In vivo genetic mutations define predominant functions of the human T-cell leukemia/lymphoma virus p12\textsuperscript{I} protein

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The human T-cell leukemia/lymphoma virus type 1 (HTLV-1) ORF-I encodes a 99–amino acid hydrophobic membrane protein, p12\textsuperscript{I}, that affects receptors in different cellular compartments. We report here that proteolytic cleavage dictates different cellular localization and functions of p12\textsuperscript{I}. The removal of a noncanonical endoplasmic reticulum (ER) retention/retrieval signal within the amino terminus of p12\textsuperscript{I} is necessary for trafficking to the Golgi apparatus and generation of a completely cleaved 8-kDa protein. The 8-kDa protein in turn traffics to the cell surface, is recruited to the immunologic synapse following T-cell receptor (TCR) ligation, and down-regulates TCR proximal signaling. The uncleaved 12-kDa form of p12\textsuperscript{I} resides in the ER and interacts with the β and γ\textsubscript{E} chains of the interleukin-2 receptor (IL-2R), the heavy chain of the major histocompatibility complex (MHC) class I, as well as calreticulin and calnexin. Genetic analysis of ORF-I from ex vivo samples of HTLV-1–infected patients reveals predominant amino acid substitutions within ORF-I that affect proteolytic cleavage, suggesting that ER-associated functions of p12\textsuperscript{I} may contribute to the survival and proliferation of the infected T cells in the host.

Introduction

Human T-cell leukemia/lymphoma virus type 1 (HTLV-1) is the etiologic agent of a rare but aggressive hematopoietic malignancy of T cells, designated adult T-cell leukemia/lymphoma (ATLL), as well as a progressive myelopathy associated with human T-cell leukemia/lymphoma virus (HTLV-I)-associated myelopathy/tropical spastic paraparesis (HAM/TSP). In addition to structural and enzymatic proteins, the viral RNA genome encodes different cellular compartments. We re-

among those, the p12\textsuperscript{I} protein encoded by open reading frame I (ORF-I) is a 99–amino acid highly hydrophobic protein containing 4 putative SH3 binding motifs, 1 leucine zipper domain, 1 leucine zipper-like domain, and 2 putative transmembrane domains (TM1, TM2). Evidence of the expression and importance of this protein in HTLV-1 pathogenesis is at present indirect. The singly spliced mRNA encoding p12\textsuperscript{I} is found in infected cells in vitro and in ex vivo samples from HTLV-1–infected patients. In addition, experimentally infected animals produce antibodies that recognize recombinant p12\textsuperscript{I} and HTLV-1–infected individuals mount a cytotoxic T lymphocyte (CTL) response to ORF-I.

Two isoforms of p12\textsuperscript{I} have previously been described, the p12\textsuperscript{I}K and p12\textsuperscript{I}R that are ubiquitinated and targeted for degradation by the proteasome, whereas the more stable p12\textsuperscript{I}R protein encodes an arginine at position 88 and is therefore not ubiquitinated. However, no specific disease association of the 2 p12\textsuperscript{I} isoforms has previously been established. Several functions have been ascribed to p12\textsuperscript{I}. Ectopically expressed p12\textsuperscript{I} resides in the endoplasmic reticulum (ER) and traffics to the cell surface, is recruited to the immunologic synapse following TCR ligation, and down-regulates TCR proximal signaling. The uncleaved 12-kDa form of p12\textsuperscript{I} resides in the ER and interacts with the β and γ\textsubscript{E} chains of the interleukin-2 receptor (IL-2R), the heavy chain of the major histocompatibility complex (MHC) class I, as well as calreticulin and calnexin. Genetic analysis of ORF-I from ex vivo samples of HTLV-1–infected patients reveals predominant amino acid substitutions within ORF-I that affect proteolytic cleavage, suggesting that ER-associated functions of p12\textsuperscript{I} may contribute to the survival and proliferation of the infected T cells in the host.

