Modifying Antiretroviral Therapy in Virologically Suppressed HIV-1-Infected Patients

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Abstract HIV-1-infected patients with suppressed plasma viral loads often require changes to their antiretroviral (ARV) therapy to manage drug toxicity and intolerance, to improve adherence, and to avoid drug interactions. In patients who have never experienced virologic failure while receiving ARV therapy and who have no evidence of drug resistance, switching to any of the acceptable US Department of Health and Human Services first-line therapies is expected to maintain virologic suppression. However, in virologically suppressed patients with a history of virologic failure or drug resistance, it can be more challenging to change therapy while still maintaining virologic suppression. In these patients, it may be difficult to know whether the discontinuation of one of the ARVs in a suppressive regimen constitutes the removal of a key regimen component that will not be adequately supplanted by one or more substituted ARVs. In this article, we review many of the clinical scenarios requiring ARV therapy modification in patients with stable virologic suppression and outline the strategies for modifying therapy while maintaining long-term virologic suppression.

1 Introduction

Advances in antiretroviral (ARV) therapy (ART) have made it possible to achieve and maintain virologic suppression in nearly all HIV-1-infected patients. However, even patients with sustained virologic suppression require ART changes to manage acute toxicities, limit long-term adverse effects, improve adherence, and avoid drug–drug interactions [1, 2]. Indeed, ART is modified more commonly for these indications than for virologic failure (VF) [3–6]. ART modification in patients with stable virologic
suppression has been reported in more than one-third of patients on first-line ART over a 7-year period in a large Canadian cohort and between 8 and 43% annually in a variety of other clinical cohorts [3–5, 7–10]. Although modifying a suppressive ART regimen may be beneficial or even required for many patients, it carries a risk of VF and the development of resistance to one or more of the ARVs in a patient’s modified regimen [9]. This risk is heightened in patients with a history of VF because potentially not all ARVs in such patients’ regimens will be fully active. Therefore, modifying or switching therapy in such patients requires a review of past and current ARV regimens.

As new ARVs with improved toxicity profiles have been developed, there have been an increasing number of clinical trials of ART modification in virologically suppressed patients, and many of these studies have been summarized in an excellent review by Van den Eynde and Podzamczer [11]. Together, these trials provide guidance for several specific clinical scenarios and outline important principles necessary to maintain virologic suppression while changing therapy. However, many clinical scenarios and ART modification strategies have not been evaluated in randomized clinical trials and are instead supported primarily by non-randomized trials, observational cohort studies, and expert opinion. Here we review many of the clinical scenarios requiring ART modification in patients with stable virologic suppression and the accompanying strategies for modifying therapy while maintaining long-term virologic suppression.

2 Indications for Antiretroviral Therapy (ART) Modification

In the earliest years of ART, several studies attempted to limit the potential toxicity of ART by reducing the number of ARVs prescribed to virologically suppressed patients [12–14]. The VF rates in these early studies were unacceptably high, in part because virologic suppression was defined by insensitive virus load assays with lower limits of detection of 400–500 copies/ml and in part because the ARVs used to attain virologic suppression were less efficacious than the ARVs used now. As a result of these early failures and the improved tolerability of current ARVs, the strategy of simply removing an ARV from the regimen of a patient with stable virologic suppression is now studied primarily in patients receiving medications with a high genetic barrier to resistance, most commonly pharmacologically boosted protease inhibitors (PIs).

There have also been several intensification studies in which an ARV is added to the regimen of a patient with stable virologic suppression. The goal of these studies was to eliminate or reduce the residual levels of viremia that can often be detected with highly sensitive single-copy HIV-1 assays [15]. The rationale for these studies was based on the hypothesis that even very low levels of virus (fewer than 50 copies/ml) may be associated with damaging systemic inflammation and replenishment of the HIV-1 proviral DNA reservoir. Intensification studies, however, generally demonstrated no clinical benefit or effect on low-level residual viremia using highly sensitive single-copy/ml assays [15–18]. As a result, the addition of an ARV to the regimen of a patient with stably suppressed viremia is not recommended.

Table 1 is a comprehensive list of the main indications for switching ARVs in patients with stable virologic suppression. Among the most common chronic toxicities and forms of intolerance leading to ARV switches are gastrointestinal intolerance, lipid or other metabolic abnormalities, coronary artery disease, neuropsychiatric symptoms, renal dysfunction, and jaundice caused by unconjugated hyperbilirubinemia. Mitochondrial toxicity and injection site reactions are currently uncommon causes of treatment switches because there has been a marked decrease in the use of the mitochondrial toxic nucleoside reverse transcriptase inhibitors (NRTIs)—stavudine (d4T), didanosine (ddI), and zidovudine (AZT)—and of the parenteral fusion inhibitor enfuvirtide (ENF).

Co-morbid conditions typically necessitate ARV regimen changes when they exacerbate a pre-existing ARV-associated toxicity or require the use of a medication that has undesired or unpredictable pharmacologic interactions with an ARV in a patient’s regimen. For example, switching ARVs to avoid interaction with rifamycins during tuberculosis (TB) therapy is common in areas in which TB is endemic. Switching ARVs to limit pharmacologic interactions with direct-acting antivirals for hepatitis C treatment or to improve absorption in patients receiving proton-pump inhibitors is common in upper-income regions (Table 1).

There are also numerous drug–drug interactions between ARVs and other commonly prescribed drugs. Although the effects of many of these interactions on drug absorption and metabolism have been well characterized, other empirically observed interactions are poorly understood. As a result, there is no substitute for reviewing the potential interactions of a new ARV with each medication a patient is receiving. The US Department of Health and Human Services (DHHS) guidelines document contains an extensive highly accessible series of tables containing the most common interactions associated with each ARV.

