

Weight Change on F/TAF Versus Placebo: Using Common F/TDF Groups to Bridge Data Across Clinical Trials

David Glidden¹, Andrew Whiteman², Yuan Tian², Andrea Marongiu³, Joshua Gruber², Cal Cohen²

¹University of California San Francisco, San Francisco, CA, USA; ²Gilead Sciences, Inc., Foster City, CA, USA; ³Gilead Sciences Ltd, Uxbridge, UK

Conclusions

- This is the first analysis to compare changes in weight among people taking F/TAF versus placebo
 - Data from two Phase 3 randomized double-blind clinical trials in people without HIV were used, with F/TDF cohorts in each clinical trial acting as bridging cohorts
- Mean change in weight with F/TDF was consistent between the two trials, allowing comparison of the F/TAF and placebo cohorts
- Changes in weight were observed in the placebo group, providing important information on the weight increase expected over time in people without HIV assigned male at birth
- There were no statistically or clinically significant differences in mean weight gain with F/TAF compared with placebo
- Additionally, top percentiles of weight gain at Week 48 were similar with F/TAF and placebo
- Weight gain through 96 weeks in participants who switched from F/TDF to F/TAF at baseline was higher than for participants on placebo with no prior F/TDF exposure; this is consistent with a reversible weight-suppressive effect of F/TDF

Plain Language Summary

- Some people taking a human immunodeficiency virus (HIV) medication containing emtricitabine/tenofovir alafenamide (F/TAF) gain weight
 - Sometimes this is because the HIV medication is helping them become healthier, but at other times the reasons are not fully known
- This analysis looked at weight gain in two studies (one called iPrEx and the other called DISCOVER) of people who did not have HIV but who took medication to protect themselves from getting HIV (also known as PrEP or prevention)
- In iPrEx, one group was taking placebo (a pill that does not contain any medicine but that makes people think they are taking the study medication), while in DISCOVER, one group was taking F/TAF
- In both studies, the second group of people was taking emtricitabine/tenofovir disoproxil fumarate (F/TDF; the standard of care at the time of the iPrEx study). This common group helped compare one study to another
- The amount of weight gained by people without HIV who took F/TAF compared with people who took placebo was similar. Any difference was too small to be meaningful

Introduction

- Emtricitabine/tenofovir alafenamide (F/TAF) and emtricitabine/tenofovir disoproxil fumarate (F/TDF) are nucleoside/nucleotide reverse transcriptase inhibitors commonly used as the backbones for many guideline-recommended treatments for HIV¹
- F/TDF has previously been associated with reversible weight suppression,² whereas the association of F/TAF with weight gain is currently debated^{3,4}
- People with HIV often gain weight during treatment, and the cause of this can be multifactorial.⁵ This means that determining the direct impact of HIV medication on weight is challenging
- Comparison of F/TAF with placebo in people without HIV may give valuable insight into the effect of F/TAF on weight

