Signal peptides (SP) are key determinants for targeting and membrane insertion of secretory and membrane proteins (reviewed in reference 25). They can be removed co- or posttranslationally by the cellular membrane-bound signal peptidase or may, if not cleaved, serve as membrane anchors for proteins with distinct membrane orientations. In general, SP are composed of three domains, of which a central 6- to 15-amino-acid (aa)-long hydrophobic domain (h-domain) is the most essential. An N-terminal polar domain (n-domain) usually of net positive charge shows high variability in overall length, ranging from 15 to more than 50 aa. The composition and structure of the n-domain influences protein orientation in the membrane. The polar C-terminal domain (c-domain) often contains helix-breaking as well as small uncharged residues in positions -3 and -1 which determine the site of SP cleavage. In most cases, SP cleavage is thought to occur cotranslationally; however, for some proteins, e.g., the human immunodeficiency virus type 1 (HIV-1) envelope glycoprotein gp160, SP cleavage occurs inefficiently and very late after translocation (21). A basic amino acid stretch in the n-domain of gp160 is responsible for this phenomenon and believed to influence folding and exit of HIV-1 Env from the endoplasmic reticulum (ER) (21). Recent studies revealed that SP bear specific information accounting for distinct functions in targeting and membrane insertion or even for defined metabolic pathways after their cleavage from the parent protein (reviewed in reference 25). The HIV-1 SP_env, for example, is further processed by the signal peptidase, leading to the release of an SP fragment into the cytosol, where it binds to calmodulin (26). The function of this process in viral replication is not known.

Foamy viruses (FV), as studied with the prototype member human foamy virus (HFV), follow a replication cycle which is characterized by several unique features setting them apart from the family of retroviruses. These are the independent expression of the Pol protein from a spliced mRNA, efficient reverse transcription prior to particle release, and intracellular retrotransposition (14, 24). The essential functions of retroviral glycoproteins are binding of the viral particle to cellular receptors and subsequent fusion of viral and cellular lipid membranes to release the viral capsid into the cytoplasm (reviewed in reference 19). The FV Env protein is unique among all retroviral glycoproteins since its expression is essential for the FV particle budding and release process (3, 7). Similar to B- or D-type retroviruses, FV particles assemble in the cytoplasm of infected cells. However, unlike the case for all other retroviruses, FV capsids do not bud across cellular membranes in the absence of FV Env, and heterologous viral glycoproteins cannot complement FV Env to enable particle release (3, 7, 28). The particle-associated FV Env glycoprotein is synthesized as a 130-kDa precursor. Analogous to other retroviral Env proteins, FV Env is cleaved during its transport to the cell surface by a cellular protease, yielding a 80- to 90-kDa surface (SU) and a 48-kDa transmembrane (TM) subunit (11, 23). However, the cytoplasmic domain (CyD) of the TM subunit contains an ER retrieval signal, leading to accumulation of FV Env in the ER when other FV structural proteins are absent (10, 11, 29). Thus, the export of FV capsids requires the coexpression of cognate Env protein, and vice versa, the surface localization of Env depends on the presence of cognate capsids. This implies inherent specific interactions between the two partners.
We have shown previously that the membrane-spanning domain (MSD) but not the CyD of Env TM is essential for the particle release process (28, 29). Since the C terminus of Env does not appear to mediate the interaction with Gag, we investigated whether the N-terminal SP sequence, besides targeting the Env protein to the secretory pathway, might have additional functions in the particle release process.

MATERIALS AND METHODS

Expression constructs. The eukaryotic expression constructs for various FV envelopes depicted in Fig. 4 and 5 are based on a previously described plasmid, pchFwE-wt (see Fig. 2A), which expresses only gp130 due to inactivation of the internal splice donor and splice acceptor pair within the FV Env coding region (EM02 mutation) (23, 28). All deletion and point mutants within the N region of the SP (pchFwEnv EM41 to 44 and EM66 to 76, described below) were generated by recombinant PCR techniques (18) with pchFwE-wt as the template and primers introducing the desired codon changes or deletions. The amplimers were cloned into the NheI/EcoRI sites of plasmid pDNA3.1 and xbaI KpnI fragments together with an KpnI/EcoRI fragment of pchFwE-wt. For the N-glycosylation and signal peptide complex (SPC) cleavage site mutants (EM58 and EM78 to EM84), a 3'KpnI/2I fragment of pcHFE-wt was replaced by the corresponding mutated PCR fragments. All PCR-derived inserts were completely sequenced to verify the presence of only the desired mutations.

