Title page

The characteristics of the HIV-1 Env glycoprotein contribute to viral pathogenesis

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1 Abstract

2 The understanding of HIV-1 pathogenesis and clinical progression is incomplete 3 because of the variable contribution of host, immune and viral factors. The 4 involvement of viral factors has been investigated in extreme clinical phenotypes 5 from rapid progressors to long-term non-progressors (LTNPs). Among HIV-1 6 proteins, the envelope glycoprotein complex (Env) has concentrated many 7 studies for its important role in the immune response and in the first steps of viral 8 replication. In this study, we analyzed the contribution of 41 Envs from 24 patients 9 with different clinical progression rates and viral loads (VLs), LTNP-Elite 10 Controllers (LTNP-ECs); Viremic LTNPs (vLTNPs), and non-controller's 11 individuals contemporary to LTNPs or recent, named Old and Modern 12 progressors. We analyzed the Env expression, the fusion and cell-to-cell transfer 13 capacities as well as viral infectivity. The sequence and phylogenetic analysis of 14 Envs were also performed. In every functional characteristic, the Envs from 15 subjects with viral control (LTNP-ECs and vLTNPs) showed significant lower 16 performance compared to those from the progressor individuals (Old and 17 Modern). Regarding sequence analysis, the variable loops of the gp120 subunit 18 of the Env (i.e., V2, V4 and mainly V5) of the progressor individuals showed 19 longer and more glycosylated sequences than controller subjects. Therefore, 20 HIV-1 Envs presenting poor viral functions and shorter sequences were 21 associated with viremic control and the non-progressor clinical phenotype. 22 whereas functional Envs were associated with the lack of virological control and 23 progressor clinical phenotypes. These correlations support the central role of Env 24 genotypic and phenotypic characteristics in the in vivo HIV-1 infection and 25 pathogenesis.

26 Words: 250

IMPORTANCE

 investigated in isolates from individuals with different progression rates. In this work, we studied the properties of the envelope glycoprotein complex (Env) in individuals with different progression rates to elucidate its role in pathogenesis. We estimated the Env expression, the CD4 binding, the fusion and cell-to-cell viral transfer capacities that affect the infectivity of the viral Envs in recombinant viruses. The Envs from individuals which control viral replication and lack clinical progression (LTNP-ECs and vLTNPs) showed lower functional capacities than from subjects with clinical progression (Old and Modern). The functional infectivity and in increased length of variable loops and the number of glycosylation sites of the Env (gp120/SU). These results support the concept that viral characteristics contribute to viral infection and pathogenesis. Words: 148 44 45 46 47 48 49 50 	28	The role of the virus in the pathogenesis of HIV-1 infection has not been
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51 Introduction

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52 Pathogenesis of viral infections is the result of complex interactions between 53 host genetics, immune responses and viral factors. In human immunodeficiency virus tye 1 (HIV-1) infection and pathogenesis, the role of host (1-6), immune (6-54 55 15) and viral factors (16-20) has been widely investigated. The interactions of 56 these factors have been primarily studied in extreme clinical phenotypes like 57 rapid progressors (RPs) (21, 22) or long-term non-progressors (LTNPs), LTNP-Elite Controllers (LTNP-ECs), HIV controllers or Elite suppressors (ES) (17-19, 58 59 23, 24). 60 Due to these entangled interactions, the investigation of the role of viral proteins 61 and their specific properties in HIV-1 pathogenesis is challenging. Among the 62 viral proteins, the envelope glycoprotein complex (Env) has attracted numerous 63 studies because its essential role in the immune response and in the initial events of the HIV-1 biological cycle (25-29), i.e the binding to the cellular 64 65 receptors (29-42). The binding efficiency of the viral Env to the CD4 receptor 66 determines further steps of the viral cycle: virus-cell signaling, fusion and cell-to-67 cell virus transfer capabilities (18, 19, 43). HIV-1 Envs unable to stabilize microtubules (i.e., increasing post-transductional acetylation of Lys⁴⁰ residue in 68 69 α -tubulin), to reorganize F-actin for the delineation of pseudopod-entry virus hot 70 zones present low CD4 binding, restricted fusion and low early infection (18, 19, 71 43-45). 72 There are few reports investigating the characteristics of viral Envs from HIV

74 efficiency of viral Envs from ES individuals relative to chronically infected

individuals with different clinical characteristics. Lassen et al. studied the entry

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- viremic and chronic progressors. Envs from ES showed decreased entry
- refficacy and slower entry kinetics than those of chronic progressors (20). Our

77 group studied the CD4 binding, signaling capacity, fusogenicity of viral Envs 78 from viremic non-progressors (VNPs) that were similar to those of progressors 79 individuals (19). In previous reports, deficient viral Env glycoproteins, because 80 of poor CD4 binding, low transfer and signaling capacity (18) were identified in a 81 cluster of poor replicating viruses from a group of LTNP-ECs without clinical 82 progression for more than 20 years (17, 18). Thus, these works have stablished 83 that viral Env play an important role in the pathogenesis control in LTNPs (17-84 20, 46, 47). 85 To further investigate the role of viral Env in HIV-1 infection and pathogenesis, 86 in this work, we expanded our previous studies to viral Envs from other sets of 87 LTNP-ECs and Viremic LTNPs (vLTNPs) in comparison with groups of chronic 88 progressors. Clonal full-length env genes derived from viruses of individuals in 89 these distinct clinical groups were analyzed for expression, CD4 dependent-90 Env-mediated fusion, cell-to-cell viral transfer and infection efficiency. This 91 analysis permitted the establishment of a relationship between the initial events 92 of the viral replication cycle, mediated by the viral Env characteristics, with the 93 VL control and the clinical outcome and pathogenesis of the HIV-1 infection. 94

96 **Results**

97 Analysis of the characteristics of viral envelopes of viruses from different

98 risk groups.

99 For the investigation of the potential role of the HIV-1 Env in virological control 100 and pathogenesis, we studied the phenotypic characteristics of 41 Envs from 24 101 individuals without antiviral therapy and different VLs (Table 1). We analyzed 10 102 Envs from 6 LTNP-EC individuals with undetectable VL and infected in the late 103 80's and 90's; 10 viral clones from 6 Viremic LTNPs (vLTNPs) with VL <10,000 104 viral copies/mL and infected in the 90's. To ascertain that the characteristics of 105 the Envs from these LTNPs were not due to the sampling time, we compared 106 them with 10 Envs obtained from 6 HIV-1 individuals also infected in the same period (90's), but with high VL>10⁵ viral copies/mL and chronic infection; these 107 Env were designated Old. Finally, we studied 11 viral clones from 6 chronic 108 individuals infected between 2013-2014 with VL>10⁴ viral copies/mL and named 109 110 Modern. The main characteristics of the participants are summarized in Table 111 1.

We first analyzed the potential differences in the expression between the Env clones from the clinical groups, by measuring their cell-surface expression levels in HEK-293T cells (**Figure 1A**, *shows study scheme*, and **Figure 2**).

Although we observed a progressive augmentation of Env expression in viral clones derived from patients that do not control viremia (i.e., Old and Modern patients) compared to LTNPs (EC and Viremic), this increase did not reach statistical significance (**Figure 2**). Thus, the expression capability of the viral Envs appears to not contribute to the differences in VL and pathogenesis between groups.

121

122 Analysis of cell-to-cell membrane fusion and viral transfer capacity of viral

123 envelopes.

