Long-term Safety Profile of Tenofovir Alafenamide in Chronic Hepatitis B Patients: Final 8-Year Results of 2 Phase 3 Studies

Young-Suk Lim¹, Henry Lik Yuen Chan², Kosh Agarwal³, Patrick Marcellin⁴, Maurizia R Brunetto⁵, Wan-Long Chuang⁶, Harry LA Janssen^{7,8}, Scott K Fung⁹, Namiki Izumi¹⁰, Maciej S Jablkowski¹¹, Frida Abramov¹², Hongyuan Wang¹², Leland J Yee¹², John F Flaherty¹², Calvin Pan¹³, Dr Shalimar¹⁴, Wai-Kay Seto¹⁵, Edward J Gane¹⁶, Maria Buti^{17,18}

¹Asan Medical Center, University of Ulsan College of Medicine, the Chinese University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Hong Kong; ⁴ Hepatology Department, Hôpital Beaujon, APHP, INSERM, University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Hong Kong; ⁴ Hepatology Department, Hôpital Beaujon, APHP, INSERM, University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Paris, Clichy, France; ⁵ Azienda Ospedaliero-University of Paris, Clichy, France; ⁵ Azienda, Pisa, Italy; ⁶ Kaohsiung Medical University of Paris, Clichy, France; ⁵ Azienda, Pisa, Italy; ⁶ Kaohsiung Medical University, Italy; ⁶ Kaohsiung Medical Heat City, Canada; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino Hospital, Tokyo, Japanese Red Cross Musashino Hospital, Tokyo, Japanese, University of Lodz, Lodz, Poland; ¹² Gilead Sciences, Inc., Foster City, CA, USA; ¹³ NYU and Hepatology, Japanese Red Cross Musashino Hospital, Tokyo, Japanese Red Cross Musashino, Netherlands; ⁹ University of Lodz, Lodz, Poland; ¹⁴ Belth Network, Toronto, Canada; ¹⁰ Department of Medical University of Lodz, Lodz, Poland; ¹⁴ Belth Network, Toronto, Canada; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Netherlands; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Department of Gastroenterology, Japanese Red Cross Musashino, Poland; ¹⁰ Departm Langone Health, New York University of Hong Kong; ¹⁶Auckland, New Zealand; ¹⁷Hospital Universitario Vall d'Hebron, Barcelona, Spain; ¹⁸CIBEREHD del Instituto Carlos III, Madrid, Spain (18) and the set and School of Clinical Sciences, New Delhi, India; ¹⁹Department of Medicine, New Zealand; ¹⁷Hospital Universitario Vall d'Hebron, Barcelona, Spain; ¹⁸CIBEREHD del Instituto Carlos III, Madrid, Spain (18) and the set and School of Clinical Sciences, New Delhi, India; ¹⁹Department of Medicine, New Zealand; ¹⁹Auckland, New Zealand; ¹⁹Auckland,

Key Findings

- Through 8 years of treatment, no new safety signals were identified for TAF
- Increases in fasting lipids and body weight were observed, which plateaued after year 5
- Minimal declines in eGFR_{cc} and in hip and spine BMD occurred among patients treated with TAF over 8 years
- Among those treated with DB TDF, the early declines in renal function and BMD steadily improved after switching to TAF

Conclusions



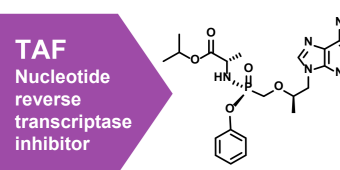
Over 8 years, treatment with TAF was safe and well tolerated by patients with chronic HBV; switching from TDF to TAF after 2 or 3 years resulted in improvements in renal and bone safety parameters



These results provide further support for use of TAF as a preferred treatment for chronic **HBV** infection

Introduction

- Hepatitis B virus (HBV) infection affects approximately 296 million individuals globally and is associated with cirrhosis, hepatic decompensation, and hepatocellular carcinoma (HCC)^{1,2}
- Tenofovir alafenamide (TAF):
 - A novel tenofovir (TFV) prodrug with greater plasma stability, enhanced hepatic delivery of active drug (TFV-diphosphate) to



