The N-Terminus of Murine Leukaemia Virus p12 Protein Is Required for Mature Core Stability

Darren J. Wight1, Virginie C. Boucherit1, Madushi Wanaguru1, Efrat Elis2, Elizabeth M. A. Hirst3, Wilson Li1, Marcelo Ehrlich2, Eran Bacharach2, Kate N. Bishop1*

1 Division of Virology, MRC National Institute for Medical Research, The Ridgeway, Mill Hill, London, United Kingdom, 2 Department of Cell Research and Immunology, George S. Wise Faculty of Life Sciences, Tel Aviv University, Tel Aviv, Israel, 3 Division of Developmental Neurobiology, MRC National Institute for Medical Research, The Ridgeway, Mill Hill, London, United Kingdom

Abstract

The murine leukaemia virus (MLV) gag gene encodes a small protein called p12 that is essential for the early steps of viral replication. The N- and C-terminal regions of p12 are sequentially acting domains, both required for p12 function. Defects in the C-terminal domain can be overcome by introducing a chromatin binding motif into the protein. However, the function of the N-terminal domain remains unknown. Here, we undertook a detailed analysis of the effects of p12 mutation on infected viral cores. We found that both reverse transcription complexes and isolated mature cores from N-terminal p12 mutants have altered capsid complexes compared to wild type virions. Electron microscopy revealed that mature N-terminal p12 mutant cores have different morphologies, although immature cores appear normal. Moreover, in immunofluorescent studies, both p12 and capsid proteins were lost rapidly from N-terminal p12 mutant viral cores after entry into target cells. Importantly, we determined that p12 binds directly to the MLV capsid lattice. However, we could not detect binding of an N-terminally altered p12 to capsid. Altogether, our data imply that p12 stabilises the mature MLV core, preventing premature loss of capsid, and that this is mediated by direct binding of p12 to the capsid shell. In this manner, p12 is also retained in the pre-integration complex where it facilitates tethering to mitotic chromosomes. These data also explain our previous observations that modifications to the N-terminus of p12 alter the ability of particles to abrogate restriction by TRIM5α or Fv1, factors that recognise viral capsid lattices.

Introduction

Retroviruses initially assemble as immature viruses containing a core of Gag and Gag-Pol polyproteins. During maturation these are cleaved into mature proteins by the virally encoded protease (PR). Cleavage of the gammaretrovirus Gag polyprotein produces four mature proteins: matrix (MA), p12, capsid (CA) and nucleocapsid (NC). A mass rearrangement follows cleavage, forming the mature CA core surrounding the condensed ribonucleoprotein complex [1]. Cryogenic electron microscopy studies on the maturation intermediates of HIV-1 have indicated that maturation is a step-wise and regulated process [2]. Maturation is essential for infectivity and blocking maturation using PR inhibitors has been heavily utilised in the control of HIV-1 infection [3]. Resistance to PR inhibitors remains a significant problem [4], and a greater understanding of the viral and cellular factors involved in maturation could yield new therapeutic targets. Furthermore, the CA shell of the core is beginning to be implicated in many early events from reverse transcription to integration [5–7], and understanding how the core is formed and maintained during an infection is of central importance.

The Gag protein p12 has important roles during both the early and late stages of murine leukaemia virus (MLV) infection [8]. It harbours the PPPV late-domain (L-domain), essential for recruiting HECT ubiquitin ligases to manipulate the ESCRT pathway for efficient budding [8,9]. Additionally, seven mutants have been defined in Mo-MLV p12, four in the N-terminal and three in the C-terminal, which have a potent block during the early stages of infection [Fig. 1A] [8,10]. The replication defects of these mutants fall into three groups: (i) mutants defective in reverse transcription in vivo (mutant 6), (ii) mutants defective in reverse transcription in certain cell lines (mutant 8) and (iii) mutants competent for reverse transcription but failing to integrate their viral DNA (mutant 5, 7, 13, 14 and 15) [8,10]. We have recently shown that the N- and C-terminal regions of p12, mapped out by these mutants, are actually two sequentially acting domains, both of which are required to be active on the same p12 molecule for the transduction of target cells [10].

Biochemical analysis showed that isolated pre-integration complexes (PICs) from C-terminal p12 mutants were comparable to wild type, and were functional to integrate their DNA in vitro...
All retroviral genomes contain a gag gene that codes for the Gag polyprotein. Gag is cleaved upon viral maturation to release individual proteins, including matrix, capsid and nucleocapsid, providing the structural components of the virion. In murine leukaemia virus (MLV), Gag cleavage releases an additional protein, named p12, required for both early and late stages of the viral life cycle. The role of p12 during early events is poorly understood, and it is the only MLV protein without a function-associated name. Here, we show that p12 binds to the capsid shell of the viral core and stabilises it. Mutations that give rise to N-terminally altered p12 proteins result in a rapid loss of both p12 and capsid from viral cores, leading to abnormal core morphologies and abolishing the ability of particles to abrogate restriction by cellular factors that target viral capsid lattices. Understanding how the mature retroviral core forms and how it disassembles during infection is important as this determines the infectivity of all retroviruses, including HIV-1. Furthermore, altering core stability has recently become a novel target for HIV-1 therapeutics.

Results

Only the N-terminal domain of p12 is required for abrogation of TRIM5alpha activity

One way of testing whether non-infectious viral particles can be recognised by the CA-targeting restriction factors Fv1 and TRIM5alpha is to perform a saturation, or abrogation, assay. Briefly, target cells expressing low levels of restriction factor are pre-exposed to increasing amounts of tester virus. If these tester virus particles can be recognised by the restriction factor, they will bind to and saturate the factor, abrogating restriction and allowing a GFP reporter virus, that would normally be restricted, to infect the cells. We previously demonstrated that p12 contains two domains that act in concert, one towards the N-terminus of the protein and the second towards the C-terminus [10]. Using restriction factor abrogation assays, we showed that N-tropic MLV (N-MLV) particles containing N-terminally mutated p12 lost the ability to abrogate Fv1 and TRIM5alpha whilst C-terminal p12 mutants did not (mutations shown in Fig. 1A) [10]. Interestingly, virus particles containing mixtures of both N- and C-terminally altered p12 were non-infectious showing that these mutations do not complement each other [10]. However, we wondered whether the inability of the N-terminal p12 mutants to abrogate restriction factors could be rescued in trans by C-terminal p12 mutant molecules. VSV-G pseudotyped LacZ-encoding virus-like particles (VLPs) were prepared by transfecting 293T cells with varying ratios of plasmids expressing either p12 mutant 6 or mutant 14 N-MLV Gag-Pol. Serial dilutions of these “tester” virus particles were added to TE671 cells, expressing humanTRIM5alpha, followed by a fixed and equal amount of GFP-encoding “reporter” N-MLV, and the number of GFP positive cells was measured after 72 hours. As expected [10], p12 mutant 6 VLPs were unable to abrogate restriction by TRIM5alpha, in contrast to the C-terminal mutant 14 and wild type particles (Fig. 1B, green triangles versus purple and black triangles, respectively). However, despite being non-infectious (Fig. 1C), all of the mixed particles tested, containing decreasing amounts of p12 mutant 14 from 90–10%, were able to abrogate TRIM5alpha restriction (Fig. 1B, coloured lines with open coloured circles). This phenotype was also observed when p12 mutant 6 was mixed with either of the other two C-terminal p12 mutants, 13 or 15 (Fig. S1 A and B). This suggests that, unlike for infectivity, C-terminal p12 mutant proteins can rescue the ability of particles containing N-terminal p12 mutant proteins to saturate TRIM5alpha, even when only 10% of the p12 molecules in the particle have a wild type N-terminus. Thus, a small amount of the N-terminus of wild type p12 is required to enable VLPs to interact with restriction factors.

Mutations in the N-terminus of p12 alter the biophysical properties of the MLV CA core in target cells

TRIM5alpha targets the CA shell of N-MLV. The failure of N-terminal p12 mutants to abrogate restriction factors therefore suggests an intriguing possibility that the CA shell is perturbed in these mutants. Thus, we investigated what effect N-terminal p12 mutations had on CA. We previously showed that CA expressed from gag containing an upstream mutation in p12 could be incorporated into the CA core and be recognised by restriction factors, indicating that mutations in p12 do not affect the function of CA molecules per se [10]. Therefore, we studied the effect of p12 mutations on CA complexes using a modified faye-of-CA assay. D17 cells were challenged with wild type or p12 mutant
Mo-MLV VLPs for 4 hours and cell lysates were separated on 10–42% (w/w) linear sucrose gradients by velocity sedimentation. The presence of CA in each gradient fraction was assessed by immunoblotting (Fig. 2).

