.gro.esnilebiugvih.www at Selecting an Initial ART Regimen. The full guideline is available State Department of Health AIDS Institute guideline go directly to a mobile-friendly version of the guideline. This  $^1/\!\!\mu$ -Folded Guide is a companion to the New York ← Use this code with your phone's QR code reader to



TDF: tenofovir disoproxil fumarate

RAL: raltegravir; RPV: rilpivirine; RTV: ritonavir; TAF: tenofovir alafenamide; DTG: dolutegravir; EFV: efavirenz; EVG: elvitegravir; FTC: emtricitabine; ATV: atazanavir; BIC: bictegravir; COBI: cobicistat; DOR: doravirine; DRV: darunavir; DRUG NAME ABBREVIATION KEY: 3TC: lamivudine; ABC: abacavir;

a documented negative HLA-B\*5701 test result.

JABC-containing regimens should not be used for rapid start without

results are available (such as for renal function or evidence of resistance). and any necessary adjustments to therapy should be made as soon as the available. Laboratory tests should be ordered at the time of initiation of ART, necessary to have the results of baseline laboratory tests immediately • When initiating ART at the time of diagnosis ("rapid start"), it is not

# ART-Initiation Laboratory Testing

- baseline viral load is very high (HIV RNA level >750,000 copies/mL). Consultation with an experienced HIV care provider is advised when a patient's
- incidence of hyperlipidemia than unboosted integrase strand transfer inhibitors.
- Boosted protease inhibitors and COBI-boosted EVG are associated with a higher studies, although not in others. No clear causal link has been established.
  - ABC has been associated with a higher risk of myocardial infarction in some
  - may cause a slight increase in measured creatinine. • Both COBI and DTG can cause decreased tubular excretion of creatinine and
- interactions (see NYSDOH AI guideline Rapid ART Initiation). possible) may be appropriate while addressing adherence issues and possible initiating ART. In some special cases, delay of initiation (for as short a time as · Neither mental health nor substance use disorders are contraindications to

### Special Considerations

**KEY POINTS** 

### Selected Drug-Drug Interactions to Discuss Before Initiating ART in **Treatment-Naive Patients Drug Class** ARV(s): Comments **ATV**: In treatment–naive patients on boosted ATV, $\rm H_2$ –blockers should be taken simultaneously with ATV/RTV with food. If simultaneous dosing with food is not possible, ATV/RTV should be taken at least 10 hours *after* the $\rm H_2$ –blocker. $\rm H_2$ –blocker H,-blockers doses should not exceed the equivalents of 40 mg famotidine twice daily for ART-naive patients or 20 mg famotidine twice daily for ART-experienced patients. $\mbox{RPV}:$ Use with caution; administer $\mbox{H}_{\mbox{\tiny 2}}\mbox{-blockers}$ at least 12 hours before or at least 4 hours after RPV. Inhaled steroids COBI; RTV: Alternatives or dose adjustments may be needed. Consult the package inserts for drug-drug interactions between specific statins and ARVs. Statins **BIC; DTG**: Take 2 hours before or 6 hours after DTG; calcium-containing antacids or iron supplements may be taken simultaneously if taken with food. Polyvalent cations [a] **RAL**: Magnesium- or aluminum-containing antacids are contraindicated; calcium-containing antacids are acceptable. RAL HD: Magnesium- or aluminum-containing antacids are contraindicated; coadministration of calcium-containing antacids is not recommended. EVG: Separate dosing by 2 hours, either before or after dose of EVG **ATV**: Contraindicated with ATV in treatment–experienced patients; in treatment naive patients, use no more than equivalent of 20 mg of omeprazole with ATV, Proton pump inhibitors separated by 12 hours RPV: Contraindicated. **DTG**: Metformin levels are significantly raised when coadministered with DTG. If used concomitantly, total daily dose of metformin should not exceed 1,000 mg without clinical evaluation of efficacy and adverse events. Metformin ATV/COBI; DRV/COBI; DRV/RTV; EFV: Use alternative or additional (e.g., barrier) contraceptive methods or choose alternative ART regimen. Ethinyl estradiol and norethindrone ATV; ATV/RTV: Use with caution; see manufacturer's package insert for specific dosing information. Factor Xa Apixaban: Reduce dose by 50% if patient is on 5 mg twice daily; avoid use if the indicated dose is 2.5 mg twice daily (based on age, weight, creatinine level). Dabigatran: No adjustment needed if CrCl ≥50 mL/min; avoid if CrCl <50 mL/min.</li> inhibitors Rivaroxaban: Avoid use. Platelet COBI; RTV: inhibitors Clopidogrel: Avoid use. - Prasugrel: No adjustment needed. Ticagrelor: Avoid use.