There are also several up-to-date authoritative websites containing even more extensive information on drug–drug interactions such as the one maintained by the University of Liverpool: http://www.hiv-druginteractions.org.
| Type of indication | Specific indication | Main implicated ARVs | Comments and key references |
|--------------------|---------------------|---------------------|----------------------------|
| Chronic toxicity   | Lipoatrophy         | d4T > AZT           | Lipatrophy is an NRTI-induced mitochondrial toxicity strongly associated with d4T and to a lesser extent AZT. Patients who discontinue one of these NRTIs—including those who switch to ABC or TDF—prevent a worsening of their condition and may experience clinical improvement [19, 20] |
|                    | Lipid abnormalities and coronary artery disease | LPV/r | LPV/r and several less frequently used PI/r’s increase triglyceride and total cholesterol levels often increasing a patient’s predicted risk of coronary artery disease and requiring the use of lipid lowering drugs [21]. ATV/r, ATV/c, DRV/r, and DRV/c have much more modest effects than LPV/r on lipid levels [22, 23] |
| Renal dysfunction  | TDF                 |                     | TDF can be associated with a slow progressive reduction in GFR and renal tubular dysfunction that typically improves upon TDF discontinuation [26–28]. It is also necessary to discontinue TDF or adjust its dose in patients with an eGFR <60 ml/min regardless of the cause of renal insufficiency [1]. TAF is associated with reduced markers of renal tubular dysfunction. Long-term effects of TAF on eGFR as well as guidelines on its dosing with impaired renal function are not yet available [29, 30] |
| Osteopenia         | TDF                 |                     | TDF is associated with slightly reduced bone density in a large proportion of patients and requires discontinuation in patients with moderate or severe osteopenia [31]. TAF is associated with less short-term loss of bone mineral density compared to TDF but long-term data are not available [29, 30] |
| Intolerance        | Gastrointestinal    | LPV/r               | Diarrhea and nausea are among the most common clinical adverse reactions associated with LPV/r and several older PIs. These symptoms are much less severe with ATV/r, ATV/c, DRV/r, and DRV/c [32] |
| Neuropsychiatric   | EFV                 |                     | Headache, confusion, impaired concentration, amnesia, and sleep abnormalities occur commonly during the first few days of treatment and lead to discontinuation in 5–10 % of patients [33]. In most patients, these CNS symptoms decline to baseline levels within weeks with continued therapy [33, 34]. In an analysis of four ACTG studies of 3,241 EFV-recipients and 2,091 comparators, EFV use carried a hazard ratio of 2.6 for attempted or completed suicide (17 events vs. five events) [35]. A subsequent larger meta-analysis, however, did not demonstrate this association [34]. In combination with TDF/FTC, 400 mg has been shown to be as effective as 600 mg of EFV and to be associated with fewer treatment discontinuations and may be an option where such formulations are available [36] |
| Jaundice           | ATV or ATV/r        |                     | ATV causes indirect hyperbilirubinemia in some patients related to its inhibition of the UGT1A1 enzyme [37]. In some studies, this has been a major reason for changing therapy [38] |
| Acute toxicity     | Hypersensitivity reactions | ABC | Screening for HLA-B*5701 should prevent nearly all cases of ABC hypersensitivity [39, 40] |
|                    |                     | NVP                 | NVP is associated with an increased risk of severe rash-associated liver toxicity especially in women with elevated baseline transaminase levels or CD4 counts greater than 250 [1, 19, 41] |
|                    |                     | Other ARVs          | Hypersensitivity reactions occur in <1 % of patients receiving ARVs other than ABC and NVP. The DHHS guidelines specifically cite DTG as being associated with a hypersensitivity reaction in less than 1 % of patients during registration trials and an uncommon MVC-associated hypersensitivity reaction associated with transaminase elevations. Mild rashes have been reported in about 5 % of patients receiving DRV/r. These can be managed by close follow-up because they often resolve spontaneously [1, 19] |
| Type of indication                     | Specific indication | Main implicated ARVs | Comments and key references                                                                                                                                 |
|---------------------------------------|---------------------|----------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Parenteral administration             | Injection site reactions | ENF                  | Shortly after its approval, ENF became a key component of successful salvage therapy for patients with few other options. However, with the subsequent approval of RAL, DRV, and ETR, it became possible to create an equally suppressive ART regimen in a high proportion of patients receiving ENF |
| Drug dosing                           | Twice daily         | LPV/r, RAL, ETR, DRV/r | Twice-daily LPV/r dosing is recommended for patients with ≥3 LPV-resistance mutations, pregnant women, or patients receiving EFV, NVP, or any drug that reduces LPV levels [1]. RAL and ETR are approved and recommended for twice-daily dosing. DRV/r and DRV/c can be administered once daily in patients without DRV-resistance |
| Food requirement                      | RPV                 |                      | RPV absorption has a greater dependency on being taken with food than other ARVs [42]                                                                           |
| Risk of non-adherence                 | Complicated ARV combinations |                      | Fixed-dose combinations (FDCs) improve adherence but reduce dosing flexibility when adjustments are required. Five FDCs suitable both for first-line therapy and maintenance scenarios have been approved: TDF/FTC/EFV, TDF/FTC/EVG/c, TDF/FTC/RPV, ABC/3TC/DTG, and TAF/FTC/EVG/c. Two additional FDCs containing TAF in place of TDF are nearing FDA approval |
| Pregnancy                             | Fetal issues        | EFV                  | There are conflicting data on whether EFV is associated with an increased risk of neural tube defects [43]. The DHHS panel recommends that EFV not be initiated during the first 8 weeks of pregnancy [44]. The WHO makes no such recommendation as pregnancy is rarely recognized during its first 8 weeks [45] |
| Maternal issues                       | NVP                 |                      | NVP hepatotoxicity and rash are more common during pregnancy particularly in patients with pre-existing liver disease or CD4 count >250 cells/mm³ [46] |
| Co-infections                         | Tuberculosis        | NNRTIs               | Standard EFV dose is acceptable. RPV and ETV are not recommended in patients receiving rifampin due to CYP3A induction of NNRTI metabolism [1] |
|                                       | PI                  |                      | Not recommended in patients receiving rifampin due to CYP3A induction of PI metabolism by rifampin. May be used with close monitoring in patients receiving rifabutin [1] |
|                                       | INSTI               |                      | Twice the standard RAL dose (800 mg twice daily) is recommended but has not been evaluated in clinical studies. DTG must be used twice daily in patients receiving rifampin. In patients with documented or suspected INSTI resistance, DTG should be used with rifabutin [1] |
| HCVa                                  | Ledipasvir          |                      | Ledipasvir increases TDF levels, which is enhanced further with ritonavir or cobicistat-boosted PIs or EVG/c [47]. Patients receiving ledipasvir should therefore be monitored for TDF toxicity. TPV is contraindicated in patients receiving ledipasvir |
|                                       | Sofosbuvir          |                      | TPV co-administration is not recommended because it may lower sofosbuvir concentration by inducing P-glycoprotein |
|                                       | Simeprevir          |                      | PIs, EFV, NVP, ETR, and EVG/c not recommended. Expert consultation is recommended |
|                                       | Ombitasvir/paritaprevir/dasabuvir/ritonavir |                      | PIs other than ATV, NNRTIs, and EVG/c are contraindicated. Expert consultation is recommended |

[1] DHHS Panel on Antiretroviral Guidelines for Adults and Adolescents, 2018 guidelines. The Henry Ford Health System, 2018. [42] Ritzel T, et al. J Clin Pharmacol. 2017. [43] Lu J, et al. Neurology. 2014. [44] DHHS Panel on Antiretroviral Guidelines for Adults and Adolescents, 2018 guidelines. The Henry Ford Health System, 2018. [45] WHO Guidelines for the use of antenatal and intrapartum HIV prophylaxis, 2018. [46] Centers for Disease Control and Prevention. Guidelines for the prevention of opportunistic infections among HIV-exposed and HIV-infected children. 2018. [47] Rader JD, et al. J Infect Dis. 2013.
Pre-conception counseling coupled with prospective ART modification for women likely to become pregnant will reduce the likelihood that ART will need to be changed during pregnancy. The DHHS preferred regimens for treatment-naive pregnant patients include ritonavir-boosted atazanavir (ATV/r), darunavir (DRV/r), or raltegravir (RAL) in combination with one of the following NRTI backbones: abacavir (ABC)/lamivudine (3TC), tenofovir disoproxil fumarate (TDF)/emtricitabine (FTC), TDF/3TC, or AZT/3TC [44]. The initiation of efavirenz (EFV) is not recommended during the first 8 weeks after conception in treatment-naive patients. However, as pregnancy is frequently not detected until several weeks after conception, this recommendation is relevant primarily for the choice of therapy in women planning to conceive.

Pharmacokinetic changes may lower PI plasma levels during the third trimester [44]. Therefore ritonavir-boosted lopinavir (LPV/r) and DRV/r should be administered twice daily and unboosted ATV is not recommended. Although there are fewer data on the safety to the fetus of both the most recently approved ARVs such as elvitegravir (EVG), dolutegravir (DTG), cobicistat, and tenofovir alafenamide (TAF) and the more rarely used ARVs such as tipranavir (TPV), maraviroc (MVC), and enfuvirtide (ENF), no ARVs, other than the combined use of d4T and ddI, are specifically contraindicated [44]. ARV switches during pregnancy may also be associated with a greater risk of virologic failure compared with continued unchanged therapy, supporting the recommendation that pregnant patients be managed in consultation with specialists experienced in treating HIV in pregnancy [48].

Adherence to ART is essential for both attaining and sustaining virologic suppression. Adherence can be influenced by several factors including the likelihood of forgetting to take prescribed medications. Treatment-naive patients are at higher risk of non-adherence. It is not possible to provide specific recommendations on which regimens are associated with the lowest cost to a patient and if these regimens are associated with the highest adherence. It is also not possible to provide specific recommendations on the cost of an ART regimen either through co-payment or drug company financial assistance programs, and third-party payers. Nonetheless, health-care providers should be aware of the direct patient costs when choosing therapy.