In an attempt to reconcile the seemingly disparate functions of this viral protein, we found that the p12\textsuperscript{I} protein undergoes complex posttranslational modifications that include proteolytic cleavage between amino acid positions 9 and 10 followed by a second cleavage between the amino acids 29 and 30. The first proteolytic cleavage removes a noncanonical ER retention/retrieval signal at the amino terminus of p12\textsuperscript{I} and allows for...
further trafficking of this viral protein to the Golgi apparatus and the lipid rafts. Importantly, we found a high frequency of genetic mutations in the ORF-1 of provirus from HTLV-1–infected individuals causing ER retention of p12I, suggesting an important role for p12I functions in the ER in vivo.

Methods

Expression plasmids and antibodies

The pME18S p12Δ3SL expression plasmid has been described previously. The p12I and its mutants were generated by polymerase chain reaction (PCR) or the QuickChange Site-Directed Mutagenesis Kit (Stratagene, La Jolla, CA) using site-specific mutagenic oligonucleotides following the manufacturer’s instructions. The pAB-GTG molecular clone molecule was generated by ligation of the Cla-I Sal-I DNA fragment from the pBST molecular clone into the pACH equivalent restriction sites and the initiation codon of p12 mutated as previously described.19

The following oligonucleotides were used and the sequence of plasmid clones was analyzed to confirm the mutations: L9A-F: 5′-GCTCTTCTTACG-CCCGGATCCACTTGAGGAGC-3′; L9A-R: 5′-AGGGCCATGTCGAGCG-CCGGGATCCGGAAGGAGG-3′; S10A-F: 5′-CCTCTTACAGGCGATTGCGTC- TACCTGGCGCTA-3′; S10A-R: 5′-TGAGGAAGAGCAGGAGC-3′; S29F: 5′-TGCTCTTCCGGGAGGCTGAGCGACTCTG- TCTTCTC-3′; S29R: 5′-GCCGGAAAGAGGGCGTCTGAGCTGCC-3′; S29-D24: 5′-AGGCTCTTGGAGGGCCGTCAGGCGCTTCTC-3′; S29-D29: 5′-TGCTCTTGAGAACAACAAACTGTCATTATTTGTTC- AGGTTCA-3′; S29P-F: 5′-GCTCTTCTTACG-CCCGGATCCACTTGAGGAGC-3′; S29P-R: 5′-GCCGGAAAGAGGGCGTCTGAGCTGCC-3′; S29P-D24: 5′-AGGCTCTTGGAGGGCCGTCAGGCGCTTCTC-3′; S29P-D29: 5′-TGCTCTTGAGAACAACAAACTGTCATTATTTGTTC- AGGTTCA-3′; S29P-D29: 5′-GCCGGAAAGAGGGCGTCTGAGCTGCC-3′; and S29P-R: 5′-GCCGGAAAGAGGGCGTCTGAGCTGCC-3′; S29P-D29: 5′-GCCGGAAAGAGGGCGTCTGAGCTGCC-3′. The DNA fragments were cloned into the pCR2.1 vector (Invitrogen, Carlsbad, CA) and sequenced.

Because the forward primer overlaps with ORF-1, the first 7 amino acids of p12I do not reflect the actual amino acid sequence in those samples. The remaining 295 clinical samples were amplified and sequenced as described.19,21,28

The anti-HA antibody clones 12CA5 and 3F10-hrp were obtained from Roche Applied Science (Indianapolis, IN); the e6E2 clone with the following primers: p300-ORF(+): 5′-CAGTGCAGATATGCGACATGC- GCTTTCGCTTACGAC-3′; p300-ORF(-): 5′-ATGGATCGAGTCTTCGAGGAGGAGGAGG-3′. The DNA fragments were cloned into the pCR2.1 vector (Invitrogen, Carlsbad, CA) and sequenced.

Figure 1. Mutations of amino acids at positions 10 and 29 affect p12 cleavage and cellular localizations. (A) Single-letter amino acid code of p12 from HTLV-1. The arrows indicate the computer-predicted cleavage sites. (B) Schematic representation of the p12 mutants. Changes in amino acid positions are indicated on the top. The L9 and S10 were changed to alanines. Glycine 29 was changed to serine. The Δ symbol signifies deletion of amino acids at the amino terminus for the truncated mutants D24 and D29. (C) Cell lysates from the 293T cells transfected with the p12I mutants were analyzed by Western blot with anti-HA antibody. (D) Localization of p12I, p12IG29S, and p12I1-32 in Jurkat T cells stained with an anti-PDI antibody. (E) Localization of p12I, p12IG29S, and p12I1-32 in HeLa cells compared with ER and Golgi markers. Red-ER is used as the ER marker, whereas an antibody to mannosidase II (Mann II) is used to label the Golgi apparatus.
Alexa-488 conjugation was obtained from Cell Signaling Technology (Danvers, MA). Rabbit polyclonal mannosidase II antibody—Golgi marker, mouse monoclonal (RL90) to PDI–ER marker, and mouse monoclonal antibody (FMC 75) to calreticulin were all obtained from Abcam (Cambridge, MA).

