Identification of amino acid substitutions associated with neutralization phenotype in the human immunodeficiency virus type-1 subtype C gp120

Kirchherr, J. L., Hamilton, J., Lu, X., Gnanakaran, S., Muldoon, M., Daniels, M., Kasongo, W., Chalwe, V., Mulenga, C., Mwananyanda, L., Musonda, R.M., Yuan, X., Montefiori, D.C., Korber, M.T., Haynes, B.F., & Musonda, R. M. (2011). Identification of amino acid substitutions associated with neutralization phenotype in the human immunodeficiency virus type-1 subtype C gp120. Virology, 409(2), 163-174. DOI: 10.1016/j.virol.2010.09.031

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- The Kirchherr paper studied the role of neutralizing antibodies in attaching to common signature sequences in the HIV virus.
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HIV is a Highly Diverse Virus Due to Constant Adaptation in Variable Regions

- HIV-1 undergoes rapid genetic mutation.
 - Env gene is highly variable.
- HIV is a retrovirus which has 9 subtypes (A-K).
 - Subtype C accounts for 1/2 of HIV-1 infections globally.
- The Env gene codes for gp120 and gp41 which are expressed on the surface of the virus.
 - gp120 is an envelope protein which helps the virus avoid detection during the HIV life cycle.
 - V4 region of gp120 on the env gene is thought to contribute to patterns of neutralization susceptibility in subtype C viruses (Kirchherr et al. 2010).

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Kirchherr et al. Identified Potential Neutralizing Signature Sequences In The HIV Virus.

- Focus of this paper was to identify what induces Nabs in the HIV virus.
 - Nabs are neutralizing antibodies that are released in response to the HIV-1 infection.
- 15 subjects infected with the subtype C of the HIV virus were studied for neutralization.
 - Neutralization susceptibility was studied in autologous and heterologous plasma.
 - Heterologous neutralization was divided into strong, cross reactive neutralization and weak neutralization.
- The V4 region in the HIV virus was found during analysis of plasma samples to have the greatest neutralization potency.
 - This identification can have further reaching implications for the development of a HIV vaccine.

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Summary of Subjects SGA's and Functional *Env* Genes

Table 1Summary of SGAs and functional *env* genes from HIV-1-infected individuals.

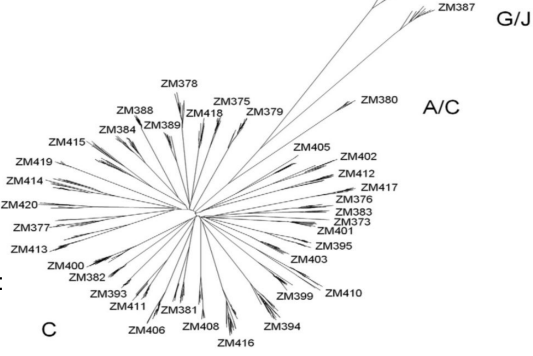
ID	Gender	Age	Viral load	Subtype	No. of SGAs	No. of functional env genes	Functional env genes (%)
ZM373	M	25	44,800	С	5	4	80
ZM375	F	31	52,800	C	10	8	80
ZM376	F	30	105,600	C	15	13	87
ZM377	M	41	248,800	C	16	15	94
ZM378	F	36	67,200	C	13	11	85
ZM379	M	35	97,600	C	14	11	79
ZM380	M	37	268,800	A/C	10	10	100
ZM381	F	36	487,200	C	13	12	92
ZM382	M	39	557,600	C	10	10	100
ZM383	F	31	20,240	C	10	5	50
M384	M	41	15,440	C	11	10	91
ZM387	M	30	1,120	G/J	9	9	100
ZM388	F	28	21,360	C	11	6	55
ZM389	M	33	1,520,000	C	10	10	100
ZM393	F	21	<384	C	11	2	18
M394	F	30	197,600	C	19	12	63
ZM395	F	22	776,800	C	21	20	95
M399	M	36	68,240	C	14	14	100
M400	M	35	103,200	C	18	15	83
M401	M	34	84,000	C	11	9	82
ZM402	F	37	3,920	C	9	6	67
ZM403	F	45	48,080	C	11	10	91
ZM405	F	36	13,920	C	12	9	75
ZM406	M	27	104,000	C	10	10	100
M407	M	38	283,200	D	11	9	82
ZM408	F	26	42,400	C	10	8	80
M410	M	42	143,200	C	11	6	55
ZM411	F	41	61,600	C	14	6	43
ZM412	F	30	260,000	C	10	7	70
ZM413	F	34	155,200	C	16	14	88
M414	F	25	213,600	C	22	14	64
ZM415	M	38	64,720	C	21	18	86
M416	M	45	80,800	C	23	23	100
ZM417	M	32	96,000	C	9	9	100
ZM418	F	21	532,000	C	10	8	80
ZM419	F	33	433,600	C	10	1	10
ZM420	F	24	286,400	C	14	13	93
Total	1000		The state of the state of		474	377	80