- hepatocytes, and lower circulating levels of TFV compared with tenofovir disoproxil fumarate (TDF)³⁻⁶
- In comparative trials, TAF demonstrated noninferior antiviral efficacy and improved renal and bone safety compared with TDF at weeks 48 and 96 among viremic and virally suppressed hepatitis B e antigen (HBeAg)-negative and HBeAg-positive patients7-9
- Patients from these trials were eligible to receive openlabel (OL) TAF, and favorable renal and bone safety were observed during a 5-year interim analysis¹⁰

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Acknowledgments: We extend our thanks to the patients and their families and all participating investigators. These studies were funded by Gilead Sciences, Inc. Medical writing support was provided by Charlotte Bavley, PhD, of AlphaScientia, a Red Nucleus company, and was funded by Gilead Sciences. Inc.

Disclosures: HLYC served as an advisor for Aligos, GSK, Gilead Sciences, Inc., Roche, Vaccitech, Vir Biotechnology, Inc., and Virion Therapeutics and reports speaker fees from Gilead Sciences, Inc., Roche, and Viatris; KA served as a speaker, consultant, and/or advisory board member for Assembly Biosciences, Aligos Therapeutics, Arbutus Biopharma, Boehringer Ingelheim, Bristol Myers Squibb, Drug Far, Gilead Sciences, Inc., GSK, Janssen, Roche, Saigmet, and Sobi, and his institution received research support from Gilead Sciences, Inc.; PM received grants from AbbVie, Aligos, Assembly Biosciences, Bristol Myers Squibb, Gilead Sciences, Inc., Humedics, Madrigal, Novo Nordisk, Pfizer, Roche, and Intercept; **MRB** reports speaker and consultancy fees from AbbVie, Eisai-MSD, Gilead Sciences, Inc., Janssen, and Roche; HLAJ received research grants from Gilead Sciences, Inc., GSK, Janssen, Roche, and Vir Biotechnology, Inc., and served as a consultant for Aligos, Antios, Eiger, Gilead Sciences, Inc., GSK, Janssen, Roche, and Vir Biotechnology, Inc.; SKF served as an advisor for AbbVie, Gilead Sciences, Inc., Novo Nordisk, and Pfizer, reports speaker fees from AbbVie, Gilead Sciences, Inc., and Lupin, and received research support from Gilead Sciences, Inc.; FA, HW, LJY, and JFF are employees and stockholders of Gilead Sciences, Inc.; CP received research support from Gilead Sciences, Inc.; WKS served as an advisor for Abbot, AbbVie, and Gilead Sciences, Inc., and reports speaker fees from AbbVie, AstraZeneca, Gilead Sciences, Inc., and Mylan; EJG served as an advisor for AbbVie, Aligos, Arbutus, Gilead Sciences, Inc., Janssen, Roche, Vir Biotechnology, and Virion Therapeutics; **MB** reports speaker fees, research support, and consulting fees from AbbVie, Gilead Sciences, Inc., and Janssen; YSL, WLC, NI, MSJ, and DS report no conflicts of interest.

Objective

Methods

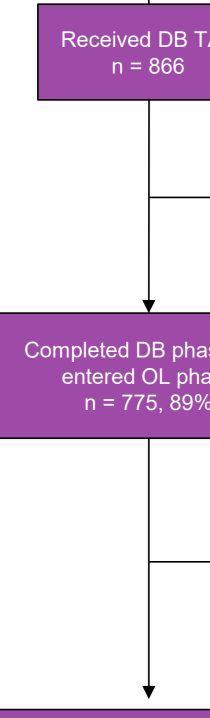
Study Design

Key inclusion criteria

- HBV DNA ≥20,000 IU/mL
- ALT >60 (males) and >38 U/L (females)
- With/without compensated cirrhosis
- eGFR_{CG} ≥50 mL/min
- Treatment naïve or reatment experience

