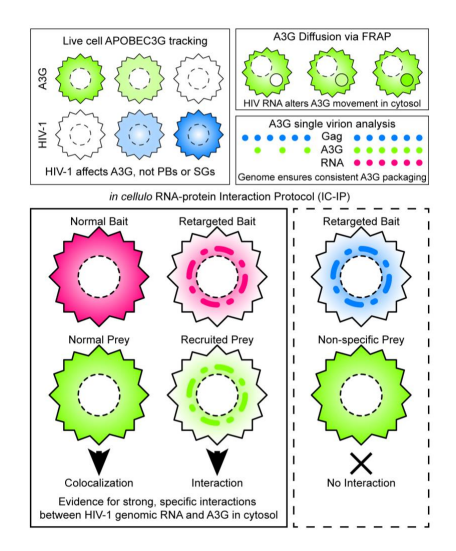
1 HIV-1 genome trafficking initiates APOBEC3G packaging in the cytosol

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17 ABSTRACT

18 HIV-1 RNA genomes interact with diverse RNA binding proteins in the cytoplasm 19 including antiviral factor APOBEC3G (A3G) that, in the absence of viral Vif proteins, is 20 packaged into virions. When and where genome-A3G interactions are initiated in the host 21 cell is unknown. Here we use quantitative long-term (>24 h) live cell fluorescence video 22 microscopy and a new in-cell RNA-protein interaction assay (the "IC-IP") to describe 23 subcellular viral and A3G trafficking behaviors over the entire HIV-1 productive phase. 24 Among other findings, we demonstrate that genome-A3G interactions are initiated in the 25 cytosol soon if not immediately after genome nuclear export; that A3G-genome 26 interactions are sufficiently strong so that tethering either factor to membranes inhibits 27 trafficking of the reciprocal binding partner; and that selective recognition of genomes 28 promotes consistent delivery of A3G to sites of virion assembly. Further elucidation of 29 RNA signature(s) detected by A3G may inform development of RNA-targeted antivirals.

30 INTRODUCTION

31 The human immunodeficiency virus type 1 (HIV-1) hijacks a diverse set of host 32 cellular RNA binding proteins (RBPs) to carry out viral RNA transcription, nuclear export. 33 translation, and trafficking₁₋₄. Select host RBPs are packaged into virions and exhibit 34 antiviral properties, with the best-characterized example being members of the 35 Apolipoprotein B mRNA editing enzyme, catalytic polypeptide-like 3 (APOBEC3) family. 36 A subset of APOBEC3 proteins (A3F, A3G, and A3H) are packaged into virions to abolish 37 infectivity by deaminating cytidines on the nascent minus-sense DNA strand of the viral 38 genome, thereby generating G-to-A mutations in the DNA provirus₅₋₈. During productive 39 infection, however, APOBEC3 proteins are counteracted by viral infectivity factor (Vif) 40 proteins that facilitate their proteasome-mediated degradation prior to the onset of virus 41 particle production₉₋₁₁.

42 How A3G proteins is delivered to virions in the absence of Vif remains a poorly 43 understood aspect of the cell-intrinsic host defense. HIV-1 virion assembly is driven by 44 viral Gag capsid proteins multimerizing on an RNA scaffold consisting of host-derived 45 RNA molecules and two dimerized viral RNA genomes, with Gag-RNA binding mediated 46 by Gag's C-terminal Nucleocapsid (NC) domain [reviewed in 12]. In the absence of viral 47 genomes, Gag expression is sufficient to drive formation of non-infectious particles 48 containing cellular RNAs, packaged in proportion to their relative abundance in the 49 cell_{13,14}. A3G's incorporation into virions has been shown, for most cell types, to be both 50 RNA- and NC-dependent₁₅₋₂₀. However, the relative contributions of genomes vs. host 51 RNAs to this process remains controversial. On the one hand, A3G-RNA binding is 52 relatively promiscuous, with A3G incorporated into virus particles even when packageable

53 genomes are not expressed^{17,19}. On the other hand, A3G incorporation levels are 54 moderately enhanced when genomes are present^{16,21}; and A3G has been shown to 55 exhibit selective RNA-binding characteristics^{19,22} including a reported preference for G-56 rich segments of the HIV-1 genome₂₀.

57 Where and when A3G interfaces with Gag and/or genomes in the cell has also been 58 under investigation for some time with conflicting results. At steady-state, A3G is 59 distributed throughout the cytosol (the aqueous phase of the cytoplasm) and accumulates 60 to high levels at non-membranous cytoplasmic sites of mRNA decay known as processing 61 bodies (PBs)_{23,24}. Cytosolic A3G is also rapidly re-localized to sites of translational 62 repression known as stress granules (SGs) in response to heat shock or oxidative 63 stress_{23,24}. Initial studies proposed a functional link between PBs and A3G's antiviral 64 activity_{23,25,26}. By contrast, more recent work has shown visible PBs to have little to no 65 discernible relevance to the HIV-1 life cycle_{27,28}.

66 The dynamic and complex nature of cytoplasmic RNA trafficking during HIV-1 virion 67 genesis may have obscured consistent prior observations made in fixed cells, 68 emphasizing a need for real-time characterization the host RBP response to HIV-1 69 infection at subcellular resolution. Accordingly, herein we set out to comprehensively 70 define the behaviors of HIV-1 RNA genomes, Gag, A3G, and additional PB and SG 71 markers over the entire viral productive phase using long-term (>24h) live cell video 72 microscopy. Our results expose single-cell A3G degradation and trafficking behaviors in 73 the presence or absence of Vif; and show that HIV-1 infection has little to no net effect on 74 cytoplasmic PBs or SG formation. Fluorescence recovery after photobleaching (FRAP) 75 and a new in-cell RNA-protein interaction assay indicate that A3G-genome interactions

76 are strong, specific, Gag-independent, and initiated in a non-localized fashion throughout 77 the cytosol long before the onset of virus particle assembly. Importantly, A3G-genome 78 interactions were sufficiently strong so that artificially tethering A3G to membranes 79 sequestered genomes in the cytoplasm and inhibited HIV-1 virus particle production. 80 Finally, using single virion analysis, we demonstrate that while genomes are not essential 81 for A3G packaging, their presence promotes a more consistent per virion delivery of A3G 82 to sites of virus particle assembly. We discuss the potential implications of A3G's selective 83 recognition of RNA in the host cytosol, and how these principles might be exploited to 84 suppress genome trafficking and virus replication in vivo.

85

86 **RESULTS**

87 Tracking HIV-1 replication using a YFP-A3G biosensor cell line. To study real-time 88 RBP responses to HIV-1 in single cells over an entire round of viral replication, we 89 generated "biosensor" HeLa cell lines stably expressing fluorescent protein-tagged 90 versions of A3G (YFP-A3G) (Fig 1a) or relevant cytoplasmic RBPs including TIA1-YFP 91 (a marker of SGs) (Fig. 2a) and CFP-DCP1A (a marker of PBs) (Fig. 2e)_{24,29}. Cells were 92 infected with either of two versions of an HIV-1NL4-3 reporter virus modified to express 93 cyan fluorescent protein (CFP) from the *nef* locus (as a marker of early viral gene 94 expression); either; (1) Vif-competent (Vif+) virus; or (2) a virus rendered Vif-minus (Vifxx) 95 due to insertion of two stop codons in the vif reading frame (see cartoon depiction in Fig 96 1b). In the absence of virus, YFP-A3G was localized to the cytoplasm and accumulated 97 in bright foci consistent with PBs, as anticipated (Fig. 1a, left panel). Induction of oxidative 98 stress using 250 µM sodium arsenite (Ars)₃₀ caused YFP-A3G to accumulate in SGs, 99 confirming an intact stress response (Fig. 1a, right panel). We also confirmed that our 100 YFP-A3G fusion protein was packaged into Vifxx virus particles and retained antiviral 101 activity using single-round assembly and infectivity assays (Fig. 1c).

In our first set of live cell imaging experiments, we infected Hela.YFP-A3G cells with Vif+ virus at a MOI of <1 in order to visualize both infected and uninfected cells side-byside in the same field. Viral gene expression (based on detection of CFP) was initiated 12-16 hours post-infection (hpi), followed by a gradual loss of YFP-A3G signal consistent with Vif-dependent YFP-A3G proteasomal degradation (decay rate of 24.5% / h +/-6.0%; n=33; Fig. 1d, Video 1, and quantification in Fig. 1e). We observed YFP-A3G to be depleted from both the diffuse cytosolic and PB pools simultaneously, with levels 109 stabilized at near-background with little to no recovery, even at the latest time points 110 (Fig.1d shows a 48h time point, with Fig.1e showing kinetics over the entire 48h time 111 course). We also observed cell rounding at the latest time points (Fig. 1d and Video 1, 112 see 48 h time point) consistent with real-time visualization of Vif-induced cell cycle 113 arrest₃₁. Taken together, these movies validated our use of YFP-A3G as a proxy sensor 114 for detecting HIV-1 infection and tracking multivariate single-cell responses to Vif. Further, 115 they exposed Vif-A3G kinetic details including: (1) that Vif's capacity to degrade A3G is 116 potent in every single infected cell; (2) that Vif operates ubiguitously throughout the 117 cytoplasm; and (3) that Vif's effects are remarkably persistent on a per cell basis, *i.e.*, the 118 YFP-A3G biosensor demonstrated no discernible fluctuations to single-cell Vif activity 119 post-YFP-A3G degradation, even over time frames of greater than 24 h.