The resultant constructs contain the mutations and deletions in parentheses: N-terminal deletions, EM42 (Δ2–4, M16I, M72I), EM43 (Δ2–16, M72I), EM70 (Δ2–25), EM71 (Δ2–40), EM72 (Δ2–50), EM73 (Δ2–66), and EM44 (Δ2–72); internal deletions, EM66 (Δ16–25), EM67 (Δ16–40), EM68 (Δ16–50), EM69 (Δ16–66), EM74 (Δ41–50), EM75 (Δ41–66), and EM76 (Δ45–66); C-terminal deletion, EM50 (Δ47–989); and point mutations EM41 (M4I, M16I, M72I), EM52 (N25Q), EM77 (N109Q), EM78 (N141Q), EM79 (N25Q, N109Q), EM80 (N25Q, N141Q), EM81 (N25Q, N109Q, N141Q), EM82 (N25Q, N109Q, N141Q), EM83 (C69R), EM84 (G19R).

The replication-deficient pMH62 vector (see Fig. 2A) was described previously (28). It expresses the FV Gag/Pol proteins and contains an internal splice focus forming virus U3 promoter-directed enhanced green fluorescent protein (EGFP) marker gene expression cassette.

The parental human cytomegalovirus (CMV) immediate-early promoter-driven infectious proviral clone pCH2 (see Fig. 3A) has been described previously (27). The pCHSRV2 mutant clones (M61 and M60) were generated by replacing a 1.35-kb SmaI/FacI fragment with respective PCR amplons (18).

The replication-deficient murine leukemia virus (MuLV) vector pczCFG2 EGN is based on SFG GFPs65T (22). In pczCFG2 EGN, the CMV enhancer/promoter replaces the U3 region of the 5' long terminal repeat (LTR) and drives transcription in the producer cell; however, expression of the EGN marker protein is driven by the reconstituted wild-type MuLV LTR upon reverse transcription and integration into the genome of the target cell. The IEGN marker gene was generated by recombinant PCR and contains the neomycin resistance gene fused in frame to the C terminus of the EGFP gene.

FV SP<sup>Env</sup>-specific polyclonal antiserum. The prokaryotic expression construct for the fusion protein of maltose binding protein (MBP) and HFV SPEnv was generated by inserting a Klenow enzyme-blunted BamI/EcoRI fragment of pCHFwEnv EM50 into the XmnI/EcoRI sites of pcMAL-C2 (New England Biolabs). The soluble fusion protein was generated in Escherichia coli TB1 cultures after induction with 0.5 mM isopropylthiogalactopyranoside (IPTG) for 3 to 6 h and affinity purified according to the manufacturer’s instructions. Envelope protein expression at the cell surface was analyzed by autoradiography. The resultant expression was determined by flow cytometry of the recipient cells 48 h later. Absolute percentages of EGFP-positive cells ranged from 30 to 60% and 2 to 4% for wild-type FV Env (EM02) with FV and MuLV vectors, respectively. Mock-transduced cells gave values of maximal 0.1% positive cells. All transduction experiments were performed at least three times; in each independent experiment, the values obtained with wild-type FV Env (EM02) were arbitrarily set to 100 in the case of FV capsids and to 1 in the case of MuLV capsids.

In some experiments involving FV vectors, intracellular viral particles were artificially released by a freeze-thawing of the transfected 293T cells and subsequent centrifugation and filtration of the supernatant through 0.45-μm-pore-size filters to remove cellular debris. The resulting supernatants were then assayed as described above.

