HIV-1 drug resistance and resistance testing

Dana S. Clutter, MD a,⁎, Michael R. Jordan, MD, MPH b,c, Silvia Bertagnolio, MD d, Robert W. Shafer, MD a

a Division of Infectious Diseases and Geographic Medicine, Stanford University School of Medicine, 300 Pasteur Drive, L-134, Stanford, CA 94304, United States
b Tufts University School of Medicine, Boston, MA 02111, United States
c Tufts Medical Center, Boston, MA 02111, United States
d HIV Department, World Health Organization, Avenue Appia 20, Geneva, Switzerland

Abstract

The global scale-up of antiretroviral (ART) therapy (ART) has led to dramatic reductions in HIV-1 mortality and incidence. However, HIV drug resistance (HIVDR) poses a potential threat to the long-term success of ART and is emerging as a threat to the elimination of AIDS as a public health problem by 2030. In this review we describe the genetic mechanisms, epidemiology, and management of HIVDR at both individual and population levels across diverse economic and geographic settings. To describe the genetic mechanisms of HIVDR, we review the genetic barriers to resistance for the most commonly used ARVs and describe the extent of cross-resistance between them. To describe the epidemiology of HIVDR, we summarize the prevalence and patterns of transmitted drug resistance (TDR) and acquired drug resistance (ADR) in both high-income and low- and middle-income countries (LMICs). We also review two categories of HIVDR with important public health relevance: (i) pre-treatment drug resistance (PDR), a World Health Organization-recommended HIVDR surveillance metric and (ii) and pre-exposure prophylaxis (PrEP)-related drug resistance, a type of ADR that can impact clinical outcomes if present at the time of treatment initiation. To summarize the implications of HIVDR for patient management, we review the role of genotypic resistance testing and treatment practices in both high-income and LMIC settings. In high-income countries where drug resistance testing is part of routine care, such an understanding can help clinicians prevent virological failure and accumulation of further HIVDR on an individual level by selecting the most efficacious regimens for their patients. Although there is reduced access to diagnostic testing and to many ARVs in LMIC, understanding the scientific basis and clinical implications of HIVDR is useful in all regions in order to shape appropriate surveillance, inform treatment algorithms, and manage difficult cases.

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Keywords:
HIV-1
Drug resistance
Antiretroviral therapy
Diagnostic test
Surveillance
Mutations

Infection, Genetics and Evolution 46 (2016) 292–307

⁎ Corresponding author.
E-mail addresses: dclutter@stanford.edu (D.S. Clutter), mjordan@tuftsmedicalcenter.org (M.R. Jordan), bertagnolios@who.int (S. Bertagnolio), rshafer@stanford.edu (R.W. Shafer).
1. Introduction

The world has embarked on a fast-track strategy to achieve the Joint United Nations Programme on HIV/AIDS 90-90-90 target and reduce the number of new HIV infections to 500,000 per year by 2020. Achieving the 90-90-90 target requires that 90% of all people living with HIV will have been diagnosed, 90% of all people with known HIV infection will be receiving antiretroviral (ARV) therapy (ART), and 90% of all people receiving ART will have a suppressed viral load. To meet the targeted reduction in new infections, the incidence must decline over four-fold from 2.1 million in 2015 (UNAIDS, 2014, 2016). To help achieve these goals, the World Health Organization (WHO) in 2016 recommended that all HIV-infected individuals should begin ART as soon after diagnosis as possible (Treat All) and that pre-exposure prophylaxis (PrEP) should be considered for persons at high risk of HIV infection (World Health Organization, 2016b). While the successful implementation of these recommendations will reduce the number of new HIV infections, the dramatic increase in ARV use will likely increase the prevalence of acquired drug resistance (ADR) in treated individuals and transmitted drug resistance (TDR) in newly infected individuals (Phillips et al., 2014b).

Whereas the principles of drug resistance are similar in all populations, differences in access to diagnostic testing and to many ARVs in low- and middle-income countries (LMICs) necessitate different approaches to the management of TDR and ADR. In most LMICs, HIV drug resistance (HIVDR) testing is neither feasible nor routinely recommended for patients receiving ART. Nonetheless, understanding the principles and mechanisms of HIVDR is useful to developing appropriate surveillance, informing treatment algorithms, and managing particularly difficult clinical cases. In high-income countries where drug resistance testing is part of routine care, such an understanding can also help clinicians prevent virological failure (VF) and accumulation of further HIVDR by selecting the most efficacious regimens for individual patients infected with drug-resistant viruses.

Although twenty-seven ARVs belonging to six classes have been approved for HIV-1 treatment, eleven of these ARVs are no longer available patients infected with drug-resistant viruses. We will instead focus on the 16 more commonly used ARVs proved for HIV-1 treatment, eleven of these ARVs are no longer available patients infected with drug-resistant viruses. Additionally, specific fixed dose combinations (FDCs) will not be addressed due to the rapid evolution of generic ARV formulations that currently dominate the ARV marketplace in LMICs (Waning et al., 2010).

2. Mechanisms of HIV-1 drug resistance

HIV-1 genetic variability results from the high rate of HIV-1 reverse transcriptase (RT) processing errors, recombination when more than one viral variants infect the same cell, and the accumulation of proviral variants during the course of infection (Abram et al., 2010; Coffin, 1995; Levy et al., 2004; Mansky, 1996). Although most HIV-1 infections are initiated by a single viral variant (Keele et al., 2008), innumerable variants (or quasispecies) related to the initial transmitted virus emerge within weeks following infection (Coffin, 1995; Perelson and Ribeiro, 2013).

The selection of drug-resistant variants depends on the extent to which viral replication continues during incompletely suppressive therapy, the ease of acquisition of a particular drug resistance mutation (DRM), and the effect of DRMs on drug susceptibility and virus replication. Although naturally occurring drug-resistant viruses arise every day in untreated patients (Perelson and Ribeiro, 2013), these variants rarely rise to detectable levels because they are less fit than drug-susceptible viruses in the absence of selective drug pressure. Indeed, nearly all clinically significant DRMs arise only as a result of selective drug pressure and are otherwise non-polymorphic.

For some ARVs, multiple DRMs are required to reduce susceptibility, while for others a single DRM is sufficient. The number of DRMs required and the effect of each DRM on virus fitness contribute to the ARV's genetic barrier to resistance. DRMs can be categorized as primary DRMs that directly reduce drug susceptibility and accessory DRMs that enhance fitness of variants containing primary DRMs or that contribute