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121 HIV-1 infection in the absence of Vif re-localizes YFP-A3G to virion assembly sites 122 at the plasma membrane. We next visualized YFP-A3G responses after HeLa.YFP-A3G 123 infection with Vif-deleted (Vifxx) virus. As expected, YFP-A3G expression was maintained 124 in the cytoplasm and at PBs throughout the entire course of infection, and cell cycle arrest 125 was not observed (Fig. 1f and Video 2). Strikingly, however, in infected cells we observed 126 YFP-A3G coalescing in large (up to 4 µm diameter) clusters at 30-48 hpi (Fig. 1f, white 127 arrows highlight growing clusters). These clusters were unlikely to be PBs or SGs based 128 on their proximity to the cell surface and real-time observations of cluster release and 129 transfer to neighboring cells (Fig. 1f, 32h time point, and tracked in Video 2). Instead, we 130 hypothesized that clusters represented recruitment of YFP-A3G from the cytoplasm to 131 sites of virus particle assembly at the plasma membrane. Consistent with this hypothesis,

132 YFP-A3G clusters stained positive for both HIV-1 Gag/Gag-Pol and viral genome but not 133 YFP-A3G mRNA (as a negative control) detected using a 4-color combined anti-p24Gag 134 immunofluorescence (IF) and RNA fluorescence in situ hybridization (FISH) protocol (Fig. 135 1g). Thin section electron microscopy also confirmed clustering of virus particles at the 136 surface of HeLa.YFP-A3G cells at 48 hpi for both the Vif+ and Vifxx conditions (Fig. 1h). 137 Based on these results, we concluded that even in the absence of Vif, YFP-A3G serves 138 as a useful proxy biosensor for detecting HIV-1 virus particle assembly through monitoring 139 YFP-A3G's virus-driven translocation from the cytoplasm to the plasma membrane.

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141 HIV-1 replication has little to no effect on PB integrity or SG formation. Because 142 HIV-1 drastically alters YFP-A3G subcellular trafficking behaviors in both the presence 143 and absence of Vif expression, we thought it important to ascertain if HIV-1 impacted the 144 net integrity of A3G's major cytoplasmic sites of accumulation, *i.e.*, SGs and PBs. To 145 address this question, we monitored Vif+ and Vifxx HIV-1 infection of two biosensor cell 146 lines engineered to stably express either YFP-TIA1 (Fig. 2a-2c) or CFP-DCP1a (Fig. 2e); 147 validated markers of PBs and SGs, respectively [reviewed in 29]. We first confirmed that 148 our YFP-TIA1 "stress" biosensor responded to stress similar to YFP-A3G (see Fig. 1a) by 149 treating cells with 250 µM Ars 30 (Fig. 2a, right panel) or infecting cells with rhinovirus A16 150 (RVA16), a positive-strand RNA virus we had studied previously₃₂ and discovered to 151 cause rampant SG formation as early as 4h post-infection (Fig. 2b and Video 3).

Unlike RVA16, neither Vif+ nor Vifxx HIV-1 triggered SG formation in either YFP-TIA1 or YFP-A3G cells over 48h of continuous imaging, a time window sufficient to encompass an entire round of viral replication (Fig. 2c with quantification in 2d). HIV-1 155 also had no discernible effects on PB number or morphology in CFP-DCP1A biosensor 156 cells; with PB number being remarkable consistent and stable for all individual cells and 157 conditions (typically 6-9 PBs per cell, Fig. 2e with quantification in 2f). Based on these 158 largely negative results, we concluded that despite HIV-1's marked effects on YFP-A3G 159 as shown in Fig. 1, net modulation of PBs or induction of SGs are unlikely to be intrinsic 160 features of the HIV-1 replication cycle.

161 Because HIV-1 Gag was recently reported to inhibit SG formation_{33,34}, we also 162 tested the effects of treating HeLa.YFP-A3G or HeLa.YFP-TIA1 biosensor cells with Ars 163 at 31 hpi when NL4-3 Gag expression in HeLa cells is relatively high (e.g., see 31). 164 Contrary to our expectations, both YFP-A3G and YFP-TIA1 accumulated in SGs in Ars-165 treated cells infected with either Vif+ or Vifxx viruses (Fig. 2g with quantification in 2h). To 166 localize Gag and genome under these conditions, we again performed combined Gag IF 167 and genome FISH, finding that the bulk of the HIV-1 genome signal co-localized with 168 YFP-A3G in Ars-induced SGs at this time point (Fig. 2i, gag-pol FISH). By contrast, the 169 cytoplasmic pool of Gag was largely excluded from these complexes (Fig. 2i, anti-p24).

SGs are induced in response to protein kinase R-mediated phosphorylation of translation initiation factor eIF2a, triggering aggregation of scaffolding proteins that include TIA1, TIAR, and G3BP1 in complex with mRNAs bound to translation initiation factors and polysome-associated proteins [reviewed in 29,35]. Accordingly, co-localization of YFP-A3G with genomes in SGs could suggest stronger recognition of genomes by YFP-A3G relative to Gag in association with free ribosomes in the cytosol.

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HIV-1 RNA genomes but not Gag modulate A3G subcellular mobility; evidence for 177 178 genome-A3G interactions in the cytosol. So that we could also monitor Gag and 179 genome trafficking dynamics in living HeLa. YFP-A3G cells, we next engineered two-color, 180 HIV-1 genomes that encoded CFP-tagged Gag from a "self-tagging" genome bearing 24 181 copies of MS2 RNA binding loop (located in the gag-pol open reading frame; ORF) and 182 expressing an RFP-tagged MS2 bacteriophage (MS2-RFP) coat protein from the viral nef 183 locus (Gag-CFP/MS2-RFP virus, see cartoon depictions in Figs. 3a and 3c). Gag-CFP 184 allowed for single cell measurements of viral late gene expression, with genomes tracked 185 based on their binding to the MS2-RFP protein. Expression of Vif+ or Vifxx Gag-186 CFP/MS2-RFP viruses recapitulated effects seen during infection with CFP reporter 187 viruses (Figs. 1 and 2); including rapid, Vif-dependent down-regulation of YFP-A3G 188 (decay rate = 25.9% per hour +/- 7.0%; n = 31, Fig. 3b, green, with quantification in Figs. 189 3e and 3f, Video 4) and, for Vif-deficient (Vifxx) conditions, co-clustering of YFP-A3G, 190 MS2-RFP-tagged genomes, and Gag-CFP to assembly sites at the plasma membrane 191 (Fig. 3d, arrows, Video 5; and see Fig. 4d for an image recorded at higher magnification). 192 We observed Vif-mediated YFP-A3G degradation to occur prior to the onset of virus 193 particle assembly (Fig. 3b, compare 2 h and 3 h time points), consistent with a prior 194 report₃₆, but at >1h after genomes had populated the cytoplasm (Fig. 3d, compare 1h and 195 4h time points).

Both prior to and during virus particle assembly, Vifxx genomes and YFP-A3G were co-distributed in a non-localized fashion throughout the cytoplasm, with YFP-A3G but not genome or Gag-CFP accumulating at PBs (Fig. 3d, 2 and 3 h time points post-detection of MS2-RFP). To test if genomes and YFP-A3G were interacting in the fluid compartment, 200 we performed fluorescence recovery after photobleaching (FRAP) analysis to measure 201 rates of YFP-A3G recovery with or without genomes and in the presence or absence of 202 Gag (Figs. 3g, 3h, and quantification in 3i). Infection with Vifxx HIV-1 mCherry reporter 203 virus reduced rates of YFP-A3G recovery ($t_{1/2} = 34.4 \text{ s} + -5.3 \text{ s}; n=8$) relative to an 204 uninfected "YFP-A3G alone" control ($t_{1/2} = 27.0 \text{ s} + -13.5 \text{ s}; n=14$). Interestingly, we also 205 observed incomplete recovery of YFP-A3G at the latest time points in infected cells (~75% 206 recovery compared to nearly 90% for untreated control cells Fig. 3i); suggesting virus-207 induced immobilization of YFP-A3G 37.