For wild type particles, CA was present in two populations (Fig. 2A, top immunoblot). The first, in fraction 1 at the top of the gradient, presumably represented CA not in complexes, most likely released from viral particles after membrane fusion and/or during the poorly defined uncoating process. The second population was distributed through fractions 3 to 7 in the middle of the gradient, with the peak of material found in fraction 6. Importantly, this material co-migrated with viral cDNA, as measured by qPCR, (Fig. 2A, bottom immunoblot), implying that the CA complexes observed were reverse transcription complexes (RTCs). We have previously reported that in genetic assays, approximately only 10% of the p12 protein in the viral particle needs to be wild type for full infectivity [10]. CA and p12 are present at a 1:1 ratio in viral particles as both are formed when Gag is cleaved. By directly comparing the ratio of CA to p12 in wild type viral particles and cell lysates four hours after infection (Fig. 2B), we could clearly see that much less p12 was present in the cell than CA. This suggests that most of the p12 is degraded rapidly after infection, presumably because it is not associated with the RTC, with no detrimental consequences for the virus.

When we analysed cell lysates following infection with our panel of p12 mutants, we observed that the pattern of CA migration through velocity sucrose gradients for the C-terminal p12 mutants was similar to wild type (Fig. 2C) with the peak in fractions 6 and 7. However, the N-terminal p12 mutants had a notably different CA distribution (Fig. 2C). Interestingly, overall, the CA complexes from the N-terminal mutants did not travel as far through the
velocity gradient as the wild type complexes, indicating a reduction in their apparent S values. The phenotype of mutant 6 was the most strikingly different. Here, most of the CA was present in fractions 1 to 3 suggesting that the rate of sedimentation of these CA-containing complexes was much slower than wild type complexes. Notably, mutant 6 exhibits a 10-fold reduction in reverse transcription in cells [8,10], and reverse transcription is known to be sensitive to the state and stability of the viral core. To quantify these observations and compare multiple experiments, the density of each fraction was measured and all gradients were found to be comparable. The intensity of the CA signal on the immunoblot was determined and the relative shift in the peak CA position compared to wild type RTCs was calculated. The mean and range of at least three independent experiments are displayed.

doi:10.1371/journal.ppat.1004474.g002

Figure 2. The effect of p12 mutations on the biophysical properties of the Mo-MLV RTC. (A) D17 cells were infected for four hours with wild type Mo-MLV, and cell lysates were separated using velocity sedimentation through a 10–42% (w/w) sucrose gradient. Fractions were analysed by qPCR with primers targeted to strong stop viral cDNA and proteins were precipitated and analysed by immunoblotting using anti-CA (top blot) and anti-p12 monoclonal (bottom blot) antibodies (Fraction 1 is the top of gradient). The graph displays the copies of viral DNA per microliter (black line) and the percentage of total CA in each fraction (grey line). (B) Wild type Mo-MLV was used to challenge D17 cells and after four hours the cells were lysed (cell lysate). An aliquot of the Mo-MLV VLPs used for the infection was also lysed (virus) and both samples were analysed by immunoblotting with anti-CA (top panel) and anti-p12 (bottom panel) monoclonal antibodies. Two repeats of this experiment are shown in the immunoblot. (C) Infected cell lysates from D17 cells challenged with wild type or p12 mutant Mo-MLV VLPs were separated using velocity sedimentation through a 10–42% (w/w) sucrose gradient and analysed by immunoblotting for CA. (D) For each experiment, the intensity of the CA signal on the immunoblot was determined and the relative shift in the peak CA position compared to wild type RTCs was calculated. The mean and range of at least three independent experiments are displayed.
gradient for each p12 mutant was compared to that of wild type RTCs, and the relative shifts in peak positions are plotted in Fig. 2D. Clearly, the N-terminal p12 mutant RTCs had reduced sedimentation rates, suggesting that they have altered CA complexes. Interestingly, there was a slight, but reproducible, increase in the rate of sedimentation of the C-terminal p12 mutant CA complexes compared to wild type (Fig. 2D). Unfortunately, it was difficult to detect p12 in these samples due to the low levels of p12 in complexes and the fact that our monoclonal antibody to p12 does not recognise the N-terminal p12 mutants.

N-terminal p12 mutants have intrinsically different CA complexes

Both the abrogation and biophysical data presented thus far have highlighted that N-terminal p12 mutant RTCs are altered in infected cells. However, to assess whether virions themselves are intrinsically different, we attempted to isolate CA cores from whole VLPs. Concentrated wild type and p12 mutant Mo-MLV VLPs were spun through a layer of detergent into a 10–42% (w/w) linear sucrose gradient. Fractions were harvested and analysed by immunoblotting using an anti-CA antibody (note: fractions 1–3 are diluted by 1:13 to reduce the immunoblot signal; *: this input was run on a separate gel from the gradient fractions). Representative immunoblots are shown for wild type and each of the p12 mutants. For each experiment, the sucrose density of the fraction containing the peak CA signal was measured, and the change in density compared to peak CA fraction for wild type virions was calculated. The mean and range from two independent experiments are displayed in the histogram.

doi:10.1371/journal.ppat.1004474.g003

Figure 3. The effect of p12 mutations on the biophysical properties of the Mo-MLV intra-virion CA core. (A) Purified wild type or p12 mutant Mo-MLV VLPs were spun through a layer of detergent into a 10–42% (w/w) linear sucrose gradient. Fractions were harvested and analysed by immunoblotting using an anti-CA antibody (note: fractions 1–3 are diluted by 1:13 to reduce the immunoblot signal, *: this input was run on a separate gel from the gradient fractions). Representative immunoblots are shown for wild type and each of the p12 mutants. (B) For each experiment, the sucrose density of the fraction containing the peak CA signal was measured, and the change in density compared to peak CA fraction for wild type virions was calculated. The mean and range from two independent experiments are displayed in the histogram.
cores are less stable or they are not correctly formed. To investigate core formation, we analysed the morphology of p12 mutant particles by transmission electron microscopy (TEM). Large batches of purified p12 mutant 6, 7, or 8 VLPs were pelleted and prepared for TEM alongside a matching wild type control. Thin (50 nm) sections were sliced through the virus pellet, stained and images acquired throughout the section at 20,000× magnification. Representative images are shown in Fig. 4 and further images are included in Fig. S3. The infectivity of the VLPs used was assessed in D17 cells (Fig. S4) and the three p12 mutants displayed the characteristic infectivity defect (100–1000-fold reduction in infectivity) as seen in Fig. 1C and previously described [10]. TEM images were selected at random, and the core morphologies of all VLPs with a diameter of 80–120 nm were classified into one of four categories according to a standard set of morphologies (Fig. 4E and F). At least 93 particles were scored for each sample. As expected, most wild type Mo-MLV virions contained a roughly circular electron-dense mature core filling up most of the intra-virion space (Fig. 4A and F, Fig. S3A). Strikingly, only 13% of p12 mutant 6 virions contained such cores compared to 78% of the corresponding wild type control (Fig. 4B and F, Fig. S3B). The majority of these particles either had no identifiable mature or immature core (28%) or contained cores with a grossly aberrant morphology (57%) such as a small electron dense spot or

Figure 4. Analysis of the Mo-MLV p12 mutant intra-virion CA core by transmission electron microscopy. Purified Mo-MLV VLPs were pelleted and prepared for TEM. (A–D) Representative electron micrographs of wild type Mo-MLV and p12 mutant VLPs are shown. (E) Representative examples of the different core morphologies used to classify particles observed in the micrographs. (F) The morphology of all Mo-MLV particles (within 80–120 nm in diameter) were scored from multiple micrographs (at least 93 particles scored for each) and the results are displayed as a percentage of the total particles scored. All scale bars are 50 nm.
doi:10.1371/journal.ppat.1004474.g004
asymmetrical restricted density (Fig. 4B and F, Fig. S3B). The p12 mutant 6 particles manifest their defect earlier in the viral life cycle than the other p12 mutants; failing to synthesize wild type levels of cDNA in the target cell [8,10]. In contrast, p12 mutant 7 is competent to reverse transcribe normally [8,10]. Somewhat intuitively, therefore, p12 mutant 7 had less of an effect on core morphology than p12 mutant 6, with 26% and 17% of virions containing an aberrant core or lacking a core, respectively (Fig. 4C and F, Fig. S3C). A much higher number of p12 mutant 7 virions contained a mature core (56%) when compared to p12 mutant 6, although this was still considerably lower than the corresponding wild type control (73%) (Fig. 4F). These observations suggest that mutation of the N-terminal domain of p12 is detrimental to the structure of the mature Mo-MLV CA core, and the severity of the defect correlates with the ability of particles to reverse transcribe in D17 cells.

Alteration of the CA core is only evident after maturation

Gag and Gag-Pol are incorporated into the immature Mo-MLV particle as polyproteins, forming an immature Gag lattice. During maturation, the viral protease (PR) cleaves the polyproteins into individual Gag and Pol proteins, allowing CA to rearrange and assemble into a structurally distinct mature CA lattice [25–30]. Since p12 mutant 6 failed to assemble mature CA cores correctly, we asked whether p12 mutant 6 immature Gag lattices, the precursor of the mature core, were also altered.

