

## Variability at Human Immunodeficiency Virus Type 1 Subtype C Protease Cleavage Sites: an Indication of Viral Fitness?

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**Naturally occurring polymorphisms in the protease of human immunodeficiency virus type 1 (HIV-1) subtype C would be expected to lead to adaptive (compensatory) changes in protease cleavage sites. To test this hypothesis, we examined the prevalences and patterns of cleavage site polymorphisms in the Gag, Gag-Pol, and Nef cleavage sites of C compared to those in non-C subtypes. Codon-based maximum-likelihood methods were used to assess the natural selection and evolutionary history of individual cleavage sites. Seven cleavage sites (p17/p24, p24/p2, NC/p1, NC/TFP, PR/RT, RT/p66, and p66/IN) were well conserved over time and in all HIV-1 subtypes. One site (p1/p6<sup>gag</sup>) exhibited moderate variation, and four sites (p2/NC, TFP/p6<sup>pol</sup>, p6<sup>pol</sup>/PR, and Nef) were highly variable, both within and between subtypes. Three of the variable sites are known to be major determinants of polyprotein processing and virion production. P2/NC controls the rate and order of cleavage, p6<sup>gag</sup> is an important phosphoprotein required for virion release, and TFP/p6<sup>pol</sup>, a novel cleavage site in the transframe domain, influences the specificity of Gag-Pol processing and the activation of protease. Overall, 58.3% of the 12 HIV-1 cleavage sites were significantly more diverse in C than in B viruses. When analyzed as a single concatenated fragment of 360 bp, 96.0% of group M cleavage site sequences fell into subtype-specific phylogenetic clusters, suggesting that they coevolved with the virus. Natural variation at C cleavage sites may play an important role, not only in regulation of the viral cycle but also in disease progression and response to therapy.**

One of the most dramatic changes in the human immunodeficiency virus type 1 (HIV-1)-AIDS epidemic has been the rapid emergence and devastating spread of subtype C viruses (16; <http://www.unaids.org/epidemicupdatedec01/report/index.html>). HIV-1 C now accounts for >56% of all circulating viruses and is the most commonly transmitted subtype worldwide (9). Subtype C predominates in southern Africa (11, 20, 40) and India (29) and is increasing in frequency in China (10, 28) and Brazil (4, 10a, 32). The disproportionate increase in C viruses relative to other HIV-1 strains suggests that subtype C may be more easily transmitted or that it has a higher level of “fitness” at the population level. One possible explanation is that founder effects relating to the ongoing introduction of subtype C into new population groups with different host factors, or different social and sexual practices, may be responsible for the rapid spread. However, founder and host effects cannot account for the fact that C viruses are overtaking pre-existing virus subtypes in several different geographical regions, including Yunnan Province in China and the southern region of Brazil (4, 10, 28, 32). It is increasingly evident that additional (nonhost) viral factors are also contributing to the rapid spread of HIV-1 C.

Viral studies indicate that subtype C has distinct genetic and phenotypic properties that differentiate it from other HIV-1 subtypes. Various studies have postulated that an extra NF- $\kappa$ B binding site in the long terminal repeat (28), a prematurely truncated Rev protein (10, 28), or a 5-amino-acid insertion in Vpu (18) may influence viral gene expression, altering the transmissibility and pathogenesis of C viruses (34). Factors related to viral entry and pathogenesis, such as the CCR5 and non-syncytium-inducing properties of C isolates (2, 21, 25), may also contribute to the increased spread of C viruses. One area of research that is receiving consideration is the possibility that C viruses have a more active, catalytically efficient protease (41).

The C protease is highly conserved at the amino acid level and has a distinct signature sequence that differentiates it from those of subtypes A, B, and D (11, 41). A subset of these signature residues, present in the hinge (M36I/R41K/H69K) and  $\alpha$ -helix (I89M) of C (and A) proteases, has been linked to increased catalytic activity (41). Another signature pattern, identified in >80% of C viruses from South Africa, is T12S/T15V/L19I/I93L (11). The 12S, 15V, and 19I residues of this motif are located near the N terminus of the protease in an extended  $\beta$ -chain. The 93L polymorphism is located within a hydrogen-bonded turn immediately upstream from the protease-reverse transcriptase (RT) cleavage site, in close proximity to 12S/15V/19I and the dimerization domain (22, 30, 33, 35). All of these polymorphisms lie outside the catalytic site of the protease, in regions that would be expected to alter the