Table 1

<table>
<thead>
<tr>
<th>Drug class</th>
<th>Drug name</th>
<th>Abbreviation</th>
<th>LMIC use</th>
<th>High-income country use</th>
</tr>
</thead>
<tbody>
<tr>
<td>NRTIs</td>
<td>Lamivudine</td>
<td>FTC</td>
<td>First-line, second-line</td>
<td>First-line, second-line</td>
</tr>
<tr>
<td></td>
<td>Emtricitabine</td>
<td>FTC</td>
<td>First-line, second-line</td>
<td>First-line, second-line</td>
</tr>
<tr>
<td></td>
<td>Tenofovir</td>
<td>TDF</td>
<td>First-line, second-line</td>
<td>First-line, second-line</td>
</tr>
<tr>
<td></td>
<td>Abacavir</td>
<td>ABC</td>
<td>Not commonly used (HLA testing required)</td>
<td>First-line, second-line</td>
</tr>
<tr>
<td></td>
<td>Zidovudine</td>
<td>AZT</td>
<td>Alternative first-line, second-line</td>
<td>Salvage</td>
</tr>
<tr>
<td></td>
<td>Elvireniwe</td>
<td>EFV</td>
<td>First-line</td>
<td>Alternative first-line</td>
</tr>
<tr>
<td></td>
<td>Nevirapine</td>
<td>NVP</td>
<td>Alternative first-line</td>
<td>Not commonly used</td>
</tr>
<tr>
<td>PIs</td>
<td>Atazanavir</td>
<td>ATV</td>
<td>Second-line</td>
<td>Alternative first-line</td>
</tr>
<tr>
<td>INSTIs</td>
<td>Darunavir</td>
<td>DRV</td>
<td>Alternative second-line, third-line</td>
<td>First-line, second-line, salvage</td>
</tr>
<tr>
<td></td>
<td>Lopinavir</td>
<td>LPV</td>
<td>Second-line</td>
<td>Not commonly used</td>
</tr>
<tr>
<td></td>
<td>Elvireniwe</td>
<td>EVG</td>
<td>Not commonly used</td>
<td>First-line</td>
</tr>
<tr>
<td>CCR5 Antagonist</td>
<td>Raltegravir</td>
<td>RAL</td>
<td>Alternative second-line, third-line</td>
<td>First-line</td>
</tr>
<tr>
<td></td>
<td>Doluzgravir</td>
<td>DTC</td>
<td>Alternative first-line</td>
<td>First-line, second-line, salvage</td>
</tr>
</tbody>
</table>

Abbreviations: NRTIs: nucleoside reverse transcriptase inhibitors; NNRTIs: non-nucleoside reverse transcriptase inhibitors; PIs: protease inhibitors; INSTIs: integrase strand transfer inhibitors; LMIC: low- and middle-income countries (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016; World Health Organization, 2015a).
susceptibility in cell culture. Susceptibility is usually reported as the ARV concentration that inhibits HIV-1 replication by 50% (IC₅₀). The IC₅₀ of a patient’s virus is compared to that of a drug-susceptible reference strain, and expressed as a ratio, referred to as fold change, of the IC₅₀ of the patient’s virus relative to the reference control. Nearly all susceptibility assays use recombinant viruses created by inserting PCR-amplified patient virus gene segments (protease/RT, integrase, envelope) into the backbone of a wild type laboratory clone (Petrosoulos et al., 2000). Due to cost and lengthy turnaround time, phenotypic resistance testing is typically reserved for drug development, drug resistance research, or complex clinical cases.

The clinical significance of the reductions in in vitro susceptibility reported from phenotypic testing varies markedly among ARVs, owing to both pharmacokinetic and pharmacodynamic factors. For example, the NRTIs must be triphosphorylated to their active form, a process that often occurs at different rates in vivo compared with the cell lines typically used for drug-resistance testing. Additionally, the dynamic susceptibility range, or the fold change in IC₅₀ between wild type and the most NRTI-resistant isolates, is as low as 5-fold for TDF and >200-fold for zidovudine (AZT), lamivudine (3TC), and emtricitabine (FTC) (Melikian et al., 2012; Whitcomb et al., 2003). Similar but less pronounced differences in dynamic susceptibility ranges exist within the NNRTI, PI, and INSTI classes.

Standard genotypic resistance testing (SGRT) involves the use of dideoxynucle terminator Sanger sequencing of protease, RT, and/or integrase to identify established clinically significant DRMs. SGRT is performed on PCR products directly amplified from cDNA that has been reverse-transcribed from plasma RNA. Because the virus population in an individual is typically heterogeneous, it is common for more than one nucleotide (i.e., a mixture) to be present in a given position on Sanger sequence electropherograms. Standardization between laboratories for studies of drug resistance has been facilitated through the use of software tools that apply a consistent approach to identify mixed bases (Woods et al., 2012) and to assess sequence quality (Shafer et al., 2000). Indeed, the information present in an electropherogram provides intrinsic quality control data including the level of background “noise” as indicated by low signal intensities and numerous poorly defined peaks. In addition, a high number of electrophoretic mixtures, atypical mutations, and APOBEC mutations observed in a nucleotide sequence can signal quality control problems (Rhee et al., 2016; Shafer et al., 2000). Furthermore, phylogenetic analysis and distance measurements of sequences can be used to detect possible laboratory contamination or specimen mislabeling at the point of collection.

Although the lower limit of SGRT detection is generally agreed to be around 20%, both point mutation assays (PMAs) and next generation deep sequencing (NGS) technologies often detect drug-resistant variants at levels well below a 20% threshold. However, the frequency with which these ultra-sensitive PMA and NGS assays reliably detect variants present at levels >20% varies widely between studies. Some of the low-abundance DRMs detected by highly sensitive assays may arise from PCR or sequencing errors (Jabara et al., 2011; Keys et al., 2015). Although the use of Primer IDs, random sequence tags incorporated into primers to generate templates with unique primer identification sequences, can reduce such errors (Jabara et al., 2011), this approach is used primarily in research settings. Low-abundance DRMs have been associated with a reduced response to NNRTI-based ART regimens, and the effect is greatest when the DRM-bearing variants are present at frequencies ≥1% (Li et al., 2011). Despite the potential clinical implications of low-abundance DRMs in patients initiating certain regimens, guidelines for the use of these more sensitive technologies have not yet been developed.

In high-income countries, SGRT, which has a retail cost of 400–500 USD, is considered cost-effective at time of diagnosis or treatment initiation and at VF. However, in LMIC settings, models have produced conflicting results regarding the cost-effectiveness of SGRT (Levison et al., 2013; Phillips et al., 2014a; Phillips et al., 2014b; Rosen et al., 2011).
Platforms that are more economical than SGRT may play a role in expanding access to genotypic drug resistance testing, which is currently limited or non-existent in most LMICs.

Due to the high number of sequences obtained with NGS, the cost per sequence obtained with this platform is lower than with SGRT. In NGS deep sequencing the depth of coverage refers to the number of viral variants sequenced, typically several hundred. Researchers have shown that harnessing the cost savings of NGS using a “wide sequencing” strategy, wherein many samples are processed in a single run with a lesser depth of coverage, could be a viable method of genotypic testing in LMICs (Dudley et al., 2012; Lapointe et al., 2015). Cost-effectiveness models using NGS platforms have not been published, but the estimated cost per sample is much lower than SGRT at 1 to 20 USD (Dudley et al., 2012; Lapointe et al., 2015).

Whereas wide sequencing with NGS would require processing batched samples in few centralized laboratories, point-of-care (POC) assays aim to be deployable in a variety of settings, from hospitals to remote clinics, and to deliver actionable results during a single clinical encounter. The platforms in development that appear the most amenable to POC or near-POC use in LMICs are PMAs capable of detecting key DRMs at costs below $5 USD per locus (MacLeod et al., 2015; Mutsvangwa et al., 2014; Panpradist et al., 2016; Zhang et al., 2016; Zhang et al., 2013). A major challenge to the successful development of a PMA is the extreme genetic variability of HIV-1 (Clutter et al., 2016b). The variability surrounding DRM positions will necessitate extensive validation studies using diverse subtypes and may require multiple primers, thus driving up the cost of assay development and production. However, recent efforts to mitigate the effect of genetic variability on the success of a PMA have shown promising results (MacLeod et al., 2015). Notably PMAs do not produce sequences; thus they cannot be used for phylogenetic analyses or the discovery of novel DRMs.

