Recognition of dileucine-based sorting signals from HIV-1 Nef and LIMP-II by the AP-1 \(\gamma-\sigma^1\) and AP-3 \(\delta-\sigma^3\) hemicomplexes

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The sorting of transmembrane proteins to endosomes and lysosomes is mediated by signals present in the cytosolic tails of the proteins. A subset of these signals conform to the \([DE]XXXL[LI]\) consensus motif and mediate sorting via interactions with heterotetrameric adaptor protein (AP) complexes. However, the identity of the AP subunits that recognize these signals remains controversial. We have used a yeast three-hybrid assay to demonstrate that \([DE]XXXL[LI]-type signals from the human immunodeficiency virus negative factor protein and the lysosomal integral membrane protein II interact with combinations of the \(\gamma\) and \(\sigma^1\) subunits of AP-1 and the \(\delta\) and \(\sigma^3\) subunits of AP-3, but not the analogous combinations of AP-2 and AP-4 subunits. The sequence requirements for these interactions are similar to those for binding to the whole AP complexes in vitro and for function of the signals in vivo. These observations reveal a novel mode of recognition of sorting signals involving the \(\gamma/\delta\) and \(\sigma\) subunits of AP-1 and AP-3.

Introduction

The sorting of many transmembrane proteins at the plasma membrane, the TGN, and the endosomes is mediated by interactions of signals present in the cytosolic domains of the transmembrane proteins with adaptor proteins (APs) that are components of membrane coats (Mellman, 1996; Hirst and Robinson, 1998; Bonifacino and Traub, 2003). Most of these sorting signals consist of short, linear arrays of amino acid residues that fit one of several consensus motifs (Bonifacino and Traub, 2003).

“Tyrosine-based” sorting signals contain a critical tyrosine residue and conform to either NPXY or YXXØ motifs (Bonifacino and Traub, 2003) (N, asparagine; P, proline; X, any amino acid; Y, tyrosine; Ø, bulky hydrophobic amino acid). NPXY signals were originally discovered in the cytosolic tail of the low density lipoprotein receptor (Chen et al., 1990) and are now known to interact with the phosphotyrosine-binding domain of a family of monomeric clathrin adaptors that includes the autosomal recessive form of hypercholesterolemia protein (He et al., 2002; Mishra et al., 2002) and Disabled 2 (Morris and Cooper, 2001). YXXØ signals (Canfield et al., 1991; Collawn et al., 1991) are typically found in another subset of endocytic receptors such as the transferrin receptor, as well as in many transmembrane proteins targeted to endosomes and lysosomes (Bonifacino and Traub, 2003). Recognition of these signals is a function of the \(\mu\) (medium) subunits of the heterotetrameric AP complexes AP-1, AP-2, AP-3, and AP-4 (Ohno et al., 1995, 1998; Owen and Evans, 1998; Ohno et al., 1998; Fig. 1 A). AP-1 and AP-2 have long been known to be components of clathrin coats associated with the TGN/endosomes and the plasma membrane, respectively (Hirst and Robinson, 1998). AP-3 appears to exist as part of both clathrin and nonclathrin coats localized to endosomes, whereas AP-4 is most likely part of a nonclathrin coat associated with the TGN (Robinson and Bonifacino, 2001).
Another major group of sorting signals is referred to as “dileucine-based” and contains critical leucine–leucine or leucine–isoleucine residues. It has recently become apparent that there are at least two distinct types of dileucine-based sorting signals defined by the consensus motifs DXXLL and [DE]XXXLI] (D, aspartate; E, glutamate; L, leucine; I, isoleucine) (Bonifacino and Traub, 2003). DXXL signals occur within the cytosolic tails of intracellular sorting receptors such as the mannose 6-phosphate receptors (MPRs) and mediate sorting between the TGN and endosomes (Johnson and Kornfeld, 1992; Chen et al., 1997). It is now well established that DXXL signals bind to the Vps27, Hrs, and Stam (VHS) domain of the Golgi-localized, γ-carboxy-containing, ARF-binding proteins (GGAs; Nielsen et al., 2001; Puertollano et al., 2001; Takatsu et al., 2001b; Zhu et al., 2001), which is another family of monomeric adaptors associated with clathrin coats at the TGN and endosomes.

[DE]XXXLI] signals (Letourneur and Klausner, 1992; Pond et al., 1995) are present in the cytosolic tails of numerous transmembrane proteins targeted to endosomes, lysosomes, and lysosome-related organelles (Bonifacino and Traub, 2003). A well-characterized example of proteins having [DE]XXXLI] signals (e.g., ERAPLI) is the lysosomal integral membrane protein II (LIMP-II; Barriocanal et al., 1986; Fig. 1 B). The leucine–leucine or leucine–isoleucine pairs are critical elements of the signals, whereas the aspartate or glutamate residue at position –4 from the first leucine is often important but not always essential for function. Interestingly, sequences resembling the canonical [DE]XXXLI] motif are also found in the negative factor (Nef) of all isolates of the human immunodeficiency virus (HIV) and simian immunodeficiency virus (SIV; Fig. 1 B). Unlike LIMP-II, Nef is not a membrane-spanning protein but an N-myristoylated protein tethered to the cytosolic leaflet of membranes (Geyer et al., 2001). A variable EXXXLL sequence is located in a solvent-exposed, unstructured loop near the COOH terminus of HIV-1 Nef (Lee et al., 1996; Grzesieck et al., 1997; Fig. 1 C). This sequence has proven critical for the ability of Nef to down-regulate the CD4 coreceptor from the surface of helper T cells, and has therefore been proposed to function in a manner similar to conventional [DE]XXXLI] signals (Bresnahan et al., 1998; Craig et al., 1998; Greenberg et al., 1998).