**Cell culture and DNA transfection**

Jurkat T cells were grown in RPMI 1640 and 293T, COS-7, and HeLa cells were grown in Dulbecco modified Eagle medium (DMEM) and both media were supplemented with 10% FCS, 2 mM penicillin-streptomycin, and 5 mM l-glutamine (Invitrogen), respectively.

Transfections were performed using DMRIE-C (Invitrogen) or the Cell Line Nucleofector Kit V (Amaxa, Gaithersburg, MD) for Jurkat T cells, Effectene (QIAGEN, Valencia, CA) for 293T-cells, and FuGENE-6 or FuGENE HD (Roche, Indianapolis, IN) for COS-7 and HeLa cells following the manufacturers’ protocols.

For the TCR stimulation, Jurkat T cells were centrifuged and treated with 1 μg/mL anti-CD3e (UCH-T1) from EMD Biosciences (La Jolla, CA) in fresh media. The cells and culture media were harvested at the indicated times.
Immunofluorescence and confocal microscopy

For the formation of the immunologic synapse, Jurkat T and Raji cells were prepared separately at the concentration of 10^6 cells/mL in the fresh medium. Raji cells were prepulsed with or without 10 μg/mL SEE from Toxin Technology (Sarasota, FL) for 1 hour at 37°C. Cells were combined at a 1:1 ratio and fixed by adding 4% paraformaldehyde (PFA) at indicated time point. Cells (10^5) were analyzed on a glass slide coated with poly-L-lysine. Cells were washed in PBS and permeabilized in ice-cold methanol for 20 minutes at −20°C followed by staining with indicated antibodies.

HeLa cells and COS-7 cells were seeded on 12-mm coverslips and transfected the following day by FuGENE HD or FuGENE-6. Cells on coverslips were fixed 48 hours after transfection with 4% PFA in PBS for 20 minutes, washed in PBS, and permeabilized in ice-cold methanol for 20 minutes at −20°C. For immunofluorescence the fixed cells were washed 3 times in PBS before being inverted onto a nescofilm containing a 25-μL droplet of 0.5% BSA in PBS blocking buffer for 15 minutes. The cells were further incubated for 1 hour with 25-μL droplets containing primary antibodies in appropriate dilutions, washed 3 × 5 minutes in PBS, and incubated in anti–mouse or anti–rabbit Alexa-conjugated secondary antibodies for 1 hour in the dark followed by 3 × 5 minutes washes in PBS. Finally, the cells were rinsed quickly in water before being mounted in 5 μL mounting media on a glass slide before being examined by Zeiss laser scanning confocal microscope LSM 510 (Carl Zeiss MicroImaging, Thornwood, NY) with a 63 ×/100 ×/1.3 NA Plan Apochromat oil objective.

Luciferase assay

The Dual-Luciferase Reporter Assay System was obtained from Promega (Madison, WI) and all the measurements were performed according to the manufacturer’s protocol. Luciferase values were normalized to protein content and protein concentration was determined using the Bio-Rad (Hercules, CA) Protein Assay. All experiments were repeated at least 3 independent times and representative data are shown as average and standard deviation of triplicates.

Quantification of HTLV p19 Gag

The HTLV p19 Gag enzyme-linked immunosorbent assay (ELISA) kit from ZeptoMetrix (Buffalo, NY) was used to quantify p19 Gag in the supernatant. Direct or diluted cell-culture supernatants were used for the analyses to obtain concentrations within the standard range. Experiments were repeated at least 3 times and all the measurements performed according to the manufacturer’s protocol are shown as average and standard deviation of duplicates.