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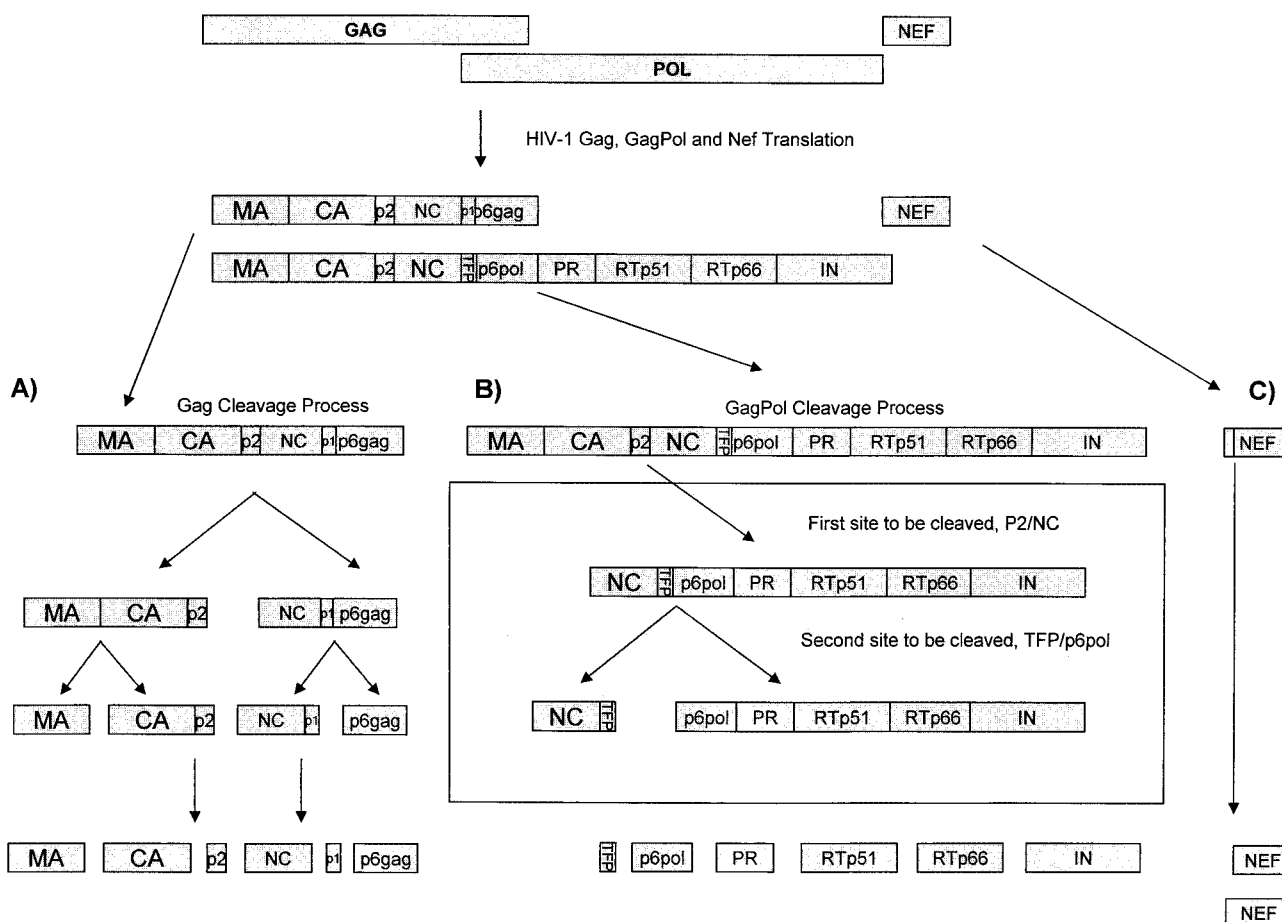


FIG. 1. Schematic of the Gag and Gag-Pol processing sites showing the 12 individual protease cleavage sites: 5 cleavage sites in Gag (p17/p24, p24/p2, p2/NC, p7/p1, and p1/p6<sup>gag</sup>), 6 cleavage sites in Gag-Pol (NC/TFP, TFP/p6<sup>pol</sup>, p6<sup>pol</sup>/PR, PR/RT, RT/p66, and p66/IN), and a single site in Nef. The frequency of amino acid substitution at each of these cleavage sites is shown in Fig. 2 and Tables 3 and 4.

enzyme's activity toward its natural cleavage sites, leading to adaptive (compensatory) changes in the cleavage site itself.

Since protease inhibitors (PIs) are currently the most active antiretroviral drugs used for the treatment of HIV-1 (44), it is important to collect information, not only on the C protease but also on its drug responsiveness, substrate-inhibitor specificities, and cleavage site characteristics. This information is needed to design PIs that are maximally effective against C viruses and to obtain new insights into the mechanism of drug resistance. Studies have shown that resistance mutations in the B protease are associated with impaired proteolytic processing, decreased enzymatic activity, and a failure to produce mature infectious virions (6, 7, 44). Compensatory cleavage site mutations can partially compensate for these defects (47). In this report, we describe the natural variability of subtype C cleavage sites in viruses from Africa, India, and Brazil and compare the results to cleavage site patterns in representative B and group M viruses.

**MATERIALS AND METHODS**

**Cleavage site characteristics.** The HIV-1 protease is a small, 99-amino-acid aspartic enzyme that mediates the cleavage of Gag, Gag-Pol, and Nef precursor polyproteins. These reactions occur late in the viral life cycle, during virion assembly and maturation at the cell surface. The process is highly specific,

temporally regulated, and essential for the production of infectious viral particles (13–15, 33). As shown in Fig. 1, the main structural proteins are formed by cleavage of the Pr55<sup>gag</sup> polyprotein into matrix (MA; p17), capsid (CA; p24), nucleocapsid (NC; p7), p6<sup>gag</sup>, and two spacer peptides, p2 and p1. The viral enzymes are formed by cleavage of Pr160<sup>gag-pol</sup>, a fusion protein derived by ribosomal frame shifting (13). Although Pr160<sup>gag-pol</sup> also contains p17, p24, and p2, its C-terminal cleavage products are NC, a transframe protein (TFP), p6<sup>pol</sup>, protease (PR), reverse transcriptase (RTp51), RNase H (RTp66), and integrase (IN) (12, 35). In total, 12 proteolytic reactions are required to generate a mature infectious virion. Each reaction occurs at a unique cleavage site that differs in amino acid composition (3). Some cleavage sites contain phosphorylated Ser/Thr or Tyr residues that alter the sites' susceptibilities to cleavage (39). P6<sup>gag</sup>, the major phosphoprotein of HIV-1, plays an essential role in the release of virus from the membranes of infected cells (19).