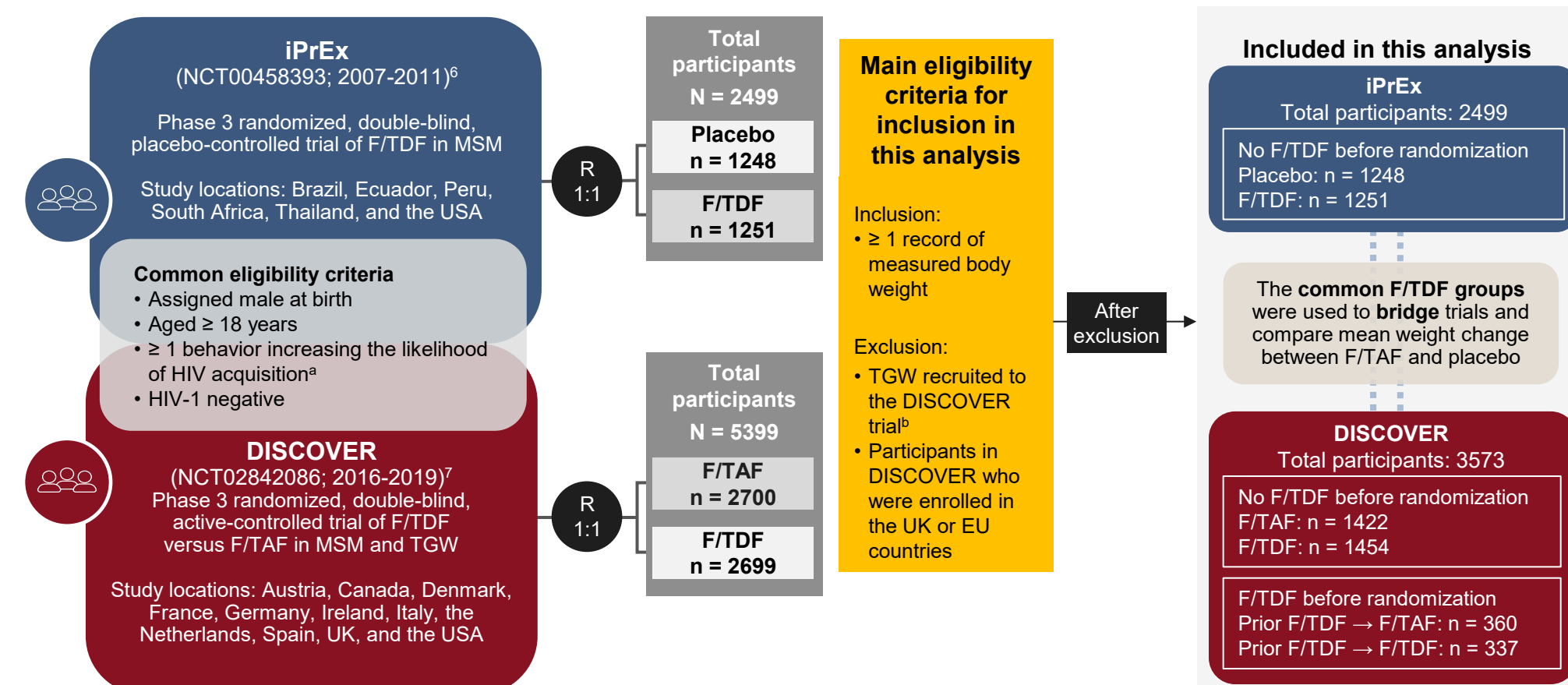
Objective

- To compare weight gain with F/TAF versus placebo among people without HIV, using data from two large Phase 3 randomized double-blind pre-exposure prophylaxis (PrEP) efficacy and safety clinical trials

Methods

- This was a retrospective analysis using data from two large Phase 3 randomized clinical trials of F/TDF versus placebo (iPrEx) and F/TDF versus F/TAF (DISCOVER) as PrEP in people without HIV (Figure 1)
- Both trials enrolled adults assigned male at birth who had an increased likelihood of HIV acquisition
- Body weight was measured during regular clinic visits
- To analyze the mean change in weight with F/TAF compared with placebo, common F/TDF groups in iPrEx and DISCOVER were used to bridge the trials
 - Participants in DISCOVER were stratified based on prior F/TDF use at baseline
- A linear mixed model, adjusting for baseline covariates and time-varying indicators of weight change-associated medication, was used to analyze mean weight change
 - Restricted cubic splines were used to model how effects of treatment may change over time
- Quantile regression was used to compare top percentiles of weight gain at Week 48, adjusting for the same baseline covariates (complete case analysis)

Figure 1. Study Design



⁶iPrEx: No condom use during anal intercourse with a male HIV-positive partner or a male partner of unknown HIV status during the last 6 months; anal intercourse with > 3 male sex partners during the last 6 months; exchange of money, gifts, shelter, or drugs for anal sex with a male partner during the last 6 months; sex with a male partner and diagnosis of a sexually transmitted infection during the last 6 months or at screening; sexual partner of an HIV-positive man with whom condoms were not consistently used in the last 6 months. DISCOVER: Condomless anal intercourse with ≥ 2 unique male partners in the past 12 weeks (partners must be either HIV-positive or unknown HIV status); documented history of syphilis or rectal gonorrhea or chlamydia in the past 24 weeks. ⁷TGW were not excluded from iPrEx in this study; further analyses of this subpopulation are ongoing. F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate; MSM, men who have sex with men; R, randomization; TGW, transgender women.

