## Identification and Characterization of Previously Described Epitopes in HIV-1 Subtypes B, C, F and BF in Brazil

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Genetic analysis of HIV-1 is essential to improve treatment strategies and select epitopes for vaccine programs. The objective of this study was to determine whether known CD4<sup>+</sup> and CD8<sup>+</sup> epitopes were present in Brazilian HIV-1 strains. We used previously described CD8<sup>+</sup> and CD4<sup>+</sup> epitopes from the Los Alamos laboratory to search for these epitopes in the Brazilian sequences using the HIVbase program and we compared the frequency results with the analyses using physical-chemical profile tools from Network Protein Sequence Analysis (NPSA), and the SYFPEITHI program. Furthermore, this analysis was carried out with the Prosite tool using the GeneDoc program and ds/dn analyses using the Synonymous Nonsynonymous Analysis Program (SNAP). The HIVbase epitope mapping demonstrated that 30 CD8<sup>+</sup> and 6 CD4<sup>+</sup> epitopes were present in the Brazilian sequences at a high frequency. Only two of these epitopes were heavily glycosylated. Interestingly, ds/dn analyses showed evidence of purifying selective pressure. These types of analyses could be useful for the assessment of possible vaccine efficiency in populations. Key-Words: HIV-1, *env*, vaccine, epitope.

HIV-1, identified as the etiological agent of AIDS [1], contributes to the development of immunodeficiency. Its biological complexity and high mutation rate have made the design of a vaccine to control the pandemic difficult. CD8<sup>+</sup> T lymphocyte responses are largely responsible for controlling viral replication during both acute and chronic infection [2-4]. The antibody responses appears much later and select a mutant virus [5]. Mutations in CTL epitopes and sites recognized by antibodies and acquisition of glycosylation sites are escape mechanisms that allow the virus to replicate and infect more cells [5-7].

The mapping of epitopes in diverse virus proteins and the identification of possible modifications can provide useful information to aid vaccine development. For example, mutations in the TW10 and SL9 Gag epitopes result in a fitness cost to the virus [7], and this kind of information is very useful for the vaccine development process. The use of bioinformatics software can identify escaping epitopes. Bioinformatics programs have been developed for diverse areas including protein analysis, physical-chemical characteristics, MHC binding databases, and posttranslational modification. The use of these programs to study the molecular epidemiology and genetic variation of the HIV-1 epidemic in Brazil may provide important information

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about the epidemic in this country. This information could be important in the design of an effective vaccine as well as for antiretroviral treatment. The objective of this study was to characterize genetic variation in previously identified epitopes in the Brazilian HIV-1 *env* sequences.

#### **Materials and Methods**

Sequences of all of the Brazilian HIV-1 strains (3,813) were collected from GenBank and added to the HIVbase Database [8]. Epitope mapping analyses were performed as described in the HIV Immunology and HIV/SIV Los Alamos Vaccine Databases [9]. We selected and analyzed only the env region, consisting of 2,644 sequences from gp120: C1V1=51; C2=59; C3=200; V3loop=515; and gp41=50. The alignment of these sequences was carried out using the ClustalX [10] software. Genedoc [11] software was used to edit and translate the alignment, and the potential site analyses were performed using the Prosite [12] tool. For selective pressure analysis, we used the Synonymous Nonsynonymous Analysis Program (SNAP) [13] from Los Alamos. The proportion of synonymous substitutions per potential synonymous site and the proportion of nonsynonymous substitutions per potential nonsynonymous site were calculated using the Nei and Gojobori method [14]. Prediction of cellular epitopes was made with the SYFPEITHI online database [15]. Identification of Antibody epitopes and their physical-chemical characteristics were carried out using the Physico-chemical profiles program of the Network Protein Sequence Analysis (NPSA) [16-20].

#### Results

HIVbase epitope mapping showed that thirty CD8<sup>+</sup> (Table 1) and six CD4<sup>+</sup> (Table 2) epitopes had a high frequency and showed varying degrees of conservation in the Brazilian HIV sequence. However, the SYPETEITHI analysis was restricted to the following HLA alleles: HLA-A\*03, -A\*6801, -A\*2402 and -A\*0201. Two of the epitopes in this program had several

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# Table 1. Mapping of the frequency of the Los Alamos CD8 epitopes in the Brazilian HIV-1 sequences

