIC/Bystander gene  
382 sets have a generally strong concordance between themselves and there is a modest to strong  
383 non-zero mean trend in logFC among genes that changed in at least one of the contrasts among  
384 replicates (FDR 5%). In every comparison, a significant positive correlation was obtained from  
385 the commonly detected, significantly differentially expressed genes of Proivirus or PIC/Bystander  
386 clusters in the two biological repeats when compared to the Control samples. The weakest  
387 correlations were observed in comparing PIC/Bystander to Control cell DEGs, especially Control  
388 experiment 2 (probably because there is more commonality between genes detected in  
389 PIC/Bystander cells and Control cells than there is between Proivirus cluster cells and Control  
390 cells), but still the data between HIVreplicate1 and HIVreplicate2 were concordant. Therefore,  
391 the transcriptome data obtained from the 2 biological repeat experiments were not different. In  
392 other words, statistically identical representative transcriptomes for Proivirus and PIC/Bystander  
393 clusters were obtained in independent biological repeat experiments.

394 As was the case for HIVrepeat1, a clear Provirus cluster was detectable in the  
395 HIVrepeat2 (Fig. 12, S-8). However, because the level of Provirus infection in HIVrepeat2 was  
396 lower than in HIVrepeat1, the frequency of DHIV3 transcript detection in the Provirus cluster  
397 cells was proportionately lower, while the absolute number of detectable PIC cells was relatively  
398 higher. As was observed for HIVrepeat1, PIC cells with detectable HIV-1 transcripts, were  
399 randomly distributed throughout the Bystander cluster. Unsupervised clustering (Fig. 12, S-9)  
400 generated 10 clusters at a K nearest neighbor values of 10. Clusters 1, 2 and 4-10 contained most  
401 of the cells in the semi supervised PIC/Bystander from Figure 12. Cluster 3 contained 135 of the  
402 227 cells in the semi supervised Provirus cluster (circled in red, Fig. 12), and was compared to  
403 the combined transcriptomes of the remaining clusters to generate the violin plots in Figure 13.  
404 The patterns of transcription observed in HIVrepeat1 were confirmed in this experiment. The  
405 transcripts of gag, vif, vpr, vpu, and mCherry were detectable in PIC cells at frequencies and  
406 levels of expression similar to those observed in the Provirus cells. In contrast, even correcting  
407 for the Provirus/PIC ratio of 0.17, the transcripts of gag-pol, tat, env, and nef were again  
408 detectable in proportionally higher number of PIC cells. Again, there was clear overlap in the  
409 loads of HIV-1 gene transcripts detectable in Provirus and PIC cells. No cells expressing rev  
410 were detected in the Provirus cluster in this repeat, due either to the low efficiency of detecting  
411 this transcript in the 10X system, or because rev is expressed only at low levels in relatively few  
412 cells, or both.

413

414 **Psupertime analysis indicates progression of cluster transcriptomes  
415 from Control to PIC/Bystander to Provirus.**

416 To understand the transcriptome transitions needed to move from unexposed and  
417 uninfected “Control” cells to PIC/Bystander cells, and on to Provirus cluster cells, we performed  
418 a psupertime analysis [41-43] of the respective cell cluster transcriptomes. Psupertime is a  
419 supervised pseudotime [41] technique. It explicitly uses sequential condition labels as input.  
420 Psupertime is based on penalized ordinal logistic regression that places the cells in the ordering  
421 specified by the sequence of labels. This allows for targeted characterization of processes in  
422 single cell RNA-seq data.

423 One thousand cells were randomly selected from each transcriptome cluster (Control,  
424 PIC/Bystander and Provirus) and their transcriptomes combined for psupertime analysis.  
425 Imposition of Cluster identity yielded the image shown in Figure 14. The psupertime-type  
426 analysis showed closer similarity between Control and PIC/Bystander transcriptomes than  
427 between Control and Provirus transcriptomes, and closer similarity between PIC/Bystander and  
428 Provirus transcriptomes than between Control and Provirus. The GSEA list of the DGEs that  
429 contributed to this faux-progression from Control to PIC/Bystander to Provirus are presented in  
430 Appendix II.

431 When we examined the expression of DHIV3 transcripts through the psupertime  
432 progression, the analysis showed no obvious preference for early gene transcription in cells  
433 belonging to the PIC/Bystander versus the Provirus clusters (Fig. 14B).

434 When questioning which transcription factors were regulating the transcriptome  
435 transitions, we searched the contributory psupertime DGE transcripts for transcription factors.  
436 Many transcription factors that differed in expression in the contrasted transcriptomes were  
437 identified, Appendix II. However, this yielded a complicated picture and did not clarify which  
438 transcription factors might be most important for controlling the transcriptome transitions from

439 Control to PIC/Bystander to Provirus clusters. However, because the activity of most  
440 transcription factors is regulated by activation of proteins already present within the cell, and not  
441 at the transcription level, we speculated that Transcription Factor Targeting analysis might be  
442 more informative as to which transcription factors were key to cluster transitions.

443

444 **E2F, NF- $\kappa$ B and AP1 control phenotype transitions between**  
445 **PIC/Bystander cells and Provirus cells**

446 We used Transcription Factor Targeting analysis to identify transcription factors that  
447 controlled DGEs in our Control, PIC/Bystander and provirus clusters. This analysis was  
448 consistent with the aforementioned Hallmark and REACTOME analyses. The E2F family of  
449 transcription factors predominate in regulating the Provirus cluster DGEs (Table II). Twenty out  
450 of the 29 possible promoter-associated transcription factor interactions positively associated with  
451 the transition from the PIC/Bystander to the Provirus transcriptome identified with the E2F  
452 transcription factor family (Table 2). Thus, E2F is clearly associated with pathways that  
453 determine the phenotype of cells in the Provirus cluster. Conversely, 19 possible promoter-  
454 associated transcription factor interactions were negatively associated with DGEs reflecting the  
455 transition from PIC/Bystander to Provirus cells. Of these 19 possible promoter complexes, 8  
456 different promoter interactions were identified to be associated with NF $\kappa$ B and AP1 transcription  
457 (Table 2) suggesting that downregulation of NF $\kappa$ B and AP1 also plays a key role in shaping the  
458 Provirus cluster cell transcriptome.

459 AP-1 and NFkB appear to play roles in maintenance of the PIC/Bystander cell  
460 transcriptome as well. Correspondingly, Transcription Factor Targeting identified differences  
461 between Control and PIC/Bystander cell transcriptomes. Simple exposure of activated THP-1  
462 cells to DHIV3 was sufficient to decrease E2F signaling in PIC/Bystander cells, compared to  
463 Control cells, and to increase AP-1 and NFkB signaling (Appendix III). We presume this effect  
464 on the PIC/Bystander cells is through PAMP and/or interferon signaling. Again, these results are  
465 consistent with the results obtained with Hallmark and REACTOME analysis of the DGEs.

466

467 **Western Blot analysis of Retinoblastoma protein phosphorylation  
468 from Control, PIC/Bystander and Proivirus cells.**

469 To confirm a role for E2F and NFkB in regulating the transcriptomes of Proivirus and  
470 PIC/Bystander clusters (respectively), we sorted Proivirus (mCherry positive cells) from PIC  
471 /Bystander cells using the FACS Canto. Rb phosphorylation is associated with activation of E2F  
472 promoter family transcription. We hypothesized Proivirus cluster cells would exhibit  
473 retinoblastoma (Rb) phosphorylation consistent with E2F activation [27, 42]. We used anti-T821  
474 Phospho-Rb antibody. Phosphorylation of Rb at threonine-821 (T821) blocks pocket protein  
475 binding, including E2F family proteins, and activates E2F family promoter gene transcription  
476 [44]. Isolated Proivirus cells had the greatest phosphorylation of Rb (pRb), compared to Control  
477 and PIC/Bystander cells (Fig. 15). Interestingly, phosphorylation of Rb in PIC/Bystander cells  
478 was actually lower than that detected in Control cells (Fig. 15). We also probed these blots for  
479 mCherry protein as was shown in Figure 7. In this case, with cells being sorted before protein  
480 preparation, mCherry protein detection was detectable and only in the Proivirus cluster cells.

481 These findings are consistent with the Transcription Factor Targeting analysis, which  
482 showed that E2F promoter family transcription was predominant in Proivirus cluster cells. It also  
483 agreed with the Transcription Factor Targeting analysis in that PIC/Bystander cells exhibited  
484 reduced Rb phosphorylation, and presumably E2F driven transcription, when compared to either  
485 Proivirus cluster of Control cells. We did not find a difference in NFkB deactivation in the  
486 Proivirus cells as would be implied by phospho-IkB S32, (Fig. 15), and hypothesize that other  
487 mechanisms must account for the relative decrease in NFkB driven transcription in Proivirus  
488 cluster cells.

489

490 **Cells transcribing from provirus are more likely to produce viral  
491 proteins upon second infection than PIC or Bystander cells.**

492 If Proivirus cluster cells have already committed to the production of virus, represented by  
493 the switch of their background transcriptome to favor E2F transcription factor interactions, we  
494 hypothesized that they should be more efficient at producing virus upon second infection. We  
495 sequentially infected of activated THP-1 cells with DHIV3-mCherry followed at 24 hr by an  
496 infection with DHIV3-GFP (DHIV3-mCherry infection at 0 hr and DHIV3-GFP infection at 24  
497 hr). We found a higher percentage of cells positive for mCherry and GFP after 48 hr compared to  
498 GFP alone (Fig. 16). At this time point, which was 24 hours after DHIV3-GFP infection, about  
499 half the mCherry positive cells were also GFP positive. Whereas, less than one quarter of the  
500 mCherry negative cells were expressing GFP protein. This trend continued out to 72 hours post  
501 DHIV3-mCherry infection, 48 hours after DHIV3-GFP addition, where about 60% of the  
502 mCherry cells were also GFP positive, compared to about 40% GFP positive in mCherry  
503 negative cells. In repeat experiments and in experiments using primary macrophage and T-cell

504 cultures (Fig. S-12) the results were the same. Provirus cluster, mCherry positive cells, were 2X  
505 to 5X more likely to make DHIV3-GFP protein upon second infection than PIC/Bystander  
506 cluster cells. Thus, cells already committed making virus were more likely to make virus from a  
507 second infection than PIC/Bystander cells on first infection.

508

## 509 **Discussion**

510 Early research into the HIV-1 life cycle identified transcription of HIV-1 PIC cDNA in  
511 T-cells and macrophages [5-12]. The unintegrated viral DNA can take several forms, linear and  
512 the 1-LTR- and 2-LTR circles [7, 45], with much of the transcription thought to emanate from 1-  
513 LTR circles. In macrophages it has been shown that HIV-1 PIC cDNA can persist and be  
514 actively transcribed for months. However, it is generally agreed that PIC HIV-1 transcription in  
515 macrophages does not routinely produce infectious virus [12]. The use of single cell techniques  
516 has enabled us to quantify both the numbers of cells expressing given HIV transcripts in a mixed  
517 culture, and also the relative transcript loads of each HIV-1 gene in each infected cell [13-15]. It  
518 also allows us to put the cells containing viral transcripts into the context of their background  
519 transcriptomes. This provides the opportunity to compare cells producing late viral proteins to  
520 those producing transcript but not late viral proteins in order to better understand the cellular  
521 metabolic background necessary for viral production.

522 As reported by Marsh and Wu and colleagues [10], “transcription in the absence of  
523 integration is selective and skewed towards certain viral early genes such as nef and tat, with  
524 highly diminished rev and vif”. In general, our single cell analysis agrees with this conclusion,  
525 but provides a more nuanced picture. In our data, compared to cells in the same co-culture  
526 transcribing from provirus, many more PIC cells are producing tat, nef, gag-pol, and env

527 transcripts. However, the level of a given transcript per cell is not detectably different from  
528 Provirus cluster cells. In contrast, the prevalence of PIC cells that make rev, gag, and accessory  
529 gene transcripts constitute a smaller fraction of the PIC cells. Nevertheless, those few PIC cells  
530 that are producing rev, gag and accessory gene transcripts are doing so at levels equivalent to  
531 Provirus cells. In Provirus cells, the numbers of cells detectably making spliced or un-spliced  
532 transcripts are even. The levels of these transcripts is on average similar to the levels of  
533 transcripts produced by PIC cells. Thus, it is difficult to make the generalization that high overall  
534 transcript levels are a trigger for virus production.

535 Only the Provirus cells appear to be making p24, mCherry or Vpu protein, while Nef  
536 protein can be detected in protein form both Provirus and PIC cluster cells. Using Western blot  
537 analysis, we only see the production of p24, mCherry and Vpu in protein samples containing  
538 Provirus cells. It is not clear whether our failure to detect other HIV proteins in preparations  
539 enriched for PIC /Bystander cells is because very few cells in this cluster were making the  
540 proteins (and thus below our level of detection), or if there was some restriction mechanism  
541 preventing their production, or both. Single cell protein analysis technology might be able to  
542 address this point. The robust detection of mCherry, p24 and Vpu in protein samples enriched for  
543 Provirus cells confirms that late protein synthesis is an attribute of the Provirus cells. Nef protein  
544 is not made in cells infected with the DHIV3-mCherry, due to the mCherry sequence replacing  
545 the 5' portion of the nef gene, and we could not detect any in proteins from DHIV3 infected  
546 cultures.

547 Integrated provirus transcription is required for virus production [12]. It is regulated by  
548 promoter elements in the HIV LTR. Thierry and co-workers have recently shown that PIC cDNA  
549 and provirus are differentially responsive to NFkB promotion [45]. As others have found, they

550 report provirus transcription is enhanced by NFkB and AP1 binding. However, they find PIC  
551 cDNA transcription to be inhibited by NFkB activation. Our data adds an additional layer of  
552 complexity to this picture. We can show that early and late gene transcripts are detectable in  
553 Provirus cells, at levels equal to the respective transcripts detected in PIC cells. We also show  
554 that, in cells that have transitioned from PIC/Bystander to Provirus, the background  
555 transcriptome reflects an overall down regulation of NFkB and AP1 transcripts, and upregulation  
556 by E2F family promoted transcripts. E2F is not a promoter in the HIV-1 LTR, and we do not  
557 suggest that E2F regulation of provirus transcription is the key to the PIC to Provirus transition.  
558 However, we do propose that an E2F family promoter-dominated transcriptome is required for  
559 virus production.

560 This proposition appears counter to literature, in which E2F is thought to suppress viral  
561 transcription [25, 46]. Nevertheless, the pathways upregulated by E2F are those consistent with  
562 what one would intuitively anticipate a being required for viral production. The singling out of  
563 E2F promoter family proteins in virus production is not new and is consistent with literature  
564 citing a role for Rb phosphorylation and E2F activation in HIV-1 linked tumorigenesis [47].  
565 Indeed, we show that levels of phospho-Rb are higher in Provirus cells. Consistent with this was  
566 our observation in our THP-1 system, and in primary cultures of macrophage and T-cells, that  
567 cells that have already committed to making virus are more likely to make virus upon second  
568 infection. If this interpretation is correct, then study of genome-wide interactions that accompany  
569 provirus integration and amplify E2F signaling might be key to understanding the switch in  
570 transcriptome necessary for viral protein production. In any case, this new understanding makes  
571 it possible to discriminate between active and latent HIV-1 infected cells from patient tissues,  
572 using single cell analysis and description of the cell's background transcriptomes.

573

574 **Materials and methods**

575 **Reagents**

576 THP-1 cells, a monocytic cell line, were obtained from ATCC (Cat#TIB-202). HyClone™  
577 RPMI 1640, kanamycin sulfate, Corning™ Accutase™ detachment solution and phorbol 12-  
578 myristate 13-acetate (PMA) were obtained from Fisher Scientific (cat. # SH30011.03, BP906-5,  
579 MT25058CI, and BP685-1, respectively). Fetal bovine serum was purchased from Atlanta  
580 Biologicals (cat. # S11150). BD Horizon™ Fixable Viability Stain 450 was obtained from BD  
581 Biosciences (cat. # 562241).

582

583 **Cell culture**

584 THP-1 cells were grown in RPMI 1640 supplemented with 10% FBS and KAN (50  
585 µg/mL) at 37°C, 5% CO<sub>2</sub>.

586

587 **Generation of DHIV3-mCherry**

588 DHIV3-mCherry virus was generated by calcium phosphate transfection (25). In brief,  
589 HEK293FT cells were grown to 80% confluence. DHIV3-mCherry plasmid and VSVg plasmid  
590 were mixed with calcium chloride (2.5 M) and HEPES buffered saline solution. The calcium  
591 phosphate-DNA suspension was added dropwise to the cells. Chloroquine (100 mM) was  
592 subsequently added. HEK293FT cells were incubated overnight with solution. The medium was  
593 replaced with fresh DMEM and incubated for an additional 48 hours. Supernatant was collected  
594 and filtered (0.45 µm). Optimal viral titers were determined by titrating the virus in THP-1 cells.  
595 A titer volume of 100 µL in 500 µL total (~5 x 10<sup>6</sup> TU/mL in THP-1s) was chosen due to its

596 high DHIV infection and minimal effects on viability. Titer volume (up to 400  $\mu$ L of 500  $\mu$ L)  
597 was increased as infectivity fell off in stocks over time.

598

## 599 **DHIV3-mCherry and THP-1 co-culture**

600 For the THP-1s, cells were preincubated overnight in PMA (20 ng/mL) at 500,000  
601 cells/well in order to generate differentiated macrophages. Medium was replaced for THP-1 cells  
602 1 hour prior to the addition of DHIV3-mCherry. Cells were incubated at 37°C for 24 hours  
603 followed by preparation for analysis by flow cytometry and cell imaging.

604

## 605 **Flow cytometry**

606 Adherent THP-1 cells were incubated with Accutase™ for 15 minutes at 37°C. THP-1  
607 cells were transferred to 5-mL tubes and washed with PBS. Cells were resuspended in BD  
608 Horizon™ Fixable Viability Stain 450 (0.25  $\mu$ g/mL) and incubated at 4°C for 30 minutes. Cells  
609 were then fixed in 2% formaldehyde for 30 minutes at 4°C. Cells were analyzed using a FACS  
610 Canto. Percent infection by DHIV was quantified as a subset of the live population  
611 (FSC/V450/50-). Gates for infection were set according to the uninfected “mock” THP-1 cell  
612 controls. Population analysis was then done using FlowJoTM v10.7.