4. Drug resistance test interpretation

Understanding the results of drug resistance tests is one of the most difficult tasks facing HIV care providers. There are many DRMs and they arise in complex patterns that cause varying levels of drug resistance. Some DRMs are responsible for just low levels of resistance to a drug but would not be a contraindication to using that drug if the remaining components of a patient’s regimen were fully active. Further complicating genotypic resistance test interpretation is the recognition that SGRT and even NGS deep sequencing may not detect all of the resistant variants within an individual patient and that inference into the possible presence of these variants will depend on a patient’s past treatment regimens.

A DRM can be characterized according to whether it is selected by therapy and whether it is polymorphic or nonpolymorphic in untreated individuals. The development of a mutation while on ART suggests that the mutation is associated with reduced susceptibility to an ARV in the regimen. This type of evidence is particularly important for DRMs associated with drugs that exhibit only subtle in vitro susceptibility changes. Most major DRMs are non-polymorphic in that they do not occur naturally in the absence of selective drug pressure. An important corollary of this observation is that HIVDR to the current NRTIs, NNRTIs, PIs, and INSTIs rarely occurs in the absence of prior selective drug pressure.

DRMs can also be characterized by their effects on in vitro susceptibility and in vivo response to therapy. The difference in ARV susceptibility between a wild type laboratory clone and a clone bearing a DRM allows for direct assessment of that DRM’s phenotypic effect. The reduction in susceptibility caused by a particular DRM can also be studied in clinical isolates, in which multiple DRMs may be present, through the use of regression analyses where the presence of the DRM is an explanatory variable and the fold reduction in susceptibility is the outcome variable. The regression coefficients obtained from these models indicate the relative contribution of a DRM to reduced ARV susceptibility while controlling for other DRMs in a virus sequence (Melikian et al., 2012; Melikian et al., 2014; Rhee et al., 2010).

The clinical significance of DRMs for ART-naive individuals was first demonstrated by the poor responses to therapy observed in the earliest reports of patients with TDR (Grant et al., 2002; Little et al., 2002). It was then re-enforced in subsequent studies in which initial therapy was selected without the availability of genotypic resistance testing but for which the presence of resistance mutations prior to initiation of therapy was detected retrospectively through the analysis of cryopreserved blood samples (Borroto-Esoda et al., 2007; Chaix et al., 2007; Chung et al., 2014; Hamers et al., 2012; Kuritzkes et al., 2008; Lee et al., 2014; Little et al., 2002; Wittkop et al., 2011).

The predictive value of specific mutations on the response to a new ARV has been shown most convincingly in large clinical trials in which individuals were randomly assigned to receiving a new ARV or placebo in combination with an optimized background regimen (Castagna et al., 2014a; de Meyer et al., 2008; Eron et al., 2013; King et al., 2007; Lanier et al., 2004; Miller et al., 2004; Vingerhoets et al., 2010). Additionally, observational studies have also shown that the predicted activity of a new ARV regimen (often called the genotypic susceptibility score) is a strong independent predictor of the response to that regimen (Rhee et al., 2009; Wittkop et al., 2011).

The Stanford HIV Drug Resistance Database (HIVDB) provides an online genotypic resistance interpretation program to help clinicians and laboratories interpret HIV-1 genotypic resistance tests (http://hivdb.stanford.edu/). The program accepts RT, protease and/or integrase sequences submitted by the user and returns a list of penalty scores for each DRM and an estimate of reduced susceptibility for each ARV. The estimate of reduced susceptibility for each ARV is obtained by adding each of the DRMs’ penalty scores for the ARV. The DRM penalty scores (http://hivdb.stanford.edu/DR/) are based upon the DRM characteristics and the consensus on the clinical significance of a DRM among experts in the field, such as the IAS-USA Drug Resistance Mutations Group (Wensing et al., 2014).

5. NRTI resistance mutations

There are two genetic mechanisms of NRTI resistance: (i) discriminatory mutations that enable RT to distinguish between deoxy-NRTI chain terminators and the cell’s own dNTPs, thus preventing NRTIs from being incorporated into viral DNA; and (ii) primer unblocking mutations that facilitate the phosphorylcytic excision of an NRTI-triphosphate from viral DNA. Unblocking mutations are also referred to as thymidine analog mutations (TAMs) that they are selected by the thymidine analogs AZT and stavudine (d4T) (Tang and Shafer, 2012). Even though d4T has not been used for years in high-income countries and AZT’s use has declined, TAMs have persisted in the circulating viruses this setting due to the relatively higher virus fitness associated with these mutations compared to the discriminatory DRMs. This higher degree of fitness slows reversion to wild type, and allows for ongoing transmission of these DRMs.

The most common NRTI-resistance mutations are M184V/I (Fig. 2). M184 V/I are selected by and cause high-level (~200-fold) reduced susceptibility to the cytosine analog NRTIs, 3TC and FTC. M184V/I is also selected by, and causes low-level resistance to abacavir (ABC). During VF on a regimen containing either 3TC or FTC, M184I often emerges before M184V. However, in most patients M184V will outcompete M184I within several weeks (Frost et al., 2000; Keulen et al., 1996). Despite the high-levels of 3TC and FTC phenotypic resistance associated with M184V/I, it is common practice to continue to use these NRTIs to treat viruses with M184V/I because maintenance of these DRMs reduces virological fitness (Campbell et al., 2005; Castagna et al., 2006; Diallo et al., 2003; Eron et al., 1995) and increases susceptibility to AZT and TDF (Kuritzkes et al., 1996; Melikian et al., 2012; Miller et al., 2004; Nijhuis et al., 1997; Whitcomb et al., 2003). In addition, 3TC and FTC are well tolerated with few associated side effects or toxicities.
The next common discriminatory mutations include K65R, K70E/G/Q, L74V/I, Y115F and the Q151M complex of mutations. K65R is selected primarily by TDF and to a lesser extent by ABC and d4T. K65R increases susceptibility to AZT. Although K65R reduces TDF susceptibility in vitro by only two-fold, it is, after M184V, the most common DRM in patients with VF on a TDF-containing regimen.

L74V/I and Y115F are selected primarily by ABC. However, L74I and Y115F are also often selected by TDF, particularly in LMMCs where resistance tests are often performed after prolonged VF (Skhosana et al., 2015; Van Zyl et al., 2013). K70E/G/Q are also selected by TDF and ABC-containing regimens and are associated with minimal reductions on in vitro susceptibility to these NNRTIs (Tenores Study, 2016). Q151M is an uncommon DRM that usually occurs only in heavily treated patients with prolonged VF. It usually occurs in combination with several accessory DRMs (A62V, V75I, F77L and F116Y) and is associated with high-level resistance to AZT and ABC and intermediate resistance to TDF, 3TC, and FTC.