### Table 1 continued

| Type of indication | Specific indication | Main implicated ARVs | Comments and key references |
|--------------------|---------------------|----------------------|-----------------------------|
| Additional drug–drug interactions | Drug absorption | Acid reducing agents | There are many interactions between ARVs and between ARVs and other drugs commonly used in HIV-1-infected patients. The DHHS guidelines and several websites contain comprehensive tables and expert recommendations on which drug combinations are contraindicated and which require dosage modifications of the ARV or interacting drug [1] |
| | Hepatic induction and inhibition | Polyvalent cations | |
| | | p-glycoprotein effects | |
| | | CYP450 enzymes | |
| | | UGT enzyme | |

ABC abacavir, ACTG AIDS Clinical Trials Group, ART antiretroviral therapy, ARV antiretroviral, ATV atazanavir, AZT zidovudine, c cobicistat, d4T stavudine, ddI didanosine, DHHS US Department of Health and Human Services, DTG dolutegravir, EFV efavirenz, ENF enfuvirtide, eGFR estimated glomerular filtration rate in ml/min, ETR etravirine, EVG elvitegravir, FDA US Food and Drug Administration, FDC fixed-dose combination, FTC emtricitabine, HCV hepatitis C virus, HDL high-density lipoprotein, INSTI integrase strand transfer inhibitor, LDL low-density lipoprotein cholesterol, LPV lopinavir, NRTI non-nucleoside reverse transcriptase inhibitor, NRRTI nucleos(t)ide reverse transcriptase inhibitor, NVP nevirapine, PI protease inhibitor, PI/r ritonavir-boosted protease inhibitor, PK pharmacokinetic, r low-dose ritonavir, RAL raltegravir, TAF tenofovir alafenamide, TDF tenofovir disoproxil fumarate, TPV tipranavir, WHO World Health Organization

Available HCV medications and data on interactions are rapidly emerging. Up-to-date detailed drug interactions can be found at: [http://www.hep-druginteractions.org/](http://www.hep-druginteractions.org/)
3 Clinical Trials of ART Modification

ART modification trials can be divided into those enrolling patients with no history of VF who are receiving an initial or early ART regimen and those enrolling patients who may have a history of VF with documented or suspected drug resistance. In patients with stable virologic suppression and no history of VF, many ART regimens are likely to be effective options when a change in therapy is necessary. Indeed, each of the regimens classified by the DHHS into Recommended, Alternative, or Other options for first-line therapy (Table 2) appear to be acceptable options for patients with stable virologic suppression and no history of VF. For example, TDF/FTC/rilpivirine (RPV) is categorized as an Alternative option for first-line therapy because it is not recommended for patients with plasma HIV-1 RNA levels above 5.0 log copies/ml. However, it has been shown to be highly effective in patients with stable virologic suppression regardless of their pre-therapy plasma HIV-1 RNA level [49].

It is more challenging to modify therapy in virologically suppressed patients with a history of VF. In such patients it is difficult to know whether the discontinuation of one of the ARVs in a suppressive regimen constitutes the removal of a key regimen component that will not be adequately supplanted by a substituted ARV. In such patients it is necessary to consider past drug resistance test results and past ARVs to which drug resistance may have developed but may not have been documented by resistance testing [50, 51].

Table 3 summarizes the design and main findings of most of the randomized controlled trials of ART modification in virologically suppressed patients. The trials are grouped according to the ARV classes substituting for one another. In most trials, virologic suppression was defined as a VL below 50–75 copies/ml for 12–24 weeks. Histories of VF or drug resistance were exclusion criteria for many of the studies.

The primary endpoint in most trials was the proportion of patients with VF by week 48, where VF was defined as the development of HIV RNA levels above a particular threshold ranging from 50 to 400 copies/ml. HIV RNA levels were assessed at either a single time point as in the FDA Snapshot analysis [77] or had to be confirmed with a repeat test. To preserve the principles of intention-to-treat, several studies distinguished primary VF from treatment failure, which included primary VF, treatment discontinuation because of intolerance or drug toxicity, and losses to follow-up. The difference among treatment groups was often assessed using a non-inferiority analysis in which the lower bound of the confidence interval for the difference in proportions of successfully treated patients was compared using a pre-specified margin of difference usually between 10 and 20 %.

4 ART Modification Strategies

The following sections organize ART modifications in patients with stably suppressed virus according to the ARV class of a drug that requires discontinuation and the ARV class of its replacement drug. Three sections address ART modifications that involve discontinuing NRTIs, PIs, and NNRTIs regardless of whether discontinuation is required to manage toxicity, address drug–drug interactions, or improve adherence. Additional sections address the potential roles for boosted PI monotherapy, boosted PIs plus lamivudine, and boosted PIs plus an integrase strand transfer inhibitor (INSTI), several unorthodox ART modification approaches, and the discontinuation of ENF.

4.1 Discontinuation of a Nucleoside Reverse Transcriptase Inhibitor (NRTI)

d4T, ddl, and to a lesser extent AZT are associated with a wide range of toxicities as a result of their off-target inhibition of human mitochondrial gamma polymerase [19,
| Classes | Study | Study design | Key inclusion criteria | Findings |
|---------|-------|--------------|------------------------|----------|
| Within NRTI | BICOMBO [52] | 2 NRTIs → ABC/3TC (n = 190) vs. TDF/FTC (n = 290) | • VL <200 × 24 W  
• Cr <2 mg/ml | • Trend towards increased TF by ITT with ABC (19 %) vs. TDF (13 %;  
\( p = 0.06 \))  
• 4 patients on ABC vs. 0 on TDF developed VF (\( p = 0.04 \))  
• Median CD4 increase was greater with ABC (44 cells/mm\(^3\)) vs. TDF (−3 cells/mm\(^3\)) |
| | STEAL [53] | ≥2 NRTIs → ABC/3TC (n = 180) vs. TDF/FTC (n = 180) | • VL <50 × 12 W  
• eGFR ≥70 | • Similar rates of VF (confirmed VL >400) for ABC/3TC (5.6 %) vs. TDF/FTC (3.9 %)  
• ↑ Cardiac events with ABC/3TC (2.2/100 patient years) vs. TDF/FTC (0.3/100 patient years;  
\( p = 0.05 \))  
• ↓ Bone mineral density (hip T-score) with TDF/FTC |
| | SWIFT [54] | ABC/3TC + PI/r → TDF/FTC + PI/r (n = 155) vs. no change (n = 156) | • VL <200 × 12 W  
• No resistance to TDF/FTC or PI/r  
• eGFR ≥50 | • ↑ VF (confirmed VL >200) with ABC/3TC (11/156; 7.1 %) than TDF/FTC (3/155; 1.9 %;  
\( p = 0.03 \))  
• ↓ Cholesterol, triglycerides and Framingham risk with TDF/FTC  
• ↓ Bone mineral density (hip T-score) with TDF/FTC  
• ↓ eGFR (8.3 vs. 4.5 ml/min) with TDF/FTC  
• ↓ Total cholesterol (26 mg/dl,  
\( p < 0.001 \)) and LDL cholesterol (15 mg/dl,  
\( p = 0.005 \)) with TDF/FTC, ↓ HDL cholesterol with TDF/FTC (4 mg/dl) compared to control (1 mg/dl,  
\( p = 0.026 \)) |
| | Behrens et al. [55] | ABC/3TC + LPV/r → TDF/FTC + LPV/r (n = 43) vs. no change (n = 42) | • VL <50 × 12 W  
• Total cholesterol ≥200 mg/dl | • 87 % of patients in both arms had VL <50 at week 24  
• Adverse reactions were low in both arms |
| | ASSURE [56] | TDF/FTC + ATV/r → ABC/3TC + ATV (n = 199) vs. no change (n = 97) | • VL <75 × 2 tests  
• No prior VF | |
| | ROCKET [57] | ABC/3TC + EFV → TDF/FTC + EFV (n = 78) vs. no change (n = 79) | • VL <50 × 12 W  
• Hyperlipidemia  
• eGFR ≥60 | • ↓ LDL cholesterol with TDF/FTC (22 mg/dl), ↓ HDL cholesterol with TDF/FTC (5 mg/dL) |
| | SWAP [58, 59] | AZT/3TC → ABC/3TC (n = 20) vs. TDF/FTC (n = 20) | • VL <40 × 12 W | • ↓ Bone mineral density by 2 % at 24 and 48 W with TDF/FTC |
| Classes   | Study       | Study design                                                                 | Key inclusion criteria                                                                 | Findings                                                                                                                                 |
|-----------|-------------|-----------------------------------------------------------------------------|----------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------|
| Within PI | SWAN [60]  | • PI or PI/r-regimen → ATV (n = 278) (ATV/r if on TDF, n = 26) vs. no change (n = 141)  
• 48 W | • Stable PI ≥12 W  
• VL <50 × ≥12 W  
• No prior VF on a PI-containing regimen | • ↓ VF for those on ATV or ATV/r (19/278; 7 %) vs. control (22/141; 16 %,  
   p = 0.004)  
• ↓ Total cholesterol, non-HDL cholesterol, and fasting TG for those on ATV or ATV/r |
|          |             |                                                                             |                                                                                         |                                                                                                                                            |
|          | ATAZIP [61]| • LPV/r + NRTIs → ATV/r + NRTIs (n = 121) vs. no change (n = 127)  
• 48 W | • VL <200 × 24 W  
• ≤2 episodes of VF on a PI-regimen  
• ≤5 PI-DRMs | • Similar rates of TF with ATV/r (21/121; 17 %) vs. LPV/r (25/127; 20 %)  
• Similar rates of VF (confirmed VL >200) with ATV/r (6/121; 5 %) vs.  
   LPV/r (9/127; 7 %)  
• Similar rates of overall TF (14 vs. 19 % by TLOVRb) and VF (1/210; 0.5 %  
   vs. 7/209; 3.5 %) with ATV vs. ATV/r, respectively  
• ↓ TG (−80mg/dl) and total cholesterol (−19mg/dl; p < 0.001) in the ATV group |
|          | ARIES [62] | • ATV/r + ABC/3TC induction × 36 W → ATV + ABC/3TC (n = 210) vs. no change (n = 209)  
• 48 W | • ART-naïve  
• VL <50 × 36 W  
• HLA-B*5701 | • Similar rates of TF with ATV/r (21/85; 25 %) vs. ATV (19/87; 22 %)  
• ↓ hyperbilirubinemia (26 % vs. 15 %) with ATV  
• Adverse reactions were low in both arms  
• No prior VF  
• No pre-ART TDF or FTC resistance | |
| InduMa [63] | • ATV/r + 2 NRTIs induction × 24 W → ATV + 2 NRTIs not including TDF (n = 87) vs. no change (n = 85)  
• 48 W | • ART-naïve  
• VL <50 × 24 W  
• No prior VF | • 87 % of patients in both arms had VL <50 at week 24 |
|          | ASSURE [56]| • TDF/FTC + ATV/r → ABC/3TC + ATV (n = 199) vs. no change (n = 97)  
• 24 W | • VL <75 × 2 tests  
• No prior VF | • Similar rates of TF with ATV/r (21/85; 25 %) vs. ATV (19/87; 22 %)  
• ↓ TG with ATV but no differences in LDL or HDL levels  
• ↓ hyperbilirubinemia (26 % vs. 15 %) with ATV  |
| SLOAT [64] | • LPV/r-containing regimen → ATV (n = 102) (ATV/r if TDF used n = 53) vs. no change (n = 87)  
• 48 W | • VL <50 × 24 W | • Similar rates of VF with ATV (5/49; 10 %), ATV/r (7/53; 13 %), and LPV/r (9/87; 10 %). Of patients with VF, 5/12 on ATV or ATV/r vs 1/9 on LPV/r  
   failed with PI resistance  
• ↓ TG (−80mg/dl) and total cholesterol (−19mg/dl; p < 0.001) in the ATV group |
|          | SWITCHMRK  [65]| • LPV/r + ARVs → RAL + ARVs (n = 350) vs. no change (n = 352)  
• 24 W | • VL <50–75 × ≥12 W  
• No CAD, DM, lipid-lowering drugs | • ↑ TF with RAL (16 %) vs. control (9 %; p < 0.05)  
• ↓ 13 % reduction in total cholesterol and 42 % reduction in TG with RAL  
• Similar proportion with TF for RAL (15/139; 11 %) vs. control (18/134; 13 %). 4 patients in RAL arm vs 6 in control arm with VF |
|          | SPIRAL [66] | • PI/r + ARVs → RAL + ARVs (n = 134) vs. no change (n = 139)  
• 48 W | • VL <50 × 24 W | |
|          | STRATEGY-PI [67]| • PI/r + TDF/FTC → EVG/c + TDF/FTC (n = 293) vs. no change (n = 140)  
• 48 W | • No prior VF  
• No pre-ART TDF or FTC resistance | • ↑ TF with TDF/FTC/EVG/c (6 %) vs. control (13 %; p = 0.025) driven by  
   non-VF discontinuations in control arm |