[DE]XXXLI] signals do not bind to the VHS domain of the GGAs (Puertollano et al., 2001), this is in part due to the unfavorable position of the acidic residue upstream of the dileucine pair (–4 vs. –3 relative to the first leucine; Misra et al., 2002; Shiba et al., 2002). Instead, some [DE]XXXLI] signals, including those of LIMP-II and HIV-Nef, bind to AP-1, AP-2, and AP-3 complexes in vitro (Dietrich et al., 1997; Bresnahan et al., 1998; Höning et al., 1998; Fujita et al., 1999; Hofmann et al., 1999; Peden et al., 2001; Kongsvik et al., 2002; Rodionov et al., 2002; Janvier et al., 2003). However, the identity of the subunits of the AP complexes that harbor the binding site for [DE]XXXLI] signals remains controversial. Using various in vitro binding and yeast two-hybrid assays, [DE]XXXLI] signals have been shown to interact with the μ subunits of AP-1, AP-2, and/or AP-3 in some studies (Bremnes et al., 1999; Rodionov and Bakke, 1998; Hofmann et al., 1999; Craig et al., 2000), and with the β subunits of AP-1 and AP-2 in other studies (Greenberg et al., 1998; Rapoport et al., 1998; Geyer et al., 2002).

Our lab has made extensive use of the yeast two-hybrid system to characterize interactions of YXXO signals with AP μ subunits (Ohno et al., 1995, 1998) and of DXXL signals with the VHS domains of the GGAs (Puertollano et al., 2001; Kato et al., 2002). However, our attempts to demonstrate interactions of [DE]XXXLI] signals with single subunits of AP complexes using the yeast two-hybrid system have yielded only negative or marginally positive results. We reasoned that our failure to detect strong [DE]XXXLI]–AP subunit interactions using the yeast two-hybrid system could reflect a requirement for more than one subunit. This would be the case if the binding site involved residues on more than one subunit or if the iso-
whereas the strains coexpressing Nef with either described in D. Values are the mean by optical density at 600 nm after 48 h in culture. The value 1.0 on

![Figure 2](image.png)

**Figure 2. Three-hybrid analysis of the interaction of HIV-1 Nef (NLA4-3 variant) with different AP subunits.** (A) GAL4BD-Nef and the μ subunits were expressed from pBridge, whereas the β subunits and γ1 were expressed as fusions with GAL4AD from pGADT7. (B) GAL4BD-Nef and the σ subunits were expressed from pBridge, whereas the β1, β2, β3A, β4, γ1, αC, δ, and ε were expressed as fusions with GAL4AD from pGADT7. (C) Immunoblot analysis (using anti-HA) of AP subunits in yeast cells cotransformed with γ1–1A, αC–σ2, δ–σ3A, or ε–σ4. The subunits were expressed as in B, and all subunits were tagged with the HA epitope. (D) GAL4BD-Nef and the σ subunits were expressed from pBridge, whereas γ1, αC, δ, and ε were expressed as fusions with GAL4AD from pGADT7. In A, B, and D, interactions were evidenced by growth on agar plates made with medium without histidine (−His) or by expression of β-galactosidase (β-Gal) activity. (E) Time course of growth of yeast strains coexpressing Nef with either γ1–σ1A, γ1–σ1A, or δ–σ3A (δ–σ3A) was measured by optical density at 600 nm (OD<sub>600</sub>). (F) Inhibition by 3AT of growth of yeast strains coexpressing Nef with either γ1–σ1A, γ1–σ1A, or δ–σ3A (δ–σ3A). Growth was measured by optical density at 600 nm after 48 h in culture. The value 1.0 on the y axis corresponds to the growth of each strain in the absence of 3AT. The expression of Nef and AP subunits from plasmids was as described in D. Values are the mean ± SD of triplicate determinations.

lated subunit containing the binding site did not fold properly in the absence of another subunit. To test this hypothesis, we resorted to a yeast three-hybrid system in which [DE][XX][L][I] signals from HIV-1 Nef and LIMP-II were examined for interactions with combinations of two subunits from all four AP complexes.