613

## 614 **10X Genomics library construction and sequencing**

615 Two biological replicate cultures of HIV(+) THP-1 cells (HIVreplicate1 and  
616 HIVreplicate2) and HIV(-) THP-1 cells (here after referred to as Control, or WT1 and WT2 for  
617 wild type ) were processed through the 10X Genomics Chromium Single Cell Controller with

618 Single Cell Gene Expression 3' Solution (v2 chemistry). Sequencing was done on an Illumina  
619 HiSeq 2500 instrument.

620 Cell suspensions were partitioned into an emulsion of nanoliter-sized droplets using a  
621 10X Genomics Chromium Single Cell Controller and RNA sequencing libraries were  
622 constructed using the Chromium Single Cell 3' Reagent Kit v2 (10X Genomics Cat#PN-  
623 120237). Briefly, droplets contained individual cells, reverse transcription reagents and a gel  
624 bead loaded with poly(dT) primers that include a 16 base cell barcode and a 10 base unique  
625 molecular index. Lysis of the cells and gel bead enables priming and reverse transcription of  
626 poly-A RNA to generate barcoded cDNA molecules. Libraries were constructed by End Repair,  
627 A-Tailing, Adapter Ligation and PCR amplification of the cDNA molecules. Purified cDNA  
628 libraries were qualified on an Agilent Technologies 2200 TapeStation using a D1000 ScreenTape  
629 assay (Agilent Cat#5067-5582 and Cat#5067-5583). The molarity of adapter-modified molecules  
630 was defined by quantitative PCR using the Kapa Biosystems Kapa Library Quant Kit (Kapa  
631 Biosystems Cat#KK4824).

632 HiSeq 125 Cycle Paired-End Sequencing v4: Sequencing libraries (25 pM) were  
633 chemically denatured and applied to an Illumina HiSeq v4 paired end flow cell using an Illumina  
634 cBot. Hybridized molecules were clonally amplified and annealed to sequencing primers with  
635 reagents from an Illumina HiSeq PE Cluster Kit v4-cBot (PE-401-4001). Following transfer of  
636 the flowcell to an Illumina HiSeq 2500 instrument (HCS v2.2.38 and RTA v1.18.61), either a  
637 26x100 cycle or 125 cycle paired-end sequence run was performed using HiSeq SBS Kit v4  
638 sequencing reagents (FC-401-4003). Basic html (notebook) files describing coding and QC data  
639 are provided in Appendix IV.

640

## 641 Data analysis for UMAP (Fig. 2) of HIVrepeat1

642 Raw FASTQ files from 10x Genomics from were processed by 10x Genomics' Cell  
643 Ranger software. Each library was processed with 'cellranger count' pipeline with a common  
644 genomic reference made up of human and HIV genomes as well as mCherry. The human  
645 genomic reference was GRCh38 with gene annotation from Ensembl release 91, where only  
646 features with gene\_biotype:protein\_coding were kept. The HIV genome and annotation was  
647 acquired from NCBI genome (RefSeq ID NC\_001802.1). No warnings were issued by 10x  
648 Genomics regarding sequencing, alignment, or cell-based QC metrics; however, the samples  
649 could have been sequenced deeper as reflected in the sequencing saturation statistics.

650 In attempt to recover those (perhaps lower quality) GEM partitions, the raw gene-barcode  
651 matrices from 'cellranger count' (located in 'outs/raw\_gene\_bc\_matrices') was processed with  
652 the EmptyDrops algorithm (R package DropletUtils v1.2.2) to discriminate cells from  
653 background GEM partitions at a false discovery rate (FDR) of 0.001% [48]. GEM partitions with  
654 2000 UMI counts or less were considered to be devoid of viable cells, while those with at least  
655 10,000 UMI counts were automatically considered to be cells.

656 For each technical replicate, additional quality control measures were taken to filter out  
657 low-quality cells. Cell-based QC metrics were calculated with R package scater (v1.16.2) using  
658 the perCellQCMetrics function [49]. Cells with extremely low UMI counts, extremely low gene  
659 counts or extremely high percentage of expression attributed to mitochondrial genes were also  
660 flagged as low quality. Extremeness in any of these three measures was determined by 3 median  
661 absolute deviations from the median with the scater isOutlier function. These cells suspected of  
662 being low quality were removed from downstream analysis with one exception in the HIV  
663 replicates; if cells exhibited above median HIV gene expression, they were not discarded as these

664 were thought to hold potential value as examples of cells in which viral replication suppressed  
665 other gene expression (not observed). Further analysis of the HIV biological replicates showed  
666 there were remaining low quality cells as marked by unusual mitochondrial gene expression or  
667 low library size, which were removed to improve the signal to noise ratio. Specifically, HIV-  
668 infected cells with mitochondrial expression of 7.23% and above were removed as well as cells  
669 that have less than 3242 UMI counts. To ensure no cell type was discarded due to filtering,  
670 average gene expression was compared gene-wise between discarded and kept cells in scatter  
671 plots. There were no genes of interest that exhibited markedly different average gene expression  
672 between the discarded and kept cells, suggesting the filtering did not remove interesting sub-  
673 populations. After filtering low quality cells, technical replicates were combined into biological  
674 replicates HIVreplicate1, HIVreplicate2, wt1 and wt2.

675 For each biological replicate, cells were normalized [48] and scored for a number of  
676 important attributes. Each cell from HIV biological replicates was assigned a HIV activity score  
677 with Seurat's (v3.2.2) AddModuleScore function [50], where a high score indicates HIV gene  
678 expression was stronger in the cell relative to randomly selected genes of similar expression  
679 strength in the biological replicate. Cell cycle phases and scores were assigned cells with the  
680 cyclone method [51]; Cells were scored against a simulated doublet population of cells with  
681 scran's (v1.10.2) doubletCells function; those cells with extremely high doublet scores (5 median  
682 absolute deviations) were removed and remaining cells were re-normalized again with scran's  
683 quickCluster and scater's normalize methodology [48] for differences in sequencing depth  
684 between libraries.

685

686 **Data analysis for t-Sne insert (Fig. S-3)**

687 10x Cell Ranger raw sequencing data was processed into UMI counts with ; using the  
688 ‘mkfastq’, ‘count’, bioinformatic modules. Cell Ranger de-multiplexed cDNA libraries into  
689 FASTQ files with Illumina’s bcl2fastq and aligned reads to a hybrid genomic reference  
690 composed of human (Ensemble GRCh38), HIV (NCBI ID: NC\_001802.1), and mCherry  
691 genomic references with STAR aligner [48, 54]. CellRanger filtered cell barcodes and unique  
692 molecular identifiers (UMIs) in estimation of gene-cell UMI counts using only reads that  
693 mapped uniquely within the transcriptome. We specified an ‘expected cell number’ of 3000 per  
694 library based on reported cell recovery rates.

695 The QC metrics reported by Cell Ranger indicated that our library construction was a  
696 success; the libraries averaged 97.9% valid cell barcodes, 60.6% of reads mapping to the  
697 transcriptome, and reported a median of 2402 genes detected per cell (mean of 15262.2 genes per  
698 library). Only in HIV-infected samples did reads map to the HIV genome. Cell Ranger also  
699 evaluated dimension reduction, clustering, and differential gene expression analysis under  
700 default parameters. For further details of the Cell Ranger data processing and analysis pipeline,  
701 see <https://support.10xgenomics.com/single-cell-gene-expression/software/pipelines/latest/algorithms/overview>. Some interactive data analysis, i.e. t-Sne visualization, was conducted with  
703 10x Genomics’ Cell Loupe Browser.

704 Due to cost, this study was limited by low sequencing depth (mean of 41,433 reads per  
705 cell per library). This limitation was mitigated by removing genes with low sequencing coverage;  
706 specifically, a gene was filtered out if it did not have 1 percent of cells reporting at least 3 UMI;  
707 cells were filtered out if they did not have at least 200 genes with a UMI count. Cells with  
708 exceedingly high (top 2%) ribosomal and/or mitochondrial content were filtered out. To reduce

709 mutliplets contaminating analysis, cells with the top 2.3% total UMI were removed (see 10x  
710 Genomics benchmarks).

711

## 712 **Dimension reduction, clustering, and differential expression**

713 Highly variable genes were identified with scran's trendVar and decomposeVar.  
714 Specifically, loess smoothing was applied to the gene variance (dependent variable) and the  
715 mean gene expression (independent variable) after having corrected for the % mitochondrial  
716 expression and cell cycle effects on the cells. Genes with average expression below the first  
717 quartile were filtered out of consideration. Gene variance was decomposed into biological and  
718 technical components, where genes with variance above the mean trend (loess fit) were assumed  
719 to possess biological variation [52] . This process was repeated for the HIV biological replicates  
720 separately followed by scran's combineVar function applied to the combined HIV replicates to  
721 identify genes estimated to have positive biological variation and controlled with a false  
722 discovery rate of 0.05.

723

## 724 **Gene set enrichment analysis**

725 A gene set enrichment analysis (GSEA) was conducted with R package fgsea (v1.8.0)  
726 [53]. The log2 fold change vector of strong-HIV vs. weak-HIV was evaluated for enrichment  
727 against 3 different collections of MSigDB gene sets; namely, Hallmark, REACTOME, and  
728 Transcription Factor Targets. The Benjamini-Hochberg false discovery rate (FDR) was  
729 controlled at 10%.

730 A DGE and GSEA analysis was conducted on HIV biological replicates in comparison to  
731 Control biological replicates. The wild type replicate expression data was subject to (nearly)

732 identical quality control and identical pre-processing steps. As described above, quality control  
733 on HIV cells was subject to a greater degree of scrutiny.

734 To mitigate biases due to potential batch-specific variation, DGE and GSEA analyses  
735 contrasting Provirus -HIV and PIC/Bystander-HIV populations to Control cells leveraged  
736 consensus between various pairwise contrasts. For instance in the HIV- Provirus vs. Control  
737 (WT) contrast, t-tests were evaluated in 4 distinct contrasts: (i) HIVreplicate1-Active vs. WT1;  
738 (ii) HIVreplicate1-Act Proivirus ive vs. WT2; (iii) HIVreplicate2- Provirus vs. WT1; (iv) and  
739 HIVreplicate2- Provirus vs. WT2. The scran function combine Markers performed a meta-  
740 analysis across the 4 contrasts with the Simes method. The Simes meta-analysis tested whether  
741 any of the 4 contrasts manifest either a change a gene-wise expression for DGE analysis; that is  
742 to say the meta-analysis p-value encodes the evidence against the null hypothesis, which assumes  
743 the gene is not changed in any of the 4 comparisons. Similarly in the GSEA analysis, the log2  
744 fold change statistics from the 4 comparisons were tested for enrichment of the 3 previoulsy  
745 mentioned MSigDB gene set collections (Hallmark, REACTOME, Transcript Factor Targets).  
746 The results were also merged with the Simes meta-analysis. This same strategy for the HIV-  
747 Provirus vs. Control contrast was repeated in the HIV- PIC/Bystander vs. Control comparison.

748 There was interest in modeling the progression of infection from wild type to  
749 PIC/Bystander-HIV to Proivirus -HIV clusters. Specifically, there was interest in identifying  
750 what genes exhibit a variable expression profile or non-constant trend when ordered from  
751 Control to PIC/Bystander-HIV to Proivirus -HIV. Macnair and Claassen [41] developed a  
752 supervised psuedotime R package, called psupertime, that is tailored to this express purpose. In  
753 particular, a penalized logistic ordinal regression model was fit to the combined HIV and Control  
754 data. The input data was a subset of highly variable genes identified in the same manner as

755 described above, but including Control data as well. The gene expression data had been  
756 normalized, log2 transformed and followed by linear correction of effects due to percent  
757 mitochondrial expression and cell cycle phase. The model was able to clearly order cells that  
758 reflects the expected order of Control then PIC/Bystander-HIV then Provirus -HIV. The  
759 psupertime method also reports a small set of genes which strongly associate with the expected  
760 progression, which is based on the magnitude of the penalized coefficients in the logistic ordinal  
761 regression.

762

## 763 **Western protocol**

764 The stored THP-1 (uninfected, HIV-1 infected, HIV-1 infected with MK-2048 treatment)  
765 cells were pelleted by centrifugation for 5 min at 10,000 x g. Precipitates were then washed twice  
766 in cold PBS. Afterwards, cells were lysed using 50 mM Tris pH 7.4, 100 mM NaCl and 2 mM  
767 EDTA, complete® protease inhibitor, 2 mM NaF, 2 mM sodium orthovanadate and 10% SDS.  
768 Protein concentrations were determined using bovine serum albumin standard and Coomassie  
769 Plus Protein Reagent from Pierce Biotechnology (Rockford, IL). 10 µg of the whole cell lysate  
770 was separated using NuPAGE 4–12% Bis-Tris gradient gels (Invitrogen Life Sciences, Carlsbad,  
771 CA) and transferred to a PVDF membrane (Millipore, Billerica, MA). These were then blocked  
772 in 2% BSA in TBST for 20 min at room temperature, incubated overnight with primary  
773 antibodies at 4°C in blocking buffer solution and with the secondary antibody for 45 minutes at  
774 room temperature. Protein was detected using chemiluminescence and blots were visualized  
775 using a Protein Simple FluoroChem M system.

776

## 777 **Antibodies and fluors**

778 P24 and Gag protein precursor production was detected with monoclonal mouse IgG-  
779 AG3.0, (NIH AIDS Res. Reagent Prog., Germantown, MD; Cat. # 4121), 1:500, and using an  
780 AlexaFlour 633nm or 700nm goat anti-mouse IgG (H+L) secondary anti-body (Fisher Scientific,  
781 Pittsburgh, PA) for flow cytometry. Anti-HIV-1 NL4-3 VPU rabbit polyclonal, also from the  
782 NIH AIDS Res. Reagent Prog. (Cat. # 969), was used at 1:20 dilution for Western Blots. Goat  
783 anti-mCherry, OriGene (TR150126), was used at a 1:5000 dilution for Western Blots.  
784 ThermoFisher Scientific Anti-Rb (LF-MA0173, 32C8) was used at a 1:1000 dilution. Invitrogen  
785 antibodies: anti-pRB (T821, 710314) was used at a 1:500 dilution, anti-pIkB (S32, 701271) was  
786 used at a 1:500 dilution, and anti-B-actin (PA5-85291) was used at a 1:5000 dilution.

787 Other antibodies obtained from the NIH AIDS Res. Reagent Prog., Germantown, MD  
788 include: anti-HIV-1 IIIB gp120 Polyclonal (Cat. # 57), anti-HIV-1 RF gp160 Polyclonal (HT7)  
789 (Cat. # 189), anti-HIV-1 Tat Polyclonal (Cat. # 705), anti-Nef Monoclonal (EH1) (Cat. # 3689),  
790 anti-HIV-1 Nef Polyclonal (Cat. # 2949), anti-HIV-1 Vpr 1-50 aa Polyclonal (Cat. # 11836);  
791 anti-HIV-1 HXB2 Vif Polyclonal (Cat. #12256), anti-HIV-1 HXB2 IN Polyclonal (Antigen 2)  
792 (Cat. # 12877), anti-HIV-1 RT Monoclonal (MAb 21) (Cat. # 3483), anti-HIV-1 HXB2 RT  
793 Polyclonal (Antigen 2) (Cat. # 12881), anti-HIV-1 Protease Polyclonal (Cat. # 13564), anti-HIV-  
794 1 Rev Monoclonal (1G7) (Cat. # 7376), which were tested at various dilutions.

795 HRP-conjugated secondary antibodies were used to develop Western Blots, anti-rabbit  
796 A0545 (1:5000) and anti-mouse A9044 (1:5000) from Sigma Chem. Co., and anti-goat 401515  
797 (1:10,000) from CalBiochem. Propidium Iodide was obtained from Molecular Probes, Eugene,  
798 OR. DAPI blue was obtained from Acros Organics, Geel, Belgium.

799

800 **Real-time PCR methods**

801 For detection of integrated proviral DNA, DNA from control, DHIV3-mCherry infected,  
802 DHIV3-mCherry infected plus integrase inhibitor treated THP1 cells was purified using Qiagen  
803 Blood and Tissue DNeasy kits. The PCR evaluation of integrated HIV was performed using the  
804 primers and PCR conditions described from Chun et al. [34]. Briefly, the primers were:

805 Alu->LTR 5, 5'-TCCCAGCTACTCGGGAGGCTGAGG-3'

806 LTR->Alu 3', 5'-AGGCAAGCTTATTGAGGCTTAAGC-3'

807 Nested secondary PCR primers (generating a 352 bp amplicon):

808 5', 5'-CACACACAAGGCTACTTCCCT-3'

809 3', 5'-GCCACTCCCCIGTCCCGCCC-3'

810 We used Ranger polymerase and buffer conditions (Meridian Bioscience, Thomas  
811 Scientific, Swedesboro, NJ) for the long-range PCR with Alu-LTR primer sets, and BioTaq  
812 polymerase and buffer conditions (Meridian Bioscience, Thomas Scientific, Swedesboro, NJ) for  
813 the nested PCR. The integrated HIV PCR was performed on an MJ PTC-200 with an MJE 2X48  
814 and a Chromo-4 alpha unit for the long-range and nested PCR, respectively. The nested PCR was  
815 performed as a real-time assay using SYBR Green I to detect the amplicon progression curves  
816 and evaluate the melting curve.

817 For detection of total HIV DNA, in order to determine if comparable total HIV DNA was  
818 present in the samples the same samples described above, we utilized the 5' nested primer (5'-  
819 CACACACAAGGCTACTTCCCT-3') along with the LTR->Alu 3' primer (5'-  
820 AGGCAAGCTTATTGAGGCTTAAGC-3') using PCR conditions similar to the nested PCR

821 described above but with a 30 sec extension time for the 484 bp amplicon. This PCR was  
822 performed on a Roche LightCycler 480 instrument using BioTaq polymerase and buffer  
823 conditions with SYBR Green I detection of the 484 bp amplicon.