208 Expression of genomes in the absence of Gag using a "genome only" Vifxx MS2-209 RFP genome, wherein we mutated the gag start codon to abolish Gag synthesis (see Fig. 210 3g), yielded markedly slower rates of YFP-A3G recovery ($t_{1/2} = 70.8 \text{ s} + -27.6 \text{ s}; n=6$) 211 relative to both the "YFP-A3G Alone" condition, and a "Gag Only" condition wherein we 212 expressed Gag-mCherry from an mRNA modified to lack 5' and 3' viral regulatory RNA 213 sequences and codon-optimized to further reduce potential contributions of viral cis-214 acting RNA elements located in the gag ORF (COGag-mCh; $t_{1/2} = 30 \text{ s} + -8.9 \text{ s}; n=7$) 215 (Fig. 3i). Taken together, because the expression of genomes but not Gag altered YFP-216 A3G movements in the cytoplasm based on FRAP analysis, genome-YFP-A3G 217 interactions are likely initiated in the cytoplasm independently of virus particle assembly; 218 and potentially even prior to the onset of Gag synthesis.

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Genome-A3G interactions are selective, Gag-independent, and initiated prior to the
 onset of virus particle assembly. Because FRAP was unable to distinguish between
 direct or indirect genome-A3G interactions, we next developed a single cell assay to allow

223 us to measure the strength and specificity of RNA-RBP interactions in the cytoplasm head 224 on; dubbed the In-Cell RNA-Protein Interaction Protocol (IC-IP) (depicted in Fig. 4a). The 225 IC-IP was based on the principle of immobilizing genomes at unnatural sites in the 226 cytoplasm (*e.g.*, membranes or the actin cytoskeleton) using an MS2-based tethering 227 strategy that we have described previously₃₈. Should an RBP such as YFP-A3G be 228 strongly associated with the tethered genome, its localization will be also shifted with clear 229 evidence of RNA-RBP co-localization at the unnatural site of tethering.

230 For genome-A3G IC-IPs, MSL-bearing genomes served as bait for YFP-A3G, as 231 depicted in Fig. 4a. Genome retargeting was achieved by co-expressing "targeter" MS2 232 coat proteins that were modified to tether MSL-tagged genomes to either membranes 233 (using Src-MS2; bearing a 10 amino-acid membrane targeting signal) or the actin 234 cytoskeleton (using Lifeact-MS2; bearing a 17 amino acid F-actin targeting signal) (Fig. 235 4b). The Src-MS2 and Lifeact-MS2 targeters were fused to iRFP670 so that we could co-236 visualize both the targeter and the genome (labelled with the self-tagging MS2-RFP 237 "tracker" protein, see cartoon depiction in Fig. 4c) without experiencing spectral overlap 238 (see Fig. 4d).

239 Co-expression of two-color Vifxx Gag-CFP/MS2-RFP genomes with the Src-MS2-240 iRFP or Lifeact-MS2-iRFP targeters caused moderate re-localization of Gag-CFP 241 proteins from the cytosolic pool to intracellular vesicles (Fig. 4d, compare panels iv. and 242 ix.) or F-actin filaments (Fig. 4d, compare panels iv. to xiv.), respectively; consistent with 243 a subset of Gag proteins co-trafficking with its genome substrate in the cytosol (as we 244 have previously shown₃₈). Remarkably, however, re-localization of YFP-A3G was more 245 striking, with >50% of the net YFP-A3G per cell fluorescent signal now associated with 246 intracellular vesicles or F-actin filaments (Fig. 4d, compare panels ii., viii., and xiii., and 247 see YFP-A3G regions-of-interest; ROI; with quantification in 4e and 4f). Consistent with 248 our results from FRAP analyses (Figs. 3g and 3h), a "genome only" bait was sufficient to 249 recruit >50% of YFP-A3G to intracellular vesicles as directed by Src-MS2, suggesting that 250 Gag plays no role in genome-A3G cytosolic interactions (Figs. 5a and 5b). By contrast, a 251 control MSL-bearing mRNA encoding codon-optimized Gag-CFP ("Gag Only") had no 252 effect on YFP-A3G cytoplasmic distribution (Figs. 5c and 5d). Collectively, these results 253 indicated that HIV-1 genome-A3G interactions in the cytoplasm are strong, specific, and 254 occur in the presence or absence of Gag.

255 A particularly useful feature of the IC-IP is that it could be coupled to video 256 microscopy in order to determine the timing of YFP-A3G-genome interactions timed to 257 the onset of viral gene expression. Live cell imaging of cells co-expressing Src-MS2-258 iRFP670 with a "Genome Only" RNA bait demonstrated that YFP-A3G was relocalized to 259 vesicles within two hours of first detecting genome expression in the cytoplasm (Fig. 5e, 260 compare 1h and 2h time points, and see Video 6 for additional detail). Accordingly, this 261 experiment indicated that strong genome-A3G interactions are initiated concomitantly 262 with or very soon after genome nuclear export.

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Tethering A3G to membranes arrests HIV-1 genome trafficking and reduces virus particle production. We reasoned that if A3G's interactions with genomes were sufficiently strong, then tethering A3G to membranes would, as a corollary, arrest HIV-1 genome mobility in the cytosol and abolish genome trafficking to sites of virus particle assembly. To test this idea, we engineered a "reciprocal" IC-IP experiment (Fig. 6a) 269 wherein YFP-A3G was modified so that it would bind membranes due to its bearing the 270 same N-terminal Src-derived myristoylation signal we used for our Src-MS2 genome 271 retargeting constructs (Figs. 2b-2e). This fusion protein (Src-YFP-A3G) was co-272 expressed with two-color (Gag-CFP/MS2-RFP) Vifxx (Fig. 6b) or a CFP-tagged variant 273 (Src-CFP-A3G) co-expressed with one-color (MS2-RFP) "Genome Only" Vifxx genomes 274 (Fig. 6c). As would be anticipated for a strong interaction, Src-FP-A3G expression 275 induced the marked clustering of MS2-RFP-tagged genomes to perinuclear vesicles, with 276 genomes co-localizing with Src-FP-A3G (Figs. 6b and 6c). To address the more relevant 277 scenario of infected cells, we also generated a cell line constitutively expressing Src-YFP-278 A3G (HeLa.Src-YFP-A3G cells) and infected these cells with a Vifxx reporter virus (Fig. 279 6d). As expected, we observed genomes (and, to a lesser extent, Gag) co-clustering with 280 Src-YFP-A3G at cytoplasmic vesicles at 24 hpi, detected using combined IF/FISH (Fig. 281 6d, bottom, white arrows).

282 We hypothesized that arrest of genome subcellular trafficking due to Src-YFP-A3G 283 expression would reduce Gag synthesis and net virus particle production. Consistent with 284 this hypothesis, co-expression of Src-YFP-A3G or a Src-MS2 control with MSL-tagged 2-285 color genome (Gag-BFP in this experiment) yielded a dose-dependent reduction to 286 cytosolic Gag level and net efficiency of virus particle release (Fig. 6e). Taken together, 287 these experiments indicated that A3G-genome interactions in the cytosol are sufficiently 288 strong that tethering A3G to membranes can effectively repurpose this protein as an 289 inhibitor of genome trafficking and net virion production.

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291 The presence of genomes promotes more consistent per virion delivery of A3G to 292 sites of virus particle assembly. Combined, the above experiments suggested that 293 A3G has evolved to preferentially recognize one or more HIV-1 genome RNA signatures. 294 However, A3G is encapsidated by Gag into virus particles even in the absence of 295 packageable genomes₁₇₋₁₉. In an effort to better rationalize prior observations of A3G 296 genome specificity vs. promiscuity, we performed a comparative single virion analysis 297 (SVA) of A3G delivery into virus particles; based on a technique originally pioneered by 298 the Hu and Pathak groups wherein fluorescently-labeled virus-like particles (VLPs) are 299 harvested from HEK293T cells (that produce a greater quantity of particles relative to 300 HeLa) and subjected to sub-micron quantitative multicolor fluorescence imaging to 301 measure relative levels of per particle genome and/or A3G incorporation_{39,40} (Fig. 7). For 302 our analysis, we compared per particle levels of YFP-A3G encapsidation for four 303 independent Gag/genome scenarios (depicted in Fig. 7a); "Gag-Only" mRNAs encoding 304 codon-optimized Gag-CFP (COGag-CFP = "Gag-Only"), viral Vif+ or Vifxx 2-color HIV 305 Gag-CFP/MS2-RFP genomes, or mRNAs encoding codon-optimized Gag-CFP wherein 306 the NC RNA-binding domain was replaced by a leucine zipper (Δ NCzip) to abolish Gag-307 RNA binding (based on 41). As expected, YFP-A3G packaging was only observed for 308 COGag and the Vifxx HIV-1 conditions (Fig. 7b).

Using SVA, Vif+ and Vifxx viruses exhibited high efficiency MS2-RFP incorporation (~89 and ~88% of Gag-CFP particles scoring positive for MS2-RFP, respectively), very consistent with a prior report of HIV-1 genome packaging efficiency³⁹. Vifxx Gag-CFP/MS2-RFP particles exhibited a similar frequency of YFP-A3G incorporation (86% of total Gag-CFP particles) (Fig. 7c and quantification in 7d). However, for "Gag-Only" 314 (COGag-CFP) particles, the frequency of detectable YFP-A3G incorporation was lower 315 (57% efficiency), intermediate to Vifxx HIV-1 vs. the ∆NCzip "no-RNA" negative control 316 (Fig. 7c and quantification in 7d). COGag particles also exhibited reduced per virion levels 317 of YFP-A3G fluorescence when compared to full-length, Vifxx HIV (Fig. 7c and 318 quantification in 7e). In sum, these data indicated that, although genomes are not 319 essential for YFP-A3G delivery to virus particles, selective genome-A3G interactions 320 promote a more consistent and more enriched per virion delivery of A3G to virus particle 321 assembly sites.