**Results**

**Yeast three-hybrid analysis of the interactions of HIV-1 Nef with combinations of two AP subunits**

The yeast three-hybrid assay consisted of coexpressing the following: (a) the GAL4 DNA binding domain (GAL4BD) fused to HIV-1 Nef (NLA4-3 variant, herein referred to as Nef unless otherwise specified; cloned in multiple cloning site 1 of pBridge), (b) one of the AP subunits (cloned in multiple cloning site 2 of pBridge), and (c) the GAL4 transcription activation domain (GAL4AD) fused to another AP subunit (cloned in pGAD424 or pGADT7; Fig. 1D). The NLA4-3 variant of Nef contains the dileucine-based signal ENTSLL (Fig. 1B), which is required for induction of CD4 down-regulation in vivo (Bresnahan et al., 1998; Craig et al., 1998) and for interactions with the AP-1 and AP-3 complexes in vitro (Bresnahan et al., 1998; Janvier et al., 2003). Interactions in the three-hybrid system were detected by growth in histidine-deficient medium or expression of β-galactosidase activity. Using this assay, we were unable to detect interactions of Nef with different combinations of μ with either μ (Fig. 2 A) or σ subunits (Fig. 2 B) from any of the four complexes. However, these assays revealed relatively strong interactions of Nef with γ1–σ1A and δ–σ3A, but not αC–σ2 and ε–σ4 (Fig. 2, B and D), pairs of subunits that are known or expected to interact with one another in the intact complexes (Page and Robinson, 1995; Takatsu et al., 2001a; Collins et al., 2002; Fig. 1 A). All of the homologous subunits were expressed at comparable levels (Fig. 2 C), indicating that the failure of the αC–σ2- and ε–σ4-transformed strains to grow was not due to the absence of these subunits. All other combinations of γ1/αC/β/e with σ1–4 tested negative in this assay (Fig. 2 D). Interactions of Nef with γ1–σ1A appeared stronger than those with δ–σ3A, as judged from the following: (a) the expression of β-galactosidase activity only by the γ1–σ1A-expressing strain (Fig. 2 D), (b) the faster growth in histidine-deficient liquid medium of the γ1–σ1A-expressing strain (Fig. 2 E), and (c) the higher concentrations of 3-aminotriazole (3AT) needed to inhibit growth of the γ1–σ1A-expressing strain (Fig. 2 F). These observations are entirely consistent with previous reports of interactions of HIV-1 Nef with the intact AP-1 and AP-3, but not AP-2, in vitro (Bresnahan et al., 1998; Janvier et al., 2003). Thus, the ability of Nef to interact with AP-1 and AP-3 appears to be a function of the γ1–σ1A and δ–σ3A hemicomplexes.

**Interactions of Nef from different HIV-1 and SIV variants with γ1–σ1A and δ–σ3A**

Both human and simian immunodeficiency viruses are highly variable genetically, owing to their error-prone reverse transcriptase (Roberts et al., 1988). However, the critical elements of the [DE][XX][L][I]-type signal in Nef are conserved in all isolates of HIV-1, HIV-2, and SIV (Fig. 1 B; see supplemental material in Greenberg et al., 1998). The acidic residue is invariant, and the two bulky hydrophobic positions are occupied by leucine–leucine, leucine–valine, or
leucine–methionine pairs in all Nefs. In contrast, the sequences flanking the [DE]XXXL[LI] signal and the X residues are variable (Fig. 1 B). To determine the generality of Nef–AP subunit interactions, we compared the binding of Nef gene products from four isolates of HIV-1 (NLA4-3, 248, ELI, and DH123) and one isolate of SIV (RQ). All five Nefs bound to γ1–σ1A, whereas only the NLA4-3 and 248 Nefs bound to δ–σ3A (Fig. 3). Thus, binding to γ1–σ1A appears to be the most general feature of all Nefs. Another general property revealed by these assays is the inability of all Nefs tested to bind to αC–σ2 and e–σ4 (Fig. 3).

Determinants of interactions of Nef with γ1–σ1A and δ–σ3A

The γ1 and δ subunits are organized into “trunk,” “hinge,” and “ear” domains (Fig. 1 A; Robinson and Bonifacino, 2001). Yeast three-hybrid analyses showed that interactions with Nef were mediated by the trunk, but not the hinge–ear domains, of γ1 and δ in combination with σ1A and σ3A, respectively (Fig. 4 A). To determine whether the interactions of Nef with AP subunits were mediated by the ENTSLL sequence (Fig. 4 B), and to delineate the sequence requirements for these interactions, we performed an alanine-scan mutagenesis. We observed that both leucine residues (L164 and L165) of the ENTSLL sequence were strictly required for interactions with γ1–σ1A and δ–σ3A (Fig. 4 C, asterisks). Mutation of the acidic residue (E160) of the signal diminished but did not completely abrogate the interaction with γ1–σ1A and abolished the interaction with δ–σ3A, as shown in both plate (Fig. 4 C, open triangle) and liquid medium growth assays (Fig. 4 D). These observations are in agreement with previous GST pull-downs assays, which showed that L164 and L165 are critical for binding to AP-1 and AP-3, and that E160 is only partially important for AP-1 but absolutely critical for AP-3 (Bresnahan et al., 1998; Janvier et al., 2003). Mutation of eleven residues other than E160, L164, and L165 had little or no effect on the interactions of Nef with both γ1–σ1A and δ–σ3A (Fig. 4 C).

CD4 down-regulation induced by Nef mutants

To correlate the sequence requirements for Nef–AP subunit interactions and Nef function, we examined the ability of
Down-regulation of CD4 by Nef or Nef mutants.

