## Decoding the biogenesis of HIV-induced CPSF6 puncta and their fusion with the nuclear speckle.

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### 18 Highlights

- The formation of HIV-induced CPSF6 puncta is critical for restoring HIV-1 nuclear
   reverse transcription (RT).
- CPSF6 protein lacking the FG peptide cannot bind to the viral core, thereby failing to form HIV-induced CPSF6 puncta.
- The FG peptide, rather than low-complexity regions (LCRs) or the mixed charge
   domains (MCDs) of the CPSF6 protein, drives the formation of HIV-induced CPSF6
   puncta.
- HIV-induced CPSF6 puncta form individually and later fuse with nuclear speckles
   (NS) via the intrinsically disordered region (IDR) of SRRM2.
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### 30 Summary

Viruses rely on host cellular machinery for replication. After entering the nucleus, the HIV genome accumulates in nuclear niches where it undergoes reverse transcription and integrates into neighboring chromatin, promoting high transcription rates and new virus progeny. Despite anti-retroviral treatment, viral genomes can persist in these nuclear niches and reactivate if treatment is interrupted, likely contributing to the formation of viral

36 reservoirs. The post-nuclear entry dynamics of HIV remain unclear, and understanding these

37 steps is critical for revealing how viral reservoirs are established.

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In this study, we elucidate the formation of HIV-induced CPSF6 puncta and the domains of CPSF6 essential for this process. We also explore the roles of nuclear speckle scaffold factors, SON and SRRM2, in the biogenesis of these puncta. Through genetic manipulation and depletion experiments, we demonstrate the key role of the intrinsically disordered region of SRRM2 in enlarging nuclear speckles in the presence of the HIV capsid.

We identify the FG domain of CPSF6 as essential for both puncta formation and binding to the viral core, which serves as the scaffold for CPSF6 puncta. While the low-complexity regions (LCRs) modulate CPSF6 binding to the viral capsid, they do not contribute to puncta formation, nor do the disordered mixed charge domains (MCDs) of CPSF6. These results demonstrate how HIV evolved to hijack host nuclear factors, enabling its persistence in the host.

50 Of note, this study provides new insights into the underlying interactions between host

51 factors and viral components, advancing our understanding of HIV nuclear dynamics and

52 offering potential therapeutic targets for preventing viral persistence.

#### 53 Introduction

54 Since the discovery of HIV (Barre-Sinoussi et al., 1983), the initial stages of the viral life cycle 55 have been understood to primarily occur within the host cytoplasm. Only the pre-integration 56 complex, carrying the fully reverse-transcribed viral DNA, was believed to enter the nucleus 57 for integration into the host chromatin (Suzuki and Craigie, 2007). Recent studies highlighted 58 that the viral genome is transported in the nucleus via a shuttle that shields it from the hostile 59 cellular environment (Rasaiyaah et al., 2013). This shuttle is constituted by the viral capsid 60 (Blanco-Rodriguez and Di Nunzio, 2021; Blanco-Rodriguez et al., 2020; Chen et al., 2016; Selyutina et al., 2020; Yamashita and Emerman, 2004; Zila et al., 2021), which comprises 61 62 250 hexamers and 12 pentamers (Pornillos et al., 2009). Within the structure of the capsid, 63 hydrophobic pockets exist between hexamers, which serve as targets for various 64 nucleoporins, especially the ones carrying FG repeats, facilitating the translocation of the 65 capsid through the nuclear pore complex (NPC) (Buffone et al., 2018; Di Nunzio, 2013; Di 66 Nunzio et al., 2012; Lelek et al., 2015; Matreyek et al., 2013; Price et al., 2014). Recent 67 studies suggest that HIV uses multiple FG regions of several nucleoporins to translocate 68 through the NPC, acting as a chaperone by itself (Dickson et al., 2024; Fu et al., 2024). 69 Using a reductionist system of Nup98 condensates, it has been demonstrated that FG-70 mediated phase partitioning identifies specific sites on the capsid that allow it to interact 71 autonomously with these phases. These findings complement the evolving understanding of 72 the early stages of HIV infection, which has been revisited in recent years. They unveil that 73 crucial stages of early viral infection occur within the host nucleus (Burdick et al., 2020; 74 Dharan et al., 2020; Francis et al., 2020; Scoca et al., 2023; Selyutina et al., 2020). Notably, 75 it has been shown that the pre-integration complex forms within the host nucleus (Muller et 76 al., 2021; Scoca et al., 2023), and incoming viral RNA genomes accumulate in nuclear 77 niches containing, such as like cleavage and polyadenylation specificity factor subunit 6 78 (CPSF6), RNA-binding protein SON and Splicing component, 35 kDa (SC35, also known as 79 serine/arginine-rich splicing factor 2 (SRSF2) (Francis et al., 2020; Rensen et al., 2021; 80 Scoca et al., 2023). However, the mechanistic requirements that govern post-nuclear entry 81 phases, which are crucial for a successful viral infection and the establishment of viral 82 reservoirs, remain enigmatic.

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Specifically, the mechanism behinds the formation of nuclear niches containing viral components and nuclear speckle (NS) factors, such as SC35—traditionally used as a marker for nuclear speckles remains unclear. However, recent studies have revealed that SON and serine/arginine repetitive matrix protein 2 (SRRM2) are essential for NS biogenesis (Fu and Maniatis, 1990). Particularly, intrinsically disordered regions play a critical role in the NS formation (Ilik et al., 2020). These membraneless organelles (MLOs) fulfil various cellular

90 functions besides splicing. Recent evidence indicates a direct role of NSs in cellular 91 transcription regulation, as their spatial proximity correlates with gene expression 92 amplification, as demonstrated by live-cell imaging of heat-shock responsive genes (Chen et 93 al., 2018; Zhang et al., 2021). HIV, being a virus capable of generating new particles through 94 splicing and integrating into active host genes, finds NSs highly conducive for viral 95 replication. HIV particles, along with their RNA genome, accumulate within nuclear MLOs 96 enriched in NS factors (Rensen et al., 2021; Scoca et al., 2023). Notably, CPSF6, a 97 paraspeckle factor first identified as a viral partner by KewalRamani's laboratory (Lee et al., 98 2010), has been clearly detected in HIV-induced CPSF6 puncta (Francis et al., 2020; Lee et 99 al., 2010; Luchsinger et al., 2023; Rensen et al., 2021; Scoca et al., 2023). These puncta 100 serve as hubs for nuclear reverse transcription (RT) and the formation of pre-integration 101 complexes, which generate active proviruses detected outside but in close proximity to NSs 102 (Li et al., 2021; Scoca et al., 2023).

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In our study, we aim to elucidate how these HIV-induced CPSF6 form and identify the NS factors involved in their formation. CPSF6, along with NS factors, contains intrinsically disordered regions that can guide HIV-1 to the correct nuclear location for successful infection or allow the virus to remain sequestered during drug treatment, forming reservoirs. Importantly, viral reservoirs are the major bottleneck for curing the infection.

In this study, we investigate which disordered domain of CPSF6 is responsible for tracking the viral core and generating HIV-induced CPSF6 puncta in the host nucleus. Simultaneously, we elucidate the key component of NSs that, through its intrinsically disordered regions (IDRs), enable fusion with HIV-induced CPSF6 puncta, likely stabilizing them. Notably, we observed that the virus rebounds when anti-reverse transcription drugs are removed, but only if nuclear niches containing HIV, NS factors, and CPSF6 are present. If these niches are pharmacologically dismantled, viral rebound does not occur.

116 Overall, studying the biogenesis of HIV-induced nuclear niches is crucial for understanding 117 how the virus navigates and persists in the nucleus of infected cells, and for designing new 118 antiretroviral strategies.

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#### 120 Results

121 Critical Role of HIV-Induced CPSF6 Puncta in Restoring Nuclear RT After Anti-RT

122 Therapy Discontinuation.

Upon nuclear entry, HIV enhances the formation of CPSF6 clusters, where RT ends. The treatment with the reversible RT inhibitor nevirapine (NEV) can trap the viral RNA genome in these nuclear niches (Rensen et al., 2021; Scoca et al., 2023). Once NEV is removed, the trapped vRNA can resume RT entirely within the nucleus, a process we term nuclear RT . Here we demonstrate that this phenomenon is dependent on the presence of CPSF6 puncta, as their disruption by high doses of PF74 (25 µM) significantly impairs nuclear RT, shown by

the absence of luciferase expression, similar to full NEV treatment (Figure 1A-B).

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#### 131 CPSF6 FG Domain is Required for HIV-Induced Puncta Formation.

