6TH IAS CONFERENCE ON HIV PATHOGENESIS, TREATMENT AND PREVENTION (IAS 2011)

PROTEASE INHIBITORS AFTER 15 YEARS: REPORT FROM THE IAS

Jean-Guy Baril, MD Clinique Médicale du Quartier Latin Montreal, QC

> Jason Brunetta, MD Maple Leaf Medical Clinic Toronto, ON

In 1996, a protease inhibitor (PI)-based antiretroviral combination led to the first opportunity for long-term survival for most patients living with human immunodeficiency virus (HIV). On the 15th anniversary of that breakthrough, PIs are no longer a mandatory component of an effective antiretroviral regimen, but they remain an important and reliable option in HIV control. As they are among the most potent antiretroviral agents, PIs remain a reasonable choice in a wide variety of clinical scenarios, including first-line treatment. Although some of the initially introduced PIs remain available, the common PIs in most formularies are ritonavir-boosted atazanavir (ATV/r), darunavir (DRV/r), and lopinavir (LPV/r). Relative to first generation PIs, these have few dosing restrictions, a low pill burden, and less relative risk of adverse events. Despite the broad experience with all of these agents, optimal use of PIs remains a dynamic exploration of relative strengths and weaknesses, particularly in the context of an aging HIV-infected population. This update assembles key studies on PIs from the 2011 IAS meeting.



Jean-Guy Baril, MD & Jason Brunetta, MD 6th IAS Conference on HIV Pathogenesis, Treatment and Prevention (IAS 2011)

Protease Inhibitors Compared: Renal Function

Protease inhibitors (PIs) are potent and versatile antiretroviral agents that were the foundation of the first effective combination therapies for suppression of human immunodeficiency virus (HIV). While they continue to be used in a traditional three-drug regimen with two nucleoside reverse transcriptase inhibitors (NRTIs), they are being increasingly used in novel strategies, particularly in combinations that permit sparing of NRTIs which is an appealing approach to reducing the risk of adverse events and to simplify therapy. NRTI-sparing regimens may play an important role in reducing co-morbidities common to aging HIV patients, such as renal impairment. In traditional or novel applications, the current efforts to compare the relative characteristics of the most frequently prescribed PIs focus almost entirely on relative safety. In fact, one of the areas of greatest focus at this year's IAS conference was relative effects on renal function, a rapidly-emerging concern in countries, including Canada, where the average age of individuals infected with HIV is rising rapidly.

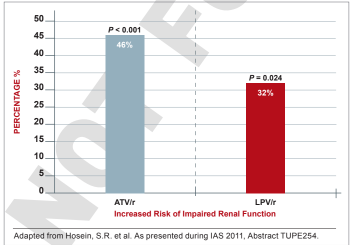
Of a set of new studies evaluating renal toxicity with Pls, one came from Canada. Drawn from the Canadian

Data indicated that PIs are associated with modest renal toxicity, but that risk may not be shared equally among agents within this class.

Observational Cohort Collaboration (CANOC), the data indicated that atazanavir (ATV) and lopinavir (LPV) were associated with modest renal toxicity compared to

non-PI regimens. In this study, factors associated with impaired estimated glomerular filtration rate (eGFR) were assessed in 985 patients being followed at eight collaborating centres in British Columbia, Ontario and Quebec. Of the two PIs evaluated, ATV was associated with a hazard ratio (HR) of 1.46 (P<0.001) while LPV posed a HR of 1.32 (P=0.024) in a multivariate analysis when both were compared to antiretroviral therapy without a PI (Figure 1).

FIGURE 1 | CANOC Study



The authors, led by Sean R. Hosein of Canadian AIDS Treatment Information Exchange (CATIE), in Toronto, Ontario, suggested that the PI data are consistent with previous studies, including a cross-sectional analysis of the French Aquitaine cohort (Dauchy FA et al. *Kidney Int* 2011;80(3):302-9) and the collaborative EuroSIDA cohort study (Mocroft A et al. *AIDS* 2010; 24(11):1667-78). In a multivariate analysis conducted in the Aquitaine study, the odds ratio (OR) for developing proximal renal tubular dysfunction on ATV was 1.28 per year of exposure. The OR for the same outcome for tenofovir (TDF) in the study was 1.23 per-year of exposure. In the EuroSIDA cohort, the OR among PIs for two consecutive measures of abnormal eGFR (<60 ml/min per 1.73 m²) was 1.21 for ATV (*P*=0.0003), 1.12 for indinavir (IDV) (*P*<0.0001), and 1.08 for LPV/r (*P*=0.03).