Amino acid sequence analysis

A total number of 304 p12I sequences for subtypes A and B (cosmopolitan A and B) were analyzed. These sequences were derived from ex vivo proviral DNA samples from HTLV-1–infected individuals (continued). (B) Unrooted neighbor-joining tree for 65 variants. Cosmopolitan type B variants are represented by open circle and highlighted in blue, and cosmopolitan type A variants are represented by red triangles. The closed red triangles indicate samples with a frequent glycine to serine mutation at position 29. (C) Lysates from 293T cells transfected with plasmids encoding the p12V51, p12V52, and p12V2 variants were analyzed by Western blot with the anti-HA antibody.
DNA samples of HTLV-1–infected patients. The 203 subtype B samples could be grouped into 24 different variants and the remaining 101 subtype A samples were grouped into 41 different variants (Figure 2A). We used European Molecular Biology Open Software Suite (EMBOSS) for sequence handling and formatting. A The package for the R project for statistical computing (Foundation For Statistical Computing, http://www.R-project.org) was used to calculate genetic distance (Jukes-Cantor genetic distance model) and to generate the unrooted neighbor-joining tree (Figure 2B).

Results

Proteolytic cleavage dictates the cellular localization of the 12-kDa and 8-kDa forms of the p12I complex

Expression of the singly spliced ORF-I cDNA (Figure 1A) consistently results in the expression of 2 or more protein bands that are usually referred to as p12I (Figure 1C lane 2), whose nature and derivation are unknown. Computer program analysis (PSORT; http://psort.nibb.ac.jp; Kenta Nakai, Human Genome Center, Institute for Medical Science, University of Tokyo, Tokyo, Japan) and ELM (http://elm.eu.org) of the p12I amino acid sequence predicted 2 possible signal peptide cleavage sites at positions L9, S10 and VSG29, L3004 (Figure 1A top). Indeed mutation of serine at position 10 to alanine resulted predominantly in the production of the 12-kDa protein (Figure 1B,C lane 4), whereas mutation of leucine at position 9 to alanine appeared to favor the generation of the 8-kDa protein (Figure 1C lane 3). In addition, the G29→S mutant yielded a single uncleaved protein product of 12 kDa (Figure 1B,C lane 7), whereas substitution of leucines 30 and 31 with alanine did not affect the generation of the 8-kDa protein band (data not shown). Additional substitution of leucine with alanine at position 9 in the p12G29S mutant partially restored cleavage (Figure 1B,C lane 8), whereas a mutant lacking the first 29 amino acids (p12IΔ29) yielded a protein of 8 kDa (Figure 1B,C lane 10), and a mutant lacking the first 24 amino acids (p12IΔ24) (Figure 1B) resulted in similar migration (data not shown). Altogether these data are consistent with the finding that, at steady state, the p12I complex is constituted by an uncleaved 12-kDa precursor molecule and a completely cleaved 8-kDa protein.

The p12I protein complex has been reported to localize to the ER, Golgi, and the lipid rafts. To assess how the proteolytic cleavage of p12I affects its cellular localization, we transfected Jurkat T, HeLa, or COS-7 cells with the wild-type p12I, the p12G29S mutant, as well as the deletion mutant p12IΔ29 (Figure 1B). In Jurkat T cells, the p12I wild-type protein and the p12IΔ29 mutant localized diffusely and in dots in the cytoplasm (Figure 1D). This localization was also observed for the p12IΔ24 mutant (data not shown). In contrast, when proteolytic cleavage was abrogated by mutation at amino acid position 29, p12I accumulated near the nucleus (Figure 1D). To further study the cellular localization of this mutant, we compared the distribution of p12I and the p12G29S mutant to the localization of the ER-resident proteins, protein disulphide isomerase (PDI) and calreticulin. We noticed a much lower staining intensity of PDI (Figure 1E) and calreticulin (data not shown) in cells expressing p12I and the p12G29S mutant (Figure 1E). Similar observations have been reported for the BPV E5 protein, a functional analog of p12I that alters the ER distribution. To circumvent this problem, HeLa cells were cotransfected with pDsRed2-ER (Red-ER). The Red-ER construct expresses the red fluorescent protein from Discosoma sp which has the ER targeting sequence of calreticulin fused to the 3′ and the ER retention signal KDEL fused to the 3′ of the pDsRed2. To define the Golgi apparatus, we used an antibody to mannosidase II, which is a medial Golgi localized protein. The wild-type p12I and the p12IΔ29 mutant colocalized with Red-ER and partially colocalized with mannosidase II (Figure 1F,G). The expression of the p12G29S mutant caused a redistribution of the Red-ER protein to a dotted pattern (Figure 1F). Furthermore, the p12G29S mutant appeared to be excluded from the Golgi apparatus since no colocalization was observed with mannosidase II (Figure 1G).