Metabolic labeling and analysis of particle release. For radioimmunoprecipitation analysis (RIPA), transiently transfected 293T cells were metabolically labeled with [35S]methionine and [35S]cysteine for approximately 20 h. Alternatively, cells were pulse-labeled for 30 min and chased for various time periods in fresh growth medium containing a 10-fold excess of cold methionine and cysteine. Subsequently the cells were lysed in RIPA buffer (20 mM Tris [pH 7.4], 0.3 M NaCl, 1% Triton X-100, 0.1% [wt/vol] sodium dodecyl sulfate [SDS]) containing protease inhibitors. Viral proteins were precipitated as described earlier (7, 23), using rabbit antiserum directed against recombinant FV proteins and specific for Env (23) and Gag (13). For endoglycosidase treatment, protein A-Sepharose cluates were denatured by boiling in 0.5% SDS-1% β-mercaptoethanol and subsequently incubated with endoglycosidase H (endo H) peptide N-glycosidase or (PNGase F) in the appropriate incubation buffer as suggested by the manufacturer (New England Biolabs) prior to loading on gels for SDS-polyacrylamide gel electrophoresis (PAGE). Particle-associated proteins were analyzed after centrifugation through a 20% sucrose cushion as described previously (7, 23).

For Western blot analysis, transiently transfected 293T cells were lysed in RIPA buffer, and purified viral particles were obtained by ultracentrifugation as described above. Further purification by equilibrium sedimentation centrifugation using a 8.5 to 40% iodixanol (Optiprep; Gibco BRL) step gradient was essentially performed as described elsewhere (2). The protein samples were subjected to SDS-PAGE and semidry blotted onto nitrocellulose membranes (Amersham). The blots were incubated with rabbit antiserum raised against recombinant HFV Gag (13) or the MBP-HFV SP<sup>Env</sup> fusion protein described above and were developed with the Amersham ECL (enhanced chemiluminescence) detection system.

Cell surface biotinylation of 293T cells transiently transfected with the individual Env expression constructs was carried out essentially as described recently (29). Briefly, 293T cells were transiently transfected and metabolically labeled with [35S]methionine and [35S]cysteine. At 36 h after addition of the DNA, cell surface protein was labeled with N-hydroxysuccinimidobiotin (Calbiochem) at 1 mg/ml in phosphate-buffered saline for 30 min. Subsequently, the biotinylation reaction was stopped by adding phosphate-buffered saline containing 100 mM glycine prior to cell lysis in RIPA buffer. lysates were precipitated with a FV SP Env-specific polyclonal antiserum (7, 23) and incubated with Protein A-Sepharose. Particle-associated proteins were separated by SDS-PAGE, and blotted onto nitrocellulose membranes (Hybond ECL; Amersham). Envelope protein expression at the cell surface was analyzed using streptavidin conjugated to horseradish peroxidase (Pierce), followed by detection with ECL (Amersham). The chemiluminescent biotin signal was allowed to fade overnight. Thereafter, the blot was exposed to X-ray film, and total cellular envelope expression was detected by autoradiography.

RESULTS

The FV SP<sup>Env</sup> is cleaved posttranslationally, and cleavage products are viral particle associated. A special structural feature of the FV Env protein is an unusually long N-terminal SP sequence with an SP cleavage site predicted to be located after aa 86 (Fig. 1) (8, 35). The majority of secretory SP are cotranslationally cleaved; however, for some retroviral glycoproteins, e.g., HIV-1 and feline immunodeficiency virus (FIV), cleavage takes place late during intracellular transport (21, 34). To determine whether and when FV SP cleavage occurs we generated a FV SP<sup>Env</sup>-specific polyclonal antiserum by immunizing rabbits with a protein containing the N-terminal 86 aa of HFV Env fused to MBP. The FV Env precursor glycoprotein
gp130 was efficiently detected by this antiserum in immunoprecipitation analysis experiments using lysates of 293T cells cotransfected with the Gag/Pol-expressing FV vector pMH62 (28) and an expression vector for wild-type FV Env (Fig. 2A), indicating that gp130 still contains the SP (Fig. 2B, lane 1 to 3). Furthermore, upon longer exposure a specific faint band about 18 kDa in size was detected, but not the 80 kDa SU subunit (Fig. 2B, lanes 1 and 2).