824 Detection of circular 2-LTR DHIV3-mCherry PIC DNA was performed as described in  
825 Brussel and Sonigo [33]. Briefly, the primers used were:

826 HIV F, 5' GTGCCGTCTGTTGTGACT 3'

827 HIV R1, 5' ACTGGTACTAGCTTGTAGCACCATCCA 3'.

828 Initially we performed the PCR conditions used by Brussel and Sonigo [33], using  
829 BioTaq polymerase and buffer conditions with a 25 sec extension time and SYBR Green I  
830 detection, but we were unable to detect any amplicon 2LTR circle PIC product. Following the  
831 detection of the total PIC HIV data, we hypothesized that the 2-LTR content in these samples  
832 could be considerably lower at this 24 hr time point. Therefore, we reran the PCR again a second  
833 time and were able to detect an expected 231 bp amplicon. To verify this approach, we  
834 synthesized new PCR primers:

835 RU5 forward: 5'-GCTTAAGCCTCAATAAAGCTTGCCT-3' (this is the compliment of the  
836 LTR->Alu primer described above from Chun et al. [34]).

837 U3 reverse: 5'-ACAAGCTGGTGTCTCTCCT-3'.

838 This primer set also did not generate the 2-LTR circular PIC amplicon within 50 PCR cycles but  
839 was designed to encompass the amplicon generated by Brussel and Sonigo primers above.  
840 Therefore, we used the HIV F and R1 primers as a nested set and ran a 1:20 dilution of this

841 amplification for another 50 PCR cycles to obtain the expected 231 bp amplicon. This  
842 demonstrates the 2-LTR circular form of PIC cDNA was present at low levels in all the  
843 conditions where DHIV3-mCherry was used, but not in the control samples.

844

## 845 **Sequential DHIV3-mCherry, DHIV3-GFP infection**

846 PBMCs from healthy human donors were isolated using lymphocyte separation medium  
847 (Biocoll separating solution; Biochrom) or lymphoprep (Stemcell). CD4+ T cells were  
848 negatively isolated using the RosetteSep<sup>TM</sup>Human CD4+ T Cell Enrichment Cocktail (Stem Cell  
849 Technologies) or the EasySep<sup>TM</sup> Human Naïve CD4+ T Cell Isolation Kit (Stem Cell  
850 Technologies) according to the manufacturer's instructions. Primary rCD4s were cultured to a  
851 density of 5 x 10<sup>6</sup>/mL in RPMI-1640 medium containing 10% FCS, glutamine (2 mM),  
852 streptomycin (100 mg/mL), penicillin (100 U/mL) and interleukin 2 (IL-2) (10 ng/mL).

853 Monocyte-derived macrophages (MDMs) were obtained by stimulation of PBMC  
854 cultures with 15 ng/mL recombinant human M-CSF (R&D systems) and 10% human AB serum  
855 (Sigma Aldrich) in DMEM supplemented with glutamine (2 mM), streptomycin (100 mg/mL)  
856 and penicillin (100 U/mL) for 6 days.

857

## 858 **Statistical analysis**

859 The pairwise TTests function from Scran was used to determine statistically significant  
860 differential expression of genes between groups. This was performed for all comparison sets.  
861 Only those genes which were significantly different were included in Hallmark, REACTOME,  
862 pseudotime, psupertime and TFT analyses. Other default statistical standards were adopted from  
863 the various software recommendations during data analyses unless otherwise specified.

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1115

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1117

1118 **Figure Captions**

1119

1120 **Figure 1. Flow cytometry analysis of DHIV3-mCherry infected THP-1 cells. Panel A)**

1121 mock infection of PMA activated THP-1 cells. Panel **B**) PMA activated THP-1 cells infected

1122 with DHIV3-mCherry. Absicca mCherry (Texas Red) emission. Ordinate, GFP (FITC) emission.

1123 MCherry positive cells equal approximately 8.5% of total viable cell population.

1124

1125 **Figure 2. UMAP projection of scRNA seq data from experiment HIVreplicate1.**

1126 Greater than 14,000 different cellular genes were detected in this analysis, including the 9 viral

1127 genes and mCherry message originating from DHIV3-mCherry. A semi supervised two cluster

1128 model was adopted, the smaller “Provirus cluster” (cluster A) was 8.1% of the total cell

1129 population, approximately equivalent to the percentage of mCherry positive cells from Fig. 1.

1130 The two semi supervised clusters are circled in red. PIC/Bystander cluster is indicated as cluster

1131 B. The HIV activity scale presents the Seurat module score that is described in methods. Input

1132 data in this analysis included 33,819 PCA entries. The number of cells with detected genes

1133 following Seurat QC in the PIC/Bystander cluster was 12% of the total population. Bar codes of

1134 same cells tracked to the Provirus clusters, regardless of whether the clusters were generated

1135 using UMAP or Seurat-tSne tools (Fig. S-3).

1136

1137 **Figure 3. UMAP Feature plot of experiment HIVreplicate1. Panel A)** Distribution of

1138 HIV-1 transcript positive cells shown in Fig. 2. Provirus cluster circled in red, PIC/Bystander

1139 cluster in blue square. Panels **B**), **C**) and **D**), respectively, show no influence of cell cycle,

1140 number of genes detected per cell, or percent mitochondrial transcripts (positively correlated

1141 with cell stress) on the distribution of PIC cells (HIV-1 transcript containing cells) throughout  
1142 the PIC/Bystander cluster.

1143

1144 **Figure 4. Unsupervised clustering of UMAP shown in Fig 2. Panel A)** shows  
1145 unsupervised clustering obtained at K equals 10. Panel **B**) Violin plot of HIV-1 transcripts/cell in  
1146 the 10 clusters identified at K10 (Scran's buildSSNGraph using the PCA as input). PIC cells with  
1147 detectable HIV-1 transcripts, were distributed throughout clusters 1-5, 7 and 9-10. Clusters 6 and  
1148 8 contained 372 of the 381 cells included in the semi supervised Provirus cluster (circled in red).  
1149 Stipulation of lower K values means that during analysis any one given cell is clustered with a  
1150 smaller number of cells with similar transcriptomes.

1151

1152 **Figure 5. Flow cytometry analysis of DHIV3-mCherry infected THP-1 cells using p24**  
1153 **antibody.** Panel **A**) mock infection. Panel **B**) mCherry expression was positively correlated with  
1154 p24/Gag antigen detection by flow cytometry. Abscissa mCherry (Texas Red) emission.  
1155 Ordinate, GFP (FITC) emission. mCherry/p24 positive cells equal approximately 18% of total  
1156 viable cell population in this experiment.

1157

1158 **Figure 6. Integrase inhibitor treatment selectively reduces mCherry positive cells.**  
1159 Panel **A**) flow cytometry analysis of DHIV3-mCherry infected THP-1 cells, versus viability  
1160 stain. Abscissa shows viability stain intensity, ordinate shows mCherry intensity. Infected  
1161 (Active), mCherry-producing cells account for approximately 12 % of the cell population. Panel  
1162 **B**) Same as A except with the addition of 25nM MK-2048 integrase inhibitor at time of

1163 infection. Integrase inhibitor effectively reduces number of mCherry producing cells, without  
1164 decreasing cell viability.

1165

1166 **Figure 7. Effect of integrase inhibitor on mCherry, p24, Gag and Vpu protein**  
1167 **production in cultures containing DHIV3-mCherry infected cells.** MW, molecular weight  
1168 markers. Lane 1, Control cell protein; Lane 2, protein from DHIV3 infected culture; Lane 3,  
1169 protein from DHIV3 infected cultures treated with integrase inhibitor (25nM MK-2048) as  
1170 shown above in Figure 9. **A)** Lane 2, p24 and Gag precursor proteins visualized with p24  
1171 antibody used above in Fig. 6, and HRP linked anti-mouse secondary antibody. The p24 band in  
1172 lane 3 is residual from infection as reported in the literature [10]. The presence of precursor  
1173 proteins in lane 2 shows Provirus synthesis in cultures containing Provirus cells. **B)** Antibody  
1174 used was goat anti-mCherry, developed with HRP linked anti-goat secondary. mCherry protein  
1175 clearly visible in preparations containing Provirus cell protein. **C)** Lane 2, Vpu detected with  
1176 rabbit antibody, visualized using HRP linked anti-rabbit secondary. The resolution of image  
1177 slightly compromised due to small size of Vpu protein. **D)** Lanes 1 and 2, Control cell protein at  
1178 24 and 48 hrs respectively; lanes 3 and 4, protein from DHIV3 infected culture at 24 and 48 hrs  
1179 respectively; lanes 5 and 6, protein from DHIV3 infected cultures treated with integrase inhibitor  
1180 (as above) at 24 and 48 hrs respectively. At 24 hrs post infection, we only found both p24 and  
1181 precursor Gag proteins in the protein samples from DHIV3 infected cells in the absence of  
1182 integrase inhibitor. At 48 hrs post-infection, in the absence of integrase inhibitor, the amounts of  
1183 detectable p24 and Gag proteins were dramatically increased from levels at 24 hrs post infection.  
1184 As seen initially (Panel A), some p24 protein was detectable in integrase inhibitor treated  
1185 cultures at 24 hrs post infection, however Gag is not detectable at this time. At 48 hrs post

1186 infection in the integrase inhibitor treated cultures, some Gag protein does becomes detectable,  
1187 reflecting production in cells that escaped complete integrase inhibition. This is in agreement  
1188 with our flow cytometry analysis that showed suppressed, but still detectable numbers of  
1189 mCherry positive cells in the integrase inhibitor treated cultures. The Gag precursor proteins only  
1190 appear in the integrase inhibitor treated culture proteins 48 hrs after treatment. All antibodies,  
1191 sources and dilutions are provided in Methods.

1192

1193 **Figure 8. UMAP analysis of integrase-inhibitor treated DHIV3-mCherry infected**  
1194 **THP-1 cells.** Experiment performed as shown in Figure 6 B, with 25nM MK-2048 added at the  
1195 time of DHIV3 addition. Data were analyzed identically to data shown in Figure 2. Panel **A**)  
1196 Feature plot showing distribution of cells containing HIV-1 transcript, generated as described.  
1197 Panel **B**) K10 unsupervised clustering generated 7 clusters (Scran's buildSSNGraph using the  
1198 PCA as input). HIV-1 transcripts were distributed equally throughout all of them. No cluster  
1199 corresponding to the "Provirus" cluster detected in Figure 4 was detected, regardless of K value  
1200 used (see Fig. S-6). These data agree with the concept that integrase inhibitors selectively target  
1201 and reduce the number of Provirus cluster cells.

1202

1203 **Figure 9. Distribution of gene transcripts exhibiting high levels of differential**  
1204 **expression between Provirus and PIC/Bystander clusters.** Feature plot showing distribution  
1205 of cells from Figure 2, containing transcripts of 10 of the 20 most highly differentially expressed  
1206 transcripts in Provirus versus PIC/Bystander GSEA data sets. APOE, IFI6 and EIF5 were also  
1207 included because they were highly expressed in the PIC/Bystander cluster. As above, these  
1208 UMAP projections were made with Seurat's FeaturePlot function. They are colored by

1209 expression of individual genes (normalized log2 values). Highly expressed genes in Proivirus  
1210 cluster cells include PHIP (Pleckstrin Homology Domain Interacting Protein), CDKN2C (Cyclin  
1211 Dependent Kinase Inhibitor 2C), COMMD3 (COMM Domain Containing Protein 3), REEP3  
1212 (Receptor Accessory Protein 3) and PCLAF (PCNA Clamp Associated Factor). Highly  
1213 expressed transcripts detected in the PIC/Bystander cell transcriptome include FABP5 (Fatty  
1214 Acid Binding Protein 5), CTSL (Cathepsin L), FTH1 (Ferritin Heavy Chain 1), MMP9 (Matrix  
1215 Metallopeptidase 9), LIMS1 (LIM Zinc Finger Domain Containing 1), APOE (Apolipoprotein  
1216 E), IFI6 (Interferon Alpha Inducible Protein 6) and EIF5 (Eukaryotic Translation Initiation  
1217 Factor 5).

1218

1219 **Figure 10. The Distribution of HIV-1 transcripts throughout Proivirus and**  
1220 **PIC/Bystander clusters.** Panel **A**) Feature plot showing distribution of cells from UMAP in  
1221 Figure 2 that contain detectable DHIV3-mCherry transcripts. As described above, these UMAP  
1222 projections were made with Seurat's FeaturePlot function. They are colored by expression of  
1223 individual genes (UMAP projection colored by walktrap, normalized log2 values). ASP is a  
1224 negative control, bacterial gene transcript sequence. **B**) Violin plots of DHIV3-mCherry  
1225 transcript/cell in cells from the Proivirus and PIC clusters showing transcript level and cell  
1226 number. The provirus cluster contained transcriptomes of 371 cells, the number of PIC cells in  
1227 the PIC/Bystander cluster was 569 cells, thus the Proivirus/PIC cell number ratio was 0.65. The  
1228 plots were made with Seurat's VlnPot function. They show normalized log2 transcript levels.  
1229 Two patterns of transcript distribution are evident. The first pattern is seen with gag-pol, tat, env  
1230 and nef, in which relatively high numbers of cells in the PIC/Bystander cluster express the  
1231 transcripts, with the transcripts being detected in fewer numbers of Proivirus cluster cells. The

1232 second pattern is seen with gag, vif, vpr, rev, vpu, and mCherry, in which relatively equal  
1233 absolute numbers of cells in the Proivirus and PIC/Bystander clusters are detected with the  
1234 transcript sequences, remembering that there are more PIC cells than Proivirus cells. The relative  
1235 transcript loads per PIC cell versus the Proivirus cells overlap. Negative control sequence (asp)  
1236 shows no distribution.

1237

1238 **Figure 11. Differential gene expression comparison of Proivirus and PIC/Bystander**  
1239 **cluster gene transcripts versus 2 independent biological repeat control (wt) experiments.** In  
1240 every comparison, a significant positive correlation was obtained from the common detected  
1241 differentially expressed genes of Proivirus or PIC/Bystander clusters in the two biological repeats  
1242 when compared to the Control samples. Consistent positive correlation in this 8-way comparison  
1243 confirmed statistical identity between biological repeat experiments. The trend line in the plot is  
1244 the result of the function: stats::loess (R Package Documentation) [49], using default parameters.  
1245 The fitted curves shown with 95% confidence band.

1246

1247 **Figure. 12 Unsupervised clustering of HIVrepeat2.** Panel A) shows unsupervised  
1248 clustering obtained at K euqals 10. Panel B) Violin plot of HIV-1 transctipts/cell in the 10  
1249 clusters identified at K10 (Scran's buildSSNGraph using the PCA as input). PIC cells with  
1250 detectable HIV-1 transcripts, were distributed throughout clusters 1, 2 and 4-10. Cluster 3  
1251 contained 135 of the 227 cells in the semi supervised Proivirus cluster (circled in red).

1252

1253 **Figure 13 The Distribution of HIV-1 transcripts throughout Proivirus and**  
1254 **PIC/Bystander clusters in HIVrepeat2.** Violin plots of DHIV3-mCherry transcript/cell in cells

1255 from the Proivirus and PIC clusters showing transcript level and cell number. As described above,  
1256 these were made with Seurat's VlnPot function. They show normalized log2 transcript levels.  
1257 The two patterns of transcript distribution observed in HIVrepeat1 are evident. The first pattern  
1258 is seen with gag-pol, tat, env and nef, in which high numbers of cells in the PIC/Bystander  
1259 cluster detectably express the transcripts. The second pattern is seen with gag, vif, vpr, vpu, and  
1260 mCherry, in which fewer Proivirus or PIC cluster cells are detected expressing the transcripts, but  
1261 those cells expressing the transcripts are doing so at slightly higher average levels of transcripts  
1262 per cell. It is difficult to compare transcript loads in the Proivirus cluster cells to the results in  
1263 HIVrepeat1 (Fig. 10) due to the lower number of Proivirus cells detected in this HIVrepeat2  
1264 experiment. In this experiment the ratio of Proivirus cells to PIC cells was 0.17. Nevertheless, the  
1265 relative patterns observed in HIVrepeat1 are observed here. Following Seurat QC, no Proivirus  
1266 cells expressing rev were detected. Negative control sequence (asp) shows no distribution.

1267

1268 **Figure 14. Psupertime analysis of Control, PIC/Bystander and Proivirus cell**  
1269 **transcriptomes.** Psupertime analysis is a supervised pseudotime approach that explicitly uses  
1270 the sequential labels as input. It uses a regression-based model that acknowledges the cell labels  
1271 to identify genes relevant to the process. Panel **A**) one thousand Control (WT), PIC/Bystander  
1272 (PIC/B) and Proivirus (Pro) cell transcriptomes were randomly selected and analyzed. Imposition  
1273 of identity revealed a pseudo-evolution of Control to PIC/Bystander to Proivirus cell  
1274 transcriptomes. Panel **B**) distribution of HIV-1 transcripts through these clusters agrees with  
1275 results shown in Figure 5, showing no bias toward early or later gene transcripts.

1276

1277 **Figure 15. Western blot analysis for phospho- Rb or IkB in protein from mCherry**  
1278 **negative versus mCherry positive cells.** Cells infected with DHIV3-mCherry were purified by  
1279 FACS sorting based on their expression of mCherry fluorescence. Lane 1) Protein from Control  
1280 cells; Lane 2) Protein from PIC/Bystander cells; Lane 3) Protein from Provirus cells. Phospho-  
1281 Rb (Phospho-T821 Rb antibody) was used to quantify Rb pocket phosphorylation, anti- Rb  
1282 control antibody was used to quantify Rb protein levels relative to actin (visualized with beta-  
1283 actin antibody). MCherry protein confirmed with anti-mCherry antibody used in Figure 7.  
1284 PIC/Bystander cells show lowest level of Rb phosphorylation, Provirus show the highest, in  
1285 close agreement with Transcription Factor Targeting results. Panel B, Lane 1) Protein from  
1286 Control cells; Lane 2) Protein from PIC/Bystander cells; Lane 3) Protein from Provirus cells.  
1287 Phospho-IkB S32 antibody was used to quantify activated IkB. Control cells show lowest level  
1288 of IkB phosphorylation, no difference was detectable between Provirus and PIC Cluster cells.