The 12-kDa uncleaved form of p12I predominates in ex vivo samples from HTLV-1–infected individuals

The data presented here demonstrate that mutation at the cleavage sites of p12I interferes with its cellular localization. To investigate the relevance of these findings to HTLV-1 infection, we studied genetic polymorphism at the protein level of p12I obtained from 304 ex vivo proviral DNA samples from HTLV-1–infected individuals with either subtype A (nA = 101, 33.2%) or subtype B (nB = 203, 66.8%). The subtypes are characterized by the amino acid at position 23 (proline in subtype B and serine in subtype A). The 203 samples for subtype B could be grouped into 24 different variants, whereas the 101 samples for subtype A were more diverse and could be grouped into 41 different variants (Figure 2A). Seventy percent of all variants are represented by only one sequence sample, whereas V2 with 167 sequences is the most frequent variant in our dataset (Figure 2A). The average pairwise genetic distance was 4.38 for subtype A variants and 1.91 for subtype B variants using Jukes-Cantor correction method. The difference in the number of variants between the subtypes could be due to a sampling bias. Amino acid sequence near the first putative proteolytic cleavage site at position 9 shows a low level of genetic polymorphisms for both subtypes (Figure 2A), whereas the second cleavage site around position 29 is more variable for subtype A and more conserved for subtype B. In addition, position 29 itself is conserved within subtype B but not in subtype A. Twenty-two percent of the variants for subtype A have a serine instead of a glycine at this position. Although the low level of polymorphism among the 65 variants does not provide great statistical power for a detailed phylogenetic analysis, in a best tree analysis the different variants group according to their subtypes except for variant V25 (Figure 2B). It also shows that samples with a serine instead of the glycine at position 29 cluster together. These results suggest that the sequence differences might reflect emergence of functionally distinct variants.

Prototype expression plasmids were generated from variant 51 (V51) that carries the G29S mutation, variant 52 (V52) that has no mutations in this region, and variant 2 (V2) that carries the S23P subtype determining polymorphism. Expression in 293T cells of these prototype p12I variants demonstrated that the p12IΔ29 yielded the expression of the 12-kDa and 8-kDa form as observed for p12I, whereas p12IΔ29 yielded a single uncleaved protein product of 12 kDa. Similar observations have been reported for the BPV E5 protein, a functional analog of p12I that alters the ER distribution. To circumvent this problem, HeLa cells were cotransfected with pDsRed2-ER (Red-ER). The Red-ER construct expresses the red fluorescent protein from Discosoma sp which has the ER targeting sequence of calreticulin fused to the 3′ and the ER retention signal KDEL fused to the 3′ of the pDsRed2. To define the Golgi apparatus, we used an antibody to mannosidase II, which is a medial Golgi localized protein. The wild-type p12I and the p12IΔ29 mutant colocalized with Red-ER and partially colocalized with mannosidase II (Figure 1F,G). The expression of the p12G29S mutant caused a redistribution of the Red-ER protein to a dotted pattern (Figure 1F). Furthermore, the p12G29S mutant appeared to be excluded from the Golgi apparatus since no colocalization was observed with mannosidase II (Figure 1G).
A 2-step proteolytic cleavage is required to remove a noncanonical ER retention/retrieval signal within the first 5 amino acids of p12I

The data presented in Figure 1 are consistent with the hypothesis that a portion of the ORF-I protein product, the 12-kDa form, is retained/retrieved in the ER, whereas the other, the 8-kDa form, may be generated by a 2-step proteolytic cleavage and traffics to the cell surface. To provide further support to the notion that there are 2 cleavage sites within p12I, we generated constructs whereby different portions of the amino terminus of p12I were juxtaposed at the 5’ of the GFP-Mem construct. The GFP-Mem contains 20 amino acids of the plasma membrane targeting signal from neuromodulin fused to the 5’ of green fluorescent protein (GFP) and is targeted to the plasma membrane, as demonstrated in HeLa cells (Figure 3A,B).