Figure 5. The effect of p12 mutant 6 on the immature Mo-MLV core. (A and B) A mutation was introduced into the viral protease (PR) to inactivate its activity (D32L, called PR-). PR- (A) wild type and (B) p12 mutant 6 Mo-MLV VLPs were prepared for TEM and representative electron micrographs of each are shown. The colored boxes show enlargements of individual virions. (C and D) Wild type and p12 mutant 6 Mo-MLV producer cells were pelleted and prepared for TEM. Representative electron micrographs of (C) wild type and (D) p12 mutant 6 immature VLPs budding from producer cells are shown. Immature VLPs are highlighted by the red boxes. All scale bars are 50 nm.

doi:10.1371/journal.ppat.1004474.g005

mutant 8 virions contained an aberrant core and 30% contained no identifiable core at all. Less p12 mutant 8 virions contained a mature core than p12 mutant 7 (43% vs 56%), although the corresponding wild type control also had a reduced number of virions with mature cores (64% vs 73%) (Fig. 4F).
Immature VLPs were produced by transfecting 293T cells with a Gag-Pol expressing construct that contained the D32L mutation in PR, which abolishes enzymatic activity [31]. Protease minus (PR-) wild type and p12 mutant 6 VLPs were concentrated and analysed by equilibrium sedimentation (Fig. S5). This analysis revealed no difference in the density of whole immature p12 mutants compared to the wild type control, as was seen for whole mature p12 mutant VLPs (Fig. S2), confirming that no gross abnormalities occur during particle assembly. Large batches of PR- wild type and p12 mutant 6 VLPs were then synthesised and prepared for TEM analysis as described above. Strikingly, PR- wild type and p12 mutant 6 VLPs were almost identical (Fig. 5A vs B). For both, nearly all particles contained an electron dense ‘train track’-like ring directly underneath the lipid bilayer, indicative of an immature core morphology [26]. Moreover, when cells producing either wild type or p12 mutant 6 VLPs with unmodified protease were analysed by TEM, “natural” immature particles could be seen budding from both p12 mutant 6 and wild type producer cells that were indistinguishable from one another (Fig. 5C vs D). Taken together, these data suggest that the p12 mutant 6 immature Gag lattice is not grossly altered; indicating that the core defect either manifests during maturation or is too subtle to be detected in immature particles.

The N-terminal domain of p12 is required for retention of p12 in the MLV PIC

Although p12 mutant 6 has a clear defect in mature core formation, p12 mutants 7 and 8 appeared to produce significant amounts of particles (approximately half) with wild type-like core morphologies (Fig. 4). Therefore, the more than 99.5% reduction in infectivity of these mutants (Fig. S4) could not be explained by the complete absence of correctly formed mature cores. The data from our biophysical studies (Fig. 2 and 3) determined that the CA complexes are altered, so if this is not reflected by core appearance, it might suggest that the core is less stable. Moreover, mutations that alter the stability of the HIV core lead to aborted infections [32]. To observe cores in vivo, we followed infections by indirect immunofluorescence. Ecotropic Mo-MLV VLPs containing a previously described single myc-tag in p12 [12] were synthesised by transient transfection in 293T cells. U/R cells (U20S cells stably expressing mCAT-1 [12]) were challenged with equal doses of these wild type (MOI 3) or p12 mutant Mo-MLV-myc VLPs by spinoculation (1000g at 4°C). Cells were extensively washed, to remove unbound VLPs, returned to the incubator and fixed at various times post-infection. All the infections were done in duplicate with one sample stained for the myc-tagged p12 and the other for the CA protein as the fixing protocols necessary for each antibody were incompatible.

For wild type Mo-MLV, an apparent decrease in the number of p12 puncta could be seen with time (Fig. 6A, top row). Traditionally, it has been thought that the rate of CA uncoating for MLV is slower than HIV-1, with significant amounts of CA remaining with the MLV PIC during transit to the nucleus [24,33]. In line with these observations, we could see that cells challenged with wild type Mo-MLV contained large numbers of CA puncta, which remained for the length of the time course (Fig. 6B, top row). Thus, during infection with wild type Mo-MLV, some proportion of the viral p12 is shed from the PIC faster than CA, mirroring our earlier immunoblot observations (Fig. 2B).

Interestingly, all of the N-terminal p12 mutants exhibited a differing phenotype to wild type Mo-MLV: A significant reduction in the number of p12 puncta could be observed very early, 0.5–1 hour post-infection (Fig. 6A and S6A). Furthermore, the number of CA puncta in N-terminal p12 mutant challenged cells also decreased with time (Fig. 6B and S6B). Consistent with our biophysical data, cells infected with the C-terminal p12 mutant 14 showed a similar pattern of puncta to wild type infections (Fig. S6A and B, bottom row).

To quantify these observations, cells were chosen at random using the nuclear counterstain and the entire cell body was imaged using a spinning disk confocal microscope. Outlines of the cells were drawn and the number of p12(myc) and CA puncta within each cell determined. Table S1 shows the mean number of puncta measured at time zero and two hours post-infection. The mean number of p12(myc) and CA puncta at each time point for each infection was normalised to the mean number of puncta at the zero hour time point and plotted against time post-infection. Figure 7 shows the analysis from 10 cells containing high numbers of puncta (~250–300 puncta per cell at the zero hour time point). Importantly, very similar results were obtained from analysis of 10–16 cells from separate infections containing lower numbers of puncta; 30–60 puncta per cell at the zero hour time point (unpublished data). These analyses clearly demonstrate that there was a rapid reduction of p12 puncta from all N-terminal p12 mutant infected cells (Fig. 7A–D, coloured dashed lines), with more than 75% of the p12 puncta lost by two hours post-infection (Fig. 7F). In contrast, less than 50% of the p12 puncta were lost in cells infected with either wild type Mo-MLV (Fig. 7A–E, black dashed line, 7F) or p12 mutant 14 (Fig. 7E, purple dashed line, 7F). Notably, the number of CA puncta was also reduced by 65–75% by two hours post-infection in cells infected with the N-terminal p12 mutant 7 (Fig. 7F), while there was only a minor reduction of CA puncta for wild type and p12 mutant 14 infections (Fig. 7F). Statistical analysis (t-test) of the number of puncta in cells two hours after infection (Fig. 7F) showed highly significant differences for N-terminal p12 mutants compared to wild type infections. Specifically, comparing p12 puncta with wild type gave p values of 0.01, 0.002, 0.0038 and 0.0029 for p12 mutants 5, 6, 7 and 8 respectively, and comparing CA puncta with wild type gave p values of 0.0028, 0.0023, 0.0029 and 0.0036 for p12 mutants 5, 6, 7 and 8, respectively. Taken together, these results suggest that alteration to the N-terminus of p12 results in a rapid loss of both p12 and CA itself from incoming viral cores. This suggests that the N-terminal domain of p12 is required for the retention of p12 within the RTC, and for conservation of the MLV CA core in the target cell.

The N-terminus of p12 is responsible for binding to CA

For p12 to be incorporated into the RTC, one would expect p12 to interact with core components. Indeed, a small amount of CA was previously immunoprecipitated from cells challenged with Mo-MLV using an antibody against myc-tagged p12, although this could not be recapitulated by immunoprecipitation of p12 from virions [12]. Given that p12 appears to influence the stability of the core, it is logical to predict that p12 binds directly to CA. However, a direct binding has never been shown, and most CA binding assays are hindered by the fact that the CA in the RTC is present in the form of a lattice, so monomeric CA may not recapitulate the binding surface present in an array. Fortunately, a protocol to form mature MLV CA lattice arrays on lipid nanotubes was previously established to study CA-Fv1 interactions [34]. We therefore used this approach to investigate whether p12 directly binds the CA lattice. Briefly, purified His-tagged N-MLV CA was immobilised on lipid nanotubes comprising the Ni2+-chelating lipid, DGS-NTA. These tubes were then incubated with purified p12 protein, and bound complexes were separated from unbound p12 by centrifugation through a sucrose cushion. The pellet fraction was analysed for the presence of His-tagged CA.
or p12 proteins by immunoblotting with anti-His tag and anti-p12 polyclonal antibodies respectively. Fig. 8 shows representative immunoblots from 4 independent experiments that demonstrate detectable binding of wild type p12 protein to CA-coated lipid nanotubes (lane 2) but not p12 mutant 6 (lane 6). Importantly, we did not detect binding of either p12 protein to a version of CA that cannot form high density, regular arrays, CA-P1G [34] (lanes 3 and 7) showing there was little non-specific binding. Nor did either p12 protein pellet in the absence of CA-coated nanotubes under these conditions (lanes 4 and 8). In addition, cell lysates expressing either Fv1<sup>a</sup> or Fv1<sup> mutants were also incubated with the same CA-coated tubes as a positive and negative control for CA binding respectively [34]. Fig. S7 shows that we could detect binding of Fv1<sup> mutants to the nanotubes, but Fv1<sup>a</sup> had much reduced binding as expected (compare lanes 2 and 6), confirming that the CA was arranged in regular arrays that mimic true viral cores. Both Fv1 proteins showed weak binding to CA-P1G (lanes 3 and 7) indicating some non-specific binding. Together, this indicates that p12 does bind directly to the CA lattice and that the N-terminus of p12 is necessary for this interaction.