PIs to INSTI

82 S. E. Collins et al.
| Classes       | Study            | Study design                                                                 | Key inclusion criteria                                                                 | Findings                                                                                                                                                                                                 |
|--------------|-----------------|------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| PI to NNRTI  | NEFA [68]       | • PI or PI/r-regimen to 2 NRTIs + NVP (n = 155), EFV (n = 156), or ABC (n = 149) | • NNRTI and ABC-naïve • VL < 200 × 24 W                                                | • Kaplan–Meier estimates of death, AIDS progression, or VL > 200 were 6, 10, and 13 % for EFV, NVP, and ABC, respectively. Both EFV and NVP were superior to ABC. No significant difference between EFV and NVP |
|              | AI266073 [69]   | • PI, PI/r, NNRTI-regimen to TDF/FTC/EFV (n = 203) vs. no change (n = 97)     | • VL < 200 × 12 W • No prior VF                                                        | • Similar proportion with TF (confirmed VL > 200) for TDF/FTC/EFV (11 %) vs. unchanged therapy (12 %)                                                                                                       |
|              | SPIRIT [49]     | • PI/r + 2 NRTIs → TDF/FTC/RPV (n = 317) vs. no change (n = 159)             | • VL < 50 × 24 W • No prior VF • No NRTI or RPV-resistance                            | • Similar results when stratified by baseline PI, PI/r, or NNRTI-regimen                                                                                                                                 |
|              | Waters et al    | • 2 NRTIs + EFV → 2 NRTIs + ETR 400 mg OD (intermediate vs. delayed switch)  | • VL < 50 • Ongoing CNS symptoms after 12 W                                             | • EFV in 3/317 (0.9 %) on TDF/FTC/RPV and 8/159 (5 %) in the PI/r arm • None of 18 with a pre-ART K103N developed VF on TDF/FTC/RPV |
| Within NNRTI | SWITCH-EE       | • EFV + 2 NRTIs → EFV vs. ETR + 2 NRTIs × 12 W (double-blind crossover)      | • VL < 50 × 12 W • ETR administered 400 mg OD                                         | • EFV remaining therapeutic for 2–4 W post switch. RPV was therapeutic by 1–2 W post switch, because of EFV CYP450 3A4 induction • VF without drug resistance in 2/49 patients by 48 W |
|              | [70]            | • TDF/FTC/EFV → TDF/FTC/RPV                                                  | • VL < 50 × 12 W • EFV intolerance • No prior ART or RPV resistance                   | • Patient preference for RAL in 41 %, EFV in 23 %, and neither in 36 % based on CNS symptomatology                                                                                                       |
|              | SWITCH-ER       | • EFV + 2 NRTIs → RAL + 2 NRTIs vs. no change × 4 W (double-blind crossover) | • VL < 50 × 12 W • No prior VF • No TDF or FTC resistance • eGFR ≥ 70                 | • Approximately 75 % were on TDF/FTC/EFV at entry • Similar proportion with TF, TDF/FTC/EVG/c (7 %) vs. no change (12 %; p = 0.07). VF in 1 % in each arm, none failed with drug resistance |
|              | Mills et al     | • NVP or EFV + TDF/FTC → EVG/c + TDF/FTC (n = 292) vs. no change (n = 147)    | • VL < 50 × 24 W • No prior VF • No TDF or FTC resistance • eGFR ≥ 70                 | • Virologic non-inferior, 85 vs. 88 % had VL < 50 • More discontinuations due to adverse events in switch arm 10/275 (4 %) vs. 0/277 in the continuation arm |
|              | STRATEGY-NNRTI  | • NVP or EFV + TDF/FTC → EVG/c + TDF/FTC (n = 292) vs. no change (n = 147)    | • VL < 50 × 24 W • No prior VF • No TDF or FTC resistance • eGFR ≥ 70                 | • Virologic non-inferior, 85 vs. 88 % had VL < 50 • More discontinuations due to adverse events in switch arm 10/275 (4 %) vs. 0/277 in the continuation arm |
|              | STRIVING        | • PI, PI/r, NNRTI, or INSTI (RAL or EVG)-regimen → ABC/3TC/DTG (n = 274) vs. no change (n = 277) | • VL < 50 × 2 tests • Stable ART × 24 W • No prior VF                                  | • Virologic non-inferior, 85 vs. 88 % had VL < 50 • More discontinuations due to adverse events in switch arm 10/275 (4 %) vs. 0/277 in the continuation arm |
Classes | Study | Study design | Key inclusion criteria | Findings
--- | --- | --- | --- | ---
GS Study | 109[76] | • EFV, EVG/c, ATV/r or ATV/c + TDF/FTC → TAF/FTC/EVG/c (n = 959) vs. no change (n = 477) | • VL <50 × 24 W | • Switch superior, 97 vs. 93% had VL <50 at 48 W (p < 0.001).
• eGFR ≥50 | • VF 1% in both arms
• Successful completion of prior clinical study for ARV initiation | | • Small improvement in bone density and markers of renal tubular dysfunction (measured as % change) in the switch arm

