(up to 6 samples per subject).

HIV-1 RNA values over time.

Linear mixed effects regression models were used to assess plasma



sensitivity.

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(all data)



Figure 6. Distribution of individual subject plasma

HIV-1 RNA slopes

(Wooks 96-360)

▲ Week 96

Week 144

• Week 204

Week 252

Figure 7. Correlations between baseline and during-study

Baseline HIV-1 RNA (log₁₀ copies/mL)

Background

Persistent viremia can be detected in most HIV-1 infected patients on antiretroviral therapy despite suppression of plasma RNA to <50 copies/ml. Our previous studies have shown diverse antiretroviral regimens suppress plasma viremia to a new setpoint that correlates with pretherapy viremia1. These studies could not detect a significant decline in the viremia setpoint over 60-110 weeks o therapy (Figure 1). The current analysis

assesses plasma HIV-1 RNA levels in subjects

on suppressive therapy for 7 years, using a

real-time RT-PCR assay with single copy

Assav

used to test all samples.

copies/mL.

* An internally controlled real-time RT-PCR assay

with single-copy sensitivity (single-copy assay, SCA)2 was

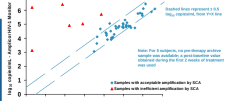
Study Entry Criteria

❖ In Study 720, antiretroviral-naive subjects received lopinavir/ritonavir (400/100 mg twice daily) with stayudine and lamiyudine twice daily for up to 7 years. Subjects remaining on study for 7 years (360 weeks) who never demonstrated detectable viremia (>50 or >400 copies/mL) during weeks 96-360 were included (Figure 2).

Subjects with comparable baseline assay results (SCA vs. Amplicor) were included in the longitudinal analysis (Figure 3). Figure 2. Sample selection · On-study samples were tested 100 subjects enrolled vearly from week 96 to week 360 in study 720

Figure 1. No decline in viremia setpoint over 60-110 47 subjects had no viral "blips . Between year 6 and year 7. weeks of therapy subjects were allowed to switch from stavudine to tenofovir DF3. oplification of baseline sample Primary analyses described above 4 HIV-1 RNA <assay limit (by SCA) excluded values after switch to 1 subject with no valid SCA secondary analyses assessed changes in HIV-1 RNA quality issues with each sample values after the switch to tenofovir using a 1-sample t test. 40 subjects included in

> Figure 3. Baseline results: SCA vs. Amplicor HIV-1 Monitor assay



Results Approximately 76% of samples

obtained between weeks 96 and 360 had detectable low-level viremia ranging from 1-39 copies/mL (median 1.7 copies/mL). *Based on all data from weeks 96-

360, a statistically significant decrease in plasma HIV-1 RNA level was (half-life=239 observed weeks. p=0.003, Figure 4).

* However, when data from week 96 were excluded, no decrease in plasma HIV-1 RNA over time was observed (half-life=971 weeks, p=0.53, Figure

* The distributions of individual subject slopes based on all data and on data from weeks 144-360 are

displayed in Figure 6. ❖Pre-therapy plasma HIV-1 RNA levels were significantly associated with week 96 levels (p=0.002) and week 252 levels (p=0.005) but not with levels at other timepoints (Figure 7).

No evidence of a change in plasma

HIV-1 RNA values was observed in 18 subjects who replaced stavudine with tenofovir DF.

Median value immediately prior to the switch was 1.37 copies/mL, compared to 1.33 copies/mL 12-24 weeks after the switch (p=0.82, 1-

Conclusions: These results are consistent with our prior finding that persistent viremia on treatment may originate from virus produced by cells that are infected before initiation of therapy. The apparent biphasic decay in persistent viremia implies that relatively short-lived cells contribute to viremia through 96-144 weeks, and very long-lived cells contribute thereafter. Testing of additional samples between weeks 60-120 may help to elucidate distinctions between phases of decay of persistent viremia.

971 weeks (0.53)

A HIV-1 RNA by SCA

A HIV-1 RNA by SCA

(value < assay limit)

Fitted model (all data)

t_{1/2} (p-value)

239 weeks (0.003

References: 1) Palmer S. et al. 12th Conference on Retroviruses and Opportunistic Infections, Boston, MA. February 2005, Abstract 163,

Note: Values obtained after switch to tenofovir DF not shown

2) Palmer S, et al. J Clin Microbiol 2003;41(10):4531-36.

Figure 4. Longitudinal plasma HIV-1 RNA values

Figure 5. Longitudinal plasma HIV-1 RNA values

A HIV-1 RNA by SCA

△ HIV-1 RNA by SCA (value < assay limit)

Fitted model (excluding Week 96)

(excluding week 96)

3) da Silva B, et al. 7th International Workshop on Adverse Drug Reactions and Lipodystrophy in HIV, Dublin, Ireland, November 2005. Abstract L957.

sample t test).

. Based on sample volumes available in this study, the lower limit of assay sensitivity ranged from 0.4 to 1.0