132 Formation of CPSF6 puncta upon HIV-1 infection hinges into two key events: the entry of the 133 HIV-1 core into the nucleus and the binding of CPSF6 to the HIV-1 core (Blanco-Rodriguez 134 and Di Nunzio, 2021; Blanco-Rodriguez et al., 2020; Buffone et al., 2018; Zila et al., 2021). 135 To determine the contribution of CPSF6's disordered domains for the formation of CPSF6 136 puncta upon HIV-1 infection, we correlated the binding of CPSF6 to the HIV-1 core with the 137 formation of CPSF6 puncta. To this end, we first generated CPSF6 knockout (KO) THP-1 138 cells (Figure 2A-B) to eliminate the interference from the endogenous protein, which could 139 affect the interpretation of results regarding the role of the analyzed CPSF6 domains. CPSF6 140 depletion in THP-1 cells was performed by CRISPR Cas9 technology. To completely 141 eliminate the expression of the CPSF6 gene we selected single clones by limiting dilution. 142 We identified a clone that was completely KO for CPSF6, confirmed through western blot and 143 immunofluorescence (Fig. 2A-B) and we infected this clone and the control clone with HIV. 144 CPSF6 puncta were detected only in the control-infected cells and not in the KO clone (Fig. 145 2B; Suppl. Fig.3A). The viral integrase (IN) was observed within CPSF6 puncta, consistent 146 with previous studies (Francis et al., 2020; Rensen et al., 2021; Scoca et al., 2023), but 147 absent in CPSF6 KO cells where viral IN was predominantly observed in the cytoplasm (Fig. 148 2B; Suppl. Fig.3A). Thus, we used KO cells for CPSF6 to assess the role of selected CPSF6 149 domains in HIV-induced condensates. We designed various CPSF6 deletion mutants (Figure 150 2C) to specifically assess the significance of the main disordered regions of CPSF6 protein 151 (Suppl. Figure 2A) such as, the FG motif, the low complexity regions (LCRs), and the mixed 152 charge domain (MCD), in the ability of CPSF6 to bind to the core and facilitate the formation 153 of CPSF6 puncta. We investigated the role of the FG peptide by generating a mutant that 154 exclusively lacks the FG peptide ( $\Delta$ FG). Previous *in vitro* studies have shown that the FG 155 peptide binds to the hydrophobic pocket formed between capsid hexamers (Buffone et al., 2018; Price et al., 2014). Here, we want to investigate the role of FG peptide in the context of 156 157 the protein.

158 To further explore this, we developed an alternative plasmid by expanding the FG peptide 159 deletion to include surrounding prion-like LCRs ( $\Delta$ FG  $\Delta$ LCR). These regions, outside the 160 CPSF6 context, have been identified as crucial for facilitating strong CPSF6 binding to 161 capsid lattices (Wei et al., 2022). In our study, we aim to evaluate their role within a more 162 physiological setting. Additionally, we assessed a CPSF6 variant that carries the 15-mer FG 163 peptide flanked by non-LCR sequences, such as those derived from Beta-adducin (ADD2), 164 kindly provided by Mamuka Kvaratskhelia (△LCR+ADD2). These protein segments are 165 known for their high flexibility, akin to the LCR of CPSF6. Furthermore, to elucidate the 166 contribution of the LCRs of CPSF6 in the formation of CPSF6 puncta, we generated a mutant 167 lacking both LCRs ( $\Delta$ LCR) (Figure 2C). Analysis of the MCD contribution to both the ability of 168 CPSF6 to bind to the core and formation of CPSF6 puncta was achieved by deleting the 169 MCD and adding 3 nuclear localization signals (3xNLS  $\Delta$ MCD) since the deletion of the MCD 170 results in a protein that localizes mainly into the cytoplasm (Figure 2C; Suppl. Figure 2B).

171 To correlate the ability of CPSF6 to bind to the HIV-1 core with formation of CPSF6 puncta, 172 we expressed wild type and mutant CPSF6 constructs in THP-1 cells knockout for CPSF6. 173 Subsequently, we infected these cells with HIV-1 and analyzed the presence or absence of 174 CPSF6 clusters at 24 hours post-infection. Importantly, for the imaging experiment we 175 expressed CPSF6 WT and mutants without tags to avoid the formation of aggregates that 176 could interfere with our conclusions. Our data show that HIV-induced CPSF6 clusters can 177 form extremely rarely with the deletion mutant CPSF6 ADD2ALCR and with the mutant 178 lacking the FG or both the FG peptide and the LCRs (Fig. 2D-F; Suppl. Fig.3B). However, 179 when we analyzed the role of the MCD domain in CPSF6 puncta formation, which was 180 indicated to be important for condensing CPSF6 in NS (Greig et al., 2020), comparing the 181 number of CPSF6 WT puncta induced by HIV infection with CPSF6 mutants revealed that 182 the MCD domain does not play a critical role in HIV-induced CPSF6 puncta formation (Fig. 183 2D-F). In addition, we observed that the majority of analysed CPSF6 3xNLS∆MCD puncta 184 contain vRNA inside, similar to CPSF6 WT puncta (Figure 2G), thus corroborating the lack of 185 a role for this intrinsically disordered domain in HIV-induced CPSF6 puncta. Since the NLS 186 domain from SV40, which replaces the MCD, is highly basic and could potentially induce 187 condensates, we fused CPSF6 with a non-basic NLS (PY-NLS) or removed the NLS entirely. 188 Even though these two proteins do not efficiently enter the nucleus, the few that do manage 189 to reach the nucleus can host viral particles, as evidenced by the presence of IN. Many 190 viruses are typically blocked in the cytoplasm due to the presence of these mutants that are 191 mainly cytoplasmic. However, because we used a high viral dose, the blockage in the 192 cytoplasm was not complete. As a result, the viruses that successfully entered the nucleus 193 induced the formation of clusters associated with CPSF6-deleted mutants, indicating that the

MCD is not critical for the formation of HIV-induced CPSF6 puncta (Fig.2H). Similar to the MCD, when we compared CPSF6 truncated for the LCRs with CPSF6 WT, we observed that the LCRs do not contribute to CPSF6 puncta formation. Therefore, the FG peptide alone, without the LCRs, is the only CPSF6 domain required for their formation (Fig. 2D-F; Suppl. Fig.3B).

199 Next, we tested the ability of the different CPSF6 deletion mutants for their ability to bind the 200 viral core using a previously described capsid binding assay (Selyutina et al., 2018). Wild 201 type and mutant CPSF6 proteins were expressed in human 293T cells at similar levels 202 (INPUT) (Figure 3A). Extracts containing wild type and mutant CPSF6 proteins were 203 incubated with stabilized HIV-1 capsid tubes for 1 h at 25° C in the presence of 10 µM of 204 PF74, which is a small molecule that competes with CPSF6 for binding to the hydrophobic 205 pocket formed between hexamers that constitute the viral core (Buffone et al., 2018: Price et 206 al., 2014). HIV-1 capsid stabilized tubes were washed, and the bound proteins were eluted 207 using Laemli buffer (BOUND). For every construct, the percentage of bound protein relative 208 to input in the presence or absence of PF74 is shown (Figure 3B). Our results revealed that 209 the absence of the FG peptide ( $\Delta$ FG) entirely abolished CPSF6's ability to bind to the viral 210 core. In agreement, simultaneous deletion of the FG motif and LCRs ( $\Delta$ FG  $\Delta$ LCR) resulted in 211 a construct unable to bind to the viral core. Similar outcomes were observed when the LCRs 212 were replaced with sequences derived from ADD2, even if the FG was present.

213 LCR-FG is notably more disordered than ADD2-FG, containing a high proportion of prolines 214 (48 out of 98 residues), which makes it mostly non-foldable (Figure 3C-L). Since proline is a 215 structure-disrupting residue, LRC-FG is not expected to adopt any secondary structure. In 216 contrast, ADD2-FG contains fewer prolines (15 out of 98 residues) but has many charged 217 residues. It is predicted to form two short  $\alpha$  helices and a ß strand, arranged as:  $\alpha$  helix - FG 218 - ß strand -  $\alpha$  helix. ADD2-FG may form a flexible collapsed state, as its oppositely charged 219 residues are evenly distributed, potentially allowing polyelectrostatic compaction. This 220 suggests that FG within ADD2-FG may be less accessible for the interaction with the viral 221 core's hydrophobic pocket due to its involvement in this collapsed conformational ensemble 222 (Figure 3C-G, suppl. Fig.4). This aligns with the inability of CPSF6 carrying ADD2 in place of 223 the LCRs to induce CPSF6 puncta (Figure 2F). On the other hand, the deletion of only the 224 two LCRs, while keeping the FG peptide intact, resulted in unexpected findings. The  $\Delta$ LCR 225 mutant exhibited a stronger binding affinity for the viral core when compared to the wild-type 226 protein (Figure 3B). These results suggest that the LCRs surrounding the FG motif are 227 modulating the affinity of CPSF6 to the viral core, which might be important for function. By 228 contrast, deletion of the MCD ( $\Delta$ MCD) but retention of other regions, such as the FG peptide 229 and the LCRs, demonstrated a binding affinity to the viral core similar to that of the wild-type

protein. These results suggest that the MCD domain is not involved in the binding of CPSF6to the viral core, which is not surprising since the CPSF6 (1-358), which does not have an

MCD, binds to the viral core (Lee et al., 2010).

Thus, the viral capsid, through the FG peptide of CPSF6, constitutes the scaffold of HIVinduced CPSF6 puncta.

235 In summary, our results suggest that the FG peptide is the main determinant involved in the 236 binding of CPSF6 to the viral capsid. Interestingly, our work implies that the LCRs may be 237 modulating the affinity of the FG motif for the viral core (Figure 3B). Recognition motifs that 238 mediate protein-protein interactions, such as the FG motif of CPSF6, are usually embedded 239 within longer intrinsically disordered regions that can modulate affinity of the interaction 240 (Karlsson et al., 2022). Taken together, our data show that the FG peptide coordinates both 241 the binding to the viral core and the induction of CPSF6 puncta. This coordination suggests 242 that the FG peptide plays a critical dual role in recognizing the viral capsid and facilitating the 243 cellular clustering of CPSF6, which may be part of the cellular response to viral entry.

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#### 245 HIV-induced CPSF6 mutants puncta and Nuclear Speckles.

246 The NS factor SC35, commonly used as a marker of NS, has been detected in HIV-induced 247 CPSF6 puncta (Figure 4A). In this study, we investigated whether HIV-induced CPSF6 248 mutant puncta are associated with SC35. We infected cells expressing CPSF6 wild-type 249 (WT), CPSF6 3xNLS AMCD, CPSF6 ALCR, CPSF6 AMCD, and CPSF6 PY NLS AMCD, and 250 examined whether the nuclear puncta formed by the various CPSF6 proteins associate with 251 SC35. Imaging analysis revealed no significant difference in the association of SC35 with 252 CPSF6 WT or the CPSF6 mutants (Figure 4B), confirming that the disordered domains, 253 LCRs, and MCDs are dispensable for the formation of HIV-induced CPSF6 puncta that 254 localize in the canonical nuclear niche marked by SC35.