We fused the GFP-Mem protein with either the first 15 amino acids of wild-type or of the S10A mutant that is not cleaved (Figure 1C). Expression of p12I1-15-GFP-Mem in HeLa cells demonstrated plasma membrane and intracellular distribution that colocalized with the Red-ER (Figure 3A). In contrast, the p12S10A1-15-GFP-Mem was predominantly localized to the ER (Figure 3A), suggesting that the introduction of alanine at position 10 in the p12I1-15-GFP-Mem construct caused ER retention/retrieval. Similarly, fusion of the first 32 amino acids of either the wild-type p12I or the p12G29S mutant to the GFP-Mem protein resulted in retention/retrieval of the p12I1-32-GFP-Mem and p12G29S1-32-GFP-Mem protein in the ER of both HeLa cells (Figure 3B). These results were also confirmed in COS-7 cells (data not shown).

Thus, these data support the existence of 2 cleavage sites at the amino terminus of p12I and suggest the presence of an ER retention/retrieval signal within the first 9 amino acids of the protein. Indeed, computer analyses predicted a putative noncanonical ER retention/retrieval signal within the first 5 amino acids at the amino terminus: MLFRL. To investigate this, the first 5 amino acids of p12I were fused at the 5’ of the GFP-Mem construct, generating p12I1-5-GFP-Mem. As a positive control, the canonical ER retention/retrieval signal KDEL was fused to the C-terminus of the GFP-Mem construct. The GFP-Mem contains 20 amino acids of the plasma membrane targeting signal from neuromodulin fused to the 5’ of green fluorescent protein (GFP) and is targeted to the plasma membrane, as demonstrated in HeLa cells (Figure 3A,B).

Figure 3, p12I contains a noncanonical ER retention/retrieval signal at its amino terminus. (A) Localization of the GFP-Mem, p12I1-32-GFP-Mem, and p12S10A1-32-GFP-Mem and Red-ER in HeLa cells. (B) Localization of the GFP-Mem, p12I1-32-GFP-Mem, and p12G29S1-32-GFP-Mem compared with Red-ER in HeLa cells. (C) Localization of GFP-Mem-KDEL and p12I1-5-GFP-Mem in COS-7 cells. (D) Localization of the p12I1-5-GFP-Mem compared with Red-ER in HeLa cells.

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We fused the GFP-Mem protein with either the first 15 amino acids of wild-type or of the S10A mutant that is not cleaved (Figure 1C). Expression of p12I1-15-GFP-Mem in HeLa cells demonstrated plasma membrane and intracellular distribution that colocalized with the Red-ER (Figure 3A). In contrast, the p12S10A1-15-GFP-Mem was predominantly localized to the ER (Figure 3A), suggesting that the introduction of alanine at position 10 in the p12I1-15-GFP-Mem construct caused ER retention/retrieval. Similarly, fusion of the first 32 amino acids of either the wild-type p12I or the p12G29S mutant to the GFP-Mem protein resulted in retention/retrieval of the p12I1-32-GFP-Mem and p12G29S1-32-GFP-Mem protein in the ER of both HeLa cells (Figure 3B). These results were also confirmed in COS-7 cells (data not shown).

Thus, these data support the existence of 2 cleavage sites at the amino terminus of p12I and suggest the presence of an ER retention/retrieval signal within the first 9 amino acids of the protein. Indeed, computer analyses predicted a putative noncanonical ER retention/retrieval signal within the first 5 amino acids at the amino terminus: MLFRL. To investigate this, the first 5 amino acids of p12I were fused at the 5’ of the GFP-Mem construct, generating p12I1-5-GFP-Mem. As a positive control, the canonical ER retention/retrieval signal KDEL was fused to the C-terminus of the GFP-Mem protein. Addition of KDEL at the carboxy terminus of GFP-Mem resulted in accumulation of GFP-Mem in the ER (Figure 3C), which was confirmed by comparison and colocalization with Red-ER (Figure 3D). Thus, the amino terminus of p12I contains a noncanonical ER retention/retrieval signal that is removed by a 2-step cleavage to allow trafficking of the 8-kDa form to the cell surface.