- Table1: This table lists each individual subject from which plasma was taken for this experiment.
 - Notable sections include the subtype, number of SGAs and the percent of Env genes.

Unrooted Trees Were Formed To Show Genetic Diversity.

 Fig. 1:All subjects within the circle of the unrooted tree represent subtype C subjects.

The only outliers are of other subtypes:
 D, G/J, and A/C



0.02

ZM407

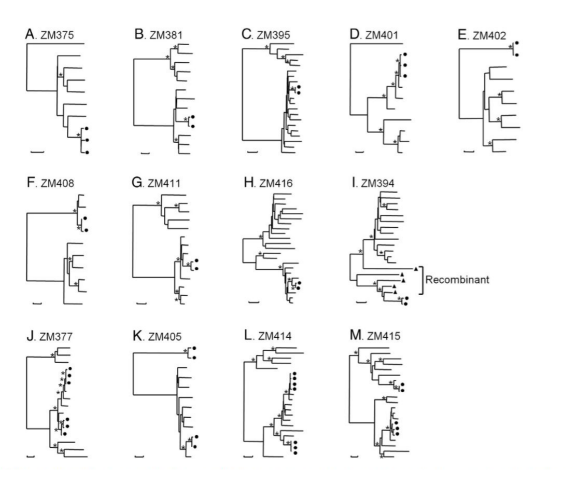
Table 2Characterization of clonal expansion *env* sequences in HIV-1-infected individuals.

ID	SGA No.	Luciferase activity (RLU)	Total viral population (%)	No. of amino acid differences among env sequences
ZM375	1	434,052	30	3, 4, 5
	9	437,851		
	11	610,112		
ZM377	10	175,205	19	1, 2, 3
	13	310,710		
	14	53,926		
	11	354,745	19	1, 4, 5
	12	30,765		
	16	77,427		
ZM378	7	802,968	15	3
	9	50,041		
ZM394	11	398,245	10	2
	18	21,273		
ZM395	14	327,566	10	1
	15	192,620		
ZM401	2	164,195	27	2
	19	66,826		
	20	161,349		
ZM402	8	48,981	20	0
	12	8,985		
ZM405	25	205,738	17	4
	44ª	1,393		
	43	92,501	17	0
	53	118,441		
ZM408	15	118,979	20	3
	23 ^a	2,218		
ZM411	6	92,343	14	3
	9	203,082		
ZM413	5	229,782	13	3
	13	348,872		
ZM414	1	47,219	18	0, 1
	10	70,754		
	23	218,271		
	28	4,134		
	9	141,022	14	1
	20	240,243		
	25	126,764		
ZM415	1	510,266	16	4
	2	114,320		
	26	553,315		
	15	496,933	10	1
	27	297,937		
ZM416	3	391,763	9	2
	16	296,184		
Control	SG3∆env	1,127		