1289  
1290 **Figure 16 Sequential infection of THP-1 cells with DHIV3-mCherry followed 24 hrs**  
1291 **later with GFP DHIV3.** Abscissa, mCherry signal, Ordinate, GFP signal. Provirus cluster,  
1292 mCherry positive, cells were 2 to 5 times more likely to make HIV-1 encoded GFP protein upon  
1293 second infection than PIC/Bystander cells upon second infection. Panel **A**) time equal 0 hrs;  
1294 addition of DHIV3-mCherry. Panel **B**) time equal 24 hrs; addition of DHIV3-GFP. Panel **C**) time  
1295 equals 48 hrs after DHIV3-mCherry addition, 24 hrs after DHIV3-GFP addition. Panel **D**) time  
1296 equals 72 hrs after DHIV3-mCherry addition, 48 hrs after DHIV3-GFP addition. Percentage of  
1297 mCherry cells also producing GFP, compared to cells producing mCherry only, is always 2 to 5  
1298 times higher than the percentage of cells making only GFP, compared to those cells not  
1299 producing either mCherry or GFP.

1300

1301

1302 **Supporting Information Figure Captions (S figures)**

1303

1304 **S-1 DHIV3-mCherry map.** Snapgene [50] map of DHIV3-mCherry plasmid.

1305

1306 **S-2 Seurat analysis of biological repeats HIVreplicate1 and HIVreplicate2.**

1307 HIVreplicate1a and HIVreplicate1b, and HIVreplicate2a and HIVreplicate2b, are technical repeat

1308 data. Technical repeats were conducted with each experiment. Figure shows PC analysis of

1309 biological repeat experiments. Technical duplicates were not different and so were combined for

1310 each repeat

1311

1312 **S-3 tSne projection of scRNA seq data from experiment HIVreplicate1.** Seurat analysis

1313 and t-SNE projection of data shown in Figure 2. Viral transcript numbers (**h**) were determined for

1314 cells containing any detected HIV-1 transcript, as described in methods. Orange dots represent

1315 high-level transcript load per cell, greater than 8 transcripts mapping to HIV-1 genes per cell, green

1316 dots indicate cells with lower transcript loads detected per cells, and blue dots indicate cells with

1317 no detectable HIV-1 transcripts. Barcodes of cells in Proivirus Cluster (A) tracked to Proivirus

1318 Cluster cells in UMAP analysis (Fig. 2).

1319

1320 **S-4 Unsupervised clustering of UMAP shown in Fig 4.** Left panels shows unsupervised

1321 clustering obtained at K values from 10 to 130. Right panels show Violin plots of HIV-1

1322 transcripts/cell in the clusters identified at the specified K values (Scran's buildSSNGraph using

1323 the PCA as input). Clusters 6 and 8 at K equal to 10 contained most of the cells in the semi

1324 supervised Proivirus cluster (circled in red) and were used to define Proivirus transcriptome,

1325 versus the remaining cells making up the semi supervised PIC/Bystander cluster. Stipulation of  
1326 lower K values means that during analysis any one given cell is clustered with a smaller number  
1327 of cells with similar transcriptomes.

1328

1329 **S-5 Real-time PCR analysis of DNA samples from Control, DHIV3-mCherry infected,**

1330 **and DHIV3-mCherry infected, integrase inhibitor treated THP-1 cells.** MW, molecular

1331 weight markers. Lanes 1, Control cell DNA; Lane 2, DNA from DHIV3-mCherry infected

1332 culture cells; Lane 3, DNA from DHIV3-mCherry infected cultures treated with integrase-

1333 inhibitor (25nM MK-2048). 100 ng of DNA was tested in each amplification unless noted.

1334 Primers used are described in Methods. Panels **A, B** and **C**) PCR demonstration of integrated

1335 proviral HIV DNA. Panel **A**) examples of the progression curves; upper curve represent lanes 2

1336 at 200 and 100 ng DNA respectively, middle curves reflect lanes 3, respectively, bottom 2 curves

1337 were generated by Control DNA . Panel **B**) melting curves; the upper curves represent lanes 2 at

1338 200 and 100 ng and lane 3 at 200 ng DNA respectively. Panel **C**) shows the amplicons generated

1339 from the integrated DNA using the nested PCR strategy described by Chun et al.[34] on a 1%

1340 agarose gel. The amplicon product sizes matched the predicted product size of 352 bp. These

1341 examples were from two biological replicates, one using 200 ng and one starting with 100 ng of

1342 starting DNA purified using Qiagen Blood and Tissue DNeasy kits. The agarose gel shows the

1343 integrated proviral DNA, assessed using an MJ PTC-200 thermal cycler and the nested PCR was

1344 evaluated using a Chromo-4 alpha unit. Note that the 200 ng samples with integrase inhibitor

1345 (Lane 3) show a small amount of integrated provirus DHIV3-mCherry DNA, demonstrating that

1346 the inhibitor did not completely inhibit the DHIV-mCherry integration. This is consistent with

1347 the 48 hr gag/p24 protein production seen in the immunoblot analysis (Fig. 7), and flow

1348 cytometry analysis (Fig. 6). Panels **D**, **E**, and **F**) show the same DNA samples used to detect 2-  
1349 LTR circle PIC cDNA from the second or two consecutive PCR runs. Panel **D**) Lower 3 curves  
1350 show progression curves with lack of 2LTR primer products in Control DNA, while all 6  
1351 biological repeats, 3 from DHIV3-mCherry infected cultures and 3 from DHIV3-mCherry  
1352 infected cultures treated with integrase inhibitor show amplification of PIC cDNA p2LTR  
1353 products. Panel **E**) shows the melting curves for these amplification products with the lower 3  
1354 curves representing Control DNA, the lowest curve representing Control DNA from the nested  
1355 PCR approach (see Methods). Panel **F**) shows the amplicons generated run out on a 1% agarose  
1356 gel. In these experiments the PCR was assessed using a Roche LightCycler 480. Biological  
1357 replicates of 100 ng starting DNA are represented as “rep 1” and “rep 2”, using the HIV F and  
1358 R1 primers of Brussels and Sonigo [33]. In the confirmation experiment lanes, under the  
1359 “nested” label in the agarose gel, wider bracketing primers were used in the first amplification  
1360 followed by the HIV F and R1 primers “nested” in the second run. Perhaps not surprising after  
1361 100 cycles, there are contaminating PCR products in the Control lanes; however, the expected  
1362 231 bp amplicon is not detectable in the Control cell DNA, while it is the predominant product in  
1363 DNA from either infected, or infected and integrase inhibitor treated cell DNAs. Panels **G**, **H**  
1364 and **I**) show the same DNA samples used to detect total DHIV-mCherry DNA, also assessed  
1365 using the Roche LightCycler 480. Primers used are described in Methods. Panel **G**) shows the  
1366 progression curves generated in this experiment, the lowest 2 curves represent Control cell DNA.  
1367 Panel **H**) shows the melting curves for these products, again the lowest 2 curves are from Control  
1368 cell DNA. Panel **I**) shows the amplicons generated, of predicted size, run out on a 1% agarose  
1369 gel. The real-time PCR results show roughly equivalent total amounts of DHIV3-mCherry in  
1370 infected culture DNAs, whether in the presence of integrase inhibitor or not. This indicates that

1371 overall, the total amounts of PIC cDNA are similar in integrase inhibitor treated and untreated  
1372 DHIV3-mCherry infected cultures.

1373

1374 **S-6 Unsupervised clustering of integrase inhibitor treated DHIV3 infected cells from**  
1375 **Figure 8.** No cluster corresponding to the Provirus cluster identified in HIVreplicate1 or  
1376 HIVreplicate2 could be identified, regardless of K value specified. Data were analyzed as in  
1377 Figure 4. Left panels shows unsupervised clustering obtained at K values from 10 to 130. Right  
1378 panels show Violin plots of HIV-1 transcripts/cell in the clusters identified at the specified K10  
1379 values (Scran's buildSSNGraph using the PCA as input). Stipulation of lower K values means  
1380 that during analysis any one given cell is clustered with a smaller number of cells with similar  
1381 transcriptomes.

1382

1383 **S-7 Feature plots of integrase inhibitor treated cultures.** Transcripts of DHIV3-mCherry  
1384 infection readily detectable in the presence of integrase inhibitor Panel **A**. Effects of cell cycle  
1385 (Panel **B**), mitochondrial gene expression (Panel **C**) and number of genes detected per cell  
1386 (Panel **D**) shown for comparison to Figure 3.

1387

1388 **S-8 Biological repeat experiments HIVreplicate1 and HIVreplicate2.** UMAP projections  
1389 of Seurat analysis of biological repeat experiments HIVreplicate1 (Panel **A**) and HIVreplicate2  
1390 (Panel **B**). Seurat analysis yielded 8.1% of cells in Provirus cluster from experiment  
1391 HIVreplicate1, 6% of cells in Provirus cluster in repeat HIVreplicate2, in agreement with

1392 percentages of mCherry positive percentages obtained for duplicate cultures analyzed by flow  
1393 cytometry.

1394

1395 **S-9 Unsupervised clustering of HIVrepeat2 UMAP projection.** Left panels shows  
1396 unsupervised clustering obtained at K nearest neighbor values from 10 to 130. Right panels show  
1397 Violin plots of HIV-1 transcripts/cell in the clusters identified at the specified K values (Scran's  
1398 buildSSNGraph using the PCA as input). Cluster 3 at K equal to 10 contained most of the cells in  
1399 the semi supervised Provirus cluster (circled in red) and was used to define Provirus  
1400 transcriptome, versus the remaining cells making up the semi supervised PIC/Bystander cluster.  
1401 Stipulation of lower K values means that during analysis any one given cell is clustered with a  
1402 smaller number of cells with similar transcriptomes.

1403

1404 **S-10. The Distribution of HIV-1 transcripts throughout Provirus and PIC/Bystander**  
1405 **clusters of HIVreplicat2.** Feature plot showing distribution of cells from UMAP in Fig. S-8  
1406 containing detectable DHIV3-mCherry transcripts. As describe above, these UMAP projections  
1407 were made with Seurat's FeaturePlot function. They are colored by expression of individual  
1408 genes (UMAP projection colored by walktrap, normalized log2 values). ASP is a negative  
1409 control, bacterial gene transcript sequence. The distribution repeats the results obtained in  
1410 HIVrepeat1 (Fig. 10A).

1411

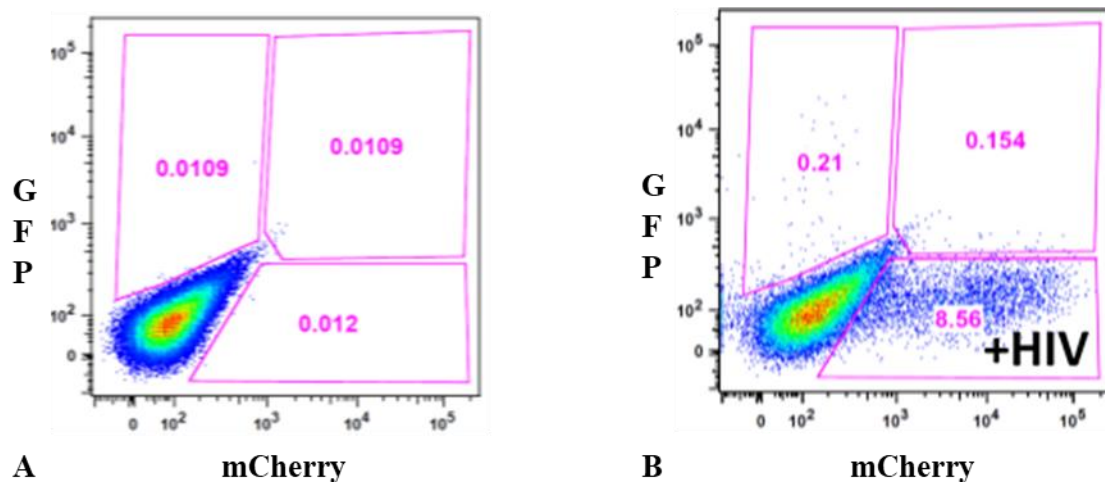
1412 **S-11 Differential gene expression comparison of Provirus and PIC cluster gene**  
1413 **transcripts from biological repeat experiments.** Consistent positive correlation of common

1414 DGE in HIVreplicate1 (abscissa) versus HIVreplicate2 (ordinate) repeat experiments  
1415 (Spearman's rank correlation coefficient of all common genes 0.384), agreed with Hallmark and  
1416 REACTOME analyses that showed similar pathways up- or down- regulated in the Proivirus  
1417 versus PIC/Bystander clusters of the biological repeat experiments.

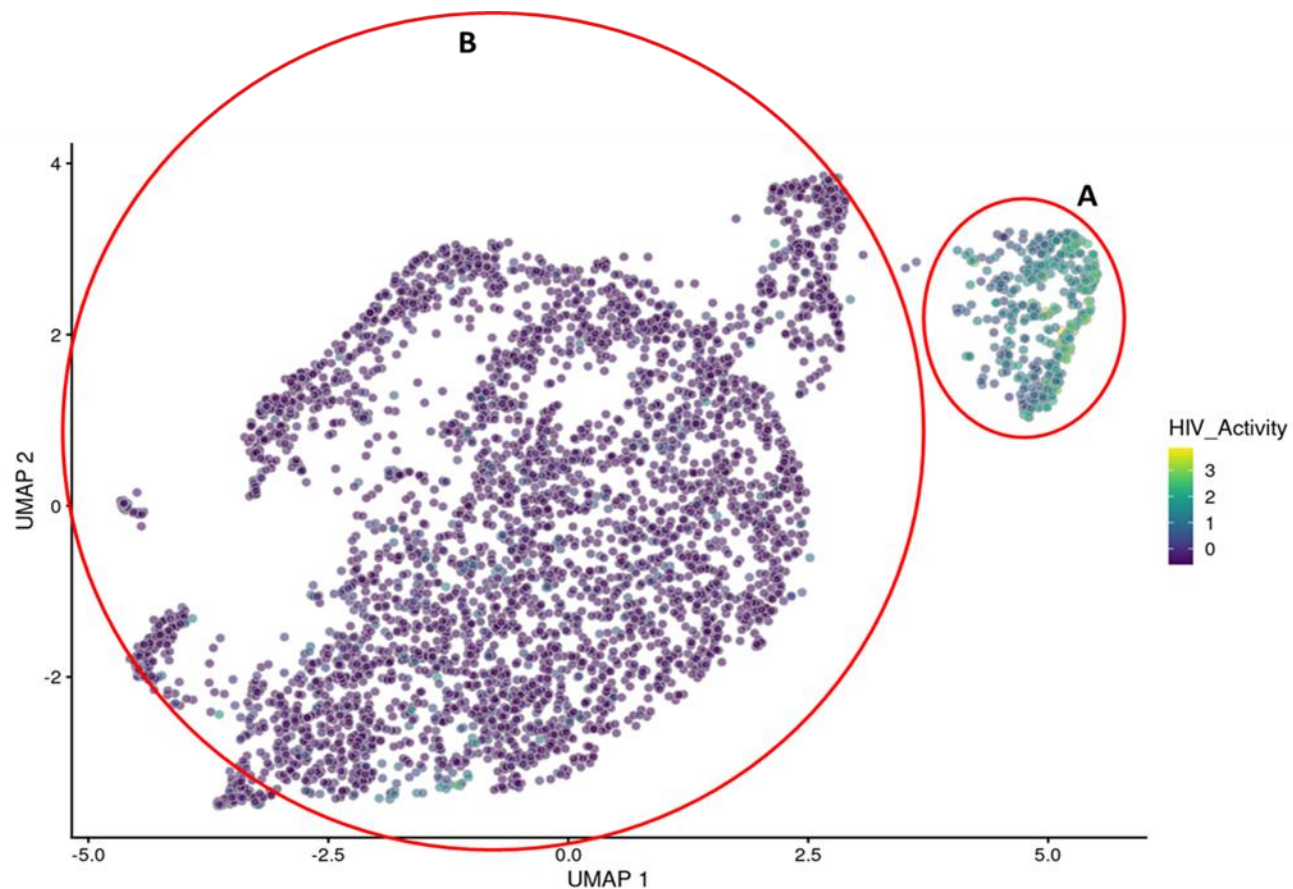
1418

1419 **S-12 Sequential infections of primary human lymphocyte and macrophage cultures.**  
1420 Active, mCherry positive, cells were 2 to 5 times more likely to make HIV-1 encoded GFP  
1421 protein upon second infection than PIC/Bystander cells upon second infection. Panels **A, H)**  
1422 primary cultures of T-lymphocytes and macrophages at time equals 0 hrs, respectively. Panels **B-**  
1423 **D)** primary lymphocytes infected with low titer DHIV3. Panels **E-G)** primary lymphocytes  
1424 infected with high titer. Panels **H-K)** primary macrophages. Percentage of mCherry cells also  
1425 producing GFP, compared to cells producing mCherry only, is always 2 to 5 times higher than  
1426 the percentage of cells making only GFP, compared to those cells not producing mCherry.  
1427 Panel **A** and **H)** Time equal 0 hrs; addition of DHIV3-mCherry. Panel **B, E** and **I)** time equal 24  
1428 hrs; addition of DHIV3-GFP. Panel **C, F** and **J)** time equals 48 hrs after DHIV3-mCherry  
1429 addition, 24 hrs after DHIV3-GFP addition. Panel **D, G** and **K)** time equals 72 hrs after DHIV3-  
1430 mCherry addition, 48 hrs after DHIV3-GFP addition.