**Discussion**

The p12 protein of MLV is essential during the early stages of viral replication [8,10,11,35]. We previously showed that viruses bearing p12 proteins with alterations to their N-terminus were unable to abrogate restriction activity of TRIM5alpha or Fv1. There are various potential reasons for this (discussed in [10]), but as these restriction factors target the viral CA shell, this immediately suggested the intriguing possibility that mutations in p12 affected the viral core structure or stability. Influencing the properties of the core would explain the genetic interdependency between p12 and CA previously observed [10,36]. Furthermore, as maintaining the appropriate core stability is vital for retroviral replication, through myriad plausible mechanisms (reviewed in [5–7]), this might also explain the infectivity defects of N-terminal p12 mutants. Therefore, we set out to investigate the interplay between the N-terminal domain of p12 and the MLV CA core.

First, we confirmed that only the N-terminus of p12 is important for the abrogation phenotype (Fig. 1B and S1). Previously, we discovered that mixing a small proportion (approximately 10%) of wild type p12 into a N-terminal p12 mutant particle was enough to restore infectivity and abrogation capability [10]. However, we were unable to rescue the infectivity of N-terminal p12 mutants using increasing ratios of C-terminally altered p12 (Fig. 1C and [10]). Despite this, these particles were able to saturate restriction factors (Fig. 1B and S1), confirming that neither infectivity nor the C-terminus of p12 is necessary for this effect.

The p12 protein is a functional constituent of the MLV RTC [12]. Accordingly, wild type p12 co-sedimented on sucrose gradients with CA and reverse transcribed cDNA from infected cells (Fig. 2A). In addition, there was a population of p12 that co-sedimented with un-complexed CA at the top of the gradient, suggesting that either a proportion of the p12 in the particle is not incorporated into the core, as is likely for CA, or that p12 is lost from the core after entry into the cell. CA and p12 are incorporated into the virus as part of the Gag polyprotein and are thus present in particles at a 1:1 ratio. However, we consistently found it difficult to detect p12 by immunoblotting following velocity sedimentation of cell lysates. Consistent with our virological data that only a fraction of the p12 in the particles need be active [10], we observed a clear reduction in the amount of p12 present, compared to CA, in infected cell lysate (Fig. 2B). This suggests that RTCs contain less p12 than CA, and implies that any p12 not associated with the RTC is rapidly degraded. This presumed instability of free p12 is consistent with the failure to detect ectopically expressed p12 molecules in the absence of other viral proteins, unless tagged with a large, stable protein such as cherry fluorescent protein (unpublished data).

In a variety of different assays, we observed striking alterations in the characteristics of CA following mutation of p12. Analysis of N-terminal p12 mutant RTCs after separation of infected cell lysates on linear sucrose gradients revealed that they have a slower rate of sedimentation than wild type RTCs (Fig. 2C and D). The sedimentation coefficient of a particle, s, depends on its mass, density and shape, that impact on the frictional forces retarding the particle’s movement. Thus, from our experiments, we can state that CA is present in different complexes in N-terminal p12 mutants and wild type viruses. However, this could reflect a difference in composition (a different collection of proteins), conformation (a different arrangement of proteins) or the relative amounts of individual proteins in the complexes. Notably, virions with reduced core stability would likely result in RTCs with slower sedimentation rates. Importantly, when we studied viral particles directly, the N-terminal p12 mutants had altered CA assemblies (Fig. 3), implying that this is not due to target cell factors or an effect on reverse transcription. Furthermore, analysis of p12 mutant viral particles by TEM also revealed differences in their CA core when compared to wild type particles (Fig. 4 and S5). It should be noted that wild type Mo-MLV particles displayed considerable heterogeneity in virus particle morphology (Fig. 4A and F, and S3A), as previously observed [37] and evident in the distribution of CA throughout the sucrose gradients (Fig. 2 and 3) [38]. This presumably reflects biological variation but is possibly enhanced by preparation of the samples for TEM. However, there was a considerable increase in particles with absent or aberrant cores for the p12 mutants, most strikingly for p12 mutant 6 (Fig. 4). Initial consideration of the TEM data suggested that p12 mutant 6 fails to form electron dense mature CA cores. However, approximately half the mutant 7 and 8 particles have formed mature cores, and so two alternative mechanisms for the function of p12 mutant 6 versus mutants 7 and 8 would need to be proposed. This does not seem credible, especially when all the N-terminal p12 mutants behave in a similar way in the biophysical assays (Fig. 2 and 3). Alternatively, mutations in p12 could result in a less stable core, with variation in the absolute stability depending on the N-terminal alteration. In the case of mutant 6, the core may be so unstable that it falls apart during formation or is more sensitive to disruption by preparation for TEM, while the other N-terminal mutants have a less severe core stability defect.

Further evidence for a core stability defect in p12 mutants comes from our immunofluorescence data. We could detect both p12 and CA containing complexes in cells challenged with wild
type virus and N-terminal p12 mutants (Fig. 6 and S6). This implies that at least some fraction of CA and p12 are in complexes, even for p12 mutant 6 where the majority of particles contained minimal electron dense material (Fig. 4). In wild type particles, CA signal was slowly lost with time (unpublished data), likely due to uncoating and integration events. In keeping with our earlier observations, the p12 signal was lost faster than CA. However, we observed a more rapid loss of both p12 and CA puncta for the p12 mutants compared to wild type virus (Fig. 7). This suggests that the N-terminal domain of p12 is required for retention of p12 within

Figure 7. Kinetics of p12 and CA loss from cells infected with Mo-MLV. Ten cells were randomly chosen (based solely on nuclear stain) and imaged using a spinning disk confocal microscope. The numbers of p12(myc) or CA puncta in each cell were determined. The mean number of puncta per cell was normalised to the mean number of CA or p12(myc) puncta present at time point zero for each infection. The amount of p12 (dashed lines) and CA (solid lines) puncta are displayed against time in the graphs for: (A) p12 mutant 5, (B) p12 mutant 6, (C) p12 mutant 7, (D) p12 mutant 8 and (E) p12 mutant 14. Wild type Mo-MLV data (p12, black dashed and CA, black solid lines) are also plotted in each graph as a reference. Each point indicates the mean with SEM error bars. (F) The normalised numbers of CA and p12(myc) puncta in cells two hours post-infection are displayed as a bar chart showing the mean with SEM error bars.
doi:10.1371/journal.ppat.1004474.g007
SDS-PAGE sample buffer and probed for CA and p12 by immunoblot-through a sucrose cushion. The pelleted material was resuspended in incubated with lipid nanotubes coated with WT CA (lanes 2 and 6), P1G NTA. Purified wild type p12 (lanes 2–4) or p12 Mutant 6 (lanes 6–8) were coated lipid nanotubes.

Figure 8. Binding of purified p12 to N-MLV CA-coated lipid nanotubes. Multimeric arrays of wild type (WT) and P1G mutant N-MLV CA were generated by immobilising the His-tagged purified proteins on lipid nanotubes comprising the N2+-chelating lipid, DG-NTA. Purified wild type p12 (lanes 2–4) or p12 Mutant 6 (lanes 6–8) were incubated with lipid nanotubes coated with WT CA (lanes 2 and 6), P1G CA (lanes 3 and 7) or no tubes (lanes 4 and 8) prior to centrifugation through a sucrose cushion. The pelleted material was resuspended in SDS-PAGE sample buffer and probed for CA and p12 by immunoblotting, using appropriate antibodies. Input, lanes 1 and 5, represents 1/40th dilution of the purified p12 proteins before incubation with CA-coated lipid nanotubes.

doii:10.1371/journal.ppat.1004474.g008

the RTC, and indicates that the presence of p12 prevents premature loss of CA complexes. Thus, the core formed in the presence of N-terminal mutant p12 appears to be less stable.

As p12 has a role in nuclear retention and integration, and only a small proportion of the p12 in the virion is required for infectivity, it is possible that p12 is retained in the PIC by virtue of an interaction with integrase or the viral cDNA. Interestingly, allosteric integrase inhibitors have been shown to affect HIV-1 core morphology [39], suggesting that various viral components are involved in mature core formation. However, given the striking effects p12 mutation has on CA complexes, and the genetic evidence from chimeric viruses [10,36], CA is the obvious binding partner for p12, and in particular the mature CA lattice. Using CA-coated lipid nanotubes, we were able to show that p12 does indeed bind to CA lattices but that there was no detectable binding for p12 mutant 6 (Fig. 8). A reduction in CA binding therefore correlates with the rapid loss of p12 puncta from cells and with the core morphology defects observed. Not only does this imply that p12 stabilises the CA lattice directly, but this is the first demonstration of direct protein binding to both MLV CA and p12.

As Gag proteins initially assemble into immature Gag lattices, we analysed the effect of p12 mutations on immature particles by equilibrium sedimentation and TEM. We observed no differences in the density (Fig. S5) or particle morphology between wild type and p12 mutant 6 virions, using either PR- mutants or imaging budding particles from producer cells (Fig. 5). Taken together with the fact that the mutants all produce similar numbers of particles to wild type MLV, and that individual Gag protein content is the same between wild type and p12 mutants by immunoblotting and ELISA [10], this implies that assembly and Gag processing are unaffected. However, there may be subtle alterations to these processes that we cannot detect, although the phenotype is certainly exaggerated post-maturation.