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#### 256 Biogenesis of HIV-Induced CPSF6 Puncta carrying Nuclear Speckle factors.

257 Nuclear speckle factors, particularly those involved in their biogenesis, such as SON and 258 SRRM2 (Ilik et al., 2020), have been identified as constituents of HIV-induced CPSF6 puncta 259 . However, the specific role of NSs in these puncta remains unclear. Most of the existing 260 results have been obtained through immunofluorescence experiments conducted several 261 days post-infection. In this study, we conducted a time course experiment to investigate the 262 biogenesis of HIV-induced CPSF6 puncta, which contain NS factors. Our goal was to capture 263 the fusion event between the HIV-induced CPSF6 puncta and NS, providing insights into the 264 dynamics of how HIV manipulates host nuclear structures during infection. We hypothesized 265 that NS factors might either be recruited during the initial formation of HIV-induced CPSF6 266 puncta, shortly after the virus is released from the nuclear basket of the NPC, or later via the 267 fusion between NSs and HIV-induced CPSF6 puncta. To investigate this, we performed live 268 imaging to track CPSF6-mNeonGreen and NSs in cells expressing endogenous SRRM2 269 fused with a Halo tag, using CRISPR Paint (courtesy of Roy Parker) (Lester et al., 2021).We 270 fixed and labeled samples at different time points post-infection, ranging from 6 h.p.i. to 30 271 h.p.i. (Figure 5A, Suppl. Fig.1G). At 6 h.p.i., 27% of HIV-induced CPSF6 puncta were still 272 individual, compared to only 9% at 30 h.p.i. (Figure 5B). Concurrently, 61% of HIV-induced 273 CPSF6 puncta were fused with NS at 6 h.p.i., rising to 75% at 30 h.p.i. (Figure 5B). This 274 indicates a progressive increase in the number of HIV-induced CPSF6 puncta fusing with NS 275 over time (Figure 5C). Overall, we detected individual CPSF6 clusters (green) and NSs (red) 276 that quickly fused, confirming that this fusion occurs within the two independent puncta, 277 CPSF6 and SRRM2, rather than during the formation of HIV-induced CPSF6 puncta (Figure 278 5A-C, movies 1A, 1B).

Taken together, these results suggest that HIV-induced CPSF6 puncta first form independently of NS and later fuse with NS, causing an enlargement of NS as part of the hijacking process by HIV-1.

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## Role of SON and SRRM2 in the Fusion and Stabilization of HIV-Induced CPSF6 Puncta within Nuclear Speckles.

Macrophage-like cells, THP-1, were depleted for SON or SRRM2 using AUMsilence<sup>™</sup> ASO 285 286 technology (Gao et al., 2024; Marasca et al., 2022; Mazzeo et al., 2024; Zhang et al.). The 287 level of depletion of SON and SRRM2 was evaluated by immunofluorescence using 288 antibodies against SON and SRRM2 (Figure 6A). Both depleted cells were analyzed also for 289 the presence of NS, labelled by SC35 (Suppl. Fig.1A-E). However, recent findings suggest 290 that the primary target of the SC35 mAb is SRRM2. To confirm this, cells depleted of 291 SRRM2 were labelled with antibodies against both SRRM2 and SC35 (Suppl. Figure 5). We 292 have observed that the reduction of SRRM2 resulted in a slight decrease in the mean 293 intensity of SC35, whereas the depletion of SON (Suppl. Fig.1F) did not have the same 294 effect.

Subsequently, we infected THP-1 depleted cells for SRRM2 and SON or control cells with HIV-1 and we fixed them at 48 h post-infection for immunofluorescence. We calculated the percentage of CPSF6 clusters in HIV infected THP-1 control cells (approximately 78%) and in HIV infected THP-1 cells depleted for SRRM2 (about 43%) and SON (around 66%). The results indicate that the partial depletion of SRRM2 affects the formation of HIV-induced puncta, while the depletion of SON slightly reduces their establishment (Figure 6B).

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### The IDR of SRRM2 is a crucial element for the fusion of HIV-Induced CPSF6 puncta to the nuclear speckles.

304 CPSF6 contains several disordered regions (Di Nunzio et al., 2023) as well as NS 305 factors(llik et al., 2020). Previous studies have established the role of IDR of SRRM2 in the 306 biogenesis of NS (llik et al., 2020).

Here, we investigate the role of IDRs within NS factors, with a specific emphasis on SRRM2, in the fusion and stabilization of HIV-induced CPSF6 clusters. To address this inquiry, we utilized previously published HEK293 cell lines generated using the CRISPaint system (Lester et al., 2021), comprising two distinct lines: HaloTag SRRM2, SRRM2 full-length (FL) (1-2748 aa) fused with halo tag(insertion at amino acid [aa] 2,708), and a cell line lacking the C-terminal IDR of SRRM2, known as  $\triangle$ IDR HaloTag SRRM2(1-429 aa, with halo insertion at aa 430).

As anticipated, the truncated form of SRRM2 displayed a more diffuse distribution within the nucleus, without recruitment to NS (suppl. Fig.1A-E), and consequently lacked nuclear puncta (Figure 6C). On the other hand, the number of SON puncta was highly similar between the two cell lines (Figure 6D).

Subsequently, we quantified the formation of CPSF6 puncta in both cell lines infected with HIV-1. No significant difference in HIV-induced CPSF6 puncta formation was observed between HEK293 and HEK293 carrying the SRRM2 halo tag (~27%). However, a substantial reduction in HIV-induced CPSF6 puncta was evident in the cell line carrying the SRRM2 form that lacks the IDR (~11%) (Figure 6E; Suppl. Fig.1). Collectively, these results underscore the pivotal role of the SRRM2 IDR in the stabilization of HIV-induced CPSF6 puncta through their fusion with NSs (Figure 5A-C).

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#### 328 Discussion

329 HIV-1 capsid has transformed long-standing assumptions in the field. Previously considered 330 an undruggable viral target due to the belief that it disassembled shortly after HIV entered 331 target cells, whereas recent findings have revealed its essential role in nuclear import (Ay 332 and Di Nunzio, 2023; Blanco-Rodriguez and Di Nunzio, 2021; Blanco-Rodriguez et al., 2020; 333 Chen et al., 2016; Taylor and Fassati, 2024; Yamashita and Emerman, 2004; Zila et al., 334 2021). Notably, the first-in-class antiretroviral capsid inhibitor, Lenacapavir, has shown 335 remarkable results in patients, demonstrating that the capsid, contrary to previous belief, can 336 indeed be a viable therapeutic target. While Lenacapavir improves patients' quality of life with 337 only two injections per year (Link et al., 2020; Segal-Maurer et al., 2022), no current 338 antiretroviral drugs provide a cure. This may be due to our incomplete understanding of 339 certain aspects of HIV biology. In this study, we shed light on the post-nuclear entry steps, a 340 critical phase for the establishment of viral reservoirs, which represent the main barrier to a 341 cure. Recent findings have shown that not only the HIV viral capsid translocates through the 342 NPC, but that the viral nuclear entry also enhances the formation of CPSF6 puncta. 343 Additionally, it has been revealed that RT is completed within the nucleus (Burdick et al., 344 2020; Dharan et al., 2020; Rensen et al., 2021; Scoca et al., 2023; Selyutina et al., 2020). 345 Furthermore, incoming viral RNA has been observed to be sequestered in nuclear niches in 346 cells treated with the reversible reverse transcriptase inhibitor, NEV. When macrophage-like 347 cells are infected in the presence of NEV, the incoming viral RNA is held within the nucleus 348 (Rensen et al., 2021; Scoca et al., 2023). This scenario is comparable to what is observed in 349 patients undergoing antiretroviral therapy. Interestingly, we found that if CPSF6 puncta are 350 pharmacologically dismantled, the nuclear reverse transcription cannot be restored, 351 indicating that HIV-induced CPSF6 puncta play a crucial role in the viral life cycle, particularly 352 for their potential role in forming viral reservoirs. Understanding the biogenesis of these 353 puncta could be a significant step towards deepening our knowledge of HIV biology and 354 providing additional tools to combat this pandemic virus.

355 Here we identify the disordered FG peptide essential for the binding with the viral capsid as 356 the inducer of HIV-induced CPSF6 puncta. Notably, CPSF6 protein lacking the FG peptide is 357 incapable of forming nuclear puncta. Moreover, we discovered that the two major intrinsically 358 disordered regions of CPSF6, the MCDs and the LCRs, are dispensable for the formation of 359 viral nuclear puncta. The MCDs of CPSF6 have been shown to provide cohesion for NS 360 condensation (Greig et al., 2020). In the context of HIV infection, the viral capsid induces the 361 formation of CPSF6 puncta that are depleted of MCDs carrying the viral RNA genome. These 362 CPSF6 puncta are highly similar to those formed by wild-type CPSF6, suggesting that MCDs 363 do not play a significant role in this process. Additionally, the LCRs of CPSF6 also do not 364 appear to influence puncta formation, as their depletion does not reduce the number of 365 CPSF6 puncta. This indicates that neither the MCDs nor the LCRs are involved in CPSF6 366 puncta formation during HIV infection. Surprisingly, when we assessed the ability of CPSF6 367 domains to bind to the viral capsid, we observed that the deletion of LCRs increases 368 CPSF6's ability to bind to the viral capsid. We hypothesize that a change in charges may 369 alter the binding mechanism of CPSF6 when LCRs are absent. In scenarios where the FG 370 motif is depleted, we observed a dramatic inhibition of HIV-induced CPSF6 puncta formation 371 and a lack of binding to the viral core in vitro. On the flip side, the linkage between CPSF6 372 entities is facilitated by the FG peptides' interaction with certain hydrophobic CA pockets 373 along adjoining hexamers (Wei et al., 2022). Therefore, it's conceivable that FG peptides, not 374 involved in the capsid's binding, could coalesce similarly to FG-Nups. These undergo phase 375 separation, forming condensates with nuclear pore complex-like permeability barrier features 376 (Hülsmann et al., 2012).