The TAMs include M41L, D67N, K70R, L210W, T215F/Y and K219Q/E. Although they are selected only by AZT and d4T, they confer lower levels of cross-resistance to TDF and ABC. TAMs occur in two distinct but overlapping patterns denoted as type I and type II (Marcelin et al., 2004; Tang and Shafer, 2012). The type I TAMs, which include M41L, L210W and T215Y, cause higher levels of phenotypic and clinical cross resistance to TDF and ABC than the type II TAMs (D67N, K70R, T215F and K219Q/E). The presence of all three type I TAMs markedly worsens clinical response to TDF- and ABC-containing regimens (Lanier et al., 2004; Miller et al., 2004).

The TAMs and discriminatory mutations other than M184V/I occur through two separate, opposing pathways (Boucher et al., 2006; Parikh et al., 2006; Rhee et al., 2007). Viruses already containing multiple TAMs are more likely to develop additional TAMs than K65R when treated with TDF or ABC. Some viruses with multiple TAMs will also develop a double-aminoc acid insertion at RT position 69 referred to as T69S,SS (indicating a wildtype threonine replaced by three serines) or simply T69insertion. In combination with multiple TAMs this DRM causes high-level resistance to AZT, ABC, and TDF, and intermediate resistance to 3TC and FTC (Masquelier et al., 2001; Winters et al., 1998).

### 6. NNRTI resistance mutations

The NNRTIs have a relatively low genetic barrier to resistance. To develop high-level resistance, one DRM in the case of nevirapine (NVP), one to two DRMs in the case of efavirenz (EFV), and two DRMs in the case of etravirine (ETR), are required (Melikian et al., 2014; Sluis-Cremer and Tachedjian, 2008; Vingerhoets et al., 2010). Although rilpivirine (RPV) has a similar structure to ETR, its genetic barrier to clinically significant resistance is lower than ETR because it is administered at a much lower dose (Cohen et al., 2011; Rimsky et al., 2012).

There is a high level of cross-resistance within the NNRTI class as a result of two mechanisms: (i) most NNRTI-resistance mutations reduce susceptibility to two or more NNRTIs (Melikian et al., 2014); and (ii) the low genetic barrier to NNRTI resistance allows for the emergence of multiple independent NNRTI-resistant lineages, even if not all are detectible by SGR (Antinori et al., 2002; Ruxrungtham et al., 2008; Tang and Shafer, 2012; Varghese et al., 2009).

The most common NNRTI mutations are L100I, K101E/P, K103N/S, V106A/M, Y181C/I/V, Y188C/H/L, G190A/S/E, and M230L (Fig. 3). Each of these can cause intermediate or high-level phenotypic resistance to NNRTIs. With the exception of Y181C/I/V, they cause intermediate- or high-level phenotypic resistance to EFV (Melikian et al., 2014). Although Y181C alone reduces EFV susceptibility by only 2-fold, it has been associated with a reduced response to EFV in patients with virological failure on an NVP-containing regimen possibly as a result of minority variant mutations 15608S22. With the exception of K103N/S, V106A/M, Y188C/H, and G190A/S, they cause phenotypic ETR and RPV resistance (Melikian et al., 2014).

In addition, the previously unrecognized mutations E138K/G/Q/A are among the most common mutations to emerge in patients receiving RPV (Rimsky et al., 2012). Although E138K reduces RPV susceptibility only minimally, it is among the most common DRMs to emerge in patients with VF on a first-line RPV-containing regimen. E138A is a notable DRM for RPV because it occurs in up to 5% of ARV-naïve patients, depending on virus subtype (Sluis-Cremer et al., 2014), and was an exclusionary criterion for enrollment in the clinical trials of Complera (TDF/FTC/RPV).

Because it has the highest genetic barrier to resistance in its class, ETR is the most commonly used NNRTI in patients previously failing other NNRTIs and is often combined with other drugs to which the virus is likely to be susceptible (e.g. boosted PIs). In studies of treatment-experienced patients initiating boosted darunavir (DRV/r), NNRTIs, and ETR, a scoring system based on the number and weight of ETR DRMs was shown to be useful in predicting VF (Vingerhoets et al., 2010).

### 7. PI resistance mutations

Boosted lopinavir (LPV/r), boosted atazanavir (ATV/r), and DRV/r are the three most commonly used PIs. ATV/r and DRV/r are the two most commonly recommended PIs in high-income countries (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016). LPV/r is widely used in second-line regimens in LMICs. LPV/r and DRV/r have high genetic barriers to resistance in that multiple DRMs are required before antiviral activity is compromised (de Meyer et al., 2008; King et al., 2007). LPV/r and DRV/r have also been shown to be effective at maintaining virological suppression in ARV-experienced patients undergoing regimen simplification (Bartlett et al., 2012; Collins et al., 2012).
Several lines of evidence suggest that ATV/r has a lower genetic barrier to resistance than LPV/r and DRV/r: (i) fewer mutations are needed for reductions in in vitro susceptibility; (ii) smaller reductions in in vitro susceptibility are required to increase rates of clinical virological failure (Monogram Biosciences, 2016; Naeger and Struble, 2006); (iii) ATV/r monotherapy has also been less effective than LPV/r and DRV/r for regimen simplification (Castagna et al., 2014b; Guiguet et al., 2012; Karlstrom et al., 2007).

PI DRMs develop much less frequently in patients receiving an initial LPV/r, ATV/r, or DRV/r-containing regimen than do NRTI and NNRTI DRMs in patients receiving NRTI/NNRTI-containing regimens (Barber et al., 2012; Dolling et al., 2013; El Bouzidi et al., 2014; Lathouwers et al., 2011; Molina et al., 2008; Mollan et al., 2012; Van Yzel et al., 2013). The narrow drug concentration range over which PI levels are both high and low enough to allow viral replication yet high enough to exert selective drug pressure likely accounts for the reduced risk of resistance associated with boosted PIs (Rosenbloom et al., 2012; Siliciano and Siliciano, 2012). Indeed, most patients without PI DRMs who experience VF while on an initial PI-containing regimen achieve virological suppression with improved adherence (Lopez-Cortes et al., 2016; Zheng et al., 2014). Nonetheless, the possibility that mutations outside of protease may also cause VF is actively being investigated (Fun et al., 2012; Rabi et al., 2013). Specifically, it has been proposed that mutations in gag matrix (MA) and the gp41 cytoplasmic domain (gp41-CD) contribute to PI resistance but none consistently responsible for PI resistance have been identified (Gupta et al., 2010; Jinnopat et al., 2009; Sutherland et al., 2014).

Although PI-associated DRMs rarely develop in PI-naïve patients receiving one of the currently recommended boosted PIs, the historical use of unboosted PIs in high-income countries has led to the evolution of many highly PI-resistant viruses. Indeed, >80 non-polymorphic PI-selected mutations have been reported (Rhee et al., 2016; Shahriar et al., 2009). The majority of these reduce in vitro susceptibility to one or more PIs (Rhee et al., 2010). The DRMs with the greatest impact on susceptibility are in the substrate cleft including V32I, G48V/M, I50V/L, 173RT, 190QA, 190TS, 200TV, 205TV, 243RA, 244LA, 245IL, 276TV, 280VR, 282VC, and 286TV. In particular, mutations that reduce susceptibility to one or more PIs are selected with increased frequency in boosted PI therapy compared to unboosted PI therapy (Gupta et al., 2010; Jinnopat et al., 2009; Sutherland et al., 2014).