After proteolytic cleavage, the immature Gag lattice is thought to break down completely allowing the CA proteins to undergo a huge rearrangement [2,25]. The current dogma is that CA alone determines the correct formation of the hexameric lattice structure seen in the mature core. Certainly, CA proteins from many retroviruses are able to form ordered hexameric lattices in vitro. However, in most systems, extra elements are present. For example, CA has often been fused to viral NC, and RNA has been introduced to seed the lattice formation [40,41]. Arguably, CA fusions would represent immature lattices and not biologically mature core structures. Alternatively, CA arrays have been formed on lipid scaffolds using His-tagged CA proteins [34,42,43]. This has the advantage of increasing the avidity of CA interactions, perhaps relieving the need for additional catalysis or stabilisation. In addition, HIV-1 and RSV CA can be induced to form higher order assemblies by adding salt or crowding agents and/or altering pH [40,44–47]. Nevertheless, many of these in vitro formed CA lattices have higher order structures, wide tubes or sheets, not observed in virions [41,42,45]. For HIV, a disulphide crosslinking strategy was deployed to enable purification and crystallization of soluble HIV-1 CA hexamers [48,49], reinforcing the fact that stabilisation of the CA lattice requires some assistance. Perhaps the requirement for disassembly, or uncoating, of the CA shell necessitates the need for an inherently unstable mature CA lattice.

However, it seems plausible that in virions, additional viral factors cooperate with CA to form a mature CA core of optimal stability. In the case of MLV, our data would suggest that p12 is required for this function. Moreover, p10 from Rous sarcoma virus (RSV), which is positionally analogous to p12 in Gag, has been shown to alter the morphology of CA-NC constructs formed in vitro from cylinders into spherical particles [41,50]. Significantly, although viral production was somewhat reduced, alterations to the C-terminus of p10 also altered the mature RSV core morphology in vivo [51,52].

Alpha-, beta-, gamma- and epsilon-retroviral genera all encode additional Gag cleavage products between MA and CA. Most of these are poorly characterised, but as mature CA lattices are thought to have similar arrangements in all retroviruses [42], it is tempting to speculate these additional Gag cleavage products function in an equivalent manner to MLV p12. Interestingly, one feature that they seem to have in common is that they harbour the late (Lc) domain essential for viral budding [53,54]. Although lentiviruses do not have an analogous protein between MA and CA, they too contain an L-domain. In the case of HIV-1, the L-domain is found in another protein, p6, cleaved from the end of Gag [53,54]. L-domains therefore represent excellent examples of functional conservation despite little positional or sequence similarity. Moreover, despite different L-domains having alternative primary binding partners [53,54], they have been shown to be functionally interchangeable [55]. Interestingly, HIV-1 p6 has been reported to alter core assembly by regulating CA processing [56]. Whether HIV-1 also contains a factor that stabilises the mature CA lattice remains to be seen; there is still a lot to learn about the formation and subsequent breakdown of the mature retroviral core structure. Nevertheless, the CA shell of the HIV-1 core has become an attractive drug target and alterations to core stability possibly influence immune responses to infection as well as local particle infectivity [57].

Overall, we have shown that p12 binds directly to CA lattices and that mutations in p12 that disturb this association have debilitating effects on the CA core, before the virus even infects a target cell. At its most striking, this defect manifests as failure to produce stable electron dense cores (mutant 6), and this correlates with a reduction in the ability of virions to reverse transcribe. Given the current notion that CA is vital for HIV integration events [5–7], it is tempting to speculate that an unstable core is responsible for the lack of integration seen for the N-terminal p12 mutants that can reverse transcribe normally (mutants 5, 7, and 8).

Whilst this may be partly true, p12 is also required for chromatin tethering of the MLV PIC [10,13,14], and so we cannot discern
whether the concomitant loss of p12 from the RTC or core instability itself is the cause of the infectivity defect and this will require further experimentation. Curiously, mutations to the C-terminus of p12 appeared to stabilise the CA complexes somewhat, particularly in virions (Fig 2 and 3). Although the p12 proteins from C-terminal p12 mutants are still associated with the viral PIC and are present in the target cell nucleus, the viral cDNA is also unable to integrate. Determining the precise function of the C-terminus of p12 and identifying any C-terminal interaction factors may shed light on why CA assemblies from these mutants have altered phenotypes in our biophysical assays. Future work will endeavour to determine the interaction interface between p12 and the CA core to understand how the interaction affects core stability.

Materials and Methods

Plasmids and cloning

Three plasmids were co-transfected to synthesize retroviral VLPs: An envelope expression plasmid for either vesicular stomatitis virus G protein (pczVSV-G) [58] or the Mo-MLV ecotropic envelope (pMoSAF) [59]; a Mo-MLV-based retroviral vector encoding LacZ (pczLTR-LacZ) [60] or eGFP (pLNCG) [58,61]; and either Mo-MLV (pKB4) [10] or N-tropic MLV (pCI G3N) [58] Gag-Pol expression plasmids. The generation of p12 mutations in these Gag-Pol expression plasmids has been described previously [10]. To create Mo-MLV Gag-Pol gene expression plasmids containing a previously described myc-tag in p12 [12], a BsrGI-Xhol fragment from pNCS p12 1×MyeR was swapped into pKB4 creating pKB4-mycE. This was also done for p12 mutant 5 by cloning the same fragment from pNCS-PM5 p12 1×MyeR into pKB4 creating pKB5-mycE. To create the other p12 mutant Gag-Pol plasmids containing myc-tagged p12, site directed mutagenesis was performed on pKB4-mycE using the Quik-Change kit (Stratagene). The following primers were used: p12 mutant 6 for 5′-gcaacctaaaacactaagttcttttctgacagtgcggcggcggccgccgcactgtca-9 5′-gccaaacctaaacctcaagctgctgctgccgctggggggccgctcatcga-3 and reverse primer 5′-tcttgggtccctataaggcgggggtgctgctgcagctgcggcgatgagcggcccccc-3.

Infections

VLP infectivity was determined as previously described [10]. Briefly, D17 cells were challenged with equivalent RT-units of LacZ-encoding VLPs. After 48–72 hours, the β-galactosidase activity in the cell lysate was measured using the Galacto-Star system (Applied Biosystems).

Restriction factor saturation assays

Restriction factor saturation assays were performed as previously described [10]. Briefly, TE671 cells expressing endogenous human TRIM5alpha were infected with 2-fold serial dilutions of freshly harvested 293T cell supernatants containing LacZ-encoding VLPs. Cultures were incubated for 4–6 hours before adding a fixed amount of GFP encoding N-MLV. After 72 hours, infected cells were harvested and the percentage of GFP positive cells was determined by flow cytometry using a FACS Calibur analyzer (Becton Dickinson).

Modified Mo-MLV fate-of-capsid assay

D17 cells were seeded at 1×10⁶ cells per well in a 6-well plate one day prior to infection. Each well was infected with 2 ml LacZ-encoding Mo-MLV VLPs (with 10 μg/ml polybrene) by spinoculation at 4°C (1600×g for 30 minutes). After 4 hours infection at 37°C, cells were washed and resuspended in 700 μl hypotonic buffer (10 mM Tris-HCl pH 8.0, 10 mM KCl, 1 mM EDTA supplemented with complete protease inhibitors). After 15 minutes incubation on ice, the cell suspension was applied to a QIAshredder column (Qiagen) and the subsequent cell lysate was layered on top of 10–42% (w/w) linear sucrose gradient using a syringe pump-driven gradient fractionator (Brandel). For each fraction, protein and viral DNA content were analysed using immunoblotting and quantitative PCR (qPCR) respectively. Proteins from each sucrose fraction were isolated and subjected to trichloroacetic acid precipitation and resuspended in 1× protein loading buffer.