(A) Analysis by immunofluorescence microscopy. HeLa cells were cotransfected with pCMV-CD4, pEGFP encoding GFP-histone H2B, and pCI-neo encoding either wild-type Nef (NLA4-3 variant) or Nef mutants. At 24 h after transfection, cells were fixed, permeabilized, and immunostained for CD4 (red). Cotransfected cells were identified by the expression of GFP-histone H2B in the nucleus (green). Images corresponding to selected Nef constructs are shown. Bar, 25 μm. (B) Analysis by FACS®. HeLa cells were cotransfected with pCMV-CD4, pCMV-CD8, and pCI-neo encoding either wild-type Nef (NLA4-3 variant) or Nef mutants. At 24 h after transfection, cells were immunostained with allophycocyanin-conjugated anti-CD4 and phycoerythrin-conjugated anti-CD8 antibodies. The percentage of CD8+ cells that also expressed CD4 on the surface was quantified by FACS® analysis, as described in Materials and methods. The results obtained in different determinations for each Nef construct are indicated by the individual dots. Bars represent the mean values for each Nef construct.

Interactions of the [DE]XXXL[LI] signal from LIMP-II with AP subunits

The results obtained with Nef beg the question of whether other [DE]XXXL[LI] signals also bind to the same combinations of AP subunits. To address this question, we examined the interaction of the four AP complexes with the cytosolic tail of LIMP-II, which has a well-characterized signal, ERAPLI (Ogata and Fukuda, 1994; Sandoval et al., 1994; Pond et al., 1995). We initially performed GST pull-downs using as sources of adaptors either HeLa cell cytosol or a mixture of AP complexes purified from bovine brain clathrin-coated vesicles. This latter preparation contained large quantities of AP-1 and AP-2, moderate quantities of AP-3, and traces of AP-4. Binding to the immobilized GST constructs was detected by immunoblotting with antibodies to specific AP subunits. We observed that GST–LIMP-II, like GST–Nef (Bresnahan et al., 1998; Janvier et al., 2003), bound to both AP-1 and AP-3, but not AP-2 and AP-4 (Fig. 6 A). The ability of the LIMP-II tail to bind AP-1 in this pull-down assay is at variance with previous findings made using surface plasmon resonance spectroscopy, which revealed binding only to AP-3 (Höning et al., 1998; Rodionov et al., 2002). As specificity controls, we showed that neither AP complex bound detectably to GST or to GST appended with the cation-independent MPR (CI-MPR) tail, which contains a DXXLL-type signal (Fig. 6 A). In addition, we found that mutations of the glutamate, leucine, or isoleucine residues to alanine residues completely abolished the interactions with both AP-1 and AP-3, thus, demonstrating that these interactions are mediated by the ERAPLI signal.

In agreement with the GST pull-down assays, yeast three-hybrid analyses showed that the cytosolic tail of LIMP-II interacted with γ1–σ1A, δ–σ3A, and δ–σ3B (σ3A and σ3B are two isoforms of σ3), but not with αC–σ2, ε–σ4, μ3A,
The use of the yeast three-hybrid system has allowed us to detect relatively strong interactions of the [DE]XXXL[LI] signals from HIV-1 Nef and LIMP-II with specific combinations of two subunits from the AP-1 and AP-3 complexes. Strikingly, the subunits that interact with these signals when expressed together, γ̃1–σ̃1 and δ–σ̃3, are precisely the two subunits from each complex that had not been previously implicated in the recognition of [DE]XXXL[LI] signals. A previous yeast two-hybrid study had indicated a role for the μ subunits of AP-1, AP-2, and AP-3 in the recognition of the EXXXLL sequence from HIV-1 Nef (Craig et al., 2000). Although we were able to confirm the interaction of Nef with μ1A, this was much weaker than the interaction of Nef with γ̃1–σ̃1 and δ–σ̃3 (unpublished data). Other studies involving competition of labeling with photoactivatable peptides (Greenberg et al., 1998), yeast two-hybrid assays (Geyer et al., 2002), and binding of in vitro translated proteins to GST fusions (Geyer et al., 2002) had indicated that the EXXXLL sequence from Nef interacts with the β1 and β2 subunits of AP-1 and AP-2, respectively. These studies are representative of others in which [DE]XXXL[LI] signals from several proteins were shown to interact with either μ or β subunits (Bremsnes et al., 1998; Rapoport et al., 1998; Rodionov and Bakke, 1998; Hofmann et al., 1999).

In view of the aforementioned discrepancies, what gives us confidence that the interactions with γ̃1–σ̃1 and δ–σ̃3 reported herein reflect the actual mode of recognition of [DE]XXXL[LI] signals by AP complexes? We think that the weight of our evidence comes from the extensive correlational analyses performed in our study. The fact that the [DE]XXXL[LI] signals from both Nef and LIMP-II interact with γ̃1–σ̃1 and δ–σ̃3 but not αC–σ̃2 and e–σ̃4 correlates well with the interaction of those signals with the intact AP-1 and AP-3, but not AP-2, in GST pull-down assays (Bresnahan et al., 1998; Janvier et al., 2003; this paper). This correlation holds true even in the one instance in which the

or the VHS domain of GGA1 (Fig. 6 B). In contrast, the cytosolic tails of the cation-dependent MPR (CD-MPR) and of the CI-MPR, which contain DXXLL-type signals, bound specifically to the VHS domain of GGA1 (Fig. 6 B; Gough et al., 1999). This experiment thus shows the specific recognition of three different types of sorting signals by their cognate recognition proteins.

