

# Safety, PK, and Efficacy of Low-Dose B/F/TAF in Children ≥2 Years Old Living With HIV

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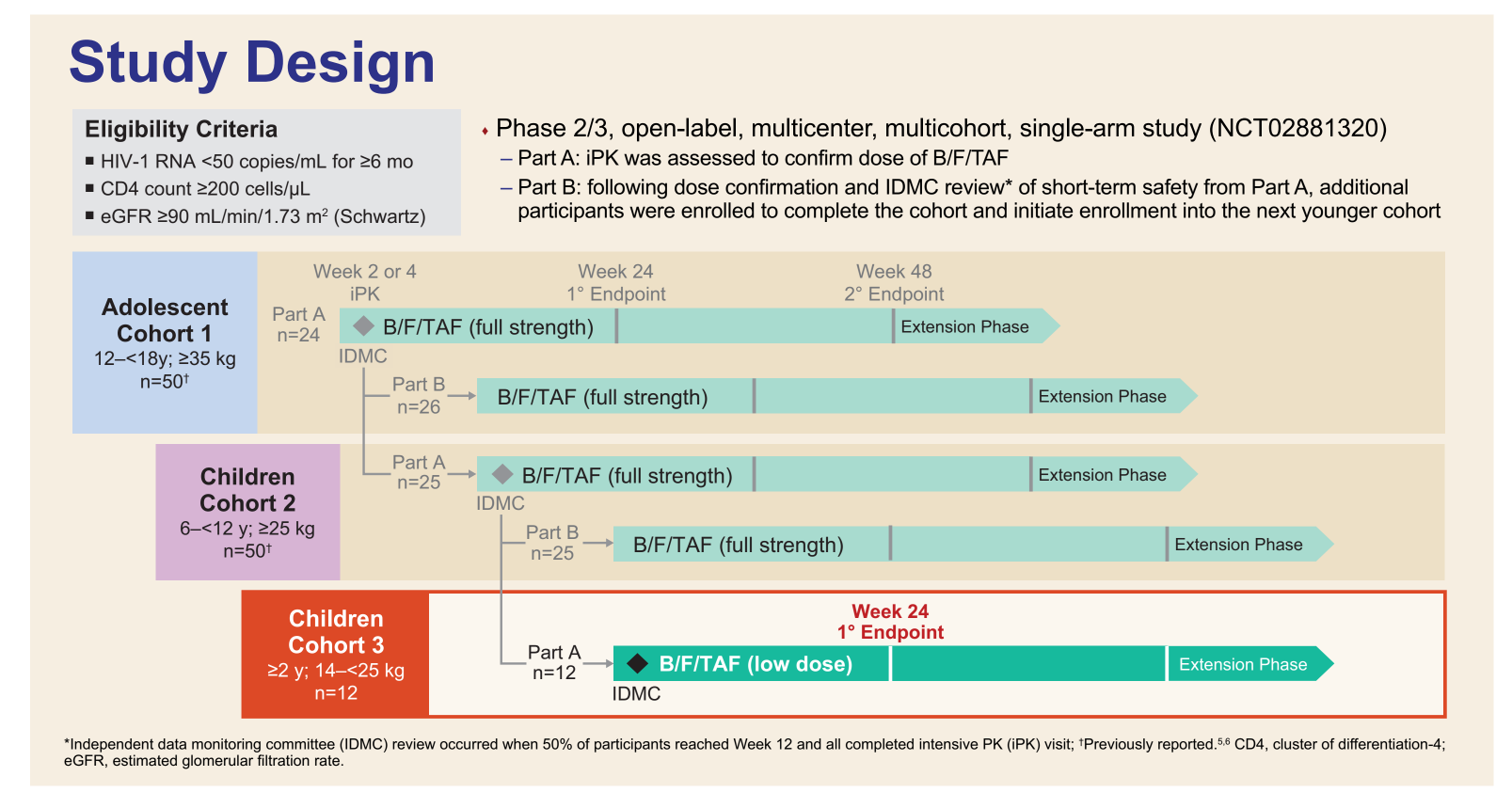
## Introduction