Biophysical analysis of virions in sucrose gradients

VLP containing cell supernatant was concentrated through a 20% (w/v) sucrose cushion in an SW41 rotor (Beckman Coulter) at 28,300 rpm for 90 minutes, 4°C. Supernatant and sucrose were aspirated and the viral pellet was resuspended in 250 μl PBS on ice for 4–5 hours. Linear 10–42% (w/v) sucrose gradients were prepared by co-transfection of 293T cells with a 1:1:1 ratio of three plasmids encoding the appropriate wild type or mutant Gag-Pol, VSV-G or MLV ecotropic Env, and a reporter gene (β-galactosidase or GFP) respectively, using polyethyleneimine (PEI, PolySciences) as a transfection reagent. To make mixed mutant viral particles, two mutant N-MLV Gag-Pol expression plasmids were added to the transfection mix at different ratios, keeping the total concentration of Gag-Pol plasmid constant. After ~24 hours, cells were washed and fresh media was added for a further ~15 hours. Virus containing supernatants were harvested, filtered, and viral titres were quantified using a modified ELISA for reverse transcriptase activity (Cavidel). In the restriction factor saturation assays, Turbofect (Fermentas) was used as transfection reagent. At approximately 18 hours after transfection, cells were washed and sodium butyrate media (0.01 M sodium butyrate, 10% FCS and 1% penicillin/streptomycin in DMEM) was added for 6 hours before replacing with fresh media. VLPs were then harvested after ~15 hours, as above. For qPCR experiments, viruses were treated with RQ1-DNase (Promega) prior to infection.
formed in 14 × 89 mm polycarbonate ultracentrifuge tubes (Beckman Coulter) with either: 250 μl of 1% Triton X-100 in 5% (w/v) sucrose PBS or just 5% (w/v) sucrose PBS (for intact VLP analysis) on top; followed by 250 μl of 2.5% (w/v) sucrose PBS. Concentrated virus was gently layered on top of the gradients (5 μl was kept for an input control), and spun in an SW41 rotor at 28,500 rpm for 16 hours at 4 °C. Fractions were collected using a syringe pump-driven fractionator (Brandel) and each fraction was diluted in 4× protein loading buffer.

**Immunoblot analysis**

All VLPs analysed by immunoblotting were concentrated by centrifugation through a 20% (w/v) sucrose cushion for 1 hour at 16,000 × g, 4 °C and resuspended in 1× protein loading buffer. Proteins were separated by SDS polyacrylamide gel electrophoresis (SDS-PAGE) and transferred onto polyvinylidene fluoride (PVDF, Millipore) membrane. Immunoblotting was performed with a rat anti-p30CA (hybromeda CRL-1912, ATCC), mouse anti-p12 monoclonal (hybromeda CRL-1890, ATCC), goat anti-p12 polyclonal (a gift from J. Stoye), mouse anti-His (Penta-His Antibody, Qiagen) or rabbit anti-Fv1 (a gift from J. Stoye) followed by anti-rat, anti-goat, anti-mouse or anti-rabbit HRP-conjugated secondary antibodies. Detection was performed using the Immobilon chemiluminescent substrate (Millipore) and hyperfilm processed through a Fujifilm FPM-3800A developer.

**Quantitative PCR (qPCR) analysis**

The quantity of viral DNA in each sucrose fraction was analysed using qPCR detection of minus strand strong stop reverse transcription products as described previously [10]. Briefly, reactions were performed in triplicate using Taqman Gene Expression Master Mix (Applied Biosystems) with 900 nM of each primer: oJWB45 (5'-cctgacgggtagtcaatcactcag); oJWB47 (5'-ctgaaggctgatgataagaag); and 250 nM of probe oJWB38 (5'-FAM-WWGCATCGTGGTCTTTCT-TAMRA) [62]. The PCR reactions were performed with a Fast 7500 PCR system (Applied Biosystems) using standard cycling conditions: 50 °C for 2 minutes, 95 °C for 10 minutes followed by 40 cycles of 95°C for 15 seconds and 60°C for 1 minute. Relative cDNA copy number was determined by comparison to a dilution series of the LacZ-LTR plasmid in D17 cellular DNA.

**Transmission electron microscopy**

Large batches of wild type and p12 mutant Mo-MLV VLPs were synthesised by transient transfection in 293T cells as described above with one modification: Transfected cells were washed and incubated in DMEM without added serum before viruses were harvested. VLP-containing 293T cell supernatant was pelleted by centrifugation at ~100,000 × g in an SW32 rotor (Beckman Coulter) using an Optima L-90K ultracenrifuge (Beckman Coulter). Pelleted VLPs were resuspended in 20 μm-filtered PBS on ice for 30 minutes. Resuspended viral pellets (300 μl total) were pooled and 300 μl of 5% gluteraldehyde/0.2 M sodium cacodylate buffer (Fisher Scientific and Sigma Aldrich, respectively) was added, mixed and fixed for 1 hour on ice. Virus was pelleted at 17,800 × g for 1 hour at room temperature. The fixative was removed and 250 μl warm (37 °C) 2% low melting point (LMP) agarose (Fisher Scientific) placed on top and mixed. Virus was then centrifuged at 16,200 × g for 20 minutes in a centrifuge heated to 38 °C, immediately followed by incubation on ice. After 30 minutes, ice-cold 2.5% gluteraldehyde/0.1 M sodium cacodylate buffer was layered on top of the solid agarose (up to the top of the tube). This was left overnight on ice to completely set the agarose. For VLP producer cell samples, approximately 2×10^7 cells were washed from the culture dish with PBS and gently pelleted at 500 × g for 5 minutes in a bench top centrifuge (4 °C). The cell pellet was gently resuspended in 2.5% gluteraldehyde, 0.1M sodium cacodylate and pelleted at 600 × g for 10 minutes (4 °C). Fixation was continued overnight on ice.

Both the cell pellet and the VLP pellet set in LMP agarose were post-fixed with 1% osmium tetroxide for 90 minutes and washed with 0.1M sodium cacodylate. Samples were then stained with 1% aqueous uranyl acetate for 90 minutes and dehydrated in an ethanol series before propylene oxide. All samples were then embedded in medium Agar 100 resin and polymerised overnight at 70 °C. 50 nm thick sections were stained with saturated ethanolic uranyl acetate and Reynolds’s lead citrate. Samples were viewed on a Jeol 1200EX transmission electron microscope (Jeol Ltd) operating at 60 kV and a magnification of ×20,000. Images of cells and purified particles were captured on an Orion CCD camera (Gatan) using the auto exposure mode. Quantification of core morphology from purified VLPs was performed on randomly selected micrographs and only particles between 90–120 nm in diameter were scored (at least 93 individual particles were scored for each sample).

**Indirect immunofluorescence**

U/R cells were seeded at 4×10^4 on sterile 13 mm coverslips (VWR) in a standard 12-well plate (Corning). Cells were challenged 16 hours later with ecotropic wild type or p12 mutant Mo-MLV VLPs at an MOI 3 (mutants were normalised to wild type by the level of RT activity) by spinoculation (1000 × g for 2 hours). Infections were done in duplicate from a single batch of diluted virus due to the differing antigen retrieval conditions required for the p12(myc) and CA antigens. Infected cells were then washed three times with pre-warmed DMEM complete and fixed at the indicated time points as follows: (i) for p12(myc) detection; 4% paraformaldehyde (AlfaAesar) for 20 minutes then 0.1% triton X100 in PBS for 10 minutes or (ii) for CA detection; 4% paraformaldehyde 2 minutes followed by −20 °C methanol for 5 minutes. Cells were washed three times with PBS and blocked in 5% bovine serum albumin (BSA, Fisher Scientific) for 1 h. Primary monoclonal antibodies (hybriodima supernatants) were diluted 1:6 in 1% BSA: mouse anti-myc 9E10 [12] or rat anti-p30CA (CRL-1912, ATCC); and incubated on the cells for 1 hour. Coverslips were then washed with PBS three times for 10 minutes each and secondary antibodies diluted in 1% BSA were added for 1 hour. The goat anti-mouse Cy3-conjugated or goat anti-rat FITC-conjugated antibodies (Jackson ImmunoResearch laboratories) were diluted 1:300 or 1:100, respectively. DAPI stain was added to the slides together with the secondary antibodies.

**Fluorescence microscopy**

Images were acquired with either a spinning disk confocal (Yokogawa CSU-22 Confocal Head) microscope (Axiovert 200 M, Carl Zeiss MicroImaging) or an Ultraview spinning disk confocal microscope (Perkin Elmer) equipped with a C9100-13 electron multiplying charged-coupled device (EMCCD, Hamamatsu). For quantification, the entire cell volume was imaged by confocal microscopy and the picture was deconvoluted using the Nearest Neighbors deconvolution algorithm of SlideBook. Subsequently, three dimensional acquisitions were projected on to a two dimensional plane. After this, the specific signals of the p12-based and CA-based staining were identified through intensity based segmentation and the number of objects (puncta) in the inspected cells was determined. Approximately 250 dots of p12 or CA signal per cell at time zero were analysed. All the above steps were
performed using the SlideBook software (Intelligent Imaging Innovations).

Recombinant protein expression and purification

Wild type and mutant p12 sequences from N-tropic MLV were cloned into pGEX6.1 using the BamHI and XhoI restriction endonuclease sites. The N-terminally GST-tagged fusion proteins were expressed in E. coli Rosetta 2(DE3)pLysS by inducing a mid-log culture grown in the presence of 1% glucose with 1 mM isopropyl-β-D-thiogalactopyranoside (IPTG). For lysis, cells were resuspended in 50 mM Tris pH 8, 500 mM NaCl, 0.5 mM TCEP, 0.1% Triton X-100 (Buffer A) in the presence of protease inhibitors (Roche) and incubated with Lysozyme (Sigma Aldrich) and Benzonase nuclease (Sigma Aldrich) for 1 hour at 4°C. The crude lysates were then sonicated twice for 5 minutes at 40% amplitude and centrifuged at 48,000 x g for 45 minutes to remove debris. The clarified lysates were applied to 1 ml GST-trap columns (GE Healthcare). After washing with Buffer A, untagged-p12 was eluted from the resin by digestion with 3C precision protease. The eluate was then heated at 65°C for 10 minutes and centrifuged at 40,000 x g for 20 minutes to remove precipitates. Acetic acid (pH ~3) was added to the supernatant which was then centrifuged at 40,000 x g for 20 minutes to remove DNA. The supernatant was then applied to a Superdex 75 (16/60) size exclusion column equilibrated in 200 mM Ammonium bicarbonate. Eluate fractions containing p12 were pooled and lyophilised. The purity of the protein preparations were assessed by SDS-PAGE and the concentrations were determined from the absorbance at 280 nm.