124 A key process for HIV Env-mediated infection is the interaction of the Env 125 complex with the CD4 receptor. When this interaction is functionally efficient. 126 viral transfer through synaptic contacts or fusion pore formation are triggered 127 during cell-to-cell or virus-to-cell contacts, repectively (18, 19, 43, 45, 48). We 128 examined the viral Env/CD4 interaction and the efficiency of subsequent 129 functions, measuring the membrane fusion capacity of the Envs (Figure 1B, 130 shows study scheme) in co-cultures between Env-expressing HEK-293T and 131 HIV-permissive target TZM-bl cells (Figure 3). To fully characterize our 132 experimental models, we used the Envs from reference HIV-1_{Bal} (CCR5-tropic) 133 and HIV-1_{NL4.3} (CXCR4-tropic) viruses (Figure 3 and 4). This fusion assay 134 yielded lower fusion values for Envs of viruses from LTNP-ECs and from 135 vLTNPs than for Old and Modern progressors, and attaining statistical 136 significance between LTNPs (EC and Viremic) and Modern Envs glycoproteins 137 (Figure 3B). 138 Next, we assayed the CD4-dependent cell-to-cell virus transfer capacity of the 139 viral envelopes. This experiment was performed co-culturing Env-expressing 140 HEK-293T cells with unstimulated primary CD4+ T lymphocytes as target cells 141 (Figure 1C, shows study scheme, and Materials and methods). In this assay, 142 we forced the formation of virological synapses between virus-effector HEK-143 293T cells expressing the different Envs together with the structural HIV Gag 144 polyprotein, and fresh primary CD4+ T cells from healthy donors (Figure 1C, 145 shows study scheme). The Envs from the LTNPs (EC and Viremic) individuals 146 displayed a lower ability to transfer viral particles to primary CD4+ T

147 lymphocytes than Envs from Old individuals and significantly lower than from

148 Modern participants (p<0.0022 between all groups) (Figure 4). These data 149 suggest that the Envs from LTNP-EC viruses had an impaired binding to the 150 cell-surface CD4 receptor and that this impairment was progressively overcome 151 in the Envs from individuals from the other groups with less control of viral 152 replication, and higher VL. 153 Thus, the phenotypic characterization of the Envs of viruses from subjects with 154 distinct progression rates confirmed that LTNP-ECs and vLTNPs presented 155 viruses with an impaired Env CD4-associated functions and a significant lower 156 fusogenic and transfer capacity, in comparison with viruses from the viremic 157 groups: These lower characteristics were also linked with the low VL detected in 158 these subjects (Figures 3 and 4). We also observed a functional improvement 159 in the viral Envs from the LTNP-EC and vLTNP individuals to those of chronic 160 Modern glycoproteins: These data support that the deficient Env fusion and 161 transfer capacities observed in the Envs of viruses from LTNP-EC and vLTNP 162 phenotypes have been enhanced in the viruses from individuals with 163 progressive infection, particularly in those of the Modern group.

164

165 Infectivity of recombinant viruses with the analyzed envelopes.

166 For the exploration of the potential consequences of these Env properties in 167 virus biology, we estimated the infectivity of recombinant viruses bearing the 168 Env from the different HIV+ phenotypic groups in TZM-bl cells (Figure 5 and 169 Figure 1D, shows study scheme). Viral Envs from the LTNP-EC group showed 170 the lowest infectivity values, whereas the Modern Envs produced the higher 171 titers. The viruses from vLTNPs displayed higher titers than LTNP-ECs but 172 lower than those from Old individuals. Recombinant viruses from individuals 173 with high VL and progressive infection (Old and Modern) have higher infectivity

174 rates than those with viral control (EC and Viremic). These results explain why

the viral properties analyzed (binding, fusion and transfer) have a significant

impact in viral infectivity with an important effect in the biology of HIV-1 and viral

177 pathogenesis.

178

179 Correlation between viral characteristics of the envelopes.

180 A significant correlation was observed between the HIV-1 Env-triggered cell-to-

181 cell transfer data, which is directly mediated by Env/CD4 binding, with Env-

182 mediated infectivity and fusogenicity (Figure 6). In all viral characteristics, the

183 Envs from subjects with virological control (EC and Viremic) showed the lower

values, whereas those from the non-controlling individuals (Old and Modern)

185 had the higher values. Therefore, HIV-1 Envs displaying poor viral functions,

186 because of the poor binding of the viral Env to the CD4, correlated with viremic

187 control and non-progressor clinical phenotypes. In contrast, functional Envs are

associated with the lack of viremic control and the progressor clinical

189 phenotypes. These statistical correlations support the role of viral properties in

190 the viral phenotype that contributes to HIV-1 infection, disease progression and

191 pathogenesis.

192

193 Analysis of the viral envelope sequences.

194 For the search of potential mechanisms involved in the changes of the

195 characteristics among the different Envs sets, we analyzed the Env amino-acid

196 (aa) sequences that could be associated with the distinct clinical phenotypes.

197 Initially, we performed a phylogenetic reconstruction from *env* aa sequences

198 together with other aa sequences obtained from HIV-1 Spanish individuals. All

aa sequences analized correspond to HIV-1 subtype B. This analysis did not

reveal phylogenetic relationships between the different groups analysed and no
clustering except for those aa sequences obtained from the same individual
(Figure 7). Envs from LTNP-ECs and one vLTNPs grouped in short branches,
as a consequence of the viral and evolutionary control, whereas long branch
length was observed in the sequences obtained from non-controller patients
(Old and Modern), because of the higher replication and viral evolution in these
individuals.

207 We then carried out a comprehensive study of the protein sequences focusing 208 in the variable loops and their associated potential N-linked glycosylation sites 209 (PNGs) in the gp120 subunit of the Env. In general, as previously reported, 210 there is a trend in the HIV-1 viral Env to gain length and glycosylation sites 211 along the epidemic (49-51). This increasing trend is also found in our work 212 where viruses from the LTNPs (EC, Viremic) and Old Envs isolated in the 90's 213 showed shorter lengths than those of the Modern group obtained in 2013-2014 214 (Table 2). The V3 loop was the most conserved and constant region in length 215 and glycosylation sites (Table 2 and Figure 8), while the other loops showed 216 length increases predominantly in the V2 and V5 loops that were reproduced in 217 the total length (Table 2 and Figure 8). The only statistical differences were 218 noticed between the total length in the LTNPs (EC and Viremic) versus Old and 219 Modern Envs in the V2 and V5 regions (Figure 8). 220 Regarding the PNGS in the sequences, many of the 24 relevant sites previously

described (52-55) were present in these set of viral glycoproteins. However,

222 major differences were observed in the aa extension of the loops with a

progressive acquisition of more PNGS in the Modern Envs (Table 2). Glycan at

224 N289 site was more present in LTNP-ECs, vLTNPs and Old viruses but is not

present in Modern ones. Position N362 which is N proximal to the CD4 binding

- ²²⁶ "DPE" motif (positions 368-370HXB2 sequence) was conserved in LTNP-EC,
- 227 Viremic and Old but was only present in two of the Modern Envs. It is interesting
- to highlight that changes also occurred in the viral transmembrane gp41 protein
- in glycan N816 that was dominant in LTNPs but not in chronic individuals (Old
- and Modern).
- 231 It is interesting to mention that the trend in Env length increase follows the same
- 232 pattern that the functional growth of the Env shown in the distinct viral
- characteristics (see Figures 3 to 6). We observed a good correlation between
- the genetic distance to the subtype B ancestor sequence obtained from Los
- 235 Alamos National Laboratory HIV Database (LANL database,
- http://www.hiv.lanl.gov) and the functionality of viral Env proteins analysed
- 237 (Figure 9). In general, the lower evolutionary sequences (less genetic distance
- to subtype B ancestor) are those with lower functionality (LTNP-ECs) and the
- higher evolutionary sequences are those with higher functionality (Moderns). In
- summary, the viral Envs with the most efficient characteristics are found within
- the Envs of the Modern group that also show the longer gp160 proteins, with
- 242 more glycosylated sites and higher distance to the subtype B ancestor.

243

245 **Discussion**

- HIV-1 infected individuals display a wide spectrum of clinical progression rates.
- 247 The causes of this dispersion are multiple and associated with the operation of
- 248 numerous combinations of host genetic, immunological and viral factors. In this
- 249 work, we studied the potential contribution of viral Env glycoprotein
- 250 characteristics to the clinical outcome of HIV-1 infection in HIV+ individuals with
- 251 different clinical status.
- 252 The different groups of patients were defined by their clinical characteristics,
- 253 distinct VLs and isolation dates because several studies have described a clear

254 correlation between patients' VL and the likelihood of virus transmission,

- disease progression and pathogenesis (56-63).
- 256 Although viral control in HIV-1 individuals has been linked to the host-immune
- responses (10, 64), other researchers and our group, however, stablished, in

258 previous works, a direct connection between deficiencies in HIV-1 Env-

- associated functions and long-term viremia control in LTNP-ECs (17, 18, 20).
- 260 The Envs from these LTNP-EC individuals were ineffective in the CD4 binding
- and in the subsequent functions: viral signaling, fusion and cell entry. These
- 262 Env characteristics ensued in low replication and transmissibility of the virus
- 263 (18, 19, 43, 45). All these data strongly support the role of the viral Env in the
- 264 LTNP-EC phenotype and viral pathogenesis.
- 265 In the present work, we extended these observations to more Env from non-
- progressor subjects, which are not associated with a cluster of infection, in
- 267 comparison to different sets of progressor chronic individuals. The Envs
- 268 characteristics from LTNP individuals (EC and Viremic) were compared with
- those of individuals with progressive infection (Old and Modern). We

