- · Addition or removal of pharmacokinetic "boosters" or "inducers" can cause adverse effects associated with elevated exposure or withdrawal of concomitant medication. These adverse effects may be falsely attributed to a new ART regimen rather than the need for dose adjustment or modification of the coadministered medication.
- production from a single infected clone.
- a period of at least 1 month may be the cause or result of chronic immune activation and should prompt a clinician to assess for adherence, preexisting resistance, or drug-drug interactions.

· Virologic failure is defined as a confirmed HIV viral load ≥200 copies/mL despite a

rebounds to ≥200 copies/mL after a patient achieves viral suppression.

· Persistent low-level viremia (HIV RNA 50 to 199 copies/mL) confirmed over

patient's use of recommended ART for at least 24 weeks or an HIV viral load that

- Once underlying drug resistance, potiential drug-drug interactions, and adherence have been addressed, persistent low-level viremia may reflect a large viral reservoir size or the consequence of constitutive, post-integration virus

- Identifying and addressing adherence problems causing virologic failure can prevent unnecessary ART intensification. Treatment intensification can further complicate adherence and expose additional classes of ARVs to the risk of resistance development.

- after suppression has been achieved, the clinician should confirm the result with a repeat HIV RNA test within 4 weeks of the original test. (A3) • When a patient's viral load test result indicates virologic failure (HIV RNA ≥200 · If a patient had a detectable viral load before a treatment interruption of <4 weeks, copies/mL) or low-level viremia (HIV RNA 50 to 199 copies/mL) confirmed over the clinician should obtain a plasma genotypic resistance test as soon as possible. (A2) a period of at least 1 month, the clinician should assess for and address the
- resistance, the clinician should recommend resumption of an appropriate ART regimen (based on assessment above) as soon as possible. (A2)
- · If the factor(s) related to interruption confer a low likelihood of emerging
- whether they are still present (A2)
- Existence of any barriers to adherence before the treatment interruption, and
- Pattern of adherence prior to discontinuation (A2)
- are being used (A2)
- Duration of the interruption, particularly if agents with long clearance half-lives
- The patient's plasma HIV-1 RNA level (viral load) at the time of ART interruption (A2)
- Reason for a treatment interruption, i.e., strategic or unplanned (A3)
- · Although drug resistance may not be present in all cases, when reinitiating ART after an interruption, clinicians should identify factors that may have contributed to potential selection of drug resistance, including:
- **Resumption of ART After a Treatment Interruption**

8- SELECTED KEY POINTS

P.3

ALL RECOMMENDATIONS (continued from P.3)

P.4

ALL RECOMMENDATIONS (continued from P.2)

Changes to Address Adverse Effects continued

- especially when switching from or to a regimen that may induce or inhibit medications, including nonprescription and over-the-counter medications, - Account for potential drug-drug interactions with chronically used concomitant - Account for the adverse effect profiles of ARVs, including cross-class toxicities.
- impaired renal function, or chronic anemia. underlying chronic medical conditions, such as cardiovascular disease or risk, - Minimize the potential for negative effects of a new ARA regimen on any shared metabolic pathways.

Changes to Address Drug-Drug Interactions

- :(2A) bluode neicinilo • When changing a patient's ART regimen to address drug-drug interactions, the
- medications planned for treatment of a comorbid condition before constructing - Acquire a current list of all medications that a patient is taking or any
- Account for the drug-clearance mechanisms and pharmacokinetic drug .n9mig91 TAA n6
- Pay particular attention to the effect of starting or stopping specific ARVs, interactions of ARVs to select optimal regimens.
- .instment. such as COBI or RTV, on concurrent medications that may require dose

Changes Due to Pregnancy

to Reduce Perinatal HIV Transmission in the United States. (A3) Recommendations for the Use of Antiretroviral Drugs During Pregnancy and Interventions pregnancy, the clinician should follow the recommendations of the DHHS: When changing an ART regimen for a patient who is pregnant or planning.

Changes for Regimen Simplification

- · Clinicians should not prescribe single-agent ART. (A1)
- · When changing an ART regimen for simplification, i.e., to improve adherence,
- reduce cost, improve quality of life, or respond to a patient's request, clinicians
- should construct a new regimen based on an assessment of:
- Prior resistance testing results (A1)
- History of ART failure (A2)
- Tolerability (A2)
- Evidence of clinical effectiveness (A2)

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HIV CLINICAL RESOURCE

ALL RECOMMENDATIONS

- Adherence (A2)

John's wort) (A*)

genotype test. (A2)

(B3)

weeks after discontinuation. (A2)

and its clinical effect is unclear. (A3)

Identifying and Managing Virologic Failure

following factors that may reduce ART efficacy:

when switching from a suppressive regimen.

ALL RECOMMENDATIONS (continued from P.1)

Changes to Address Drug Resistance

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P.1

Continued on P.2 \rightarrow

- :bluods sneibinib · When choosing a new ART regimen for a patient with drug-resistant virus,
- Choose a regimen that is likely to fully suppress viral replication, even if it may

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NYSDOH AIDS INSTITUTE HIV CLINICAL GUIDELINE

SECOND-LINE ART AFTER TREATMENT

• When a patient's plasma HIV–1 RNA level (viral load) is not suppressed to <200

copies/mL by 24 weeks after ART initiation or if it rebounds to ≥200 copies/mL

- Interactions between ART agents and concomitant medications, including over-the-counter medications and supplements (e.g., divalent cations, St.