3TC lamivudine, ABC abacavir, ART antiretroviral therapy, ARV antiretroviral, ATV atazanavir, AZT zidovudine, c cobicistat, CNS central nervous system, Cr serum creatinine, d4T stavudine, ddI didanosine, EFV efavirenz, eGFR estimated glomerular filtration rate in ml/min, ETR etravirine, EVG elvitegravir, FDA US Food and Drug Administration, FTC emtricitabine, HDM high-density lipoprotein, INSTI integrate strand transfer inhibitor, ITT intention-to-treat, LDL low-density lipoprotein cholesterol, LPV lopinavir, N number in overall study, n number in study subgroup, NNRTI non-nucleoside reverse transcriptase inhibitor, NRTI nucleos(t)ide reverse transcriptase inhibitor, NFV nelfinavir, NVP nevirapine, PI protease inhibitor, r low-dose ritonavir, RAL raltegravir, TDF tenofovir disoproxil fumarate, TF treatment failure, TG triglycerides, TLOVR time to loss of virologic response, VF virologic failure, VL plasma viral load, vs. versus, W week

a Treatment failure (TF) definitions differ slightly between studies. TF typically encompasses virologic breakthrough, confirmed virologic failure, and medication switches due to intolerance. In ITT analyses participants lost to follow-up are considered TFs. Individual study differences are noted in the table.
b TLOVR analysis is a time-dependent method of assessing antiviral treatment efficacy in study subjects with virologic suppression. Subjects with two consecutive viral load measurements over the threshold (typically 50 copies/ml) are considered failures. The FDA Snapshot algorithm has now become the preferred method of assessment because it gives similar results but is easier to calculate and interpret. Details can be found at the FDA’s website [77].

Table 3 continued
4.2 Discontinuation of a Protease Inhibitor (PI)

Ritonavir-boosted PIs are among the most common ARVs requiring substitution because they are more likely than NNRTIs and INSTIs to be associated with gastrointestinal intolerance, elevated lipids, and drug–drug interactions [21]. Although each of the most commonly used PI/r’s—DRV/r, ATV/r, and LPV/r—can have these side effects, they are usually more severe with LPV/r. This is due in part to LPV’s uniform requirement for 200 mg rather than 100 mg of ritonavir for boosting.

It is not yet known whether adverse effects associated with DRV/cobicistat (DRV/c) and ATV/cobicistat (ATV/c) are less frequent than those associated with DRV/r and ATV/r, but initial data suggest that these new formulations have a similar side-effect profile [91]. Jaundice from unconjugated hyperbilirubinemia is the most common side effect of pharmacologically boosted ATV and it often requires a treatment change in patients who find the condition disturbing [38]. ATV/r is also associated with an increased risk of nephrolithiasis [92]. There are three main approaches to modifying therapy in virologically suppressed patients with PI-associated toxicity including changing therapy to a different boosted PI or unboosted ATV (at an increased dose of 400 mg daily), changing therapy to an INSTI, or changing therapy to an NNRTI.

4.2.1 Switching Among PIs

Table 3 summarizes six clinical trials that studied the effect of changing PI therapy in virologically suppressed patients. In three trials, previously treated patients receiving LPV/r or an older PI were randomized to either continued therapy or to a change from LPV/r to ATV/r or ATV while continuing the other ARVs in their regimen [60, 61, 64]. In one trial, previously treated patients on ATV/r were randomized to either continued ATV/r or a change to ATV [56]. In two trials, previously ART-naive patients received an induction regimen of ATV/r plus two NRTIs and were then randomized to continued therapy or change to unboosted ATV [62, 63]. Previous VF on a PI-containing regimen was an exclusion criterion in all but one trial. Because TDF reduces ATV levels, studies of unboosted ATV did not include patients with stable virologic suppression on a TDF-containing regimen.

The SWAN trial compared a switch to ATV/r or ATV with continued therapy with one of the older, generally less well tolerated PIs. It demonstrated that a change to ATV/r or ATV was associated with a lower risk of VF than continued therapy (7 vs. 16 %; p = 0.004) [60]. One-third of the patients in the SWAN study were initially receiving LPV/r and two-thirds were receiving PIs that are now rarely used, including nelfinavir (NFV), indinavir (IND), and saquinavir (SQV). Two of the five trials reported an improved lipid profile associated with a switch to ATV/r or ATV, and one reported a modest reduction in hyperbilirubinemia in patients switching from ATV/r to ATV [63].

LPV/r and DRV/r have a strong track record for salvage therapy in patients with high-level NRTI resistance [93, 94]. Two lines of evidence suggest LPV/r and DRV/r have a higher genetic barrier to resistance than boosted ATV. First, these boosted PIs usually require three or more drug-resistance mutations (DRMs) in the protease before most of their inhibitor activity is lost [95, 96]. In contrast, a single protease DRM is often sufficient for the loss of ATV/r activity. Second, LPV/r and DRV/r have significantly higher clinical phenotypic cut-offs compared with ATV/r [97]. Despite the favorable outcome of a change to ATV/r or ATV in these trials, they do not provide evidence supporting the switch from LPV/r or DRV/r to ATV/r in stably suppressed patients with a history of previous VF [98, 99]. However, changes from ATV or ATV/r to either LPV/r or DRV/r would be expected to maintain virologic suppression, regardless of a patient’s past treatment history.

4.2.2 Switching from PIs to Integrase Strand Transfer Inhibitors (INSTIs)

There are five randomized clinical trials in which stably suppressed patients have switched therapy from a PI/r to an INSTI (Table 3). The STRATEGY-PI, STRIIVING, and GS Study 109 trials enrolled patients without a history of VF or resistance to TDF or FTC [67, 75, 76]. The SWITCHMRK and SPIRAL studies included patients with a history of VF [65, 66].

In the STRATEGY-PI study, there were significantly more treatment failures in patients continuing an LPV/r, ATV/r, or DRV/r-containing regimen than those switching to TDF/FTC/elvitegravir/cobicistat (EVG/c) (13 vs. 6 %; p = 0.025), the difference driven by discontinuations for non-virologic reasons. Preliminary results from the GS Study 109, which randomized 1,436 suppressed participants on TDF/FTC plus either EVG/c (32 %), EFV (26 %), ATV/r, or ATV/c (42 %) to a FDC of TAF/FTC/EVG/c or continued therapy was superior at 48 weeks with 97 versus 93 % suppressed (p < 0.001) [76]. Small improvements were seen in short-term measures of bone density and renal function in participants in the switch arm.

The STRIIVING study also switched virologically suppressed patients without a history of VF who were HLA*B-5701 negative from a variety of first-line regimens to the FDC of ABC/3TC/DTG and compared them to those who continued their current treatment. This switch was non-inferior to continued therapy at 24 weeks with 85 versus 88 % maintaining virologic suppression [75].

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**Table 4** Phase II/III trials of boosted protease inhibitor mono- and dual therapy

| PI   | Study                  | Study design                                                                 | Key inclusion criteria                                                | Findings                                                                                                                                                                                                 |
|------|------------------------|------------------------------------------------------------------------------|-----------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| LPV/r | OKO4 [81, 82]          | • LPV/r + 2 NRTIs → LPV/r (n = 100) monotherapy vs. no change (n = 98)    | • VL <50 × 24 W                                                     | • LPV/r was non-inferior to LPV/r + 2 NRTIs according to the primary TF analysis which allowed for the readdition of NRTIs. The rates of maintaining HIV VL <50 in the more strict intention to treat analysis were also similar among the two arms (77 vs. 78 %). |
|      |                        | • 96 W                                                                       | • No prior VF on a PI-containing regimen                            |                                                                                                                                                                                                           |
|      | OLE [83]               | • LPV/r + 2 NRTIs → LPV/r + 3TC (n = 123) vs. no change (n = 127)         | • VL <50 × 24 W                                                     | • Discontinuation due to adverse events less common with LPV/r (0 vs. 8 %; p = 0.003)                                                                                                                  |
|      |                        | • 48 W                                                                       | • No resistance to LPV/r or 3TC                                     |                                                                                                                                                                                                           |
| DRV/r| MONOI-ANRS [84]        | • DRV/r + 2 NRTIs induction × 8 W followed by DRV/r (n = 112) vs. continued DRV/r + 2 NRTIs (n = 113) | • VL <400 × 72 W and <50 × 12 W                                   | • TF defined as confirmed VL >50, clinical progression, or discontinuation occurred in 3 % of each arm                                                                                                     |
|      |                        | • 48 W                                                                       | • No history of VF on a PI-containing regimen                      |                                                                                                                                                                                                           |
|      | PROTEA [87]            | • Stable ART → DRV/r (n = 127) vs. DRV/r + 2 NRTIs (n = 129)               | • VL <50 × 24 W                                                     | Rates of TF for DRV/r were not non-inferior to DRV/r + 2 NRTIs in either the per-protocol (6 vs. 1 %) or ITT analysis (12.5 vs. 8 %)                                                                              |
|      |                        | • 96 W                                                                       | • No prior VF                                                       |                                                                                                                                                                                                           |
|      | SALT [90]              | • ART × 48 W → ATV/r + 3TC (n = 143) vs. ATV/r + 2 NRTI (n = 143)         | • VL <50 × 24 W                                                     |                                                                                                                                                                                                           |
|      |                        | • 48 W                                                                       | • No prior VF                                                       |                                                                                                                                                                                                           |
|      |                        |                                                                              | • No resistance to ATV/r or 3TC                                     |                                                                                                                                                                                                           |
|      |                        |                                                                              |                                                                       |                                                                                                                                                                                                           |