377 We have also identified HIV-induced CPSF6 puncta formation independent of NS at early 378 stages post-infection, with a progressive increase in CPSF6 puncta co-localizing with NS 379 over time. These results were obtained by focusing our studies on two scaffold proteins 380 involved in NS biogenesis: SON and SRRM2. Over the past approximately 0.6–1.2 billion 381 years of metazoan evolution, these two factors have undergone significant lengthening, 382 unlike many other proteins involved in splicing. This extension primarily occurred within their 383 IDRs, which are commonly associated with liquid-liquid phase separation (LLPS) and the 384 formation of biomolecular condensates (Rai et al., 2018). Co-depleting SRRM2 with SON, or 385 depleting SON in a cell line where it is deleted the intrinsically disordered C-terminus of 386 SRRM2, abolished the formation of NSs (Ilik et al., 2020). However, the depletion of only one 387 factor does not abolish NSs. Consistent with this finding, we observed that the depletion of 388 SRRM2 does not affect the presence of SON nuclear clusters. Similar results were obtained 389 with cells genetically modified to express the truncated form of SRRM2 lacking the IDRs. 390 When the *\DR* SRRM2-halo tag was detected by the halo ligand, there was no recruitment 391 of the truncated SRRM2 form in NSs. However, if SC35 is used as a target for antibodies, it 392 can still be detected, albeit with a much lower intensity signal than in cells expressing the full-393 length SRRM2. Additionally, we observed a significant reduction in the detection of HIV-394 induced CPSF6 puncta. The few CPSF6 clusters detected in these cells colocalized with the 395 weak SC35 signal. These results suggest that the IDR of SRRM2 plays an important role in 396 HIV-induced CPSF6 puncta stabilization, but rare clusters can appear, likely induced by 397 redundant NS factors. This indicates that further investigation is needed to better understand 398 the hijacking of NS by HIV.

Taken together, our results reveal the intricate interplay between individual CPSF6 domains and the viral capsid in dictating the nuclear fate of the virus. Concurrently, the IDR domain of SRRM2 contributes to the enlargement of particular nuclear speckles with the fusion of HIVinduced CPSF6 puncta. Lastly, these HIV-induced CPSF6 puncta necessitate of FG peptide to engage the viral core.

404 Overall, this study could provide insights into the understanding of viral invasion and 405 persistence within the host.

- 406
- 407
- 408 Materials and methods
- 409

#### 410 Cell lines

411 THP-1 cells are immortalized monocytic cells, which, once seeded, differentiate into 412 macrophage-like cells under phorbol 12-myristate 13-acetate (PMA) treatment (160 nM). 413 THP-1 cells were also engineered knocking out CPSF6. These cell populations These cells 414 were cultivated in RPMI 1640 medium supplemented with 10% fetal bovine serum (FBS) and 415 1% penicillin-streptomycin solution (100 U/ml). HEK293T cells are human embryonic kidney cells used to produce LVs. For figure 2, we used two engineered HEK293 strains, Halo 416 417 tagged SRRM2 HEK293 cells and Halo tagged SRRM2 ∆IDR HEK293 cells, kind gifts from 418 Roy Parker's lab (Lester et al., 2021), characterized by the Halo tagged SRRM2 protein. In 419 Halo tagged SRRM2 ∆IDR HEK293 cells Parker's lab deleted also the SRRM2 sequence 420 encoding for aa 430-2748. All the HEK293 strains were cultivated in Dulbecco's modified 421 Eagle medium supplemented with 10% FBS and 1% penicillin-streptomycin (100 U/ml).

422

#### 423 Bacteria strains

All E. coli bacteria strains were grown in Luria-Bertani (LB) medium at 37°C. DHα competent
cells and Stellar Competent Cells were used for molecular cloning, while E.coli One-Shot
BL21star (DE3) cells were exploited for protein production.

427

#### 428 Plasmids

To express the WT CPSF6/WT CPSF6-mNeonGreen and the mutant CPSF6/mutant CPSF6mNeonGreen clones, the correspondent coding sequences were engineered in pSICO plasmids. The two original plasmids used were pSICO CPSF6-mNeonGreen and pLPCX

432 CPSF6 ADD2, gift from Mamuka's lab. HIV-1 $\Delta$ EnvINHA $\Delta$ Nef plasmid encodes the  $\Delta$ EnvHIV-433 1 LAI (BRU) viral genome where the IN protein is fused to the HA tag while pNL4.3  $\Delta$ env 434  $\Delta$ Nef IRES GFP plasmid encodes the  $\Delta$ EnvHIV-1 NL4.3 viral genome and contains also a 435 GFP sequence headed by an IRES. The pNL4.3  $\Delta$ env  $\Delta$ NefLuc has the Luciferase cDNA as 436 reporter gene.

437

#### 438 CRISPR-Cas9 knockout in THP-1 cells

439 To target CPSF6, three different crRNAs were used simultaneously (specific sequence: 5'-440 TCGGGCAAATGGCCAGTCAAAGG-3', 5'-AGGACGGGGCCGTTTTCCAGGGG-3', and 5'-441 CATGTAATCTCGGTCTTCTGGGG-3', all ordered from Integrated DNA Technologies, IDT). 442 Pre-designed unspecific crRNA was used as control (IDT). crRNA and tracrRNA were 443 resuspended in IDT Duplex Buffer according to the manufacturer's instructions. On the day of 444 the nucleofection, duplexes were formed by mixing equimolar concentration of crRNA and tracrRNA, followed by 5-min annealing at 95°C. RNA duplexes were then mixed (1:2) with 445 TrueCut<sup>™</sup> Cas9 Protein v2 for 10 min at RT to generate ribonucleoprotein (RNP) complexes. 446 2×10<sup>5</sup> THP1 cells were resuspended in P3 Primary Cell Nucleofector<sup>™</sup> Solution, mixed with 447 RNP and Alt-R<sup>®</sup> Cas9 Electroporation Enhancer (90 pmol, IDT), and nucleofected in a 4D-448 Nucleofector<sup>™</sup> System using the P3 Primary Cell 4D-Nucleofector<sup>™</sup>X Kit S (program FI-110). 449 450 After nucleofection, cells were seeded in complete RPMI medium with 20% FBS. Three days 451 after nucleofection, cells were plated for clonal selection.

452

#### 453 Clonal selection of KO cell lines

454 Seventy-two hours post nucleofection, cells were diluted in RPMI medium containing 20% 455 FBS and plated in five 96-well plates at 1 and 5 cells/well condition. After one-month, 456 selected microcolonies (50-100) are placed into 24- wells plate. Once that wells were near 457 confluence, cells were transferred into the well of a 6- well plate. After growing for another 458 one-month, cells were proceeded for western blot.

459

#### 460 Western blot

Proteins were extracted on ice from THP-1 cells using RIPA buffer (20 mM HEPES, pH 7.6,
150 mM NaCl, 1% sodium deoxycholate, 1% Nonidet P-40, 0.1% SDS, 2mM EDTA,
complete protease inhibitor), and protein concentration was quantified using a detergentcompatible (DC) protein assay (Pierce<sup>™</sup> BCA Protein Assay Kit) with bovine serum albumin

(BSA) as a standard. 90 µg of total protein lysate were loaded onto SDS-PAGE 4- 12% BisTris gels (Invitrogen); an Ab rabbit anti-CPSF6 (1:500) and an anti-rabbit HRP-conjugated
(1:5000) were used for the detection of CPSF6, whereas the normalization was done by an
Ab anti- actin HRP-conjugated (1:3000). Visualization was carried out using an ECL solution .

#### 469 **AUMsilence<sup>™</sup> ASO**

470 AUMsilenceTM 352 ASOs were synthesized by AUM BioTech, LLC (Philadelphia, USA). 471 THP-1 negative control, THP-1 KD SRRM2 and THP-1 KD SON cells were differentiated with 472 PMA (160 nM) for 48 h then incubated 72 h with 10  $\mu$ M of a AUMsilenceTM ASOS 473 complementary to the mRNA of SRRM2 and SON, respectively (scramble control 474 AUMscrambleTM ). All cells were kept in incubator at 37°C and 5% CO2. These cells were 475 cultivated in RPMI 1640 medium supplemented with 10% fetal bovine serum (FBS) and 1% 476 penicillin–streptomycin solution (100 U/ml).

#### 477 Cloning

pSICO CPSF6-mNeonGreen was used to generate deletion mutants by exceeding different regions from the original sequence according to Table2. The mutants were produced with and without the mNeonGreen tag, except for pSICO CPSF6- $\Delta$ MCD and pSICO CPSF6- $\Delta$ MCD PY NLS. All primers are specified in Table1.