270 investigated the defects in the association of Envs with the CD4, membrane fusion impairment and the cell-to-cell virus transfer and viral infection capacities. 271 272 Viral Envs from LTNPs showed the lower binding capacity to the CD4 receptor 273 and this initial inefficient Env/CD4 interaction led to a deficiency in membrane 274 fusion and virus cell-to-cell transfer capabilities. The properties of the Env from 275 LTNPs were not due to the ancestral origin of the LTNPs viruses isolated in the 276 late 80's and 90's, because the chacteristics of the Old viruses which were 277 contemporary to the LTNPs did not showed these limited functional 278 characteristics. On the contrary, Envs from progressors (Old and Modern) 279 presented efficient CD4-mediated viral functionality that triggered an effective 280 membrane fusion and viral transfer. Thus, we disclosed that there is a clear 281 correlation between the level of viral fusion, the transfer capacity of the viral Env 282 and viral infectivity. The observed differences between the characteristics of the 283 Envs from these groups could not be associated with viral tropism, because all 284 the env nucleotide sequences from the studied viruses, showed an R5 tropism 285 (Web PSSM, https://indra.mullins.microbiol.washington.edu/webpssm/). 286 In summary, viral Envs from LTNPs exhibited non-functional characteristics 287 (Figures 3-6) in comparison with those from viruses of the progressive infection 288 groups, supporting the concept that the properties of the Envs were associated 289 with viral control and the clinical progression rate of the HIV-1 individuals. 290 In spite of the limited sampling, because of the difficult and laborious viral 291 characterization of the viral phenotypes, we observed statistically significant 292 differences between the characteristics of the Envs of viruses from LTNP-ECs 293 and the Moderns. Also, if we consider the Env characteristics from all clinical 294 groups, there is a consistent and recurrent tendency, although with no statistical 295 power in some cases, to gain functionality in the viral Envs from the LTNP

296 individuals (LTNP-ECs and vLTNPs), to those of the progressive groups (Old 297 and Modern).

298 Remarkably, the increase in Env functionality also correlated with longer and 299 more glycosylated proteins. The aa length and PNGs' profile of the Envs from 300 the individuals of the distinct clinical groups showed that the studied Envs tend to increase length and glycosylation over the course of the epidemic as 301 302 previously described (see (49, 51)). We observed that Env changes 303 accumulated essentially in the V1, V2, V4 and V5 loops, as previously shown in 304 works relating the role of V1 and V4 loops in the CD4 binding and neutralization 305 (65-68) and viral cell-to-cell transfer capacity (50, 69, 70). Regarding specific 306 changes detected in our study, the loss of the N362 PNGs (position in the HXB2) 307 isolate; group M, subtype B (HIV-1 M:B HXB2R: NCBI:txid11706)) which was 308 prevalent in the EC, Viremic and Old but not in the Modern Envs groups could 309 be associated with the gain of functionality in the Envs. However, the opposite 310 effect with more efficient fusion and transfer capacity was found in Australian 311 viruses with the N362 glycosylation site (55). The potential role of the other 312 changes in PNGs detected in our study need to be further investigated. Besides 313 these important changes, it is clear that point mutations could have a significant 314 impact in the viral characteristics and HIV pathogenesis (71, 72). The variants 315 of concern (VOCs) of the pandemic severe acute respiratory syndrome 316 coronavirus (SARS-CoV-2) unfortunately are reminding us (73, 74). Thus, the 317 contribution of the individual mutations deserves further studies but it is now out 318 of the scope of the present work. 319 In contrast with the more significative changes detected in the V2 and V5 loops, 320

it is important to point the stability in length and glycosylation of the V3 loop.

321 This structure is key for viral tropism (75-79) and for the correct CD4 Env

binding as revealed with anti-V3 neutralizing antibodies that abrogate Env-CD4
interaction (80, 81).

324 In this study, we confirmed the inefficient functionality of the Envs from LTNP-325 EC individuals previously described for a cluster of viruses (18, 20), but 326 extended to HIV+ individuals controlling viremia which are not clustered by the 327 same transmitted/founder (T/F) virus. Also, a gain of Envs functionality from 328 those of the LTNP individuals to the chronic not controlling individuals was 329 identified. This improvement was detected in every Env characteristic analyzed; 330 expression, fusion, virus transfer and infectivity. Interestingly, this functional 331 growth of viral Env was associated in this study with length and PNGs increases 332 in the variable loops. This increase was also reported in studies analyzing the 333 susceptibility, neutralization sensitivity, co-receptor binding, host range and viral 334 phenotype (49). This increase in the V1-V2 length and PNGs has also been 335 detected thorough chronic infections from early to late viral Env sampling like in 336 our work (49). Likewise in a group of individuals infected with closely related 337 viruses higher PNGs density has been observed in the V1-V5 region of the 338 gp120 during chronic infection compared to those oberved during the early 339 acute infection phase (82). In viruses from the HIV-1 subtype B, it seems that 340 early after viral transmission to a new host a selection for viral variants with 341 shorter variable regions and a reduced degree of PNGs occurs (83). The growth 342 in functionality of the viral characteristics was also correlated with the genetic 343 distance of the sequences to the subtype B ancestor. Genetic variability in env 344 gene has been is associated with an increase in viral infectivity and replication 345 capacity (84-89). These changes could facilitate viral replication by increasing 346 viral fitness that favors the escape from the immune response and anti-retroviral 347 therapy (ART) failure (90-99).

348 The non-functional characteristics of the primary Envs of LTNP individuals (ECs 349 and Viremics) resulted in poor viral replication and very limited evolution that 350 could allow the efficient immune control of HIV-1 infection and pathogenesis. It 351 has been reported that in a LTNP-EC patient that followed discontinued ART, 352 the V1 domain of his HIV-1 strain that retained good infectivity and replicative 353 capacity included two additional N-glycosylation sites and was placed in the top 354 1% of lengths among the 6,112 Env sequences analyzed in the Los Alamos 355 National Laboratory online database (100). 356 Therefore, it is conceivable that the functional characterization of the inefficient 357 HIV-1 Envs could be significant in the development of a new generation of

358 immunogens. Indeed, attenuated HIV or simian immunodeficiency virus (SIV)

359 vaccines (LAHVs or LASVs) have been postulated as therapeutic vaccine

360 strategies (101-107). However, further antigenic and immunogenicity work is

361 needed to disclose the potential implications of these non-functional HIV Envs

in the vaccine/cure field.

363

In summary, in this work, we exposed that the characteristics of the viral Envs from different groups of HIV-1 infected individuals could be associated with the short or long-term VL control and the clinical progression rate of the infection. The non-functional HIV-1 Envs could help in the development of new strategies for functional cure and virus eradication. Our data support the hypothesis that the functionality of viral Envs is a crucial characteristic for the control of viral infection, replication and pathogenesis.

371

372

373 Material and methods

374 Viral envelopes.

- 375 Forty-one viral envelopes (Envs) were obtained from samples of different
- 376 origins: the HIV HGM BioBank integrated in the Spanish AIDS Research
- 377 Network (RIS-RETIC, ISCIII) (samples 1,2,3,6,7,8,13,14,15,16,17,18,19), the
- 378 Centro Sanitario Sandoval, Hospital Clínico San Carlos (samples
- 379 21,22,24,28,30,31,32,33,36,37,38,39,40,42,43,44,45,46,49,50,51,52), the
- irsiCaixa Research Foundation (samples 9,10,11,12) and from Hospital Xeral
- de Vigo (samples 26,27). Samples were obtained in three different phases of
- the Spanish epidemic from 1993-94, 2004-2005 and 2013-2014. Samples were
- 383 processed following current procedures and frozen immediately after their
- 384 reception. All patients participating in the study gave their informed consent and
- 385 protocols were approved by institutional ethical committees. Identification
- 386 numbers and characteristics are found in Table1.
- 387

388 Ethics Statement.

389 Samples were obtained from participants who gave informed consent for 390 genetic analysis studies and they were registered as sample collection in the 391 Spanish National Registry of Biobanks for Biomedical Research with number 392 C.0004030. The consents were approved by the Ethical and Investigation 393 Committees of the "Centro Sanitario Sandoval" (Madrid) and the samples were 394 encoded and de-identified in these Centers. All clinical investigations were 395 conducted according to the principles expressed in the Declaration of Helsinki. 396 The studies were approved by the Comité de Ética de la Investigación y de 397 Bienestar Animal of the Instituto de Salud Carlos III with CEI PI 05 2010-v3 and 398 CEI PI 09-2013 numbers.