To corroborate the specificity of the three-hybrid interactions, we performed an alanine-scan mutagenesis of the LIMP-II cytosolic tail. Of 12 residues that were mutated, only three were essential for interactions with γ̃1–σ̃1A, δ–σ̃3A, and δ–σ̃3B, the leucine-isoleucine pair and the glutamate residue at position −4 (Fig. 6 C). Mutation of other residues had no effect on the interactions. Together, these experiments demonstrate that the LIMP-II cytosolic tail binds to the AP-1 and AP-3 complexes through γ̃1–σ̃1A and δ–σ̃3 (A or B isoforms). Binding to both complexes is mediated by the ERAPLI signal, with only the glutamate, leucine, and isoleucine residues being absolutely required for interactions. These sequence requirements are in agreement with the requirements for function of the ERAPLI signal in sorting to lysosomes (Ogata and Fukuda, 1994; Sandoval et al., 1994; Pond et al., 1995), which provides strong support for the physiological significance of these interactions.

**Discussion**

**Identity of the AP subunits that bind [DE]XXXL[LI] signals**

The use of the yeast three-hybrid system has allowed us to detect relatively strong interactions of the [DE]XXXL[LI] signals from HIV-1 Nef and LIMP-II with specific combinations of two subunits from the AP-1 and AP-3 complexes. Strikingly, the subunits that interact with these signals when expressed together, γ̃1–σ̃1 and δ–σ̃3, are precisely the two subunits from each complex that had not been previously implicated in the recognition of [DE]XXXL[LI] signals. A previous yeast two-hybrid study had indicated a role for the μ subunits of AP-1, AP-2, and AP-3 in the recognition of the EXXXLL sequence from HIV-1 Nef (Craig et al., 2000). Although we were able to confirm the interaction of Nef with μ1A, this was much weaker than the interaction of Nef with γ̃1–σ̃1 and δ–σ̃3 (unpublished data). Other studies involving competition of labeling with photoactivatable peptides (Greenberg et al., 1998), yeast two-hybrid assays (Geyer et al., 2002), and binding of in vitro translated proteins to GST fusions (Geyer et al., 2002) had indicated that the EXXXLL sequence from Nef interacts with the β1 and β2 subunits of AP-1 and AP-2, respectively. These studies are representative of others in which [DE]XXXL[LI] signals from several proteins were shown to interact with either μ or β subunits (Bremsnes et al., 1998; Rapoport et al., 1998; Rodionov and Bakke, 1998; Hofmann et al., 1999).

In view of the aforementioned discrepancies, what gives us confidence that the interactions with γ̃1–σ̃1 and δ–σ̃3 reported herein reflect the actual mode of recognition of [DE]XXXL[LI] signals by AP complexes? We think that the weight of our evidence comes from the extensive correlational analyses performed in our study. The fact that the [DE]XXXL[LI] signals from both Nef and LIMP-II interact with γ̃1–σ̃1 and δ–σ̃3 but not αC–σ̃2 and e–σ̃4 correlates well with the interaction of those signals with the intact AP-1 and AP-3, but not AP-2, in GST pull-down assays (Bresnahan et al., 1998; Janvier et al., 2003; this paper). This correlation holds true even in the one instance in which the

Figure 6. Interactions of the dileucine-based sorting signal from LIMP-II with AP complexes. (A) GST pull-down assays. GST fused to the cytosolic tail of the CI-MPR, LIMP-II, or mutants of this latter construct were incubated with either a cytosolic extract prepared from HeLa cells or a mixture of AP complexes prepared from bovine brain clathrin-coated vesicles. The LIMP-II tail mutants contain the amino acid substitutions indicated in the figure relative to the LIMP-II sequence shown in Fig. 1 B. Bound AP complexes were revealed by immunoblot analysis with antibodies specific for each complex. Some of the AP subunits (most notably α3 and α in bovine brain) appeared as doublets. (B) Three-hybrid analysis of the interaction of the LIMP-II, CD-MPR, CI-MPR, and LAMP-2a cytosolic tails fused to GAL4BD (pBridge) with γ̃1, α, δ, and e (pBridge) and different ε subunits; μ3A and the VHS domain of GGA1 fused to GAL4AD (pGAD424). Interactions were detected by growth in the absence of histidine (−His). (C) Alanine-scan mutagenesis analysis of the interaction of LIMP-II cytosolic tail mutants with γ̃1–σ̃1A (γ̃1 + σ̃1A), δ–σ̃3A (δ + σ̃3A), or δ–σ̃3B (δ + σ̃3B) using the yeast-three-hybrid system. Mutated residues are underlined. Plasmid constructs were as specified in B. The SV40 large T antigen (TAG) and p53 were used as controls. Interactions were detected by growth in the absence of histidine (−His).
results of our GST pull-down assays are at variance with previously published results, as is the case for LIMP-II, which was previously reported not to bind AP-1 (Höning et al., 1998). A second type of correlation concerns the residues of the signals required for interactions in the yeast three-hybrid system and in GST pull-downs. Again, there is excellent correspondence between these two sets of data: the key glutamate and leucine–leucine or leucine–isoleucine residues are required for interactions in both assays. Even the fact that the key glutamate is less important for Nef than for LIMP-II interactions is reflected quantitatively in both assays (compare the results in Fig. 4 C in this paper with those in Fig. 7 B of Janvier et al. [2003], and in Fig. 6, A and C, in this paper). Thus, binding to γ1–σ1 and δ–σ3 recapitulates the binding to the intact complexes, implying that both are manifestations of the same molecular recognition event.