8. INSTI resistance mutations

Fig. 5 shows the most common INSTI-associated DRMs. More information is known about the DRMs selected by raltegravir (RAL) because it was the first FDA-approved INSTI. RAL resistance occurs by three main, often overlapping mutational pathways with the following signature DRMs: N155H ± E92Q; Q148H/R/K ± G140S/A; and Y143C/R (Fig. 4). For instance, in addition to several mutations in the enzyme flipping M46L and I54V/M/L/T/S/A, and in the enzyme core L33F, L76V, and N88S also markedly reduce susceptibility. Most PI-resistant viruses also require one or more additional protease mutations and one or more compensatory gag cleavage-site mutations (Foukakis-Murzyczki et al., 2007; Fun et al., 2012). The requirement for multiple accessory mutations outside of protease almost certainly contributes to the high genetic barrier to resistance of boosted PIs.

The ATV-selected DRM, IS0L, is notable for causing high-level resistance to just one PI. Each of the remaining major mutations reduces susceptibility to two or more PIs. Due to the large number of PI DRMs and the fact that different DRMs at the same position can have markedly different effects on PI susceptibility, PI cross resistance is complex (Tang and Shafer, 2012). This is particularly the case for DRMs at positions 50, 54 and 82 (Rhee et al., 2010). Due to their high genetic barriers to resistance, DRV/r and LPV/r are often used in patients failing therapy, even those who are PI-experienced. In such patients, multiple models incorporating information from 10 or more PI DRM positions have been shown to predict virological responses to LPV/r (Kempf et al., 2002; King et al., 2007). In studies evaluating the effect of baseline DRMs on response to DRV/r in PI-experienced patients, reduced responses were associated with 3 to 5 of the following DRV DRMs: V11I, V32I, L33F, L47V, I50V, I54L/M, T74P, L76V, I84V, and L89V (de Meyer et al., 2008; Pozniak et al., 2008).
much lower levels of phenotypic resistance, they are likely a harbinger of high-level resistance.

Whereas the published data on RAL-resistant isolates are from patients receiving first-line RAL-containing regimens and from patients receiving RAL for late salvage therapy, most published experience with elvitegravir (EVG) resistance is from virus isolates obtained during first-line EVG-containing regimens. EVG shares two mutational pathways with RAL: N155H ± E92Q and Q148H/R/K ± G140SA. E92Q alone is particularly common in patients receiving EVG and is often the most common EVG-resistance DRM in clinical trials (White et al., 2011). There are also two non-polymorphic DRMs – T66I/K and S147G – that occur more commonly with EVG and cause lower-levels of cross-resistance to RAL (Afram et al., 2013).

Considering the extensive overlap between the genetic mechanisms of resistance to RAL and EVG, there are no scenarios in which patients with VF while receiving one of these INSTIs should be switched to the other INSTI. Moreover, even viruses with DRMs that do not appear to cause resistance to both INSTIs when they occur in isolation (e.g. Y143C/R for RAL) may be associated with higher than expected levels of cross-resistance if one or more of the accessory DRMs in theFig. 5legend are present (Huang et al., 2013).

In contrast to RAL and EVG, multiple DRMs are required for clinically significant dolutegravir (DTG) resistance. Most data on the genetic mechanisms of DTG resistance have been obtained from RAL-experienced patients who received DTG salvage therapy in the VIKING trials (Castagna et al., 2014a; Eron et al., 2013). In these trials, patients with viruses containing a Q148 mutation in combination with E138 ± G140 mutations were at increased risk of VF. A number of accessory INSTI-resistance DRMs and N155H also appear to increase the risk of cross-resistance if one or more of the accessory DRMs in the Fig. 5 legend are present (Huang et al., 2013).

In vitro susceptibility data have been highly inconsistent possibly because viruses with this mutation replicate poorly.

9. HIV-1 tropism

Maraviroc allosterically inhibits gp120 Env of CCR5 (R5)-tropic HIV-1 strains from binding to the seven-transmembrane G protein-coupled R5 receptor (Tan et al., 2013). Whereas HIV-1 gp120 binds to the N-terminus and second extracellular loop region of R5, maraviroc binds to a pocket formed by the transmembrane helices (Tan et al., 2013). In patients receiving R5 inhibitors, the most common mechanism of VF is the expansion of pre-existing CXCR4 (X4) tropic viruses that are intrinsically resistant to R5 inhibitors (Gulick et al., 2008; Westby et al., 2006). HIV-1 can also develop maraviroc resistance mutations that allow HIV-1 gp120 to bind to an inhibitor-bound R5 receptor. However, reports of such resistance have been documented primarily in vitro and in only a small number of clinical viral isolates (Jiang et al., 2015; Westby et al., 2007).

More than 80% of patients are initially infected with exclusively R5 tropic viruses. X4 tropic viruses usually emerge during later stages of infection, and about 50% of patients chronically infected with HIV-1 harbor X4 tropic viruses (Brumme et al., 2005; Hunt et al., 2006; Melby et al., 2006; Poveda et al., 2006). X4-tropic viruses are often present in low proportions relative to R5-tropic viruses when they emerge (Brumme et al., 2005; Wilkin et al., 2007).

A tropism assay should be performed to confirm that a patient is not infected with X4-tropic variants before using maraviroc. Tropism can be detected phenotypically or genotypically. The phenotypic Trofile test assesses the tropism of complete env genes amplified from patient samples using reporter cell lines expressing R5 or X4 (Whitcomb et al., 2007). Given a sufficiently high plasma virus level, it has the sensitivity to detect X4 variants at populations < 1% (Su et al., 2009).

Genotypic testing for tropism involves sequencing the envelope V3 loop and using algorithms to predict tropism from the sequence. Positively charged residues at positions 11 and 25 of the V3 loop of gp120 and several less common combinations of mutations primarily but not exclusively within the V3 loop are associated with X4 tropism (Hartley et al., 2005). SGRT has a specificity of ~ 85% and sensitivity of 50–70% for detecting the presence of X4 tropic variants. The low sensitivity of SGRT arises from the fact that X4 tropic variants are often present at levels below the 20% sensitivity of SGRT and because some of the determinants of tropism lie outside of the V3 loop. However, sequencing multiple independent aliquots of viral cDNA or using NGS deep sequencing technology can increase the sensitivity of genotypic tropism testing (Archer et al., 2012; Swenson et al., 2010). Several clinical laboratories in high-income countries offer one or more genotypic approaches for detecting minor X4 tropic viruses.

10. Effect of subtype on HIV-1 drug resistance

Subtype B viruses comprise ~ 95% of HIV-1 strains in the U.S. and >60% of strains in Europe but only about 10% of strains worldwide. Subtype B viruses played an outsized role in the earliest ARV development efforts and in the earliest studies of ARV resistance. Nonetheless, existing ARVs are equally effective at treating subtype B and non-subtype B viruses (Bannister et al., 2006; Geretti et al., 2009; Scherrer et al., 2011). Moreover, each of the mutations that cause resistance in subtype B viruses also cause resistance in non-B subtypes.