- Few antiretroviral (ARV) options exist for very young children living with HIV and no single-tablet regimen (STR) is used or approved for this population
- Bictegravir (BIC; B) is a novel, unboosted integrase strand transfer inhibitor (INSTI), with a high resistance barrier and low potential for drug-drug interactions
- BIC has been coformulated with emtricitabine (FTC; F) and tenofovir alafenamide (TAF) into a once-daily STR (B/F/TAF)
- B/F/TAF is approved for use and is a guideline-recommended regimen in children weighing ≥25 kg living with HIV¹-⁴
- ◆ B/F/TAF has been formulated as a low-dose STR for children aged
   ≥2 y and weighing 14—<25 kg</li>
   Low-dose STR strength is B/F/TAF 30/120/15 mg (60% of full-strength)
- STR)
- Can be taken without regard to food
- This is the 1st study to report the pharmacokinetics (PK), safety, and efficacy of B/F/TAF in young children aged ≥2 y living with HIV

# Objectives

- Primary: to determine the plasma PK of BIC, and evaluate the safety and tolerability of B/F/TAF through 24 wk of treatment in virologically suppressed adolescents (Cohort 1) and children (Cohorts 2 and 3) living with HIV
- Secondary: to evaluate the safety and tolerability of B/F/TAF for 48 wk, and its antiviral activity at 24 and 48 wk, in virologically suppressed adolescents (Cohort 1) and children (Cohorts 2 and 3) living with HIV
- Cohort 3 is the focus of the present analyses

