Treatment Response among HIV Patients Co-enrolled in the Etravirine (ETR) and Raltegravir (RAL) Expanded Access Programs (EAPs) at Kaiser Permanente

H-1263

48th Annual ICAAC/46th Annual IDSA Joint Meeting October 25-28, 2008 Washington, DC

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Background

- In HIV treatment-experienced patients, DHHS guidelines recommend the inclusion of at least two, and preferably three, fully active antiretroviral agents when constructing drug regimens.
- Etravirine (ETR) is a next-generation non-nucleoside reverse transcriptase inhibitor (NNRTI) with demonstrated virologic activity against NNRTI resistant virus as well as a favorable safety profile.²⁻³
- Raltegravir (RAL) is an oral integrase inhibitor with established safety, efficacy, and patient tolerability. As a member of a novel drug class, it is unexpected for treatmentexperienced patients to have significant resistance to RAL.4-5
- In the clinic setting, the concurrent ETR and RAL Expanded Access Programs (EAPs) provided an opportunity to examine the efficacy and safety of ETR + RAL + background therapy (BT) in treatment-experienced patients.

Objectives

Primary Objective

 To assess the virologic effect of ETR + RAL + BT in HIV-1 infected treatmentexperienced patients.

Secondary Objectives

- To study the immunologic effect of ETR + RAL + BT in HIV-1 infected treatmentexperienced patients.
- To evaluate the effect of cumulative and baseline ETR resistance on virologic outcomes.
- *Cumulative resistance defined as baseline resistance + all available historic resistance data.
- To summarize the safety profile of ETR + RAL + BT.

Methods

Inclusion Criteria

- Patients initiating ETR and RAL simultaneously via EAP enrollment at Kaiser Permanente Northern and Southern California.
- Key EAP inclusion criteria included:
 - *Patient has limited or no treatment options due to virologic resistance or intolerance to multiple antiretroviral regimens.
 - Documented resistance to at least 1 drug in each of the 3 classes of oral ARVs (NRTI, NNRTI, and PI) by genotype or phenotype testing.
 - Intolerance is defined as having had a clinically significant adverse event which in the opinion of the investigator provides a contraindication to the use of any drug in that class.
 - *Patient has experience to at least 3 antiretroviral classes (NRTI, NNRTI, and PI).
 - *Patient is not achieving adequate virologic suppression on his/her current regimen and at risk of clinical or immunologic progression.
- *Patient is unable to use currently approved NNRTIs due to resistance (primary or acquired) and/or intolerance.
- *Patient has not received RAL or any other integrase inhibitor prior to EAP enrollment.

Methods, continued

Study Design

- A multicenter, retrospective study evaluating patients concurrently enrolled in ETR and RAL EAPs at 10 Kaiser Permanente HIV clinics in Northern and Southern California.
- •HIV-1 RNA, CD4 cell count, LFTs, serum creatinine, and lipid panel were collected at Screening, Baseline, Weeks 4, 12, and 24.
- All available genotype tests prior to ETR + RAL + BT initiation were collected for study analysis.
- ●ETR resistance was calculated utilizing the 2008 ETR weighted score method⁶. The 17 ETR mutations included were:
 - *V90I, A98G, L100I, K101E, K101H, K101P, V106I, E138A, V179D, V179F, V179T, Y181C, Y181I, Y181V, G190A, G190S, M230L
- Baseline ETR mutation score was calculated for each patient using their most recent genotype, prior to initiating ETR + RAL + BT regimen.
- Cumulative ETR mutation score was calculated using the baseline genotype plus all available historic genotypes.

Results

| No. of patients enrolled, n | 53 |
|---|--------------------------------|
| Gender, male, n (%) | 50 (94%) |
| Mean age, years | 49 |
| Median baseline CD4 count, cells/mm ³ (IQR) | 171 (74-290.5) |
| Received boosted protease inhibitor (PI) as part of BT, n (%) | 47 (89%) |
| darunavir/ritonavir de novo use not de novo use | 44 (83%) 43 (81%) 1 (2%) |
| lopinavir/ritonavir de novo use not de novo use | 3 (6%) 1 (2%) 2 (4%) |
| atazanavir/ritonavir de novo use not de novo use | 1 (2%) 0 (0%) 1 (2%) |
| Received enfuvirtide as part of BT, n (%) de novo use not de novo use | 6 (11%) 4 (8%) 2 (4%) |
| Mean length of time with known HIV-1 positive diagnosis, years | 17 |
| Mean length of time on prior ARV therapy, years | 14 |
| Mean total number of past ARVs | 11 |
| Median number of primary PI mutations at baseline ⁷ | 2 |