C-terminally His-tagged N-MLV CA WT and P1G mutant proteins were expressed and purified as previously described [34].

Binding assays with CA-coated lipid nanotubes

The assays were performed essentially as previously described [34]. Lipid nanotubes were generated by combining the tube-forming lipid, d-galactosyl-β-1,1’ N-neurovenosyl-d-erythro-sphingosine (GalCer) (Avanti) with the Ni²⁺-chelating lipid, DGS-NTA (Avanti) in a 7:3 ratio. After mixing the lipids, residual cholesterol and methanol were removed under a gentle stream of nitrogen. The tubes were then heated at 80°C for 20 minutes to remove DNA. The supernatants were collected and analysed by immunoblotting using an anti-CA antibody. Representative immunoblots are shown (Fraction 1 is the top of the gradient). (B) For each experiment, the sucrose density of the fraction containing the peak CA signal was measured, and the change in density compared to peak CA fraction for wild type virions was calculated. The mean and range of two independent experiments are displayed in the histogram. (TIF)

Figure S2 Migration profile of intact mature p12 mutant VLPs in an equilibrium gradient. (A) Purified VLPs were subjected to equilibrium sedimentation through a 10–42% (w/w) sucrose gradient (without detergent). Fractions were collected and analysed by immunoblotting using an anti-CA antibody. Representative immunoblots are shown (Fraction 1 is the top of the gradient). (B) For each experiment, the sucrose density of the fraction containing the peak CA signal was measured, and the change in density compared to peak CA fraction for wild type virions was calculated. The mean and range of two independent experiments are displayed in the histogram. (TIF)

Figure S3 Additional electron micrographs of purified wild type and p12 mutant VLPs. Mo-MLV VLPs were purified and prepared for TEM, as in Fig. 4. Additional electron micrographs of (A) wild type, (B) p12 mutant 6, (C) p12 mutant 7 and (D) p12 mutant 8 VLPs are shown. All scale bars are 50 nm. (TIF)

Figure S4 The infectivity of p12 mutant VLPs used for transmission electron microscopy. D17 cells were challenged with equivalent RT-units of the wild type and p12 mutant LacZ-encoding VLPs used in the TEM analysis (Fig. 4 and S3). Infectivity was measured by detection of beta-galactosidase activity in a chemiluminescent reporter assay and plotted as a percentage of wild type N-MLV infectivity. (A) Wild type and p12 mutant 6 VLPs, (B) Wild type and p12 mutant 7 VLPs and (C) Wild type and p12 mutant 8 VLPs. (TIF)

Figure S5 Migration profile of intact immature p12 mutant VLPs in an equilibrium gradient. A mutation was introduced into PR to inactivate its activity (D32L, called PR-). (A) PR- wild type and p12 mutant VLPs were subjected to equilibrium sedimentation through a 10–42% (w/w) sucrose gradient (without detergent). Fractions were collected and analysed by immunoblotting using an anti-CA antibody. Representative immunoblots are shown (Fraction 1 is the top of the gradient). (B) For each experiment, the sucrose density of the fraction containing the peak CA signal was measured, and the change in density compared to peak CA fraction for wild type virions was calculated. The mean and range of three independent experiments are displayed in the histogram. (TIF)

Figure S6 Immunofluorescence of p12 and CA in cells infected with Mo-MLV p12 mutant 5 and 14. U/R cells were challenged with ecotropic wild type, p12 mutant 5 or p12 mutant 14 Mo-MLV VLPs, containing a myc-tag in p12, by cold spinoculation (MOI 5). Cells were fixed at various times post-infection and stained with either an (A) anti-myc or (B) anti-CA antibody followed by a Cy3 (A) or FTTC (B) -conjugated secondary antibody. The nuclear DNA was counterstained using DAPI.
Multimeric arrays of wild type (WT) and P1G mutant N-MLV CA were generated by immobilising the His-tagged purified proteins on lipid nanoparticles comprising the Ni2+-chelating lipid, DGS-NTA. Cell lysates containing Fv1n (lanes 2–4) or Fv1b (lanes 6–8) were incubated with lipid nanotubes coated with WT CA (lanes 2 and 6), P1G CA (lanes 3 and 7) or no tubes (lanes 4 and 8) prior to centrifugation through a sucrose cushion. The pelleted material was resuspended in SDS-PAGE sample buffer and probed for CA and Fv1 by immunoblotting, using appropriate antibodies. Input, lanes 1 and 5, represents 1/10th dilution of cell extracts before incubation with CA-coated lipid nanoparticles.

Table S1 Numbers of CA and p12 puncta counted in immunofluorescent studies. The mean number of CA or p12 puncta present per cell at time zero or 2 hours post-infection is shown ± standard errors. The numbers of puncta at 2 hours post-infection were normalised to the numbers of puncta at time zero and these data are represented graphically in the bar chart in Figure 7F.

Acknowledgments
We thank Neil Ball for help with protein purification and Jonathan Stoye and Ian Taylor for helpful discussions. We would also like to acknowledge the confocal microscopy facility at NIMR for their advice and expertise, especially Donald Bell and Kate Sullivan.

Author Contributions
Conceived and designed the experiments: DJW VCB MW EE KNB. Performed the experiments: DJW VCB MW EE KNB. Contributed reagents/materials/analysis tools: WL ME. Wrote the paper: DJW VCB MW EE KNB.