HXB2 Location Protein region	% Similarity	Epitopes sequence	Frequency	HLA	HXB2 Location Protein region	% Simi <b>l</b> arity	Epitopes sequence	Frequency	HLA
gp120/C1	Total=72.5	33NLWVTVYYGV42	37/51	A02	gp120 /C1	Total=98.0	34LWVTVYYGV42	50/51	A*0201
subtype B	30	 К	3/10 4/10 3/10		subtype B	100		10/10	
subtype F	100	Q	10/10		subtype F	100		10/10	
subtype C	100		12/12		Subtype C	100		12/12	
B/F	63		12/19		B/F	94.7		18/19*	
recombinant		Q	2/19		recombinant				
		К	2/19						
		D	2/19*						
gp 120/C1	Total=100	36VTVYYGVPV44	51/51	A02	gp 120/C1	Total=92.1	36VTVYYGVPVWK46	47/51	A*6801
Subtype B	100		10/10		Subtype B	100		10/10	
Subtype F	100		10/10		Subtype F	100		10/10	
Subtype C	100		12/12		Subtype C	83.3		10/12	
							R	2/12	
B/F	100		19/19		B/F recombinant	89.5		17/19	
recombinant	T ( 1 00 4		47/54	4*0004	400/04	T ( 1 00 0	R NA(O) (D) AA((C)A	2/19	0.7
gp 120/01	lotal=92.1	37TVYYGVPVWK46	47/51	A*0301,	gp 120/C1	lotal=90.2	38VYYGVPVWKEA48	46/51	Cw7
Culatura D	100		10/10	A 6601,	Culatura D	100		10/10	
Subtype B	100		10/10		Subtype B Subtype E	100		0/10	
Subtype C	83.3		10/12		Subtype C	83.3		10/12	
oubtype o	00.0	R	2/12		oubtype o	00.0	B	2/12	
B/F	89.5		17/19		B/F recombinant	89.5		17/19	
recombinant		R	2/19				R	2/19	
gp 120/C1	Total=62.7	42VPVWKEATTT 51	32/51	B*5501,	gp 120/C1	Total=62.7	42VPVWKEATTTL 52	32/51	B*3501
				B55	•••				
Subtype B	80		8/10*		Subtype B	80		8/10*	
Subtype F	90		9/10*		Subtype F	90		9/10*	
Subtype C	0	K	10/12*		Subtype C	0	K	10/12*	
B/F	78.9		15/19		B/F recombinant	78.9		15/19	
recombinant		R N	2/19*		(00/0)	<b>T</b> ( ) 00 (	R.N.	2/19*	
gp 120/C1	Iotal=90.2	50TTLFCASDAK59	46/51	A3supertype	gp 120/C1	Iotal=92.1	51TLFCASDAK59	47/51	A3supertype
Subtype B	100	••••	10/10		Subtype B Subtype F	100	••••	10/10	
Subtype F	75		0/10		Subtype F	83.3		10/10	
Subtype C	75	R	9/12 2/12*		Subtype C	03.5	R	2/12	
B/F	89.5		17/19*		B/F recombinant	89.5		17/19*	
recombinant									
gp 120/C1	Total=90.2	108 SLWDQSL 116	46/51	A2.1	gp 120/C1	Total=94.1	109 SLWDQSLK 117	48/51	A11
Subtype B	100		10/10		Subtype B	100		10/10	
Subtype F	100		10/10		Subtype F	100		10/10	
Subtype C	83.3		10/12*		Subtype C	91.6		11/12*	
B/F	84.2		16/19		B/F recombinant	89.5		17/19*	
recombinant		V	2/19*		(00/0.0.//				
gp 120/C1V1	Iotal=94.1	110SLWDQSLKP 118	48/51	A03	gp 120/C1V1	Iotal=92.1	117KPCVKLTPLC 126	47/51	B/
Subtype B	100		10/10		Subtype B	100		10/10	
Subtype F	100		10/10		Subtype F	90	•••••	9/10	
B/F	91.0		17/10*		B/E recombinant	84.2		16/10*	
recombinant	05.5		17/15		D/1 100011011ant	04.2		10/13	
ap120/C2	Total=61.0	252KPVVSTQLLL261	36/59	B07.B08	ap120/V3loop	Total=54.9	296CTRPNNNTRK305	283/515	A03:A02
Subtype B	28.6		4/14		Subtype B	47.7		187/392	
21 · -		R	10/14		<i>,</i> , –	••	G	31/392	
Subtype F	100		11/11		Subtype F	84.4		38/45	
							Y	2/45	
Subtype C	50		7/14		Subtype C	55.2		16/29	
			3/14				E	8/29	
D/F	70	M	2/14*		D/F 11 1	05 7	G	2/29	
B/F	70		14/20		B/F recombinant	85.7		42/49	
recombinant	Total=60 F		121/200	D1616	an120/02	Total=60.0		2/49	120
gp120/03	10181-00.5	3755FINCGGEFF383	121/200	B15-B63	gp120/C3	101al-00.0	3755FINCGGEFF 1 384	120/200	AZ9
Subtype B	68.4		117/153	010,000	Subtype B	68.4		116/153	
Oubtype b	00.4	т	24/153		Oubtype D	00.4	т	23/153	
		А	4/153				Α	4/153*	
		R	2/153*						
Subtype F	0	R	10/14		Subtype F	0	R.	10/14	
		Ν. Μ.	4/14				ΝΜ	4/14	
Subtype C	0	R	7/9*		Subtype C	0	R	7/9*	
B/F	16.6	· · · · <u>·</u> · · · ·	4/24		B/F recombinant	16.6	· · · · <u>·</u> · · · · ·	4/24	
recombinant	T-1-1 70 5	K.	18/24*			T-+-1 70 5	K	18/24*	400
gp120/C3	Iotal=73.5	376ENCGGEFF383	147/200	Cw4	gp120/C3	lotal=/2.5	376FNUGGEFFY384	145/200	A29

