

PREDICTORS OF VIROLOGIC FAILURE IN POSTPARTUM WOMEN ON ART IN PROMISE 1077HS

. The probabilities of VF were (see top panel of Figure 2):

Risa M. Hoffman¹, Meredith Warshaw², K. Rivet Amico³, Jose Pilotto⁴, Gaerolwe Masheto⁵, Jullapong Achalapong⁶, Elizabeth Machado⁷, Kulkanya Chokephaibulkit⁸, Geraldo Duarte⁹, Anne Coletti¹⁰, Renee Browning¹¹, Nahida Chakhtoura¹², Karin L. Klingman¹¹, and Judith S. Currier¹ for the PROMISE 1077HS Team

ABSTRACT

virial (ART) adherence can be challenging for postpartum women and may result in virologic failure (VF). We examined predictors of VF and viral re-sion in postpartum women randomized to continue ART in PROMISE 1077HS.

METHODS
Appropriate FMY- non-breastfeeding women with pre-ART CD4 cell counts 1-800 cellsimm* who started ART during pregnancy were randomized up to 42 days plate delively to continue or describes. Perform who started that T2001-T10014. The preferred register are LEPIRTY will TEPFTC.

ART Vari are appropriate was defined as 2 consortive was tools. T300-General artifacts of the application of the appropriate was defined as 2 consortive was loads. T300-General, after VF, FT of an appropriate was defined as instance, was defined as 2 consortive was loads. T300-General, after VF, FT of an appropriate was sentenced as relatively extracted and table for present and the table for the sentence as of contention of the table for the sentence as the content of the present of the table for the present of the table for the table

RESULTS
Arroy the DDD section condensed to confuse ART condense again at early are '1' years' (1972 32-3) and neclear 1211 424 cours till discharge from 1922 for the confuse of the confu

suppression was 0.37 by 48 weeks, 0.46 by 99 Weeks, we us or yet yet.

OCICCUISON

CHICAGO STATE OF ST

BACKGROUND

More than 1 1/2 million HIV-infected women become pregnant and deliver babies annually, and the majority of these women now receive antiretroviral therapy (ART) antepartum and are expected to continue lifelong AR after delivery¹. The benefits of postpartum ART were recently reported from PROMISE 1077HS; hor high rates of virologic failure (VF) were seen in this study².

Previous studies have shown high rates of nonadherence and VF among postpartum HIV-infected women^{3,4}. Predictors of poor adherence and VF have included younger age^{4,5}, nondisclosure/stigma⁶, and low levels of HIV, ART and prevention knowledge⁶.

The PROMISE 1077HS study design provides a unique opportunity to explore predictors of virologic failure among women randomized to continue ART postpartum, including whether self-report about missed doses of ART is an accurate measure of adherence and risk for VF.

METHODS

PROMISE 1077HS was an open-label, randomized clinical trial evaluating two strategies for the management of ART among postpartum women with high CD4 T-cell counts (>400 cells(mm²); continuing ART or discontinuing ART and restarting when clinically indicated (Figure 1). The preferred ART regimen supplied by the study was Lopinavir/Ritonavir (LPV/RTV) plus fixed dose combination Emtricitabine/Tenofovir (FTC/TDF) and was chosen because it was the preferred regimen for use in pregnancy according to DHHS guidelines at the time the study was designed. Women enter within 42 days of delivery.

Participants randomized to the discontinue ART arm re-started ART if they met any of the following criteria:

- 1) Developed an AIDS-defining/WHO Stage 4 illness Had a confirmed CD4+ T-cell count <350 cells/mm
- Developed a clinical condition (other than pregnancy) considered an indication for ART by country-specific guidelin otherwise required ART as determined by the clinical manage

For women who continued ART, viral load and self-reported adherence were collected every 12 weeks. For these analyses

- VF was defined as 2 consecutive viral loads >1 000 conies/ml_after 24 weeks on ART
- Viral re-suppression was defined as 2 consecutive viral loads ≤1,000 copies/mL after first VF.
- · Self-reported adherence was dichotomized as missing
- versus not missing any ART doses in the prior 4 weeks. Predictors of VF and re-suppression were examined using Cox proportional hazards regression, with adherence as a time-varying covariate. Other predictors were values at baseline.
- We compared regional differences among the US, Asia (China/Thailand), Africa (Botswana), and South America and the Caribbean (Argentina, Brazil, Haiti, Peru).
- · Exploratory analyses were performed comparing the probability of VF using a cut-off of 400 copies/mL

Of the 827 women randomized to continue ART, 825 had HIV-1 RNA and adherence data, of whom 802 were on ART for at least 24 weeks and were included in the analysis. Complete characteristics of women are included in Table 1.