However, several DRMs preferentially occur in certain HIV-1 subtypes. The majority of these subtype-specific differences in DRM distribution are attributed to differences in codon usage. The NNRTI DRM V106M occurs more often in subtype C viruses from patients treated with NVP or EFV because V106M requires a single base-pair change in subtype C viruses – GTG (V) → ATG (M) – but a two base-pair change in all other subtypes – GTA (V) → ATG (M) (Bremer et al., 2003). CRF01_AE viruses preferentially develop the NNRTI DRM V175M (Ariyoshi et al., 2003), subtype G viruses preferentially develop the PI DRM V82M (Palma et al., 2012), and subtype A viruses from the former Soviet Union (AFSU) preferentially develop the NNRTI DRM G190S (Kolomeets et al., 2014) by a similar mechanism.

In subtype B viruses, integrase G140 is encoded primarily by GGT or GGC whereas for nearly all other subtypes, G140 is encoded primarily by
GCG or GGA (Doyle et al., 2015). Therefore, a single G to A change at the first position of the G140 codon results in a serine in subtype B viruses, but not in other subtypes. G140S is an important compensatory mutation for Q148 mutations and this may explain why Q148 mutations occur significantly more commonly among subtype B patients compared to non-B patients with RAL VF (Boyd et al., 2015; Doyle et al., 2015; Fouriati et al., 2015).

Several lines of evidence suggest that K65R is more likely to emerge in subtype C viruses than viruses belonging to other subtypes. First, biochemical studies have shown that the unique subtype C sequence in the region of K65R, specifically five consecutive adenosines preceding the adenosine at the second position in the K65 codon, facilitates mutation during reverse transcription (Coutsinos et al., 2009; Invernizzi et al., 2009). Second, K65R emerges more rapidly during in vitro passage of subtype C viruses than during passage of subtype B viruses in the presence of tenofovir (Brenner et al., 2006). Third, three retrospective studies have suggested that in patients who develop VF on a TDF-containing regimen are at a somewhat higher risk of developing K65R if they are infected with a subtype C virus (TenoRes Study, 2016; Theys et al., 2013; White et al., 2016). Despite the reported tendency to acquire the K65R DRM, a recent study showed that when demographic and clinical factors are controlled for among patients receiving at TDF-containing regimen, infection with a subtype C virus does not increase the risk of developing VF (White et al., 2016).

11. Transmitted drug resistance

TDR is a type of HIVDR that occurs when primary HIV infection is caused by a DRM-bearing virus. The prevalence of TDR by SGRT is about 12% to 24% in the U.S. and 10% in Europe (Bodin et al., 1999; Booth and Geretti, 2007; Kassaye et al., 2016; Li et al., 2016; Little et al., 2002; Parker et al., 2007; Rhee et al., 2015a; Shet et al., 2006; Simon et al., 2002; Weinstock et al., 2000; Weinstock et al., 2004). While limited data suggest that the prevalence is below 5% in most of the LMICs in sub-Saharan Africa and South/Southeast Asia (Frentz et al., 2012; Rhee et al., 2015a; Wheeler et al., 2010), intermediate TDR rates of 5 to 10% have been reported from Latin America, Eastern Europe, and high-income Asian countries (Frentz et al., 2012; Rhee et al., 2015a). Upward temporal trends in TDR continue to be reported from many regions, largely attributable to increases in NNRTI TDR (Bertagnolio et al., 2016; Gupta et al., 2012; Jordan et al., 2016; Rhee et al., 2015a).

Because DRMs reduce viral fitness, many transmitted drug-resistant viruses gradually revert to wild type over a period of several years (Brenner et al., 2002; Castro et al., 2013; Jain et al., 2011; Little et al., 2008; Pao et al., 2004). DRMs associated with the greatest reductions in viral fitness revert to wild type more quickly than those with less effect on viral fitness. Conversely, those DRMs with a low fitness cost have been shown to persist for a prolonged period following transmission (Mourad et al., 2015).

In the absence of selective drug pressure, PI and NNRTI DRMs gradually revert to wild type after an average of approximately 3 years. Within the NRTI class, the rate of reversion is more variable: M184V usually fades within one year; T215FY also fades rapidly; but T215 revertants persist for more than ten years (Castro et al., 2013; Jain et al., 2011). It is because of this tendency to fade to lower levels over time that the rates of detectable TDR are higher in recently compared with chronically infected patients, and that studies using more sensitive methods for detecting low-abundance drug-resistant viruses (see section on Diagnosis and Management of Drug Resistance) will detect higher rates of TDR than those using SGRT (Buckton et al., 2011; Cozzi-Lepri et al., 2015; Ellis et al., 2009; Goodman et al., 2011; Metzner et al., 2014).

The most common transmitted NNRTI-associated DRMs include K103N, Y181C, G190r, and K101E which combined accounted for >80% of NNRTI-associated TDR in a recent global meta-analysis. The most common NRTI-associated DRMs include M184V; the TAMs, the T215 revertants, T69D, and F77L (Rhee et al., 2015a). Not surprisingly, M184V occurs more commonly in recently infected patients compared with chronically infected patients. The most common PI SDRMs were L90M, M46I/L, I85V, I54V, N88D, I84V, D30N, G73S, and they occurred less frequently (Bertagnolio et al., 2016; Rhee et al., 2015a). Although transmitted INSTI-associated resistance has been practically absent in ongoing surveillance efforts (Casadella et al., 2015; Doyle et al., 2015; Stekler et al., 2015), there have been at least four case reports of INSTI-associated TDR (Boyd et al., 2011; Varghese et al., 2016; Volpe et al., 2015; Young et al., 2011).

TDR arises from two distinct sources: (i) from patients failing therapy who have developed ADR, or (ii) from an ART-naive person with TDR, a phenomenon termed “onward transmission”. With a greater number of intervening transmission events in the case of onward transmission, the prevalence of DRMs with high fitness costs, such that M184V, is likely to decrease. Onward transmission is therefore responsible for established drug-resistant lineages that typically contain fit drug-resistant strains. Such established lineages are often the major sources of TDR in European cohorts (Mourad et al., 2015; Yerly et al., 2009).

In high-income countries, SGRT is recommended at the time of a patient’s initial diagnosis. If therapy is delayed, SGRT may be repeated at the time of ART initiation if the patient is considered to have been at risk for superinfection (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016). If transmitted INSTI drug resistance is suspected, SGRT of the integrase gene should also be performed (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016).

Despite its increasing frequency, there are few published guidelines for treating patients with TDR. In patients with isolated NNRTI resistance, multiple studies indicate that boosted PIs are highly effective for initial treatment (Chaux et al., 2007; Clutter et al., 2016a; Oette et al., 2006; Wirtzop et al., 2011; Zu Knyphausen et al., 2014) and several studies have indicated that standard first-line INSTI-containing regimens are also likely to be highly effective (Clutter et al., 2016a; Kulkarni et al., 2014). In patients with transmitted NRTI resistance, the selection of first-line therapy should be based on the extent of NRTI resistance. Both LPV/r and DRV/r-containing regimens have been shown to be highly effective for treating patients with multiple NRTI and NNRTI-associated DRMs following the failure of a first-line NRTI/NNRTI-containing regimen. Ajoje et al., 2012; Bartlett et al., 2007; Madruga et al., 2007; Paton et al., 2014). Therefore, they are each also likely to be highly effective at treating ARV-naive patients harboring multiple NNRTI-resistance mutations. However, first-line ATV/r, RAL, or ETV-containing regimens should be used only for treating patients with minimal transmitted NRTI resistance. The use of DTG-containing regimens in this scenario may be useful but needs further study.