322 DISCUSSION

Herein we studied the HIV-1 life cycle from the host cell perspective, combining several complementary live cell imaging approaches with functional assays to characterize the coordinated behaviors A3G, PB or SG proteins, and viral elements (Gag and genome) over an entire round of viral replication (summarized in Fig. S1).

327 Our YFP-A3G biosensor allowed for indirect measurements of HIV-1 replication 328 kinetics in single cells and exposed A3G behaviors including; (1) that Vif-mediated A3G 329 degradation occurs from all subcellular pools simultaneously (Fig. 1d and Movie 1); (2) 330 that A3G suppression by Vif is remarkably stable and maintained over the entire late 331 phase of infection (lasting hours to days, see Figs. 1d, Movie 1, and 1e); and (3) that A3G 332 degradation is completed prior to the onset of virus particle assembly (very consistent 333 with prior report from Holmes et al., 36 but shown here to be after genome nuclear export) 334 (Fig. 3b). In the absence of Vif, we could also monitor virus particle assembly, observing 335 large quantities of YFP-A3G being re-localized from the cytoplasm to the plasma 336 membrane during this process (Fig. 1f and Video 2). Taken together, on a technical level 337 there should be great utility in the general strategy of tracking RBP biosensors to detect 338 virus replication dynamics and study host cell responses (e.g., see also RVA16-induced 339 stress in Fig. 2b). Moreover, now validated, HeLa.YFP-A3G cells should serve as useful 340 multivariate reporter system for further dissection of the viral and host machineries 341 governing A3G degradation and packaging.

Despite HIV-1's marked effects on YFP-A3G behaviors, we observed little to no discernible effects of HIV-1 on SGs or PBs monitored using our YFP-TIA1 or CFP-DCP1A biosensors, with or without Vif expression (Fig. 2). These movies reinforce that 345 modulation of PB number or architecture or SG induction is unlikely to be an intrinsic 346 feature of the HIV-1 life cycle_{27,28}. Overall, the underlying significance of A3G's localization 347 to PBs and SGs remains is unknown; but predicted to spatially coupled A3G to RNP 348 complexes with active roles in viral mRNA surveillance (*i.e.*, PBs are sites miRNA-349 dependent mRNA decay and SGs can suppress viral translation)_{23,24,26}. In this context, 350 we speculate that all successful retroviruses and endogenous elements must adapt to 351 subvert, suppress, or avoid PB- or SG-associated activities. Indeed, HIV-142,43, other 352 retroviruses_{44,45}, and endogenous retroelements including LINEs₄₆ and yeast Ty 353 elements₄₇ are thought to exploit non-membranous RNP complexes related to PBs/SGs 354 in order to compartmentalize activities including mRNA translation and genome 355 packaging. Regarding suppression, intriguing recent work from Mouland and colleagues 356 has shown HIV-1 to actively inhibit SG formation, proposed to reduce the potential for 357 stress signaling triggered by Gag NC interacting with host RNA species_{33,34}. While we did 358 not observe overt SG suppression in this study (Figs. 2g-i), the dynamics of this process 359 warrant additional investigation and it seems reasonable to posit that retroviruses prevent 360 SG formation to promote a fluid flow of genomes from the nucleus to polysomes and then 361 on to capsid assembly sites. Overall, however, our data support the notion that HIV-1's 362 general modus operandi is to largely avoid association with visible PBs or induction of 363 SGs.

At a more molecular level, our imaging assays also suggest that A3G is able to selectively compete for genome binding in the cytosol. First, HIV-1 genomes but not Gag rapidly accumulated at YFP-A3G-enriched SGs in infected cells after Ars treatment (Fig 2i). Second, based on FRAP analysis, the presence of genomes, but not Gag, reduces 368 A3G mobility in the cytoplasm (Fig. 3g and 3h). Third, and most convincingly, our IC-IP 369 experiments demonstrated that >50% of the cytosolic YFP-A3G is recruited to 370 membranes or F-actin by Src- or Lifeact-MS2-tethered genomes, respectively, (Fig. 4f); 371 effects that were far less evident for Gag-CFP; not observed in the absence of genomes 372 (Fig. 5d); and detected soon if not immediately after genome nuclear export (Fig. 5e). As 373 to why A3G competes for genome binding, single virion analyses (Fig. 7) revealed an 374 ~90% frequency of A3G incorporation into genome-containing virions relative to "Gag-375 only" particles wherein the frequency of A3G incorporation was less than 60%. This result 376 suggests to us that, although A3G-RNA binding is, overall, relatively promiscuous 19, it has 377 evolved to preferentially target genomes in a way that ensures consistent and efficient 378 delivery to virions, largely consistent with a model recently proposed by Bieniasz and 379 colleagues wherein Gag and A3G are adapted to compete for similar RNA sequences₂₀. 380 In this way, A3G maximizes its antiviral potential and thus necessitated evolution of the 381 Vif antagonist. Endeavors to further delineate the relevant protein constituents of A3G-382 genome "surveillance" complexes, and to determine whether or not A3G exhibits 383 differential detection of genomes destined for packaging vs. translation (as gag-pol 384 mRNAs), are ongoing.

To summarize, our studies reinforce that the core, essential nature of genome trafficking in the cytosol is diffusion in dynamic association with RNP complexes, with some proteins (*e.g.*, A3G) more strongly associated with genomes than others. While we have yet to define the RNA signature(s) that specify genome detection by A3G, our observations may be informative in the context of recent CLIP-seq studies demonstrating A3G's RNA-binding preference to be relatively sequence non-specific_{19,20} coupled to 391 compelling recent work showing that HIV-1 genomes are selectively enriched in selective 392 post-transcriptional regulatory marks including N₆ methyladenosine (m6a)₄₈₋₅₀, 5-393 methylcytosine (m5c), and 2'O-methylation₅₁. Artificially tethering A3G to membranes 394 restricts HIV-1 genome trafficking and virus particle assembly (Fig. 6). Based on these 395 effects, we predict it will be feasible to design inhibitory molecules or biologicals capable 396 of achieving similar strong, selective inhibition of HIV-1 RNA trafficking, translation, and 397 genome packaging in the context of antiviral strategies.

398 ACKNOWLEDGEMENTS

399 We thank Randall Massey at the University of Wisconsin SMPH Electron Microscopy 400 Facility for assistance with EM sample preparation. We are grateful to Michael Malim 401 (King's College London), Chad Swanson, (King's College London), and Sarah Gallois-402 Montbrun (Institut Cochin) for plasmid reagents and advice. The following reagent was 403 obtained through the NIH AIDS Reagent Program, Division of AIDS, NIAID, NIH: HIV-1 404 p24 hybridoma (183-H12-5C) (from Bruce Chesebro) 52. We thank Kelly Watters, Marchel 405 Hill, and Ann Palmenberg for RVA1632. This study was supported by NIH grants 406 RO1AI110221 and U54AI150470; the Wisconsin Partnership Program New Investigator 407 Program (ID 2830); the Greater Milwaukee Foundation's Shaw Scientist Program; and 408 the UW-Madison UW2020 Program to N.M.S. J.T.B. was supported by National Science 409 Foundation Graduate Research Fellow Program grant DGE-1256259 and both a 410 Research Competition Award and a Dissertation Completion Fellowship from the UW— 411 Madison office of the Vice Chancellor of Research and Graduate Education. Any 412 opinions, findings, and conclusions or recommendations expressed in this material are 413 those of the authors and do not necessarily reflect the views of the National Science 414 Foundation. E.L.E. III received support from NIH training grant CA009135 and an 415 Advance Opportunity Fellowship from the UW-Madison SciMed/GRS program. B.E.B. 416 was supported by National Science Foundation Graduate Research Fellow Program 417 grant DGE-1256259 and NIH training grant GM008349. S.L.F. received support from a 418 UW-Madison Hilldale Undergraduate Research Fellowship.

419 AUTHOR CONTRIBUTIONS:

- 420 JTB and NMS conceived the project, designed reagents, and established methods. JTB
- 421 carried out all experiments and wrote the first version of the manuscript. ELE, BEB, SLF,
- 422 LES, and AEB contributed reagents and generated preliminary data. JTB, ELE, and NMS
- 423 processed and analyzed data. All authors contributed to the final revised manuscript and
- 424 approved submission.