Additional abbreviation: CrCl, creatinine clearance.

**Notes:** a) Aluminum, calcium, magnesium, or iron in some antacids or vitamin preparations. b) For emergency contraception, other oral combinations, and patch, ring, or injectable formulations, please refer to package insert for specific ARV for dosing instructions and safety information.

to assess initial response to therapy. (A3)

· Clinicians should obtain a viral load test within 4 weeks after ARA initiation tolerance and adherence; adherence should be reinforced at regular intervals. (A3) methods, preferably within 2 weeks after treatment initiation, to assess

Clinicians or clinical support staff should follow up by telephone or other

#### Follow-up

through their toll-free Line: 1-866-637-2342.

- -The NYSDOH Clinical Education Initiative provides access to HIV specialists virus coinfection; or active opportunistic infections. (B3)
- comedications; impaired renal function; hepatitis B virus or hepatitis C - Extensive comorbidities, including metabolic complications and obesity;
- other than the available preferred or alternative regimens. (A3) - Baseline genotypic testing results indicating the need for an FAA regimen
  - selecting an initial ART regimen for a patient who has: Clinicians should consult with an experienced HIV care provider when

# Expert Consultation

resistance-associated mutations. (A1) resistance mutation. DTG/3TC is contraindicated in patients with these not have a major reverse transcriptase mutation, including the M184V/I - Genotypic resistance testing results have confirmed that a patient does

- HIV resistance and hepatitis B virus status are known. (A1)
- regimens as initial ART. (A3) Clinicians should prescribe DTG/3TC only after: • With the exception of DTG/3TC, clinicians should not prescribe 2-drug

Regimen Selection, cont.

ALL RECOMMENDATIONS (continued from P.1)

# HIV CLINICAL RESOURCE # 1/4-FOLDED GUIDE

VISIT HIVGUIDELINES.ORG TO LEARN MORE OR VIEW COMPLETE GUIDE



# **SELECTING AN INITIAL ART REGIMEN**

### NYSDOH AIDS INSTITUTE HIV CLINICAL GUIDELINE

Note: The recommendations in this guideline pertain to initial ART regimens for adults with HIV who are not pregnant.

### Dolutegravir (DTG) Safety Statement, May 2021

The recommendation regarding discussion of the small risk of teratogenicity with DTG in the first trimester and the need for birth control while using DTG was removed. DTG has been shown to be safe throughout pregnancy.

For more information, see Use of Dolutegravir in Individuals of Childbearing Capacity at hivguidelines.org/dtg

# **ALL RECOMMENDATIONS**

P.1

### Regimen Selection

- · When selecting an initial ART regimen for treatment naive-patients, clinicians should:
- Perform genotypic HIV resistance testing results for protease (A2), reverse transcriptase (A2), and integrase (B2) genotypic resistance if the testing has not already been performed or results are not otherwise available.
- Inform patients of the options and engage in shared decision-making to optimize the likelihood of adherence. (A3)
- Assess for comorbidities and chronic coadministered medications that may affect the choice of regimen for a patient's initial ART. (A3)
- Choose a preferred ART regimen unless one of the alternative regimens is a better choice based on individual patient factors. (A1)
- Recommend a single-tablet regimen or a regimen with once-daily dosing unless those regimens are contraindicated by HIV resistance, drug-drug interactions, intolerance, allergy, or access. (A2)
- Ask patients about their reproductive plans and discuss the use of contraception.(A3)
  - In choosing an initial ART regimen for a patient who is pregnant or planning a pregnancy, refer to DHHS Recommendations for the Use  $\,$ of Antiretroviral Drugs During Pregnancy and Interventions to Reduce Perinatal HIV Transmission in the United States. Continued on P.2 →