The 8-kDa form of p12I, but not the uncleaved 12-kDa form, is recruited to the immunologic synapse and mediates proximal TCR signaling down-regulation.

Ligation of the TCR by peptides from MHC to antigen-presenting cells (APCs) leads to the formation of a complex structure that is termed the immunologic synapse (IS), which includes the TCR, protein tyrosine kinases, and LAT. Since we previously have shown that p12I is recruited to the IS upon TCR ligation, we hypothesized that the 8-kDa but not the 12-kDa form down-regulates TCR signaling. To test this, we used Jurkat T cells and Raji cells, as an in vitro model of CD4+ T-cell and APC interaction, whereby polarization of the TCR can be easily visualized with anti-CD3 following cocultivation of Jurkat T cells with *Staphylococcus aureus* enterotoxin E (SEE) pre-pulsed B cells. Within 15 minutes of coculture, p12I and the p12Δ29 mutant were recruited to the IS formed between the T cells and the B cells (Figure 4A), whereas the p12G29S mutant was not observed in the IS, suggesting that the 8-kDa protein is the isoform recruited to the IS.

To investigate how cellular localization affects p12I function, the abilities of the mutants p12G29S, p12S10A, p12Δ24, p12Δ29, and wild-type p12I to down-regulate proximal TCR signaling and viral replication were assessed in Jurkat T cells. The HTLV-1 LTR-driven luciferase construct was cotransfected with the HTLV-1 molecular clone pACH-GTG, whereby p12I expression was eliminated by a genetic mutation together with p12I and p12I mutants. In this experimental model, ligation of TCR with the anti-CD3 antibody increases Tax activity and viral replication. Because p12I dampens TCR signaling, a readout of Tax activity was measured by the luciferase assay (Figure 4B). Thus, ligation of TCR by the anti-CD3 antibody increased Tax activity by p12I mutant p12G29S. Similarly, viral replication by p19 Gag antigen capture assay of the culture supernatant was assessed (Figure 4C). The expression of p12I mutants was verified by Western blot (Figure 4D). Decreased
Discussion

Viruses that carry limited genetic information exploit complex posttranslational modification to generate protein products with diverse functions, and HTLV-1 is no exception. Here, we show that the HTLV-1 ORF-I 99–amino acid product undergoes complex posttranslational modification that yields 2 protein forms: one that mostly remains in the ER and the other that traffics to the cell surface. Since both mutations at serine 10 and glycine 29 result in loss of the 8-kDa form of p12I, our results suggest that the production of the 8-kDa protein occurs through a 2-step proteolytic cleavage, where the first cleavage occurs between amino acids 9 and 10 (PL ↓ SP) and the second between amino acids 29 and 30 (VSG ↓ LL). Our data support the hypothesis that the first cleavage is necessary to remove the amino terminus ER retention signal LFRL and allow trafficking of the partly cleaved protein to another cellular compartment, likely the Golgi, where the second cleavage can occur.

We provide 3 separate lines of evidence demonstrating that proteolytic cleavage is essential for plasma membrane localization of p12I. First, modification of serine at position 10 and/or glycine at position 29 results in a loss of the 8-kDa form of p12I and neither of these p12I mutants localize to the plasma membrane. Second, fusion of the GFP-Mem to the noncleavable first 15 or 32 amino acids from the p12I S10A or p12IG29S mutants, respectively, impaired trafficking of the GFP-Mem protein to the plasma membrane, demonstrating that cleavage at both positions is necessary for plasma membrane localization of these chimeric proteins. Third, we have identified a noncanonical ER retention/retrieval signal within the 5 amino acids of p12I.