In a separate study at the IAS meeting, the same issue was addressed when electronic medical records were reviewed for 2,115 patients starting antiretroviral therapy with efavirenz (EFV), ATV/r, LPV/r, or darunavir (DRV)/r over a recent nearly four-year period. The study was designed to identify the rate of renal toxicity, defined as eGFR <60 ml/min per 1.73^2 . All participants had an eGFR >60ml/min at baseline. Compared to EFV, the HR elevation for renal impairment was not statistically significant for DRV (P=0.108), but was statistically significant for LPV (P=0.017) and ATV (P=0.004) after adjusting for gender, age, baseline eGFR, baseline CD4 count, hepatitis B and C status, prior exposure to TDF and IDV.

The authors, led by Dr. Neesha Rockwood, Chelsea and Westminster Hospital, London, UK, noted that 50% of those who developed renal impairment did show recovery over follow-up, but they suggested PIs may not be interchangeable for this risk. The mechanism of potential kidney toxicity of ATV and LPV is not clear. Of course, caution is required when interpreting the results of retrospective cohort studies. In particular, patients in non-randomized studies may have received different prescriptions based on baseline eGFR. Therefore, any associations observed in retrospective analyses are best investigated in prospective, randomized, and controlled trials.

New information about renal stones, another potential complication of antiretroviral therapy, was also presented at the IAS meeting. In this study, also led by Dr. Rockwood of London's Chelsea and Westminster Hospital, investigators systematically looked for renal stones in patients taking ATV, LPV, DRV, or the non-nucleoside reverse transcriptase inhibitor (NNRTI) EFV. The rate of stones per 1000 patient years of exposure on ATV was 7.3, or more than three times higher, than the 1.9 rate associated with any of the other agents evaluated (*P*<0.001). The relative increase in

risk remained similar after excluding patients previously exposed to IDV. This led the authors to conclude that ATV exposure is associated with significantly increased rate of renal stones when compared to the other therapies evaluated (EFV, LPV/r, and DRV) with and without adjusting for prior IDV exposure (Table 1).

TABLE 1 | Rates of Renal Stones Associated with PI Therapies

				Renal stone sub-analysis excluding IND/r/ATV/r		
	ATV/r n=1206	EFV/DRV/LPV n=4449	P value	ATV/r n=1000	EFV/DRV/LPV n=3293	P value
No. of patients with RS	24	24		15	15	
Prevalence renal stones: per 1000 patients (95% CI)	20 (13-30)	5.4 (3.2-7.6)	<0.001	15 (8.4-24.6)	4.6 (2.5-7.5)	<0.001
Rate of renal stones per 1000 patient years of ARV exposure (95% CI)	7.3 (4.7-10.8)	1.9 (1.2-2.8)	<0.001	5.67 (3.18-9.36)	1.51 (0.85-2.49)	<0.001

PI Risks and Potency

In direct comparisons of traditional PI-anchored threedrug regimens, there was little attention paid at the 2011 IAS meeting to relative effects on cardiovascular (CV) risk. Although renal impairment is a CV risk factor, it is well recognized that aging HIV patients should be

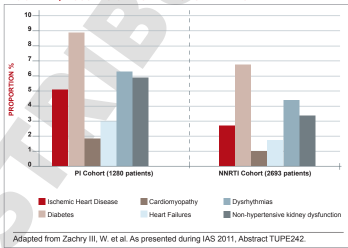
Although renal impairment is a CV risk factor, it is well recognized that aging HIV patients should be closely monitored for all CV risk factors, regardless of regimen.

closely monitored for all CV risk factors, regardless of regimen. However, one study, led by a team of investigators from Abbott Laboratories

did find higher baseline CV risk in patients treated with a PI compared to those treated with a NNRTI. In data derived from a medical claims database, those in the PI cohort (1280 patients) had a greater baseline prevalence of ischemic heart disease (5.08% vs. 2.67%; P<0.001), diabetes (8.83% vs.