399 Generation of *env* gene expression plasmids.

400 The env genes were amplified at limiting dilution by nested PCR from proviral 401 DNA. The products were cloned into the pcDNA3.1D/V5-His's Topo expression 402 vector (Invitrogen) and NL4.3. The R5-tropic BaL.01-env (catalog number 403 11445) glycoprotein plasmid was from the NIH AIDS Research and Reference 404 Reagent Program. Ten viral Envs were derived from 6 LTNP-EC patients, 10 405 clones from 6 Viremic LTNPs, 10 clones from 6 "Old" individuals (contemporary 406 to LTNPs) and 11 clones from 10 recent "Modern" patients and NL4.3 and 407 BaL.01 reference clones expression plasmids were transformed in DH5 α cells, 408 and clones sequenced to check the correct insertion of the env gene. 409 Env expression and fusion assays. 410

411 The Env expression plasmids were used to transfect HEK-293T cells with X-

412 tremeGENE HP DNA Transfection Reagent (Sigma) in combination with either

413 a Tat expression plasmid pTat for Env expression and fusion assays, or with the

414 *env* defective HIV-1 backbone pSG3 plasmid for viral transfer assays (18, 19,

415 **108**). As a negative control, HEK-293T cells were transfected only with pTat

416 and as a positive control we use the BaL and NL4.3 Envs. HEK-293T cells were

417 chosen as effector cells since they provide sensitive measures of fusion even

418 when using low fusogenic Env. 24 hours post-transfection, cells were collected,

419 and tested for Env surface expression and also fusion activity.

420 To test Env expression, 1×10^5 Env/Tat co-transfected HEK-293T cells were

421 incubated with 2G12 and IgGb12 monoclonal antibodies (mAbs; Polymun,

422 Viena, Austria) at 6 μg/mL each for 45 minutes at RT. After washing the cells,

423 the PE-labeled goat anti-human IgG (Jackson ImmunoResearch Laboratories)

424 was added and incubated in the dark at room temperature for 15 minutes, as

425	similarly reported (18, 19). Cells were washed, fixed in formaldehyde 1%,
426	acquired in a Celesta flow cytometer (BD FACS Celesta) and analyzed using
427	the Flow-Jo software (Tree Star Inc.) The percentage of Env-positive cells and
428	the Mean Fluorescence Intensity (MFI) of these cells were used to evaluate Env
429	expression.
430	To test fusion activity, 1x10 ⁴ Env/Tat-transfected or control Tat-transfected
431	HEK-293T cells were mixed (ratio 1:1) in 96-well plates with
432	CD4 ⁺ CXCR4 ⁺ CCR5 ⁺ TZM-bl reporter cells for 6 hours at 37°C. Luciferase
433	activity was measured (Fluoroskan Accent, Labsystems) using Brite-Lite
434	(PerkinElmer) and normalized to BaL-Env-mediated fusion. NL4.3 and BaL-Env
435	expression plasmids were used as positive controls for Env staining and as
436	reference value for fusion activity (BaL = 100%), as similarly reported (19, 108)
437	(summarized in the scheme of Figure 1B).
438	

....

439 HIV-1 transfer/CD4 binding

440 To test viral transfer activity, which exclusively depends on the binding of gp120 441 to the CD4 molecule, Env expression plasmids were co-transfected with the 442 Env-defective pSG3 plasmid in HEK-293T cells, as similarly reported (18, 19, 108). One day after transfection, 1×10^5 HEK-293T cells were mixed at a 1:1 443 444 ratio in 96-well plates with primary CD4+ T lymphocytes freshly isolated from 445 healthy donors by negative selection (CD4+ T-Cell Isolation Kit II, human, 446 Miltenyi Biotec). Viral transfer was assessed after 24 hours of incubation at 447 37°C in permeabilized (FIX & PERM Cell Permeabilization kit, Invitrogen Life 448 Technologies) and stained cells with the anti-HIV-1 p24 KC57 mAb (anti HIV 449 core antigen RD1 labelled, IZASA) for 20 minutes in the dark at RT. Then, the cells were washed and fixed in formaldehyde 1%, and acquired in a Celesta 450

451 flow cytometer (BD FACS Celesta) and the content of p24 in gated CD4+ T

452 cells and gated HEK-293T cells was analyzed using the Flow-Jo software (Tree

453 Star Inc.). The percentage of p24+ HEK-293T cells was used as a control for

454 transfection efficiency and was similar among all experiments. Since co-

455 receptor binding or fusion activity are not required for viral transfer, the

456 frequency of p24+/CD4+ T cells was a direct measure of the amounts of HIV-1

457 virions bound to or taken up by target cells (summarized in the scheme of

458 **Figure 1C**).

459

460 Infectivity assay

461 Cloned viral Envs were used to generate pseudoviruses by co-transfection with

462 pSG3 plasmid of HEK-293T cells as indicated above and tested in TZM-bl cells

to determine the infectivity capacity. Serial Dilutions of the pseudoviruses

464 generated with the different Envs of the different groups of patients were made

in a 96-well plate. Then, 1×10^5 TZM-bl cells were added to the pseudoviruses

466 with DEAE dextran hydrochloride (Sigma) at 18 µg/mL. After 48 hours of

467 incubation at 37°C, luciferase activity was measured (Fluoroskan Accent,

468 Labsystems) using Brite-Lite (PerkinElmer). Uninfected TZM-bl cells were used

469 as a negative control. The TCID₅₀ (Median Tissue Culture Infectious Dose)

470 value was calculated with Montefiori template and normalized with the viral

471 concentrations (summarized in the scheme of Figure 1D).

472

473 **Phylogenetic Analysis.**

474 The evolutionary history was inferred by using the "maximum likelihood" (ML)

475 method and JTT matrix-based model (109). The tree with the highest log

476 likelihood (-49687,86) is shown. The percentage of trees in which the

477	associated taxa clustered together is shown next to the branches. Initial tree(s)
478	for the heuristic search was(were) obtained automatically by applying Neighbor-
479	Join and BioNJ algorithms to a matrix of pairwise distances estimated using the
480	JTT model, and then selecting the topology with superior log likelihood value. A
481	discrete Gamma distribution was used to model evolutionary rate differences
482	among sites (5 categories (+G, parameter = 0,6825)). The rate variation model
483	allowed for some sites to be evolutionarily invariable ([+I], 18,05% sites). The
484	tree is drawn to scale, with branch lengths measured in the number of
485	substitutions per site. This analysis involved 140 aa sequences. All positions
486	with less than 95% site coverage were eliminated (i.e., fewer than 5% alignment
487	gaps), and missing data and ambiguous bases were allowed at any position
488	(partial deletion option). There were a total of 829 positions in the final dataset.
489	Evolutionary analyses were conducted in MEGA X (110).
490	Nucleotide sequences have been deposited in GeneBank under the following
491	numbers: KC595156, KC595162, KC595225, KC595227, KC 595189,
492	MH605987, MH605986, KC595190, MH605988, MH605992, MH605991,
493	MH605970, MH605971, KC595223, KC595222, MH605972, MH605975,
494	MH605976, MH605978, MH605973, MH605979, MH605980, MH605981,
495	MH605982, MH605983, MH605984, MK394184, MK394185.
496	

497 Statistical analysis.

498 Data and statistical analyses were performed using GraphPad Prism, version

499 6.07 (GraphPad Software). Significance when comparing groups was

500 determined with a nonparametric Kruskal-Wallis or by nonparametric Dunn's

- 501 test for multiple comparisons. A nonparametric Spearman test was used to
- 502 calculate correlations.

503 Data Availability

504 All "accession numbers" and "data" of this work are available.

505

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544

546 Figures legends

547	Figure 1. Outline of the experimental model used for the analysis of Env
548	expression, Env-mediated cell-to-cell fusion, viral transfer and viral
549	infectivity. (A) Env expression: HEK-293T cells will be co-transfected with
550	primary of reference full-length viral <i>env</i> and a ptat Δenv HIV-1 expression
551	plasmid, allowing Env cell-surface expression in a viral production context. Cell-
552	surface Env expression will be then analyzed by flow cytometry using specific
553	anti-Env antibody. (B) Env-mediated fusion activity: after 24 hours, effector
554	HEK-293T cells producing HIV-1 particles bearing primary or reference Envs
555	will be co-cultured with TZM-bl cells to force synapsis formation and CD4-
556	mediated binding of budding particles to target cells. (C) Env-mediated viral
557	transfer: HEK-293T cells producing HIV-1 particles carrying primary or
558	reference Envs will be co-cultured with primary CD4+ T cells. Then, HIV-1
559	transfer will be analyzed by flow cytometry using specific anti-p24 antibody in
560	target CD4+ T cells. (D) Env-mediated viral infection: TZM-bl cells will be
561	infected with serial dilutions of viral particles obtained from transfected HEK-
562	293T and carrying the different primary or reference HIV-1 Envs. After 48 hours,
563	infectivity capacity will be analyzed by quantifying luciferase assay in infected
564	TZM-bl cells.
565	

566 Figure 2. Analysis of the expression of the different HIV-1-Env

567 glycoproteins from LTNP-EC, Viremic LTNP and control progressors
568 patients.