#### Table 1. (continued)

							1		
Subtype B	89.5		141/153		Subtype B	88.6		139/153	
		А	4/153				А	4/153*	
		P	2/152*						
		ĸ	2/100						
Subtype F	0	K	10/14		Subtype F	0	K.	10/14	
		M	4/14				M	4/14	
Subtype C	0	R	7/9*		Subtype C	0	R	7/9*	
D/E	25		6/04		B/E recombinent	25		6/04	
D/F	25	· · · · · · · · · · · ·	0/24		B/F recombinant	25	· · · · · · · · · · ·	0/24	
recombinant		R	18/24				R	18/24	
ap120/C3	Total=67.5	377NCGGEFFYCN386	135/200	ND**	ap 41	Total=74.0	529TMGAASITL537	37/50	A2
Subtype B	99.6		120/152		Subtype B	29.6		2/7	
Subtype B	00.0		129/133		Subtype B	20.0		2/1	
		D	10/153				<b>L</b>	2/7	
		A	4/153*				VA.	2/7*	
Subtype F	0	R	9/14		Subtype F	90.9		10/11*	
	-	M	1/1.1*						
			4/14					10/10	
Subtype C	0		7/9*		Subtype C	100		13/13	
B/F	25		6/24		B/F recombinant	63.2		12/19	
recombinant		P	19/24		_,		M	2/10*	
Tecombinant			10/24				· · · · ·	3/19	
_gp 41	Total=52.0	565LLQLTVWGI573	26/50	A2	gp 41	Total=70.0	584ERYLKDQQL592	35/50	B14,A32
Subtype B	42.9		3/7		Subtype B	42.9		3/7	
31		М	4/7		31		R	2/7	
			-1/1					0/7	
							G	2/1	
Subtype F	100		11/11		Subtype F	63.6		7/11	
							Q	4/11	
Subtype C	0	М	13/13		Subtype C	73.9		10/13	
eabiype e	Ū	141	10/10		Cubtype C	10.0		0/40	
							<del>K</del>	2/13	
B/F	63.2		12/19		B/F recombinant	78.9		15/19	
recombinant		M	6/19*				Q	3/19*	
ap 41	Total=70.0	584 FRYLKDOOLLG594	35/50	ND**	ap 41	Total=70.0	585RYLKD00LL593	35/50	A*23 A24
				110					
Cubture D	40.0		2/7		Culture D	12.0		2/7	
Subtype B	42.9		3/7		Subtype B	42.9		3/7	
Subtype B	42.9	R	3/7 2/7		Subtype B	42.9		3/7 2/7	
Subtype B	42.9	R G	3/7 2/7 2/7		Subtype B	42.9	R G	3/7 2/7 2/7	
Subtype B	42.9	R G	3/7 2/7 2/7 7/11		Subtype B	42.9	· · · · R. · · · · · · · · · · · · · · ·	3/7 2/7 2/7 7/11	
Subtype B Subtype F	42.9 63.6	R G	3/7 2/7 2/7 7/11		Subtype F	42.9 63.6	· · · · R. · · · · · · · · · · · · · · ·	3/7 2/7 2/7 7/11	
Subtype B Subtype F	42.9 63.6	R	3/7 2/7 2/7 7/11 4/11		Subtype B Subtype F	42.9 63.6	R G Q	3/7 2/7 2/7 7/11 4/11	
Subtype B Subtype F Subtype C	42.9 63.6 73.9	R	3/7 2/7 2/7 7/11 4/11 10/13		Subtype B Subtype F Subtype C	42.9 63.6 73.9	R G Q	3/7 2/7 2/7 7/11 4/11 10/13	
Subtype B Subtype F Subtype C	42.9 63.6 73.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13*		Subtype B Subtype F Subtype C	42.9 63.6 73.9	R G Q R	3/7 2/7 2/7 7/11 4/11 10/13 2/13*	
Subtype B Subtype F Subtype C B/F	42.9 63.6 73.9 78.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19		Subtype B Subtype F Subtype C B/E recombinant	42.9 63.6 73.9 78.9	R 	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19	
Subtype B Subtype F Subtype C B/F	42.9 63.6 73.9 78.9	Q	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 2/40*		Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9		3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 2/10*	
Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9	RQ	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19*		Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19*	
Subtype B Subtype F Subtype C B/F recombinant gp 41	42.9 63.6 73.9 78.9 Total=90.0	R Q 676WLWYIKIFI666	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41	42.9 63.6 73.9 78.9 Total=82.0	R Q R esoWYIKIFIMIess	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B	42.9 63.6 73.9 78.9 Total=90.0 100	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 4/5/50 7/7	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B	42.9 63.6 73.9 78.9 Total=82.0 100	R Q R Q 680WYIKIFIMI688	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9	С	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 7/0/11*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=82.0 100 90.9	R Q R Q R Q 680WYIKIFIMIese	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11*	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 40/42*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=82.0 100 90.9	R Q Q Q Q	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 7/7 10/11* 10/11*	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3	R G R  678WLWYIKIFI686	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10//11* 11/13*	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2	R.     G.       Q.     R.       R.     G.       G.     <	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4	R Q Q Q Q 	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2	R G R Q 678WLWYIKIFI686	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 <u>Total=82.0</u> 100 90.9 84.6 68.4		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19*	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2	R R Q R R G G R G	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19*	A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 2/19*	A*2402