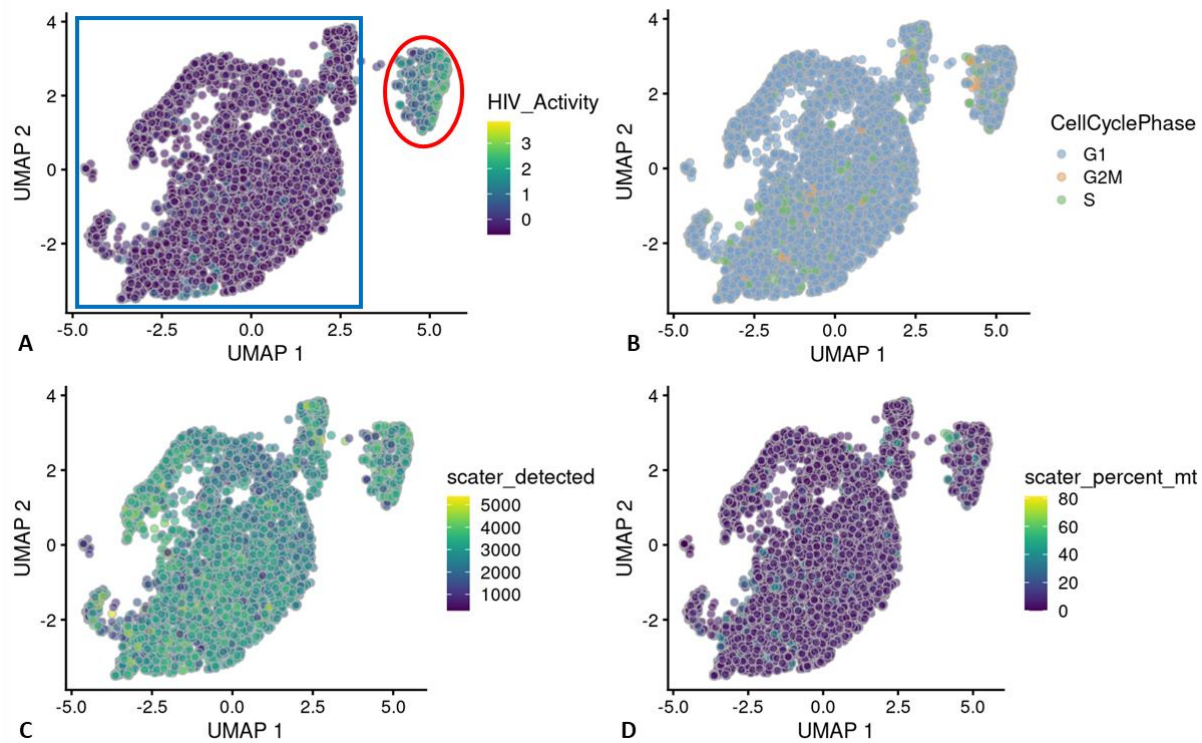
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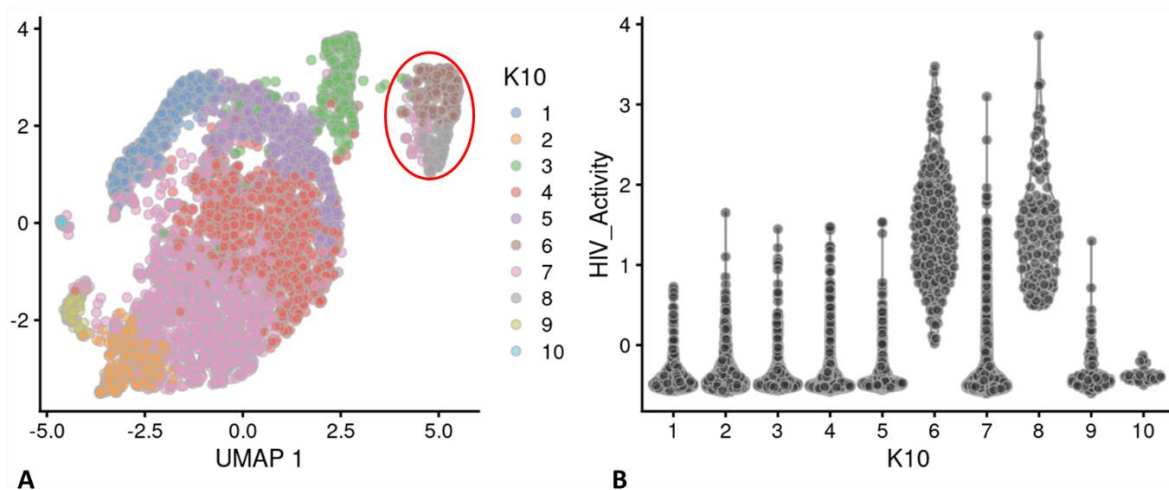
**Figure 1. Flow cytometry analysis of DHIV3-mCherry infected THP-1 cells.** Panel A) mock infection of PMA activated THP-1 cells. Panel B) PMA activated THP-1 cells infected with DHIV3-mCherry. Abscissa, mCherry (Texas Red) emission. Ordinate, GFP (FITC) emission. mCherry positive cells equal approximately 8.5% of the total viable cell population.



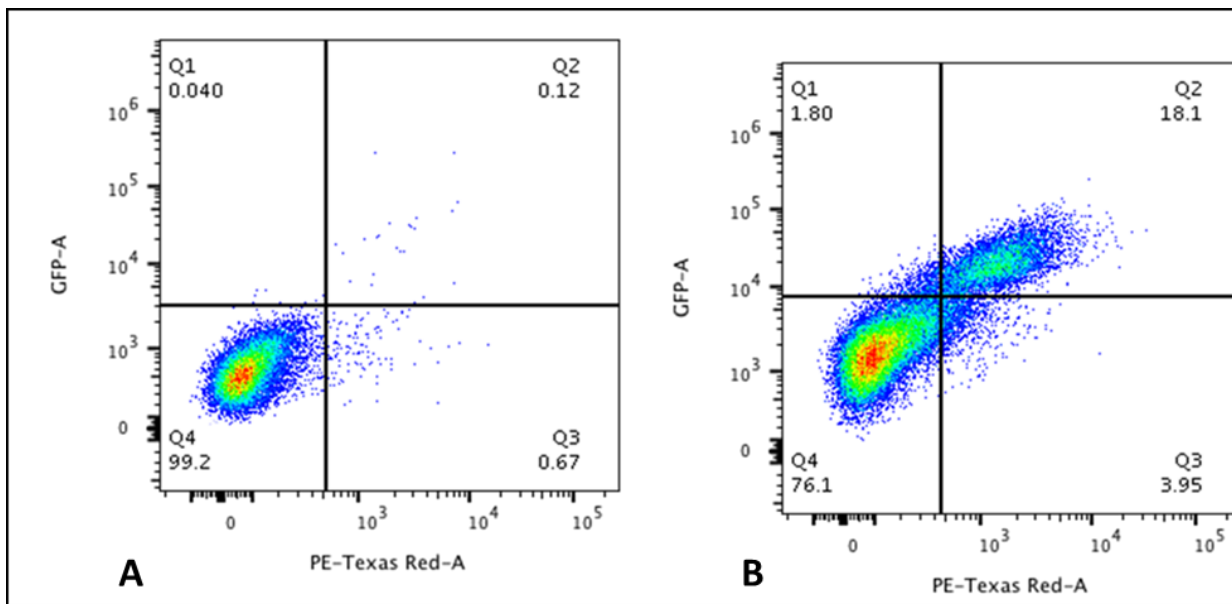
**Figure 2. UMAP projection of scRNA seq data from experiment HIVreplicate1.** Greater than 14,000 different cellular genes were detected in this analysis, including the 9 viral genes and mCherry message originating from DHIV3-mCherry. A semi-supervised two cluster model was adopted, the smaller “Provirus cluster” (cluster A) was 8.1% of the total cell population, approximately equivalent to the percentage of mCherry positive cells from Fig. 1. The two semi-supervised clusters are circled in red. PIC/Bystander cluster is indicated as cluster B. The HIV activity scale presents the Seurat module score that is described in methods. Input data in this analysis included 33,819 PCA entries. The number of cells with detected genes following Seurat QC in the PIC/Bystander cluster was 12% of the total population. Bar codes of the same cells tracked to the Provirus clusters, regardless of whether the clusters were generated using UMAP or Seurat-tSne tools (Fig. S-3).



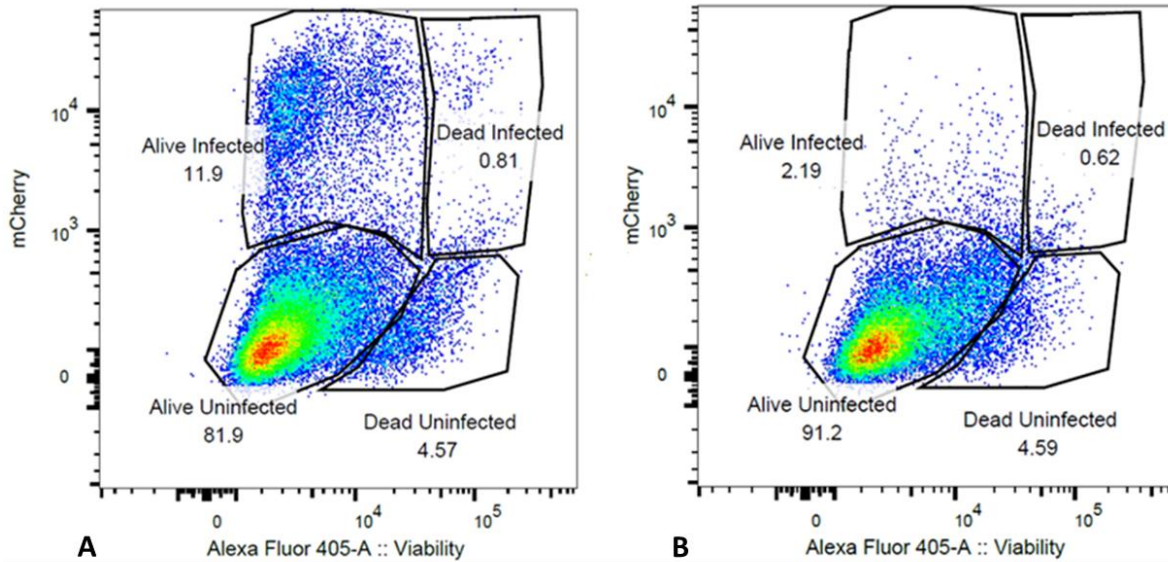
**Figure 3. UMAP Feature plot of experiment HIVreplicate1.** Panel A) Distribution of HIV-1 transcript positive cells shown in Fig. 2. Provirus cluster circled in red, PIC/Bystander cluster in the blue square. Panels B), C) and D), respectively, show no influence of cell cycle, number of genes detected per cell, or percent mitochondrial transcripts (positively correlated with cell stress) on the distribution of PIC cells (HIV-1 transcript containing cells) throughout the PIC/Bystander cluster.



**Figure. 4 Unsupervised clustering of UMAP shown in Fig 2.** Panel **A**) shows unsupervised clustering obtained at K equals 10. Panel **B**) Violin plot of HIV-1 transcripts/cell in the 10 clusters identified at K10 (Scran's buildSSNGraph using the PCA as input). PIC cells with detectable HIV-1 transcripts, were distributed throughout clusters 1-5, 7 and 9-10. Clusters 6 and 8 contained 372 of the 381 cells included in the semi-supervised Provirus cluster (circled in red). Stipulation of lower K values means that during analysis any one given cell is clustered with a smaller number of cells with similar transcriptomes.

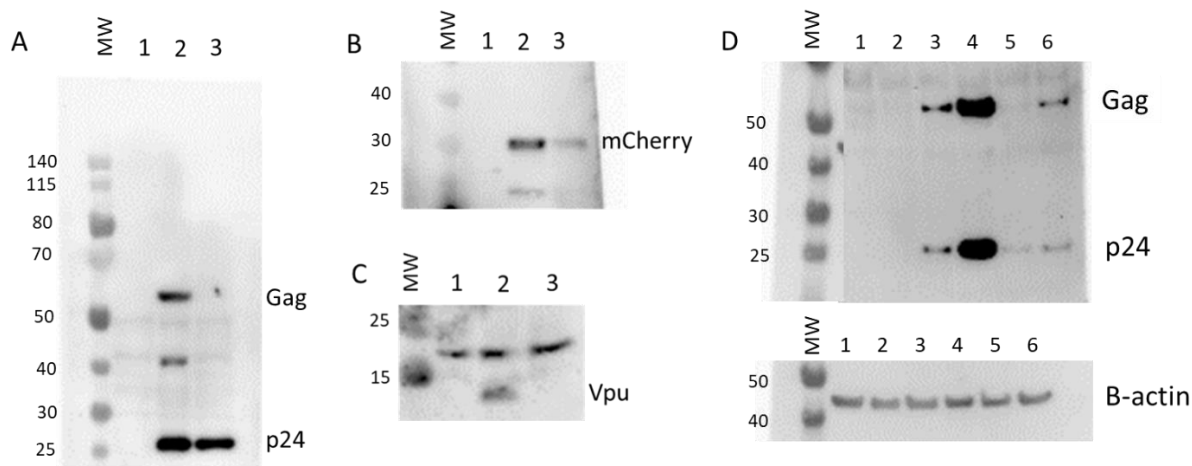


**Figure 5. Flow cytometry analysis of DHIV3-mCherry infected THP-1 cells using p24 antibody.** Panel A) mock infection. Panel B) mCherry expression was positively correlated with p24/Gag antigen detection by flow cytometry. Abscissa mCherry (Texas Red) emission. Ordinate, GFP (FITC) emission. mCherry/p24 positive cells equal approximately 18% of the total viable cell population in this experiment.

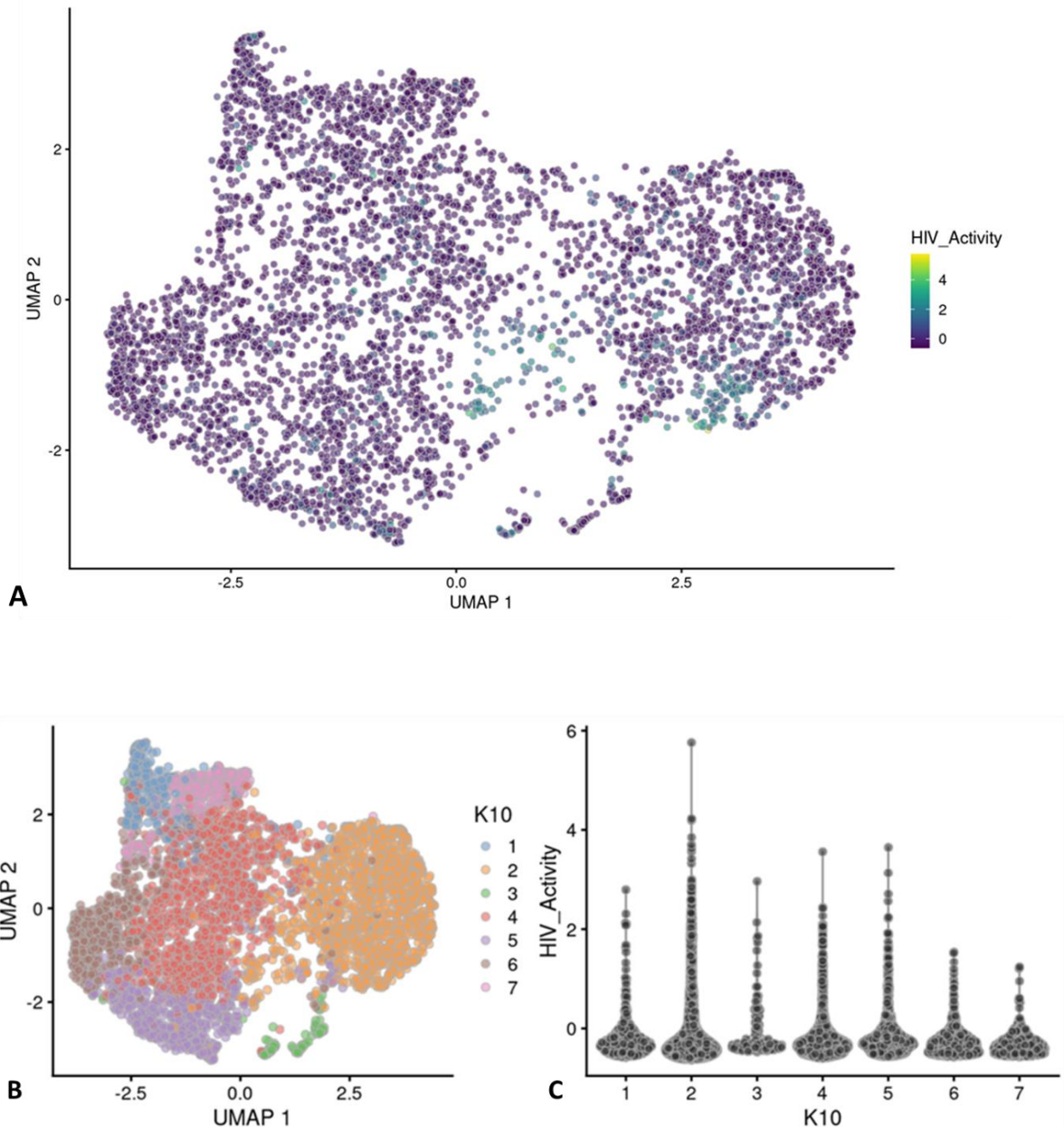


**Figure 6. Integrase inhibitor treatment selectively reduces mCherry positive cells.**

Panel A) Flow cytometry analysis of DHIV3-mCherry infected THP-1 cells, versus viability stain. Abscissa shows viability stain intensity, ordinate shows mCherry intensity. Infected (Active), mCherry-producing cells account for approximately 12 % of the cell population. Panel B) Same as A except with the addition of 25nM MK-2048 integrase inhibitor at the time of infection. Integrase inhibitor effectively reduces the number of mCherry producing cells, without decreasing cell viability.