569 Flow cytometry analysis of the cell-surface expression level of the assayed HIV-

570 1 Envs in HEK-293T cells from LTNP-EC (gray bars), vLTNP (green bars), Old

571 (orange bars) and Modern individuals (red bars) or reference HIV-1 viral strains

572	(ptat, No Ab2, NL4.3 and BaL, black bars). Env protein expression for each
573	patient (A) and Env protein expression in each group of patients comparing
574	mean values between each group (Kruskal-Wallis, Dunn's Multiple
575	Comparisons Test) (B); p value for comparison between all groups is shown,
576	top left. Values are mean ± S.E.M. of three independent experiments.
577	
578	Figure 3. Analysis of membranes fusion-phenotypic features of HIV-1
579	Envs isolated from LTNP-EC, viremic LTNP and P individuals. Analysis of
580	the ability to induce cell-to-cell fusion of HIV-1 Env proteins obtained from
581	LTNP-EC (gray bars), vLTNP (green bars), Old (orange bars) and Modern
582	individuals (red bars) or reference HIV-1 viral strains (ptat, NL4.3 and BaL,
583	black bars). (A) Env fusogenic activity for each patient in each group. (B)
584	Relative fusion activity of the full Env collection compared to the BaL control
585	established at 100% and grouped in the different groups of patients. Values are
586	mean ± S.E.M. of three independent experiments. Statistical analysis was
587	performed using Kruskal-Wallis, Dunn's Multiple Comparisons Test; p value for
588	comparison between all groups is shown, top left.
589	
590	Figure 4. Analysis of HIV-1 Env-mediated cell-to-cell viral transfer.
591	Analysis of the ability to induce cell-to-cell virus transfer of HIV-1 Env proteins obtained

592 from LTNP-EC (gray bars), vLTNP (green bars), Old Patients (orange bars), recent

593 patients (Moderns) (red bars) or reference HIV-1 viral strains (pSG3, CD4+ cells, NL4.3

and BaL, *black bars*). Analysis of HIV-1 Env-mediated cell-to-cell viral transfer for each

- 595 patient (A) and in each group where P values compare medians between groups using
- 596 a nonparametric Kruskal-Wallis Test (Kruskal-Wallis, Dunn's Multiple Comparisons

- 597 Test) (B); p value for comparison between all groups is shown, top left. Values
- 598 are mean ± S.E.M. of two independent experiments.
- 599

600 Figure 5. Viral infectivity of the viral Envs.

- 601 Analysis of the infectivity (TCID₅₀ value normalized by viral p24 input) of the different of
- 602 HIV-1 Env proteins obtained from LTNP-EC (gray bars), vLTNP (green bars), Old
- 603 (orange bars) and Moderns (red bars) patients or reference HIV-1 viral strains (pSG3,
- 604 NL4.3 and BaL, *black bars*). Analysis of Env infectivity for each patient (A) and in each
- 605 group where P values compare medians between groups using a nonparametric
- 606 Kruskal-Wallis, Dunn's Multiple Comparisons Test (B); p value for comparison
- 607 between all groups is shown, *top left*. Values are mean ± S.E.M. of three
- 608 independent experiments.
- 609

610 Figure 6. Analysis of the correlation of the fusion, transfer and viral

611 infectivity Env characteristics between groups.

612 (A) Correlation between Relative fusion and HIV Transfer of all Envs of the

613 different groups LTNP-EC (gray circle), vLTNP (green circle), Old patients

614 (orange square) and Modern patients (red square). The correlation was

- 615 calculated with a nonparametric Spearman test. (B) Correlation between
- 616 Relative fusion and Infectivity (TCID₅₀ value normalized by viral p24 input) of all
- 617 Envs of the different groups LTNP-EC (gray circle), vLTNP (green circle), Old
- 618 patients (*orange square*) and Modern patients (*red square*). The correlation was
- 619 calculated with a nonparametric Spearman test. (C) Correlation between
- 620 Infectivity and HIV Transfer of all Envs of the differents groups LTNP-EC (gray
- 621 *circle*), vLTNP (green circle), Old patients (orange square) and recent patients
- 622 Moderns) (red square) is shown. The correlation was calculated with a

- 623 nonparametric Spearman test. Values are mean ± S.E.M. of three independent
- 624 experiments; p value for comparison between all groups is shown, top left.
- 625
- 626 Figure 7. Phylogenetic analysis of the vial Envs.
- 627 The evolutionary history of the Env aa sequences was inferred as described in
- 628 Materials and Methods using the Maximum Likelihood method and JTT matrix-
- based model (109). The tree with the highest log likelihood (-49687,86) is
- 630 shown. The percentage of trees in which the associated taxa clustered together
- 631 is shown next to the branches. Evolutionary analyses were conducted in MEGA
- 632 X (110).
- 633

Figure 8. Analysis of the length and glycosylation sites in the loops of the
Envs from the different groups.

- 636 Analysis of the length of each variable loops V1 (A), V2 (B), V3 (C), V4 (D), V5
- 637 (E) and all variable loops together (F). The results were grouped (LTNP-ECs:
- 638 gray bar, vLTNPs: green bar, Old patients: orange bar, and recent patients
- 639 (Moderns): red bar) and compared using a nonparametric Kruskal-Wallis,
- 640 Dunn's Multiple Comparisons Test; p value for comparison between all groups
- 641 is shown, *top left*. Values are mean ± S.E.M. of three independent experiments.
- 642

643 Figure 9. Correlation of the expression, fusion, transfer and viral

644 infectivity Env characteristics with the nucleotide genetic distance to

- 645 subtype B ancestor.
- 646 Correlation between genetic distance to subtype B ancestor of all Envs of the
- 647 different groups and Env expression (A), Relative fusion (B), HIV Transfer (C)
- and Infectivity (D). LTNP-ECs (gray circle), vLTNPs (green circle), Old patients

- 649 (orange square) and Modern patients (red square). The correlations were
- 650 calculated with a nonparametric Spearman test (p and r values are shown, *top*
- 651 *left*). Values of Env expression, Relative fusion, HIV transfer and Infectivity are
- 652 mean ± S.E.M. of three independent experiments.

653

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656

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Clinical Group	Sub-group	Env code ^a	Patient Identification code	Viral Load ^g (at sampling)	Diagnostic time	Sampling time	Viral dating ^b	HLA B
LTNP	EC	1	2057906-3	< 50	1993	2004	1989	4901/5701
		2	3227050	< 50	1988	2004	1991	0702/5201
		3	3227058-3	< 50	1992	2004	1991	1402/1402
		6	20044616-3	< 50	1998	2004	1999	1501/5703
		* ۲	10246788	< 50	1992	2005	1993	4402/5701
		L 8		< 50	"	u	"	u
		٢9	MDM ^c	507	1998	1996	1987	4402/3501
		10	с	< 50	"	2011	1996	u
		11	с	< 50	"	2005	"	u
		L12	с	< 50	u	2005	"	u
	Viremic	13	4022834	3.710	1994	2004	ND	1401/4403
		14	9684	2.557	1998	2005	1994	1302/4001
		15	2988465	2.286	1993	2004	1999	1402/2705
			38 17 5	418	1996	2014	1999	2705/5801
		17			u	u	u	u
		18			"	u	"	u
		L19			u	"	u	"
		21	30	7.597	1989	1998	2000	1501/3501
		22	64	11.926	1989	1999	1999	4402/4901
		L24		u	"	2002	"	u
Progressor	Old	 [26	V10 ^d	N.D.	1993	1994	1999	4002/4402
		L27				"	"	
		28	V13	N.D.	1992	1994	1990	0702/1402
		□ 30	L10	89.000		1993	1993	1501/4901
		_31		u	u	u	"	u
		□ ³²	L 11	42.000	1993	1993	2000	1801/5101
		L33		u	"		"	"
		□ 36	I14 ^d	130.000	1987	1994	2002	0702/3502
		L37	d	u	u	"	u	"
		38	118	170.000	1991	1994	1990	1402/4403
	Modern	□ 39	ESI 17A	156.300	2014	2013	N.A ^f	4201/4402
		L40		"	"	u		u
		□ 42	ESI 39A	137.700	2012	2014	N.A.	1517/3801
		43		u	u	"		"
		□ 44	ESI 41A	129.700	2012	2014	N.A.	3503/5701
		45		u	u	u		u
		46	ESI 5A 2	49.107	2004	2007	N.A.	4102/4402
		□ 49	ESI 42 A	11.510	2011	2014	N.A.	1402/4403
		L50		u	и	u		u
		∑ 51	ESI 42 B	41.090	2011	2014	N.A.	0702/1501
		L52		u	u	"		"

 $\begin{array}{c} 1009 \\ 1010 \end{array}$ ^aHIV-1 Env number used in this study and identification codes.