The data presented here suggest a model on how this small viral protein exerts its pleiotropic functions (Table 1; Figure 5). The p12I protein binds to and down-regulates surface expression of the β and γc chains of IL-2R by retaining both IL-2R chains in the ER, and increases signal transducer and activator of transcription 5 (STAT5) phosphorylation. By doing so, p12I decreases the threshold for T-cell activation and IL-2 requirement. The p12I protein also interacts with and down-regulates the cell surface expression of the heavy chain (Hc) of MHC class I. Indeed, the fraction of Hc-MHC class I and IL-2R chains that binds p12I is endoglycosidase H (EndoH) sensitive, demonstrating that this interaction occurs before the trafficking of p12I to the Golgi apparatus. Similarly, the interaction of p12I with the ER-resident proteins calreticulin and calnexin most likely occurs with the uncleaved form of p12I.

In contrast, as we demonstrate here, membrane localization of p12I is essential for TCR proximal signaling down-regulation. Indeed, it is the 8-kDa protein, but not the uncleavable p12I G29→S mutant, that is recruited to the immunologic synapse following TCR engagement and down-regulates proximal TCR signaling. The results reported here explain the contrasting effects of p12I on TCR signaling. It is likely that the uncleaved form of p12I, together with PMA, increases cytoplasmic calcium and NFAT activity, and this effect is LAT-independent. In contrast, after ligation of TCR, p12I decreases proximal signaling in a LAT-dependent manner. Here, we show that this effect is mediated exclusively by the 8-kDa protein.

Collectively our results suggest a model whereby the effects of p12I on IL-2R chains, calreticulin, and calnexin as well as MHC-I-Hc may be mediated by the uncleaved form of p12I that resides in the ER, whereas down-regulation of the TCR is mediated

| Table 1. Comparison of functions and localization of the 12-kDa and 8-kDa isofoms of p12I |
|-----------------|-----------------|-----------------|-----------------|-----------------|
| p12I isoform    | TCR signaling  | Viral replication | Recruitment to IS | Localization    |
| 12 kDa          | –               | –                | –                | ER              |
| 8 kDa           | +               | +                | +                | Plasma membrane |
by the completely cleaved 8-kDa protein (Figure 5). The availability of the p12I mutants generated in the present work will enable us to dissect the role of the uncleaved and the cleaved form of p12I on other molecules affected by p12I, such as LFA-1, ICAM-1, ICAM-2, and the vacuolar ATPase.13,17,20

Importantly, we found that both the naturally occurring mutation S23→P and G29→S observed in HTLV-I–infected individuals inhibit cleavage and the generation of the 8-kDa protein. Interestingly, within the HTLV-I provirus both of these mutations do not result in amino acid sequence changes in the overlapping ORF-II or in the antisense reading frame encoding HBZ. A predominance of the uncleaved form of p12I in vivo would cause increased STAT activation, calcium release, and T-cell proliferation with resulting increase in the number of infected T cells. In turn, the uncleaved form of p12I, by interacting and down-regulating the MHC class II molecules, would favor escape from immune recognition of virus infected cells. Lastly, whether the uncleaved form of p12I also mediates down-regulation of ICAM-1 and ICAM-2 remains to be investigated.

In conclusion, the findings reported here provide the rationale for the design of a prospective study that will attempt to correlate different functions of p12I with proviral load and disease development in HTLV-I–infected individuals.

Acknowledgments

We thank Tom Misteli for helpful discussions; Robyn Washington Parks, Jonathan T. Magruder, Talisa A. Goss, Joanna Ostas, and Tatiana Karpova for helpful suggestions with some of the studies; Tatiana Karpova for helpful suggestions with some of the studies; Tatiana Karpova for helpful suggestions with some of the studies; Tatiana Karpova for helpful suggestions with some of the studies. This research was supported by the Intramural Research Program of the National Institutes of Health (NIH), National Cancer Institute, Center for Cancer Research. R.F. was supported in part by the Japan Society for the Promotion of Science (Tokyo, Japan).

Authorship

Contribution: R.F., V.A., I.B., and V.C. designed and performed research and analyzed data; J.M.N., C.N., and V.W.V. performed research; A.G. contributed samples; J.-C.W. analyzed the protein sequences; and G.F. designed research, analyzed data, and wrote the paper.

Conflict-of-interest disclosure: The authors declare no competing financial interests.

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In vivo genetic mutations define predominant functions of the human T-cell leukemia/lymphoma virus p12 protein

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