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Title	Integrase-independent HIV-1 infection is augmented under conditions of DNA damage and produces a viral reservoir.
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3	DNA damage and produces a viral reservoir
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Integrase-independent HIV-1 infection is augmented under conditions of

17 Abstract

18	HIV-1 possesses a viral protein, integrase (IN), which is necessary
19	for its efficient integration in target cells. However, it has been reported
20	that an IN-defective HIV strain is still capable of integration. Here, we
21	assessed the ability of wild type (WT) HIV-1 to establish infection in the
22	presence of IN inhibitors. We observed a low, yet clear infection of
23	inhibitor-incubated cells infected with WT HIV which was identical to cells
24	infected with IN-deficient HIV, D64A. Furthermore, the IN-independent
25	integration could be enhanced by the pretreatment of cells with
26	DNA-damaging agents suggesting that integration is mediated by a DNA
27	repair system. Moreover, significantly faster viral replication kinetics with
28	augmented viral DNA integration was observed after infection in irradiated
29	cells treated with IN inhibitor compared to nonirradiated cells. Altogether,
30	our results suggest that HIV DNA has integration potential in the presence
31	of an IN inhibitor and may serve as a virus reservoir.

- 33 Keywords
- 34 HIV-1, integration, integrase inhibitor, provirus, HIV reservoir, DNA repair

Retroelements, 38such long terminal repeat as (LTR)-retrotoransposons, 39non-LTR-retrotransposons, and retroviruses, 40 insert their reverse transcribed cDNA into the host chromosome during viral replication. To carry out efficient replication, the retrovirus family has robust 4142integration machinery consisting of the retrovirus integrase (IN), which 43executes the insertion of viral cDNA into the genome of host cells. The IN protein consists of three distinct domains (Craigie, 2001): the N-terminal 4445domain contains an HHCC motif; the catalytic core domain in the center of 46 INs possesses a DDE motif that mediates catalysis; and the C-terminal 47domain of IN has little sequence conservation yet possesses nonspecific DNA binding activity. 48

It has been reported that human immunodeficiency virus type 1
(HIV-1) cDNA is preferentially inserted into the gene coding region of the
host genome (Schroder et al., 2002). The host lens epithelium-derived growth

52factor/p75 (LEDGF/p75) protein directly binds IN (Llano et al., 2004; Maertens et al., 2003) and recruits a pre-integration complex consisting of 53viral cDNA, host proteins and viral proteins to the gene coding region 5455(Cherepanov et al., 2005; Engelman and Cherepanov, 2008; Shun et al., 2007). The genome position of inserted HIV-1 provirus is thought to 5657determine the magnitude of viral gene expression. In fact, transcription of 58provirus integrated into gene coding regions is extremely high and produces 59large amounts of viral particles (Wang et al., 2007). In contrast, the transcription of proviruses inserted outside of the gene coding region is 60 61 relatively dormant and has a potential for persistent and latent infection (Brady et al., 2009; Skupsky et al., 2010). 62

INs dramatically increase the efficiency of viral nucleic acid insertion into the host DNA, however, other mechanisms for exogenous nucleic acid incorporation may also exist. Accumulating lines of evidence indicate that foreign nucleic acids, not related retroelements, can be inserted into the genome of host cells. For instance, transfected plasmid DNA (Suzuki

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68	et al., 2010), a DNA virus genome (Dall et al., 2008), mitochondorial DNA
69	fragments (Nitz et al., 2004) and borna virus cDNA (Horie et al., 2010) have
70	been reported to integrate into host chromatin DNA despite the absence of
71	an IN enzyme. The host DNA repair system which involves homologous
72	recombination or non-homologous end joining appears to be involved in the
73	insertion (Horie and Tomonaga, 2011).
74	In this study, we first tested the integration of HIV with a mutated
75	IN lacking catalytic activity. We could detect a low but significant amount of
76	integrated HIV cDNA. Because integration is essential for HIV-1 replication,
77	IN inhibitors have been developed and used as an antiviral therapy for HIV
78	(Summa et al., 2008). We tested the illegitimate integration of WT HIV in
79	the presence of IN inhibitors. A low frequency of integration was observed in
80	the presence of an IN inhibitor which capable of producing infectious virus
81	particles, and the level of integration was clearly enhanced under
82	DNA-damaged condition. Our results suggest that retroviral cDNA is
83	inserted into the host chromosome through host DNA repair machinery via

84 an IN-independent pathway and serves as a virus reservoir.