3TC lamivudine, ART antiretroviral therapy, ATV atazanavir, ATV/r ritonavir-boosted atazanavir, ITT intention to treat, LDL low-density lipoprotein cholesterol, LPV/r ritonavir-boosted lopinavir, n number in study subgroup, NRTI nucleos(t)ide reverse transcriptase inhibitor, PI protease inhibitor, TF treatment failure, TLOVR time to loss of virologic response, VF virologic failure, vs. versus, W week

\(^a\) Treatment failure (TF) definitions differ slightly between studies. TF typically encompasses virologic breakthrough, confirmed virologic failure, and medication switches due to intolerance. In ITT analyses participants lost to follow-up are considered TFs. Individual study differences, including clinical progression of disease, are noted in the table
Analysis of adverse events noted that ten participants randomized to ABC/3TC/DTG (4 % of the total) but 0 participants who did not change therapy stopped ARVs due to various forms of intolerance.

In the SWITCHMRK trials (SWITCHMRK 1 and SWITCHMRK 2), 702 patients stably suppressed on LPV/r plus two NRTIs for 12 weeks were randomized to either continued therapy or a switch to RAL + 2 NRTIs [65]. Approximately 60 % of patients in the trials had received one or more previous ART regimens and approximately 30 % of patients had a history of VF on an earlier regimen. Despite having screening plasma HIV-1 RNA levels below 50 copies/ml, 5 % had detectable viremia (median RNA level: 101 copies/ml; IQR: 63–193) immediately prior to starting study drugs.

Within 12 weeks, patients who had switched to RAL experienced reductions in their non-HDL cholesterol and triglyceride levels of 15 and 42 %, respectively. However, the trial was terminated prematurely because, by 24 weeks, VF was significantly more likely in patients who had switched to RAL than in those who had continued their previous regimen (15.6 vs. 9.4 %). Among those patients with VF who underwent genotypic resistance testing, most had developed RAL resistance. Of the 32 RAL recipients with VF, 27 (84 %) reported that the LPV/r regimen at study entry was not their first ART regimen, and 18 of these 27 had a history of previous VF. Of the 17 RAL recipients with a baseline RNA level above the 50 copies/ml threshold of detection, seven developed VF.

The SPIRAL trial enrolled 273 patients most of whom were stably suppressed for 24 weeks on LPV/r (44 %), ATV/r (35 %), or FPV/r (12 %), then randomized them to either continue therapy or to switch to RAL [66]. Nearly 90 % of these patients had received one or more previous ART regimens and nearly 40 % had a history of VF. At 48 weeks, treatment failure had occurred in 11 % of patients receiving RAL and in 13 % of those remaining on PI/r’s. Possible reasons for the discordant results between SWITCHMRK and SPIRAL may be the longer period of virologic suppression prior to randomization in SPIRAL or genotypic resistance to the NRTI backbone, which was detected in half of SWITCHMRK participants who failed RAL but not in SPIRAL.

The significant increase in VF in patients switching to RAL in the SWITCHMRK studies underscores the importance of an uncompromised NRTI backbone in patients receiving RAL. Although RAL and EVG/c may be as potent antiviral inhibitors as the currently used PI/r’s, they have lower genetic barriers to resistance. Just one or two drug resistance mutations may be sufficient to cause high-level RAL and EVG resistance [100], whereas four or more drug resistance mutations are usually required for high-level LPV/r or DRV/r resistance [95, 96]. Three or more drug-resistance mutations in the integrase gene region are usually required for DTG resistance [101–103], and DTG has been shown to be more effective than RAL at treating ARV-experienced patients with VF [104]. An ongoing study of ABC/3TC/DTG for second-line therapy in patients with NRTI-resistant viruses will provide insight into the ability of DTG-containing regimens to maintain virologic suppression in PI/r-treated patients with underlying NRTI resistance [105].

4.2.3 Switching from PIs to Non-Nucleoside Reverse Transcriptase Inhibitors (NNRTIs)

There have been two studies in which stably suppressed patients without a history of VF switched therapy from a PI/r to an NNRTI. The first of these studies, AI266073, enrolled patients receiving a wide range of PI- and NNRTI-containing regimens and randomized them to either continued therapy or a switch to TDF/FTC/EV [69]. Over a period of 48 weeks, 89 % of patients receiving TDF/FTC/EV and 88 % whose therapy was unchanged maintained virologic suppression.

The second study, SPIRIT, enrolled patients receiving mainly ATV/r, LPV/r, and DRV/r who had no history of VF or prior ART and no evidence for NRTI or RPV resistance. These patients were randomized to switch to TDF/FTC/RPV or to continue their current therapy [49]. After 24 weeks, there was no difference in VF between the two groups (6 % of those who had switched to TDF/FTC/RPV and 10 % of those who did not change therapy experienced VF; \(p = 0.15\)). Of note, none of the 18 patients with a pre-ART genotype containing K103N, the commonly transmitted nevirapine (NVP) and EFV-resistance mutation, developed VF on TDF/FTC/RPV. Plasma HIV-1 RNA levels prior to the start of their initial PI/r containing regimen did not predict VF, suggesting that once virus loads are stably suppressed the risk of VF may not be greater in patients with pre-therapy RNA levels that exceed 5.0 log10 copies/ml.

One of the earliest ART modification studies substituted EFV, NVP, or ABC for a PI in patients with plasma HIV-1 RNA levels below 200 copies/ml for 24 weeks. Approximately 90 % of these patients were receiving IDV or NFV, and approximately 50 % had a history of mono or dual NRTI therapy prior to attaining virologic suppression [68]. At 12 months, the Kaplan–Meier estimates of VF or disease progression were 6 % in the EFV group, 10 % in the NFV group, and 13 % in the ABC group. A history of suboptimal therapy prior to attaining virologic suppression was a strong risk factor for VF with 23 of 29 patients with VF having a history of mono- or dual NRTI therapy.

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4.3 Discontinuation of an NNRTI

NVP is no longer recommended as a preferred medication by the WHO, DHHS, or EACS for any specific indication and is now used infrequently in middle- and upper-income countries [1, 2, 45]. NVP is associated with severe hypersensitivity including rash and hepatitis in about 2% of patients and less severe abnormalities in up to 10% [19, 41, 46]. EFV is a key component of most WHO-recommended ART regimens. It was recently removed from the list of DHHS-Recommended ART regimens, but the FDC of TDF/FTC/EFV remains in wide use as a DHHS-Alternative regimen. Central nervous system symptoms are the main side effects of EFV [33–35]. The dose-dependency of these symptoms and the frequency with which they persist to cause non-adherence or serious morbidity is now being studied as newer FDCs have become available for comparison [34–36].

The newer NNRTIs etravirine (ETR) and RPV have fewer safety and tolerability concerns than NVP and EFV. Although ETR and RPV have similar cross-resistance profiles, ETR is considerably more potent and has a higher genetic barrier to resistance because the recommended 400-mg daily dosage of ETR is 16-fold higher than the recommended 25-mg daily dosage of RPV. The lower daily dosage of RPV is due to the fact that healthy, non-infected human volunteers experienced QT prolongations at doses of 75 mg or higher [106]. However, the high potency and lack of co-formulation of ETR mean it is primarily used as a salvage therapy option.