- Adverse effects that lead to poor adherence or cessation of treatment (A2)

- Reviews of all prior drug resistance testing results, previous treatment

experience, and reason for treatment changes or discontinuation (A3)

· For all cases of virologic failure, clinicians should perform genotypic resistance

- If the viral load is ≥500 copies/mL, clinicians should obtain a plasma RNA

archived DNA genotype test if viral suppression is not achieved after any drug-drug interactions or problems with adherence have been addressed.

· In patients with persistent low-level viremia, clinicians should consult an

experienced HIV care provider; low-level viremia can have multiple causes,

- If the breakthrough viral load is <500 copies/mL, clinicians should obtain an

testing, ideally while the patient is taking the failing regimen or no longer than 4

FAILURE OR FOR REGIMEN SIMPLIFICATION

- require multi-tablet dosing. (A)
- tolerable regimen to suppress drug-resistant HIV effectively. (A3) - Document and evaluate the importance of all RMAS and identify the most
- Clinicians should address barriers to ART adherence that may have contributed to
- failure of a patient's first-line regimen. (A2)
- Review all prior genotype or phenotype resistance assay results that are In constructing a new regimen to replace a failed ART regimen, the clinician should:
- identifying potentially active medications. (A2) retrievable and previous instances of virologic treatment failure to assist in
- Select agents to which the patient is naive or active second-generation agents within
- DRV, DTG, or BIC, if the M184V RAM is present and FTC/3TC will be used in - Select a regimen containing an agent with a high barrier to resistance, such as a previously prescribed class to avoid potential within-class cross-resistance. (A2)
- conjunction with TAF/TDF. (A*)
- Choose the equivalent of 3 fully active ARVs; a 2-drug regimen may be - Avoid monotherapy (i.e., an ART regimen with fewer than 2 fully active agents). (A1)
- resistance barrier, i.e., a boosted PI or a second-generation INSTI. (A2) prescribed when both are fully active and at least 1 is an agent with a high
- Consult with an experienced HIV care provider when planning treatment
- FTC or another agent with activity against HBV (e.g., ETV) in the patient's ART - If a patient has chronic HBV infection, include TAF/TDF in conjunction with 3TC/ regimens for patients with multiclass drug-resistant virus. (A3)
- Clinicians should closely monitor the patient's response to ART by obtaining an regimen. (A2)
- HIV RUA test within 4 weeks of a change in regimen and at least every 8 weeks
- thereafter until virologic suppression is achieved. (A3)

Changes to Address Adverse Effects

- :(ZA) bluods When changing a patient's ART regimen to address adverse effects, the clinician
- for evidence of virologic failure to inform the choice of a fully active regimen - Review all prior genotype and phenotype resistance test results and ART history

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Genotypic Resistance Testing Based on Viral Load				
HIV RNA (Viral Load)	Indicated Genotypic Resistance Test			
0 to 500 copies/mL	HIV proviral DNA genotype (RT, PR, INT) or phenotype (tropism)			
500 to 1,000 copies/mL	HIV RNA genotype (RT, PR, INT) or phenotype (tropism) at assay amplification threshold; may use HIV proviral DNA test if nonamplifiable			
≥1,000 copies/mL	HIV RNA genotype if currently or recently (within 4 weeks) on ART; DNA proviral genotype may be considered for patients who are currently not taking ART but have in the past			

Antiretroviral Medications by Level of Genetic Barrier to Resistance (for group M, subtype B HIV)

Low Resistance (single mutation)	Intermediate Resistance (1 or 2 mutations)	High Resistance (>2 mutations)
• 3TC • EFV • EVG • FTC • NVP • RAL • RPV	 ABC CAB DOR FTR T20 TAF TDF ZDV 	 ATV (with COBI or RTV) BIC DRV (with COBI or RTV) DTG ETR MVC

Types of HIV Resistance Tests (Note: All resistance assays are affected by limitations of detection; minor variants may not be present at high enough concentrations to be amplified by the assay.)