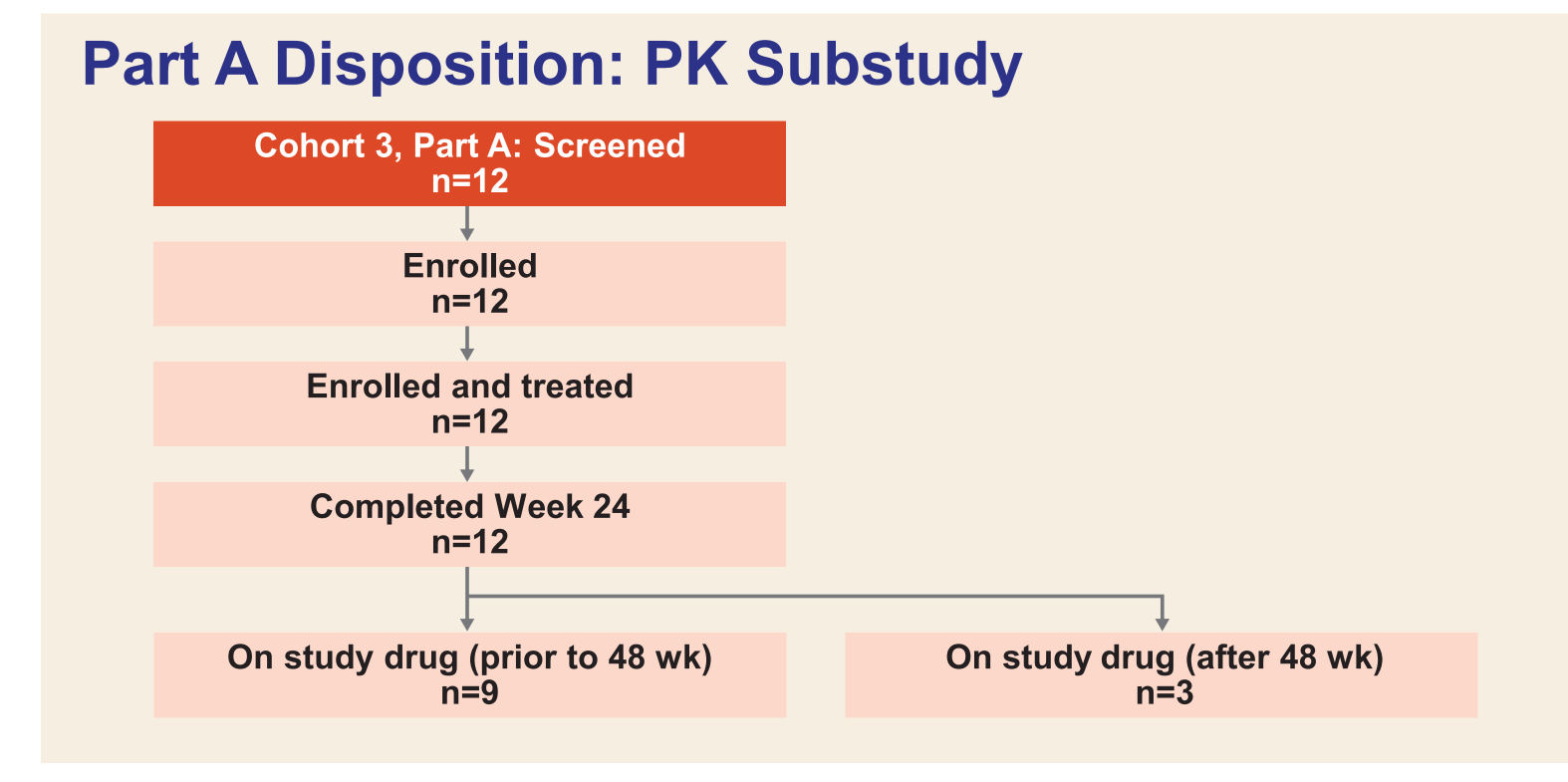
# Methods



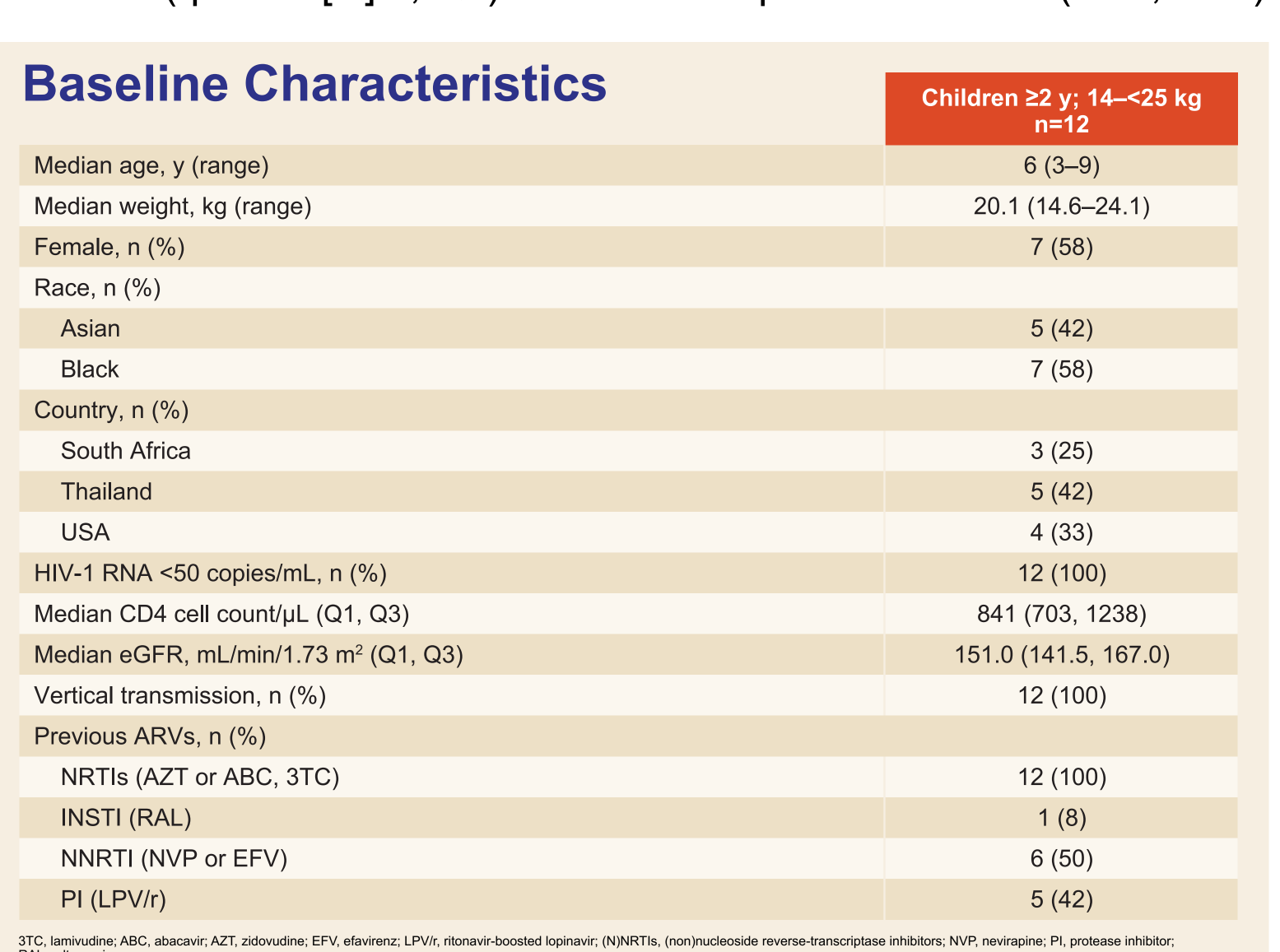
#### **Study Assessments**

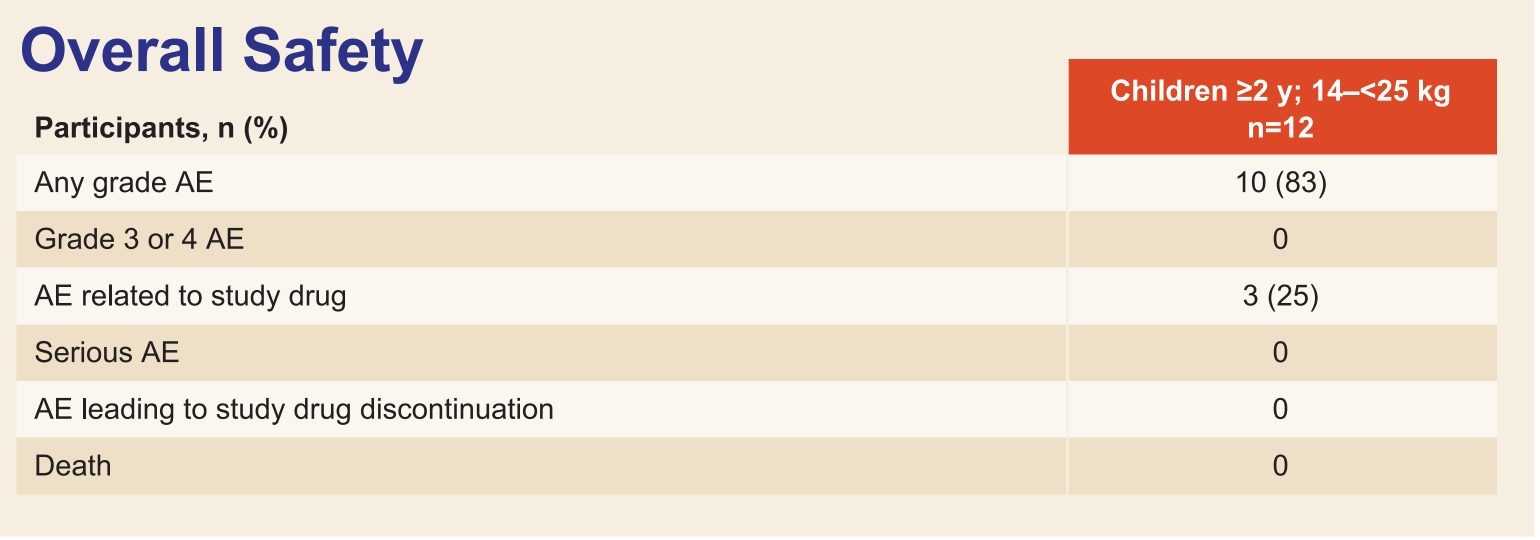
- PK: intensive and sparse PK samples collected to examine steadystate exposure of BIC, FTC, and TAF
- ◆ Safety: adverse events (AEs) and clinical laboratory abnormalities
- Efficacy: HIV-1 RNA and CD4 cell count
- Palatability and acceptability: questionnaires and facial scale
- Adherence: assessed by pill count at each visit

