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Full length HIV-1 Gag determines protease inhibitor susceptibility within *in vitro* assays

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Keywords

HIV; HAART; protease inhibitors; gag; subtype

Introduction

HIV-1 protease (PR) cleaves Gag and Gag-Pol polyproteins during virus maturation. Gag mutations involving cleavage sites, in particular p7/p1, p1/p6, have been associated with drug resistance or recovery of replicative capacity (RC) both in the absence[1] and presence [2-4] of PR drug resistance associated mutations. Polymorphisms at these positions as well as the p24/p2 and p2/p7 cleavage site have also been documented in drug naïve patients [3,

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Conflicts of interest

RG has received educational travel grants from Gilead Sciences, Bristol Myers Squibb and Boehringer-Ingelheim. DP has acted as a consultant for Bristol Myers Squibb, Johnson&Johnson, Boehringer Ingelheim, Roche, and Gilead Sciences. AK, ALM, GJT and CMP have no conflicts of interest

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5]. Furthermore, we and others have identified areas of the Gag protein from PI-treated patients that are associated with both drug resistance and recovery of RC (outside cleavage site regions) [6-8]. Polymorphisms throughout *gag* may therefore be relevant to response to PIs in naïve patients. Despite this body of data, phenotypic testing of clinical samples is still focused primarily on the *protease* gene. Phenotypic resistance test vectors have generally incorporated patient derived *Protease* and the 5' end of *gag* (including p7/p1 and p1/p6 cleavage sites) into a *gag* containing vector, typically derived from an HIV-1 subtype B molecular clone. There exists therefore the possibility of misclassifying drug susceptibility based on such an approach.

Non-B subtypes of HIV-1 account for the vast majority of infections worldwide. As the WHO public health rollout of HAART progresses substantial numbers of patients will require second line therapy with ritonavir-boosted protease inhibitors (PI). Since little *in vitro* work has been undertaken on the basis of PI susceptibility in full length non B viruses we focused on PI drug susceptibility in wild type subtype A and C laboratory and clinical strains, not previously exposed to antiretrovirals, as these different subtypes provide a wide genetic diversity within protease and *gag*. We sought to explore the role of full-length *gag* as well as *protease* in determining PI susceptibility in such wild type viruses.

Materials and Methods

Construction of resistance test vectors

A replication defective, envelope deleted subtype B clone p8.91NSX, expressing gag-pol, tat and rev was used as previously described [6]. This vector was further modified by silencing an existing EcoRI site at the end of integrase and creating a new EcoRI site just after the end of protease using site directed mutagenesis to allow cloning of gag-protease sequences. The two vectors had indistinguishable replication capacities and drug susceptibility profiles (data not shown). We obtained the following molecular clones from the NIH AIDS Research and Reference Reagent program: (subtype) p97ZA012.1 (C)[9], p8CN006.29 (C)[9], pMJ4 (C) [10], p94CY017.41 (A)[11]. Additionally the gag-protease region was amplified from two treatment naïve patients, identified by phylogenetic analysis as having subtype A virus during routine genotypic testing at first HIV diagnosis. RNA was extracted from plasma virus (QIA amp® viral RNA extraction kit, Qiagen,) and reverse transcribed using Moloney-MLV-RT and random priming (Promega). Gag-pol coding sequence was amplified from cDNA by nested PCR using outer primers 5'gagout (GTG TGG AAA ATC TCT AGC AG) and RTrev1 (CTG GRA TAA CYT CTG CTT) and an inner primer set as described[6]. Both PCR steps were carried out using Expand® High Fidelity PCR kit (Boehringer Manheim). The subtype A gag-protease sequences were cloned into the vector backbone, and representative clones (12.2 and 32G) chosen that had the same pol sequence as determined from plasma virus population sequencing. We utilized the naturally occurring ApaI restriction site towards the end of gag to generate chimeras between the test samples and the subtype B reference virus as used by commercial phenotypic systems.

Generation of virus stocks and drug susceptibility testing

293T cells were cotransfected with a *gag-pol* expression vector containing cloned *gag-protease* sequences, and plasmids encoding VSV-G envelope and luciferase reporter genes, with PI drug susceptibility testing carried out as previously described [6]. Results, derived from at least two independent experiments (each in duplicate) were analyzed and IC₅₀ determined using Graph Pad Prism 5 (GraphPad Software, Inc. La Jolla, CA, USA). Susceptibility testing for RT inhibitors differed in that supernatant containing resistance test vectors was harvested 48 hours post transfection, and used to infect fresh target 293T cells in the presence of serial dilutions of RT inhibitors in a 96 well plate format. Replicative

capacity (RC) of these viruses was assessed by comparing the luciferase activity of recombinant virus to that of the wild type subtype B control virus in the absence of drug. Equal amounts of input plasmid DNA were used and we have previously shown that % infectivity correlates well with infectivity/ ng p24 in this system[6]. The drugs used in this study (ATV, DRV, LPV, SQV, AZT and NVP) were obtained from the AIDS Research and Reference Reagent Program, Division of AIDS, NIAID, NIH.