**Sequence data and construction of cleavage site fragments.** A total of 84 full-length nucleotide sequences were selected for analysis. These sequences included two C isolates from South Africa, TV001 and TV002 (46), in addition to another 25 subtype C, 30 subtype B, and 27 representative group M reference sequences (including A [n = 3], B [n = 4], C [n = 5], D [n = 3], F1 and F2 [n = 4], G [n = 2], H [n = 2], J [n = 2], and K [n = 2] subtypes) extracted from the Los Alamos database (Table 1) (16). Sequences were selected based on the patient being treatment naive. Since the prevalence of drug resistance in untreated patients has been reported to range from 1 to 11% (17), sequences were also screened and excluded from the study if they were found to contain primary resistance mutations. The majority of the sequences were obtained by direct DNA PCR amplification and cloning of peripheral blood mononuclear cells. Nucleotide sequences were aligned by CLUSTAL W (37) and manually edited with the codon alignment of the Genetic Data Environment (GDE version 2.2) program (31). Calculation of the pairwise distance matrix, phylogenetic infer-

TABLE 1. Sequences used for the analysis of protease cleavage sites<sup>a</sup>

Group or subtype	Data set	Accession no.	Group or subtype	Data set	Accession no.
M	A1.KE.93.Q23-17	AF004885	C	B.US.87.BC	L02317
	A1.UG.85.U455	M62320		B.TW.-.TWCYS	AF086817
	A1.UG.92.92UG037	U51190		B.US.86.JRFL	U63632
	B.FR.83.HXB2	K03455		B.AU.86.MBC200	AF042100
	B.US.83.RF	M17451		B.DE.86.HAN	U43141
	B.US.86.JRFL	U63632		B.US.83.RF	M17451
	B.US.90.WEAU160	U21135		B.US.97.ARES2	AB078005
	C.BR.92.92BR025	U52953		B.AU.96.MBCC98	AF042104
	C.BW.96.96BW0502	AF110967		B.KO.97.WK	AF224507
	C.ET.86.ETH2220	U46016		B.AR.98.ARCH054	AY037268
	D.CD.83.ELI	K03454		B.AR.00.ARMS008	AY037269
	D.CD.83.NDK	M27323		B.AR.99.ARMA132	AY037282
	D.CD.84.84ZR085	U88822		B.DE.86.D31	U43096
	F1.BE.93.VI850	AF077336		B.US.88.WR27	AF286365
	F1.BR.93.93BR020.1	AF005494		B.US.86.YU2	M93258
	F2.CM.95.MP255	AJ249236			
	F2.CM.95.MP257	AJ249237		C.ET.86.ETH2220	U46016
	C.ZA.98.TV001c8.5	AY16222		C.BR.92.BR025	U52953
	C.ZA.98.TV002c12	AY16224		C.IN.93.93IN999	AF067154
	G.BE.96.DRCBL	AF084936		C.IN.93.93IN904	AF067157
	G.FI.93.HH8793-12.1	AF061641		C.IN.93.93IN905	AF067158
	H.BE.93.VI991	AF190127		C.IN.94.94IN11246	AF067159
	H.BE.93.VI997	AF190128		C.IN.94.94IN476	AF286223
	J.SE.93.SE7887	AF082394		C.IN.95.95IN21068	AF067155
	J.SE.94.SE7022	AF082395		C.BW.96.96BW01B03	AF110959
	K.CD.97.EQTB11C	AJ249235		C.BW.96.96BW0402	AF110962
	K.CM.96.MP535	AJ249239		C.BW.96.96BW0502	AF110967
				C.BW.96.96BW1104	AF110969
				C.BW.96.96BW1210	AF110972
				C.BW.96.96BW17B03	AF110980
				C.BW.96.96BW1626	AF110978
				C.BW.96.MJ4	AF321523
				C.ZM.96.ZM651	AF286224
		C.ZM.96.ZM751	AF286225		
		C.ZA.97.ZA012	AF286227		
		C.BR.98.98BR004	AF286228		
		C.IL.98.98IS002	AF286233		
		C.IN.98.98IN012	AF286231		
		C.IN.98.98IN022	AF286232		
		C.TZ.98.98TZ013	AF286234		
		C.TZ.98.98TZ017	AF286235		
		C.ZA.98.TV001c8.5	AY16222		
		C.ZA.98.TV002c12	AY16224		
B	B.US.84.NY5CG	M38431			
	B.US.-.AD8	AF004394			
	B.CN.-.RL42	U71182			
	B.US.90.WEAU160	U21135			
	B.US.-.P896	U39362			
	B.GA.-.OYI	M26727			
	B.US.-.DH123	AF069140			
	B.GB.-.CAM1	D10112			
	B.NL.86.3202A21	U34604			
	B.AU.87.MBC925	AF042101			
	B.ES.89.89SP061	AJ006287			
	B.FR.83.HXB2	K03455			
	B.US.83.SF2	K02007			
	B.US.90.WCIPR9018	U69591			
	B.US.84.MNCG	M17449			