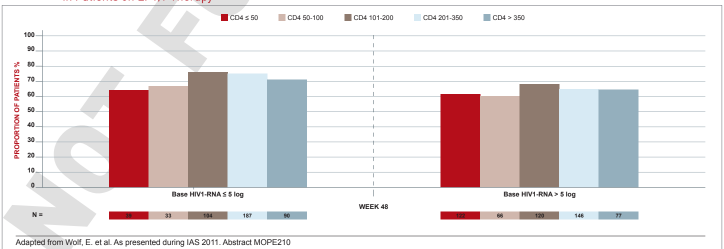
6.76%; P=0.02), cardiomyopathy (1.80% vs. 0.97%; P=0.026), heart failure (2.97% vs. 1.67%; P=0.008), dysrhythmias (6.25% vs. 4.34%; P=0.01), and non-hypertensive kidney dysfunction (5.86% vs. 3.34%; P<0.001) when compared to those in the NNRTI cohort (2693 patients) (Figure 2). Also, more patients in the PI cohort vs. the NNRTI cohort had a history of HIV-related conditions (17.73% vs. 12.33%; P<0.001) including Kaposi's sarcoma, cytomegalovirus, pneumocystosis, cachexia and cryptosporidiosis. Post-hoc (sensitivity) analysis of patients with only one diagnosis of HIV and >1 ARV prescriptions yielded similar results.

FIGURE 2 | Baseline CV risk in PIs vs. NNRTIs



Although the authors did not speculate on the reasons why patients prescribed PIs had increased baseline prevalence of CV risk factors, CV-related morbidity and HIV-related morbidity when compared with subjects receiving NNRTIs, these results may help future observational analyses better control for potential confounders when investigating antiretroviral-associated CV risk.

FIGURE 3 | STAR and STELLA Cohorts: 48-week Virologic Response Stratified by CD4 Cell Categories and Baseline HIV-1 RNA Levels in Patients on LPV/r Therapy



Again, there was no significant study comparing potency in two traditional triple-drug regimens anchored with a PI, but one study did assess the potency of a ritonavir-boosted LPV (LPV/r)-based regimen in patients with low baseline CD4 counts. Although a previous study had suggested less treatment success when CD4 counts are low at treatment initiation, data from the German STAR and STELLA cohorts reached a different conclusion. In this study, led by Dr. Eva Wolf, MUC Research, Munich, Germany, no relationship was seen between 48-week treatment success (HIV suppression at <50 HIV RNA copies/mL) and baseline CD4 counts in 984 treatment-naive patients treated with a LPV/r-based regimen. The efficacy of LPV/r across baseline strata was seen both in univariate analysis and after adjusting for variables such as age, NRTI backbone, and viral load (Figure 3).

More data have been made available to challenge the once prominent belief that EFV-based regimens may not be appropriate in patients with high viral loads or low baseline CD4 cell counts. In an open-label study, 87 treatment-naive patients with a baseline CD4 count <100 cells/mm³ were randomized to EFV, LPV/r, or ATV/r plus TDF and emtricitabine (FTC). At 48 weeks, the proportion of patients with optimal HIV suppression (<50 HIV RNA copies/mL) was 79% for EFV, 62% for LPV/r. and 53% for ATV/r. The immune reconstitution expressed as cells/mm³ was greatest on LPV/r (226), but was not significantly different from the rates on EFV (199) and ATV/r (186). The authors of the study, led by Dr. José M. Miro, Hospital Clinic i Provincial, Barcelona, Spain, confirmed that EFV should not be considered less potent than PIs in highly immunocompromised patients.