- Two Phase 3, randomized, DB, active-controlled trials
 - Study 108 (NCT01940341; N = 425 originally randomized and treated): HBeAgnegative patients
 - Study 110 (NCT01940471; N = 873 originally randomized and treated): HBeAgpositive patients
- After completion of the DB phase, all patients were eligible to receive OL TAF through year 8
- Safety endpoints: • Cumulative adverse events (AEs), serious AEs, and graded laboratory abnormalities during the OL phase
 - **Bone:** changes in hip and spine bone mineral density (BMD) by dual energy X-ray absorptiometry and serum markers of bone turnover
 - **Renal:** changes in estimated glomerular filtration rate by Cockcroft-Gault (eGFR_{cc}) and quantitative urinary markers of tubular proteinuria—ratio of retinol-binding
 - protein (RBP) to creatinine (Cr) and ratio of β_2 -microglobulin (β_2 M) to Cr Metabolic parameters: changes in fasting lipids, including total cholesterol (TC), high-density lipoprotein cholesterol (HDL), low-density lipoprotein cholesterol (LDL),

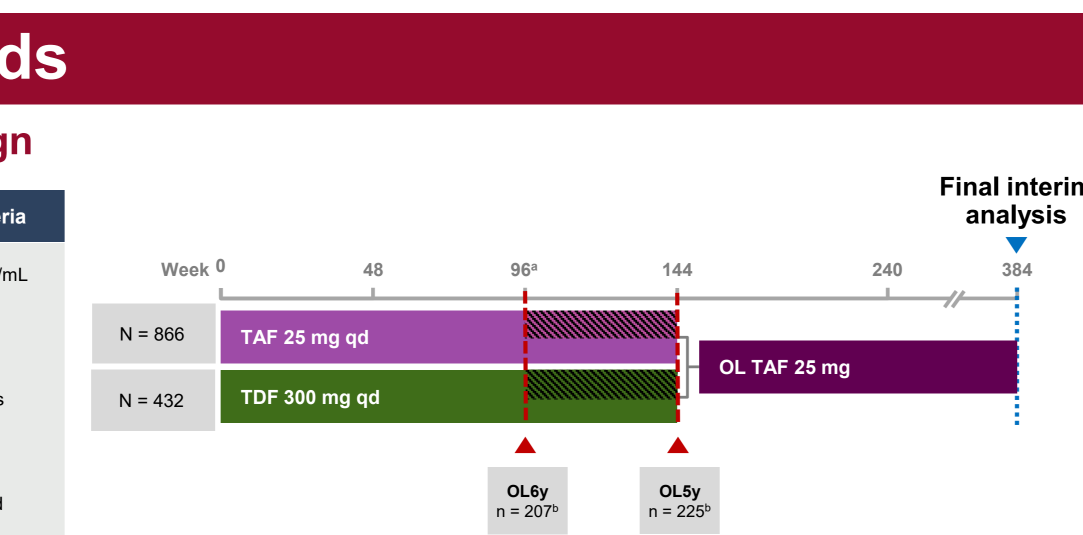
Results **Patient Disposition**



Completed C n = 647, 7

^aMetastatic HCC. HBsAg, hepatitis B surface antigen — 974 of 1298 (75%) patients completed the OL phase — Overall excellent patient retention with very few patients (n = 11; <1%) discontinuing OL TAF due to an AE

— To evaluate safety outcomes at year 8 (week 384) in patients with HBeAg-negative and HBeAg-positive chronic HBV treated with TAF (double blind [DB] and OL) or TDF (DB) followed by TAF (OL)



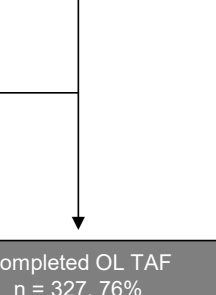
^aAmendment 3 enacted to extend DB to week 144 and OL to week 384 (year 8). Shaded areas represent patients who rolled over to OL TAF at week 96 (OL6y) or week 144 (OL5y) based upon the timing of the amendment. Patients who received DB TDF and switched to TAF. ALT. alanine aminotransferase: eGFR., estimated glomerular filtration rate by Cockcroft-Gault: gd