<sup>a</sup> Identification information for the sequences is in the following format: subtype.country.isolation year.common name. The country is represented by the two-letter country code using the international naming convention from ISO 3166. The accession numbers are from GenBank.

ence, and tree construction were performed on a dual-processor Linux computer by using the PAUP version 4.0b2a program (Sinauer Associates, Sunderland, Mass.) and a GDE for Linux HIV-1 interface (6a). Thirty-base-pair segments, consisting of 15 nucleotides (5 amino acids) on each side of the 12 cleavage sites, were extracted and concatenated into a 360-bp nucleotide sequence.

**Reconstruction of ancestral cleavage site sequences.** To examine the evolutionary histories of individual cleavage sites, Phylogenetic Analysis under Maximum Likelihood (PAML) software (26) was used to identify amino acid and nucleotide substitutions along each branch of the tree. Branch lengths were estimated using a nucleotide substitution model; amino acid sequences were deduced from the reconstructed nucleotide triplets. The analyses involved the use of maximum-likelihood methods and a time-reversible model which assume different substitution rates, base frequencies, and transition/transversion rate ratios ( $\kappa$ ) (42, 43). Using this approach, we were able to reconstruct the ancestral sequences and internal nodes for each of the 12 protease cleavage sites in the B, C, and M group data sets. The number of proximal ancestors for each data set was  $n - 1$ , which translated into 29 ancestral sequences for subtype B, 26 sequences for subtype C, and 26 sequences for the group M viruses. The most recent common ancestor (MRCA) nucleotide sequence for each virus in the three data sets was saved and translated into its corresponding amino acids.

**Diversity and cleavage site polymorphisms.** Nucleotide diversity at cleavage sites was measured using a Kimura 2- $\alpha$  parameter model with a distance matrix implemented in the MEGA program version 2.0 (Arizona State University, Tempe). Amino acid diversity was measured using a Poisson distribution method implemented in the same MEGA package. *P* values for diversity measurements were calculated by applying the *t* test to the distance matrix of each data set. To determine whether the sequences had evolved over time, amino acid profiles for individual Gag, Gag-Pol, and Nef cleavage sites were compared to the inferred MRCA for that site.

**Assessment of positive selection pressure.** Nucleotide sequences were also analyzed with Codeml, a program from the PAML software package (26). The likelihood ratio test (1) and recently developed codon-based models (42, 43) were used to assess natural selection and adaptive evolution at the amino acid level. These selection models use maximum-likelihood scores to account for variation in the  $d_n/d_s$  (nonsynonymous/synonymous) ratio ( $\omega$ ) at individual codons along the length of the sequence. High rates of synonymous mutation are indicative of conservation and a strict requirement for biological function, while high rates of nonsynonymous substitution are indicative of adaptive change in response to host selection pressure. An individual amino acid was considered to be positively selected if the  $d_n/d_s$  ratio was significantly greater than 1.0.

TABLE 2. Inter- and intrasubtype diversity at 12 cleavage sites, expressed as amino acid distances between sequences

Protease site <sup>a</sup>	Mean % distance			P value		
	Subtype B	Subtype C	M group	B vs C	B vs M	C vs M
p17/p24	2.31	5.15	2.23	<0.0001	0.806	<0.0001
p24/p2	2.41	3.07	2.67	0.077	0.454	0.267
p2/NC	18.66	42.42	39.22	<0.0001	<0.0001	0.062
NC/p1	0.40	5.42	5.22	<0.0001	<0.0001	0.7244
p1/p6 <sup>gag</sup>	8.93	9.81	14.47	0.318	<0.0001	<0.0001
NC/TFP	5.30	3.90	6.30	0.0012	0.0215	<0.0001
TFP/p6 <sup>pol</sup>	16.69	7.60	24.70	<0.0001	<0.0001	<0.0001
p6 <sup>pol</sup> /PR	17.66	16.28	16.55	0.15	0.019	0.365
PR/RT	0.70	1.56	2.93	<0.0001	<0.0001	<0.0001
RT/p66	0.81	7.49	6.03	<0.0001	<0.0001	0.004
p66/IN	2.44	2.22	2.24	0.524	0.61	0.961
Nef	11.89	24.52	25.68	<0.0001	<0.0001	0.33
Gag (501 aa)	6.99	9.75	15.72	<0.001	<0.001	<0.001
Pol (1,004 aa)	6.02	5.83	5.89	0.376	0.419	0.875
Nef (207 aa)	14.66	16.50	18.50	0.07	<0.001	<0.001
12 protease cleavage sites (360 bp; 120 aa)	4.80	10.10	12.10	<0.001	<0.001	<0.001

<sup>a</sup> aa, amino acids.