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19* 30/50	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype C B/F subtype B	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100	R     G       Q     G       R     G       G     G	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 2/19* 30/50 7/7	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 7/7 10/11*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype B Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6		3/7 2/7 2/7 7/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 2/19* 30/50 7/7 7/11	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 2/19* 30/50 7/7 7/11 4/11	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype F Subtype F	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6		3/7 2/7 2/7 7/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 30/50 7/7 7/11 4/11 7/12	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F Subtype C	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9 84.6	R.     G.       Q.     G.       R.     G.       G.     <	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11* 10/11*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype F Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype F Subtype C	42.9 63.6 73.9 78.9 <u>Total=82.0</u> 100 90.9 84.6 68.4 <u>Total=60.0</u> 100 63.6 0		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11//3* 13/19 2/19* 2/19* 30/50 7/7 7/11 4/11 7/13	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype F Subtype F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9 84.6	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11* 10/11*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype C Subtype B Subtype F Subtype F Subtype C	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6 0	R Q esoWYIKIFIMIese 	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19* 30/50 7/7 7/11 4/11 7/13 3/13*	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9 84.6 68.4	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11* 10/11* 11/13*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype F Subtype C B/F recombinant gp 41 Subtype C B/F recombinant Subtype C Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6 0 84.2		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19 2/19 30/50 7/7 7/11 4/11 7/13 3/13*	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype F Subtype C B/F Subtype C B/F	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9 84.6 68.4	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11* 11/13* 11/13*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype C B/F recombinant gp 41 Subtype B Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=82.0 100 90.9 84.6 68.4 Total=60.0 100 63.6 0 84.2	R Q esoWYIKIFIMIess  	3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19* 2/19* 30/50 7/7 7/11 4/11 7/13 3/13* 16/19 2/19*	A*2402 A*0205
Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype F Subtype C B/F recombinant gp 41 Subtype B Subtype F Subtype C B/F recombinant	42.9 63.6 73.9 78.9 Total=90.0 100 90.9 92.3 84.2 Total=82.0 100 90.9 84.6 68.4	R	3/7 2/7 2/7 7/11 4/11 10/13 2/13* 15/19 3/19* 45/50 7/7 10/11* 12/13* 16/19 2/19* 41/50 7/7 10/11* 10/11* 11/13*	A2 A2	Subtype B Subtype F Subtype C B/F recombinant gp 41 Subtype F Subtype C B/F recombinant gp 41 Subtype C B/F recombinant	42.9 63.6 73.9 78.9 <u>Total=82.0</u> 100 90.9 84.6 68.4 <u>Total=60.0</u> 100 63.6 0 84.2		3/7 2/7 2/7 7/11 4/11 10/13 2/13 * 15/19 3/19* 41/50 7/7 10/11* 11/13* 13/19 2/19 2/19* 30/50 7/7 7/11 4/11 7/13 3/13* 16/19 2/19*	A*2402 A*0205

\* Mutation with frequency lower than 5% was excluded from the table. \*\* HLA not determined. % similarity: % of sequences that have that epitope sequence. Frequency: n° of sequences/n° of sequences that have that epitope or mutation.