425

426 METHODS

427 Cell culture, plasmids, and stable cell lines. Human HeLa and HEK293T cell lines 428 (obtained from ATCC) were cultured in DMEM (Sigma-Aldrich) supplemented with 10% 429 fetal bovine serum (heat-inactivated, filter-sterilized), 1% L-glutamine, and 1% penicillin-430 streptomycin. HeLa.YFP-A3G, HEK293T.YFP-A3G, HeLa.Src-YFP-A3G, HeLa.YFP-431 TIA1, and HeLa.CFP-DCP1a were generated as previously described₃₈. Briefly, YFP-432 A3G, Src-YFP-A3G, YFP-TIA1, and CFP-DCP1a cDNA₂₄ were inserted into a MIGR1-433 derived retroviral vector (pCMS28) upstream of sequence encoding an internal ribosome 434 entry site (IRES) and a second reading frame encoding Puromycin-N-acetyltransferase48. 435 High performance clones were selected by limiting dilution and maintained in 2µg/mL 436 puromycin. YFP, YFP-A3G, and Src-YFP-A3G as well as CFP versions of construct were 437 also generated using the pcDNA3.1 transient expression vector backbone. HIV-1 reporter 438 virus plasmids were derived from the pNL4-3 molecular clone₅₃ bearing inactivating 439 mutations in *env*, *vpr*, and expressing a green fluorescent protein (GFP) reporter from the 440 nef locus (E-R-/GFP)54; with the GFP reporter replaced with either cerulean fluorescent 441 protein (CFP) or mCherry (mCh). Vif-minus HIV-1 reporter virus plasmids (Vifxx CFP and 442 Vifxx mCherry) were generated by changing vif codons 26 and 27 to stop codons55 using 443 overlapping PCR and inserted into Vif+ CFP and Vif+ mCherry plasmids. Two-color 444 fluorescent HIV-1 reporter viruses (Gag-CFP/MS2-RFP) were generated by replacing the 445 gag reading frame in Vif+ and Vifxx CFP reporter viruses with gag-CFP just upstream of 446 a cassette encoding 24 copies of the MS2 bacteriophage RNA stem loop (MSL, a kind 447 gift of Robert Singer, Albert Einstein University, New York, NY, USA)56, and then 448 subsequently replacing CFP with sequence encoding the MS2 coat protein fused to

449 mApple or mCherry and bearing an SV40 nuclear localization signal (MS2-RFP). Gag 450 was also fused to mTagBFP2 for select experiments (e.g., Fig. 6e) in order to avoid 451 CFP/YFP cross-detection when immunoblotting for Src-YFP-A3G. Mutated versions of 452 full-length HIV-1 were generated using overlapping PCR as previously described₃₈. Src-453 MS2-CFP, Src-MS2-RFP, and Src-MS2-iRFP targeting constructs encoding an amino-454 terminal membrane targeting signal derived from the Src kinase (MGSSKSKPKD) were 455 generated by overlapping PCR and subcloned into pcDNA3.1. mTurguoise2 (CFP) and 456 mApple (RFP) were gifts of Michael Davidson (Addgene # 54843 and 54747, 457 respectively). iRFP670 cDNA₅₇ was amplified from the ColorfulCell expression plasmid, 458 a gift of Pierre Neveu (Addgene plasmid # 62449)₅₈. mTagBFP2₅₉ was a gift of Michal 459 Davidson (Addgene plasmid # 55302).

460

461 Retroviral assembly and infectivity assays. Cells at 30-40% confluency were 462 transfected with 2µg DNA in six well plates using polyethylenimine (PEI; #23966, 463 Polysciences Inc.). pcDNA3.1 or pBlueScript were used as empty vector controls. Culture 464 media were replaced at 24 hours post-transfection and cell lysates and supernatants were 465 harvested for immunoblot analysis at 48 hours as previously described₆₀(Sherer et al. 466 2011). Briefly, 1mL of harvested culture supernatant was filtered, underlaid with 20% 467 sucrose (w/v) in PBS, subjected to centrifugation at >21,000g for two hours at 4° C, and 468 viral pellets were resuspended in 35µL dissociation buffer (62.5 mM Tris-HCl, pH 6.8, 469 10% glycerol, 2% sodium dodecyl sulfate [SDS], 5% β -mercaptoethanol). Cells were 470 harvested in 500µL radioimmunoprecipitation assay (RIPA) lysis buffer (10 mM Tris-HCl, 471 pH 7.5, 150 mM NaCl, 1 mM EDTA, 0.1% SDS, 1% Triton X-100, 1% sodium 472 deoxycholate), homogenized by passage through a 26G needle, subjected to 473 centrifugation at 1,500g for 20 minutes at 4°C, and liquid supernatant fraction was 474 combined 1:1 with 2X dissociation buffer. Proteins were resolved by sodium dodecyl 475 sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to 0.2 µm 476 nitrocellulose membranes. Gag was detected using a mouse monoclonal antibody 477 recognizing HIV-1 capsid/p24 (183-H12-5C; 1:1000 dilution) from Dr. Bruce Chesebro 478 and obtained from the NIH AIDS Research and Reference Reagent Program (Bethesda, 479 MD, USA)52 and anti-mouse secondary antibodies conjugated to an infrared fluorophore 480 (IRDye680LT, 1:10000 dilution, Li-Cor Biosciences) for quantitative immunoblotting. As 481 loading controls, heat shock protein 90A/B (HSP90) was detected using a rabbit 482 polyclonal antibody (H-114, 1:2500 dilution, Santa Cruz Biotechnology, Santa Cruz, CA, 483 USA) or glyceraldehyde 3-phosphate dehydrogenase (GAPDH) was detected using a 484 mouse monoclonal antibody (6C5, 1:2500 dilution, Santa Cruz Biotechnology) and anti-485 rabbit or anti-mouse secondary secondary antibodies conjugated to an infrared 486 fluorophore (IRDye800CW, 1:7500 dilution, Li-Cor Biosciences). YFP-containing proteins 487 (e.g., YFP-A3G) were detected using a rabbit polyclonal antibody recognizing GFP (FL 488 sc-8334, 1:1000 dilution, Santa Cruz Biotechnology) and anti-rabbit secondary antibodies 489 conjugated to an infrared fluorophore (IRDye800CW). For infectivity assays, 490 supernatants containing single-round infectious HIV-1 virions (pseudo-typed with VSV-G) 491 were filtered and then added to target HeLa cells in the presence of 10µg/mL polybrene. 492 Approximately 36 hours later, target HeLa cells were scanned using a BioTek Cytation5 493 to detect reporter CFP (455nm/510nm excitation/emission filter) expression following 494 infection.

495

496 Microscopy, immunofluorescence, and FISH. Cells were plated in 24-well glass-497 bottom plates (Eppendorf) or 8-well microslides (IBIDI) and transfected using PEI or 498 infected with HIV-1 reporter viruses. Transfection mixes contained 1µg (24-well) or 333ng 499 (IBIDI) total plasmid DNA, respectively. Infections were performed with VSV-G pseudo-500 typed HIV-1 reporter viruses produced in HEK293T cells and titered on HeLa cells to 501 determine MOI of 0.5-1, thereby ensuring that 33-66% of the cells would be infected per 502 experiment. Fixed-cell experiments were performed on a Nikon Ti-Eclipse inverted wide-503 field microscope (Nikon Corporation, Melville, NY, USA) using a 100x Plan Apo oil 504 objective (numerical aperture [NA], 1.45). Transfected cells were fixed 24 – 32 hours post-505 transfection and infected cells were fixed 42 hours post-infection using 4% 506 paraformaldehyde. If treated with 250 µM sodium arsenite (Sigma-Aldrich), drug or an 507 equivalent volume of dimethyl sulfoxide were added to cell culture wells at approximately 508 one hour prior to fixation and returned to incubation at 37°C. Live-cell imaging 509 experiments were also performed on a Nikon Ti-Eclipse inverted wide-field microscope 510 using a 20x Plan Apo objective lens (NA, 0.75) with images acquired every 60 minutes 511 (except where otherwise indicated) over a time course of 16 to 90 hours. Images were 512 acquired using an ORCA-Flash4.0 CMOS camera (Hammatsu Photonics) and the 513 following excitation/emission filter sets (nanometer ranges): BFP (402/455), CFP 514 (430/470), YFP (510/535), mApple (555/605), mCherry (572/632), and iRFP (645/705). 515 All images were processed and analyzed using FIJI/ImageJ2.

516 For fixed-cell experiments using FISH, cells were plated as described as above 517 and previously₃₈. At 42 hours post-infection or after 24 hours plated (for uninfected cells), 518 cells were washed, fixed in 4% formaldehyde, and permeabilized in 70% ethanol for at 519 least 4 hours at 4°C. Custom Stellaris FISH probes were designed to recognize NL4-3 520 HIV-1 gag-pol reading frame nucleotides 386-4456 using the Stellaris RNA FISH Probe 521 Designer 4.1 (Biosearch Technologies, Inc.) available online. To detect yfp mRNAs, we 522 used a DesignReady Probe set specific to eGFP. Both probe sets were labeled with CAL 523 Fluor Red 590 Dye (Biosearch Technologies, Inc.). Samples were hybridized with the 524 gag/gag-pol probe set according to the manufacturer's instructions available online. 525 Simultaneous immunofluorescence to detect Gag used a mouse monoclonal antibody 526 specific to p24 (#24-2, a gift from Dr. Michael Malim). Imaging experiments were 527 performed as describe above on a Nikon Ti-Eclipse inverted wide-field microscope using 528 a 100x Plan Apo objective lens.