RESULTS

**Viral characteristics.** No primary RT- or PI-resistant mutations were detected among the 84 full-length sequences selected for study. Although attempts were made to include only sequences amplified directly from HIV-1 proviral DNA and to match these sequences based on duration of infection, plasma viral load, and CD4 count, this proved difficult. A surprisingly small amount of full-length sequence data was available from treatment-naïve patients infected with subtype B, and when available, it was often poorly annotated. Despite these limitations, the frequency and pattern of naturally occurring polymorphisms observed in this study were remarkably similar to those reported for a control group of subtype B infections treated with nucleoside reverse transcriptase inhibitors but not with PIs or nonnucleoside reverse transcriptase inhibitors (5). Most of the non-C sequences came from regions of the world where treatment is not yet readily available.

**Genetic diversity and patterns of amino acid variability at individual cleavage sites.** Variation at the 12 cleavage sites of subtypes B and C and group M is shown in Table 2. Seven (58.3%) sites (p17/p24, p24/p2, NC/p1, NC/TFP, PR/RT, RT/p66, and p66/IN) were found to be relatively well conserved, both over time and between subtypes, with a mean intrasubtype distance ranging from 0.40 ± 0.20% to 7.49 ± 5.62%. The remaining five (41.7%) sites exhibited moderate (p1/p6<sup>gag</sup>) to extensive (p2/NC, TFP/p6<sup>pol</sup>, p6<sup>pol</sup>/PR, and Nef) variation, with mean intrasubtype diversities reaching levels as high as 42.42 ± 15.16%. For the purposes of this study, we have referred to these three patterns as conserved, moderately variable, and variable. Polymorphisms were more common among C than B viruses (*P* < 0.0001). Overall, six cleavage sites (p17/p24, p2/NC, NC/p1, PR/RT, RT/p66, and Nef) had significantly higher levels of diversity among C viruses (*P* < 0.0001); five sites (p24/p2, p1/p6<sup>gag</sup>, NC/TFP, p6<sup>pol</sup>/PR, and p66/IN) had similar levels of diversity in both subtypes, and one site, TFP/p6<sup>pol</sup>, was highly variable among B and group M viruses (mean

distances, 16.7 and 24.7%, respectively) but relatively conserved in C viruses (mean distance, 7.6%) (*P* < 0.0001).

Compared to the M data set, the level of polymorphism at C cleavage sites was as wide ranging as that observed for the entire M group, a data set containing nine different HIV-1 subtypes. One cleavage site, p17/p24, was significantly more diverse in subtype C (mean divergence, 5.15%) than in group M (mean divergence, 2.23%) and subtype B (mean divergence, 2.31%) viruses (<0.0001 for both comparisons). Seven sites (p24/p2, p2/NC, NC/p1, p6<sup>pol</sup>/PR, RT/p66, p66/IN, and Nef) exhibited similar levels of diversity in both data sets (*P* = 0.004 to 0.961). Only four group M cleavage sites (p1/p6<sup>gag</sup>, NC/TFP, TFP/p6<sup>pol</sup>, and PR/RT) had mean diversities that were significantly greater than that observed for subtype C (*P* < 0.0001). For >50% of the sites, the variability of B viruses was significantly lower than that observed for subtype C or group M (*P* < 0.0001), despite the fact that the B viruses covered a broader time frame.

The observed polymorphisms were not randomly distributed across the variable cleavage sites but were confined to specific amino acids, most of which were positively selected in C but not in B viruses (Fig. 2). The least variable residues were the P1 positions of p1/p6<sup>gag</sup>, TFP/p6<sup>pol</sup>, and Nef and the P1' positions of p6<sup>pol</sup>/PR and Nef. As shown, these positions flank the scissile bond, with P1 located immediately upstream and P1' located immediately downstream of the cleavage junction. The most variable positions were the P1, P3, P4, P5, P3', and P5' residues of p2/NC; the p3' residue of p1/p6<sup>gag</sup>; the P1' and P4' residues of TFP/p6<sup>pol</sup>; the P1 to P4 residues of p6<sup>pol</sup>/PR; and the P3 to P5 and P2' residues of Nef.

**Subtyping and phylogenetic-tree analysis.** When subjected to phylogenetic analysis (Fig. 3), the concatenated 360-bp fragments of the group M data set fell into eight subtype-specific clusters representing subtypes A, B and D, C, F, G, H, J, and K, with the cleavage sequences for subtypes B and D segregating together in the same subcluster. This pattern was sup-

p17/p24 (MA/CA) - | Conserved |

M MRCA	V	S	Q	N	Y	/	P	I	V	Q	N
M group	I(1)								L(1)		
	S(1)										
	D(1)										
B MRCA						/					
Subtype B	A(1)		R(1)								
C MRCA						/					
Subtype C	I(3)*		R(1)						A(1)		
	D(2)										

p24/p2 (CA/p2) - | Conserved |

M MRCA	K	A	R	V	L	/	A	E	A	M	S
M group				I(4)							
B MRCA						/					
Subtype B				I(2)							
C MRCA						/					
Subtype C		K(1)	I(2)				G(1)				

p2/NC (p2/p7) - | Variable |

M MRCA	S	T	A	I	M	/	M	Q	K	G	N
M group	N(11)*	A(10)*	N(7)*	V(4)	L(1)		L(1)	R(13)	S(13)		
	A(3)	S(3)	T(6)	A(1)			V(1)				
	H(2)	P(1)	S(2)	M(1)			I(1)				
	T(1)	V(1)	V(1)								
	P(1)										
	Q(1)										
	G(1)										
	V(1)										
B MRCA		A	T			/		R			
Subtype B	P(3)	N(2)	N(3)	V(3)			L(1)	K(5)	S(1)		
	T(1)	T(2)	A(3)	M(1)			I(1)	G(1)			
	A(1)	V(1)									
C MRCA	N		N			/			S		
Subtype C	S(13)*	A(6)*	S(2)*	V(2)	L(9)*		I(2)	R(16)	G(5)*		
	T(1)	N(4)	T(1)	V(2)	G(1)		V(1)		N(1)		
	M(1)										
	S(1)										