Multiple sequence alignment

was carried out using the program Muscle (www.drive5.com/muscle/) with manual alignment in Se-Al (tree.bio.ed.ac.uk).

Results

Discrepancy between gag-protease and protease alone vectors in PI susceptibility

We tested a panel of six viruses against the following PIs: Atazanavir (ATV), Darunavir (DRV), Lopinavir (LPV) and Saquinavir (SQV). Fold changes relative to subtype B are shown in Table 1. For full-length *gag-pro* constructs we observed a range of PI susceptibilities, with 5 out of 6 showing reduced susceptibility for at least one drug (using a generous biological cut-off of FC>2, which is greater than the assay variability as determined by the standard deviations of repeated IC50 values for the reference virus). Overall, there was a 5-fold reduction in susceptibility to at least one drug in half the samples tested, with the greatest reductions seen for lopinavir, with more variable changes in susceptibility for atazanavir and saquinavir.

Importantly however, when specific *proteases* from these viruses were tested in the context of the subtype B *gag* (table 1), as in most commercially available phenotypic drug resistance assays, a reversal of this reduction in susceptibility was generally observed. This use of the *protease* alone vector resulted in overestimating the LPV susceptibility of CY017 by nearly 25 times, of ZA012 by nearly 9 times and for 32G by a factor of 7, compared to the *gag-protease* vector (table 1).

Gag alone confers reduced PI susceptibility

In order to determine if the determinants of reduced susceptibility in some of the viruses were located in *gag*, we examined the profile of our panel of *gag* sequences fused not with their cognate *protease*, but with the reference subtype B *protease* (Fig 1A) and compared these to the full-length cognate *gag-protease* samples with regard to LPV susceptibility (Table 1). We confirmed that *gag* alone determined reduced susceptibility to PI. Notably, ZA012 *gag* with B *protease* showed 14-fold reduced susceptibility as compared to subtype B *gag-pro*. This experiment also demonstrates that the cognate *protease* restores susceptibility to PI and in the extreme case of MJ4, can completely reverse a partially resistant phenotype to a hyper-susceptible one, in relation to subtype B.

There was variation in replicative capacity between constructs (Figure 1B). Interestingly the cognate *gag-protease* combination had a lower RC than *gag* with a subtype B *protease* in both ZA012 and 32G. There was no direct correlation between PI susceptibility and RC; for example there was an 8.9 fold difference in LPV FC for *gag-protease* versus *protease* alone in ZA012, but no difference in RC (Figure 1B, compare white and grey bars). As a control for any potential impact of replication capacity on drug susceptibility, we performed phenotypic susceptibility testing for the panel of viral constructs against the RT inhibitors zidovudine (Figure 1C) and nevirapine (data not shown). For both drugs, the fold changes relative to the subtype B reference were less than 1.5 fold across all constructs tested. We

also confirmed that use of cognate *gag-pro* or cognate *gag-pro-RT* did not affect PI susceptibility results (Figure 1D).

Finally, we analysed *gag* and *protease* sequences from the seven viruses tested (Figures S1 and S2). Comparison of *gag* cleavage sites (in boxes) shows that most are highly conserved, with the exception of p7/p2 (Figure S1, orange box). *Protease* sequence analysis showed no major resistance mutations and typical non-B polymorphisms in subtypes A and C, some of which are classified as minor resistance mutations (Figure S2).

Conclusions

Overall these data demonstrate that virus specific protease confers variable susceptibility to PI, and more importantly suggest that consideration of protease alone may overestimate PI susceptibility in viruses not previously exposed to this class of drugs. This could lead to misclassification of susceptibility for some isolates, and potentially also limits assessment of the genotype-phenotype relationships in drug experienced patients.

Given the expanding body of evidence showing that *gag* mutations are known to contribute to PI resistance in treated patients, and in light of our data we recommend that full *gag* sequences should be included in recombinant virus assays to determine PI susceptibility in clinical isolates. We would also encourage a rapid expansion of full-length *gag* sequencing in patients failing PI-based therapy, to populate the relevant relational databases increasingly used to assess the clinical impact of HIV drug resistance. Gag genotypic determination to guide patient care can be then be evaluated once specific Gag determinants of PI susceptibility have been identified.