• Methods for Studies 108 and 110 are described elsewhere^{8,9}

triglycerides (TG), and TC/HDL ratio; change in body weight

Randomized and treated: N = 1298

٨F	N = 89	Discontinued DB TAF or TDF	N = 49	Receiv	od F
	30	Withdrew consent	21		= 43
	14	Lost to follow-up	6		
	8	Investigator's discretion	8		Т
	13	Adverse events	4		
	9	Pregnancy	2		
	6	Noncompliance with study drug	3		
	3	Protocol-specified criteria for withdrawal	1		
	1	Death	2		
	4	HBsAg seroconversion	0		
	1	Lack of efficacy	1		↓
	0	Protocol violation	1		•
se				entered n = 3	
se	N = 128	Discontinued OL TAF	N = 55		
se		Discontinued OL TAF Withdrew consent	N = 55 27		
se	N = 128 65 19	Withdrew consent			
Se	65	Withdrew consent Lost to follow-up	27		
e	65 19	Withdrew consent	27 9		
se	65 19 12	Withdrew consent Lost to follow-up Investigator's discretion	27 9 5		
Se	65 19 12	Withdrew consent Lost to follow-up Investigator's discretion Adverse events	27 9 5 3		
e	65 19 12 8 7	Withdrew consent Lost to follow-up Investigator's discretion Adverse events Protocol-specified criteria for withdrawal	27 9 5 3		
Se	65 19 12 8 7	Withdrew consent Lost to follow-up Investigator's discretion Adverse events Protocol-specified criteria for withdrawal Pregnancy	27 9 5 3		
	65 19 12 8 7 6 4	Withdrew consent Lost to follow-up Investigator's discretion Adverse events Protocol-specified criteria for withdrawal Pregnancy HBsAg seroconversion	27 9 5 3 3 1 4	n = 3	382,
AF	65 19 12 8 7 6 4	Withdrew consent Lost to follow-up Investigator's discretion Adverse events Protocol-specified criteria for withdrawal Pregnancy HBsAg seroconversion Noncompliance with study drug	27 9 5 3 3 1 4		382,



	TAF n = 866	TDF→TAF n = 432
Age, y, mean (SD)	40 (11.8)	41 (12.3)
Male, n (%)	544 (63)	275 (64)
Asian, n (%)	687 (79)	333 (77)
White, n (%)	167 (19)	87 (21)
Black or African American, n (%)	7 (1)	6 (1)
Other race, n (%)	5 (1)	6 (1)
BMI, kg/m², median (Q1, Q3)	24 (21, 27)	24 (22, 27)
Body weight, kg, median (Q1, Q3)	67 (57, 76)	66 (58, 78)
HBeAg positive, n (%)	569 (66)	290 (67)
ALT, U/L, median (Q1, Q3)	80 (56, 123)	80 (53, 130)
FibroTest score ≥0.75, n/N (%) ^b	76/846 (9)	42/421 (10)
eGFR _{cg} , mL/min, median (Q1, Q3)	106 (91, 125)	105 (90, 124)
Osteoporosis by spine BMD T-score, n (%) ^c	57 (7)	29 (7)
Osteoporosis by hip BMD T-score, n (%) ^c	12 (1)	2 (<1)
Diabetes mellitus, n (%)	57 (7)	29 (7)
Hypertension, n (%)	99 (11)	62 (14)
Hyperlipidemia, n (%)	76 (9)	44 (10)