529 FRAP experiments were performed using a Nikon Ti-Eclipse inverted A1R+ 530 resonant/galvano hybrid confocal line-scanning microscope. Images were captured using 531 a 20x Plan Apo objective lens (NA, 0.75) and a GaAsP multi-detector for 488 and 560nm 532 channels. YFP-A3G was imaged using the 488nm laser at a low arbitrary intensity and 533 photobleached using the laser's maximum intensity. MS2-RFP and COGag-mCherry 534 were imaged using the 560nm laser at a low arbitrary intensity. Cells were maintained in 535 an environmental chamber (Pathology Devices, Inc.) at 37°C, 5% CO₂, and 50% humidity. 536 Cells were imaged every 30 seconds for 4 frames prior to photobleaching (3 rapid 537 ablations of cytoplasmic ROIs 10µm in diameter) followed by imaging every 5 seconds 538 for a minute, 15 seconds for four minutes, and 30 seconds for 5 minutes. This time frame 539 was sufficient for fluorescence recovery to reach a plateau. FRAP analysis was performed

540 using the FIJI plugin FRAP Profiler₆₁ that adjusts for incidental field photobleaching 541 outside the ROI.

542 Single virion analyses (SVA) were performed as previously described₃₉. Virus 543 particles were produced as described above for western blotting in HEK293T.YFP-A3G 544 cells to maintain consistent levels of YFP-A3G. Filtered culture supernatants (750µL) 545 were purified by sucrose centrifugation as described above and resuspended in 100µL 546 1x PBS (Sigma-Aldrich). 24-well plates were pre-coated with 2% FBS diluted in 1x PBS 547 for at least 30 minutes, this solution was removed, and the resuspended concentrated 548 virus particles were added to wells. Images were acquired on a Nikon Ti-Eclipse inverted 549 wide-field microscope using a 100x Plan Apo objective lens (NA, 1.45). These images 550 were processed and analyzed using Analyze Particles plugin in FIJI/ImageJ262.

551

552 Thin-section electron microscopy. HeLa.YFP-A3G were infected with Vif+ CFP or 553 Vifxx CFP HIV-1 viruses and fixed at 48 hours for chemical processing as previously 554 described₆₃. Samples were sectioned into 100nm slices and with sections collected on 555 copper thin-bar grids. Sections were observed with a Phillips CM120 transmission 556 electron microscope, and images were collected with a MegaView III (Olympus-SIS, 557 Lakewood, CO, USA) side-mounted digital camera. All images were processed and 558 analyzed using FIJI/ImageJ2₆₂.

559

560 **Statistics.** For assembly assays (Figs. 1b and 6e), results were obtained from three 561 biological replicates as defined as cells plated in six well plates transfected and processed 562 on separate days. Graphs plot the mean value with error bars representing the standard

563 deviation of the mean with the exception of Fig. 7e, a violin plot showing all data points 564 with the mean (solid line) and quartiles (dashed lines) indicated. All statistical 565 comparisons were carried out using the two-tailed Student's *t* test and performed using 566 Microsoft Excel or Graphpad Prism.

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725 FIGURE LEGENDS

726 Fig. 1. Tracking the HIV-1 replication cycle using a YFP-A3G biosensor cell line. a 727 HeLa.YFP-A3G cells showing diffuse and punctate (processing bodies, PB, white arrows) 728 distribution of YFP-A3G at steady-state (left) and accumulation into stress granules (SGs, 729 magenta arrows) following arsenite treatment (right, Ars). b Schematic of Vif-competent 730 (Vif+) or deficient (Vifxx) HIV-1 NL4-3 Env-, Vpr-, expressing CFP as a reporter of 731 infection. c Single-round infection confirms YFP-A3G dependent restriction of HIV-1 in 732 the absence of Vif. 293T.YFP-A3G cells were transfected with the indicated HIV-1 733 reporter viruses with or without co-expression of a plasmid encoding Vif-mCherry. Cells 734 and virus particles (n=3) were harvested at 48 h post-transfection and analyzed by 735 guantitative immunoblot using anti-p24_{Gag}, anti-YFP, and anti-HSP90 (loading control) 736 antisera. d Representative images from long-term time-lapse imaging showing 737 degradation of YFP-A3G upon HIV-1 Vif+ CFP infection and eventual Vif-dependent 738 G2/M arrest. e Quantification of movies (IntDen) represented in d (Vif+, solid lines) and f 739 (Vifxx, dashed lines). Cells from 3 independent experiments (n=33) with ≥ 16 hours per 740 cell (1 image/hour). f Representative images from long-term time-lapse imaging of YFP-741 A3G upon HIV-1 Vifxx CFP infection. Magenta boxes highlight zoomed regions of interest 742 (ROIs) shown below and white arrows highlight YFP-A3G clustering at PM. g Fixed cell 743 images show Gag and gagpol mRNA (viral RNA genomes) accumulating with YFP-A3G 744 at cell periphery. White arrows highlight cell-peripheral clusters of YFP-A3G, Gag, and 745 genome. h TSEM images showing virus particles clustering in the extracellular space after budding from HeLa.YFP-A3G cells infected with either Vif+ or Vifxx viruses. All scale 746

bars in fluorescent images = 10µm and in TSEM = 100nm. Error bars represent standard
deviation of the mean.

749

750 Fig. 2. HIV-1 infection has little to no impact on P-bodies or stress granules. a 751 HeLa.YFP-TIA1 cells showing steady-state distribution (vehicle, left) and accumulation 752 into SGs (magenta arrows) following arsenite treatment (right, Ars). b Representative 753 images of time-lapse imaging showing induction of SGs during infection by RVA16 754 (MOI=10). c Time-lapse images showing no effect on YFP-TIA1 distribution following 755 Vifxx CFP infection, quantified in d for >400 HeLa.YFP-A3G (black bars) and HeLa.YFP-756 TIA1 (gray bars) cells per condition, combined from three independent experiments. e 757 Time-lapse images showing no effect on CFP-DCP1a following Vifxx mCh infection, 758 guantified in **f** for 30 cells from 3 movies at ~18 hpi (YFP-A3G, black bars; CFP-DCP1a, 759 white bars). g HeLa.YFP-A3G infected with Vifxx CFP and treated with Ars to induce SGs 760 at 24 hpi and quantified in h for at least 100 infected cells from three independent 761 experiments. Magenta arrows highlight SGs present in infected cells. i Simultaneous 762 FISH/IF to detect gagpol mRNA and Gag show HIV RNA genomes accumulating in SGs 763 in HIV infected cells following Ars treatment. Magenta arrows highlight SGs present in 764 infected cells. All scale bars in fluorescent images = 10µm. Error bars represent standard 765 deviation of the mean.

766

Fig. 3. HIV-1 RNA genomes but not Gag regulate A3G subcellular localization. a and
 c Schematic Vif+ and Vifxx two-color self-tagging virus encoding Gag-CFP, 24xMSL, and
 MS2-RFP. b and d Time-lapse images of cells expressing Vif+ and Vifxx two-color HIV

770 constructs showing nuclear export of MS2-RFP tagged genomes, YFP-A3G degradation 771 (b, Vif+) or localization with MS2-RFP (d, Vifxx), and Gag-CFP expression and 772 accumulation into virus particles at the PM (white arrows). e and f Quantification of YFP-773 A3G degradation and MS2-RFP cytoplasmic accumulation (e) and Gag-CFP expression 774 (f) during expressing of Vif+ two-color HIV construct. Cells (n=31) from three independent 775 experiments wherein individual cells that could be tracked for at least 17 h (1 image/hour). 776 Black lines highlight the ~3 h window of YFP-A3G degradation. T=0 represents onset of 777 MS2-RFP expression. g Schematic of "genome only" and "Gag only" constructs. h 778 Representative images from before (top) and after (before) photobleaching in HeLa.YFP-779 A3G cells expressing "genome only" construct. White dashed circles represent regions of 780 targeted photobleaching. i Quantification of YFP-A3G fluorescence recovery experiments 781 and zoomed inset of time period over first 80 seconds after photobleaching. All scale bars 782 in fluorescent images = $10\mu m$. Error bars represent standard deviation of the mean.