Protease Inhibitors in Novel Approaches

One of the growing trends in PI application is NRTI-sparing regimens in which a PI is used as monotherapy or combined with such therapies as integrase inhibitors or the CCR5 antagonist maraviroc. In 144-week results from the MONET trial, which compared DRV/r monotherapy to DRV/r plus 2 NRTIs in patients with HIV RNA <50 copies/mL at baseline, monotherapy showed non-inferior efficacy in the switch-included analysis, but did not show non-inferior

The presence of HCV did not significantly increase failure rates in LPV/r patients randomized to discontinue NRTIs when compared to those who remained on triple therapy. efficacy as compared to combination therapy (69.3% vs. 75.2% <50 copies/mL) when switch was counted as failure as defined by time to loss of virologic response (TLOVR) on an

intention-to-treat (ITT) analysis in which switch=failure. Additionally, monotherapy performed far less well in those who had hepatitis C infection (HCV+) (43.5% vs. 73.3%) when the same type of analysis (TLOVR, switch=failure) was used. Although DRV/r monotherapy failed to show non-inferiority

on the primary TLOVR switch=failure analysis, the authors of the study, led by Dr. José R. Arribas, Hospital la Paz, Madrid, Spain, indicated that boosted PI monotherapy may be an option in well-controlled patients when there is a good reason to avoid NRTIs. Selective application may also be appropriate.

In a monotherapy trial with LPV/r, which is among PIs frequently used in NRTI-sparing regimens, HCV+ did not significantly increase failure rates. In this study, called OKO4, patients receiving LPV/r plus two NRTIs were randomized to remain on their regimen or to discontinue the NRTIs and remain on LPV/r as monotherapy. While the proportions of patients with HIV RNA <50 copies/mL were higher on triple therapy in both HCV+ and HCV- groups (94% and 95.8%, respectively), they were not statistically different in the LPV/r monotherapy group (90.9% and 83.9%, respectively; P=0.38) in the ITT analysis (Table 2). In the per protocol analysis, differences were even smaller. The authors of the study, led by Dr. Federico Pulido, Hospital 12 de Octubre, Madrid, Spain, concluded that although a nonsignificant trend towards higher failure rates in the HCV+ population was seen in the ITT analysis, LPV/r monotherapy may be an option in selected HCV+ patients when it is well tolerated and patients remain adherent.

TABLE 2 | Monotherapy vs. Triple Therapy in HCV Infection

	Monot	herapy (n=100)	Triple Therapy (n=98)			
	HCV+ n=44 (44%)	HCV- n=56 (56%)	Р	HCV+ n=50 (51%)	HCV- n=48 (49%)	P
% <50 c/mL (M/C=F analysis)	70.5%	82.1%	0.23	74%	81.3%	0.47
% <50 c/mL (VF analysis)	90.9%	83.9%	0.38	94%	95.8%	1

Another NRTI-sparing study, called RADAR, looked at the combination of DRV/r plus the integrase inhibitor raltegravir (RAL) versus DRV/r plus TDF/FTC. Led by Dr. Roger Bedimo, VA North Texas Healthcare System, Dallas, Texas, the study team randomized 79 patients to one of the two regimens. Viral load reductions overall were far faster on the NRTI-free combination, with 65.7% vs. 5.9% reaching target suppression (<50 HIV RNA copies/mL) at 8 weeks. However, this level of viral suppression was essentially the same at 24 weeks (86.2% vs. 87.9%) (Figure 4). Treatment discontinuations, including those for severe adverse events, were not significantly different over the course of 24 weeks. In addition, efficacy was similar when patients were stratified for viral loads >100,000 copies/mL and below. These data are reassuring because the ACTG 5262 results reported at the 2011 Conference on Retroviruses and Opportunistic Infections (CROI) suggested suboptimal

■ RAL/DRV/r ◆ TDF/FTC/DRV/r 100 VL <48 at WK 24: 90 RAL: 86.2% TDF/FTC: 87.9% 80 =0.570) 67.7 65.7 70. 60. 50 50. 40. 30. 20. 11.8 10 WK 2 WK 4 WK 8 WK 16 WK 24 N (obs) RAL 39 34 35 35 31 29 TDF/FTC 40 36 37 34 32 33 VL <400 c/mL at WK 24: RAL: 93.1%; TDF/FTC: 100% (P=0.215)
LOG₄₀ VL Change at WK 24: RAL: -2.77; TDF/FTC: -3.05 (P=0.17) Statistically significant difference b/w groups: Weeks 4 & 8: P<0.0001

FIGURE 4 | RADAR Study: Proportion of Patients with HIV RNA <48 c/mL through Week 24 (NC=M)*

efficacy for this regimen in the presence of high viral loads when tested in a single-arm study. An appropriately powered double-blind study is needed to confirm that this is a reasonable alternative regimen in treatment-naive patients.