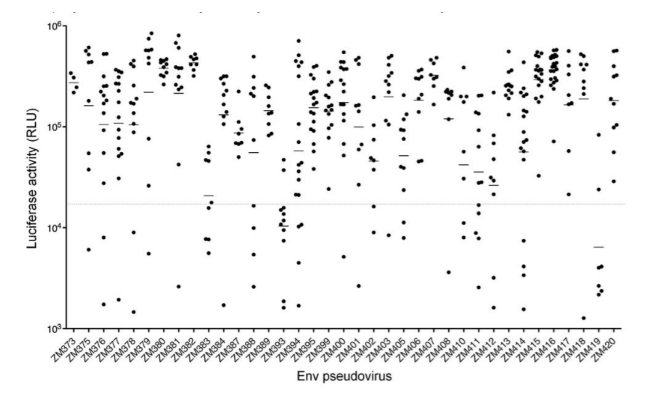
a Stop codon.

Luciferase Activity Shows Infectivity of HIV-1 Virus

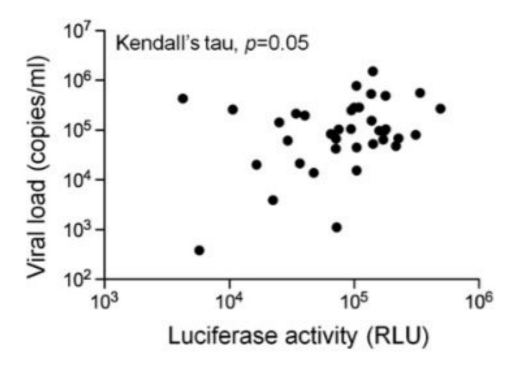
- Table 2: This table contains information that shows the infectivity of the HIV virus.
 - Using PCR, pseudoviruses were formed and then tested in the presence of TZM-bl indicator cells.
 - Luciferase activity indicates the presence of the HIV-1 virus.



Rooted phylogenetic trees of 13 subjects demonstrating clonal expansion within the *env* gene



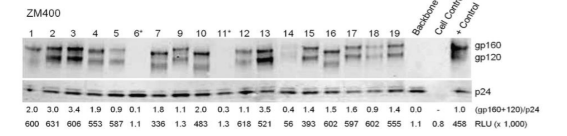
Infectivity of 474 *env* pseudoviruses from 37 individuals determined in TZM-bl cells and measured by luciferase activity.

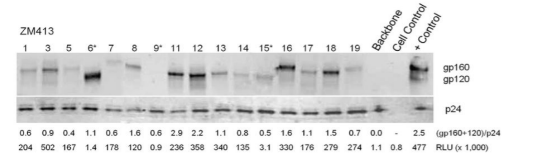


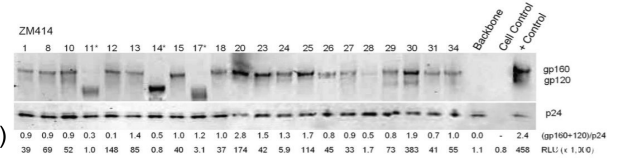
Trend supporting a positive correlation between viral load and pseudovirus infectivity. P-value of 0.05.