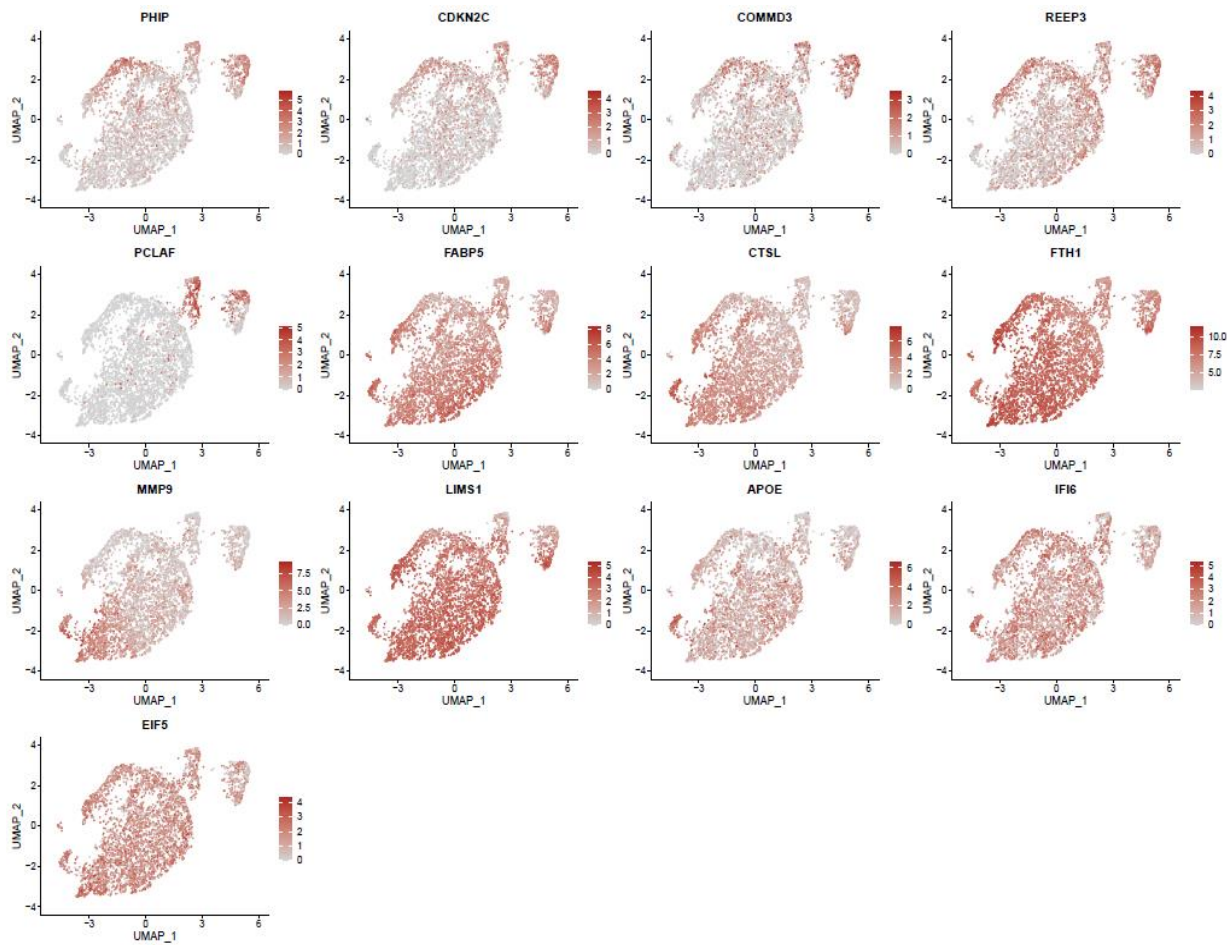
**Figure 7. Effect of integrase inhibitor on mCherry, p24, Gag, and Vpu protein production in cultures containing DHIV3-mCherry infected cells.** MW, molecular weight markers. Lane 1, Control cell protein; Lane 2, protein from DHIV3 infected culture; Lane 3, protein from DHIV3 infected cultures treated with integrase inhibitor (25nM MK-2048) as shown above in Figure 6. **A)** Lane 2, p24, and Gag precursor proteins visualized with p24 antibody used above in (Fig. 6), and HRP linked anti-mouse secondary antibody. The p24 band in lane 3 is residual from infection as reported in the literature [10]. The presence of precursor proteins in lane 2 shows Proivirus synthesis in cultures containing Proivirus cells. **B)** Antibody used was goat anti-mCherry, developed with HRP linked anti-goat secondary. mCherry protein clearly visible in preparations containing Proivirus cell protein. **C)** Lane 2, Vpu detected with rabbit antibody, visualized using HRP linked anti-rabbit secondary. The resolution of the image slightly compromised due to the small size of Vpu protein. **D)** Lanes 1 and 2, Control cell protein at 24 and 48 hrs respectively; lanes 3 and 4, protein from DHIV3 infected culture at 24 and 48 hrs respectively; lanes 5 and 6, protein from DHIV3 infected cultures treated with integrase inhibitor (as above) at 24 and 48 hrs respectively. At 24 hrs post-infection, we only found both p24 and precursor Gag proteins in the protein samples from DHIV3 infected cells in the absence of integrase inhibitor. At 48 hrs post-infection, in the absence of integrase inhibitor, the amounts of detectable p24 and Gag proteins were dramatically increased from levels at 24 hrs post-infection. As seen initially (Panel A), some p24 protein was detectable in integrase inhibitor treated cultures at 24 hrs post-infection, however Gag is not detectable at this time. At 48 hrs post-infection in the integrase inhibitor treated-cultures, some Gag protein does becomes detectable, reflecting production in cells that escaped complete integrase inhibition. This is in agreement with our flow cytometry analysis that showed suppressed, but still detectable numbers of mCherry positive cells in the integrase inhibitor treated cultures. The Gag precursor proteins only appear in the integrase inhibitor treated culture proteins 48 hrs after treatment. All antibodies, sources, and dilutions are provided in Methods.



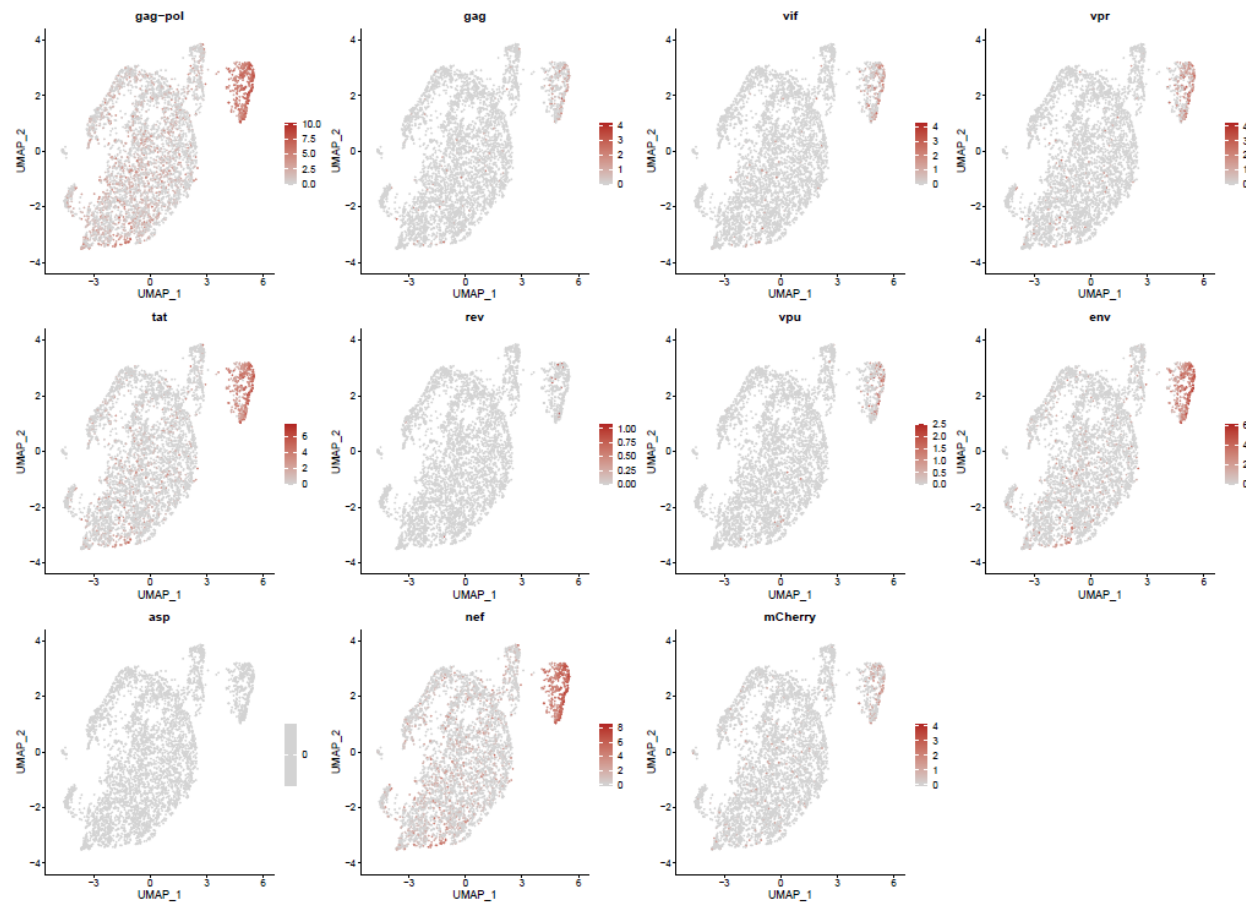
**Figure 8. UMAP analysis of integrase inhibitor treated DHIV3-mCherry infected THP-1 cells.** The experiment performed as shown in Figure 6 B, with 25nM MK-2048 added at the time of DHIV3 addition. Data were analyzed identically to data shown in Figure 2. Panel A) Feature plot showing the distribution of cells containing HIV-1 transcript, generated as described. Panel B) K10 unsupervised clustering generated 7 clusters (Scran's buildSSNGraph using the PCA as input). HIV-1 transcripts were distributed equally throughout all of them. No cluster corresponding to the "Provirus" cluster detected in Figure 4 was detected, regardless of K value used (see Fig. S-6). These data agree with the concept that integrase inhibitors selectively target and reduce the number of Provirus cluster cells.

pathway	pval	padj	ES	NES	nMoreExt	size	leadingEdge	enriched
Tnfa Signaling Via NfkB	0.00033	0.001399	-0.74028	-2.31254	0	187	NINJ1, SAT1, IER3, IL1B, NFKBIA, G0S2, SGK1, SOD2, CDKN1A, PPP1R15	negative
Complement	0.000316	0.001399	-0.67706	-2.07107	0	160	CTSL, CTSD, CTSB, LIPA, TIMP1, CD36, DUSP6, HSPA5, C3, PLEK, APOC1	negative
Inflammatory Response	0.000318	0.001399	-0.66804	-2.05591	0	166	CSAR1, IL1B, NFKBIA, TIMP1, CDKN1A, TNFRSF1B, IFNGR2, CXCL8	PTP negative
Coagulation	0.000286	0.001399	-0.70045	-2.00402	0	96	MMP9, CTSB, TIMP1, DUSP6, C3, GSN, PLEK, ISCU, APOC1, CTSH, ANXA	negative
Cholesterol Homeostasis	0.000272	0.001399	-0.71023	-1.94144	0	70	FABP5, SQLE, LPL, ATF5, S100A11, ALCAM, PNRC1, ETHE1, LGALS3, SCD	negative
Epithelial Mesenchymal Transition	0.000307	0.001399	-0.63938	-1.93759	0	147	SAT1, DAB2, VIM, TIMP1, CXCL8, EMP3, COL6A2, TPM4, PPIB, SPP1, GL	negative
Hypoxia	0.000318	0.001399	-0.6227	-1.91128	0	164	IER3, PLIN2, S100A4, CDKN1A, PPP1R15A, HSPA5, ALDOA, KLF6, PNRC	negative
Mtor1 Signaling	0.000336	0.001399	-0.54085	-1.69467	0	195	SQLE, INSIG1, CALR, CDKN1A, PPP1R15A, FKB2, HSPA5, SQSTM1, ITG1	negative
UV Response Up	0.0012	0.004614	-0.52872	-1.59832	3	141	EIF5, NFKBIA, PPIF, ATP6V1F, SOD2, SQSTM1, ARRBL2, ALDOA, MGAT1	negative
Apoptosis	0.001493	0.00533	-0.52273	-1.57706	4	139	SAT1, IER3, IL1B, LMNA, TIMP1, SOD2, CDKN1A, SQSTM1, BCAP31, GSN	negative
Kras Signaling Up	0.002443	0.008142	-0.50409	-1.52704	7	146	MMP9, IL1B, MAFB, G0S2, PPP1R15A, DUSP6, TNFRSF1B, ITGB2, GPNN	negative
IL6 Jak Stat3 Signaling	0.00326	0.010187	-0.59782	-1.63624	11	71	IL1B, CD36, TNFRSF1B, IFNGR2, HMOX1, JUN, CD14, A2M, TNFRSF12A	negative
P53 Pathway	0.003922	0.01032	-0.47746	-1.48489	11	180	CTSD, NINJ1, SAT1, IER3, S100A4, CDKN1A, PPP1R15A, ZFP36L1, PTPRE	negative
IL2 Stat5 Signaling	0.003552	0.01032	-0.47887	-1.48033	10	173	PLIN2, COL6A1, TNFRSF1B, SNX9, KLF6, ALCAM, GSTO1, CD81, PHLDA1	negative
Xenobiotic Metabolism	0.008534	0.021335	-0.4758	-1.43988	27	145	NINJ1, TDO2, APOE, CD36, PGD, GSTO1, NQO1, HMOX1, ABCC3, BLVRE	negative
Pi3k Akt Mtor Signaling	0.009983	0.022688	-0.52419	-1.49267	34	93	CALR, CDKN1A, SQSTM1, RPS6KA1, VAV3, HSP90B1, ACTR3, ARF1, ARF1	negative
Apical Junction	0.009855	0.022688	-0.47146	-1.43125	31	150	MMP9, INSIG1, RAC2, ZYX, CD276, ICAM1, ADAM9, STX4, FYB1, VASP	/negative
Angiogenesis	0.011132	0.023192	-0.72619	-1.6345	46	24	LPL, TIMP1, S100A4, SPP1, THBD, NRP1, ITGAV, LRPAP1, VEGFA	negative
Reactive Oxigen Species Pathway	0.010978	0.023192	-0.61972	-1.58502	41	47	FTL, SOD2, NQO1, JUNB, MBP, MSRA, LAMTORS, SELENOS, TXNRD1, GL	negative
Interferon Gamma Response	0.015349	0.030699	-0.43862	-1.36348	46	179	NFKBIA, SOD2, CDKN1A, LY6E, B2M, WARS, ISG20, ITGB7, IFI35, IFITM3	negative
Myogenesis	0.017438	0.033535	-0.45597	-1.37958	57	142	CDKN1A, CD36, GSN, SPHK1, COL6A2, NQO1, TPM3, SYNGR2, ATP6AP1	negative
Protein Secretion	0.022533	0.041727	-0.49591	-1.41213	78	93	CD63, ATP6V1H, ABCA1, ARF1, BNIP3, SNX2, TMED2, CLTA, AP2M1, SE	negative
Androgen Response	0.023975	0.042813	-0.50467	-1.42398	85	85	SAT1, INSIG1, SGK1, CCND1, B2M, ACTN1, SCD, TSC22D1, MYL12A	ZM1 negative
TGF Beta Signaling	0.026772	0.046158	-0.57256	-1.47921	101	50	PPP1R15A, IFNGR2, JUNB, RAB31, FKB2, ENG, SLC20A1, TGIF1, XIAP	negative
UV Response Down	0.047167	0.078612	-0.44009	-1.30127	158	122	DAB2, INSIG1, MGLL, RND3, SDC2, NRP1, SYNJ2, NOTCH2, ZM1Z1, SLC2	negative
Allotransplant Rejection	0.061941	0.099906	-0.41295	-1.25319	200	149	MMP9, IL1B, TIMP1, IFNGR2, ITGB2, SPI1, HLA-E, B2M, WARS, ICAM1	(negative
E2F Targets	0.000142	0.001399	-0.738487	-2.126021	0	199	STMN1, CDKN2C, SMC4, H2AFZ, CK51B, HMGB2, TOP2A, CDKN3, RPA3	positive
Myc Targets V1	0.000141	0.001399	-0.650748	-1.875111	0	200	H2AFZ, TYMS, DUT, RPLP0, EEF1B2, RPS3, RPSS, DEK, RPS2, RANBP1, R1	positive
G2-M Checkpoint	0.000142	0.001399	0.633325	1.818855	0	195	STMN1, CDKN2C, SMC4, H2AFZ, HMGN2, CKS1B, H2AFV, TOP2A, CDKN	positive
Spermatogenesis	0.000156	0.001399	0.692643	1.799937	0	83	CDKN3, RPL39L, PEBP1, GFI1, CCNB2, TOPBP1, AURKA, CDK1, TLE4, PO	positive
Oxidative Phosphorylation	0.003883	0.01032	0.511797	1.459765	26	183	LDHB, MPC1, UQCRC1, COX8A, SLC25A5, UQCRCQ, COX5A, SLC25A3, COX	positive