Regimen (rating)	Comments
- C - C - C - C - C - C - C - C - C - C	Available as a Single-Tablet Formulation
ABC/3TC/DTG (A1) [Triumeq]	
DTG/3TC (A1) [Dovato]	· Initiate <i>only</i> in patients with CrCl ≥30 mL/min.
	• Do not use in patients with HBV coinfection.
	• Do not initiate before HIV resistance tests results are available.
	<ul> <li>Do not initiate in patients with NRTI resistance, including the M184V/I mutation.</li> </ul>
	<ul> <li>Do not initiate in patients with baseline HIV RNA levels &gt;500,000 copies/m until additional study data are available.</li> </ul>
	Documented DTG resistance after initiation in treatment-naive patients is ran
	<ul> <li>Magnesium - or aluminum - containing antacids may be taken 2 hours before or 6 hours after DTG; calcium - containing antacids or iron supplements may be taken simultaneously if taken with food.</li> </ul>
	• Initiate <i>only</i> in patients with CrCl ≥30 mL/min.
(A1) [Biktarvy]	· Contains 25 mg of TAF, unboosted.
	<ul> <li>Magnesium- or aluminum-containing antacids may be taken 2 hours before or 6 hours after BIC; calcium-containing antacids or iron supplements may be taken simultaneously if taken with food.</li> </ul>
	$\cdot$ Documented DTG resistance after initiation in treatment-naive patients is ran
	Available as Multi-Tablet Regimen with Once-Daily Dosing
TAF 25 mg/FTC or	• For TAF/FTC, initiate <i>only</i> in patients with CrCl ≥30 mL/min.
TDF 300 mg/FTC and DTG (A1) [Descovy or Truvada and Tivicay]	· Contains 25 mg of TAF, unboosted.
	<ul> <li>For TDF/FTC, initiate only in patients with CrCl ≥50 mL/min.</li> </ul>
	• For TDF/FTC, consider bone mineral density.
	<ul> <li>Magnesium - or aluminum - containing antacids may be taken 2 hours before or 6 hours after DTG; calcium - containing antacids or iron supplements may be taken simultaneously if taken with food.</li> </ul>
	<ul> <li>Documented DTG resistance after initiation in treatment-naive patients is rar</li> </ul>
TAF 25 mg/FTC or TDF 300 mg/FTC and RAL HD (A2) [Descovy or Truvada and Isentress HD]	• For TAF/FTC, initiate <b>only</b> in patients with CrCl ≥30 mL/min.
	· Contains 25 mg of TAF, unboosted.
	• For TDF/FTC, initiate <b>only</b> in patients with CrCl ≥50 mL/min.
	<ul> <li>For TDF/FTC, consider bone mineral density.</li> <li>Administer as TAF/FTC or TDF/FTC once daily and RAL HD 1200 mg once daily, dosed as two 600 mg HD tablets.</li> </ul>
	To date, no clinical trials have been conducted with TAF and RAL; data are based on bioequivalence pharmacokinetic studies.
	Magnesium- or aluminum-containing antacids are contraindicated; coadministration of calcium-containing antacids is not recommended with RAL HD.
· Additional abbrevi	ations: CrCl, creatinine clearance; HBV, hepatitis B virus; NRTI, nucleoside reverse

- **Additional abbreviations:** CrCl, creatinine clearance; HBV, hepatitis B virus; NRTI, nucleoside reverse transcriptase inhibitor.
- ART regimens for individuals of childbearing potential: Refer to DHHS Recommendations for the Use of Antiretroviral Drugs During Pregnancy and Interventions to Reduce Perinatal HIV Transmission in the United States.
- **Substitutions:** 1) In all cases, FTC and 3TC are interchangeable. 2) TAF 10 mg and TAF 25 mg are not interchangeable.
- Dose adjustments: Refer to ARV Dose Adjustments for Hepatic or Renal Impairment section in full guideline.

CONTRAINDICATED ART Regimens Based on Routine Baseline Laboratory Parameters			
Lab Parameter	Contraindicated ART Regimens		
HIV RNA level ≥100,000 copies/mL	- ABC/3TC and ATV/COBI (Epzicom and Evotaz)     - ABC/3TC and EFV (Epzicom and Sustiva)     - ABC/3TC and ATV and RTV (Epzicom and Reyataz and Norvir)     - TAF/FTC/RPV (Odefsey)     - TDF/FTC/RPV (Complera)		
CD4 <200 cells/mm³	TAF/FTC/RPV (Odefsey)     TDF/FTC/RPV (Complera)		
CrCl <50 mL/min	ABC/3TC (Epzicom)     ABC/3TC/DTG (Triumeq)     TDF/3TC/DOR (Delstrigo)     TDF/FTC/EFV (Atripla)     TDF/FTC/RPV (Complera)		
CrCl <30 mL/min	TAF/FTC (Descovy) TAF/FTC/BIC (Biktarvy) TAF/FTC/DRV/COBI (Symtuza) TAF/FTC/EVG/COBI (Genvoya) [a] TAF/FTC/RPV (Odefsey) TDF/FTC (Truvada) DTG/3TC (Dovato)		

- Additional abbreviation: CrCl, creatinine clearance.
- $\begin{tabular}{ll} \textbf{Dose adjustments:} & \textbf{Refer to ARV Dose Adjustments for Hepatic or Renal Impairment section in full guideline.} \end{tabular}$
- Note: a) Unless CrCl <15 mL/min and on chronic hemodialysis.