85 Results

86

- 87 Generation of provirus and stable expression of HIV-1 in the absence of IN
 88 activity
- 89

We initially used a vesicular stomatitis virus (VSV)-G pseudotyped 90 91EGFP-expressing HIV vector packaged with a catalytically inactive HIV-1 92IN, which contains a D64V mutation. Jurkat cells were infected with varying 93 amounts of the D64V mutant virus corresponding to the amount of p24CA 94used at a multiplicity of infection (MOI) of 1, 5 and 10 of wt virus. Cells were 95also infected with IN-proficient WT virus at an MOI ranging from 1-10 in the absence or presence of IN inhibitors, Elvitegravir (Elv) or Raltegravir (Ral). 96 97 To ensure the removal of unintegrated HIV-1 DNA associated with cell 98 division, the cells were further cultured for 2 weeks and the levels of unintegrated, 2LTR, and integrated HIV-1 DNA were assessed. As shown 99 100 Fig.1A, the increased integrated but not unintegrated viral DNA was 101 detected with increasing MOI in WT+Elv- and D64V-infected cells. Based on 102 this result, we calculated the efficiency of integration during an IN-deficient condition to be 0.1-0.2% of WT integration (Fig.1B). We further assessed 103 green fluorescent protein (GFP) transduction efficiency by flow cytometry 104 105analysis and observed a dose dependent GFP transduction under IN-deficient conditions, in WT+Elv or WT+Ral, and D64V-infected cells 106 107 (Fig.1C). Up to 2.3% GFP positive cells were detected in D64V mutant 108 infected cells. The copy number of integrated viral DNA highly correlated with the transduction efficiency (R=0.9019 > 0.590; α =0.01) (Fig.1D). These 109 110results indicated that the transduction was from the integrated form of HIV-1 DNA. Therefore, we assumed that the level of GFP expression 2 weeks 111 112after infection using this HIV vector was representative of the level of integrated DNA. 113 114 It shown introduction has been that the of a DNA double-stranded break 115stimulate in target gene can a

116 retrotransposition of LINE-1, gene targeting and genome rearrangement

117	(Francis and Richardson, 2007; Morrish et al., 2002; Richardson and Jasin,
118	2000). To test whether DNA damage augments integration activity of HIV-1
119	lacking IN activity, cells were exposed to various doses of gamma irradiation
120	in order to induce DNA double-strand breaks (DSB) before virus infection
121	and then flow cytometry analysis was performed two weeks post infection
122	(Fig. 1E). The percentage of GFP positive cells was clearly augmented and
123	correlated with increasing doses of gamma irradiation. Furthermore, we also
124	examined the effect of chemical-induced DNA damage by hydrogen peroxide
125	in IN-deficient HIV-1 integration. Significant enhancement of GFP
126	transduction of IN-deficient HIV was observed in cells treated with hydrogen
127	peroxide in a dose dependent manner (Fig. 1F). Finally, we directly analyzed
128	the copy number of integrated HIV-1 DNA in the cells used in Fig. 1E and F.
129	(Fig. 1G and H, respectively). The efficiency of integration under IN-deficient
130	conditions was augmented from 0.1% to 1.7% depending on the dose of DNA $$
131	damage. These results suggested that the induction of DNA damage in target
132	cells enhanced the efficiency of retroviral IN-independent integration.

134	the IN-independent integration.
135	
136	Attenuation of HIV-1 gene expression from proviruses established through
137	an IN-independent pathway
138	
139	We showed that HIV-1 DNA was inserted into the host chromosome
140	without IN activity. To examine the LTR promoter activity and level of gene
141	expression from IN-independent proviruses, VSV-G pseudotyped LTIG
142	(LTR-Tat-IRES-GFP) vector was used. The HIV tat protein, an accessory
143	protein responsible for regulating HIV transcription, is expressed under the
144	regulation of an LTR promoter and the transcriptional level can be
145	monitored by GFP expression. As expected, a parallel transduction of Jurkat
146	cells with GFP-expressing WT virus in the presence of Elv and D64V mutant
147	virus in increasing doses was observed (Fig. 2A and B). However, the mean
148	fluorescence intensity (MFI) was significantly lower in cells infected with

Moreover these findings suggest that HIV uses a host DNA repair system for

133

149	LTIG virus in the presence of Elv and D64V mutant compared with that of
150	WT virus without Elv (Fig. 2A and C). Furthermore, we isolated 17, 10 and
151	20 cell clones from WT virus infected cells without IN inhibitor, WT virus
152	infected cells with inhibitor, and D64V virus infected cells, respectively. The
153	level of GFP expression in each clone was analyzed by flow cytometry as
154	shown in Fig.2D. The isolated cell clones were divided into two groups,
155	IN-dependent (WT) and IN-indendent (WT+Elv and D64V) transduction.
156	Then, the frequency of distribution based on the MFI of GFP expression is
157	shown in Fig. 2E. Chi-square distribution was assessed and statistically
158	significant difference between IN-dependent and independent groups was
159	observed (_X 2=10.927>9.488; four-degree-of-freedom, P=0.05).
1.0.0	

Given that the chromatin environment near the provirus is known to affect the level of viral gene expression, we attempted to address this possibility by analyzing the integration sites of proviruses (Table 1). In WT-infected cells, 84% of the integration sites were detected in gene coding regions as shown RefSeq, while only 68% of the events were detected in gene