^bAccording to Bello et al. (2004). J Gen Virol. Feb;85(Pt 2):399-407. doi: 10.1099/vir.0.19365-1011 0. PMID: 14769897.

- 1012
- 1013 ^cDouble infected individual (Casado et al. (2007) J Infect Dis. 2007 Sep 15;196(6):895-9. doi:
- 1014 10.1086/520885. Epub 2007 Aug 14. PMID: 17703421).
- 1015 ^dIndividuals with a short antiviral therapy (AZT (zidovudine) and DDI (didanosine) for V10
- 1016
- patient and AZT for I14 patient). ^eThe Modern Individuals have been infected within 3 years. 1017
- 1018 ^fN.A.:Not applicable.
- ^gHIV RNA copies/mL 1019
- 1020 ^{*}*Envs isolated from the same patient are indicated by brackets.*
- 1021 ": same value than above.

1022

1023Table 2. Molecular characteristics of HIV-1 Envs: sequence length and N potential1024glycosylation sites (PNGs) in the variable loops (Vn) of the gp120 subunit.

Clinical Group	Subgroup	Env code	^a V1/G	V2/G	V3/G	V4/G	V5/G	^ь ΣVn/G	^c Mean/G	^d Gp160	^e Meaı
TNP	EC	1	28/4	43/2	37/2	28/4	12/1	148/13		848	
		2	33/5	41/2	37/2	31/4	12/1	154/14		853	
		3	33/5	41/2	37/2	31/3	12/2	154/14		853	
		6	28/3	41/2	37/2	34/4	12/1	152/12		852	
		* 🗖 7	32/5	47/2	37/2	30/4	11/2	157/15	151.1/14.4	859	851.8
		8	32/5	47/2	37/2	30/4	11/2	157/15		859	
		۶٦	24/4	43/2	36/2	28/4	12/1	143/14		843	
		10	27/4	42/2	37/2	29/5	14/2	149/16		851	
		11	27/5	42/3	37/2	29/5	13/2	148/17		850	
		L12	27/4	42/3	37/2	32/5	12/1	150/14		850	
	Viremic	13	31/5	41/2	37/2	31/4	13/1	153/14		854	
		14	29/4	42/2	37/2	32/5	12/2	152/15		852	
		15	34/5	41/2	37/2	36/5	12/1	160/15		860	
		□ ¹⁶	29/5	41/2	37/1	29/5	12/1	148/14		849	
		17	29/5	41/2	37/2	29/5	12/1	148/16	150,3/14.1	849	851.5
		18	29/4	41/2	37/2	29/5	12/1	148/15		849	
		L19	29/4	41/2	37/2	29/5	12/1	148/14		849	
		21	24/3	41/2	37/1	30/5	10/0	142/11		842	
		 ²²	28/4	41/2	37/2	32/5	12/1	150/14		850	
		L24	37/7	41/2	36/2	32/5	12/1	158/15		861	
Progressor	Old	 ²⁶	31/4	41/3	37/2	39/7	14/2	160/18		862	
		L27	31/5	48/3	37/2	28/5	14/2	158/16		858	
		28	25/5	41/2	36/2	33/5	12/2	145/15		848	
		┌ 30	33/4	41/2	37/2	27/4	11/1	150/14		852	
		_ 31	33/5	41/2	37/2	36/5	13/2	158/16		860	
		□ 32	28/5	44/2	36/2	30/5	15/2	151/15	153,8/15.2	853	855.8
		L33	31/4	44/2	36/2	30/5	15/2	156/14		856	
		□ 36	28/4	46/1	37/2	34/5	14/2	157/15		859	
		_37	28/4	46/2	37/2	34/5	14/2	157/16		859	
		38	30/4	41/3	37/1	31/4	13/2	150/13		851	
	Modern ^e	┌ 39	31/4	41/2	37/2	29/4	12/2	149/14		849	
		40	31/4	41/2	37	29/4	17/2	154/14		849	
		 4 2	29/4	48/2	37	36/6	17/2	167/13		878	
		43	29/4	48/3	37	30/4	15/2	159/15		872	
		□ 44	28/4	47/3	37	31/4	15/2	158/15	158.1/14.7	859	862.0
		_45	28/4	47/2	37	31/4	15/2	158/14		859	

46	35/4	46/3	37	33/5	13/2	164/15	865
□ ⁴⁹	37/6	41/2	37	42/7	13/1	170/18	871
L50	37/6	41/2	37	42/7	13/1	170/18	871
□ ⁵¹	31/4	42/2	37	26/3	13/1	149/12	853
L52	29/4	42/2	37	32/6	12/1	152/15	856

 ^aLength in amino acid (aa) and potential glycosylation sites (PNGs) of the Env-gp120 variable regions (Vn; from V1 to V5) expressed as Vn/G ratio. ^bΣVn/G indicates the sum of the aa lengths of the Vn (n; from 1 to 5) and the potential G sites. ^cMean/G indicates the mean length and PNG value for each group of Envs. ^dGp160 shows the total length in aa of each Env including the gp41 subunit and the gp120 subunit. ^eMean gp160 length in aa for each group of Envs.

Clinical Group	Sub-group	Env code ^a	Patient Identification code	Viral Load ^g (at sampling)	Diagnostic time	Sampling time	Viral dating ^b	HLA B	
LTNP	EC	1	2057906-3	< 50	1993	2004	1989	4901/5701	
		2	3227050	< 50	1988	2004	1991	0702/5201	
		3	3227058-3	< 50	1992	2004	1991	1402/1402	
		6	20044616-3	< 50	1998	2004	1999	1501/5703	
		* ۲7	10246788	< 50	1992	2005	1993	4402/5701	
		8		< 50	"	"	u	u	
		٢9	MDM ^c	507	1998	1996	1987	4402/3501	
		10	с	< 50	"	2011	1996	u	
		11	с	< 50	"	2005	"	u	
		L12	с	< 50	u	2005	"	u	
	Viremic	13	4022834	3.710	1994	2004	ND	1401/4403	
		14	9684	2.557	1998	2005	1994	1302/4001	
		15	2988465	2.286	1993	2004	1999	1402/2705	
			38 17 5	418	1996	2014	1999	2705/5801	
		17			"	"	"	u	
		18			"	u	u	u	
		L19			"	"	"	"	
		21	30	7.597	1989	1998	2000	1501/3501	
		22	64	11.926	1989	1999	1999	4402/4901	
				"	u	2002	u	u	
Progressor	Old	26	V10 ^d	N.D.	1993	1994	1999	4002/4402	
		L27				"	"		
		28	V13	N.D.	1992	1994	1990	0702/1402	
		30	L10	89.000		1993	1993	1501/4901	
		_ 31		"	u	u	"	"	
		³²	L 11	42.000	1993	1993	2000	1801/5101	
		L 33		"	"		"	u	
		□ 36	I14 ^d	130.000	1987	1994	2002	0702/3502	
		L 37	d	"	"	"	"	u	
		38	118	170.000	1991	1994	1990	1402/4403	
	Modern	□ 39	ESI 17A	156.300	2014	2013	N.A ^f	4201/4402	
		_ 40		"	"	u		"	
		□ 42	ESI 39A	137.700	2012	2014	N.A.	1517/3801	
		43		"	u	u		u	
		□ 44	ESI 41A	129.700	2012	2014	N.A.	3503/5701	
		45		"	u	u		u	
		46	ESI 5A 2	49.107	2004	2007	N.A.	4102/4402	
		□ 49	ESI 42 A	11.510	2011	2014	N.A.	1402/4403	
		L 50		"	u	u		u	
		5 1	ESI 42 B	41.090	2011	2014	N.A.	0702/1501	
		52		"	"	"		"	

Table 1. Epidemiological, clinical and host characteristics of the viral Envs.

^a*HIV-1* Env number used in this study and identification codes.

^bAccording to Bello et al. (2004). J Gen Virol. Feb;85(Pt 2):399-407. doi: 10.1099/vir.0.19365-0. PMID: 14769897.

^cDouble infected individual (Casado et al. (2007) J Infect Dis. 2007 Sep 15;196(6):895-9. doi: 10.1086/520885. Epub 2007 Aug 14. PMID: 17703421).

^dIndividuals with a short antiviral therapy (AZT (zidovudine) and DDI (didanosine) for V10 patient and AZT for 114 patient). ^eThe Modern Individuals have been infected within 3 years.

^fN.A.:Not applicable.