References
1. Katoh I, Yoshinaka Y, Reim A, Shiibuya M, Odaka T, et al. (1985) Murine leukemia virus maturation: protease region required for conversion from “immature” to “mature” core form and for virus infectivity. Virology 145: 280–292.
2. de Marco A, Muller B, Glass B, Riches JD, Krausslich HG, et al. (2010) Structural analysis of HIV-1 maturation using cryo-electron tomography. PLoS Pathog 6: e1001213.
3. Wohulet AA, Koizumi F (2012) HIV type 1 Gag as a target for antiviral therapy. Aids Research and Human Retroviruses 28: 54–75.
4. Johnson VA, Calvez V, Gunthur HF, Paredes R, Pillay D, et al. (2013) Update of the drug resistance mutations in HIV-1: March 2013. Top Antirud Med 21: 6–14.
5. Fassati A (2012) Multiple roles of the capid protein in the early steps of HIV-1 infection. Virus Res 170: 15–24.
6. Hilditch I, Towers GJ (2014) A model for cofactor use during HIV-1 reverse transcription and nuclear entry. Curr Opin Virol 4: 32–36.
7. Ambrose Z, Aiken C (2014) HIV-1 uncoating: connection to nuclear entry and regulation by host proteins. Virology 454–455: 371–379.
8. Yuan B, Li X, Goff SP (1999) Mutations altering the moloney murine leukemia virus p12 Gag protein affect virion production and early events of the virus life cycle. The EMBO Journal 18: 4700–4710.
9. Martin-Serrano J, Eastman SW, Chung W, Bieniasz PD (2005) HECT ubiquitin ligases link viral and cellular PPXY motifs to the vacuolar protein-sorting pathway. Journal of Cell Biology 168: 89–101.
10. Wight DJ, Bouchert VC, Nader M, Allen DJ, Taylor IA, et al. (2012) The gammaretroviral p12 protein has multiple domains that function during the early stages of replication. Retrovirology 9: 83.
11. Yuan B, Fassati A, Yueh A, Goff SP (2002) Characterization of Moloney Murine Leukemia Virus p12 Mutants Blocked during Early Events of Infection. Journal of Virology 76: 10810–10810.
12. Prizan-Ravid A, Ellis E, Laham-Karam N, Selig E, Ehrlich M, et al. (2010) The Gag cleavage product, p12, is a functional constituent of the murine leukemia virus pre-integration complex. PLoS Pathog 6: e1001183.
13. Schneider WM, Brezinski JD, Ayer S, Malani N, Gryzyczka M, et al. (2013) Viral DNA tethering domains complement replication-defective mutations in the p12 protein of MuLV Gag. Proc Natl Acad Sci U S A 110: 9487–9492.
14. Ellis E, Ehrlich M, Prizan-Ravid A, Laham-Karam N, Bacharak E et al. (2012) p12 tethers the murine leukemia virus pre-integration complex to mitotic chromosomes. PLoS Pathog 8: e1002832.
15. Wu X, Li Y, Cirne B, Burgess SM (2003) Transcription start regions in the human genome are favored targets for MLV integration. Science 300: 1749–1751.
16. Mitchell RS, Beitzel BF, Schroder AR, Shin P, Chen H, et al. (2004) Retroviral DNA integration: ASLV, HIV, and MLV show distinct target site preferences. PLoS Biol 2: e234.
17. Berry C, Hannenhalli S, Leipzig J, Busman FD (2006) Selection of target sites for mobile DNA integration in the human genome. PLoS Comput Biol 2: e157.
18. Ayer S, Swapan GV, Malani N, Aramini JM, Schauder WM, et al. (2014) Altering murine leukemia virus integration through disruption of the integrase and BET protein family interaction. Nucleic Acids Res 42: 5917–5928.
19. Sharma A, Larce RC, Plumb MR, Malani N, Male F, et al. (2013) BET proteins promote efficient murine leukemia virus integration at transcription start sites. Proc Natl Acad Sci U S A 110: 12036–12041.
20. Gupta SS, Marzag T, Maertens GN, Sharif A, Rothe M, et al. (2013) Bromo- and extra-terminal domain chromatin regulators serve as cofactors for murine leukemia virus integration. J Virol 87: 12721–12736.
21. De Rijck J, de Reger C, Demuulemeester J, Vets S, El Ashkar S, et al. (2013) The BET family of proteins targets murine leukemia virus integration near transcription start sites. Cell Rep 5: 886–894.
22. Zhang F, Zang T, Wilson SJ, Johnson MC, Bieniasz PD (2011) Chlatherin facilitates the morphogenesis of retrovirus particles. PLoS Pathog 7: e1002119.
23. Popova S, Strack B, Sanchez-Merino V, Popova E, Rossin H, et al. (2011) Human immunodeficiency virus type 1 and related primate lentiviruses engage chlatherin through Gag-Pol or Gag. J Virol 85: 3792–3801.
24. Fassati A, Goff SP (1999) Characterization of intracellular reverse transcription complexes of Moloney murine leukemia virus. J Virol 73: 8919–8925.
25. Bharat TA, Davey NE, Ulbrich P, Riches JD, de Marco A, et al. (2012) Structure of the immature retroviral capsid at 8 A resolution by cryo-electron microscopy. Nature 487: 385–389.
26. Weager M, Wilson-Kubalek EM, Weigner SG, Brown PO, Reim A (1996) Supramolecular organization of immature and mature murine leukemia virus revealed by electron cryo-microscopy: implications for retroviral assembly mechanisms. Proc Natl Acad Sci U S A 95: 7299–7304.
27. Briggs JA, Riches JD, Glass B, Bartonova V, Zanetti G, et al. (2009) Structure and assembly of immature HIV. Proc Natl Acad Sci U S A 106: 11090–11095.
28. de Marco A, Davey NE, Ulbrich P, Phillips JM, Lux V, et al. (2010) Conserved and variable features of Gag structure and arrangement in immature retrovirus particles. J Virol 84: 11729–11736.
29. Yoshinaka Y, Laufft RB (1977) Murine leukemia virus morphogenesis: cleavage of P70 in vitro can be accompanied by a shift from a concentrically coiled internal strand (“immature”) to a collapsed (“mature”) form of the virus core. Proc Natl Acad Sci U S A 74: 3446–3450.
30. Yoshinaka Y, Laufft RB (1977) Properties of a P70 proteolytic factor of murine leukemia virus. J Virol 28: 769–779.
31. Fu W, Reim A (1993) Maturation of dimeric viral RNA of Moloney murine leukemia virus. J Virol 67: 5443–5449.
32. Forhsey BM, von Schwedler U, Sandquist WI, Aiken C (2002) Formation of a human immunodeficiency virus type 1 core of optimal stability is crucial for viral replication. Retrovirology 9: 83.
36. Lee SK, Nagashima K, Hu WS (2005) Cooperative effect of gag proteins p12 and capsid during early events of murine leukemia virus replication. J Virol 79: 4159–4169.
37. Auerbach MR, Brown KR, Singh IR (2007) Mutational analysis of the N-terminal domain of Moloney murine leukemia virus capsid protein. J Virol 81: 12337–12347.
38. Bacharach E, Gonsky J, Alin K, Orlova M, Goff SP (2000) The carboxy-terminal fragment of nucleolin interacts with the nucleocapsid domain of retroviral gag proteins and inhibits virion assembly. J Virol 74: 11027–11039.
39. Jurado KA, Wang H, Slaughter A, Feng L, Kesel JJ, et al. (2013) Allosteric integrase inhibitor potency is determined through the inhibition of HIV-1 particle maturation. Proc Natl Acad Sci U S A 110: 8690–8695.
40. Gross I, Hoehenberg H, Krauslich HG (1997) In vitro assembly properties of purified bacterially expressed capsid proteins of human immunodeficiency virus. European Journal of Biochemistry 249: 592–600.
41. Campbell S, Vogt VM (1995) Self-assembly in vitro of purified CA-NC proteins from Rous sarcoma virus and human immunodeficiency virus type 1. J Virol 69: 6407–6457.
42. Ganser BK, Cheng A, Sandquist WI, Yeager M (2003) Three-dimensional structure of the M-MuLV CA protein on a lipid monolayer: a general model for retroviral capsid assembly. EMBO J 22: 2086–2092.
43. Mayo K, Yama ML, McDermott J, Huebey D, Lui J, et al. (2002) Analysis of Rous sarcoma virus capsid protein variants assembled on lipid monolayers. J Mol Biol 316: 667–678.
44. Ehrlich LS, Agresta BE, Carter CA (1992) Assembly of recombinant human immunodeficiency virus type 1 capsid protein in vitro. J Virol 66: 4674–4683.
45. Li S, Hill CP, Sundquist WI, Finch JT (2000) Image reconstructions of helical assemblies of the HIV-1 CA protein. Nature 407: 409–413.
46. Bailey GD, Hyun JK, Mitra AK, Kingston RL (2009) Proton-linked dimerization of a retroviral capsid protein initiates capsid assembly. Nature 503: 402–405.
47. Kingston RL, Fitzon-Ostendorp T, Eisenmesser EZ, Schatz GW, Vogt VM, et al. (2000) Structure and self-association of the Rous sarcoma virus capsid protein. Structure 8: 617–628.
48. Pornillos O, Ganser-Pornillos BK, Banumathi S, Hua Y, Yeager M (2010) Disulfide bond stabilization of the hexameric capsomer of human immunodeficiency virus. J Mol Biol 401: 985–993.
49. Pornillos O, Ganser-Pornillos BK, Kelly BN, Hua Y, Whitby FG, et al. (2009) X-ray structures of the hexameric building block of the HIV capsid. Cell 137: 1292–1299.
50. Campbell S, Vogt VM (1997) In vitro assembly of virus-like particles with Rous sarcoma virus Gag deletion mutants: identification of the p10 domain as a morphological determinant in the formation of spherical particles. J Virol 71: 4425–4435.
51. Joshi SM, Vogt VM (2000) Role of the Rous sarcoma virus p10 domain in shape determination of gag virus-like particles assembled in vitro and within Escherichia coli. J Virol 74: 10260–10268.
52. Schicile LZ, Kenney SP, Cairns TM, Craven RC, Parent LJ (2007) Overlapping roles of the Rous sarcoma virus Gag p10 domain in nuclear export and virion core morphology. J Virol 81: 10718–10728.
53. Birnaias P (2006) Late budding domains and host proteins in enveloped virus release. Virology 344: 53–63.
54. Freed EO (2002) Viral late domains. J Virol 76: 4679–4687.
55. Yuan R, Campbell S, Bacharach E, Rein A, Goff SP (2000) Infectivity of Moloney murine leukemia virus defective in late assembly events is restored by late assembly domains of other retroviruses. J Virol 74: 7250–7260.
56. Votaweler J, Neumann L, Hahn S, Hahn F, Rauch P, et al. (2011) Highly conserved se1ina residue 40 in HIV-1 p6 regulates capsid processing and virus core assembly. Retrovirology 8: 11.
57. Rasaiyaah J, Tan CP, Fletcher AJ, Price AJ, Blondeau C, et al. (2013) HIV-1 evades innate immune recognition through specific cofactor recruitment. Nature 503: 402–405.
58. Bock M, Bishop KN, Towers G, Stoye JP (2000) Use of a transient assay for studying the genetic determinants of Fv1 restriction. J Virol 74: 7422–7430.
59. Moore HS, Boerger AL, Narayan S, Cunningham JM, Young JA (2000) Retroviral entry mediated by receptor priming and low pH triggering of an envelope glycoprotein. Cell 103: 679–689.
60. Yap MW, Nisole S, Lynch C, Stoye JP (2004) Trim5alpha protein restricts both HIV-1 and murine leukemia virus. Proc Natl Acad Sci U S A 101: 10786–10791.
61. Bruce JW, Bradley KA, Alquist P, Young JA (2005) Isolation of cell lines that show novel, murine leukemia virus-specific blocks to early steps of retroviral replication. J Virol 79: 12969–12978.