165	coding regions under IN-deficient condition. Although, we could not find
166	statistically significant differences ($P=0.098$) under these parameters, if we
167	analyzed the frequency of viral DNA insertion in respect to the presence of
168	repeat sequences, then significant differences were observed (2% vs. 19%;
169	P=0.0048). Furthermore, the deletion and insertion of nucleotides in the
170	junction of LTR-genomic DNA was a frequent feature in the IN-deficient
171	mutant as shown in a previous report (6% vs. 27%; P =0.0039) (Gaur and
172	Leavitt, 1998). These results suggest that the HIV integration pattern is
173	modified under IN-deficient conditions and it may cause reduced promoter
174	activity.
175	
176	Formation of an intact HIV-1 reservoir under IN-deficient conditions
177	
178	Here, we showed a lower level of viral gene expression from the
179	provirus generated through an IN-independent pathway. To test the ability
180	of replication-competent HIV-1 to produce infectious virus under

181 IN-suppressed conditions, we performed the experiment depicted in Fig. 3A. 182Irradiated or untreated Jurkat cells were infected with replication competent HIV-1 NL4-3 in the presence of an IN inhibitor. Three days after 183 184 infection, the IN inhibitor and free viral particles were removed with 185 extensive washing, followed by the addition of fresh Jurkat cells (as 186 indicated day 0). p24CA viral antigen in culture supernatant was monitored. 187 Under nonirradiated culture conditions, the peak of p24CA was observed at 188 days 9 and 11 after the removal of IN inhibitor. In contrast, faster viral replication was observed in the irradiated culture. The peak of viral 189 replication was observed at 5 days post infection (dpi) (Fig. 3B). The amount 190 191 of integrated and 2LTR proviral DNA at day 0 was performed by quantitative PCR and we found that the irradiated culture contained a 192193 fivefold higher amount of integrated provirus than the nonirradiated culture. 194On the other hand, there was no significant difference in the copy number of 2LTR cDNA between irradiated and non-irradiated cultures, suggesting that 195DNA damage before viral infection increased the HIV cDNA insertion under 196

197 IN-deficient condition and it promoted faster viral replication.

199 Discussion

201	Previously, Gaur <i>et al.</i> reported that mutant HIV lacking IN activity
202	due to mutations in a highly conserved DDE motif of IN is able to integrate
203	into the host chromosome (Gaur and Leavitt, 1998). Here, we showed that
204	not only an IN-deficient virus, but also WT virus in the presence of an IN
205	inhibitor is integrated into the host chromosome. The efficiency of HIV
206	integration when using an IN antagonist was only 0.1-0.2% of IN mediated
207	integration, indicating that the IN-independent integration pathway may be
208	only a minor pathway or accidental event <i>in vivo</i> . However, we showed that
209	stress inducing DNA damage enhances IN-independent infection of cells by
210	HIV and has the potential to serve as a virus reservoir, thus suggesting that
211	it may play a role in disease progression. For instance, although combination
212	antiretroviral therapy (cART) has reduced the pathogenesis of AIDS-related
213	malignancies, there has been an increase of HIV- positive patients with
214	non-AIDS-defining malignancies such as Hodgkin's lymphoma, invasive anal

215	carcinoma, lung cancer, skin cancers, and hepatocellular carcinoma (Spano
216	et al., 2008). Radiotherapy is a standard treatment procedure for many
217	individuals with cancer and HIV, even though it may adversely affect HIV
218	disease status and CD4 counts (Housri et al., 2010). Our results suggest that
219	patients receiving simultaneous medical treatment for cancer in the form of
220	radiation and anticancer drug therapy may be at a higher risk for DNA
221	repair mediated integration of proviral cDNA. In addition, it is conceivable
222	that even mental stress-induced radical oxidents may augment
223	IN-independent infection (Adachi et al., 1993; Morimoto et al., 2008).
224	Another interesting point of our study is that the level of gene expression
225	under IN-deficient conditions is reduced when compared with functional IN
226	infection. Altogether, the IN-independent integration enhanced by the stress
227	may lead to latent infection <i>in vivo</i> .
228	The mechanism of IN-independent integration remains to be
229	elucidated. Previously, Gaur et al. sequenced the host-virus junction and
230	showed that IN-independently integrated provirus do not have a duplicate

2315-bp repeat of host cell DNA which is characteristically generated by the 232staggered cleavage of host DNA during the strand transfer reaction of HIV 233(Gaur and Leavitt, 1998). They also demonstrated that the integrated DNA 234of IN defective virus includes the deletion of host DNA, LTR, and the 235insertion of unknown sequences between the virus-host DNA junction. Based 236on these results, they suggested that the integration of HIV lacking IN 237activity may be catalyzed by the host DNA repair system (Gaur and Leavitt, 2381998). In line with their findings, our results also demonstrate a significant 239increase in the deletion and insertion of nucleotides at virus-host DNA 240junctions in cells infected with an IN-deficient virus (WT: 6% vs. 241IN-deficient: 27%, P=0.0039). Furthermore, induced DNA damage in target 242cells before virus infection increased the level of integration (Fig. 1G and H). 243These findings support the idea that HIV DNA is inserted into a DNA break point by the host DNA repair system, possibly by non-homologous end 244245joining and/or homologous recombination.