^gHIV RNA copies/mL

*Envs isolated from the same patient are indicated by brackets.

": same value than above.

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Clinical Group	Subgroup	Env code	^ª V1/G	V2/G	V3/G	V4/G	V5/G	^ь ΣVn/G	۵Mean/G	^d Gp160	^e Mean
LTNP	EC	1	28/4	43/2	37/2	28/4	12/1	148/13		848	
		2	33/5	41/2	37/2	31/4	12/1	154/14		853	
		3	33/5	41/2	37/2	31/3	12/2	154/14		853	
		6	28/3	41/2	37/2	34/4	12/1	152/12		852	
		* ⁻↗	32/5	47/2	37/2	30/4	11/2	157/15	151.1/14.4	859	851.8
		L 8	32/5	47/2	37/2	30/4	11/2	157/15		859	
		٢٩	24/4	43/2	36/2	28/4	12/1	143/14		843	
		10	27/4	42/2	37/2	29/5	14/2	149/16		851	
		11	27/5	42/3	37/2	29/5	13/2	148/17		850	
		L ₁₂	27/4	42/3	37/2	32/5	12/1	150/14		850	
	Viremic	13	31/5	41/2	37/2	31/4	13/1	153/14		854	
		14	29/4	42/2	37/2	32/5	12/2	152/15		852	
		15	34/5	41/2	37/2	36/5	12/1	160/15		860	
		۲16	29/5	41/2	37/1	29/5	12/1	148/14		849	
		17	29/5	41/2	37/2	29/5	12/1	148/16	150,3/14.1	849	851.5
		18	29/4	41/2	37/2	29/5	12/1	148/15		849	
		_19	29/4	41/2	37/2	29/5	12/1	148/14		849	
		21	24/3	41/2	37/1	30/5	10/0	142/11		842	
		 <u> </u>	28/4	41/2	37/2	32/5	12/1	150/14		850	
		_24	37/7	41/2	36/2	32/5	12/1	158/15		861	
Progressor	Old	26	31/4	41/3	37/2	39/7	14/2	160/18		862	
		L ₂₇	31/5	48/3	37/2	28/5	14/2	158/16		858	
		28	25/5	41/2	36/2	33/5	12/2	145/15		848	
		۲30	33/4	41/2	37/2	27/4	11/1	150/14		852	
		_31	33/5	41/2	37/2	36/5	13/2	158/16		860	
		 []32	28/5	44/2	36/2	30/5	15/2	151/15	153,8/15.2	853	855.8
		_33	31/4	44/2	36/2	30/5	15/2	156/14		856	
		□ 36	28/4	46/1	37/2	34/5	14/2	157/15		859	
		_37	28/4	46/2	37/2	34/5	14/2	157/16		859	
		38	30/4	41/3	37/1	31/4	13/2	150/13		851	
	Modern ^e	┌ 39	31/4	41/2	37/2	29/4	12/2	149/14		849	
		40	31/4	41/2	37	29/4	17/2	154/14		849	
		□ 42	29/4	48/2	37	36/6	17/2	167/13		878	
		_43	29/4	48/3	37	30/4	15/2	159/15		872	
		□ 44	28/4	47/3	37	31/4	15/2	158/15	158.1/14.7	859	862.0
		45	28/4	47/2	37	31/4	15/2	158/14		859	
		46	35/4	, 46/3	37	33/5	13/2	164/15		865	
		49	37/6	41/2	37	42/7	13/1	170/18		871	
		50	37/6	41/2	37	42/7	13/1	170/18		871	
		 51	31/4	42/2	37	26/3	13/1	149/12		853	
		52	29/4	42/2	37	32/6	12/1	152/15		856	

expressed as Vn/G ratio. ^b $\Sigma Vn/G$ indicates the sum of the aa lengths of the Vn (n; from 1 to 5) and the potential G sites.

^c*Mean/G* indicates the mean length and PNG value for each group of Envs. ^dGp160 shows the total length in aa of each Env including the gp41 subunit and the gp120 subunit.

^eMean gp160 length in aa for each group of Envs.

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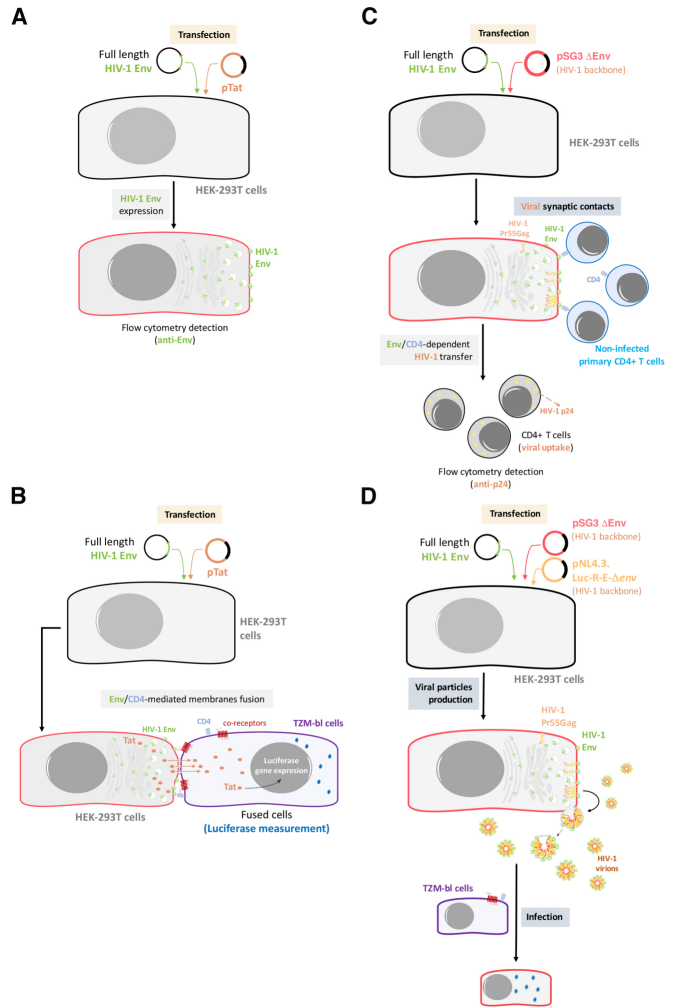
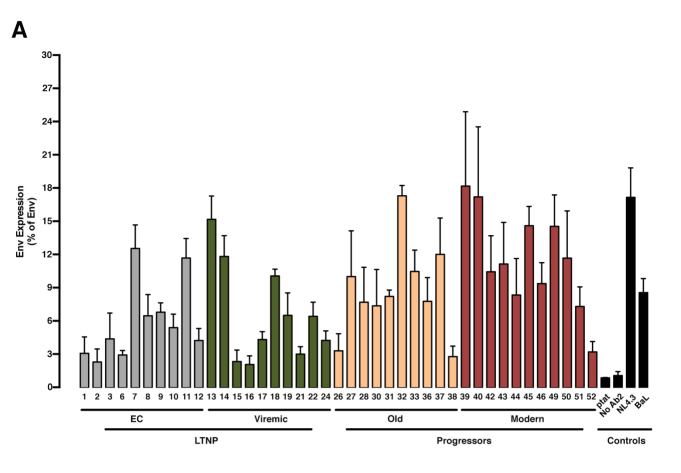
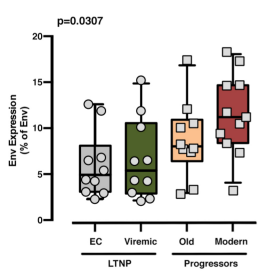
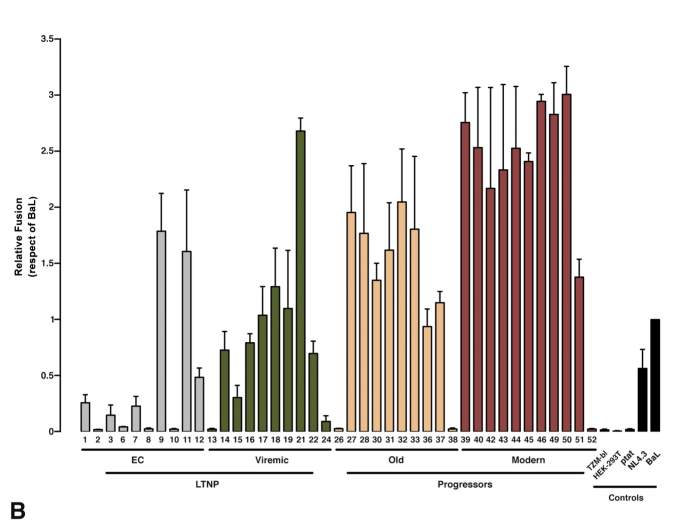