## Results



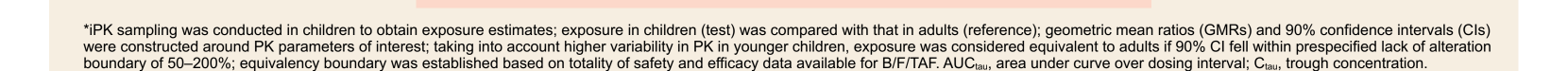
◆ Median (quartile [Q] 1, Q3) duration of exposure: 42.3 wk (40.1, 49.3)





- Most common AEs were upper respiratory tract infection (n=3 [25%]), and abdominal pain, constipation, diarrhea, vomiting, viral upper respiratory tract infection, enuresis, cough, and rhinorrhea (each n=2 [17%])
- No other AE occurred in >1 participant and all AEs were mild—moderate in severity
- ◆ 3 participants had AEs considered related to study drug: neutropenia, abdominal pain, irritability, and social avoidant behavior
- Grade 3 or 4 laboratory abnormalities: decreased neutrophils (n=2 [17%]) and increased creatinine (n=1 [8%])

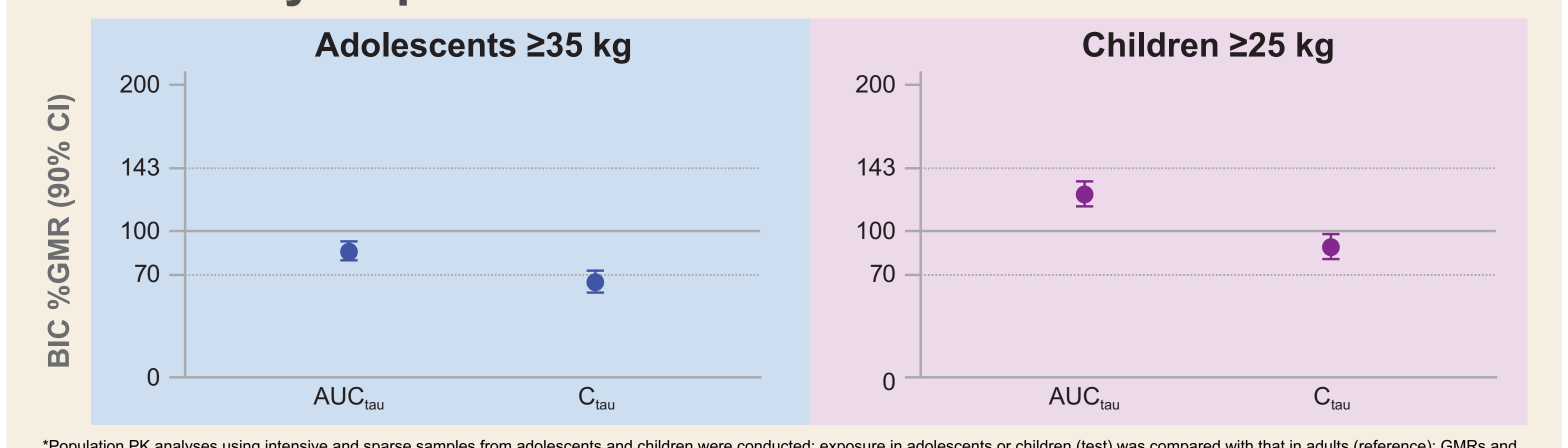
# Intensive BIC PK: B/F/TAF Low-Dose Tablet\* Children 14–<25 kg