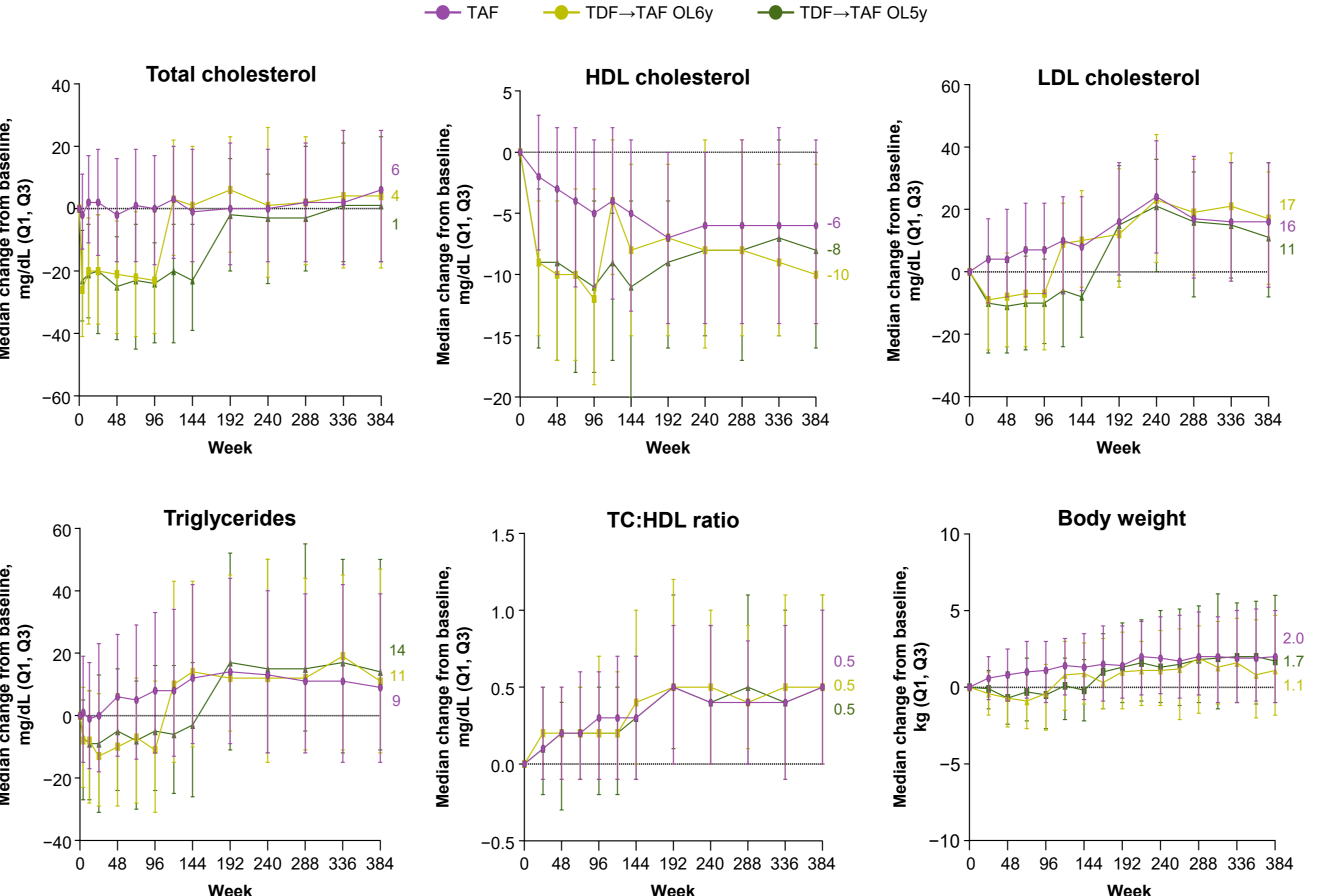
Patients, n or n/n (%)
Any AE
Grade 3 or 4 AE
Grade 3 or 4 AE related AE (1 patient each)
Serious AE
Serious AE related to st AE (1 patient each)
D/C due to AE AE (1 patient each)

Death⁵
HCC℃
Adverse events occurrin Headache Upper respiratory tra Nasopharyngitis Hypertension Arthralgia Cough Back pain
mong patients in the OL saf