NC/p1 (p7/p1) - | Conserved |

M MRCA	E	R	Q	A	N	/	F	L	G	K	I
M group	G(1)									R(2)	F(1)
	K(1)										M(1)
											L(1)
B MRCA						/					
Subtype B	K(1)										
C MRCA						/					
Subtype C	G(1)	R(1)		D(1)						R(3)	L(1)

NC/TFP - | Conserved |

M MRCA	E	R	Q	A	N	/	F	L	R	E	N
M group	G(1)										D(6)
											T(1)
											V(1)
B MRCA						/					
Subtype B	K(1)										D(21)
C MRCA						/					
Subtype C	G(1)	R(1)		D(1)							T(1)
											D(1)

p1/p6<sup>SH</sup> - | Moderately Variable |

M MRCA	R	P	G	N	F	/	L	Q	S	R	P
M group					L(1)		P(5)	N(10)*	L(3)*		
							I(1)	K(1)	S(1)		
									T(1)		
B MRCA						/					
Subtype B		R(1)					P(4)	N(3)	L(3)*		
								R(1)	T(1)		
C MRCA						/					
Subtype C				L(1)				N(11)*	T(2)		
								G(1)	S(1)		
									L(1)		

TFP/p6<sup>SH</sup> - | Variable |

M MRCA	E	N	L	A	F	/	Q	Q	G	E	A
M group		D(6)		V(1)	S(1)		P(10)*	K(2)	R(5)	K(9)	
		T(1)					L(2)				
		V(1)									
B MRCA						/	P			K	
Subtype B		D(21)	M(1)	V(1)			L(8)	R(1)	R(2)	E(4)	
							Q(1)				
C MRCA						/	P				G
		D(1)					Q(1)	E(1)		K(5)*	
Subtype C											
		T(1)									

p6<sup>SH</sup>/protease - | Variable |

M MRCA	T	S	F	S	F	/	P	Q	I	T	C
M group	V(9)*	T(4)*	L(11)*	N(14)*	C(1)						
	P(5)	A(1)	S(3)	D(1)							
	G(3)	G(1)		C(1)							
	S(3)			I(1)							
	I(1)										
	D(1)										
	Q(1)										
B MRCA	V					/					L
Subtype B	I(3)		L(7)	N(7)	L(5)			R(1)	V(1)		
				D(2)	C(1)			H(1)			
C MRCA	G	T	L	N		/					L
Subtype C		S(2)*	F(8)*	V(1)*	L(3)*		L(1)			A(1)	
		N(2)		S(1)	C(3)						
		A(1)									

protease/RT - | Conserved |

M MRCA	C	T	L	N	F	/	P	I	S	P	I
M group	R(1)			H(1)	L(1)			V(1)			
B MRCA						/					
Subtype B									C(1)		
C MRCA						/					
Subtype C	R(1)			D(1)							

RT/p66 - | Conserved |

M MRCA	G	A	E	T	F	/	Y	V	D	G	A
M group		V(4)	D(1)		Y(4)						
B MRCA						/					
Subtype B	E(1)										
C MRCA						/					
Subtype C	E(1)	V(10)*			Y(2)*						

p66/IN - | Conserved |

M MRCA	I	R	K	V	L	/	F	L	D	G	I
M group	V(1)		M(1)								
B MRCA						/					
Subtype B	V(1)		R(1)	I(1)					N(1)		
C MRCA						/					
Subtype C			R(2)								
			E(1)								

NEF - | Variable |

M MRCA	P	D	C	A	W	/	L	E	A	Q	E
M group	A(14)*	A(4)*	L(4)*	V(2)			V(1)	K(1)			T(2)
	D(2)	S(3)	S(1)	T(1)				Q(1)			Q(1)
	E(1)	T(1)	V(1)	Y(1)							
B MRCA	A					/					
Subtype B	P(1)	A(7)	I(1)	V(1)			Q(1)	K(1)		H(1)	
		N(1)		T(1)							
C MRCA	A					/					
Subtype C	P(3)*	A(7)*	L(1)*		R(1)			R(5)*	P(2)*		Q(1)
	T(1)	H(1)	G(1)					Q(4)	E(1)		K(1)
	E(1)	E(1)						K(2)	T(1)		



TABLE 3. Relationship between cleavage site signature patterns and common ancestors<sup>a</sup>