Virologic and Immunologic Outcomes at Week 24 (Intent-To-Treat Analysis)

| HIV-1 RNA BLQ*, n (%) | 50 (94%) |
|--------------------------------------|---------------------------|
| Mean CD4 cell count change | +86 cells/mm ³ |
| *BLQ = Below Level of Quantification | |

Treatment Failures

 The 3 patients with virologic failure at Week 24 had drug non-compliance with their antiretroviral regimen noted in provider clinic notes during the course of the study.

Results, continued

Virologic Outcomes at Week 24 based on Baseline and Cumulative

| ETR Weighted | Number of Patients with HIV-1 RNA BLQ at Week 24, Based on: | | |
|---|--|--|--|
| Mutation Score | Baseline Resistance Assessment, n (%) | Cumulative Resistance Assessment, n (%) | |
| 0 - 2 Highest predicted response | 35/37 (94.6%) | 28/30 (93.3%) | |
| 2.5 - 3.5 <i>Intermediate predicted response</i> | 9/10 (90.0%) | 10/10 (100%) | |
| > 3.5 Reduced predicted response | 6/6 (100%) | 12/13 (92.3%) | |

Change in Baseline and Cumulative ETR Mutation Score Among **Treatment Failures**

| Subject Number | Baseline Resistance, ETR Mutation Score | Cumulative Resistance, ETR Mutation Score |
|-------------------|--|--|
| 08929 | None None 0 | |
| 02450 | None 0 | K101E, A98G 2.0 |
| 07817 | V179D, Y181C 3.5 | K101E, V179D, Y181C 4.5 |

Baseline or cumulative ETR resistance did not predict treatment failure in these 3 pa-

 Resistance testing was not performed for these 3 treatment failures due to suspected drug non-compliance.

Table 5. Adverse Events (AEs)

| Grade 1-2 Clinical AEs occurring in | n ≥ 1 patient, n (%) | |
|---|----------------------|--|
| Rash | 10 (19%) | |
| Diarrhea | 9 (17%) | |
| Nausea | 5 (9%) | |
| Abdominal pain | 2 (4%) | |
| Fatigue | 2 (4%) | |
| Grade 3-4 Clinical AEs occurring in | n ≥ 1 patient, n (%) | |
| None | N/A | |
| Grade 1-2 laboratory abnormalities | s*, n (%) | |
| Total cholesterol | 6 (11%) | |
| Liver function tests (LFTs) | 4 (8%) | |
| Serum creatinine | 2 (4%) | |
| Grade 3-4 laboratory abnormalities | s*, n (%) | |
| Liver function tests (LFTs) | 3 (6%) | |
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^{*}Treatment-emergent laboratory abnormalities

- No patients discontinued therapy due to adverse events.
- The 3 patients experiencing Grade 3-4 liver function tests had chronic hepatitis at baseline. These LFTs subsequently decreased in all patients.

Conclusions

- Among treatment-experienced patients receiving ETR + RAL + BT, over 90% of patients achieved HIV-1 RNA below the level of quantification after 24 weeks of therapy.
- All patients were NNRTI experienced, however prior to initiating ETR + RAL + BT, more than half of patients had an ETR mutation score ≤ 2. The majority of these patients had mutations not associated with ETR resistance such as K103N, V108I, and P225H.
- When comparing virologic outcomes by baseline versus cumulative resistance, the additional historical mutations obtained from the cumulative resistance did not appear to significantly change the observed response in this limited data set.
- •ETR resistance could not predict all treatment failures. Other factors such as medication compliance and baseline PI resistance were likely to affect virologic
- The high rate of virologic success in this study population can be contributed to the use of multiple active agents in the regimen, including ETR, RAL and/or a boosted PI.
- The combination of ETR + RAL + BT was a safe and tolerable antiretroviral regimen, with minimal adverse events.

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Acknowledgements

We would like to thank all the patients who participated in this trial for their time and efforts.

A special recognition is also given to the HIV/AIDS Research Trials (H.A.R.T.) Program staff for their contributions to this study.

Appreciation provided to Tibotec Therapeutics and Tibotec for their review of this poster. Financial support for printing was provided by Tibotec Therapeutics.