Western Blot Analysis Of HIV-1 Proteins In Transfected Cells



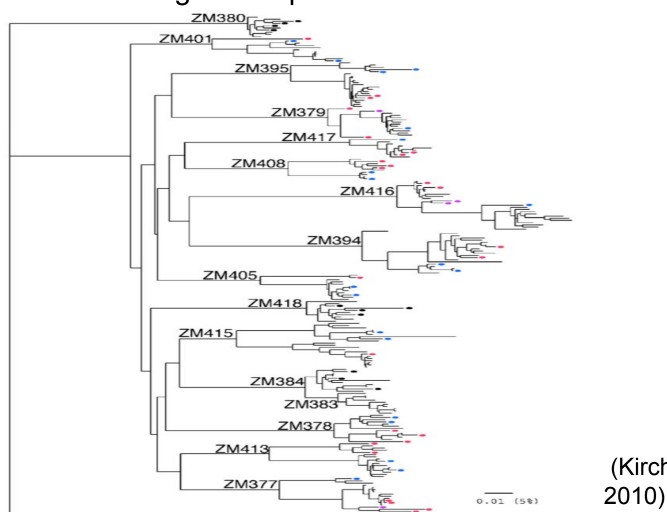






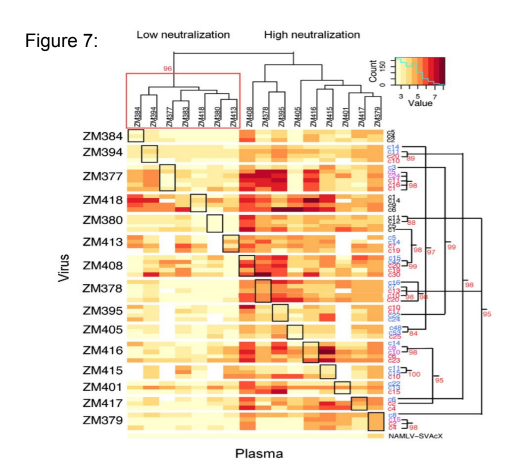
Phylogenetic tree of *Env* gene sequences

Figure 6:



(Kirchherr et al.

Hierarchical Clustering Of Virus Based On Neutralization Concentrations

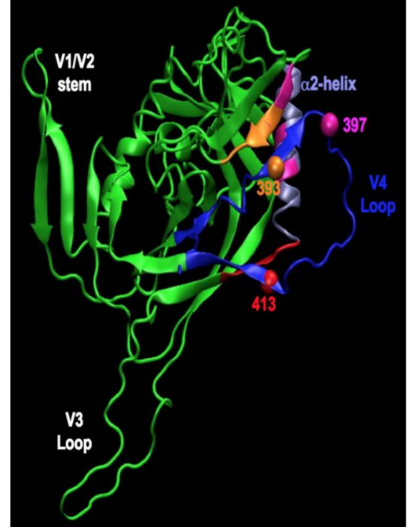


Four Sites for Signature Amino Acid Sequences Were Found; Sites Associated With Potent Neutralization Highlighted

V4	3 3 4 9 9 1 3 7 3	gp41 85
H_ZM395.CONSENSUS H_ZM401.CONSENSUS H_ZM405.CONSENSUS	GEFFYCNTsdLFngtyngtnstsnsnITlpCrIKQIINMWQKGEF-G-RNNPGNGTSDNQGKD-DKNSSKLFNTN-TDNLTIIRGKF-GYNRKD-SGNKGGGTDSN-STIKSG-VIMYDEN-STIQ	CONSENSUS-H RIRQGFEaAL1 H_ZM378.CONSENSUSL H_ZM379.CONSENSUSQ H_ZM401.CONSENSUSQ H_ZM405.CONSENSUSL H_ZM408.CONSENSUS
H_ZM416.CONSENSUS H_ZM417.CONSENSUS		H_ZM415.CONSENSUSL H_ZM416.CONSENSUSL L_ZM417.CONSENSUSL L_ZM384.CONSENSUS
L_ZM394.CONSENSUS L_ZM377.CONSENSUS L_ZM380.CONSENSUS L_ZM383.CONSENSUS L_ZM413.CONSENSUS	TNN-RLSEF-S-E-STQFNRSLSNDTEEDTDRTKNGTNMTDPQGSSWIFENGTANSTWP-GTQTNT-LLDNFISTG-S-G-STQFKFS-DNATAENATGT	L_ZM394.CONSENSUSQ L_ZM394.CONSENSUSQ L_ZM377.CONSENSUSQ L_ZM380.CONSENSUSQ L_ZM383.CONSENSUSQ L_ZM413.CONSENSUS -VQ L_ZM418.CONSENSUS -VQ

Fig. 8. The signature amino acid sequences associated with high levels of neutralization activity in plasma. H represents sequences in the group of plasma samples with high neutralizing activity against heterologous viruses, whereas L represents sequences that exhibited weak neutralizing activity. Numbers are used to show corresponding locations in the HXB2 reference strain. Dashes indicate the identical amino acids present in the reference sequence. Periods are used to designate gaps to maintain the alignment. Signature sites associated with potent neutralization are shown in red. Non-signature amino acids in key positions are shown in blue.