783

784 Fig. 4. Viral RNA Genome-A3G interactions are sufficiently strong to redistribute 785 A3G to unnatural subcellular locales. a Schematic of In Cell RNA-protein Interaction 786 Protocol (IC-IP). b Schematic of MS2-iRFP constructs used for "targeting" (top) and 787 representative images of MS2-iRFP and YFP-A3G localization in the absence of mRNAs 788 encoding MSL cassette. White arrows point to characteristic subcellular targets of Src-789 (vesicles) and Lifeact- (filamentous actin) MS2-iRFP. c Schematic of IC-IP RNA Bait -790 Vifxx two-color self-tagging HIV construct. d Representative images showing Free MS2-791 iRFP-NLS, MS2-RFP-NLS, YFP-A3G, and Gag-CFP accumulating at PM-adjacent 792 assembly sites (top). (Middle) Src-MS2-iRFP accumulates at intracellular/perinuclear 793 vesicles, recruits MS2-RFP-tagged RNA genomes, YFP-A3G, and Gag-CFP. (Bottom) 794 Lifeact-MS2-iRFP induces similar accumulation at actin filaments. Region of interest 795 (ROI) shows YFP-A3G signal in dashed white boxes with arrows highlighting assembly 796 sites (top), vesicles (middle), and linear filaments (bottom). e Quantification of re-797 localization of YFP-A3G phenotypes (n = > 90 cells per condition) in the presence of NLS. 798 Src-, and Lifeact-MS2 targeting constructs and **f** percentages of YFP-A3G per cell signal 799 (IntDen) relocalized by Src- and Lifeact-MS2 constructs (n > 6). All scale bars in 800 fluorescent images = 10µm, except in A3G ROI insets scale bars = 1µm. Error bars 801 represent standard deviation of the mean.

802

803 Fig. 5. Recruitment of YFP-A3G is specific to HIV-1 RNA genome, Gag-independent, 804 and occurs long before the onset of virus particle assembly. a and c Schematics of "genome only" and "Gag only" IC-IP bait constructs, respectively. b Representative 805 806 images showing normal co-distribution of MS2-RFP tagged RNA genomes and YFP-A3G 807 (top) and retargeting of YFP-A3G to intracellular vesicles with RNA genomes by Src-MS2-808 iRFP (bottom). d Representative images showing Src-MS2-RFP accumulating at 809 intracellular vesicles and a complete lack of YFP-A3G at these same sites (white arrow). 810 e Time-lapse images of cells expressing "genome only" construct retargeted by Src-MS2-811 iRFP over a period of 4 hours. T=0 represents the first detection of MS2-RFP genome 812 proxy. Arrow shows accumulation of YFP-A3G at Src-MS2-iRFP+ vesicles as early as 2 813 hours after the onset of MS2-RFP expression. All scale bars in fluorescent images = 814 10µm.

815

816 Fig. 6. Membrane-targeted APOBEC3G recruits HIV-1 RNA genomes and can inhibit

817 virus particle production. a Schematic of Reciprocal IC-IP using Src-YFP-A3G targeted 818 to intracellular membranes. b and c Schematic of Vifxx two-color HIV (b) and "genome 819 only" (c) constructs and representative images showing Src-YFP-A3G (b) or Src-CFP-820 A3G (c) recruiting MS2-RFP tagged genomes to intracellular vesicles (white arrows). d 821 Schematic of Vifxx Luc HIV infectious single-cycle virus used to infect HeLa.Src-YFP-822 A3G cells and representative images of FISH/IF detecting gag-po/ mRNA and Gag p-24, 823 respectively. White arrows highlight sites where Src-YFP-A3G has recruited bona fide 824 HIV RNA genomes. e Western blot showing dose-dependent inhibition of HIV virus 825 particle assembly and production from HEK293T cells transfected with Vifxx two-color 826 construct, Src-MS2-YFP (lanes 1 and 2), YFP-A3G (lane 3) and Src-YFP-A3G (lanes 4 827 and 5). Graph shows relative release factor as the ratio of Gag in virus-like particles 828 (VLPs) divided by Gag in cellular lysates quantified from three independent experiments. 829 All scale bars in fluorescent images = $10\mu m$. Error bars represent standard deviation of 830 the mean.

831

Fig. 7. Cytoplasmic A3G-genome interactions promote consistent per virion delivery of A3G into progeny virions. a Schematics of constructs used in single virion analysis (SVA) studies. b Western blot showing cellular expression, YFP-A3G packaging, and Gag assembly and release of constructs depicted in a. c Representative images of single fluorescent virion images harvested from HEK293T.YFP-A3G cell line. Scale bar = 500nm. d Quantification of SVA for each of the four constructs showing percent of virions with MS2-RFP signal (pink) and YFP-A3G (green). Error bars represent standard

- 839 deviation of the mean (n > 13,000 virions per condition). **e** Relative YFP-A3G signal per
- 840 virion or virus-like particle derived from Vifxx two-color or COGag conditions. Relative
- values are normalized to the mean value for Vifxx with means (solid lines) and 25th and
- 842 75th quartiles (dashed lines) indicated.

843 SUPPLEMENTARY MATERIAL

844

Fig. S1. Graphical abstract. Summary of single-cell live imaging approaches used andkey findings.

847

848 Video 1. Time-lapse imaging of HeLa-.FP-A3G cells infected with Vif+ HIV-1/CFP 849 virus. Multi-channel images of HeLa.YFP-A3G cells were acquired every 60 minutes for 850 49h after infection. In the highlighted cell, viral gene expression (cyan) is detected ~16hpi 851 with YFP-A3G degradation green0 at 21 hpi. This cell rounds up at 28 hpi consistent with 852 Vif-induced G2/M cell cycle arrest. 853 854 Video 2. Time-lapse imaging of HeLa.YFP-A3G cells infected with Vifxx HIV-1/CFP 855 virus. Multi-channel images were acquired every 60 minutes for 49h after infection. In the 856 highlighted cell, viral gene expression (cyan) is detected ~19hpi with detection of YFP-857 A3G (green) at virus particles (arrows) starting at ~25hpi. An example of a transfer event 858 wherein a large cluster of YFP-A3G+ virus particles is released from an infected cell is

highlighted (arrows) at 32-36hpi.

860

Video 3. Time-lapse imaging of HeLa.YFP-TIA1 cells infected with Rhinovirus A16.
Multi-channel images were acquired every 30 minutes after addition of RVA16 (MOI~10).

863 The first instances of SG formation (YFP-TIA1 in green) are evident at ~11hpi.

864

865 Video 4. Time-lapse imaging of HeLa.YFP-A3G cells expressing the two-color Vif+

866 Gag-CFP/MS2-RFP HIV-1 reporter virus. Multi-channel images were acquired every 60

867 minutes for 24h beginning ~4h post-transfection. Left panel shows genome tracked using

868 MS2-RFP (magenta), with nuclear export detected at T=4h just prior to the onset of Gag-

869 CFP (cyan) expression (right panel) and YFP-A3G (green) degradation (center panel).

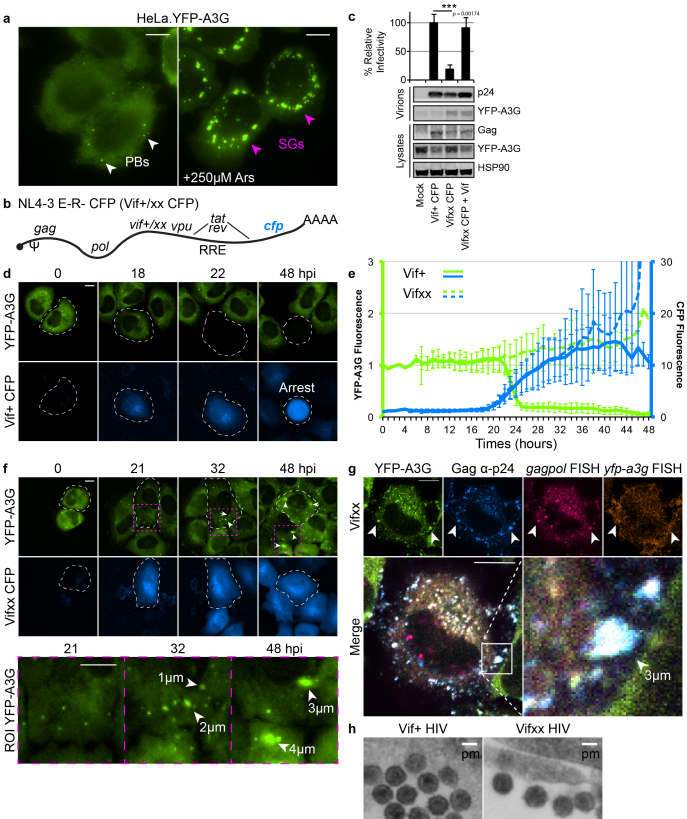
870

Video 5. Time-lapse imaging of HeLa.YFP-A3G cells expressing the two-color Vifxx Gag-CFP/MS2-RFP HIV-1 reporter virus. Multi-channel images were acquired every 60 minutes for 24 hours beginning ~4 hours post-transfection. Left panel shows genome tracked using MS2-RFP (magenta), with nuclear export detected at T=6h just prior to the onset of Gag-CFP (cyan) expression (right panel) and co-accumulation of YFP-A3G (green, center panel) with genome (MS2-RFP) and Gag-CFP at the plasma membrane (at putative sites of virus particle assembly).