Results

Baseline Characteristics

- Participants in DISCOVER (and the subgroup of participants in DISCOVER with baseline F/TDF) were more likely to be White, aged ≥ 36 years, with a body mass index of ≥ 27 kg/m² and higher use of medications associated with weight gain, compared with participants in iPrEx (Table 1)
 - Mean alanine aminotransferase levels, estimated glomerular filtration rate, and blood glucose levels (non-fasting) were similar in both trials

Table 1. Baseline Demographics and Clinical Characteristics

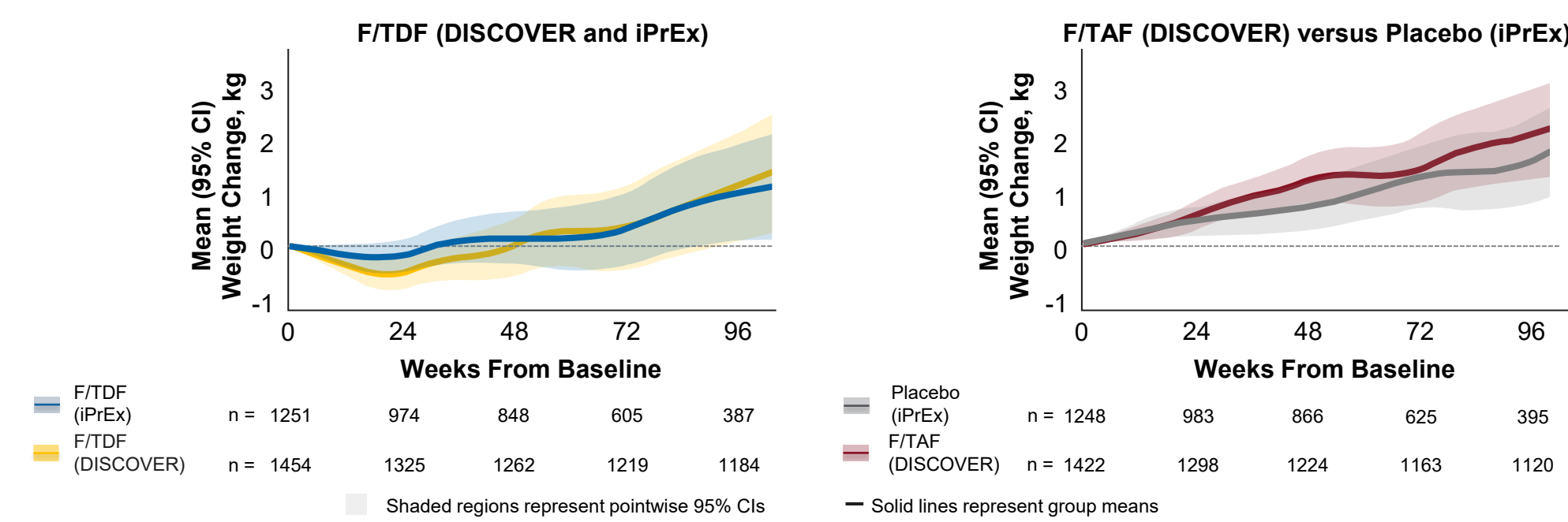
	F/TDF vs Placebo (iPrEx)		F/TAF vs F/TDF (DISCOVER)	
	All Participants N = 2499	All Participants N = 3573 ^a	All Participants N = 3573 ^a	Receiving F/TDF Before Baseline n = 697 ^b
Age, years, mean (SD)	27.1 (9)	36.1 (11)	37.5 (11)	
Race or ethnicity, n (%)				
White	431 (17)	2825 (79)	579 (83)	
Black	214 (9)	405 (11)	55 (8)	
Other ^c	1854 (74) ^c	343 (10)	63 (9)	
Hispanic or Latine	1806 (72)	871 (24)	121 (17)	
Medications associated with weight change, ^d n (%)				
Gain	63 (3)	516 (14)	131 (19)	
Loss	28 (1)	287 (8)	62 (9)	
Unknown	31 (1)	246 (7)	72 (10)	
Body weight, kg, mean (SD)	68.1 (14)	85.0 (18)	84.6 (18)	
BMI, kg/m ² , mean (SD)	23.8 (4)	27.0 (5)	26.7 (5)	
Alanine aminotransferase level, U/L, mean (SD)	26.5 (17)	28.34 (15)	29.9 (15)	
eGFR _{CG} , mL/min, mean (SD)	118.6 (23)	129.8 (37)	124.2 (36)	
Blood glucose level (non-fasting), mg/dL, mean (SD)	90.4 (14)	94.9 (19)	92.8 (18)	