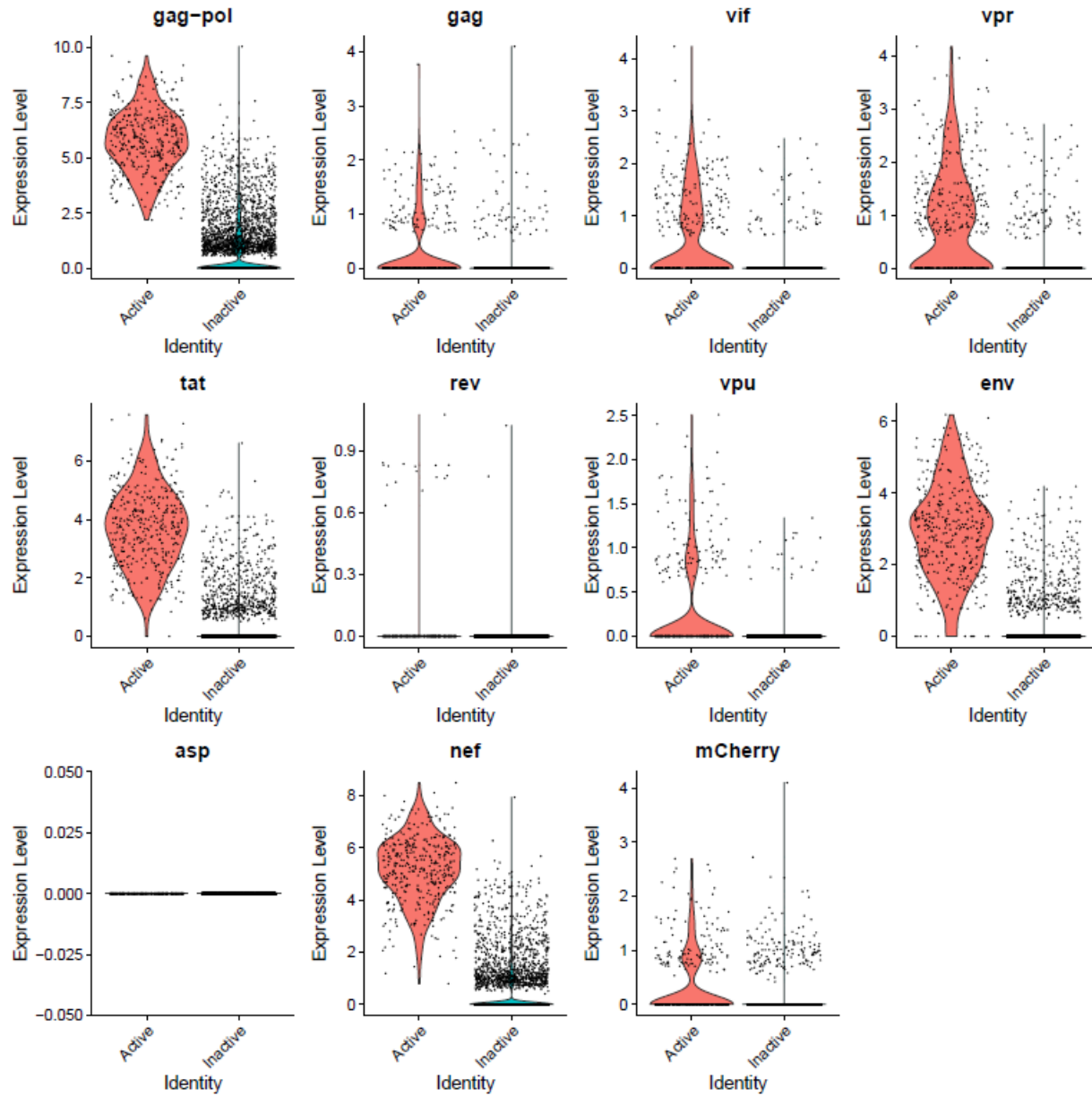
**Table I Hallmark analysis of gene pathways up or down-regulated detected in Proivirus versus PIC/Bystander cluster cells (respectively).** GSEA Hallmark analysis (fgsea R package) of metabolic pathways negatively or positively regulated ( $p<0.1$ ) in the Proivirus cluster transcriptome versus the PIC/Bystander cluster transcriptome. Pathways up-regulated in Proivirus cells include E2F, Myc targets, G2-M checkpoint, spermatogenesis, and oxidative phosphorylation. Down-regulated pathways identified are more numerous, but notably included TNF $\alpha$  signaling via NFkB, inflammatory response genes, apoptosis, and interferon  $\gamma$  response. The pairwiseTTests function from Scran was used determine the significance of genes between groups. The significant DGE subsets were used for all comparisons.



**Figure 9. Distribution of gene transcripts exhibiting high levels of differential expression between Proivirus and PIC/Bystander clusters.** Feature plot showing the distribution of cells from Figure 2, containing transcripts of 10 of the 20 most highly differentially expressed transcripts in Proivirus versus PIC/Bystander GSEA data sets. APOE, IFI6, and EIF5 were also included because they were highly expressed in the PIC/Bystander cluster. As above, these UMAP projections were made with Seurat's FeaturePlot function. They are colored by expression of individual genes (normalized log2 values). Highly expressed genes in Proivirus cluster cells include PHIP (Pleckstrin Homology Domain Interacting Protein), CDKN2C (Cyclin Dependent Kinase Inhibitor 2C), COMMD3 (COMM Domain Containing Protein 3), REEP3 (Receptor Accessory Protein 3), and PCLAF (PCNA Clamp Associated Factor). Highly expressed transcripts detected in the PIC/Bystander cell transcriptome include FABP5 (Fatty Acid Binding Protein 5), CTSL (Cathepsin L), FTH1 (Ferritin Heavy Chain 1), MMP9 (Matrix Metallopeptidase 9), LIMS1 (LIM Zinc Finger Domain Containing 1), APOE (Apolipoprotein E), IFI6 (Interferon Alpha Inducible Protein 6), and EIF5 (Eukaryotic Translation Initiation Factor 5).



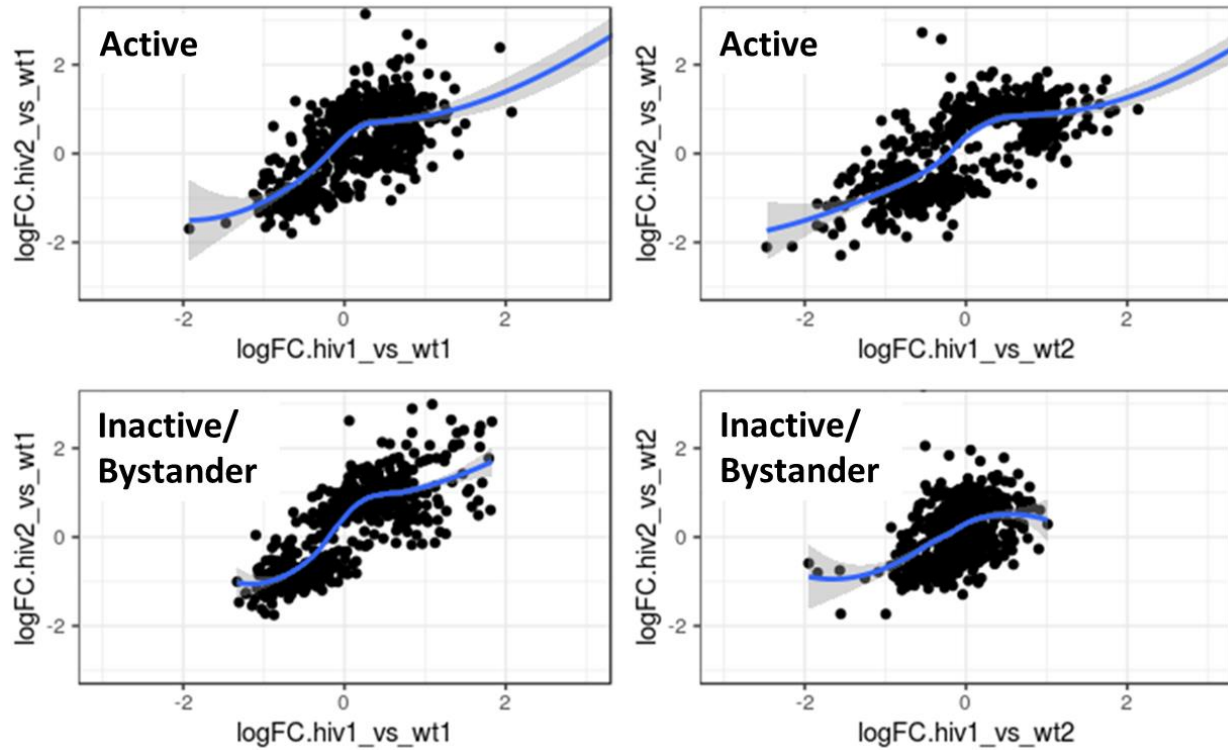
**A**



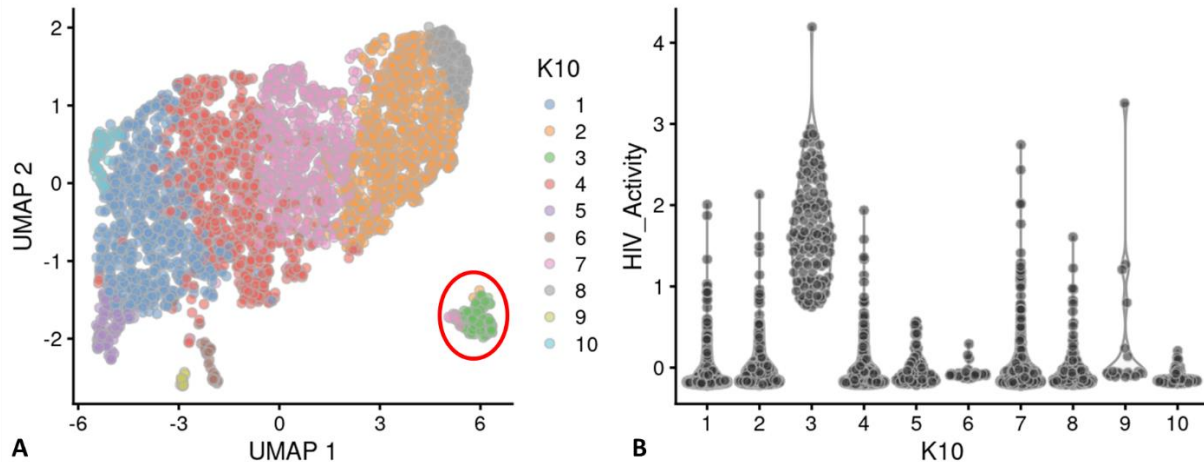
**B**

**Figure 10. The Distribution of HIV-1 transcripts throughout Provirus and PIC/Bystander clusters.** Panel A) Feature plot showing the distribution of cells from UMAP in Figure 2 that contain detectable DHIV3-mCherry transcripts. As described above, these UMAP projections were made with Seurat's FeaturePlot function. They are colored by expression of individual genes (UMAP projection colored by walktrap, normalized log2 values). ASP is a negative control, bacterial gene transcript sequence. B) Violin plots of DHIV3-mCherry transcript/cell in cells from the Provirus and PIC clusters showing transcript level and cell number. The provirus cluster contained transcriptomes of 371 cells, the number of PIC cells in

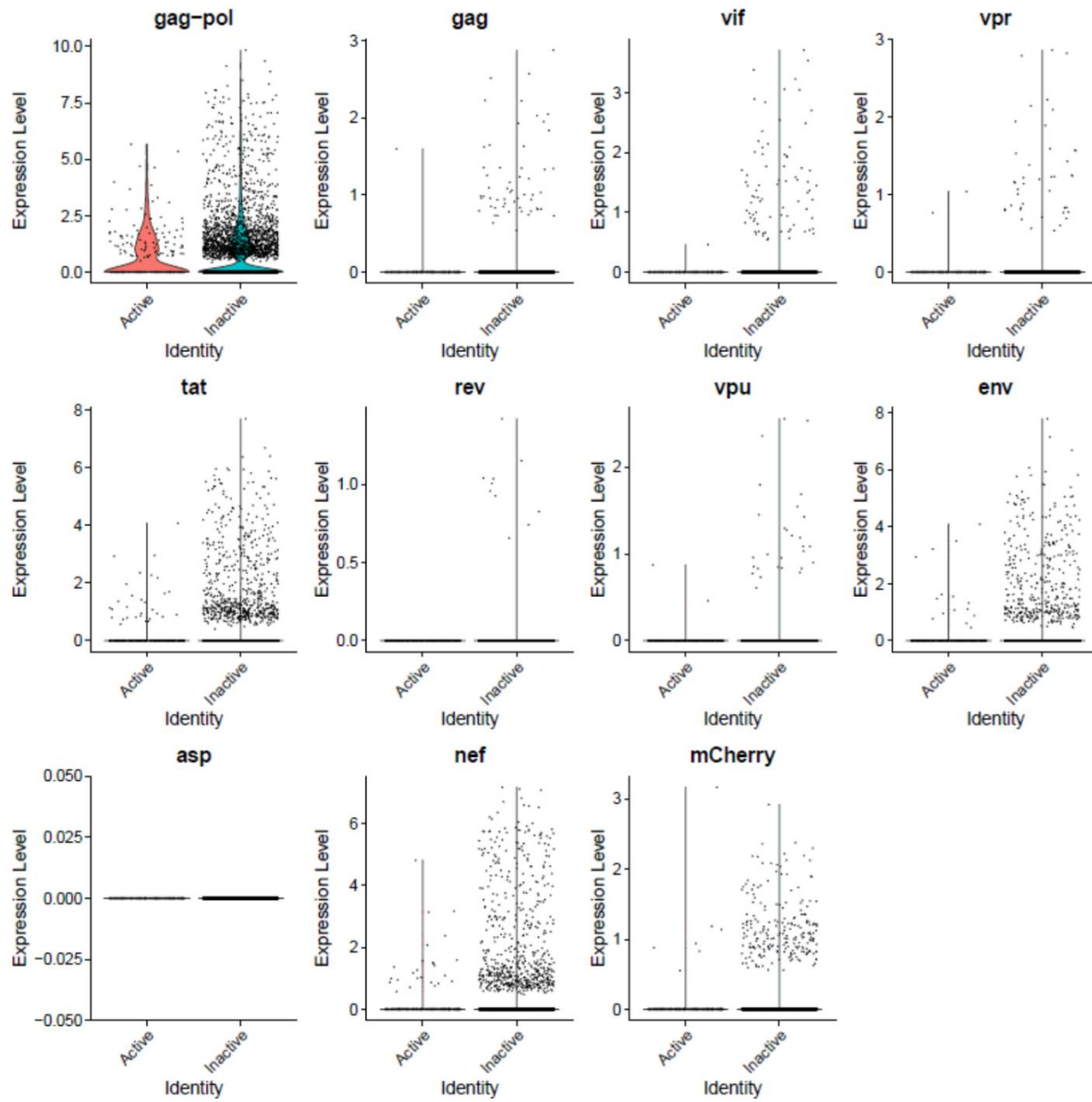
the PIC/Bystander cluster was 569 cells, thus the Proivirus/PIC cell number ratio was 0.65. The plots were made with Seurat's VlnPot function. They show normalized log2 transcript levels. Two patterns of transcript distribution are evident. The first pattern is seen with gag-pol, tat, env, and nef, in which relatively high numbers of cells in the PIC/Bystander cluster express the transcripts, with the transcripts being detected in fewer numbers of Proivirus cluster cells. The second pattern is seen with gag, vif, vpr, rev, vpu, and mCherry, in which relatively equal absolute numbers of cells in the Proivirus and PIC/Bystander clusters are detected with the transcript sequences, remembering that there are more PIC cells than Proivirus cells. The relative transcript loads per PIC cell versus the Proivirus cells overlap. Negative control sequence (asp) shows no distribution.



**Figure 11. Differential gene expression comparison of Proivirus and PIC/Bystander cluster gene transcripts versus 2 independent biological repeat control (wt) experiments.** In every comparison, a significant positive correlation was obtained from the common detected differentially expressed genes of Proivirus or PIC/Bystander clusters in the two biological repeats when compared to the Control samples. Consistent positive correlation in this 8-way comparison confirmed statistical identity between biological repeat experiments. The trend line in the plot is the result of the function: `stats:::loess` (R Package Documentation) [47], using default parameters. The fitted curves are shown with 95% confidence band.

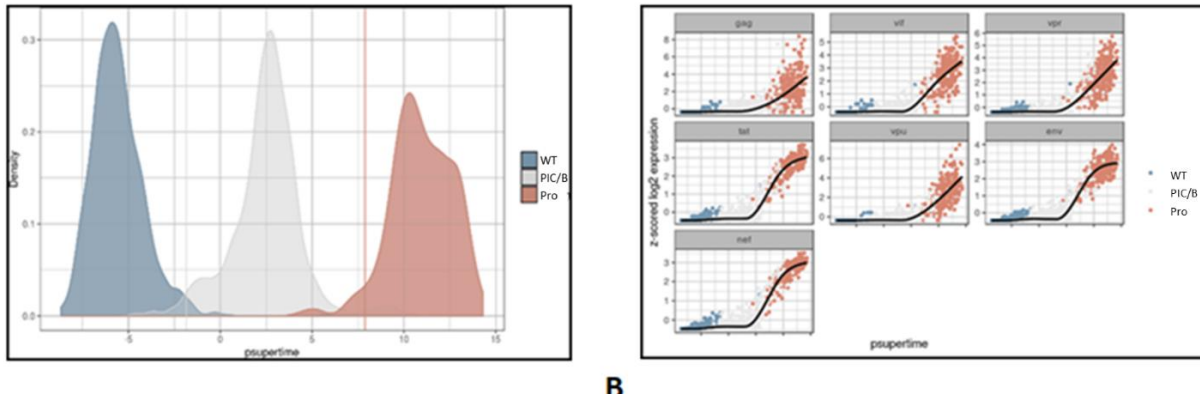


**Figure. 12 Unsupervised clustering of HIVrepeat2.** Panel **A**) shows unsupervised clustering obtained at K equals 10. Panel **B**) Violin plot of HIV-1 transcripts/cell in the 10 clusters identified at K10 (Scran's buildSSNGraph using the PCA as input). PIC cells with detectable HIV-1 transcripts, were distributed throughout clusters 1, 2 and 4-10. Cluster 3 contained 135 of the 227 cells in the semi supervised Proivirus cluster (circled in red).



**Figure 13 The Distribution of HIV-1 transcripts throughout Proivirus and PIC/Bystander clusters in HIVrepeat2.** Violin plots of DHIV3-mCherry transcript/cell in cells from the Proivirus and PIC clusters showing transcript level and cell number. As described above, these were made with Seurat's VlnPot function. They show normalized log2 transcript levels. The two patterns of transcript distribution observed in HIVrepeat1 are evident. The first pattern is seen with gag-pol, tat, env, and nef, in which high numbers of cells in the PIC/Bystander cluster detectably express the transcripts. The second pattern is seen with gag, vif, vpr, vpu, and mCherry, in which fewer Proivirus or PIC cluster cells are detected expressing the transcripts, but those cells expressing the transcripts are doing so at slightly higher average levels of transcripts per cell. It is difficult to compare transcript loads in the Proivirus cluster cells to the results in

HIVrepeat1 (Fig. 10) due to the lower number of Proivirus cells detected in this HIVrepeat2 experiment. In this experiment, the ratio of Proivirus cells to PIC cells was 0.17. Nevertheless, the relative patterns observed in HIVrepeat1 are observed here. Following Seurat QC, no Proivirus cells expressing rev were detected. Negative control sequence (asp) shows no distribution.