246

It has been reported that the provirus integration site determines

247the level of viral gene expression (Wang et al., 2007). The host protein 248LEDGF/p75 promotes HIV integration into active gene coding regions (Shun 249et al., 2007). However, silencing of the active provirus integrated at gene 250coding regions is challenging, making it unclear how latently infected cells are established. In this paper, we demonstrated that the level of gene 251expression is reduced depending on distinct integration pathways. To 252253understand the mechanism of gene silencing, we analyzed integration sites 254of HIV-1 proviruses. Unexpectedly, the integration site analysis focused on gene coding sequences did not show significant differences between the 255256targeted frequency of integration under IN-deficient and conventional 257conditions. However, HIV-1 cDNAs were frequently inserted into minisatellite-like repeat sequences of genomic DNA in IN-deficient 258conditions (P=0.0048). Minisatellites consist of 10-100 bp repeat sequences 259260and is observed near cis-acting meiotic double-strand break hotspots (Richard and Paques, 2000). If the IN-independent integration is carried out 261262by the DNA repair pathway, then double-strand break hotspots may be the target for the insertion. Moreover, an unknown determinant for these
hotspots, such as the chromatin environment, may cause reduced gene
expression.

266In our study, the integration frequency of provirus into gene coding 267regions was 68% under an IN-deficient condition and it was much higher than predicted random integration (33%). Hence, this result suggests that 268269 there is an integration preference into gene coding regions even during 270IN-independent integration. One plausible explanation for this may be 271attributed to the sensitivity of the host chromosome to DNA breaks at gene 272coding regions. If the retroviral DNA is inserted at a DNA break point under 273IN-independent conditions, our results might reflect a high frequency of 274DNA breaks that occur at gene coding regions in the host chromosome. 275Another possibility is that LEDGF/p75 may promote viral cDNA tethering to 276the gene coding region even under IN-deficient conditions the same as it would if functional IN was present. We used a lentivirus vector that has a 277278mutation in the catalytic domain of IN. It has been reported that the

279	mutation at D64 to alanine in IN does not inhibit the protein binding with
280	LEDGF/p75 (Cherepanov et al., 2005). In addition, the binding interface of
281	IN and LEDGF/p75 does not overlap with the active site of IN inhibitors
282	(Hare et al., 2010; Michel et al., 2009). Therefore, it is not surprising that
283	retroviral DNA preferentially integrated into gene coding regions even when
284	IN is devoid of function.
285	In sum, our findings serve as a caveat for an alternative infection
286	route that HIV can take during DNA damage to bypass a drug therapy
287	involving IN inhibitors. The analysis of integrated provirus obtained from
288	HIV-positive patients after radiotherapy will shed light on the practical risk
289	of the alternative infection pathway of HIV <i>in vivo</i> .

291 Materials and Methods

292

293 Virus preparation

294

295Human embryonic kidney (HEK) 293T cells were used for virus preparation. HEK293T cells were transfected by the calcium phosphate 296297 method as described previously (Kawano et al., 2004). The culture 298 supernatants at 48 hr post-transfection were centrifuged and filtrated. To 299prepare HIV-1 NL4-3, cells were transfected pNL4-3 and the 50% tissue 300 culture infective dose (TCID₅₀) was calculated (Kawano et al., 1997; Sato et 301 al., 2008). To prepare the EGFP-expressing HIV vector, pCS-CDF-EG-Pre 302 was used. pCS-CDF-EG-Pre was constructed by exchanging the CMV 303 promoter of pCS-CDF-CG-Pre (Miyoshi et al., 1998) with human elongation 304 factor 1a (EF) promoter. The pCS-CDF-EG-Pre was transfected together with packaging plasmids pMD.G, pMDLg/pRRE and pRSV Rev (Kawano et 305 al., 2004). To prepare VSV-G pseudotyped WT LTIG vector, pEV731, kindly 306

307	provided by Dr. Eric Verdin (Jordan et al., 2003), pMD.G, pMDLg/pRRE and
308	pRSV Rev were cotransfected. To prepare IN-deficient D64V mutant virus,
309	pMDKg/pRRE/D64V, kindly provided by Dr. Ikawa (Okada et al., 2009), were
310	used instead of pMDLg/pRRE. The infectious dose of GFP expressing vectors
311	was assessed as follows. Two hundreds thousand of Jurkat cells were
312	infected with serial volumes of stock virus in 1 ml culture. At 3 days after
313	infection, GFP positive cells were analyzed by flow cytometry and the
314	infectious dose were calculated based on the input volume of virus counted
315	approximate 25% of GFP positive.
316	
317	ELISA
318	
319	To quantify the viral antigen p24CA in virus solutions, an HIV-1 p24
320	antigen enzyme linked immunosorbent assay (ELISA) kit (ZetroMetrix
321	Buffalo, NY) was used.