BIC AUC<sub>tau</sub> was similar in children weighing 14–<25 kg relative to adults</li>
 BIC C<sub>tau</sub> mean estimate was lower in children weighing 14–<25 kg vs adults, but remained ~12-fold above the protein-adjusted 95%</li>

effective concentration (162 ng/mL) for wild-type virus

# Population BIC PK: B/F/TAF Full-Strength Tablet\* Previously Reported<sup>5,6</sup>

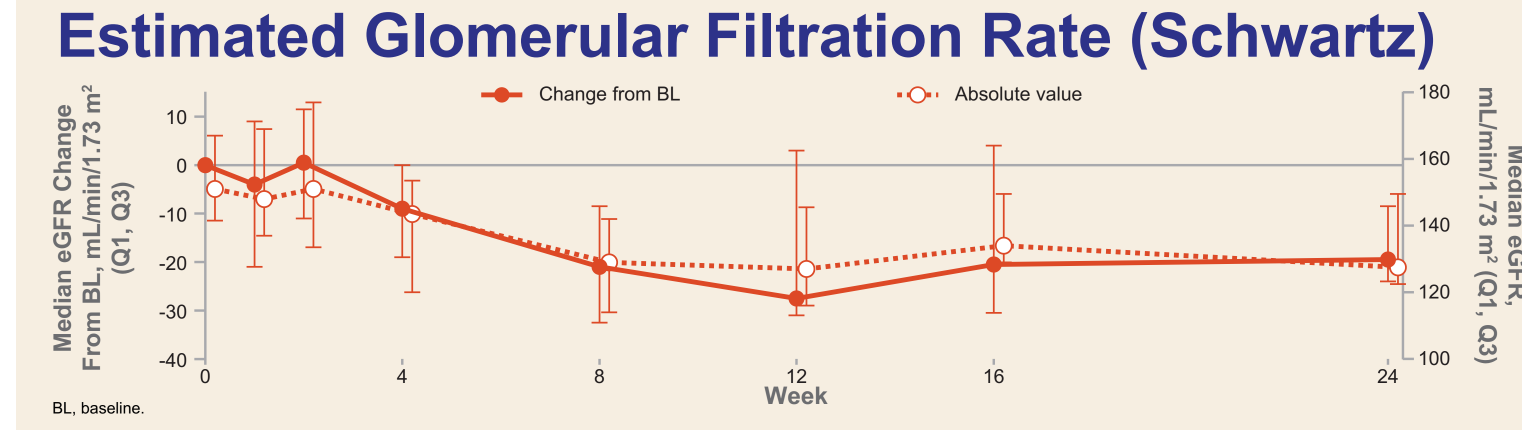


- BIC AUC<sub>tau</sub> was similar in adolescents and children weighing ≥25 kg compared with adults
- ◆ BIC C<sub>tau</sub> was similar in children and adults
- ◆ BIC C<sub>tau</sub> was lower in adolescents vs adults, but >11-fold above the protein-adjusted 95% effective concentration for wild-type virus

## **Intensive PK Data**

	PK Parameter*	Children ≥2 y; 14–<25 kg n=12 <sup>†</sup>	Adults n=74–77 <sup>‡</sup>	Child/Adult GMR% (90% CI)
FTC	AUC <sub>tau</sub> , h·ng/mL	14,576	11,790	124 (110, 139)
	C <sub>max</sub> , ng/mL	3473	2004	173 (144, 209)
	C <sub>tau</sub> , ng/mL	80§	90	89 (49, 161)
TAF	AUC <sub>tau</sub> , h·ng/mL	282	195	145 (115, 182)
	C <sub>max</sub> , ng/mL	393	227	173 (140, 214)
*Geometric mean; †1 participant was excluded from FTC summary due to noncompliance with study drug; ‡Pooled iPK data from four Phase 3 studies in adults with HIV. §n=10; C <sub>max</sub> , maximal concentration.				