Due to historically lower levels of TDR in LMICs and cost-effectiveness considerations, individual-level baseline SGRT is not recommended by the WHO in LMICs (Phillips et al., 2014a). However population-level HIVDR surveillance programs are strongly recommended and they have played a key role in tracking TDR trends and in shaping first-line ARVregimen recommendations (see Section 14).

In LMICs, ART programs are facing important challenges such as ARV stock outs and suboptimal retention in care (World Health Organization, 2016c). Patients experiencing HIV treatment interruptions are more likely to carry DRMs. When these patients re-engage into care, they must frequently re-initiate treatment with the same first-line regimen with which they were previously treated, despite the increased risk of resistance relative to the truly ART-naive population (Hamers et al., 2012). Therefore, to assess the predicted activity of first-line ART, the WHO recommends monitoring pre-treatment drug resistance (PDR) not only among individuals initiating ART for the first time but also among individuals re-initiating first-line ART (World Health Organization, 2014a). PDR is indeed the result of either TDR or ADR, the later due to prior ARV treatment among re-starters, prior exposure to PMTCT among women, or prior ARV prophylaxis.
PDR surveys will be critical given the more recently observed signals of high level HIVDR in several LMIC settings, including Angola (16%), Cuba (22%), Mexico (12%), Papua New Guinea (16%), Botswana (10%) and South Africa (14%) (Afonso et al., 2012; Avalia-Rios et al., 2016; Lavu et al., 2015; National Institute for Communicable Diseases, 2016; Perez et al., 2013). In the absence of SGRT-guided therapy, national PDR surveillance among individuals starting ART is critical to predict the activity of available first-line ART and inform the composition of these regimens. Where PDR levels remain low, patients are anticipated to do well on the WHO-recommended first-line regimen of EFV + TDF/3TC or FTC (World Health Organization, 2015a). Where PDR levels are high, alternative first-line regimens may become necessary for all starters or for a subset of those reporting prior ARV drug exposure (that is, people who defaulted from ART and restart, women exposed to PMTCT and perhaps those infected despite ARV drug prophylaxis [i.e. post-exposure prophylaxis and PrEP]). Although these surveys are strongly recommended by WHO and are essential to address data gaps in LMIC, their implementation has been slow due to lack of awareness of HIVDR as a public health threat, limited resources, and competing national and donor priorities. As discussed in Section 14, WHO is developing a coordinated and resourced global action plan for HIVDR.

12. Acquired drug resistance

In most studies, >70–80% of patients with VF develop ADR (Jordan et al., 2016; Kumarasamy et al., 2015; Paton et al., 2014; Prosperi et al., 2011; Schultz et al., 2015; Second-Line Study Group et al., 2013; TenoRes Study, 2016). M184V/I is usually selected as the initial NRTI DRM for cytidine analog (3TC or FTC) containing regimens. Among patients receiving an NNRTI-containing regimen, one or more NNRTI DRMs are also typically seen early in VF. Other NRTI DRMs usually occur after M184V/I and the NNRTI DRMs (Jordan et al., 2016; Rhee et al., 2015b). INSTI DRMs may develop early in treatment, but PI DRMs are rare and are generally a late occurrence. Resistance is detected more often in patients with infrequent viral load testing becauseVF is usually detected later, after DRMs have had time to accumulate (Gupta et al., 2009). Through this mechanism of prolonged VF, infrequent viral load testing may jeopardize the efficacy of second-line options, particularly NNRTI- and INSTI-based regimens.

Patterns of ADR at VF vary according to regimen and patient adherence levels. Indeed, the lack of any ADR in the setting of treatment failure usually indicates non-adherence. Because all recommended first-line regimens contain a cytidine analog, M184V is the most commonly acquired DRM in both high-income countries and LMICs (Rhee et al., 2015b). The later NRTI DRMs to develop depend on the second NRTI in the regimen. Although both d4T and AZT are thymidine analogs, they differ somewhat in the mutations they select. AZT selects almost exclusively TAMs whereas d4T selects primarily TAMs but may also select discriminatory mutations including K65R. TDF selects primarily for K65R and less commonly for K70E/G/Q and Y115F (Jordan et al., 2016; TenoRes Study, 2016; Van Zyl et al., 2013). Abacavir selects primarily L74V/I, Y115F, and less commonly K65R.

NNRTI DRMs are selected early and often by NNRTI-based regimens, occurring in the vast majority of patients with VF (Delaugerre et al., 2001; Molina et al., 2011; Second-Line Study Group et al., 2013; TenoRes Study, 2016). In both high-income countries and LMICs, K103 N followed by Y181C, G190A/S, Y188L, and V106M are the most commonly observed acquired NNRTI DRMs (Rhee et al., 2015b). After failing a first-line NNRTI-based regimen containing EFV or NVP, cross-resistance to the second generation NNRTIs RPV and ETR is common and has been associated with VF (Ruxrungtham et al., 2008; Teeranaipong et al., 2016; Theys et al., 2015).

In previously PI-naive patients, PI DRMs often occur in approximately 10% or fewer patients failing PI-based therapy (Barber et al., 2012; Dolling et al., 2013; El Bouzidi et al., 2014; Mollan et al., 2012). As noted previously, a high proportion of patients with VF on an initial PI-based regimen but without PI DRMs will achieve virological suppression with improved adherence alone (Zheng et al., 2014).

In patients with VF on a first-line RAL- or ETV-based regimen, INSTI resistance is not uncommon (Kulkarni et al., 2014; Rockstroh et al., 2013). In contrast, treatment-emergent INSTI resistance is rarely observed in patients failing first-line DTG-based regimens (Raffi et al., 2013; White et al., 2014b).

In high-income countries, SGRT is recommended in patients with VF when the viral load is >1000 copies/mL (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016), and should be considered when the viral load is >50 copies/mL (Gonzalez-Serna et al., 2014; Santoro et al., 2014). In patients with VF, the ability to detect ADR is much higher, if SGRT is performed while a patient is on the failing regimen or within four weeks of discontinuation (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016). Genotypic testing can indicate whether VF is due primarily to non-adherence – i.e., if no DRMs are present – and can also be used to help design second-line therapy when ADR is the cause of VF. Patients who develop VF without DRMs can often be treated successfully with adherence counseling alone (Bonner et al., 2013; Hoffmann et al., 2014). In contrast, patients with VF who have developed DRMs on a first-line regimen usually require a change in ARVs.

In high-income countries where SGRT is routinely performed in the setting of VF, a second-line regimen should include at least two, preferably three, fully active agents. As a general principle, adding a single fully active ARV to a failing regimen is not recommended due to the risk of selecting for resistance to the new ARV. A fully active agent is defined as an ARV which is expected to have activity based on the patient’s treatment history and SGRT results (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016). Managing patients with ADR who are failing second-line regimens is more challenging, and often requires expert guidance. With complex patterns of resistance, phenotypic testing may be useful to determine the activity of various ARVs (Panel on Antiretroviral Guidelines for Adults and Adolescents, 2016). The number of available fully active ARVs diminishes with each successive treatment failure. Therefore, a salvage regimen in patients who have experienced multiple VFs and who have multi-class resistant viruses is likely to be complicated and to require multiple ARVs that may have only partial residual activity.