Grade 3 or 4 Laboratory Abnormalities Occurring in ≥ 2% of Patients^a

Patients, n or n/n (%)	TAF n = 775	TDF→TAF n = 382
Maximum postbaseline toxicity grade	185/772 (24)	93/378 (25)
Amylase	15/772 (2)	10/377 (3)
Creatine kinase	11/772 (1)	8/377 (2)
Fasting cholesterol ^₅	11/767 (1)	11/373 (3)
Fasting LDL cholesterol ^b	45/760 (6)	30/373 (8)
Increased fasting glucose ^b	12/767 (2)	7/373 (2)
Fasting triglycerides	5/767 (1)	7/373 (2)
Urine occult blood ^b	26/772 (3)	12/377 (3)

Median Change in Fasting Lipids and Body Weight Over 8 Years

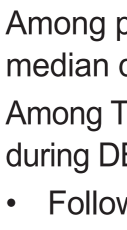


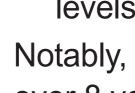
Baseline Demographics and Disease Characteristics^a

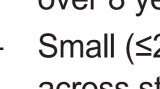
Open-Label Safety: Adverse Events^a

	TAF n = 775	TDF→TAF n = 382
	525 (68)	271 (71)
	60 (8)	27 (7)
ed to study drug	2 (<1) cerebrovascular accident; renal neoplasm	0
	97 (13)	49 (13)
study drug	4 (1) cerebrovascular accident; renal neoplasm; ALT increase; osteonecrosis	0
	9 (1) cardiopulmonary failure; myelodysplastic syndrome; HCC; pancreatic carcinoma; cerebrovascular accident; gamma-glutamyltransferase increased; osteonecrosis; osteoporosis; proteinuria	3 (1) tuberculosis; ascites; pemphigoid
	1 (<1)	0
	8 (1)	6 (2)
ing in ≥5% of patients	59 (8)	30 (8)
ract infection	55 (7) 52 (7) 37 (5) 41 (5) 28 (4) 34 (4)	27 (7) 23 (6) 26 (7) 23 (6) 27 (7) 23 (6)

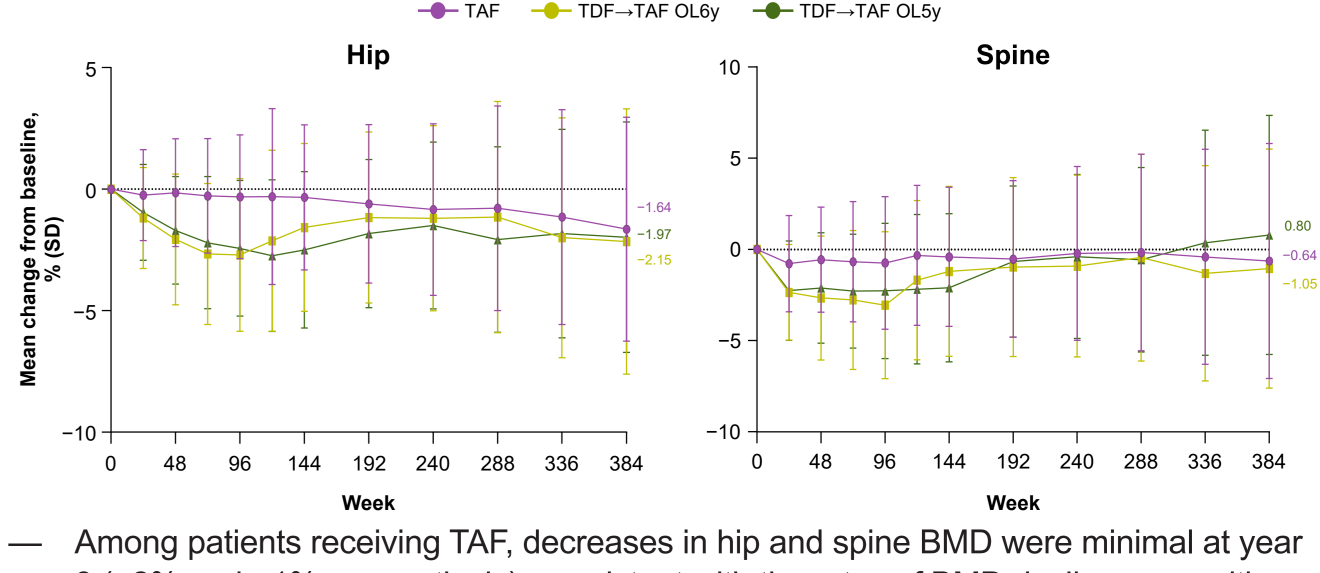
^aAmong patients in the OL safety analysis who received ≥1 dose of OL study drug (OL Safety Analysis Set); ^bTreatment-emergent death. There were 6 deaths in total (5 DB TDF 3); ^cA total of 14 HCC cases occurred during the OL phase, while overall, 21 patients developed HCC during the DB and OL phases of the study. D/C, discontinuation.