878

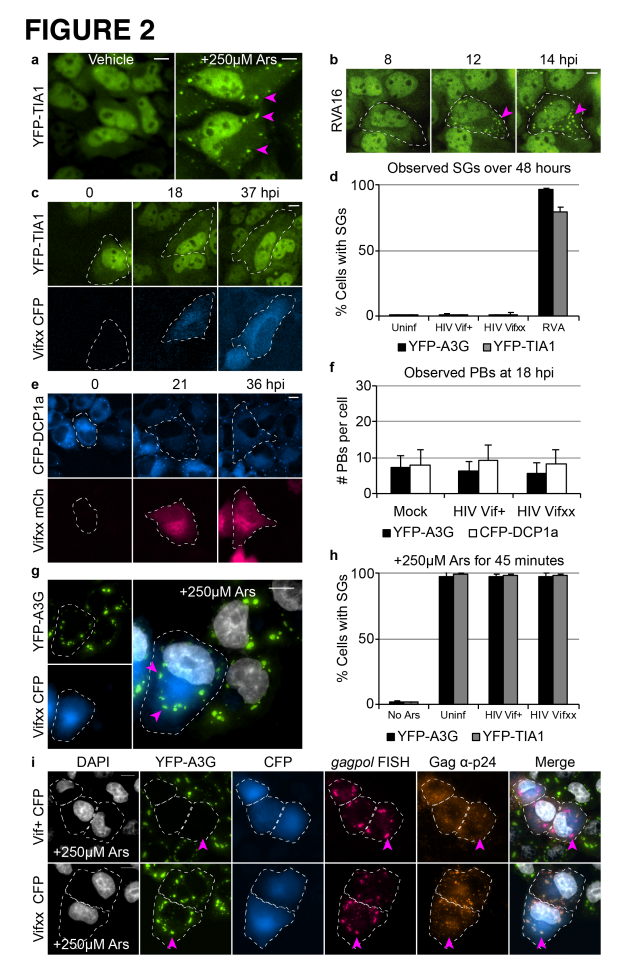
879 Video 6. Time-lapse imaging of HeLa.YFP-A3G cells co-expressing "genome only" 880 MS2-RFP HIV-1 reporter virus with the Src-MS2-iRFP targeter (orange). Multi-881 channel images were acquired every 60 minutes for 15h starting ~4h post-transfection. 882 MS2-RFP tracked genomes (magenta, central panel) are first observed at T=0, with both 883 genomes and YFP-A3G (green, right panel) immediately recruited to perinuclear vesicles, 884 co-localizing with the Src-MS2-iRFP targeter (yellow, left panel). Video demonstrates that 885 genomes have marked effects on YFP-A3G trafficking in the cytosol even at the earliest 886 time points post-genome nuclear export.

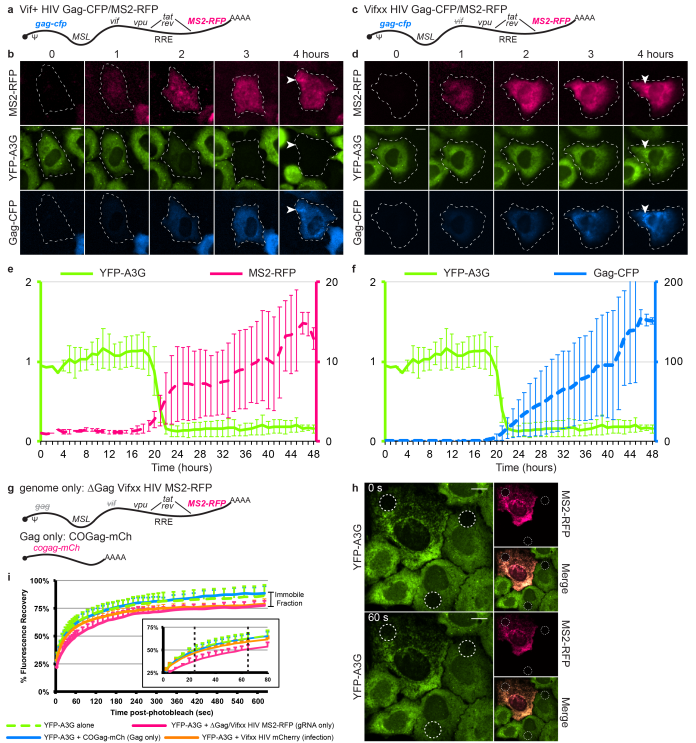
FIGURE 1

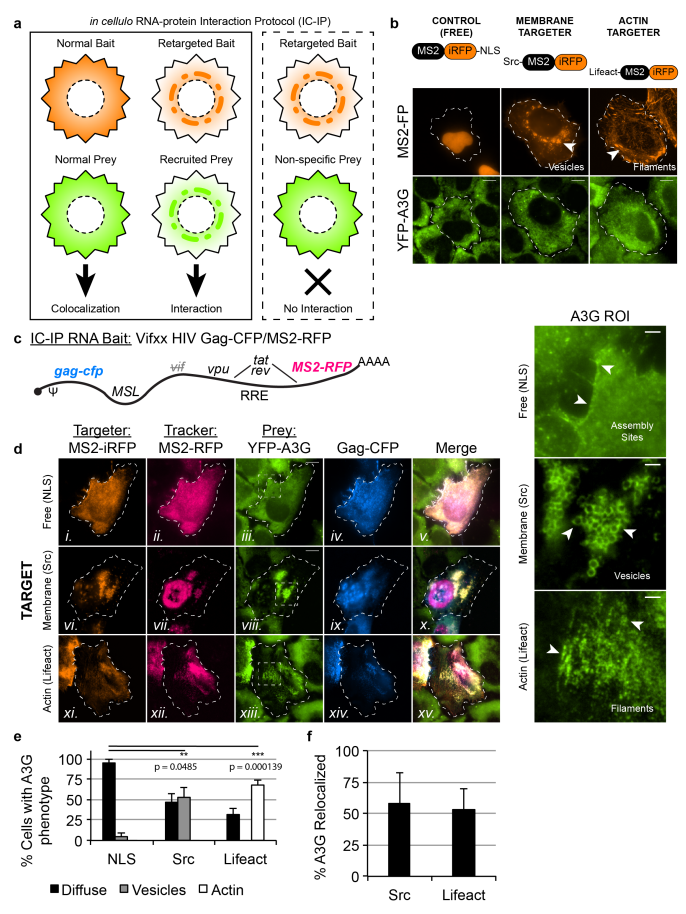


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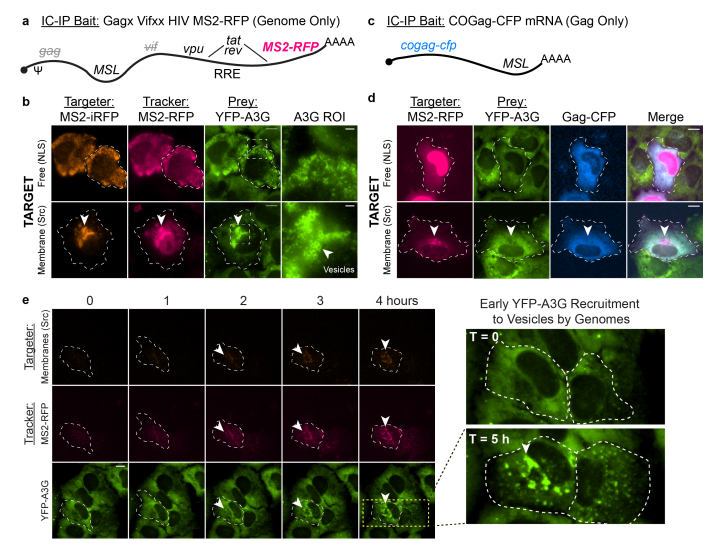
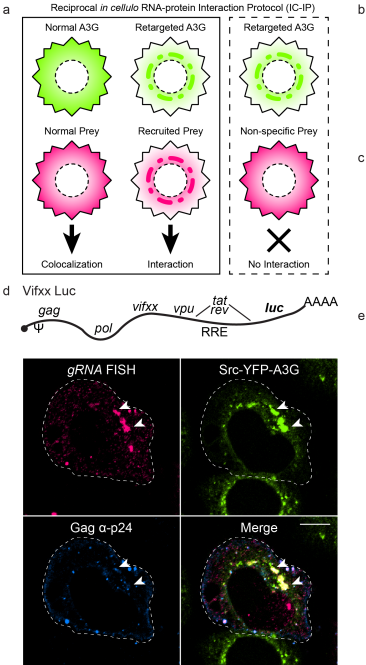


FIGURE 6



b Vifxx Gag-CFP/MS2-RFP ∕tat ∕rev AAAA MS2-RFP vpu gag-cfp MS RRE MS2-RFP Src-YFP-A3G Gag-CFP Merge c Genome Only Vifxx ∕ tat ∕ rev MS2-RFP AAAA vpu gag MSL Ψ RRE MS2-RFP YFP-A3G Src-CFP-A3G Merge *** p = 0.00141 *** p = 0.00265 Release Factor 100 50 n Gag-BFP VLPs YFP Gag-BFP Lysates YFP GAPDH 3 2 4 5 1