Among these 802 women, median age at entry was 27 years (IQR 23-32) and median CD4 T-cell count 696 cells/mm3 (IQR 576-865)

compared to protease inhibitor-based therapy (8%).

- Participants were from South America/Caribbean (38.8%), Botswana (27.9%), Asia (24.9%), and the United States (8.4%).
- At entry, 13% of women reported using lobacco in the past year, and 3% reported drinking alcohol at least 1-2 times per week in the 30 days prior to entry. At entry, 3% of women reported drug use in the past year (cocaine, heroin, amphetamines,
- and/or marijuana). At entry 9% of women reported missing ART doses over the prior 4 days. This increased to
- 11% at 48 weeks and 12% at 96 weeks Of 175 women with VF, 139 had resistance data available. Of these, 12% failed with resistance to their current regimen. VF with resistance to current regimen was more common in women on non-nucleoside reverse transcriptase inhibitor-based therapy (86%)

TABLE 1. Characteristics of women included in the analysis of VF (N=802)

eliver babies annually, and the majority of and are expected to continue lifelong ART orted from PROMISE 1077HS: however.	Characteristic	Continued ART arm (N=802)
among postpartum HIV-infected women ^{3,4} , nondisclosure/stigma ⁶ , and low levels of	Region Botswana Argentina/Brazii/Haiti/Peru Thailand/China US	224 (27.9%) 311 (38.8%) 200 (24.9%) 67 (8.4%)
by to explore predictors of virologic failure whether self-report about missed doses of the self-report about missed doses of the self-report about missed doses of the self-report self-r	Race Asian A	201 (25.1%) 54 (87%) 122 (55.2%) 1 (0.1%) 224 (27.9%) 75 (9.4%) 5 (0.6%) 71 (9.9%) 1 (0.1%) 31 (3.9%) 4 (0.5%) 1 (1.6%)
FIGURE 1. PROMISE 1077HS study design – only women in the Continue ART arm	Age at entry Min-Max Median (Q1-Q3) WHO Clinical Stage at entry	16-47 27 (23-32)
(circled) were included in the analysis of virologic failure	Clinical Stage I Clinical Stage II	785 (97.9%) 16 (2.0%)
1917 Screened	CD4 T-cell count at entry (cells/mm³) Min-Max Median (Q1-Q3) # missing	340-1800 696 (576-860) 3
1653 Enrolled	Duration on ART at entry (months) Min-Max Median (Q1-Q3)	0.0-8.6 4.0 (2.6-5.3)
1 withdrew Randomization	Viral load at entry (copies/ml) <400 400-1,000 >1,000-100,000 >100,000	724 (90.3%) 36 (4.5%) 41 (5.1%) 1 (0.1%)
n=827 n=825	Duration of study follow-up (weeks) Min-Max Median (Q1-Q3)	24.7-285.4 128.3 (84.4-181.9)
CONTINUE ART (median follow-up = 2.35 years) 79 ((8%) discontinued follow-up permaturely 15% discontinued ART	Self-rated health at entry Excellent Very Good Good Fair Poor # missing	156 (19.5%) 255 (31.9%) 316 (39.5%) 70 (8.8%) 2 (1.5%) 3 (0.4%)

RESULTS

- 0.12 by week 48
- When using a more stringent definition of VF (400 copies/mL), the estimated probability increases: 0.16 by week 48, 0.25 by week 96, and 0.29 by week 144 (Figure 2, bottom panel).