Table 2. Mapping of the frequency of the Los Alamos CD4 epitopes in the Brazilian HIV-1 sequences

HXB2 Location Protein region	% Similarity	Epitope sequency	Frequency	HLA	HXB2 Location Protein region	% Similarity	Epitope sequency	Frequency	HLA
gp 41	Total=50.0	562QQHLLQLTVWGIKQL576	25/50	ND**	gp 120 C1	Total=88.2	118IISLWDQSLKPC119	45/51	ND**
Subtype B	42.8		3/7		Subtype B	100		10/10	
		M	4/7						
Subtype F	100		11/11		Subtype F	100		10/10	
Subtype C	0	M	13/13		Subtype C	83.3		10/12*	
B/F recombinant	57.9		11/19		B/F	78.9		15/19	
		M	6/19*		recombinant		V	2/19*	
gp 41	Total=54.0	593LGIWGCSGKLIC604	27/50	ND**	gp 120 C1	Total=92.1	110SLWDQSLKPCVKLTPL125	47/51	ND**
Subtype B	100		7/7		Subtype B	100		10/10	
Subtype F	0	L	11/11		Subtype F	100		10/10	
Subtype C	100		13/13		Subtype C	91.6		11/12*	
B/F recombinant	36.8		7/19		B/F	84.2		16/19*	
		L	9/19		recombinant				
		LR	2/19*						
gp 41	Total=54.0	594GIWGCSGKLIC604	27/50	ND**	gp 41	Total=54.0	594GIWGCSGKLI603	27/50	ND**
Subtype B	100		7/7		Subtype B	100		7/7	
Subtype F	0	. L	11/11		Subtype F	0	.L	11/11	
Subtype C	100		13/13		Subtype C	100		13/13	
B/F recombinant	36.8		7/19		B/F	36.8		7/19	
		.L	9/19		recombinant		.L	9/19	
		. L R	2/19*				.LR	2/19*	

\*Mutation with frequency lower than 5% was excluded from the table. \*\* HLA not determined. % similarity: % of sequences that have the epitope. Frequency:  $n^{\circ}$  of sequences/ $n^{\circ}$  of sequences that have the epitope or mutation.

N-glycosylation sites (CTRPNNNTRK at amino acid position 296 to 305, at a frequency of 96.2%, and NCGGEFFYCN at amino acid position 377 to 386 with a frequency of 84.5%). The *ds/dn* ratio was high in some of the CD8<sup>+</sup> epitopes. This high ratio suggests that these particular epitopes may either not be under positive selection or are maintained under functional constraints.

The subtype B epitopes were the most conserved ones, especially in the C1 and C3 regions, followed by subtype F. In the V3loop region, subtype F was the most conserved one. However, none of the most frequent mutations were associated with the loss of N-glycosylation site at this position. The gp41 was the most conserved region among the subtypes and this can be explained by the absence of variable regions in this protein. Epitopes in this region showed high ds/dn ratios but these ratios were lower than the gp120 epitopes. This epitope conservation and the high ds/dn ratio suggest that these regions may be important for viral fitness. Mutations in this region might change the protein structure, reducing the infection capacity of the virus.

The epitope VPVWKEATTTL is associated with a rapid progression HLA allele, HLA-B35 [4] and exhibited low variation in non-C subtypes. Interestingly, this epitope was highly variable in B/F recombinants. This suggests that CTL may be exerting selective pressure on this subtype.

The regions involved in N-glycosylation were highly conserved. These sites are potentially important for the functioning of these proteins, and mutation in these regions might affect viral function.

#### Conclusion

An ideal vaccine would contain epitopes that would engender strong immune responses against the functionally important regions. Escape from these vaccine-induced immune responses would compromise viral fitness. Additionally, it will be important to continue analyzing epitope variability in other viral proteins in the HIV-1 strains circulating in Brazil as any eventual vaccine for use in Brazil will need to be relevant to the viruses in Brazil.

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