323 Cell culture

325	HEK293T cells were maintained in Dulbecco's Modified Eagle
326	Medium (DMEM) containing 10% fetal calf serum (FCS), 100 U/ml penicillin
327	and 100 g/ml streptomycin. Jurkat cells were maintained in RPMI 1640
328	medium containing 10% FCS, 100 U/ml penicillin and 100 g/ml
329	streptomycin.
330	
331	Pseudotyped virus infection and flow cytometry analaysis
332	
333	To infect with pseudotyped EGFP-expressing HIV vector or LTIG
334	virus, Jurkat cells (2 x 10^5 cells) were infected with a pseudotyped virus
335	solution as corresponded as the infectious dose of WT virus at indicated MOI.
336	Under IN-deficient conditions, 1 μ M Ral (NIH, Bethesda, MD) or 100 nM Elv
337	(Selleck Chemicals, Houston, TX) were added in the culture medium. Two
338	weeks after infection, the percentage of GFP positive cells were measured by

339	flow cytometry. Cells were suspended in phosphate-buffered saline (PBS)						
340	containing 1% formamide. Flow cytometry was performed with a						
341	FACSCalibur (BD Biosciences), and data were analyzed using CellQuest						
342	software (BD Biosciences).						
343							
344	Cell cloning						
345							
346	GFP+ Jurkat cells were sorted from the bulk culture 2 weeks after						
347	pseudotyped LTIG virus infection by FACSAria (BD Biosciences) and cell						
348	cloning was carried out by limiting dilution method.						
349							
350	Induction of DNA damage in target cells						
351	To induce DNA damages with ionizing radiation, Jurkat cells (2 ${\rm x}$						
352	10^5 cells/ml) were exposed to appropriate doses (5-10 Gy) of gamma						
353	irradiation at room temperature. Gamma irradiation was performed using a						
354	Faxitron RX-650 (Faxitron bioptcs, Lincolnshire, IL). To induce DNA damage						

355	with reactive oxygen species, Jurkat cells (2 x 10^5 cells/ml) were incubated
356	with medium containing 10, 50 or 100 μM of hydrogen peroxide at 37°C for 6
357	hr. The DNA damage-induced cells were immediately used for virus infection
358	after washing with medium once.

360 Analyses of HIV-1 replication kinetics of IN-independent integrated
 361 proviruses

362

The irradiated or non-treated Jurkat cells (4 x 10^4 cells) were 363 364incubated for 2 hr in HIV-1 NL4-3 solution containing 40 ng of p24CA 365 (TCID₅₀, 984419/ml) at MOI of 1 with 100 nM Elv or 1 µM Ral. After 366 extensive washing (twice with PBS, once with 5% tripsin/EDTA at 37 °C for 5 367 min, and twice with 10% FCS RPMI), the cells were resuspended and 368 cultured in 200 µl of 10% RPMI containing an appropriate concentration of IN inhibitor. After 3 dpi, the cells were extensively washed as mentioned 369 above. Then, the cells were co-cultured with fresh Jurkat cells (4 x 10^4 cells) 370

371	in the absence of IN inhibitor. The culture medium was harvested at the								
372	indicated time points and the level of p24CA antigen was measured by								
373	ELISA.								
374									
375	Quantitative analysis of retroviral DNA								
376									
377	The amount of proviral, 2LTR, and full-length forms of HIV-1 DNA								
378	was quantified by real-time PCR and the copy number of HIV-1 DNA was								
379	normalized by 6-actin as previously described (Suzuki et al., 2003).								
380	Unintegrated linear DNA was calculated by subtracting the copy number of								
381	provirus and 2LTR DNA from that of full-length viral DNA.								
382									
383	Integration site analyses								
384									
385	The integration sites were determined by the linker ligation method								
386	described previously (Ciuffi et al., 2009). Briefly, genomic DNA was extracted								

387	from the GFP positive population using a DNeasy column (QIAGEN) and
388	digested with AvrII, NheI and XbaI. The digested genomic DNA was ligated
389	with a linker adapter and then used to perform nested PCR amplification.
390	The PCR products were cloned into a pGEM-T (Promega, Madison, WI)
391	vector and sequenced using an M13 primer.
392	
393	Statistical analysis
394	
395	The student's t test was used to determine statistical significance. P
396	values of <0.05, <0.01 and <0.001 were considered significant. The
397	chi-square test was also used to determine statistical significance from the
398	frequency of distribution.
399	

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411	

413 **References**

- 415 Adachi, S., Kawamura, K., Takemoto, K., 1993. Oxidative damage of nuclear
- 416 DNA in liver of rats exposed to psychological stress. Cancer Res. 53,417 4153-4155.
- 418 Brady, T., Agosto, L.M., Malani, N., Berry, C.C., O'Doherty, U., Bushman, F.,
- 419 2009. HIV integration site distributions in resting and activated CD4+ T
 420 cells infected in culture. AIDS 23, 1461-1471.
- 421 Cherepanov, P., Sun, Z.Y., Rahman, S., Maertens, G., Wagner, G., Engelman,
- 422 A., 2005. Solution structure of the HIV-1 integrase-binding domain in
- 423 LEDGF/p75. Nat Struct Mol Biol 12, 526-532.
- 424 Ciuffi, A., Ronen, K., Brady, T., Malani, N., Wang, G., Berry, C.C., Bushman,
- F.D., 2009. Methods for integration site distribution analyses in animal cell
 genomes. Methods 47, 261-268.
- 427 Craigie, R., 2001. HIV integrase, a brief overview from chemistry to 428 therapeutics. J. Biol. Chem. 276, 23213-23216.
- 429 Dall, K.L., Scarpini, C.G., Roberts, I., Winder, D.M., Stanley, M.A.,
- 430 Muralidhar, B., Herdman, M.T., Pett, M.R., Coleman, N., 2008.
- 431 Characterization of naturally occurring HPV16 integration sites isolated
 432 from cervical keratinocytes under noncompetitive conditions. Cancer Res. 68,
 433 8249-8259.
- 434 Engelman, A., Cherepanov, P., 2008. The lentiviral integrase binding protein
- 435 LEDGF/p75 and HIV-1 replication. PLoS Pathog 4, e1000046.
- 436 Francis, R., Richardson, C., 2007. Multipotent hematopoietic cells
 437 susceptible to alternative double-strand break repair pathways that promote
- 438 genome rearrangements. Genes Dev. 21, 1064-1074.
- 439 Gaur, M., Leavitt, A.D., 1998. Mutations in the human immunodeficiency
- 440 virus type 1 integrase D,D(35)E motif do not eliminate provirus formation. J.
- 441 Virol. 72, 4678-4685.
- 442 Hare, S., Gupta, S.S., Valkov, E., Engelman, A., Cherepanov, P., 2010.
- 443 Retroviral intasome assembly and inhibition of DNA strand transfer. Nature