Test	Description and Use	
Genotype	 Assesses mutations in the HIV RNA genes that encode enzymes targeted by ARVs: RT, PR, INT Algorithms interpret the effect of mutations on ARV efficacy At diagnosis, when a patient has incomplete virologic response to ART, or when viral rebound occurs Has utility if plasma HIV-1 RNA level (viral load) is ≥500 to 1,000 copies/mL May not detect all RAMs 	
Phenotype	Assesses the effect of HIV genes on the ARV concentration required to inhibit viral growth compared with wild-type (nonmutant) virus Estimates a fold change Historically used to help assess the effect of the interplay of multiple RAMs on viral growth Supplanted by more comprehensive genotypic interpretation algorithms	
Proviral DNA genotype (archived genotype)	 Assesses genetic mutations in HIV proviral DNA genes that encode enzymes targeted by ARVs: RT, PR, INT Algorithms interpret the effect of mutations on ARV efficacy When planning ART simplification or other changes, may have a role in identifying RAMs when standard genotype testing may not yield results, i.e., in patients who have prior treatment experience, have stopped taking ARVs for >4 weeks, or have an HIV viral load <500 to 1,000 copies/mL or below the limit of quantification May not detect all RAMs or report RAMs from defective non-replication-competent proviral DNA 	
Tropism test	 Assesses the effect of HIV RNA (or proviral DNA) gp120 on the coreceptor(s) used for viral attachment: CCR5, CXCR4, or mixed/dual Treatment-experienced patients for whom a coreceptor antagonist is being imminently considered RNA tropism test can be used with viral loads ≥1,000 copies/mL; proviral DNA test can be used for viral loads <1,000 copies/mL 	

ART Options After First-Line Treatment Failure With Single-Class Drug Resistance [a]

Failed First-Line Regimen Drug Classes	Classes and Medication Options for Switch
2 NRTIS + 1 NNRTI [a]	 2 NRTIs + 1 boosted PI: TAF/FTC/DRV/COBI (single tablet) TAF/FTC + DRV/RTV 2 NRTIs + 1 INSTI: TAF/FTC/BIC (single tablet) TAF/FTC + DTG
2 NRTIS + 1 PI [a]	 2 NRTIS + 1 INSTI: TAF/FTC/BIC (single tablet) TAF/FTC + DTG 1 INSTI + 1 NNRTI: RPV/DTG (single tablet) 2 NRTIS + 1 twice-daily boosted PI
2 NRTIS + 1 INSTI [a]	 2 NRTIs + 1 boosted PI: TAF/FTC/DRV/COBI (single tablet) TAF/FTC + DRV/RTV
Multiclass	 2 NRTIS + 1 INSTI + 1 boosted PI +/- 1 NNRTI (based on genotype): Consider: MVC [b], FTR, IBA, LEN, ETR, DOR, RPV, TPV

Notes:

a. Single-class resistance, with no major NRTI RAMs other than M184V b. If current tropism assay indicates exclusive R5 tropic virus

Common Adverse Effects Associated With Antiretroviral Medications

NRTIs	NNRTIs	PIs	INSTIS
 ABC [a]: Cardio- vascular disease; hypersensitivity DDI, d4T, ZDV: Mitochondrial toxicity; lipodystrophy; lactic acidosis TAF: Weight gain; lipids TDF: Proximal renal tubule injury; decrease in bone mineral density 	 DOR: CNS effects EFV: Hepato- toxicity; vitamin D deficiency; CNS effects; skin reactions; depression; morning somnolence NVP: Hepato- toxicity; hypersensitivity RPV: CNS effects; skin reactions; effects on the measure of eGFR 	 Class effect [b]: Increased cholesterol [c]; increased triglycerides [c]; increased glucose; Lipodystrophy ATV: Nephro- lithiasis; renal insufficiency; hyperbilirubin- emia DRV: Cardio- vascular disease; skin reactions LPV/RTV: Cardiovascular disease 	 Class effect [b]: Weight gain BIC: Effects on the measure of eGFR DTG: CNS effects; effects on the measure of eGFR EVG/COBI: Increased lipids; effects on the measure of eGFR

Notes:

a. Screen to document that the patient is negative for HLA-B*5701 before use

b. Adverse effects apply to all drugs in this class

c. Especially with RTV and COBI pharmaco-enhancement

Abbreviations:

3TC, lamivudine; ABC, abacavir; ART, antiretroviral therapy; ARV, antiretroviral medication; ATV, atazanavir; BIC, bictegravir; CAB, cabotegravir; CNS, central nervous system; COBI, cobicistat; d4T, stavudine; DDI, didanosine; DOR, doravirine; DRV, darunavir; DTG, dolutegravir; EFV, efavirenz; eGFR, estimated glomerular filtration rate; ETR, etravirine; ETV, entecavir; EVG, elvitegravir; FTC, emtricitabine; FTR, fostemsavir; gp120, envelope glycoprotein 120; IBA, ibalizumab; INSTI, integrase strand transfer inhibitor; INT, integrase; LEN, lenacapavir; MVC, maraviroc; NNRTI, non–nucleoside reverse transcriptase inhibitor; NRTI, nucleoside/nucleotide reverse transcriptase inhibitor; NVP, nevirapine; PI, protease inhibitor; PR, protease; RAL, raltegravir; RAM, resistance–associated mutation; RPV, rilpivirine; RT, reverse transcriptase; RTV, ritonavir; T20, enfuvirtide; TAF, tenofovir alafenamide; TDF, tenofovir disoproxil fumarate; TPV, tipranavir; ZDV, zidovudine.



← Use this code with your phone's QR code reader to go directly to a mobile-friendly version of the guideline.

This ¹/4-Folded Guide is a companion to the New York State Department of Health AIDS Institute guideline Second-Line ART After Treatment Failure or for Regimen Simplification. The full guideline is available at www.hivguidelines.org.