In contrast, an anti-SP/SU serum raised against the N-terminal 571 aa (23) precipitated the precursor gp130 and, in addition, the p18 protein, gp80SU, and gp48TM (Fig. 2B, lanes 4 and 5). By Western blot analysis using the SP-specific antiserum, both the gp130 and the 18-kDa protein, and an additional 14-kDa protein were detected (Fig. 2C, lane 6). On autoradiograms of metabolically labeled FV particles derived from the supernatant of infected cell cultures and purified through a sucrose cushion, a faint band of 18 kDa was detectable (Fig. 2B, lane 7). Western blot analysis of virus particle lysates using the SP-specific antiserum confirmed this protein to be the 18-kDa SP cleavage product (Fig. 2C, lane 12). Besides the 18-kDa protein, two additional minor cleavage products approximately 28 and 32 kDa in size were recognized by the SP-specific antiserum in viral particles (Fig. 2C, lane 12).

To ensure the authenticity of these protein bands as integral parts of virus particles rather than inadvertently enriched cellular proteins, FV particles were further purified by equilibrium sedimentation gradient centrifugation. All three SP cleavage products described above were detected in fractions that contained the Gag proteins, demonstrating their particle association (Fig. 2D, fractions 6 to 10). Neither the gp130 precursor nor the p14 cleavage product seen in cell lysates was found in viral particle preparations (Fig. 2C, lane 12; Fig. 2D, fractions 6 to 10). Similar to what was observed with cell lysates, the gp80 SU subunit did not react with the anti-SP antiserum (Fig. 2C, lane 12; Fig. 2D).

For further analysis of the kinetics of FV SPEnv cleavage, transfected 293T cells were pulse-labeled for 30 min and then lysed immediately or chased for various time periods. Subsequently, maturation of the FV Env protein was analyzed by radioimmunoprecipitation using anti-SP or anti-SP/SU antiserum combined with glycosidase treatment. As shown in Fig. 2E, gp130 was efficiently recovered with both anti-SP/SU and anti-SP antiserum. Partially endo H-resistant forms of gp130 could be detected with both antiserum at the 1- and 3-h time points, although somewhat more efficiently with the anti-SP/SU antiserum (Fig. 2E, lanes 8 and 9). Furthermore, SU and TM subunit cleavage products were detectable only with the anti-SP/SU antiserum at the same time points. The TM subunit remained endo H sensitive during the complete chase period, indicating that the oligosaccharide chain are of high-mannose or hybrid but not of complex type.

Taken together, these data indicate that as described for the HIV-1 Env protein, no efficient cotranslational FV SPEnv processing occurs. However, in FV the SPEnv appears to be part of the mature virion. Since the SU subunit found in cells and viral particles no longer contains the SP, its removal probably occurs before or at the same time as SU/TM subunit processing.

The FV SPEnv is cleaved by a cellular protease. Some retroviral glycoproteins are processed by the viral protease. The TM subunit p15E of MuLV Env, for example, is cleaved by the cellular protease. Some retroviral glycoproteins are processed by the viral protease. The TM subunit p15E of MuLV Env, for example, is cleaved by the
MuLV protease during or shortly after capsid budding to remove a 16-aa inhibitory peptide, thereby activating the fusogenic capacity of the protein (12, 16, 30, 31). Since the n-domain of the SP is located in the cytoplasm and therefore is theoretically accessible to the viral protease, we intended to determine whether the FV protease might be involved in SP cleavage. The FV protease removes a small 27-aa peptide from the C terminus of Gag as an essential step to retain viral infectivity (6, 36). Two protease active-site mutants in the context of an infectious molecular clone were used to address this question. Mutant pcHSRV2-M61 (D24A) inactivates the protease (20), whereas pcHSRV2-M80 has, in addition, a translational stop codon introduced at the protease cleavage site in the Gag open reading frame (ORF), terminating RAVN and