- 444 464, 232-236.
- 445 Horie, M., Honda, T., Suzuki, Y., Kobayashi, Y., Daito, T., Oshida, T., Ikuta,

446 K., Jern, P., Gojobori, T., Coffin, J.M., Tomonaga, K., 2010. Endogenous

- 447 non-retroviral RNA virus elements in mammalian genomes. Nature 463,448 84-87.
- Horie, M., Tomonaga, K., 2011. Non-retroviral fossils in vertebrate genomes.
 Viruses 3, 1836-1848.
- 451 Housri, N., Yarchoan, R., Kaushal, A., 2010. Radiotherapy for patients with
- 452 the human immunodeficiency virus: are special precautions necessary?453 Cancer 116, 273-283.
- 454 Jordan, A., Bisgrove, D., Verdin, E., 2003. HIV reproducibly establishes a
- 455 latent infection after acute infection of T cells in vitro. EMBO J. 22,456 1868-1877.
- Kawano, Y., Tanaka, Y., Misawa, N., Tanaka, R., Kira, J.I., Kimura, T.,
 Fukushi, M., Sano, K., Goto, T., Nakai, M., Kobayashi, T., Yamamoto, N.,
 Koyanagi, Y., 1997. Mutational analysis of human immunodeficiency virus
 type 1 (HIV-1) accessory genes: requirement of a site in the nef gene for
 HIV-1 replication in activated CD4+ T cells in vitro and in vivo. J. Virol. 71,
- 462 8456-8466.
- 463 Kawano, Y., Yoshida, T., Hieda, K., Aoki, J., Miyoshi, H., Koyanagi, Y., 2004.
- A lentiviral cDNA library employing lambda recombination used to clone an
 inhibitor of human immunodeficiency virus type 1-induced cell death. J.
 Virol. 78, 11352-11359.
- Llano, M., Vanegas, M., Fregoso, O., Saenz, D., Chung, S., Peretz, M.,
 Poeschla, E.M., 2004. LEDGF/p75 determines cellular trafficking of diverse
 lentiviral but not murine oncoretroviral integrase proteins and is a
 component of functional lentiviral preintegration complexes. J. Virol. 78,
 9524-9537.
- 472 Maertens, G., Cherepanov, P., Pluymers, W., Busschots, K., De Clercq, E.,
- 473 Debyser, Z., Engelborghs, Y., 2003. LEDGF/p75 is essential for nuclear and
- 474 chromosomal targeting of HIV-1 integrase in human cells. J. Biol. Chem. 278,
- 475 33528-33539.
- 476 Michel, F., Crucifix, C., Granger, F., Eiler, S., Mouscadet, J.F., Korolev, S.,