482 For pSICO CPSF6-mNeonGreen & LCR ADD2, CPSF6 & LCR ADD2 sequence was amplified 483 by PCR from pLPCX CPSF6 ADD2 and specific primers were designed to add also BamHI 484 restriction sites at the extremities. Phusion Flash High-Fidelity PCR Master Mix was used 485 and the reaction was performed according to the manufacture datasheet (100 ng template 486 DNA and 56° annealing temperature). PCR products were treated with DpnI for 1h at 37°C 487 and then digested with BamHI restriction enzyme at 37°C for 1h. 2 ul of the backbone pSICO 488 CPSF6-mNeonGreen were digested with BamHI at 37°C for 3h and further treated with CIP. After gel extraction of the DNA and purification, insert and backbone were ligated with T4 489 490 Ligase for 2 h at 22°C. 5 ul of the product were used to transform 50 ul of DH5 $\alpha$  bacteria (30 min at 4°C, 45 sec at 42°C, 2 min at 4°C, incubation in SOC medium for 1h at 37°C and 491 492 plating on LB agar dishes).

All the other mutants were obtained using In-Fusion®Snap Assembly protocol, which allowed
to amplify the original plasmid deleting the small region of interest. The different primers used
are reported in Table1. As reported in the In-Fusion®Snap Assembly protocol
(https://www.takarabio.com/documents/User%20Manual/In/InFusion%20Snap%20Assembly
%20User%20Manual.pdf), 5 ng of the original plasmid were amplified with PrimeSTARMax
DNA polymerase in 35 PCR cycles (10 sec at 98°C, 15 sec at 55°C, 5sec/kb at 72°C). The
PCR products were then digested for 1h at 37°C using DpnI to get read of the original

plasmid. After plasmid cleaning up, it was circularized through a ligation of 15 min at 50°C.

501 2.5 ul of products were used in Stellar™ Competent Cells' transformation, following the same

502 procedure used for DH5 $\alpha$  transformation, previously explained.

#### 503 Lentiviral vectors and viral productions

504 LVs and HIV-1 viruses were produced by transient transfection of HEK293T cells through 505 calcium chloride co-precipitation. Co-transfection was performed as follows: for LVs, 10 µg of 506 transfer vector, 10 µg of packaging plasmid (gag-pol-tat-rev), and 2.5 µg of pHCMV-VSV-G 507 envelope plasmid; for VSV- HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef-VPX viruses and VSV-pNL4.3  $\Delta$ env  $\Delta$ Nef 508 IRES GFP-VPX, 10  $\mu$ g HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef plasmid or pNL4.3  $\Delta$ env  $\Delta$ Nef IRES GFP, 2.5  $\mu$ g 509 of pHCMV-VSV-G plasmid and 3 ug of SIVMAC Vpx (Durand et al., 2013). After the 510 collection of the supernatant 48 h post-transfection, lentiviral particles were concentrated by 511 ultracentrifugation for 1 h at 22000 rpm at 4°C and stored at -80°C. LVs and viruses were 512 tittered by qPCR in HEK293T cells 3 days post-transduction.

513

#### 514 Cell transduction and infection

515 THP-1 ctrl CRISPR clone 2 cells and THP-1 (duplex1-2-3 CRISPR) KO clone 4 cells were 516 differentiated with PMA (160 nM) for 72 h then transduced with different mutants of CPSF6 517 for 72h (MOI=1) and then infected for 30h with HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef Vpx (MOI=10) in 518 presence of Nevirapine (10uM). The medium was always supplemented with PMA (160 nM) .

For Halo tagged SRRM2 HEK 293 cells and Halo tagged SRRM2 ΔIDR HEK 293 cells,  $2 \times 10^5$  cells were seeded on coverslips coated with polylysine in complete growth medium (DMEM, GlutaMAX<sup>TM</sup>-I, 10% FBS, and 1% P/S) and incubated at 37°C (5% CO<sub>2</sub>) for 24 h. Cells were then infected with the HIV-1ΔEnvIN<sub>HA</sub>ΔNef (LAI) Bru (MOI 10) in complete growth medium supplemented with Nevirapine (10µM) for 24h.

THP-1 control (scramble), THP-1 KD SRRM2 Cells and THP-1 KD SON cells were seeded on coverslips and differentiated with PMA (160 nM) for 72 h. Then incubated 48 h with 10  $\mu$ M of a FANA ASOs (scramble control FANA (SCR-FANA), SRRM2-FANA and SON-FANA). ASOs used in this study were designed and synthesized by AUM LifeTech (Philadelphia, PA, USA). Next, cells were infected with the pNL4.3  $\Delta$ env  $\Delta$ Nef IRES GFP-VPX (MOI 25) in complete growth medium and incubated at 37°C in 5% CO<sub>2</sub> for 4 days.

In the four timepoints experiment, THP-1 cells were differentiated with PMA (160 nM) for 72 h. Cells were then infected with HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef Vpx (MOI=10) in presence of Nevirapine (10uM), in complete growth medium supplemented with PMA (160 nM), and incubated at 37°C in 5% CO<sub>2</sub> for 6 h, 9 h, 12 h, 30 h post infection.

#### 534

#### 535 Immunofluorescense Microscopy

#### 536 Immunostaining

537 On the day of fixation, the cells were washed with PBS and fixed with 4% PFA for 15 min. 538 Cells were treated with glycine 0.15% for 10 min, permeabilized with 0.5% Triton X-100 for 539 30 min, and blocked with 1% BSA for 30 min. All antibody incubations were carried out at 540 room temperature in a dark humid chamber, for 1 h with primary antibodies and for 45 min 541 with secondary antibodies. Washes between antibody incubations and antibody dilutions 542 were done in 1% BSA.

Primary antibodies were diluted as follows: anti-HA 1:500, anti-CPSF6 1:400, anti-SC35 1:200, anti-SON 1:200, anti-SRRM2 1:200, Secondary antibodies used were goat anti-rabbit Alexa-488 1:300 and donkey anti-Rabbit Cy3 1:1000 for CPSF6, SON and SRRM2, goat anti-rat Alexa-647 1:100 for IN-HA, goat anti-mouse Alexa-647 1:300 or goat anti-mouse Alexa-488 1:300 for SC35, donkey anti-mouse Alexa-647 1:300 for SRRM2.

548 Finally, cells were stained with Hoechst 33342 1:10000 for 5 min. Coverslips were mounted 549 on glass slides (Star Frost) with Prolong Diamond Antifade Mountant.

#### 550 HaloTag Labelling

551 To detect the HaloTag in Halo tagged SRRM2 HEK 293 cells and Halo tagged SRRM2 ∆IDR 552 HEK 293 cells, we used HaloTag®TMR Ligand following the Technical Manual available at www.promega.com/protocols. Specifically, cells were incubated in DMEM supplemented with 553 554 HaloTag®TMR Ligand (5 µM) for 15 min at 37°C. The ligand-containing medium was then 555 removed and replaced with an equal volume of 1X PBS, repeating the step twice and ending 556 with warm complete medium. Cells were incubated in complete culture medium for 30 min at 557 37°C. The medium was then removed and replaced with an equal volume of fresh warm 558 culture medium.

#### 559 Immuno-RNA FISH

560 On the day of fixation, fixed cells were incubated in Permeabilization/Blocking buffer (1% 561 BSA, 0.3% Triton X-100, 2mM Vanadyl Ribonucleoside complexes (VRCs) in RNase-free 562 PBS) for 1 h before the antibodies' incubations. Antibodies were diluted in 563 Permeabilization/Blocking buffer. After the primary and secondary antibody staining 564 (respectively of 1h and 45 min) with the respective washes, cells were fixed for a second 565 time in PFA 4% (in RNase-free PBS) for 10 min at RT with subsequent washes with RNase-566 free PBS. In the meantime, 40 pmol of primary smiFISH probes (Tsanov et al., 2016) (24

567 smiFISH probes designed against HIV-1 pol sequence (Rensen et al., 2021) were hybridized 568 with 50 pmol of secondary FLAP probe conjugated to a Cy5 fluorophore 569 (Cy5/AATGCATGTCGACGAGGTCCGAGTGTAA/Cy5Sp/) in 1X NEBuffer 3 (diluted in 570 RNase-free  $H_2O$ ) using a thermocycler. The program setting follows: 3 min at 85°C, 3 min at 571 65°C and 5 min at 25°C. FISH-probe solution was then diluted 1:50 in Hybridization buffer 572 (90% Stellaris® RNA-FISH Hybridization Buffer, 10% Deionized Formamide). After the 573 samples were washed in Wash A buffer (20% Stellaris® RNA-FISH Wash Buffer A, 10% 574 Deionized Formamide, in RNase-free H<sub>2</sub>O) at RT for 5 min, they were placed on parafilm, 575 covered with 50 ul of FISH-probe solution in Hybridization buffer and incubated overnight at 576 37°C. The next day, cells were washed with Wash A buffer in the dark at 37°C for 30 min. Afterwards, the samples were incubated for 10 min in Hoechst 333342 diluted 1:10000 in 577 578 RNase-free H<sub>2</sub>O and then washed with Wash B buffer for 5 min at RT in the dark. Finally, the 579 cells were washed in RNase-free H<sub>2</sub>O before the coverslips were mounted on microscopy 580 slides using ProLongTM Diamond Antifade mounting medium. The mounting medium was 581 cured overnight at RT under the chemical hood and away from light.

#### 582 Images acquisition

Images were acquired using a Zeiss LSM700 confocal inverted microscope, with a 63x
objective (Plan Apochromat, oil immersion, NA = 1.4), using diode lasers at 405 nm, 488 nm,
555 nm and 639 nm for the respective fluorophores. A pixel size of 0.07 µm was used.

586

#### 587 Live imaging on HIV-1 infected Halo tagged SRRM2 HEK 293 cells

Halo tagged SRRM2 HEK293 cells were seeded and transduced with CPSF6-mNeonGreen lentiviral vector (MOI 0.5) for 24 h. 0.3 x  $10^6$  Cells were then transferred on poly-L-Lysin coated lbidi-dishes and infected with HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef (MOI = 40) in presence of Nevirapine (10 uM) for 2 h. After having changed the medium and labeled with HaloTag®TMR Ligand (5 µM) (see previous "HaloTag labelling" section) and Hoechst 333342 diluted 1:80000 in complete DMEM medium, 4D movies were acquired.