FIG. 2. Identification of FV Env subdomains by different antisera. Cell or virus lysates of 293T cells transfected with the Gag/Pol-expressing FV vector pMH62 (Gag/Pol), wild-type FV Env expression construct pHFE EM02 (Env), or empty expression vector (pCDNA3.1) as indicated were analyzed by RIPA or Western blotting. The identity of each FV protein is indicated. The gp48TM protein often comigrates with an immunoreactive cellular protein present in negative controls. (A) Schematic outline of the transfected expression constructs. SFFV, spleen focus-forming virus; bgH pA, bovine growth hormone polyadenylation site. (B) RIPA of cells labeled for 20 h using polyclonal rabbit sera specific for Gag (α-Gag), SPEnv/SU (α-SP/SU), or SPEnv (α-SP). A longer exposure of the lower part of the SDS–10% polyacrylamide gel with the cellular lysates is shown separately, and an autoradiogram of metabolically labeled purified FV particles separated by SDS-PAGE is shown to the right. (C) Western blot analysis (SDS-PAGE [12% gel]) of cellular lysates and purified FV particles using polyclonal rabbit sera specific for Gag (α-Gag) or SPEnv (α-SP). (D) Western blot analysis (SDS-PAGE [12% gel]) of equilibrium sedimentation gradient fractions using polyclonal rabbit sera specific for Gag (α-Gag) and SPEnv (α-SP). (E) Pulse-chase analysis of FV Env maturation. Transfected 293T cells were pulse-labeled for 30 min and then chased for different time periods as indicated at the top with fresh growth medium containing an excess of cold methionine and cysteine. Equal cell lysate samples were immunoprecipitated with FV-specific antisera as indicated to the left. Equivalent aliquots of the protein A eluates were incubated with glycosidases as indicated at the top prior to separation by SDS-PAGE (7.5% gel). Identities of the different forms (g [fully glycosylated], h [endo H resistant] and p [N-deglycosylated]) of gp130Env (solid arrows), gp80SU (shaded arrows), and gp48TM (open arrows) are indicated. The bands marked with asterisks at the 1- and 3-h time point after immunoprecipitation with anti-SP/SU antiserum and PNGase F treatment represent the fully N-deglycosylated form of SU running only slightly faster than the fully glycosylated form of TM.
The n-region of the FV SP is required for FV membrane envelopment but not for targeting to the secretory pathway or envelope function. The FV SP\textsubscript{Env} contains four in-frame translation initiation codons upstream of the predicted SP cleavage site (Fig. 5A). To determine the actual translation initiation site and to examine whether certain domains of the FV SP\textsubscript{Env} might be involved in FV particle maturation, we analyzed several N-terminal truncation and point mutants with respect to their cell surface transport and ability to support FV particle release (Fig. 5A). After cotransfection of 293T cells with the FV Gag/Pol-expressing vector pMH62 and the individual mutants, protein expression was detected (Fig. 5B, lane 5). For all other mutants, as indicated by the appearance of the cellular p18 and p28, and p32 cleavage products (lanes 13 to 15). Clearly, a cellular protease rather than the viral protease is responsible for FV SP\textsubscript{Env} cleavage.

The FV SP\textsubscript{Env} is glycosylated and is cleaved beyond aa 86. The size of the major SP cleavage product of 18 kDa was larger than expected for an N-terminal peptide of 86 aa. Indeed, eukaryotic expression of the predicted 86-aa SP (EM50) yielded a protein of about 10 kDa (Fig. 4B, lane 8). To determine whether posttranslational modifications such as N-glycosylation and/or the use of an alternate cleavage site might account for this size discrepancy, several point mutants were analyzed. To test for N-glycosylation, the first three potential N-glycosylation sites in Env at N\textsubscript{109}, N\textsubscript{119}, and N\textsubscript{141} (Fig. 1) were inactivated by N-to-Q changes either individually or in combinations (Fig. 4A). All mutants were able to generate infectious FV vector particles with similar infectivities as the wild-type Env (Fig. 4A). 293T cells were cotransfected with the individual mutants and the pMH62 vector, and cellular lysates were analyzed by Western blotting with SP- and Gag-specific antisera. Only inactivation of the second potential N-glycosylation site at N\textsubscript{109} resulted in a change of the major cleavage product (Fig. 4B, lane 2). Interestingly, this change resulted in concomitantly with the 14-kDa cleavage product (Fig. 4B, lane 1). Furthermore, glycosidase treatment of cell lysates prior to Western blot analysis had the same effect (data not shown). These data show that the FV SP is removed C terminally of N\textsubscript{109} and, therefore, substantially larger than previously thought (8, 35). In addition, they imply that the 14-kDa cleavage product represents a form of gp18 lacking glycosylation at N\textsubscript{109}. As far as studied, however, SP glycosylation, does not appear to be essential for viral infectivity.