Late salvage therapy ARVs can be placed into the following categories: (i) ARVs belonging to a previously unused drug class. Enfuvirtide is a highly active treatment option for late salvage therapy. However, it is not a popular treatment choice among clinicians and patients because of its requirement for parenteral intramuscular administration. Although maraviroc is well tolerated, it is often not an option because patients with multiple VFs tend to have more advanced immunosuppression and an increased likelihood of harboring X4-tropic viruses; (ii) ARVs belonging to a previously used drug class that retain significant antiviral activity including the PIs DRV/r and tipranavir/r, the INSTI dolutegravir, and in some instances the NNRTI ETR; (iii) NRTI combinations, which often retain in vivo virological activity even in the presence of reduced in vitro NRTI susceptibility; (iv) ARVs associated with previous VF episodes and treatment-emergent drug resistance that may have regained their activity as a result of viral reversion to wild type; and (v) investigational ARVs (Tang and Shafer, 2012).

In most LMICs, a diagnosis of ADR is made empirically and the selection of second-line therapy is guided by WHO recommendations. A patient is presumed to have ADR when two viral loads performed 3 months apart are both above 1000 copies/mL, and a switch to second-line therapy is indicated (World Health Organization, 2015b). The preferred second-line options include a boosted PI, namely ATV/r or LPV/r, and two NRTIs. Subsequent options include DRV/r plus two NRTIs or LPV/r plus RAL. These alternatives are equally efficacious, but due to higher costs they are not currently preferred options in second-line regimens. Another consideration is that, as previously noted, PI-based regimens rarely select for PI DRMs, but failure of a RAL-containing...
regimen often selects for INSTI DRMs that may jeopardize INSTI activity in subsequent regimens. Second-line NRTI selection is directed by the first-line NRTIs used: if VF developed on TDF/FTC or 3TC, then AZT/3TC is recommended; if VF developed on AZT/3TC, then TDF/FTC or 3TC is recommended (World Health Organization, 2015a). In some LMICs with limited access to SGRT, resistance testing is recommended at VF of second-line therapy (Republic of South Africa Department of Health, 2014) and is managed according to the resistance detected and regimens available.

13. PrEP and drug resistance

As of September 2015, the World Health Organization has recommended TDF-containing pre-exposure prophylaxis (PrEP) for men who have sex with men, heterosexual-active men and women, and intravenous drug users considered at substantial risk for HIV-1 infection (World Health Organization, 2015c). Although the risk of HIV-1 infection is markedly reduced in patients receiving PrEP, those who become infected are theoretically at high risk of developing drug resistance. In a systematic review and meta-analysis of PrEP trials containing TDF, 11 (0.1%) of 9222 PrEP users developed FTC or TDF DRMs. However, because 7 of these 11 infections actually had acute HIV-1 infection at the time of PrEP initiation, the rate of breakthrough infection with resistance is even lower (Baeten et al., 2012; Fonner et al., 2016; Liegler et al., 2014; Thigpen et al., 2012; World Health Organization, 2015c). It has been estimated that for every case of FTC resistance attributed to starting PrEP in the setting of unrecognized acute HIV-1 infection, between 8 and 50 HIV-1 infections are prevented (Fonner et al., 2016; World Health Organization, 2015c). The most commonly reported PrEP-associated DRMs are the FTC-associated M184V/I mutations followed by the TDF-associated K65R mutation (Baeten et al., 2012; Liegler et al., 2014; Thigpen et al., 2012). The risk of PrEP-associated ADR can likely be mitigated through routine frequent laboratory testing to identify incident HIV-1 infections and initiate ART in any incident cases detected. Despite the very low levels of resistance observed in clinical trials, resistance may increase as PrEP interventions are more widely implemented, particularly in settings where unrecognized HIV infections may be exposed to PrEP for longer periods.

14. HIV drug resistance surveillance

Between 2004 and 2013, the World Health Organization recommended the surveillance of TDR among recently infected populations (Bennett et al., 2008; Myatt and Bennett, 2008; Shafer et al., 2008). The WHO-recommended method for surveillance of TDR limited enrollment to ART-naïve populations likely to have been recently infected, and assessed for the presence of 93 non-polymorphic DRMs found in all subtypes, termed surveillance DRMs (SDRMs) (Bennett et al., 2009; Shafer et al., 2008).

In 2015, WHO revised the recommendations and prioritized the surveillance of drug resistance among all populations initiating ART, regardless of duration of infection or prior ARV drug exposure(s), primarily because of its programmatic impact in guiding recommended first-line regimens and feasibility. The WHO’s HIVDR Surveillance and Monitoring approach consists of surveillance of PDR (World Health Organization, 2014a), ADR (World Health Organization, 2014b), HIVDR in infants naïve to ART, and annual monitoring of early warning indicators (EWIs) of HIVDR, defined as clinic- and program-related factors associated with the emergence of HIVDR. The surveys provide nationally representative estimates of HIVDR and aim to guide the selection of first,-second-, and third-line regimens (World Health Organization, 2015d).

The wide implementation of the newly released WHO HIV Consolidated Guidelines recommending initiation of ART to all and scale up of PrEP are expected to lead to a marked decrease in HIV incidence. However, among those patients who become infected a higher proportion of individuals will be infected primarily with a drug-resistant strain and a higher proportion of all HIV-infected patients will be at risk for ADR. To address this concern and ensure a coordinated action to monitor and respond to HIVDR, the WHO is currently developing a Global Action Plan on HIV Drug Resistance. The development of this five-year plan reflects a global consensus that HIVDR in LMICs requires a coordinated and resourced response. The Global Action Plan is an agreement by key partners about their respective roles in preventing, monitoring, and responding to HIVDR (World Health Organization, 2016a).

15. Summary and conclusions

An unprecedented scale-up of ART has been observed in the last 10 years: at the end of 2015, 17 million people were receiving antiretroviral therapy. However, the HIVDR emergence associated with increased ART use can compromise the effectiveness of antiretroviral drugs. Resistance is an inevitable consequence of any interventions that, through the use of ARVs, aim to reduce mortality, morbidity and HIV transmission. While concern for resistance should never limit ART access for all HIV-infected individuals, the long-term implications of earlier initiation on adherence and drug resistance need to be closely monitored. Knowledge of the mechanisms, implications, and patterns of HIVDR can be harnessed to both select effective SGRT-guided therapy for individuals and/or to adapt national treatment guidelines according to surveillance findings following the public health approach to HIV treatment.

Potential conflicts of interest

DSC receives research funding through the Bristol-Myers Squibb Virology Fellows Research Program. RWS is a consultant for Celebra, and receives research funding from Roche Molecular, Gilead Sciences, Bristol-Myers Squibb, and Merck. The other authors have no potential conflicts of interest to disclose.

Disclaimer

The opinions and conclusions in this article are those of the authors and do not reflect those of the respective institutions, including the World Health Organization.

Acknowledgments and funding

This work was supported by a KL2 Mentored Career Development Award of the Stanford Clinical and Translational Science Award to Spectrum [grant number NIH KL2 TR 001083 and NIH UL1 TR 001085 to DSC], by the National Institutes of Health [grant number R01 AI068581 to RWS], by CFA [grant number 1P30AI142853 to MRJ] and by the Christine E. Driscoll O’Neill and James M. Driscoll, Driscoll-O’Neill Charitable Foundation [support for MRJ].

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