3D movies were acquired using a Nikon Ti2-E Confocal Inverted Spinning Disk microscope,
using a 63x objective (Plan Apochromat, oil immersion, NA = 1.4) and a sCMOS Hamamatsu
camera, Orca Flash 4. Pixel size 6.5 μm, 2048 x 2044 pixel, QE 82%. For the Z stuck a Z

piezo stage was used, with 0.33 µm interval. For live imaging acquisitions, cells were placed
in an environmental chamber with 37°C temperature, 5% CO<sub>2</sub> and 21% O<sub>2</sub>.

599

#### 600 **Protein expression and purification.**

601 pET-11a vectors were used to express the HIV-1 capsid protein. Point mutations, A14C and 602 E45C were introduced using the QuikChange II site-directed mutagenesis kit (Stratagene) according to the manufacturer's instructions. All proteins were expressed in Escherichia coli 603 one-shoot BL21star<sup>™</sup> (DE3) cells (Invitrogen). Briefly, LB medium was inoculated with 604 overnight cultures, which were grown at 30°C until mid log-phase (A<sub>600</sub>, 0.6-0.8). Protein 605 606 expression was induced with 1 mM isopropyl-β-d-thiogalactopyranoside (IPTG) overnight at 607 18°C. Cells were harvested by centrifugation at 5,000 x g for 10 min at 4°C, and pellets were 608 stored at -80°C until purification. Purification of capsid was carried out as follows. Pellets 609 from 2 L of bacteria were lysed by sonication (Qsonica microtip: 4420; A=45; 2 minutes; 2 610 seconds on; 2 seconds off for 12 cycles), in 40 ml of lysis buffer (50 mM Tris pH=8, 50mM 611 NaCl, 100mM β-mercaptoethanol and Complete EDTA-free protease inhibitor tablets). Cell 612 debris were removed by centrifugation at 40,000g for 20 min at 4°C. Proteins from the 613 supernatant were precipitated by incubation with 1/3 of volume of saturated ammonium 614 sulfate containing 100 mM β-mercaptoethanol for 20 min at 4°C and centrifugation at 8,000g 615 for 20 min at 4°C. Precipitated proteins were resuspended in 30 ml of buffer A (25mM MES 616 pH6.5, 100 mM β-mercaptoethanol) and sonicated 2-3 times (Qsonica microtip: 4420; A=45; 617 2 minutes; 1 second on; 2 seconds off). Sample was dialyzed 3 times in buffer A (2 h, overnight, 2 h). The sample was sonicated and diluted in 500 ml of buffer A and was 618 chromatographed sequentially on a 5 ml HiTrap<sup>™</sup> Q HP column and on a 5 ml HiTrap<sup>™</sup> SP 619 620 FF column (GE Healthcare), both pre-equilibrated with buffer A. The capsid protein was eluted from HiTrap<sup>™</sup> SP FF column using a linear gradient from 0-2 M of NaCl. Absorbance 621 622 at 280 nm was checked to take the eluted fraction that had higher protein levels. Pooled 623 fractions were dialyzed 3 times (2 h, overnight, 2 h) in storage buffer (25mM MES, 2M NaCl, 624 20 mM  $\beta$ -mercaptoethanol). Sample was concentrated using centricons to a concentration of 625 20 mg/ml and stored at -80°C.

626

#### 627 Assembly of stabilized HIV-1 capsid tubes

1 mL of monomeric capsid (3 mg/mL or 1 mg/mL) was dialyzed in SnakeSkin dialysis tubing 10,000 MWCO (Thermo Scientific) against a buffer that is high in salt and contains a reducing agent (buffer 1: 50 mM Tris, pH 8, 1 M NaCl, 100 mM β-mercaptoethanol) at 4°C for 8 hours. Subsequently the protein was dialyzed against the same buffer without the reducing agent β-mercaptoethanol (buffer 2: 50 mM Tris, pH 8, 1 M NaCl) at 4°C for 8 hours. The

absence of β-mercaptoethanol in the second dialysis allows formation of disulfide bonds
between Cysteine 14 and 43 inter-capsid monomers in the hexamer. Finally the protein is
dialyzed against buffer 3 (20 mM Tris, pH 8,0, 40 mM NaCl) at 4°C for 8 hours. Assembled
complexes were kept at 4°C up to 1 month.

637

#### 638 Capsid binding assay protocol

639 Human HEK293T cells were transfected for 24 h with a plasmid expressing the specified 640 CPSF6 variant tagged with mNeonGreen. Cell media was completely removed and cells 641 were lysed in 300 µL of capsid binding buffer (CBB: 10 mM Tris, pH 8,0, 1,5 mM MgCl2, 10 642 mM KCl) by scrapping off the plate. Cells were rotated at 4°C for 15 min and then centrifuged 643 to remove cellular debris (21,000 x g, 15 minutes, 4 °C). Cell lysates were incubated with 644 stabilized HIV-1 capsid tubes for 1 h at 25 °C. Subsequently, stabilized HIV-1 capsid tubes 645 were washed by pelleting the complexes by centrifugation at 21,000 x g for 2 min. Pellets 646 were washed using by resuspension in CBB or PBS. Pellets were resuspended in Laemmli 647 buffer 1X and analyzed by Western blotting using anti-p24 or anti-mNeonGreen antibodies.

648

#### 649 Bioinformatics analysis of CPSF6

650 Intrinsic disorder propensity of CPSF6 was evaluated using the Rapid Intrinsic Disorder 651 Analysis Online platform (RIDAO) (https://ridao.app/) designed to predict disordered residues 652 and regions in a query protein based on its amino acid sequence (Dayhoff and Uversky, 653 2022). RIDAO yields results by combining the outputs of several commonly used per-residue 654 disorder predictors, such as PONDR® VLXT (Romero et al., 2001), PONDR® VL3(Peng et 655 al., 2006), PONDR® VLS2B(Peng et al., 2005), PONDR® FIT (Xue et al., 2010), as well as 656 IUPred2 (Short) and IUPred2 (Long)(Dosztanyi et al., 2005a, b). RIDAO also computes a 657 mean disorder score for each residue based on these. In the resulting intrinsic disorder 658 profile, the disorder score of 0.5 is the threshold between order and disorder, where 659 residues/regions above 0.5 are disordered, and residues/regions below 0.5 are ordered. The 660 disorder score of 0.15 is the threshold between order and flexibility, where residues/regions 661 with the disorder scores above 0.15 but below 0.5 are flexible, and residues/regions below 662 0.15 are highly ordered.

Amino acid compositions of the intrinsically disordered C-terminal domain (residues 261-358) of human CPSF6 and its different variants (CPSF6  $\Delta$ FG  $\Delta$ LCR, CPSF6  $\Delta$ LCR, CPSF6  $\Delta$ FG, and CPSF6 ADD2  $\Delta$ LCR) were analyzed to evaluate the relative abundance of prion-like low complexity region (LCR) defining uncharged, charged, and Pro residues in these protein regions. The corresponding values of the relative abundance of these residue groups were 668 calculated by dividing numbers of prion-like LCR defining uncharged (Ala, Gly, Val, Phe, Tyr, Leu, Ile, Ser, Thr, Pro, Asn, Gln), charged (Asp, Glu, Lys, Arg), and Pro residues by total 669 670 number of amino acids in the corresponding protein fragments. As references, we used the 671 corresponding data for protein sequences deposited to the UniProtKB/Swiss-Prot database 672 that provides information on the overall distribution of amino acids in nature (Bairoch et al., 673 2005); PDB Select 25 (Berman et al., 2000), which is a subset of structures from the Protein 674 Data Bank with less than 25% sequence identity, biased towards the composition of proteins 675 amenable to crystallization studies; and DisProt 3.4 that is comprised of a set of consensus 676 sequences of experimentally determined disordered regions (Sickmeier et al., 2007). Perresidue intrinsic disorder propensities of the LCR-FG and ADD2-FG sequences were 677 678 evaluated by PONDR® VLXT (Romero et al., 2001), which is sensitive to local peculiarities of 679 the amino acid sequences freely available at http://www.pondr.com/ (accessed on August 03, 680 2024). Linear distribution of the net charge per residue (NCPR) within the LCR-FG and 681 ADD2-FG sequences were evaluated by CIDER (Holehouse et al., 2017), which is a 682 webserver for the analysis of a wide range of the physicochemical properties encoded by IDP 683 sequences freely available at http://pappulab.wustl.edu/CIDER (accessed on August 03, 684 2024). Secondary structure propensities of the LCR-FG and ADD2-FG sequences were 685 evaluated by PSIPRED (McGuffin et al., 2000), which is a highly accurate secondary 686 structure prediction method freely available to non-commercial users at 687 http://globin.bio.warwick.ac.uk/psipred/ (accessed on August 03, 2024).

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689

#### 690 IMAGING AND STATISTICAL ANALYSIS

All images were analysed using Fiji Software. More in detail, for the count of the nuclear speckles a macro was computed to segment cellular nuclei and to select and count the nuclear speckles. For the nuclei segmentation a Gaussian Blur with sigma=2 was used and for the channel related to the nuclear speckles the set threshold was at 7000-46012 (min and max). For the counting, a "size =10 – Infinity summarize" was used for the nuclei, whereas for the nuclear speckles the size was reduced to "size=0-Infinity summarize".