A computer-assisted comparison of all known FV Env sequences revealed a conserved IPQG motif at aa 148 of the HFV sequence. Interestingly, this conserved motif contains a PXG sequence as seen in C termini of many cleavable SP (25). To test whether this sequence motif might be involved in FV SP\textsubscript{Env} cleavage, we analyzed the two mutants EM83 and EM84, bearing C\textsubscript{148}R and G\textsubscript{148}R mutation (Fig. 4A), respectively, in the putative −1 position of SPC cleavage sites. Similar mutations have been shown previously to inhibit SP cleavage (9). The EM83 mutation had no effect on SP cleavage (Fig. 4B, lane 10); however, infectivity was significantly reduced (Fig. 4A), indicating that C\textsubscript{148} is structurally important for Env function but not for SP cleavage. In contrast, SP cleavage of the EM84 mutant was almost completely abolished (Fig. 4B, lane 11). In addition, no infectious FV vector particles were detected in the supernatant of EM84-transfected cells (Fig. 4A) because FV particle release by the EM84 mutant was heavily impaired, at least 10-fold compared to wild type (Fig. 4C, lane 21). These results show that the SP is not cleaved after C\textsubscript{148} but support a role for SP cleavage of the conserved motif around aa 148. Furthermore, they indicate that SP cleavage is essential for efficient FV particle release and infectivity.

The FV SP\textsubscript{Env} is required for FV membrane envelopment but not for targeting to the secretory pathway or envelope function. The FV SP\textsubscript{Env} contains four in-frame translation initiation codons upstream of the predicted SP cleavage site (Fig. 5A). To determine the actual translation initiation site and to examine whether certain domains of the FV SP\textsubscript{Env} might be involved in FV particle maturation, we analyzed several N-terminal truncation and point mutants with respect to their cell surface transport and ability to support FV particle release (Fig. 5A). After cotransfection of 293T cells with the FV Gag/Pol-expressing vector pMH62 and the individual mutants, protein expression was detectable only for the EM41 mutant, which utilizes the first translation initiation codon of the FV Env ORF, thereby containing a full-length SP (Fig. 5A). In agreement with the infectivity data, FV particle release was observed only with EM41 (Fig. 5B, lane 2). For the EM44 mutant, with the first 71 aa of the SP removed, no protein expression was detected (Fig. 5B, lane 5). For all other mutants, protein expression could be detected by RIPA in cell lysates (Fig. 5B). However, some of the mutants (EM42, EM71, EM72, and EM73) showed no detectable SU/TM processing (Fig. 5B, lanes 3 and 9 to 11). The reason for this is unclear. In addition, we analyzed cell-associated virus of the mutants in this study, because FV buds predominantly intracellularly and we previously identified a mutant FV Env protein that was deficient in particle release into the supernatant but still showed budding into intracellular compartments (28).
Results obtained with the supernatants of freeze-thaw lysates of transfected cells indicated that potentially infectious FV particles were not intracellularly trapped in cases of those Env mutants that did not support release of infectious virus into the supernatant (data not shown). Electron microscopy analysis of deletion mutants revealed the presence of naked capsids in the cytoplasm of cells transfected with mutants not releasing infectious particles, while for EM41, which behaved like wild-type Env, particles associated with and budding through cellular membranes were observed (data not shown).

Interestingly, mutants with N-terminal deletions ranging from 5 up to 40 aa (EM42, EM43, EM70, and EM71) that were no longer able to support FV particle egress expressed protein at the cell surface (Fig. 5B, lanes 3, 4, 8, and 9) and gave rise to infectious pseudotyped MuLV capsids (Fig. 5A). Some of these mutants dramatically increased MuLV titers, one up to 150-fold (Fig. 5A). This observation correlated quite well with an enhanced cell surface biotinylation of the various mutants, although the EM43 and EM70 mutants, displaying similar cell surface expression levels (Fig. 5B, lane 4 and 8), showed a fivefold difference in their MuLV pseudotype titers. This indicated that the N-terminal truncations of the mutant Env proteins neither induced inherent defects in targeting the glycoprotein to the secretory pathway nor affected receptor binding and fusion capacities. The importance of N-terminal amino acid of the FV SP n-region for particle budding was further supported by analysis of point mutants in evolutionary conserved residues. Mutant EMS2 had the N-terminal W₁₀W₁₃...
motif replaced by alanines, EM53 had the C-terminal Y_{15} Y_{59} motif replaced by alanines, and EM54 was a combination of both (Fig. 5A). The EM52 mutant as well as the EM54 double mutant no longer supported FV particle release, whereas results with EM53 were similar to those for the wild-type protein (Fig. 5A and C, lanes 13 to 15).