4.3.1 Switching Among NNRTIs

There have been two randomized, blinded studies in which a small number of patients receiving an EFV-containing regimen were randomized to either switch to ETR 400 mg once daily or to continue EFV, and one non-randomized study in which a small number of patients receiving TDF/FTC/EFV were switched to TDF/FTC/RPV (Table 3). These studies suggest that despite the inductive effects of EFV on CYP450 3A4, such a switch can occur without jeopardizing virologic suppression because EFV levels remain therapeutic while RPV (and presumably also ETR) levels increase [72, 107]. These studies also suggest that those patients with persistent central nervous system symptoms on EFV will experience a reduction of these symptoms with ETR (and presumably also RPV) [70]. Because TDF/FTC/RPV has been associated with a greater risk of emergent drug resistance compared with TDF/FTC/EFV [108–110], switching from TDF/FTC/EFV to TDF/FTC/RPV should only be done in patients without a history of VF or evidence of resistance to TDF, FTC, or RPV.

4.3.2 Switching from NNRTIs to an INSTI

A randomized, double-blind, crossover study of a small number of patients switching from EFV to RAL demonstrated a significant reduction in anxiety while patients were on RAL [73]. The much larger STRATEGY-NNRTI demonstrated the non-inferiority of a switch from an NNRTI plus two NRTIs to the FDC of TDF/FTC/EVG/c and also noted a reduction in sleep disturbances in patients switching to EVG/c [74, 111]. The aforementioned GS Study 109 and STRIVIING studies both enrolled substantial proportions of participants on NNRTI-based therapy (31 and 26%) and switched them to INSTI-based therapy, supporting the efficacy of this modification strategy [75, 76].

4.3.3 Switching from NNRTIs to PIs

Pharmacologically-boosted PIs have higher genetic barriers to resistance than NNRTIs and switches from NNRTIs to boosted PIs are commonly made in patients failing NNRTI-based therapy. Changing the NNRTI component of a virologically suppressive regimen to pharmacologically boosted DRV or ATV—the most well tolerated of the DHHS Acceptable first-line PIs—is likely to be highly effective in most virologically suppressed patients. This type of switch, however, has not been studied in virologically suppressed patients because boosted PIs have been highly effective substitutes for an NNRTI even in patients with VF [37, 93, 112].

4.4 Boosted PI Mono or Dual Therapy

A large number of randomized clinical trials suggest that LPV/r and DRV/r monotherapy are effective options for stably suppressed patients without a history of PI resistance (Table 4) (reviewed in Arribas et al. [113, 114]). Although LPV/r and DRV/r monotherapy are associated with an increased risk of virologic rebound compared with triple ART, this viral rebound is rarely associated with emergent drug resistance. The re-administration of NRTIs in these patients nearly always leads to re-suppression of virus levels to below 50 copies/ml [81, 82, 84–87, 115]. The risk of VF in patients receiving PI/r monotherapy is strongly associated with a patient’s nadir CD4 count and may be confined to patients with nadir CD4 counts below 200 cells/mm³ [87, 116].

Several small studies suggest that ATV/r monotherapy is associated with a higher risk of VF than LPV/r and DRV/r monotherapy [88, 89, 117–119]. One large observational study suggests that DRV/r is also more effective than LPV/r [120]. Although patients receiving PI/r monotherapy have an elevated risk of detectable CSF virus
when compared with those receiving triple therapy, they do not appear to have an increased risk of central nervous system symptoms [116, 121–124].

Several recent and ongoing trials have shown that LPV/r, ATV/r, and DRV/r in combination with 3TC are not associated with an increased risk of virologic rebound compared with PI/r’s plus two NRTIs provided there is no baseline 3TC resistance [83, 90, 125]. Supporting this concept the GARDEL trial showed that a regimen of LPV/r plus 3TC is as effective as LPV/r plus two 2 NRTIs as initial therapy in ARV-naïve patients [126].

4.5 Boosted PIs Plus INSTIs

There have been several clinical trials in which LPV/r, DRV/r, and ATV/r have been used in combination with RAL for first- and second-line therapy or for maintenance therapy in virologically suppressed patients [22, 92–94, 127–129]. These trials have shown the following: (1) DRV/r plus RAL was comparable to DRV/r plus TDF/FTC for first-line therapy except in patients with high VL or low CD4 counts where DRV/r plus RAL was less effective in subgroup analyses [127]; (2) LPV/r plus RAL was virologically comparable to but no better than LPV/r plus two NRTIs in patients with previous VF on a first-line NRTI plus NNRTI-containing regimen [93, 94]; (3) LPV/r plus RAL and DRV/r plus RAL were comparable to LPV/r and DRV/r plus two NRTIs in virologically suppressed patients [22, 92]; and (4) ATV/r plus RAL was associated with a higher risk of virologic rebound (7/72; 10 %) compared with ATV/r plus TDF/FTC (1/37; 3 %) in patients with stable virologic suppression [128].

Regardless of the clinical scenario and boosted PI, many patients with VF developed RAL-resistance mutations that would limit future treatment options. Even under the most favorable circumstances—such as DRV/r plus RAL for first-line therapy in closely monitored patients [127]—a drug-resistance sub-study showed that 4 % of 805 patients developed RAL resistance by week 80 [130]. A likely explanation for this finding is that the prevention of RAL resistance may require a consistently high level of boosted PI adherence that is often not achieved outside of a clinical trial. Therefore switching stably suppressed patients to a boosted PI plus INSTI-containing regimen should be restricted to the use of boosted DRV—because of its combination of tolerability and high genetic barrier to resistance—plus RAL or DTG and should be done primarily for the management of NRTI toxicity in highly adherent patients without a history of PI resistance.

4.6 Discontinuation of Enfuvirtide (ENF)

The demonstration that virologically suppressed patients receiving ENF could switch to RAL without risking virologic rebound [131–134] paved the way for many of the ART modification studies that followed. As several additional, highly potent and novel ARVs were approved (including DRV, ETR, and RAL), the number of patients receiving ENF has decreased dramatically. The few patients currently receiving ENF are likely to have failed many ARV regimens and to have extensive resistance to the more commonly used drug classes. ENF discontinuation in these patients requires expert consultation to choose an individually tailored replacement ART regimen with sufficient antiviral potency and a genetic barrier high enough to replace ENF.

4.7 Miscellaneous Other Regimens

4.7.1 Maraviroc-Containing Regimens

In patients for whom MVC is being considered it is necessary to exclude the presence of CXCR4 tropic variants, which would render MVC inactive. In virologically suppressed patients, however, it is not possible to test plasma virus for CXCR4 tropism. Therefore several commercial laboratories have developed proviral DNA tropism assays. The “Appendix” summarizes the rationale for proviral drug resistance and tropism assays in virologically suppressed patients [50, 51, 135–140].

One small trial assessed the possible role of maraviroc (MVC) in virologically suppressed patients on a three-drug PI/r- or NNRTI-containing regimens by randomizing 30 such patients to either switch to MVC plus two NRTIs or to continue their therapy [141]. Additional inclusion criteria included a negative proviral DNA genotypic test for CXCR4-tropic viruses and no history of NRTI resistance. Although only one of the 15 patients randomized to MVC developed VF—a patient subsequently found to have CXCR4-tropic virus missed by the screening tropism assay—this strategy cannot be recommended until further studies are performed [1].

4.7.2 Triple NRTIs

There have been two meta-analyses of randomized controlled trials published between 2001 and 2013 in which stably suppressed patients on two NRTIs plus an NNRTI or a PI were placed on a three-drug ABC-containing regimen—usually AZT/3TC/ABC—to simplify therapy and reduce lipid levels [142, 143]. These meta-analyses concluded that AZT/3TC/ABC appeared to be as effective as
continued two-class therapy but cautioned that the ARVs used in those studies were often inferior to those currently used. However, as a result of the established inferiority of AZT/3TC/ABC for first-line therapy [144], this regimen is generally not considered a suitable option for maintenance therapy in virologically suppressed patients.

4.7.3 Non-NRTI-, Non-PI-Containing Regimens

There have been several pilot studies of combinations of ARVs including RAL plus MVC [145], RAL plus ETR [146], and the long-acting investigational INSTI cabotegravir in combination with RPV [147]. The RAL plus MVC trial was discontinued prematurely because it was associated with a high rate of virologic failure. VF occurred in only one of 25 patients receiving RAL plus ETR at 48 weeks, two others discontinued because of intolerance.