Figure 1 Silvia Pérez-Yanes *et al.* Infected cells (Luciferase measurement)

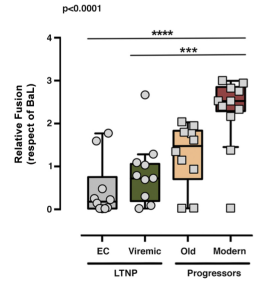


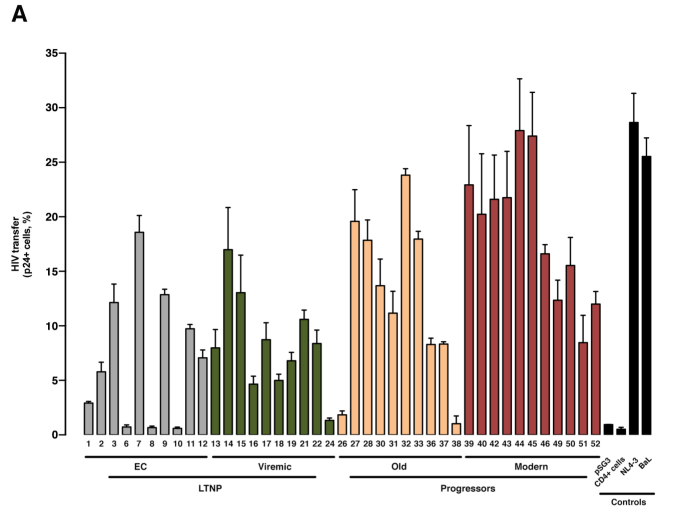




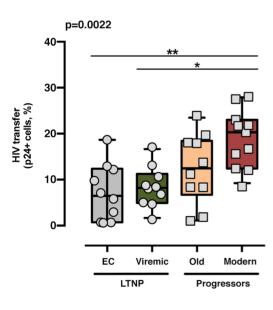


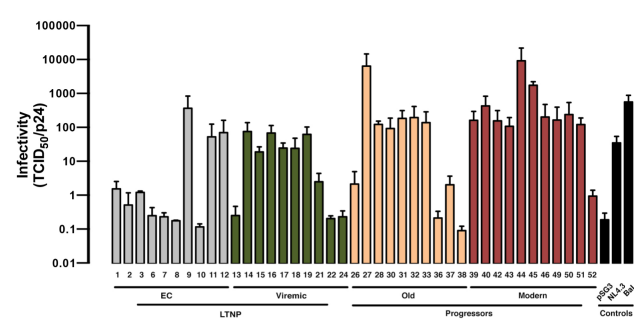
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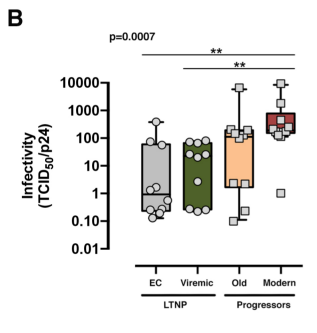




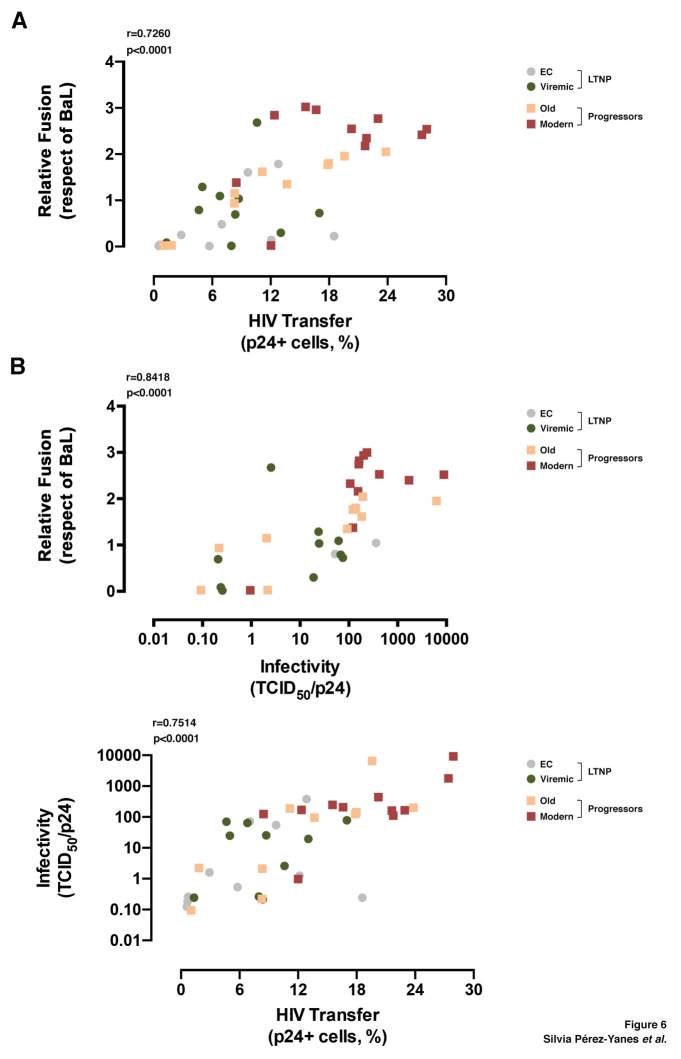


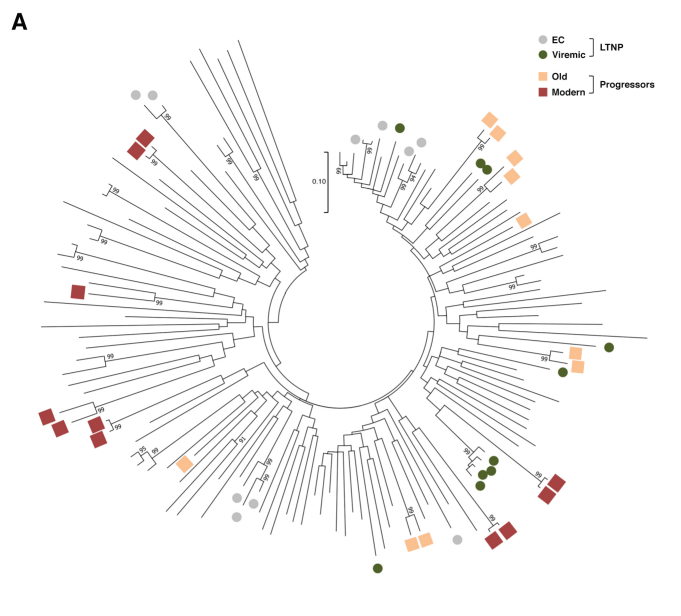


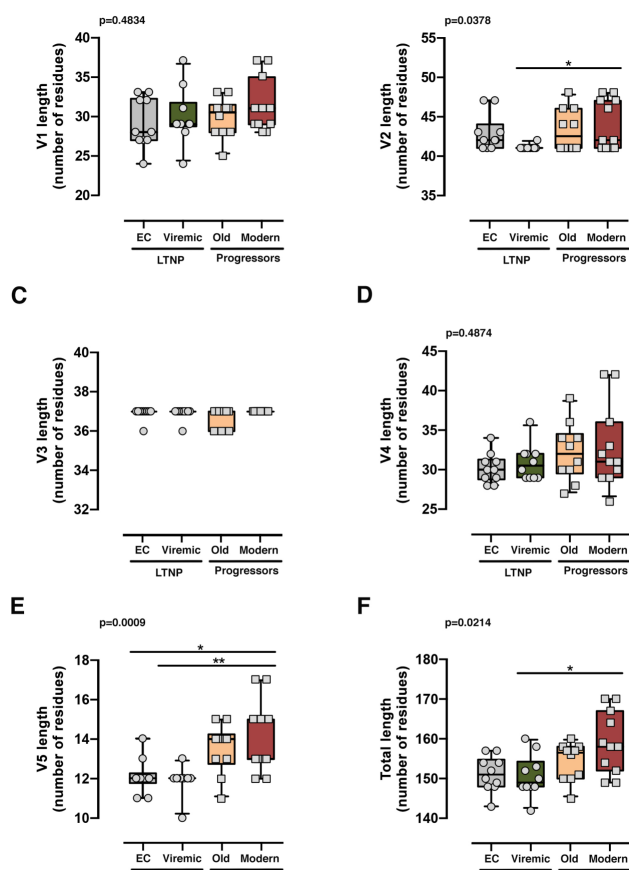




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LTNP Progressors

> Figure 8 Silvia Pérez-Yanes et al.

Progressors

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