To further delineate the region of the SP required for FV particle release and infectivity, several mutants with internal deletions in the SP n-region were analyzed. Smaller deletions of 3 to 24 aa between aa 16 and 66, such as EM66, EM67, EM74, and EM76, were tolerated, resulting at the most in 45-fold-reduced infectivity (Fig. 5A and D, lanes 19, 20, 23, and 25). However, larger deletions of 25 to 50 aa, such as in EM68, EM69, and EM75, abolished infectivity, although for EM75 particle release into the supernatant could be detected (Fig. 5A and D, lanes 21, 22, and 24). 293T cells cotransfected with the EM66 mutant showed very strong syncytium formation, indicating that it had a highly increased fusogenic activity compared to the wild-type FV Env protein (data not shown). In contrast, all other internal deletion mutants showed no obvious difference in their fusogenic activity (data not shown). This might explain why no infectious MuLV vectors pseudotyped by EM66 could be detected.

Taken together, these data show that the region comprising the N-terminal 15 aa of the FV SP_{Env}, while being dispensable for targeting to the secretory pathway and proper envelope function, is specifically involved in the FV particle budding process. Furthermore, they point to a critical role of two conserved N-terminal tryptophan residues in this process. The central part of the SP n-region on the other side does not seem to play a crucial role, as it tolerates smaller deletions. However, there are some constraints regarding spacing of the N-terminal budding domain in respect to the h-region, as larger deletions negatively influence particle release and infectivity.

**DISCUSSION**

Recently it has become clear that SP can have several additional functions apart from being responsible and essential to target glycoproteins to the secretory pathway (reviewed in reference 25). For some retroviral envelope glycoproteins, a post-translational cleavage of the SP sequence has been reported (21, 34). Similarly, we observed posttranslational cleavage of the FV SP_{Env} sequence, as full-length FV gp130 was detected (21, 34). The EM52 mutant as well as the EM54 double mutant no longer supported FV particle release, whereas results with EM53 were similar to those for the wild-type protein (Fig. 5A and C, lanes 13 to 15).

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ently think that the FV Sp Env budding domain mediates the primary interaction with the FV capsid and an interaction with MSD plays a role later during the budding process.

Interestingly, FV Gag proteins, unlike other retroviral Gag proteins, are not processed into matrix, capsid, and nucleocapsid subunits, and as mentioned earlier, wild-type FV capsids are not found to be associated with cellular membranes in the absence of FV Env expression. Therefore, it may be conceivable that the FV Sp Env performs functions in viral assembly and budding, such as membrane targeting of the capsid, analogously to the matrix subunit of other retroviruses. Furthermore, it is likely that the particle-associated Sp cleavage products have additional functions in the FV replication cycle. Even a role of Sp in binding to the cellular receptor or fusion of viral and cellular membranes should be addressed in further studies.

Surprisingly, deletion of the N-terminal budding domain of the FV Env protein but not internal Sp n-region deletions dramatically increase pseudotype titers of MuLV capsids. This shows that the budding domain is physically separable from those Sp domains required for targeting to the secretory pathway and normal envelope function, namely, receptor binding and membrane fusion. Furthermore, these results suggest an inhibitory role of this region for FV Env incorporation into heterologous retroviral particles, which may be a cause for the poor pseudotyping capacity observed for the wild-type FV Env protein (22). However, to determine if this is an active exclusion of FV Env proteins containing this domain from heterologous budding particles or simply a result of the different levels of cell surface expression observed for most of these mutants, a more detailed analysis is required.

Based on our findings, it will be interesting to analyze functions of other retroviral glycoproteins Sp for Env incorporation and particle maturation. The FIV Env (33, 34) and the mouse mammary tumor virus Env (1, 17), for example, bear also unusually long Sp sequences. FIV Env can tolerate extensive deletions in the Sp sequence with no effect on membrane targeting and intracellular transport; however, the effects of such deletions on envelope particle incorporation and infectivity have not been investigated (33). Similarly, Sp mutations and chimeras of HIV-1 Env have also been examined only with respect to their intracapsidacknowledgments

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