697 For live imaging analysis, Arivis software was used to reconstruct the 3D movies.

All data were statistically analyzed with GraphPad Prism 9 (GraphPad Software, La Jolla
California USA, www.graphpad.com). Calculations were performed and figures were drawn
using Excel 365 or GraphPad Prism 8.0. Statistical analysis was performed, with Wilcoxon
matched paired t-tests or Mann–Whitney unpaired t-tests. Spearman correlation coefficients
(r) were calculated using GraphPad Prism.

#### 703

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#### 705 Figure legends

706 Figure 1. Role of HIV-induced CPSF6 puncta in the nuclear RT upon removal of NEV. A) A) THP-1 cells, infected with VSV-G Aenv HIV-1 (NL4.3) AR LUC (MOI 10) in presence or not 707 708 of Nevirapine (10  $\mu$ M) for 5 days, or in presence of Nevirapine (10  $\mu$ M) for 2 days and then 709 the remaining 3 days without drug or in presence of Nevirapine (10 µM) for 2 days then in 710 presence of PF74 (25 µM). Confocal microscopy images, to verify the presence of CPSF6 clusters, the cells are stained with anti-CPSF6 antibody (green). Nuclei are stained with 711 712 Hoechst (blue). Scale bar 10 µm. B) Luciferase Assay, to verify luciferase expression in the 713 aforementioned samples. Luciferase values were normalized by total proteins revealed with 714 the Bradford kit. One-way ANOVA statistical test with multiple comparison was performed (\*\*\*\*=p<0.0001; \*=p<0.05; ns=p>0.05). Data are representative of two independent 715 716 experiments.

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718 Figure 2. Role of CPSF6 Domains in HIV-Induced CPSF6 Puncta. A) Western blots 719 demonstrate CPSF6 depletion using a specific antibody against CPSF6 in THP-1 cells 720 subjected to CRISPR Cas9 methods: CRISPR Cas9 bulk (left), and CRISPR Cas9 clones 721 selected by limiting dilution (right). Each condition is normalized for actin labeling. The ratio 722 between the intensity signal of CPSF6 and actin was analyzed via ImageJ and is plotted 723 below each western blot. B) Confocal microscopy images of THP-1 ctrl CRISPR clone 2 cells (Ctrl 2) and THP-1 duplex1-2-3 CRISPR clone 4 cells (CPSF6 KO 4) infected with HIV-724 725  $1\Delta$ EnvINHA $\Delta$ Nef Vpx (LAI) Bru (MOI 10) in the presence of Nevirapine (10  $\mu$ M). The cells are stained 30 h p.i. with anti-CPSF6 antibody and anti-HA antibodies to detect HA tagged 726 integrase (IN). C) Schema of CPSF6 isoform 588aa deletion mutants. D) Confocal 727 728 microscopy images of THP-1 CPSF6 KO cells, transduced with different mutants of CPSF6, 729 infected with HIV-1 [A EnvINHA ANef Vpx (LAI) Bru (MOI 10) in presence of Nevirapine (10 730 µM). The cells are stained with CPSF6 and HA antibody 30 h p.i.. Scale bar 5µm. E) Analysis 731 of the number of CPSF6 clusters in THP-1 CPSF6 KO cells transduced with different mutants 732 of CPSF6, not infected or infected in presence of Nevirapine (10 µM) (the number of analyzed cells is shown under the x-axis). Statistical test: ordinary one-way ANOVA 733 734 (\*\*\*\*=p<0.0001; \*\*\*=p<0.001; \*=p<0.05; ns=p>0.05). F) The plot compares the number of CPSF6 clusters per cell in THP-1 CPSF6 KO cells transduced with different mutants of 735 736 CPSF6, infected with HIV-1 in the presence of Nevirapine (10 µM). Statistical test: ordinary 737 one-way ANOVA (\*\*\*\*=p<0.0001; ns=p>0.05). G) Confocal microscopy images of THP-1

738 CPSF6 KO clone 4, non-transduced and non-infected or transduced with WT CPSF6 and CPSF6 3xNLSAMCD and infected with HIV-1AEnvINHAANef Vpx (LAI) Bru (MOI 10) in 739 740 presence of Nevirapine (10 µM). Immuno-RNA FISH: the cells are stained with CPSF6 741 (green) antibody and with 24 probes against HIV-1 Pol sequence (gray)(RNA-FISH) 25 h p.i. 742 Nuclei are stained with Hoechst (blue). Scale bar 10 µm. Violin plot presenting the 743 percentage of CPSF6 clusters colocalizing with the viral RNA in THP-1 CPSF6 KO clone 4 744 cells transduced with LVs expressing CPSF6 WT or CPSF6 3xNLS∆MCD (respectively n=73 745 and n=103) and infected with HIV-1 $\Delta$ EnvIN<sub>HA</sub> $\Delta$ Nef Vpx (LAI) Bru (MOI 10) in presence of 746 Nevirapine (10 µM). A total of 198 CPSF6 WT clusters and 264 CPSF6 3xNLS∆MCD 747 clusters were counted. Statistical test: unpaired t-test, ns=p>0.05. H) Confocal microscopy images of THP-1 KO CPSF6 cells transduced with WT CPSF6 and CPSF6 [AMCD without 748 NLS, with 3xNLS or with PY NLS, respectively. Cells were differentiated for 3 days, 749 750 transduced with CPSF6 lentiviral vectors (MOI 1) for 3 days and infected for 24 h with HIV-751 1ΔEnvINHAΔNef Vpx (LAI) Bru (MOI 10) in presence of Nevirapine (10 uM) (left panels). The 752 panels on the right show transduced and uninfected cells. CPSF6 and the IN tagged with the 753 HA are labeled with anti-CPSF6 (green) and anti-HA (white) antibodies, respectively. Nuclei 754 are stained with Hoechst (blue). The arrows show CPSF6 puncta in colocalization with IN-755 HA. Scale bar 10µm.

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757 Figure 3. Evaluation of CPSF6 Deletion Mutants' Binding Capacity to the Viral Core. A) 758 Ability of wild type and mutant CPSF6 proteins to bind to the HIV-1 core. Cellular extracts 759 derived from human 293T cells expressing similar levels of the indicated CPSF6 proteins 760 (INPUT) were incubated with HIV-1 capsid stabilized tubes for 1 hour at room temperatures 761 in the presence and absence of 10 µM PF74, as described in materials and methods. As a 762 carrier control, we utilized DMSO. Subsequently, HIV-1 capsid stabilized tubes were washed, 763 and the bound proteins eluted 1X Laemmli buffer 1X. The BOUND fractions were analysed 764 by Western blotting using antibodies against neon-GFP and the HIV-1 capsid. B) Experiments were repeated at least three times and the average BOUND fraction relative to 765 766 the INPUT fraction normalized to wild type binding are shown for the different CPSF6 mutants. \*\*\* indicates a p-value < 0.001; \*\*\*\* indicates a p-value < 0.0001; and ns indicates 767 768 no significant difference as determined by unpaired t-tests. C) Physicochemical 769 characteristics of the LCR-FG and ADD2-FG sequences. Intrinsic disorder predispositions 770 evaluated by PONDR® VLXT. Position of the FR segment within the LCR-FG and ADD2-FG 771 sequences is shown as gray shaded area. D) Linear distribution of the net charge per 772 residue (NCPR) within the LCR-FG sequence evaluated by CIDER. E) Linear distribution of 773 the net charge per residue (NCPR) within the ADD2-FG sequence evaluated by CIDER. F)

774 Secondary structure propensity of the LCR-FG sequence evaluated by PSIPRED. G) 775 Secondary structure propensity of the ADD2-FG sequence evaluated by PSIPRED. H) 776 Analysis of the peculiarities of the amino acid compositions of the intrinsically disordered C-777 terminal domain (residues 261-358) of human CPSF6 and its different mutants. Relative 778 abundance of prion-like LCR defining uncharged residues in analyzed protein segments. I) 779 Relative abundance of proline residues in analyzed protein segments. L) Relative abundance 780 of charged residues in analyzed protein segments. The values were calculated by dividing 781 numbers of prion-like LCR defining uncharged (Ala, Gly, Val, Phe, Tyr, Leu, Ile, Ser, Thr, Pro, 782 Asn, Gln), Pro, and charged (Asp, Glu, Lys, Arg) residues by the total number of amino acids 783 in the respective protein fragments. Corresponding values for all protein sequences 784 deposited in the UniProtKB/Swiss-Prot database, PDB Select25, and DisProt are shown for 785 comparison.

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Figure 4. A) Epifluorescence microscopy images of both infected and non-infected 293T 787 788 cells showing the presence of CPSF6 clusters only in the infected condition. CPSF6 and 789 SC35 are labeled with anti-CPSF6 (red) and anti-SC35 (gray) antibodies, respectively. Nuclei are stained with Hoechst (blue). Scale bar 5µm. B) Confocal microscopy images of THP-1 790 791 KO CPSF6 cells, differentiated for 3 days, transduced with CPSF6 lentiviral vector (MOI 1) 792 (specifically WT CPSF6, CPSF6 ALCRs, CPSF6 AMCD with 3xNLS, without NLS, or with PY 793 NLS) for 3 days and infected for 24 h with HIV-1ΔEnvINHAΔNef Vpx (LAI) Bru (MOI 10) in 794 presence of Nevirapine (10 uM). CPSF6 and nuclear speckles were labeled with anti-CPSF6 795 (green) and anti-SC35 (red) antibodies, respectively. Nuclei are stained with Hoechst (blue). 796 Scale bar 10µm. The percentage of CPSF6 puncta associated with SC35 per field of view is 797 shown in the graph. N cells were counted in each condition and a one-way ANOVA statistical 798 test with multiple comparison was performed; ns= p value > 0.05.