Structural implications for the recognition of [DE]XXXL[LI] signals

Although two subunits are necessary to detect binding of [DE]XXXL[LI] signals in the yeast three-hybrid assays, at present, we cannot ascertain whether the binding site involves residues on the surface of both subunits or is located exclusively on one subunit, with the other subunit being required for proper folding or stability of the hemicomplex. The structures of the γ1–σ1 and δ–σ3 hemicomplexes can be modeled after that of the α–σ2 hemicomplex in the context of the AP-2 core, which was recently solved by X-ray crystallography (Collins et al., 2002). The NH2-terminal trunk domain of each large subunit (γ1 or δ) would be expected to consist of a long α-helical superhelix with an “elbow-like” bend near the middle of the domain. The small chain (σ1 or σ3) would nestle at the bend, forming a tight heterodimer (Collins et al., 2002). The trunk domains of γ1 or δ seem particularly suited to harbor peptide-binding sites because of the many grooves predicted to occur on the surface of their superhelical folds. Indeed, the spatial arrangement of the NH2-terminal α-helices of the large subunits (Collins et al., 2002) resembles that of the eight α-helices that make up the VHS domain of the GGAs (Misra et al., 2002; Shibata et al., 2002), as previously proposed (Geyer et al., 2002). A groove between helices 6 and 8 of the GGA VHS domain constitutes the binding site for DXXLL signals (Misra et al., 2002; Shibata et al., 2002), and it is possible that an analogous groove might be present in the trunk domain of the large AP subunits. We would expect this binding site to occur in a membrane-proximal region of the γ1 or δ trunks because some [DE]XXXL[LI] signals (e.g., that of LIMP-II) are close to the transmembrane domain. Although it might be possible to map the peptide-binding site on the γ1–σ1 and δ–σ3 hemicomplexes by mutational analyses, its definitive identification will require X-ray crystallographic analyses.

Our observations demonstrate that [DE]XXXL[LI] and YXXØ signals (Ohno et al., 1995, 1998) bind to different subunits of the AP complexes (γ1–σ1 and δ–σ3 vs. μ subunits). This conclusion agrees with previous observations from in vivo overexpression/saturation experiments, which showed that [DE]XXXL[LI] signals compete with other [DE]XXXL[LI] signals but not with YXXØ signals for engagement of the protein trafficking machinery; the same is the case for YXXØ signals (Marks et al., 1996; Craig et al., 1998). The binding of both types of signals to the AP complexes may explain why these signals have similar functions in sorting to endosomes, lysosomes, and lysosome-related organelles. In contrast, the spatial separation of the two binding sites may afford differential regulation of signal recognition.

Possible role of AP-1 in Nef-induced CD4 down-regulation

We have shown that the ability of Nef to down-regulate CD4 correlates best with binding to AP-1, pointing to a likely role for this complex in CD4 down-regulation. AP-3 could also play a role, at least for some Nef variants. The localization of these two AP complexes to the TGN and endosomes makes it likely that the Nef-induced down-regulation of CD4 results mainly from impairment of its transport from intracellular compartments to the plasma membrane. The fact that AP-2 does not interact with Nef suggests that this adaptor is not a direct target for regulation by HIV-1. Rather, another protein such as the regulatory V1H1 subunit of the vacuolar ATPase may be responsible for the Nef-induced removal of CD4 from the cell surface (Lu et al., 1998). Alternatively, CD4 may be internalized constitutively in HeLa cells or upon dissociation of the tyrosine kinase lck in T cells (Pelchen-Matthews et al., 1991, 1998; Gratton et al., 1996). This internalization may be mediated by direct interaction of the phosphorylated CD4 tail with AP-2, which occurs even in the absence of Nef (Pitcher et al., 1999). Whenever the mechanism of endocytosis, the internalized CD4 could be captured by AP-1 and AP-3 in the presence of Nef, preventing its return to the cell surface and possibly routing it to lysosomes. An alternative scenario would involve interaction of newly-synthesized CD4 with Nef and AP-1 or AP-3, resulting in reduced transport to the cell surface and enhanced delivery to the endosomal–lysosomal system. Both scenarios would be in line with the observation that intracellular retention plays a critical role in the down-regulation of CD4 by Nef (Mangasarian et al., 1997).