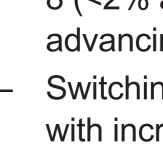


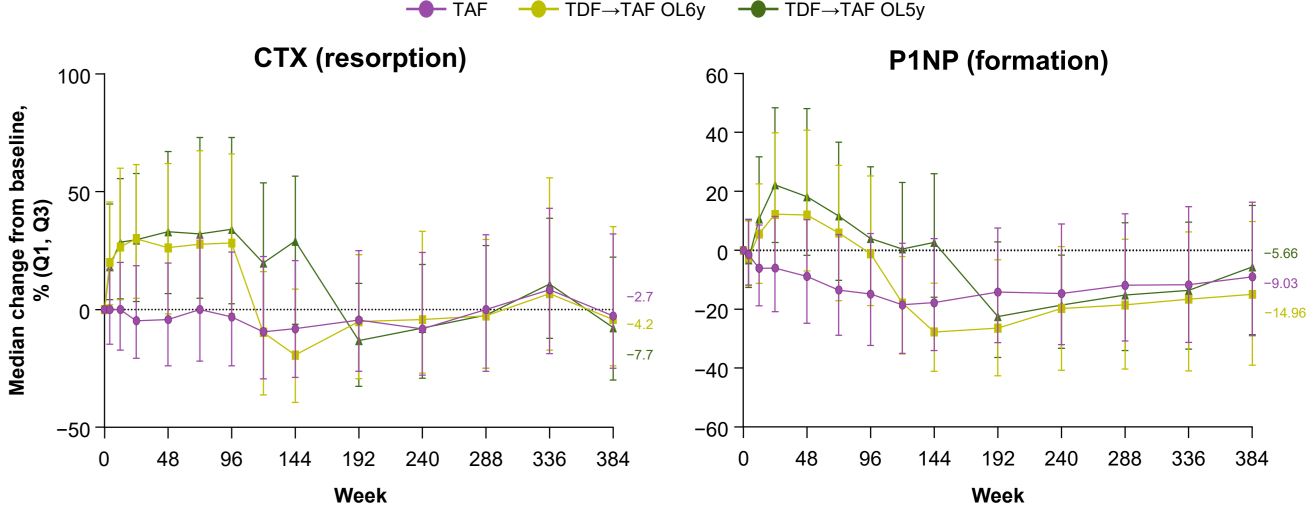


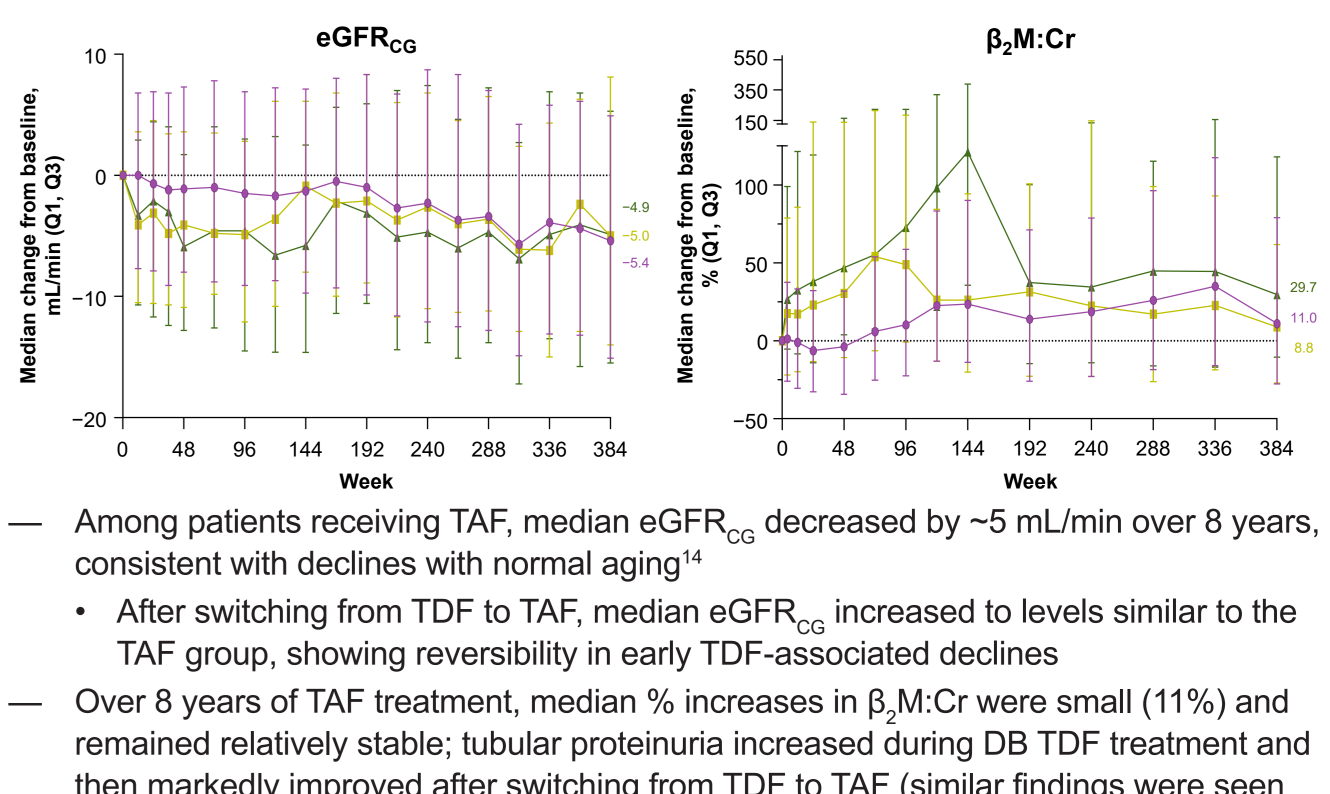












SAT-153

— Among patients receiving TAF, small median increases in TC, LDL, and TG and a small median decrease in HDL were observed

— Among TDF→TAF patients, modest decreases in TC, HDL, LDL, and TG were observed during DB TDF treatment, consistent with the known lipid-lowering effect of TDF^{11,12}

• Following the switch from TDF to TAF, levels of TC, HDL, LDL, and TG stabilized to levels observed in the TAF-only treatment group

— Notably, TC:HDL ratio, a marker of cardiovascular risk, increased minimally (≤0.5 fold) over 8 years in all groups

— Small (≤ 2.0 kg) median increases in body weight were seen after 8 years of treatment across study groups

Mean % Change in Hip and Spine BMD Over 8 Years

8 (<2% and <1%, respectively), consistent with the rates of BMD decline seen with advancing age¹³

— Switching from TDF to TAF at year 2 (week 96) or year 3 (week 144) was associated with increases in BMD, indicating that TDF-induced bone loss can be reversible

Median % Change in Bone Biomarkers Over 8 Years

CTX, C-terminal telopeptide of type 1 collagen; P1NP, N-terminal propeptide of type 1 procollagen.

Changes in Renal Parameters Over 8 Years

-- TDF \rightarrow TAF OL6y -- TDF \rightarrow TAF OL5y

then markedly improved after switching from TDF to TAF (similar findings were seen with RBP:Cr, data not shown)