- 477 Agapkina, J., Ziganshin, R., Gottikh, M., Nazabal, A., Emiliani, S., Benarous,
- 478 R., Moras, D., Schultz, P., Ruff, M., 2009. Structural basis for HIV-1 DNA
- 479 integration in the human genome, role of the LEDGF/P75 cofactor. EMBO J.
- 480 28, 980-991.
- 481 Miyoshi, H., Blomer, U., Takahashi, M., Gage, F.H., Verma, I.M., 1998.
- 482 Development of a self-inactivating lentivirus vector. J. Virol. 72, 8150-8157.
- 483 Morimoto, K., Morikawa, M., Kimura, H., Ishii, N., Takamata, A., Hara, Y.,
- 484 Uji, M., Yoshida, K., 2008. Mental stress induces sustained elevation of blood
 485 pressure and lipid peroxidation in postmenopausal women. Life Sci. 82,
 486 99-107.
- 487 Morrish, T.A., Gilbert, N., Myers, J.S., Vincent, B.J., Stamato, T.D., Taccioli,
- 488 G.E., Batzer, M.A., Moran, J.V., 2002. DNA repair mediated by 489 endonuclease-independent LINE-1 retrotransposition. Nat. Genet. 31, 490 159-165.
- Nitz, N., Gomes, C., de Cassia Rosa, A., D'Souza-Ault, M.R., Moreno, F.,
 Lauria-Pires, L., Nascimento, R.J., Teixeira, A.R., 2004. Heritable
 integration of kDNA minicircle sequences from Trypanosoma cruzi into the
- 494 avian genome: insights into human Chagas disease. Cell 118, 175-186.
- 495 Okada, Y., Ueshin, Y., Hasuwa, H., Takumi, K., Okabe, M., Ikawa, M., 2009.
- 496 Targeted gene modification in mouse ES cells using integrase-defective497 lentiviral vectors. Genesis 47, 217-223.
- 498 Richard, G.F., Paques, F., 2000. Mini- and microsatellite expansions: the 499 recombination connection. EMBO reports 1, 122-126.
- 500 Richardson, C., Jasin, M., 2000. Coupled homologous and nonhomologous
- 501 repair of a double-strand break preserves genomic integrity in mammalian
- 502 cells. Mol. Cell. Biol. 20, 9068-9075.
- 503 Sato, K., Aoki, J., Misawa, N., Daikoku, E., Sano, K., Tanaka, Y., Koyanagi,
- 504 Y., 2008. Modulation of human immunodeficiency virus type 1 infectivity 505 through incorporation of tetraspanin proteins. J. Virol. 82, 1021-1033.
- 506 Schroder, A.R., Shinn, P., Chen, H., Berry, C., Ecker, J.R., Bushman, F., 2002.
- 507 HIV-1 integration in the human genome favors active genes and local 508 hotspots. Cell 110, 521-529.
- 509 Shun, M.C., Raghavendra, N.K., Vandegraaff, N., Daigle, J.E., Hughes, S.,

- 510 Kellam, P., Cherepanov, P., Engelman, A., 2007. LEDGF/p75 functions
- 511 downstream from preintegration complex formation to effect gene-specific
- 512 HIV-1 integration. Genes Dev. 21, 1767-1778.
- 513 Skupsky, R., Burnett, J.C., Foley, J.E., Schaffer, D.V., Arkin, A.P., 2010. HIV
- 514 promoter integration site primarily modulates transcriptional burst size
- 515 rather than frequency. PLoS Comput Biol 6.
- 516 Spano, J.P., Costagliola, D., Katlama, C., Mounier, N., Oksenhendler, E.,
- 517 Khayat, D., 2008. AIDS-related malignancies: state of the art and
- therapeutic challenges. J. Clin. Oncol. 26, 4834-4842.
- 519 Summa, V., Petrocchi, A., Bonelli, F., Crescenzi, B., Donghi, M., Ferrara, M.,
- 520 Fiore, F., Gardelli, C., Gonzalez Paz, O., Hazuda, D.J., Jones, P., Kinzel, O.,
- 521 Laufer, R., Monteagudo, E., Muraglia, E., Nizi, E., Orvieto, F., Pace, P.,
- 522 Pescatore, G., Scarpelli, R., Stillmock, K., Witmer, M.V., Rowley, M., 2008.
- 523 Discovery of raltegravir, a potent, selective orally bioavailable HIV-integrase 524 inhibitor for the treatment of HIV-AIDS infection. J. Med. Chem. 51, 525 5843-5855.
- 526 Suzuki, K., Ohbayashi, F., Nikaido, I., Okuda, A., Takaki, H., Okazaki, Y.,
- 527 Mitani, K., 2010. Integration of exogenous DNA into mouse embryonic stem
- 528 cell chromosomes shows preference into genes and frequent modification at
- 529 junctions. Chromosome Res. 18, 191-201.
- 530 Suzuki, Y., Misawa, N., Sato, C., Ebina, H., Masuda, T., Yamamoto, N.,
- 531 Koyanagi, Y., 2003. Quantitative analysis of human immunodeficiency virus
- 532 type 1 DNA dynamics by real-time PCR: integration efficiency in stimulated
- and unstimulated peripheral blood mononuclear cells. Virus Genes 27,177-188.
- Wang, G.P., Ciuffi, A., Leipzig, J., Berry, C.C., Bushman, F.D., 2007. HIV
 integration site selection: analysis by massively parallel pyrosequencing
 reveals association with epigenetic modifications. Genome Res. 17,
 1186-1194.
- 539
- 540

543Fig.1. The transduction and integration efficiency of HIV under IN-deficient 544conditions. Jurkat cells were infected with a MOI of 1-10 with WT 545EGFP-expressing HIV vector in the absence or presence of 100 nM Elv or 1 µM Ral and indicated as WT, WT+Elv and WT+Ral, respectively. 546547Alternatively, the IN-deficient mutant virus (D64V) was also used. (A) Two weeks after infection, the copy number of integrated, 2LTR and unintegrated 548forms of viral DNA were analyzed by qPCR. The cell number was determined 549550quantity by qPCR detecting β -actin. (B) The efficiency of integration under 551IN-deficient conditions is shown. The values were calculated by dividing the 552integrated copy number of provirus derived from IN-deficient conditions by 553 that of WT. (C) GFP positive cells were detected by flow cytometry 2 weeks after infection. Pseudo plots of the raw data analyzed by flow cytometry are 554shown. The numbers indicated on the plot show the percentage of GFP 555556 positive cells. These results are summarized in the bar graphs depicted below.