The molecular mechanism of down-regulation likely involves the formation of a tripartite complex between Nef, CD4, and AP-1. The cytosolic tail of CD4 contains a determinant encompassing a dileucine pair at positions 413 and 414 that does not conform to the [DE]XXXL[LI] consensus (Pitcher et al., 1999). This determinant is required for Nef-induced down-regulation (Aiken et al., 1994; Hua et al., 1997; Pitcher et al., 1999) via interactions with a hydrophobic patch on Nef that includes the side chains of W57, L58, and L110 (Grzesiek et al., 1996). The [DE]XXXL[LI] signal in Nef (residues 160–165) in turn binds to the AP-1 γ1–σ1 and AP-3 δ–σ3 subunits. Thus, Nef can be thought of as a “connector” (Mangasarian et al., 1997) that bridges the tail of CD4 to AP-1. Alternatively, the interaction of Nef with AP-1 and AP-3 could modify its properties, enabling it to bind directly to the CD4 tail. In this regard, expression of Nef has been shown to stabilize the association of AP-1 and AP-3 with membranes by an Arf-independent mechanism (Janvier et al., 2003). This stabilization could enhance an otherwise weak binding of CD4 to AP-1 and AP-3.

Conclusion

In summary, our studies have uncovered a novel mode of recognition of [DE]XXXL[LI] signals from HIV-1 Nef and
LIMP-II involving two subunits of the AP-1 and AP-3 complexes, γ1–σ1 and δ–σ3, respectively. The assays developed to study these interactions are robust and should be applicable to the analysis of the interaction of other [DE]XXX[L/I] signals with AP complexes. In particular, it will be of interest to determine whether other [DE]XXX[L/I] signals are recognized by α–σ2 or ε–σ4, which tested negative in our assays despite having structures similar to γ1–σ1 and δ–σ3. Finally, our studies support a prominent role for the AP-1 complex in the Nef-induced down-regulation of CD4, implying that this down-regulation occurs mainly from endosomes or the TGN.

Materials and methods

Recombinant DNA constructs

GAL4AD fusion constructs were made by PCR amplification and in-frame cloning into yeast two-hybrid vectors (CLONTECH Laboratories, Inc.; Fig. 1 D). γ1, α3, β, and the hinge–ear fragments of γ1 and δ (BamHI–XhoI) and β1 and β3A (SmaI–SalI) were cloned into pGAD77. μ3A and σ1A (EcoRI–SalI), and α2, α3A, α3B, and σ4 (BamHI–XhoI) were cloned into pGAD424. ε and β4 in pGAD77 (Boehm et al., 2001), as well as the CC1A VHS domain and the SV40 T-antigen (Tag) in pGAD424 (Kato et al., 2002), have been described previously. β2 in pGAD424 was a gift of M.S. Robinson (University of Cambridge, Cambridge, UK).

GAL4BD fusion constructs were also made by PCR amplification and in-frame cloning into multiple cloning site 1 of pBRII (CLONTECH Laboratories, Inc.; Fig. 1 D, MCSI). Nef from different strains of HIV-1 or SIV, the cytosolic tails of LIMP-II (20 COOH-terminal residues), CI-MPR (114 COOH-terminal residues), CD-MPR (41 COOH-terminal residues), and p53 were cloned into the EcoRI–SalI sites of pBridge. The LAMP-2a cytosolic tail was cloned into the EcoRI–PstI sites of pBRII. Mutants of GAL4BD-Nef and GAL4BD-LIMP-II were obtained by PCR-directed mutagenesis.

For three-hybrid analyses, the AP subunit cDNAs were amplified by PCR and subcloned into the multiple cloning site 2 (Fig. 1 D, MCS2) of the recombinant pBridge constructs expressing the GAL4BD fusion proteins as described in the previous paragraph. The σ and μ subunits of the AP complexes were cloned into the NotI–BglII sites of the pBridge-Nef constructs. γ1, α3, and δ were cloned into the NotI site of pBridge constructs expressing the GAL4BD-LIMP-II, γ1–MI-PR, CD-MPR, and p53 fusion proteins described in the previous paragraph. ε was cloned into the BglII site of the same pBridge recombinant constructs. For expression in mammalian cells, wild-type and mutant Nef cDNA were PCR amplified and cloned into the EcoRI–SalI sites of the pcI-neo vector (Promega). The GFP-tagged histone H2B construct was made as described previously (Dey et al., 2000). The pCMV-CD4 and pCMV-CD8 expression vectors were obtained from the AIDS Reagent Program, National Institute of Allergy and Infectious Diseases.

The GST–LIMP-II and GST–CI-MPR constructs were obtained by in-frame cloning of sequences encoding the 20 COOH-terminal residues of LIMP-II and the 114 COOH-terminal residues of the CI-MPR into the EcoRI–SalI sites of pGEX-SX-1 (Amersham Biosciences). The mutant pGEX–LIMP-II constructs were obtained by PCR-directed mutagenesis. The GST fusion proteins were produced by transformation of the Escherichia coli strain BL21 (DE3) pLYS5 (Novagen) and purified according to the manufacturer’s instructions.