 There were differences by region, with participants from South America and the Caribbean having the highest
- estimated probability of VF (Figure 3).
- In univariable regression (Table 2), self-report of any missed ART doses in the prior 4 weeks, younger age, region, and shorter duration of pre-entry ART were predictive of VF.
- In the final multivariable model for VF, significant predictors included missed ART doses within the prior 4 weeks, younger age, shorter duration of pre-entry ART, and region (South America/Caribbean) (Table 2).

FIGURE 2: Estimated probability of virologic failure after the first 24 weeks on study through 144 weeks of

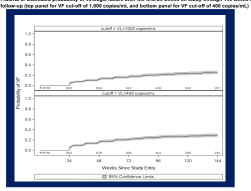


FIGURE 3: Estimated probability by region of VF through 144 weeks of follow-up

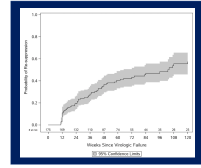


TABLE 2: Univariable and multivariable analyses for virologic failure among women who continued ART in PROMISE 1077HS (N=802)

	Univariable Analysis			Multivariable Model			
	Hazard	95% Confidence		Hazard	95% Confidence		
Variable	Ratio	Limits	p-value	Ratio	Limits	p-value	
Missed meds in last 4 weeks*	2.55	(1.89, 3.43)	<0.001	2.05	(1.48, 2.84)	<0.001	
Age at entry	0.96	(0.93, 0.98)	0.001	0.97	(0.94, 0.99)	0.01	
Pre-entry ART duration (months)	0.92	(0.85, 1.00)	0.05	0.91	(0.83, 0.99)	0.02	
Region			<0.001			0.07	
Botswana	1.07	(0.66, 1.74)	0.78	1.06	(0.65, 1.72)	0.82	
 Brazil/Haiti/Argentina/Peru 	2.06	(1.36, 3.10)	<0.001	1.69	(1.06, 2.52)	0.03	
United States	1.60	(0.87, 2.93)	0.13	1.30	(0.70, 2.43)	0.41	
· Thailand/China (reference)							
Baseline health**	0.98	(0.83, 1.15)	0.78				
ART including PI***	1.22	(0.83, 1.81)	0.31				
*Time-varying co-variate **Using a self-rated health scale (1=excellent, 5=poor) ***PI: Protease Inhibitor-based ART							

- . Figure 4 shows the probability of viral re-suppression among the 175 women with VF (two consecutive viral loads <1 000 copies/ml):
- 0.37 by 48 weeks
- 0.57 by 144 weeks
- When suppression was defined as 2 consecutive VLs <400 copies/mL, the probabilities were 0.33 by 48 weeks, 0.42 by 96 weeks, and 0.51 by 144 weeks.
- There were no statistically significant predictors of re-suppression, although analyses were limited by small

FIGURE 4: Estimated probability of re-suppression (2 consecutive VL <1.000 copies/mL) after first VF



CONCLUSIONS

- A simple 4-week ART recall question predicted first VF among women in PROMISE 1077HS.
 Regional differences in risk of VF have not been well characterized and require further study in regard to specific drivers of these differences, such as cultural, social, or health systems factors.
- Postpartum women who have VF are high risk for continued viremia; risk factors for persistent viremia
- require further study.

 Further research should explore strategies that can successfully support ART adherence for postpartum women.

REFERENCES

ACKNOWLEDGEMENTS

The 1077 PROMISE study team gratefully acknowledges the dedication and commitment of the 1652 participants without whom this study would not have been possible. Overall support for the International Maternal Pediatric Acidescent AIDS Clinical Trials (IMPAACT IDC), UMINAGESTE (IMPAACT SUBMISSES (IMPAACT IDC), UMINAGESTE (IMPAACT SUBMISSES (IMPAACT IDC), UMINAGESTE (IMPAACT SUBMISSES (IMPAACT IDC), UMINAGESTE (IMPAACT IDC), UMINAGES







IAST Databout for Monitorie Program Swarts (METC Goal, 2015; 1.6 Mp) Invasion senticid rappin contenting ATT Databout for Monitories (Program Swarts (METC) Goal, 2015; 1.6 Mp) Invasion senticid rappin contenting ATT Contenting ATT

ing ART Presented at CROI