 Exposures of FTC and TAF were within the safe and efficacious ranges of historical data in adults and adolescents following administration of approved FTC/TAF-containing products<sup>7,8</sup>

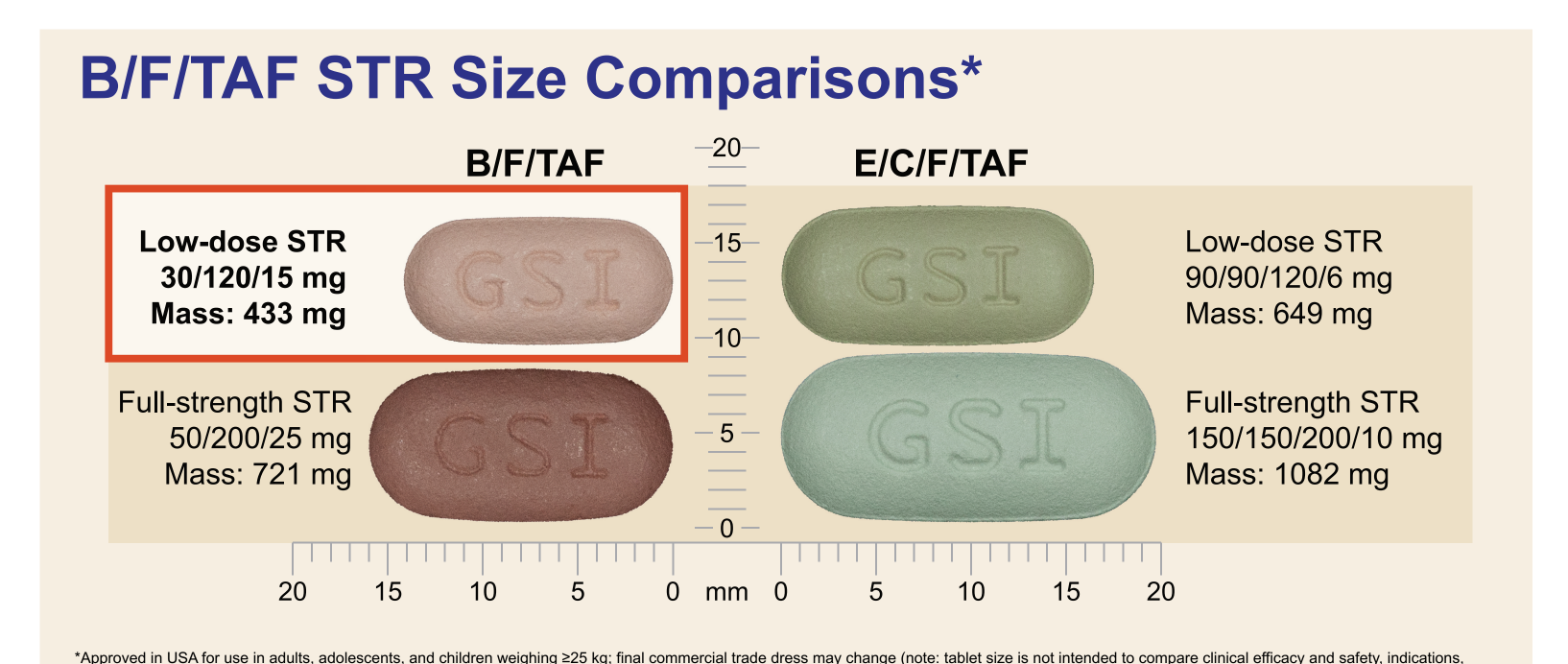


- Median changes in eGFR ranged from 0.5 to -27.5 mL/min/1.73 m<sup>2</sup> between Weeks 1 and 24
- Changes in eGFR in children weighing 14–<25 kg were consistent with the known renal creatinine transporter effect of BIC<sup>9,10</sup> and not considered clinically significant