*Non complete = missing

Adapted from Bedimo, R. et al. As presented during IAS 2011, Abstract MOPE214.

Another NRTI-sparing study, called PROGRESS, provided some insight of where this strategy may be most useful.

While nearly 20% of those on the arm that included TDF/FTC had at least a 5% decrease in total BMD. the proportion was under 5% in the RAL/LPV/r arm.

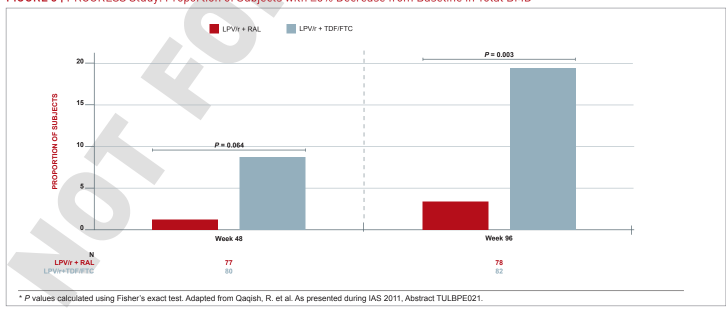
In this study, 160 treatment-naive HIV patients were randomized to LPV/r plus RAL or to a traditional PI

regimen of LPV/r plus TDF/FTC. HIV control at 96-weeks, like earlier timepoints, was similar, but new data presented

by an investigating team led by Dr. Roula B. Qagish found a difference in bone mineral density (BMD) loss. While nearly 20% of those on the arm that included TDF/FTC had at least a 5% decrease in total BMD, this level of bone loss was observed in less than 5% in the RAL/LPV/r arm. ART-naïve patients initiating LPV/r with RAL had no significant loss of total body BMD and significantly smaller reduction in spine BMD compared to those who received LPV/r plus TDF/FTC. The minimal effect of RAL in combination with LPV/r on BMD warrants confirmation through further study (Figure 5).

In a phase 2b study, a NRTI-sparing regimen containing ATV/r plus the CCR5 antagonist maraviroc was found to produce an acceptable and sustained rate of viral

FIGURE 5 | PROGRESS Study: Proportion of Subjects with ≥5% Decrease from Baseline in Total BMD



suppression. In the multicenter study, 121 patients with documented CCR5- tropic infection were randomized to once-daily regimens with either ATV/r plus maraviroc or ATV/r plus TDF/FTC. The primary endpoint was viral load <50 copies/mL. At 48 weeks, this endpoint was achieved in 74.5% of those taking the NRTI-sparing regimen and 83.6% of those taking the regimen containing NRTIs. The proportion with a viral load <400 copies/mL was 89.8% and 86.9%, respectively. Both rates were similar to those reported at 24 weeks. The authors, led by Dr. Simon Portsmouth, a researcher with Pfizer, reported that creatinine clearance, although stable in the NRTIsparing regimen, declined in those randomized to the regimen containing TDF/FTC. However, the rate of serious adverse events between groups was similar and more patients in the NRTI-sparing arm had treatment-related grade 3 or 4 adverse events (18 vs. 11). Still, due to the consistency of viral suppression over 48 weeks in the NRTIsparing arm, the authors indicated that this approach may be viable when an NRTI-sparing approach is attractive.

Conclusion

As viral suppression can now be reliably achieved with a variety of effective regimens, the attention paid to the relative safety of antiretroviral therapy has been intensified by the aging of HIV patient populations. In data from the San Francisco registry reported at the 2011 IAS meeting, the proportion of patients over the age of 50 living with AIDS had climbed above 50% for the first time. The authors of that study, led by Dr. Susan Scheer, San Francisco Department of Public Health, California, predicted that the trend will continue and suggested that the growing population of older persons with HIV will place new challenges on medical care. Pls have a well established history of potency leading to long-term HIV suppression. They remain well poised to continue to be part of strategies to sustain HIV control. Clinical studies are on-going to evaluate how these drugs can slow or circumvent the risks associated with accelerated aging.