^aParticipants in DISCOVER receiving F/TDF before baseline are counted in both DISCOVER columns and were subsequently randomized to continue with F/TDF or switch to F/TAF. ^bIncludes American Indian or Alaska Native; Asian; Native Hawaiian or Other Pacific Islander; other race; and participants with Not Permitted designation (when local regulators did not allow collection of race/ethnicity information). ^cThe majority of participants in the iPrEx study self-reported as "mixed race or other". ^dAntidepressants/psychoneuroleptics, antidiabetics, antiepileptics, antihistamines, antiulceratives, contraceptives, corticosteroids, beta-androgenic blockers, insulin. BMI, body mass index; eGFR, estimated glomerular filtration rate by Cockcroft-Gault equation; F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate.

No Significant Difference in Estimated Mean Weight Change From Baseline Between F/TAF and Placebo in Participants With No Prior Use of F/TDF

- Estimated mean weight change from baseline with F/TDF was similar in DISCOVER and iPrEx (Figure 2)
- There was no statistically significant difference across the 96-week period in mean weight change from baseline between participants receiving F/TAF or placebo at any time (P = 0.10) (Figure 2), including at Week 24, 48, 72 and Week 96 (Table 2)
- The results of sensitivity assessments (not shown) were highly consistent with primary findings. These assessments included per protocol-type analyses and robust estimation strategies

Figure 2. Estimated Mean Weight Change From Baseline in iPrEx and DISCOVER Participants With No Prior Use of F/TDF



Data were adjusted for baseline age, country of enrollment, diabetes status, glucose (non-fasting), kidney function (estimated glomerular filtration rate), liver function (alanine aminotransferase level), race (White/Black/Other), height, use of medications associated with weight change (both baseline and trial-incident), and weight. F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate.

Table 2. Mean Weight Change Contrasts (F/TAF > Placebo) At Landmark Timepoints (With No Prior Use of F/TDF)

	Difference in Mean (95% CI) Weight Change, kg
Week 24	+0.07 (-0.25, 0.39)
Week 48	+0.44 (-0.04, 0.92)
Week 72	+0.13 (-0.51, 0.77)
Week 96	+0.53 (-0.28, 1.34)

Data were adjusted for baseline age, country of enrollment, diabetes status, glucose level (non-fasting), kidney function (estimated glomerular filtration rate), liver function (alanine aminotransferase level), race (White/Black/Other), height, use of medications associated with weight change (both baseline and trial-incident), and body weight. F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate.

- Proportions of individuals experiencing large increases in body weight over 48 weeks were similar with F/TAF versus placebo (Table 3)

Table 3. Top Percentiles^a (95% CI) of Weight Change From Baseline at Week 48 in Participants With No Prior Use of F/TDF Who Were Receiving Placebo (iPrEx) or F/TAF (DISCOVER)