Danisa and American	Comments
Regimen (rating)	Comments
TAF 10 mg/FTC/DRV/COBI (B2) [Symtuza]	e as a Single-Tablet Formulation  Initiate only in patients with CrCl ≥30 mL/min. Carefully consider drug-drug interactions with COBI. Contains 10 mg TAF, boosted.
TAF 10 mg/FTC/EVG/COBI (B1) [Genvoya]	Initiate only in patients with CrCl ≥30 mL/min.     Carefully consider drug-drug interactions with COBI.     Contains 10 mg of TAF, boosted with COBI.     Separate dosing of cation-containing (Ca++, AL, Mg) antacids by 2 hours, either before or after dose of EVG.
TAF 25 mg/FTC/RPV (B3) [Odefsey]	Initiate <i>only</i> in patients confirmed to have a CD4 cell count ≥200 cells/mm³ and HIV RNA level <100,000 copies/mL. Avoid use of RPV in a rapid-start or test-and-treat regimen if a patient's viral load and CD4 cour results are not available.     Initiate <i>only</i> in patients with CrCl ≥30 mL/min.     Use with caution in patients with depression or a history of suicidality.     To date, no clinical trials have been conducted; data are based on bioequivalence pharmacokinetic studies of TAI compared with TDF.     Contraindicated with proton pump inhibitors.     Use H₂-blockers with caution and separate dosing by 12 hours.     Must take with food.     Contains 25 mg of TAF, unboosted.
TDF/3TC/DOR (B1) [Delstrigo]	<ul> <li>Initiate <i>only</i> in patients with CrCl ≥50 mL/min.</li> <li>Contraindicated when coadministered with drugs that are strong CYP3A enzyme inducers.</li> <li>Consider bone mineral density.</li> </ul>
Available as Mult	i-Tablet Regimen with Once-Daily Dosing
<b>ABC/3TC</b> and <b>DOR</b> (B2) [Epzicom and Pifeltro]	Initiate only in patients confirmed to be negative for HLA-B*5701.     When a "rapid-start" or "test-and-treat" initiation of ART occurs before baseline laboratory test results are available, avoid use of ABC until a patient's HLA-B*570 test is confirmed negative.     Consider underlying risk of coronary heart disease.     Contraindicated when coadministered with drugs that are strong CYP3A enzyme inducers.
TAF 25 mg/FTC and DOR (B2) [Descovy and Pifeltro]	Initiate only in patients with CrCl ≥30 mL/min.     Contraindicated when coadministered with drugs that are strong CYP3A enzyme inducers.
Available as Multi	-Tablet Regimen with Twice-Daily Dosing
TAF 25 mg/FTC or TDF 300 mg/FTC and RAL (B3) [Descovy or Truyada and Isentress]	For TAF/FTC, initiate <i>only</i> in patients with CrCl ≥30 mL/mir     For TDF/FTC, initiate <i>only</i> in patients with CrCl ≥50 mL/mir

[Descovy or Truvada and Isentress]

- For TFD/FTC, consider bone mineral density. · Administer as TAF/FTC or TDF/FTC once daily and RAL
- 400 mg twice daily. Magnesium - or aluminum - containing antacids are
- contraindicated; calcium-containing antacids are acceptable with RAL.
- · Additional abbreviations: CrCl, creatinine clearance; CYP, cytochrome P450.
- **ART regimens for individuals of childbearing potential:** Refer to DHHS Recommendations for the Use of Antiretroviral Drugs During Pregnancy and Interventions to Reduce Perinatal HIV Transmission in the United States.
- Substitutions: 1) In all cases, FTC and 3TC are interchangeable. 2) TAF 10 mg and TAF 25 mg are not interchangeable. 3) COBI and RTV should not be considered interchangeable because of their drug-interaction profiles.
- **Dose adjustments:** Refer to ARV Dose Adjustments for Hepatic or Renal Impairment section in full guideline.