557	(D) Coefficient of correlation. The correlation of GFP positive cells (%) and
558	HIV DNA (copies/1000 cells) are shown. Gray squares indicate WT+Elv (n=9)
559	and triangles indicate D64V (n=9). Pearson's product-moment correlation
560	coefficient was calculated from populations of WT+Elv and D64V (n=18).
561	(E-H) DNA damage enhanced IN-independent integration. Jurkat cells were
562	exposed to various doses of gamma radiation, IR (E and G) or hydrogen
563	peroxide, H_2O_2 (F and H). The cells induced with DNA breaks were infected
564	with GFP expressing HIV vector. (E and F) GFP positive cells were analyzed
565	as mentioned in Fig.1C. Pseudo plots and the summarized graph are shown.
566	(G and H) The efficiency of integration under IN-deficient conditions was
567	calculated as in 1B. The amount of integrated DNA detected in DNA
568	damage-induced cultures was divided by that obtained in the culture
569	infected with WT virus without inhibitor and under non-damaged conditions.
570	All experiments were performed in triplicate (n=3) in A, B, C, E, F and H.
571	The results of flow cytometry in C, E, and F are data from one experiment,
572	which is representative of independent experiment. The error bars in A, B, C,

573 E, F, G and H show standard deviations.

Fig. 2. HIV expression from the LTR promoter of IN-independently 575576generated provirus. Pre-irradiated Jurkat cells were infected with VSV-G pseudotyped LTIG vector. (A-C) Transduction efficiency of LTIG vector under 577 578IN-deficient conditions with either WT virus in the presence of Elv (WT+Elv) 579or D64V mutant virus (D64V). The percentage of GFP positive cells was 580analyzed by flow cytometry at 2 weeks after infection. (A) Pseudo plots of the 581raw data analyzed by flow cytometry are shown. The bottom numbers 582indicated on the plot show the percentage of GFP positive cells, while 583underlined numbers indicate the mean fluorescence intensity. The results 584are data from one experiment, which is representative of three independent 585experiments. (B) The percentage of GFP positive cells is summarized as the graph. (C) Magnitude of virus expression from LTR promoter. The MFI of the 586GFP expressing cells generated after non-irradiated (0 Gy) or irradiated (5 587 588Gy or 10 Gy) stress is shown. All experiments were performed in triplicate

589	(n=3) in B and C. The error bars in A and B show standard deviations. (D)
590	Clonal cell analysis of viral expression. The GFP positive cells were isolated
591	from cells shown C and generated clonal cell lines. The MFI of GFP in each
592	cell clones were shown. The average value of population was shown by
593	horizontal line. (E) Frequency distribution table. IN-dependent ($n=17$) and
594	IN-independent (n=30) were distributed by MFI.

596 Fig. 3. The involvement of IN-independent integration in replication597 competent HIV-1 replication.

(A) The experimental procedure is shown. Irradiated (IR(+)) or untreated (IR(-)) Jurkat cells were infected with replication competent HIV-1 NL4-3 in the presence of an IN inhibitor. At 3 dpi, the IN inhibitor and free viral particles were removed with extensive wash. To expand viral replication, fresh Jurkat cells were added and the p24CA viral antigen in culture medium was monitored over the course of 15 days. (B) The p24CA viral antigen in culture medium. (C) The copy number of integrated and 2LTR

605	form of HIV DNA at day 0. All experiments were performed in triplicate
606	(n=3) in B and C. The error bars in B and C show standard deviations. Heat
607	inactivated virus treated at 60 $^\circ\mathrm{C}$ for 2 hr was used as the negative control.
608	IN inhibitors, Ral or Elv, is indicated in top (C and D). The control
609	experiment was performed in the absence of IN inhibitor and indicated as
610	WT without inhibitor.

Fig. 1







Fig. 1



Fig. 2







Table 1. Integration sites analysis

Pathway	Total Events	In RefSeq ^{*3}	(%) *4	In repeat Seq	(%) ^{*5}	Deletion or Insertion	(%) ^{*6}
IN-dependent ^{*1}	49	40	84.4	1	2.2	3	6.1
IN-independent ^{*2}	79	54	68.4	15	19.0	21	26.6
			<i>P</i> =0.098		<i>P</i> =0.0048		<i>P</i> =0.0039

*1: This result summarizes 40 sites derived from LTIG vector infected and 9 sites derived from CS-CDF-EG-Pre infected cells.

*2: This result summarizes 9 sites derived from LTIG vector infected and 75 sites derived from CS-CDF-EG-Pre infected cells. Of these sites, 6 are the results from WT+Elv and 76 six are from D64V. Then, 13 out of 79 results were derived from preirradiated culture.

*3: Integration events counted as insertion into reference sequence (=gene coding resion).

*4: A frequency of integration into RefSeq. Random integration into RefSeq is expected to be 33% of total integration sites.

*5: A frequency of integration into repeat sequence.

*6: A frequency of integration with deletion in LTR sequence or with up to 50 bp insertion in LTR-host genome junction.