Group or subtype	No. (%) of sequence with same MRCA							
	p17/p24 (MA/CA) VSONY/PIVQN	p24/p2 (CA/p2) KARVL/AEAMS	NC/p1 (p7/p1) ERQAN/FLGKI	NC/TFP ERQAN/FLREN	p1/p6 <sup>sig</sup> RPGNF/LQSRP	PR/RT CTLNF/PISPI	RT/p66 GAETF/YVDGA	p66/IN IRKVL/FLDGI
M	23 (85.2)	23 (85.2)	21 (77.7)	17 (63.0)	13 (48.1)	22 (81.5)	19 (70.4)	25 (92.6)
B	28 (93.3)	28 (93.3)	29 (96.7)	8 (26.7)	23 (76.7)	29 (96.7)	29 (96.7)	26 (86.7)
C	21 (77.7)	23 (85.2)	21 (77.7)	22 (81.5)	15 (55.6)	25 (92.6)	14 (51.9)	24 (88.9)

<sup>a</sup> Conserved and moderately variable cleavage sites (group M MRCA is identical to subtype B and C MRCAs).

regulation of polyprotein processing and protease activation, p2/NC, TFP/p6<sup>pol</sup>, and p6<sup>pol</sup>/PR, were highly variable and differed significantly from both the subtype-specific and group M MRCAs. With the exception of the TFP/p6<sup>pol</sup> site, which was more variable in subtype B, divergence from the MRCA was greatest for C and M viruses. None of the p2/NC and p6<sup>pol</sup>/PR sequences in the group M data set was identical to the M MRCA signatures for p2/NC and p6<sup>pol</sup>/PR, STAIM/MQKGN and TSFSF/PQITC, respectively.

#### Positive selection of amino acids at protease cleavage sites.

The concatenated 360-bp cleavage site fragments were next compared internally to assess the mutational behavior of P1-P5 and P1'-P5' sites in the absence of drug therapy. As described in Materials and Methods, the analyses were performed using codon-based maximum-likelihood methods that test for the variation in selection pressure ( $d_n/d_s$ ) at individual amino acids along the length of the sequence. Application of the likelihood ratio test indicated that the best-fit model for subtype C and group M sequences was the positive-selection (discrete) model ( $\chi^2 = 54.47$ ;  $P < 0.0001$  and  $62.34$  and  $P < 0.0001$ , respectively), while for B viruses the neutral model performed as well as the positive model ( $\chi^2 = 1.2$ ;  $P > 0.05$ ). Overall, when analyzed as a single concatenated fragment, all three datasets were found to be under negative (purifying) selection, with  $d_n/d_s$  ( $\omega$ ) values ranging from 0.26 to 0.31 for all sites. Only 20 (16.6%) of the 120 amino acids within the 12 cleavage sites of subtype C were found to be under positive selection, with a  $\omega$  value of 1.7. Group M and subtype B cleavage sites had fewer positively selected amino acids: 11.1 and 6.7%, respectively (Fig. 2).

## DISCUSSION

The presence of polymorphisms in the protease of subtype C would be expected to result in adaptive (compensatory) changes in the natural cleavage sites that are recognized and cleaved by the C enzyme. To test this hypothesis, we examined the prevalences and patterns of cleavage site mutations in the Gag, Gag-Pol, and Nef proteins of subtype C compared to those of non-C viruses. Using phylogenetic and ancestral re-

construction methods, we identified two groups of natural cleavage site sequences. The first group consisted of eight sequences, seven of which were highly conserved in all HIV-1 subtypes, and an eighth sequence which was moderately variable. Mutations at these sites were uncommon and, when present, were observed at relatively equivalent frequencies among different subtypes. These cleavage sites, which defined the main structural (MA, CA, and NC) and enzymatic (RT, RNase H, and integrase) proteins of HIV-1, were under strong negative (purifying) selection pressure, had a common ancestor, and showed little genetic evolution over time. The second group consisted of five cleavage sites that were under positive (diversifying) selection pressure, exhibited extensive inter- and intrasubtype variability, and showed little (or no) resemblance to the common ancestor of group M or to the subtype-specific MRCA.

Our data showing that the majority (58.3%) of cleavage sites are highly conserved in all subtypes were not unexpected, given the narrow specificity of the HIV-1 protease relative to cellular proteases, such as pepsin (27). The strong purifying selection pressure exerted on these sites is presumably a reflection of the need to maintain the spatial configuration of the enzyme-substrate complex, conserve the hydrophobic nature of the scissile bond, and retain the biological activity of functionally important sites, such as the P1' proline of p17/p24 and the P1 and P1'-P5' residues of NC/p1. Cleavage of the P17/p24 site is known to play an important role in virion maturation, while processing of NC/p1 is required for ribosomal frame shifting and Gag-Pol expression (8, 13–15, 24). In B viruses, cleavage of p17/24, p24/p2, and NC/p1 has been shown to be suboptimal, with the NC/p1 site being rate limiting (23). It has been suggested that the slow, regulated cleavage of these structural proteins may represent a common strategy to ensure that the assembled virions have the full complement of proteins needed to bud from the cell surface, bind to a new cell, and initiate a new round of viral replication (35).