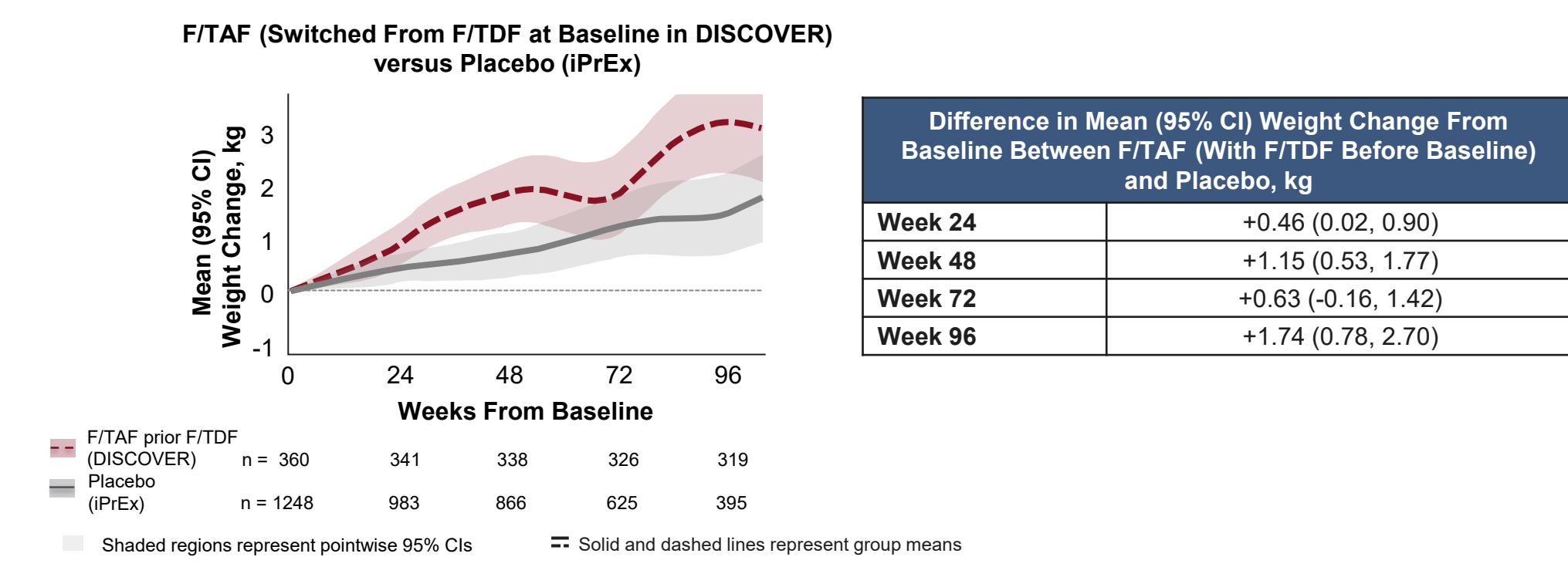
	Placebo (iPrEx), kg	F/TAF (DISCOVER), kg
80th	4.2 (3.7, 4.7)	4.1 (3.8, 4.4)
85th	5.0 (4.4, 5.7)	4.6 (4.2, 5.1)
90th	5.7 (5.1, 6.3)	5.8 (5.3, 6.2)
95th	6.9 (5.8, 8.1)	7.9 (7.0, 8.7)
99th	11.6 (9.0, 14.2)	11.1 (10.1, 12.2)

^aEstimated percentiles were adjusted for the same baseline covariates as in the primary analysis. F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate.

Average Weight Gain (Relative to Placebo) in Participants Taking F/TAF With Prior Use of F/TDF at Baseline Was Consistent With a Reversible Weight-Suppressive Effect of F/TDF

- Participants switching from F/TDF to F/TAF at baseline in DISCOVER experienced a general increase in mean (95% CI) body weight compared with those receiving placebo who had not been on F/TDF at baseline (Figure 3)

Figure 3. Estimated Mean Weight Change in Participants Taking F/TDF at Baseline Who Switched to F/TAF



Data were adjusted for baseline age, country of enrollment, diabetes status, glucose level (non-fasting), kidney function (estimated glomerular filtration rate), liver function (alanine aminotransferase level), race (White/Black/Other), height, use of medications associated with weight change (both baseline and trial-incident), and body weight. F/TAF, emtricitabine/tenofovir alafenamide; F/TDF, emtricitabine/tenofovir disoproxil fumarate.

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Correspondence: David Glidden, david.glidden@ucsf.edu.