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801 Figure 5. Dynamics of the HIV-induced CPSF6 puncta formation and their fusion with NSs. A) Time course of infection of THP-1, 6h.p.i., 9h.p.i., 12h.p.i., 30h.p.i. or non infected. 802 803 Cells were stained with antibodies against CPSF6 (green) and SC35 (red). B) The graph 804 shows the percentage of CPSF6 puncta associated with NS or adjacent to NS or isolated 805 from NS at different time post-infection. C) The graph shows the progression of CPSF6 806 puncta associated to NS during the time post-infection. N indicates the number of cells 807 analyzed. One-way ANOVA statistical test with multiple comparison was performed; ns= p 808 value > 0.05; \*\*\*\* indicates a p-value < 0.0001.

#### 809

810 Figure 6. Role of SRRM2 and SON in the Formation of HIV-Induced CPSF6 puncta. A) Depletion of SON and SRRM2 in THP-1 cells using AUMsilence<sup>™</sup> ASO technology. The 811 812 degree of depletion is quantified by mean intensity through immunofluorescence using 813 antibodies against SON and SRRM2, respectively. Scale bar 5µm. B) The percentage of 814 CPSF6 puncta formation is quantified by IF in THP-1 cells knocked down for SON, SRRM2, 815 and control (Ctrl) infected with HIV-1 (MOI 25) for 48 h. CPSF6 is stained with an antibody 816 against CPSF6 (green), and nuclei are stained with Hoechst (blue). The graph on the right 817 reports the percentage of CPSF6 clusters calculated from more than 100 cells. Scale bar 818 10μm. Experiments were performed at least twice. C) Confocal microscopy images of ΔIDR 819 HaloTag SRRM2 HEK293 and HaloTag SRRM2 HEK293 cells stained with the halo tag 820 ligand (red), and nuclei (blue). Scale bar 10µm. D) Confocal microscopy images of HaloTag 821 SRRM2 HEK293 and ∆IDR HaloTag SRRM2 HEK293 cells, both labeled with anti-SRRM2 822 (red) and anti-SON (gray) antibodies. Nuclei are stained with Hoechst (blue). Scale bar 823 10µm. Statistical studies are summarized in the violin plot which displays the distribution of 824 the number of SON puncta per cell in the two conditions. N cells were counted and 825 Kolmogorov Smirnov test was performed, ns= p>0.05. E) Confocal microscopy images of 826 HaloTag SRRM2 HEK293 and ∆IDR HaloTag SRRM2 HEK293 cells, either non-infected or 827 infected for 24 h with  $\Delta$ EnvHIV-1 LAI (BRU) (MOI 10) in the presence of Nevirapine (10  $\mu$ M). 828 CPSF6 and SC35 are labeled with anti-CPSF6 (red) and anti-SC35 (gray) antibodies, 829 respectively. Nuclei are stained with Hoechst (blue). The plot shows the mean ± SD of the 830 percentage of cells with CPSF6 clusters calculated in n fields of view (n=24, 29, 32); N is the 831 number of cells analyzed for each of the three different cell lines; an unpaired t-test was 832 performed, \*\*\*\*=p<0.0001, ns=p>0.05. Scale bar 10µm. Experiments were performed at list 833 twice.

834

#### 835 Movies 1 A and B. CPSF6 MLOs can form independently from SRRM2 MLOs in HEK293

**SRRM2 HaloTag cells.** Spinning Disk confocal images of HEK293 SRRM2 HaloTag cells acquired 9 hours after  $\Delta$ EnvHIV-1 LAI (BRU) Vpx (MOI 10) infection in presence of Nevirapine (10  $\mu$ M). Cells were previously transduced for 24h with CPSF6-mNeonGreen lentiviral vector (MOI 0.5) to detect CPSF6 and SRRM2 was visualized incubating cells with the TMR-halo tag ligand. The nuclei were stained with Hoechst. Acquisitions were performed continuously for 27 minutes.

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### 843 Supplementary Figures

844 Suppl. Figure 1. Multiple examples (A-E) of confocal microscopy images of HaloTag SRRM2 845 HEK293 and ∆IDR HaloTag SRRM2 HEK293 cells, either non-infected or infected for 24 h 846 with  $\Delta$ EnvHIV-1 LAI (BRU) (MOI 10) in the presence of Nevirapine (10  $\mu$ M). CPSF6 and 847 SC35 are labeled with anti-CPSF6 (red) and anti-SC35 (gray) antibodies, respectively. Nuclei 848 are stained with Hoechst (blue). Scale bar 10µm. F) HEK 293 HaloTag SRRM2 or AIDR were 849 labelled with Halo ligand (red) and an anti-SON antibody (gray), nuclei are stained by 850 Hoechst (blue). Scale bar 10µm. G) THP-1 infected with ΔEnvHIV-1 LAI (BRU) Vpx were 851 labelled with specific antibodies against CPSF6 (green) and SC35 (red), nuclei are stained 852 with Hoechst (blue). Scale bar 10µm.

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Suppl. Figure 2. A) Per-residue intrinsic disorder propensity of the CPSF6 isoform 588 aa 854 855 evaluated by the Rapid Intrinsic Disorder Analysis Online platform (RIDAO) (Dayhoff and 856 Uversky, 2022) that yields results for IU-Pred\_short (yellow line), IUPred\_long (blue line), PONDR<sup>®</sup> VL3 (green line), PONDR<sup>®</sup> VLXT (black line), PONDR<sup>®</sup> VSL2 (red line), and 857 858 PONDR<sup>®</sup> FIT (pink line) and computes a mean disorder score for each residue based on 859 these predictors (MDP, thick, dark pink, dashed line). Light pink shadow represents MDP 860 error distribution. The thin black line at the disorder score of 0.5 is the threshold between 861 order and disorder, where residues/regions with disorder scores above 0.5 are disordered, 862 and residues/regions below 0.5 are ordered. The dashed line at the disorder score of 0.15 is 863 the threshold between order and flexibility, where residues/regions above 0.15 are flexible, 864 and residues/regions below 0.15 are highly ordered (upper). Schema of the deletion mutants 865 of CPSF6 (bottom). B) Lentiviral Vector Transduction of PMA-Differentiated THP-1 Cells 866 expressing CPSF6 AMCD fused to mNeonGreen (left), CPSF6 NLSAMCD fused to 867 mNeonGreen (center), CPSF6 3xNLSAMCD fused to mNeonGreen (right). CPSF6 is 868 represented in green, and nuclei are stained in blue. Scale bar 5µm. C) Lentiviral Vector 869 Transduction of PMA-Differentiated THP-1 cells expressing CPSF6 deletion mutants fused to 870 mNeonGreen. CPSF6 is represented in green, and nuclei are stained in blue. Scale bar 5µm.

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**Suppl. Figure 3. A)** Multiple examples of confocal microscopy images of THP-1 ctrl CRISPR clone 2 cells and THP-1 CPSF6 KO 4 cells infected with HIV-1 $\Delta$ EnvINHA $\Delta$ Nef Vpx (LAI) Bru (MOI 10) in the presence of Nevirapine (10  $\mu$ M). The cells are stained 30 h p.i. with CPSF6 (green) and HA (red) antibodies to detect integrase (IN). **B)** Multiple examples of THP-1 CPSF6 KO clone 4 cells transduced with different LVs carrying CPSF6 WT or mutants and stained with CPSF6 and HA antibody 30 h p.i.. Scale bar 5 $\mu$ m.

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879 Suppl. Figure 4. Sequences of FG and LCRs or substituted amino acids sequences880 analyzed in figure 5.

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- 882 Suppl. Figure 5. SRRM2 depletion is confirmed using antibodies against both SRRM2 and
- 883 SC35 by IF. Statistical analysis: One-Way ANOVA (\*\*\*\*=p<0.0001; \*=p<0.05; ns=p>0.05).
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#### 885 ACKNOWLEDGMENTS

F.D.N. is supported by the Institut Pasteur and ANRS grants (ECTZ192036, ECTZ137593),
ANR-PRCI grant, Sidaction grant. C.T. is supported by fellowships Sidaction, ANR-PRCI.
S.A. is supported by ANRS fellowship ECTZ204694. F.D.-G., B.C., M.R., C.L., C.B. are
supported by NIH Grants R01AI087390 and R01AI150455. We gratefully acknowledge the
UtechS Photonic Biolmaging platform (Imagopole), C2RT at Institut Pasteur. We thank the
NIH AIDS Reagents program to support us with precious reagents and Addgene.

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1052





Β



















1 µm

Β



## **CPSF6 KO 4**











D











G









## **HIV-1** infected



## Non infected



В

Α











**SC35** 



**CPSF6** 

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HIV-1

Α

Β

## CPSF6 SC35 Merge



ns

 $CPSF6 \ \Delta LCR$ 

ΔMCD

10 µm





10 µm

**CPSF6 WT** 

HI

ר150 <sub>ר</sub> ns Percentage of CPSF6-SC35 association ns ns • • • : 50-• • • • • CPSF6 3XNI CPSF6 ALCP ANCD ANCD CPSF6 3XNI CPSF6 ALCP ANCD ANCD CPSF6 3XNI CPSF6 ALCP ANCD ANCD CPSF6 3XNI CPSF6 ALCP AND ANCD ANCD 0

Merge





## CPSF6 ΔMCD

## **CPSF6 PY NLS** ΔMCD



**HIV-1** infected

Α

## **THP-1 SON KD**

# Ctrl

Α

Β





## THP-1 SRRM2 KD

## SRRM2



5000-

4000-

3000-

2000-

1000



Mean Intensity (A.U.) of SON/nucleus









## Nuclei

10 µm

HaloTag SRRM2

HaloTag SRRM2

HEK 293 AIDR

**HEK 293** 

С

D



10 µm







10 µm

10 µm