The carboxyl terminus of NC is particularly interesting. Unlike other cleavage sites, which carry an aromatic amino acid at P1 and either a leucine or proline residue at P1', the C termini

TABLE 4. Relationship between cleavage site signature patterns and common ancestors<sup>a</sup>

Group or subtype	p2/NC (p2/p7)		TFP/p6 <sup>pol</sup>		p6 <sup>pol</sup> /PR		NEF	
	Sequence	No. (%) same	Sequence	No. (%) same	Sequence	No. (%) same	Sequence	No. (%) same
Group M	STAIM/MQKGN	0 (0)	ENLAF/QQGEA	6 (22.2)	TSFSF/PQITC	0 (0)	PDCAW/LEAQE	5 (18.5)
Subtype B	SATIM/MQRGN	13 (43.3)	ENLAF/PQGKA	5 (16.7)	VSFSE/PQITL	13 (43.3)	ADCAW/LEAQE	18 (60.0)
Subtype C	NTNIM/MQKSN	3 (11.1)	ENLAF/PQGEA	21 (77.8)	GTLNF/PQITL	10 (37.0)	ADCAW/LEAQE	6 (22.2)

<sup>a</sup> Variable cleavage sites (subtype B and C MRCAs differ from the group M MRCA).

of NC/p1 and NC/TFP carry an asparagine (N) residue at P1 opposite a phenylalanine (F) residue at P1' (5, 22–24). In this study of 84 untreated patients, no mutations were detected at the P2 or P1'-P3' positions of NC/p1 or NC/TFP and only a single N→C mutation was detected at P1. Taken together, these findings underscore the unique nature and limited mutability of the NC/p1 and NC/TFP cleavage junctions. Although these sites were strongly conserved in natural infection, recent studies have shown that an A→V substitution at the P2 positions of NC/p1 and NC/TFP is a common adaptive change, occurring in 29% of PI-resistant patients taking indinavir, saquinavir, and/or ritonavir for the treatment of subtype B (5, 7, 44, 45). This valine substitution is frequently associated with an M46 I or L mutation (and possibly a V82 mutation) in the protease and leads to altered polyprocessing and improved growth of protease-mutated viruses. Whether similar “second-locus” mutations will be observed during the treatment of non-B subtypes remains to be established. The identification of common patterns may facilitate the development of broad-based inhibitors with increased specificity and improved binding to the mutated protease. These secondary inhibitors might preempt (or delay) the emergence of resistance.

Our analyses also revealed important differences among HIV-1 subtypes. Particularly intriguing was the identification of five cleavage sites that exhibited extensive variability across all subtypes, with C viruses being significantly more variable than subtype B. Variation was restricted to a few specific amino acids, most of which were positively selected in C but not in B viruses. In contrast to conserved sites, variable cleavage sites tended to be those with regulatory rather than structural or enzymatic functions. At least four of the variable sites (p2/NC, p1/p6gag, TFP/p6pol, and p6pol/PR) are known to play major roles in the regulation of polyprotein processing and, in the case of TFP/p6<sup>pol</sup>, in the activation of the protease enzyme (22–24, 27, 30). Studies of subtype B have shown that p2/NC is the initial and most rapidly processed cleavage site, controlling both the rate and the order of Gag and Gag-Pol polyprocessing (30). Our results indicate that p2/NC is by far the most variable cleavage site, with intrasubtype diversity ranging from 18.7% in subtype B to levels of 42.4% in subtype C.

The p1/p6<sup>gag</sup> cleavage product, p6<sup>gag</sup>, is a major phosphoprotein that is critical to the release of mature, infectious virions (19). Although not well studied, phosphorylation of Gag and Gag-Pol sequences has been shown to alter susceptibility to cleavage, attenuating or even preventing the proteolytic process (38). The TFP/p6<sup>pol</sup> cleavage site, defining the N terminus of p6<sup>pol</sup>, was the only site to have a significantly higher level of diversity among B than among C viruses. TFP/p6<sup>pol</sup> is a novel cleavage site located 8 amino acids downstream from NC in the TFP domain of Gag-Pol (22, 35). Although TFP/p6<sup>pol</sup> lies outside (and upstream) of the protease, the EDL tripeptide of this cleavage site (ENL in the case of C viruses) has been postulated to have a major influence on protease activation and on the timing and specificity of Gag-Pol cleavage, delaying the release of the protease until after the viral particle has budded from the cell membrane. Such a mechanism may protect the cell from the cytotoxic effects of proteolysis (36, 38). The observed subtype variation in the cleavage sites controlling the initiation and rate of Gag and

Gag-Pol processing (p2/NC) and the activation of protease (TFP/p6<sup>gag</sup>) suggests that there may be important differences in the way that B and C viruses regulate polyprocessing and virion assembly. Subtle cleavage site differences could, over time, have a major differential impact on the pathogenesis of HIV-1 subtypes and on response to therapy. Early treatment studies suggest that C viruses give an excellent initial response to highly active antiretroviral therapy but that the duration of the response may be less than that reported for B viruses.

In summary, our results point to important inter- and intra-subtype differences in protease cleavage sites, especially in the p2/NC, TFP/p6pol, and p6pol/PR sites. The main limitations of our study relate to the cross-sectional nature of the data sets and the limited availability of well-matched pretreatment controls for use in the B data set. Despite these limitations, the potential impact of our findings on HIV-1 disease progression and response to therapy warrants further investigation, both at the patient level and in vitro using site-directed mutagenesis. The separate monophyletic clustering of B and C cleavage sites suggests that cleavage sites have evolved in a subtype-specific manner. The divergence between ancestral and contemporary sequences in the C data set and the location of an ancestral node distal to the group M MRCA suggest that variation in C cleavage sites began early, prior to the diversification of HIV-1 subtypes. A more detailed investigation of C cleavage sites, both over time and in response to therapy, is in progress. The present study forms the baseline for these ongoing studies.

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