Lower clinical cut-offs of 5.2 and 10 fold have been proposed for ATV and LPV response respectively [12, 13]. Although fold changes in our system may not be directly comparable, we were concerned to observe that some viruses approached and even exceeded these values. In the ACTG 5142 study 23% of patients starting boosted lopinavir had a viral load >50 copies/ml after 2 years, as compared to 11% for NNRTI based regimens (p=0.003)[14]. Patients with viral failure after boosted PI (bPI) in this and other studies infrequently have major protease resistance mutations[15, 16], although there is a lack of data on evolution of Gag mutations. A better understanding of determinants in Gag reducing susceptibility to PI might explain these observations.

Finally, our findings may have important implications for the public health approach to ART. Increasing numbers of patients are moving to PI-based second line therapy following the rollout of HARRT, and combined LPV/ritonavir is the most widely available PI in resource poor settings. Some of the subtype A and C constructs studied demonstrated particularly high FCs for this agent, and as resistance testing is not routinely performed in these patients, further research is clearly warranted.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Abbreviations

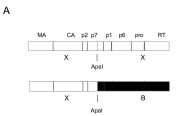
ATV	atazanavir		
DRV	darunavir		
LPV	lopinavir		
SQV	saquinavir		
AZT	zidovudine		
NVP	nevirapine		

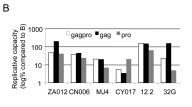
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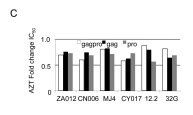
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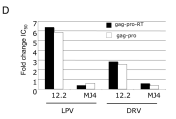


Figure 1.(A) Schematic diagram of chimeras used Key: X (test sequence), B (reference subtype B sequence), MA (matrix, or p17), CA (capsid, or p24), p2 (spacer peptide 1), p7 (nucleocapsid), p1 (spacer peptide 2), p6, pro (protease), RT (reverse transcriptase). (B) Mean replicative capacity of study viruses expressed as % of wild type subtype B reference (log $_{10}$) (C) Mean fold change in IC $_{50}$ for the panel of viruses against AZT. (D) Comparison of mean fold change in LPV and DRV IC $_{50}$ using cognate *gag-pro-RT* versus *gag-pro* for two viruses (12.2 and MJ4).

Table 1

Viral construct (subtype)	Fold change in IC50 as compared to reference subtype B [SE] Ratio of FC gagpro:pro				
	ATV	DRV	LPV	SQV	
ZAD12gagpro (C)	4.98[0.64]	2.45[1.47]	7.07[2.0]	6.30 [0.68]	
ZA012pro	1.73[0.02]	1.30[0.37]	0.80 [0.03]	0.84 [0.48]	
ZAD12gag			14.6[0.14]		
	2.88	1.88	8.85	7.5	
CN006gagpro (C)	2.18[0.22]	0.77[0.44]	2.34[0.36]	1.17[0.18]	
CN006pro	5.15[1.32]	0.51[0.08}	2.05[0.48]	1.37[0.39]	
CN006gag	-	-	4.9[0.72]		
	0.42	1.51	1.05	0.89	
MJ4 gagpro (C)	0.55[0.17]	0.31[0.01]	0.40[0.04]	0.49[0.04]	
MJ4 pro	0.62[0.19]	0.12[0.03]	0.22[0.08]	0.28[0.17]	
MJ4 gag			3.60[0.51]		
	0.88	2.58	1.82	1.75	
CY017gagpro (A)	1.19[0.53]	1.84[0.78]	5.44[1.60]	2.87[0.92]	
CY017pro	1.03[0.74]	0.59[0.14]	0.22[0.10]	0.48[0.10]	
CY017gag	-	-	8.69[0.18]		
	1.16	3.12	24.7	6.00	
12.2gagpro* (A)	7.33[0.71]	2.83[0.61]	6.38[0.60]	3.37[0.11]	
12.2pro	2.93[0.41]	0.89[0.17]	2.18[0.98]	1.32[0.15]	
	2.50	3.18	2.92	2.55	
12.2gag	-	-	6.53[0.74]	-	
32Ggagpro* (A)	0.89[0.52]	0.62[0.23]	3.13[0.43]	0.22[0.18]	
32Gpro	1.01[0.16]	0.34[0.07]	0.44[0.33]	0.50[0.15]	
32Ggag	·	-	6.15[1.8]	-	
	0.88	1.82	7.11	0.44	