**A**

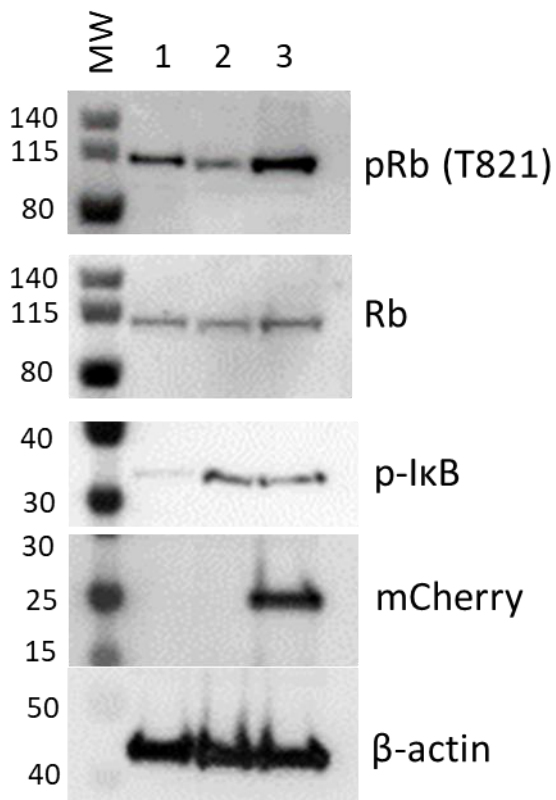
**B**

**Figure 14. Psuperstime analysis of Control, PIC/Bystander, and Provirus cell transcriptomes.** Psuperstime analysis is a supervised pseudotime approach that explicitly uses the sequential labels as input. It uses a regression-based model that acknowledges the cell labels to identify genes relevant to the process. Panel **A**) one thousand Control (WT), PIC/Bystander (PIC/B), and Provirus (Pro) cell transcriptomes were randomly selected and analyzed. Imposition of identity revealed a pseudo-evolution of Control to PIC/Bystander to Provirus cell transcriptomes. Panel **B**) distribution of HIV-1 transcripts through these clusters agrees with results shown in Figure 5, showing no bias toward early or later gene transcripts.

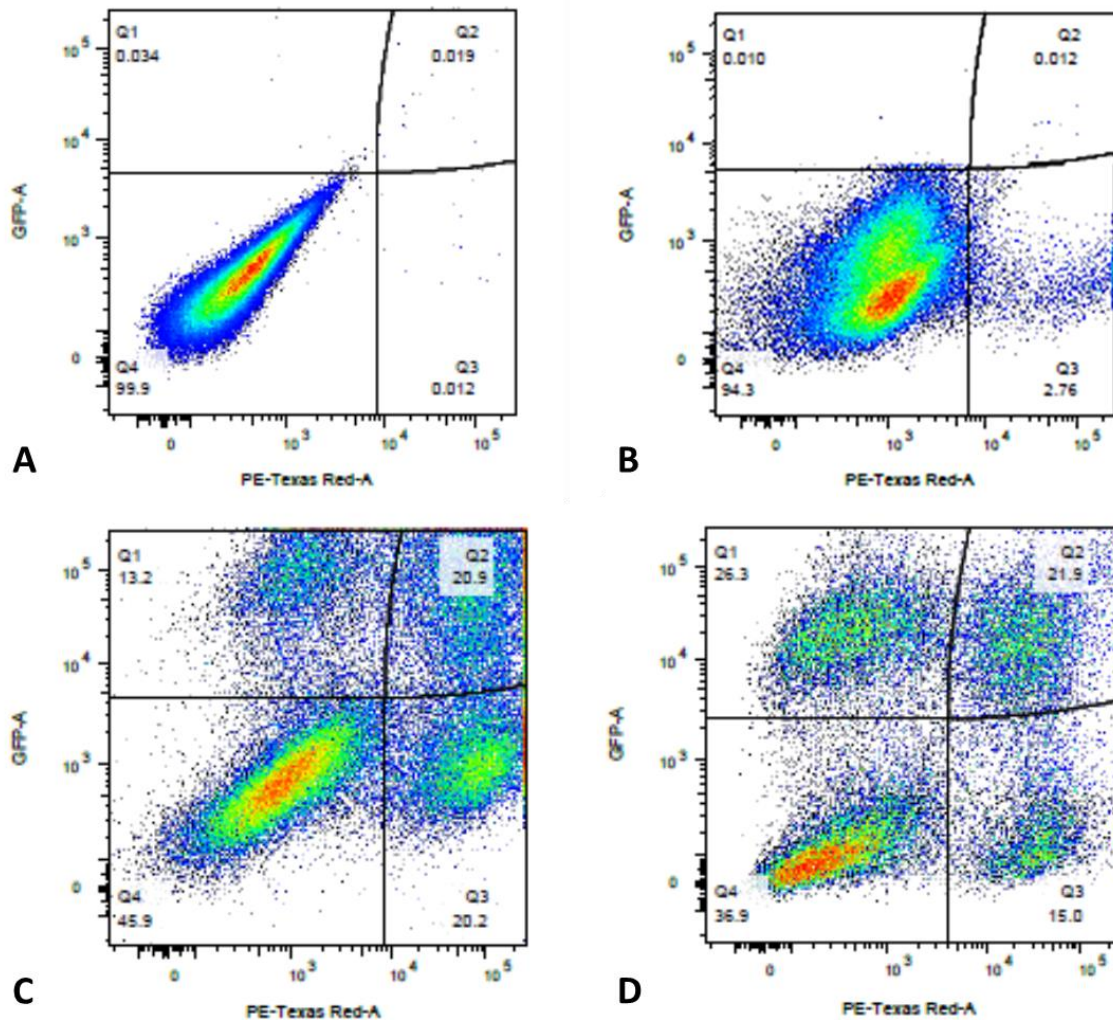
pathway	pval	padj	ES	NES	nMoreExt	size	leadingEdge	enriched
TGTYNNNNNRCARM	0.000271	0.008414	-0.67086	-1.80812	0	66	FRMD4A, ZEB2, CAMK1, BTG2, P2RX4, CDK2AP2, TSC22D1, RCAN1, negative	
NFKB.Q6_01	0.000338	0.008809	-0.553	-1.71204	0	179	MMP9, IL411, NFKBIA, MRPS6, DUSP6, SLAMF8, TNFRSF1B, MSC, R negative	
AP1.Q6	0.00035	0.008809	-0.51872	-1.63047	0	202	MMP9, LMNA, VIM, CDKN1A, LAPTM5, RTN4, YWHAZ, ALDOA, IDS negative	
ELF1.Q6	0.000352	0.008809	-0.512	-1.61358	0	206	LIMS1, TYROBP, SAT1, VIM, ARRB2, AIF1, HM13, RALA, ITGB7, CTS/ negative	
AP1.Q4_01	0.000349	0.008809	-0.5116	-1.61056	0	203	MMP9, CD68, CDKN1A, PPP1R15A, FABP4, SQSTM1, CSF1R, H3F3B negative	
TCANNTGAY.SREBP1_	0.000452	0.010535	-0.47289	-1.57952	0	386	CTSD, TM4SF19, ATP6V1F, CALR, PSAP, GPNMB, RTN4, LAMTOR1, negative	
RGAGGAARY.PU1_Q6	0.000457	0.010535	-0.42754	-1.43088	0	393	MMP9, TYROBP, IL411, VIM, PLD3, RAC2, SPI1, YWHAZ, ALDOA, CY negative	
NFKAPPAB65.01	0.001017	0.021695	-0.48751	-1.51962	2	187	MMP9, IER3, NFKBIA, SLAMF8, TNFRSF1B, CSF1R, YWHAZ, MSC, IC negative	
LXR.Q3	0.00106	0.021815	-0.65715	-1.73161	3	57	MAFB, NFKBIA, SGK1, FKBP2, APOC1, ABCA1, BLVRB, AMD1, MITF negative	
CREL.01	0.001398	0.026582	-0.4735	-1.49185	3	205	MMP9, IER3, NFKBIA, DUSP6, SLAMF8, TNFRSF1B, CSF1R, YWHAZ, negative	
AP1.Q6_01	0.001395	0.026582	-0.46803	-1.46954	3	200	LMNA, SGK1, PPP1R15A, FABP4, SDCBP, VAT1, SQSTM1, LAPTM5, negative	
TGANNYRGCA.TCF11N	0.001431	0.026582	-0.46304	-1.46874	3	216	MMP9, EIF5, SQSTM1, TPM3, RHOG, ARF4, RNF13, VASP, C8orf82, negative	
CEBP.C	0.001889	0.033994	-0.51299	-1.5463	5	141	SAT1, NFKBIA, PTPN12, H3F3B, ALDOA, CEBPB, RAB1A, FOXO3, GF negative	
BACH1.01	0.002099	0.036631	-0.46449	-1.46004	5	202	HMGAA1, LMNA, SGK1, CDKN1A, PPP1R15A, FABP4, SQSTM1, EMP3 negative	
AP1.C	0.002449	0.041493	-0.45856	-1.4439	6	204	MMP9, LMNA, CD68, PPP1R15A, FABP4, SDCBP, VAT1, LAPTM5, ID negative	
CCCNNGGAR.OLF1_C	0.002967	0.046195	-0.43587	-1.39853	7	246	IL411, ATF5, MTSS1, NFKBIA, LASP1, FKBP2, VAT1, ZEB2, H3F3B, HC negative	
AP1.01	0.004175	0.06329	-0.45533	-1.43094	11	201	LMNA, CDKN1A, VAT1, SQSTM1, EMP3, ALDOA, IDS, PHLDA2, AKIF negative	
NFKB.Q6	0.004423	0.063825	-0.45705	-1.42695	12	189	IL411, ATF5, NFKBIA, LASP1, SLAMF8, MSC, RBPJ, ICAM1, CCDC107, negative	
NRF2.Q4	0.004432	0.063825	-0.45156	-1.41242	12	191	FRMD4A, SQSTM1, H3F3B, ALDOA, IDS, PHLDA2, BLVRB, ENO1, YM negative	
E2F.Q3_01	0.000139	0.005046	0.659464	1.909187	0	208	PCLAF, STMN1, H2AFZ, HMGN2, RPS19, RANBP1, PTMA, PRKDC, C positive	
E2F.03	0.000138	0.005046	0.650807	1.893047	0	219	PCLAF, STMN1, H2AFZ, HMGN2, RPS20, RANBP1, PTMA, PRKDC, C positive	
E2F1.Q4_01	0.00014	0.005046	0.646185	1.865393	0	203	PCLAF, STMN1, H2AFZ, HMGN2, RPS19, RANBP1, PTMA, PRKDC, C positive	
E2F.06_01	0.000139	0.005046	0.642509	1.863053	0	212	PCLAF, STMN1, H2AFZ, HMGN2, RPS19, RANBP1, PTMA, PRKDC, D positive	
E2F.Q4_01	0.000139	0.005046	0.627736	1.818187	0	211	PCLAF, STMN1, H2AFZ, HMGN2, RPS19, RANBP1, PTMA, PRKDC, C positive	
E2F.Q3	0.00014	0.005046	0.620013	1.787787	0	200	STMN1, H2AFZ, HMGN2, RANBP1, PRKDC, CBX5, ATAD2, ARHGAP positive	
E2F.Q6	0.000139	0.005046	0.613171	1.774788	0	207	PCLAF, STMN1, H2AFZ, HMGN2, RANBP1, PRKDC, CBX5, ATAD2, C positive	
E2F.Q4	0.000139	0.005046	0.610694	1.768827	0	211	PCLAF, STMN1, H2AFZ, HMGN2, RANBP1, PRKDC, CBX5, ATAD2, C positive	
E2F1.Q6_01	0.000139	0.005046	0.581008	1.686153	0	215	STMN1, HMGN2, RPS19, RANBP1, PTMA, RPL18, PRKDC, CBX5, AT/ positive	
E2F1.Q6	0.000139	0.005046	0.579747	1.678268	0	209	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PRKDC, ATAD2, C positive	
E2F1DP1.01	0.000139	0.005046	0.568415	1.645244	0	207	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PTMA, PRKDC, AT positive	
E2F1DP2.01	0.000139	0.005046	0.568415	1.645244	0	207	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PTMA, PRKDC, AT positive	
E2F4DP2.01	0.000139	0.005046	0.568415	1.645244	0	207	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PTMA, PRKDC, AT positive	
E2F.02	0.000139	0.005046	0.568048	1.644182	0	207	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PTMA, PRKDC, AT positive	
E2F4DP1.01	0.000139	0.005046	0.562316	1.628381	0	210	PCLAF, STMN1, H2AFZ, H2AFV, RPS20, RANBP1, PRKDC, ATAD2, A positive	
E2F1DP1R8.01	0.00014	0.005046	0.561019	1.620205	0	204	PCLAF, STMN1, H2AFZ, HMGN2, CBX5, ATAD2, CDT1, ARHGAP11A, positive	
E2F1.Q4	0.000277	0.008414	0.554186	1.611179	1	218	STMN1, HMGN2, HMGN2, ZFP36L2, RANBP1, PRKDC, CBX5, SH3KBP1 positive	
E2F1.Q3	0.000278	0.008414	0.551892	1.601657	1	215	PCLAF, STMN1, H2AFZ, HMGN2, ATAD2, ARHGAP11A, DNMT1, SLC positive	
USF2.Q6	0.00056	0.012404	0.539697	1.558627	3	204	STMN1, COMMD3, CDKN2C, HMGN2, REEP3, H3F3A, RPL22, CBX5, positive	
SMAD.Q6	0.002545	0.041876	0.519518	1.491178	17	190	STMN1, CKS1B, BMP4, RPS14, CBX5, RPL7, SH3KBP1, NRGN, HNRN positive	
SGCGSSAAA.E2F1DP2	0.002916	0.046195	0.545308	1.522242	19	149	PCLAF, H2AFZ, H2AFV, RPS20, RANBP1, PTMA, PRKDC, ATAD2, DN positive	
CDP.02	0.004721	0.06633	0.614792	1.569087	29	73	MEF2C, RPA3, PHACTR3, BHLHE22, PTMA, GNAQ, OTX1, CDC47, M positive	
PAX2.01	0.005566	0.074992	0.678893	1.597401	33	43	HIST1H4C, HOXA10, ACTN4, MBNL1, JMJD1C, ITGB3BP, HDAC9, LRI positive	
E2F.01	0.005648	0.074992	0.647435	1.58611	34	56	SMC4, RANBP1, PRKDC, DNMT1, RMI2, AK2, TOPBP1, MCM3, MCM positive	
OCT1.02	0.005729	0.074992	0.542596	1.496063	38	132	CDKN2C, HMG2B, RPS19, HOXA10, CPNE1, MBNL1, ZNF521, ASPM positive	
COMP1.01	0.006881	0.086168	0.594609	1.536162	43	80	PCLAF, HMG81, SKA2, HOXA10, CDK6, MBNL1, ZNF281, PHF6, TLE4 positive	
CRX.Q4	0.006754	0.086168	0.508209	1.442548	46	172	CDKN2C, HMGN2, ZFP36L2, SATB1, RPA3, BM1, HIST1H4C, BHLHE positive	
E2F.Q2	0.007399	0.090675	0.516606	1.446676	50	152	STMN1, COMMD3, HMGN2, BM1, UQCRH, HLX, GFI1, ZNF281, TPR positive	
MEIS1AH0XA9.01	0.007979	0.095745	0.586922	1.521992	50	82	CDKN2C, SKA2, SATB1, PDLIM1, HLX, ONECUT2, PRR11, ADGRL2, LI positive	

**Table II Transcription Factor Targeting analysis of DGE contrasting PIC/Bystander and Provirus cells.** TFT analysis (GSEA with the fgsea R package and the C3 collection from msig) suggests that at least 3 transcription factor families control the transition from PIC/Bystander transcriptomes to Provirus cluster transcriptomes. These are E2F, NFKB, and AP1 family promoter proteins. In particular, increased E2F regulated transcription appears to correspond with the transition to the production of viral proteins. The pseudo-transition from Control to PIC/Bystander is characterized by down-regulation of E2F family regulated transcripts and up-regulation of NFKB and AP1 regulated transcripts Appendix III. In comparing Provirus to PIC/Bystander transcriptomes, E2F family promoted transcripts are up-regulated, while NFKB and AP1 transcription products are down-regulated. Comparing Provirus to

Control transcriptomes shows that overall Proviruses cells have increased E2F regulated transcripts and decreased NFkB transcripts (with no significant change detected in AP1 regulation).



**Figure 15. Western blot analysis for phospho- Rb or IκB in protein from mCherry negative versus mCherry positive cells.** Cells infected with DHIV3-mCherry were purified by FACS sorting based on their expression of mCherry fluorescence. Lane 1) Protein from Control cells; Lane 2) Protein from PIC/Bystander cells; Lane 3) Protein from Provirus cells. Phospho-Rb (Phospho-T821 Rb antibody) was used to quantify Rb pocket phosphorylation, anti-Rb control antibody was used to quantify Rb protein levels relative to actin (visualized with beta-actin antibody). mCherry protein confirmed with anti-mCherry antibody used in Figure 7. PIC/Bystander cells show the lowest level of Rb phosphorylation, Provirus show the highest, in close agreement with Transcription Factor Targeting results. Panel B, Lane 1) Protein from Control cells; Lane 2) Protein from PIC/Bystander cells; Lane 3) Protein from Provirus cells. Phospho-IκB S32 antibody was used to quantify activated IκB. Control cells show the lowest level of IκB phosphorylation, no difference was detectable between Provirus and PIC Cluster cells.



**Figure 16 Sequential infection of THP-1 cells with DHIV3-mCherry followed 24 hrs later with GFP DHIV3.** Abscissa, mCherry signal, Ordinate, GFP signal. Provirus cluster, mCherry positive, cells were 2 to 5 times more likely to make HIV-1 encoded GFP protein upon second infection than PIC/Bystander cells upon second infection. Panel **A**) time equal 0 hrs; addition of DHIV3-mCherry. Panel **B**) time equal 24 hrs; addition of DHIV3-GFP. Panel **C**) time equals 48 hrs after DHIV3-mCherry addition, 24 hrs after DHIV3-GFP addition. Panel **D**) time equals 72 hrs after DHIV3-mCherry addition, 48 hrs after DHIV3-GFP addition. Percentage of mCherry cells also producing GFP, compared to cells producing mCherry only, is always 2 to 5 times higher than the percentage of cells making only GFP, compared to those cells not producing either mCherry or GFP.