(Kirchherr et al. 2010)



CD4-bound Truncated gp120 Structures From B and C Clades Showed That Both Structures Are Similar

Plasmas With High Neutralization Of Heterologous Virus Trended Toward High Neutralization Of Autologous Viruses

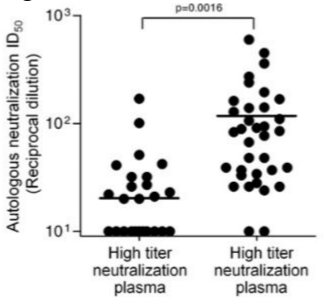


Fig. 10. Comparison of autologous neutralizing activity between low and high heterologous neutralizing plasma samples. Nab titers from low heterologous neutralization plasmas (n=6) or high heterologous neutralization plasmas (n=9) were compared. Values at Y-axis are the reciprocal plasma dilutions at which luciferase activity (RLU) was reduced by 50% relative to virus control wells by autologous plasma.

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Analysis of Data Revealed Signature Sequences of Four Amino Acids.

- This paper found variable regions in the gp120 gene were most susceptible to Nabs.
- Four signature sites were observed, three of which were in the V4 region of gp120, and the fourth being in gp41.
- The V4 region consisted of a four amino acid sequence and was the site for Nab binding in autologous and heterologous plasmas.
- Glycine residues in the V4 binding loop play a major role in binding of Nabs.
 - Loss of a gly residue would reduce negative charge in that region, preventing antibody binding.
 - Glycines have no side chains, making them more variable in terms of conforming for attachment.
- Clonal expansion could be an important role in disease proliferation.

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Identification of False Positive Sites Necessary for Future Experimentation

- Previous study by Moore et al. suggests C3V4 region may be responsible for inducing Nabs.
 - Kirchher et al.'s signature sites had amino acids located near this region, suggesting stronger Nab responses.
- The identified sites must each be tested to examine their role in susceptibility to Nabs, since three of them associate with the C3 alpha-2 helical domain, which is thought to contribute to patterns of neutralization susceptibility in subtype C viruses.
- Identification of false positives is necessary before further study
 - o One or more of the identified sites is not related to viral susceptibility.
 - Elimination of these sites will allow vaccines to target amino acid sequences in relevant sites and increase the susceptibility of the virus to Nabs.

The Kirchher et al. Paper As A Whole

- The HIV virus has the ability to mutate rapidly, protecting itself against autoimmune response.
- Kirchher et al. studied what aspect of the HIV-1 virus induced response from Nads.
- The gp120 *Env* gene was studied for it's susceptibility to Nads.
- The V4 region in gp120 contains several signature sequences that have high neutralization potency.
- The results of this study lays the groundwork for further research which can be used to develop a vaccine for the HIV-1 virus.

Acknowledgements

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References

Kirchherr, J. L., Hamilton, J., Lu, X., Gnanakaran, S., Muldoon, M., Daniels, M., Kasongo, W., Chalwe, V., Mulenga, C., Mwananyanda, L., Musonda, R.M., Yuan, X., Montefiori, D.C., Korber, M.T., Haynes, B.F., & Musonda, R. M. (2011). Identification of amino acid substitutions associated with neutralization phenotype in the human immunodeficiency virus type-1 subtype C gp120. Virology, 409(2), 163-174. DOI: 10.1016/j.virol.2010.09.031