4.7.4 Long-Acting Parenteral Therapy

Long-acting injectable ARVs may soon provide novel treatment strategies in HIV-infected patients who cannot take oral pills for medical reasons (surgery or transplant), for whom adherence to daily oral medications is poor or for prevention of HIV in uninfected patients. For example, the phase IIb LATTE study used an oral induction phase with cabotegravir (an INSTI) plus two NRTIs; participants who were suppressed after 24 weeks were switched to a maintenance regimen of dual therapy cabotegravir plus RPV. At week 96, virologic suppression for the experimental regimen was similar to EFV plus two NRTIs [147]. Both cabotegravir and RPV have the potential to be administered as long-acting injectable formulations. A study evaluating a similar switch strategy from standard ART to dual therapy oral DTG plus RPV is also underway [148].

5 Conclusions and Future Directions

In patients who have never experienced VF while receiving ART and have no evidence of drug resistance, switching to any of the acceptable DHHS first-line regimens is expected to maintain virologic suppression. An increasing number of clinical trials of once daily FDC combinations offer convenience for patients and have now been shown to be both highly effective for maintaining virologic suppression in this population. Even in these patients with little risk of virologic failure, changes can expose them to medication intolerance or adverse effects, so the indications for switching therapy should be carefully considered.

In virologically suppressed patients with a history of VF or drug resistance, it is more challenging to change therapy while still maintaining virologic suppression. In these patients, it is necessary to consider all past ARV regimens, episodes of VF, and past genotypic resistance tests. Additionally, it is usually necessary to select a regimen with a high genetic barrier to resistance and occasionally necessary to select a regimen containing ARVs from more than two drug classes.

A growing number of studies have included switches to INSTI-based regimens. The pivotal SWITCHMRK trial showed that the substitution of RAL for LPV/r carries a risk of VF and emergent INSTI resistance in patients harboring NRTI-resistant viruses. As a result of this finding, clinical trials of switches to newer INSTIs (EVG/c and DTG) have been performed almost exclusively in patients without a history of past VF or evidence of drug resistance mutations. In contrast, DTG – which has a higher genetic barrier to resistance than RAL or EVG/c and is currently being studied as an option for second-line therapy in patients who have failed a first-line NNRTI-based regimen [105] – may eventually prove to be a switch option in stably suppressed patients with prior VF. Results of a study that switched patients with prior VF who are now suppressed on a complex regimen to the FDC TAF/FTC/EVG/c plus DRV are expected in the near future [149].

Increasing data show that LPV/r or DRV/r, either alone or in combination with 3TC, are effective maintenance regimens in patients without previous PI or 3TC resistance. These regimens are likely to be useful in patients with NRTI toxicity. However, in patients without such toxicity they may be complicated to administer because they require closer follow-up for VF. LPV/r or boosted DRV plus an INSTI have also been effective maintenance regimens, and are useful options for patients with NRTI toxicity. Patients receiving such regimens must be particularly adherent to the boosted PI component of the regimen because non-adherence poses a high risk of emergent INSTI resistance and loss of future options.

Genotypic resistance and co-receptor tropism testing from peripheral blood mononuclear cells are commercially available but not frequently used. They assess the proviral DNA that is integrated into resting memory CD4+ T cells forming an “archive” of resistance mutants that can be sequenced when no virus is detected in the blood. These tests are likely to be reliable when they detect resistance mutations or CXCR4 tropism but the validity of negative assays requires further study.

6 Case Vignettes

6.1 Case 1

A 46-year-old woman with a 15-year history of ART initially with SQV/r/D4T/3TC then with EFV/TDF/FTC
presented with progressive renal dysfunction and a partial Fanconi syndrome. She had no known history of VF or drug resistance. Her CD4 count was 1,056 cells/mm³ and her viral load was consistently below 50 copies/ml. She also has a history of migraine headaches often requiring intravenous dihydroergotamine infusions.

Her progressive renal dysfunction and tubular dysfunction made it urgent to discontinue TDF. An HLA*B5701 test, which was performed to ensure that ABC was a viable option, was negative. Given her preference for a once-daily single-tablet regimen, she was changed to ABC/3TC/DTG. Her renal function has since improved and she has remained virologically suppressed. Simply substituting ABC for TDF was considered to be an inferior option because of the possibility that during her long past period of treatment she may have developed some degree of latent NRTI or NNRTI resistance that might reactivate due to the lower activity of ABC compared with TDF in the face of resistance. A boosted PI regimen was not used because of the established drug–drug interaction between boosted PIs and dihydroergotamine.

### 6.2 Case 2

A 42-year-old man with a history of hypertension had been treated initially with TDF/3TC/EFV and then TDF/FTC/EFV for a total of 9 years. He had no known history of VF and had a nadir CD4 count of 262 cells/mm³. During the past year, he has developed progressively increasing anxiety and depression requiring visits to a psychiatrist who is inquiring about the possibility of discontinuing EFV before starting an antidepressant.

Although it is unusual for EFV-associated neuropsychiatric symptoms to develop so late into therapy, it is possible that EFV could be exacerbating these symptoms even if it is not their primary cause. Considering the absence of any history of VF or drug resistance, the patient has many treatment options including each of the three DHHS-recommended single tablet regimens: TDF/FTC/RPV, TDF/FTC/EVG/c, and ABC/3TC/DTG. The newly available TAF/FTC/EVG/c FDC is also supported by evidence but not yet incorporated into treatment recommendations. In this patient, we chose to use TDF/FTC/RPV. The patient’s depression and anxiety improved dramatically within several weeks and his virus load remained undetectable.

### 6.3 Case 3

A 72-year-old man with a more than 20-year history of ART has been stably suppressed on TDF/FTC/ATV/r for 8 years. He began therapy in the early 1990s with a series of mono- and dual-NRTI regimens. He has a history of EFV and LPV/r intolerance. His nadir CD4 count was 70 cells/mm³ and his last genotypic resistance test prior to starting TDF/FTC/ATV/r contained multiple thymidine analog resistance mutations, M184V, and several NNRTI-resistance mutations. A recently ordered HLA-B*5701 test was negative. Although he has no history of coronary artery disease, he has inquired about the possibility of switching ATV/r to an INSTI to reduce that risk. Interestingly, a genotypic resistance test of proviral DNA detected the presence of three TAMs, M184V, and two NNRTI-resistance mutations that closely matched the pattern of resistance observed on his last test prior to starting his current regimen.

At the time the patient first made this inquiry DTG had not yet been approved and it was decided that a switch to RAL or to TDF/FTC/EVG/c would entail an increased risk of VF because of the partially compromised NRTI backbone. Since then DTG has been approved and, due to its higher genetic barrier to resistance, is likely to be associated with a lower risk of VF and emergent INSTI resistance than RAL or EVG/c. However, considering the relatively weak association of ATV/r with an atherogenic lipid profile and the fact that clinical trials of DTG in this scenario have not yet been completed, it was decided to continue the patient’s current therapy.

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**Appendix: Proviral DNA Testing for HIV-1 Drug Resistance and Co-Receptor Tropism in Patients with Virological Suppression**

During its replication cycle, proviral HIV-1 DNA integrates into host chromosomal DNA and is then usually expressed leading to productive infection and cell killing. In resting memory CD4+ T cells, however, integrated proviral DNA may persist for many years forming a stable reservoir of latently infected cells [135]. As a result, proviral DNA levels in peripheral blood mononuclear cells (PBMCs) remain detectable even in patients receiving ART who have undetectable plasma HIV-1 RNA levels [136, 7].
There is a strong but imperfect correlation between the drug-resistance mutations (DRMs) in PBMC proviral DNA and plasma HIV-1 RNA from the same blood samples [50, 137]. Indeed, in patients with suppressed plasma HIV-1 RNA levels, the DRMs present in PBMCs are consistent with a patient’s past ART history and previous genotypic tests [51, 138, 139]. However, PBMC sequencing does not necessarily detect all of the DRMs that were previously present in samples from patients who had past genotypic resistance tests [51, 139, 150]. Therefore, PBMC genotypic resistance should be used in conjunction with past genotypic test results and should be interpreted in light of drug resistance likely to have emerged during past episodes of virological failure.

Most studies suggest that little HIV-1 evolution occurs during ART-mediated virological suppression—particularly with most current regimens [15]. Therefore, if co-receptor tropism was determined shortly before virological suppression, it is unlikely to have changed during therapy [140]. However, if tropism was not determined previously and if a switch to maraviroc is being considered, genotypic or phenotypic PBMC tropism tests are considered reasonable but have not been validated for predicting maraviroc activity [1].

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