Antibodies

The following mouse mAbs were used: 100/3 to γ1-adapter and 100/2 to α-adapter (Sigma-Aldrich), anti-ε-adapter (Transduction Laboratories), S3.5 to human CD4, anti-CD4, allophycocyanin-conjugated anti-CD4, and phycoerythrin-conjugated anti-CD8 antibodies (Caltag Laboratories), and HA.11 to the HA tag (Covance). The following polyclonal antibodies were also used: rabbit antibody to α3 (Dell’Angelica et al., 1997), Alexa 594–conjugated anti-mouse IgG (Molecular Probes), and HRP-conjugated anti-mouse and anti-rabbit IgG (Amersham Biosciences).

Yeast culture, transformation, and two- and three-hybrid assays

The Saccharomyces cerevisiae strain H7Fc (Clontech), was maintained on dropout agar plates lacking methionine. Transformation was performed by the lithium acetate procedure as described in the instructions for the MATCHMAKER two-hybrid kit (CLONTECH Laboratories, Inc.). H7Fc transforms were selected by growth on plates lacking leucine, tryptophan, and methionine. For colony growth assays, H7Fc transforms were dotted on plates lacking leucine, tryptophan, methionine, and histidine, and allowed to grow at 30°C for 3–5 d. Quantitative growth assays were performed as described previously (Aguilar et al., 1997). Filter-based β-galactosidase assays were performed according to the instructions for the MATCHMAKER two-hybrid kit (CLONTECH Laboratories, Inc.).

Preparation of yeast lysates

Yeast were grown in synthetic medium at 30°C to an optical density of 0.6. 3 OD units of yeast cells were resuspended in 200 μl of ice-cold 10% trichloracetic acid, and then transferred to Eppendorf tubes containing 200 μl of acid-washed glass beads. The cells were broken by vigorous vortexing for 1 min, followed by chilling on ice for another 1 min. This cycle was repeated 10 times. Proteins contained in the supernatant were precipitated by centrifugation at top speed, followed by a wash in ice-cold acetone. The protein extract was resuspended in 100 μl of Laemmli loading buffer, and then analyzed by immunoblotting using an anti-HA antibody, followed by chemiluminescent detection (Amersham Biosciences).

Transfection

HeLa cells (American Type Culture Collection) were transfected using the lipofectamine reagent (Invitrogen) according to the manufacturer’s instructions. For immunofluorescent staining, cells were cotransfected with mammalian expression vectors encoding CD4 and GFP-histone H2B along with expression vectors encoding Nef or Nef mutants. For FACs® analysis, the GFP-histone H2B construct was substituted by a vector encoding CD8.

Immunofluorescence microscopy

24 h after transfection, HeLa cells grown on glass coverslips were fixed in 4% PFA in PBS, quenched for 10 min with 50 mM NH4Cl in PBS, and permeabilized for 10 min with 0.1% (v/v) Triton X-100 in PBS. After permeabilization, the cells were blocked for 30 min with 10% (v/v) normal goat serum in PBS and incubated for 1 h at RT with the primary antibody, washed with PBS, and incubated for 1 h with the secondary antibody. The coverslips were washed and mounted on slides. Images were acquired on a confocal microscope (model LSM 410; Carl Zeiss Microlmaging, Inc.).

Flow cytometry

Cotransfections of HeLa cells with plasmids encoding CD4 and CD8 were optimized ahead of time to obtain equivalent mean fluorescence value ratios, which were typically achieved with 0.5 μg CD8 and 0.8 μg CD4 plasmids. For determination of cell surface antibody binding, 106 cells transfected with CD4, CD8, and Nef (or Nef mutants) were collected by centrifugation and washed with PBS. Cells were incubated for 10 min at RT with allophycocyanin-conjugated anti-CD4 and phycoerythrin-conjugated anti-CD8 antibodies. The cells were washed three times with ice-cold PBS containing BSA or BSA, and fixed in PBS containing 4% PFA. Flow cytometric data were acquired by using a two-laser, four-color FACSCalibur flow cytometer (Becton Dickinson). Data analysis was done using CELLQuest v3.3 (Becton Dickinson) and FlowJo v3.3.2 software (Treestar Corp.).

GST pull-downs

GST pull-downs were performed using either a cysteolic extract of HeLa cells prepared in 25 mM Hepes, pH 7.4, 150 mM NaCl, 1 mM EDTA, 0.5 mM MgCl2, and 0.5% (v/v) Triton X-100 (lysis buffer) or a purified fraction of AP complexes prepared from bovine brain clathrin-coated vesicles as described previously (Ijang et al., 2000; provided by L. Greene, National Heart, Lung, and Blood Institute, NH). Purified GST, GST–CI-MPR, or GST–LIMP-II (wild-type or mutated) proteins were immobilized on glutathione-agarose beads (Amersham Biosciences), and then incubated overnight with 1 ml HeLa cell lysate (corresponding to 106 cells) or for 5 h with 0.2 ml of the purified AP complexes (in extraction buffer containing 0.5 M Tris-HCl, pH 7.0, 2 mM DTT, and 1 mM EDTA) diluted to 1 ml in lysis buffer. The beads were washed three times with lysis buffer and once with wash buffer without Triton X-100. Bound AP complexes were resolved by SDS-PAGE and revealed by immunoblotting using antibodies to γ1, α3, or ε-adapter followed by chemiluminescent detection (Amersham Biosciences).

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