# Palatability and Acceptability Ease of Swallowing If Whole Perceived Size When Swallowing If Whole Perceived Size When Swallowing If Whole Note of Split If Whole BL Week 4 Week 24 Near 1 Near 1

- Palatability and acceptability assessments:
- Questionnaire to ask the participant/parent:
- Whether tablet was broken in half to allow it to be swallowed
- If broken in half, whether both halves were taken within 10 min
- Facial scale and age-appropriate labels to rate:
- Ease or difficulty in swallowing tablet (if tablet was taken whole only)
- Acceptability of tablet shape
- Acceptability of tablet size
- Assessment of tablet taste
- ◆ Mean (standard deviation [SD]) adherence to B/F/TAF was 96.5% (6.1%)

- Nos. of children who split tablet: n=2 (BL), n=3 (Week 4), and n=1 (Week 24)
- For children who had tablet split, all were able to take both halves one right after the other (within 10 min)
- ◆ At planned assessments, all children aged 3 y (n=3) swallowed tablet whole: 1 at all time points, 1 at Week 24 only, and 1 at BL (Day 1) only



#### Efficacy: Virologic Outcome

- Using a missing = excluded analysis, virologic suppression (HIV-1 RNA <50 copies/mL) was maintained in all 12 participants (100%) at Week 24
- At Week 24, mean change (SD) in CD4 cell count was -66 cells/μL (180.7) and mean change in CD4% was -0.9 (4.64)
- No participants met the criteria for resistance analysis

# Conclusions

- In virologically suppressed children (aged ≥2 y; weight 14–<25 kg):
- The B/F/TAF low-dose STR was well tolerated
- All AEs were mild-moderate and there were no serious AEs, deaths, or AEs that led to discontinuation
- B/F/TAF demonstrated high rates of adherence and maintained virologic suppression
- Ability to swallow the low-dose STR was high, even down to age 3 y
- Exposures of BIC, FTC, and TAF were consistent with the ranges of exposures observed in adults in Phase 3 trials of B/F/TAF
- Efficacy and safety were consistent with results from Phase 3 trials of B/F/TAF in adults, which showed high proportions with viral suppression, no resistance, and good tolerability
- These data support further pediatric studies of B/F/TAF, which may be an important unboosted INSTI option for HIV-infected young children aged ≥2 y and able to swallow a tablet
- An additional 10 children have been enrolled in Cohort 3, Part B (current total n=22)
- The evaluation of other formulations of B/F/TAF in younger children who are unable to swallow tablets is planned

For information on relative bioavailability of the B/F/TAF low-dose STR, see Majeed S, et al. CROI 2020, poster 3194

**References: 1.** AIDSinfo. https://aidsinfo.nih.gov/guidelines/html/2/pediatric-arv/45/whats-new-in-the-guidelines: 12/19; **2.** Biktarvy [package insert]. Foster City, CA: Gilead Sciences, Inc.; 6/19; **3.** Gallant JE, et al. J Acquir Immune Defic Syndr 2017;75:61-6; **4.** Tsiang M, et al. Antimicrob Agents Chemother 2016.60:7086-97; **5.** Cotton M, et al. International Workshop on HIV Pediatrics 2018, oral 4; **6.** Gaur A, et al. CROI 2017, poster 424; **7.** Descovy [package insert]. Foster City, CA: Gilead Sciences, Inc.; 4/16; **8.** Genvoya [package insert]. Foster City, CA: Gilead Sciences, Inc.; 4/16; **9.** Zhang H, et al. International Workshop on Clinical Pharmacology of Antiviral Therapy 2017, poster 50; **10.** Zhang H, et al. British HIV Association 2017; oral O2. **Acknowledgments:** We extend our thanks to the participants, their families, and all participating study investigators and staff: South Africa: A Liberty, R Strehlau, R Van Zyl; Thailand: K Chokephaibulkit, P Kosalaraksa; USA: C Rodriguez, C Cunningham, E McGrath, N Rakhmanina. This study was funded by Gilead Sciences, Inc.