Viral Suppression in the Setting of Multidrug Resistance to Nelfinavir, Tenofovir and Abacavir: A Case Report

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Abstract

One major barrier to achieving the primary goals of antiretroviral therapy is the development of drug resistance mutations to antiretroviral agents which render them less effective. For patients who are treatment experienced with many drug resistance mutations, selection of an effective antiretroviral regimen becomes challenging. We report a case of a 55 year old male on tenofovir, abacavir and nelfinavir with multiple resistance mutations to each of these agents, yet able to maintain viral suppression and high CD4 T cell counts. This patient was subsequently switched to protease inhibitor monotherapy for the management of HIV in the setting of multidrug resistance.

Keywords: HIV; Drug resistant mutations; Proviral DNA genotype; Protease inhibitor monotherapy

Introduction

The primary goals of antiretroviral therapy (ART) in the treatment of HIV-1 infection are to maximally and durably suppress plasma HIV-1 RNA and to preserve immunologic function [1]. A major barrier to achieving these goals is development of resistance mutations to antiretroviral (ARV) agents which render them less effective. Drug resistance mutations (DRMs) may cause resistance to a single ARV agent or confer cross-resistance to an entire drug class, potentially limiting future therapeutic options [2].

DRMs may be identified with resistance tests such as HIV-1 genotypes, which are recommended for use in patients who are suspected to have virologic failure [1]. A limitation of the standard genotype is that the viral load of a HIV patient must be at least 500 copies/mL. For patients who have a viral load of less than 500 copies/mL or when the viral load is suppressed (<20 copies/mL), a HIV-1 proviral DNA genotype can provide HIV-1 ARV drug resistance data by next generation sequencing. This is in contrast to the standard genotype that only sequences actively replicating virus HIV-1 RNA [3].

We report here a case of a 55 year old male on ART with multiple documented resistance mutations, yet maintaining high CD4 T-cell counts and viral suppression. We also evaluated the patient's clinical response when switching from his triple ART regimen to PI monotherapy, darunavir/cobicistat (DRV/COBI).

Case Report

A 55 year old Caucasian male presents to Therapeutic Concepts (a private adult outpatient HIV clinic in Houston, TX, USA) for a 3 month follow-up appointment for management of HIV-1 infection in January 2016. The patient is here to consider a potential change in ART as discussed at an earlier appointment date.

The patient was diagnosed with HIV-1 subtype B infection in 1995 at the age of 35 years, when presenting to the hospital with symptoms of pneumocystis pneumonia with a nadir CD4 T-cell count of 60 cells/mm3. The patient’s first reported viral load was in March 1996 with a HIV-1 RNA of 6,654 copies/mL. At that time, the patient was treated with dual nucleoside reverse transcriptase inhibitor (NRTI) therapy of lamivudine and zidovudine. Because of insufficient virologic suppression, he was then switched to two new NRTIs stavudine and didanosine in combination with a non-nucleoside reverse transcriptase inhibitor (NNRTI) nevirapine. The patient did not achieve virologic suppression with this regimen and was switched again to his current ART regimen of the nelfinavir (NFV) 625 mg 2 tablets twice daily, tenofovir disoproxil fumarate (TDF) 300 mg 1 tablet once daily, and abacavir (ABC) 300 mg 1 tablet twice daily, which he reports to be tolerable.

During the past two years, the patient experienced some low level viremia periodically (Figure 1). Because the patient has had no prior history of any other type of resistance testing and low level viremia, this prompted us to order a HIV-1 proviral DNA genotype during the clinic visit in October 2015 (Figure 2), which showed complete resistance to his current ART regimen of NFV, TDF, and ABC. Upon the follow-up clinic visit in January 2016, the patient’s current ART regimen was changed from TDF, ABC and NFV to a simpler and more tolerable regimen of PI monotherapy DRV 800 mg/COBI 150 mg 1 tablet once daily, based on results from the HIV-1 proviral DNA genotype. After the patient had been on this new ART regimen of DRV/COBI for three months, we obtained a new set of labs showing a decrease in his HIV-1 viral load from 40 copies/mL to an undetectable viral load (Figure 1) (LabCorp, Burlington, NC, USA). All other standard laboratory tests including lipids, serum chemistries and complete blood count were within normal limits. A CCR5 delta 32 mutation genetic test was also ordered during the April 2016 clinic visit.
visit (Quest Diagnostics, San Juan Capistrano, CA, USA) to rule out whether the patient was a long-term non-progressor (LTNP); results were negative for this mutation.

Discussion

We report the case of an HIV-1 infected patient whose HIV-1 proviral DNA genotype showed resistance to the patient's current regimen, and yet despite this, was able to maintain an undetectable viral load for several years. The patient's HIV-1 proviral DNA genotype showed that he has four thymidine analogue mutations (TAMs) including M41L, L210W, T215Y and K219N, which together decrease susceptibility and cause intermediate to high-level resistance to ABC.
and TDF. The patient also has the primary mutation D30N, which confers high-level resistance to NFV.

Although the patient showed full resistance to his ART regimen, he was still able to maintain an undetectable viral load with only a few recent “viral blips,” defined as “intermittent episodes of detectable low-level HIV-1 viremia (viral load between 50-199 copies/mL), which are preceded and followed by an undetectable viral load without any change in therapy” [1,4]. Viral blips are a common phenomenon that occurs in about one-quarter of patients who have achieved viral suppression and may be a natural consequence of population dynamics [5]. Because of the patient’s high CD4 T-cell counts and low-level viremia, we considered that the patient may have been a LTNP; however, these results were negative.

One potential explanation for continued viral suppression in the setting of a multidrug resistant virus is poor “viral fitness” [6]. It is possible that the patient’s resistant virus has a replicative capacity that was lower than that of wild-type virus, especially, since the patient has the M184V mutation which is linked to having a reduced HIV replicative capacity [7]. Another potential explanation could be the apolipoprotein B mRNA-editing, enzyme-catalytic (APOBEC)-induced resistance mutation mechanism which may explain the persistence of the D30N mutation in archived cellular proviral DNA despite viral suppression in some patients [8].

Even though it is uncertain why this patient was able to maintain virologic suppression on his resistant ART regimen, we decided to simplify the patient’s ART regimen to an active DRV/COBI monotherapy based on his HIV-1 proviral DNA genotype results. Patients who exhibit characteristics such as high CD4 T-cell counts, sustained viral suppression, and excellent adherence to an ART regimen appear to be the most appropriate candidates for PI monotherapy [9]. This decision was also done intentionally to spare the patient of NRTIs, as he had resistance to the whole class [10]. Our concern for the patient developing drug resistance to DRV/COBI was also done intentionally to spare the patient of NRTIs, as he had the most appropriate candidates for PI monotherapy [9]. This decision of PI monotherapy suggesting no increased risk for the development of DRMs [10]. Some studies have demonstrated success with an integrase strand transfer inhibitor (INSTI) plus boosted PI in treatment-experienced patients [11,12]. We decided to not add dolutegravir at this time and potentially use it at a later time in combination with DRV/COBI plus boosted PI monotherapy: A unique patient who has resistance to his prior ART regimen (NFV, ABC and TDF), yet maintained a suppressed viral load. Potential explanations for this clinical scenario include that the patient might be a LTNP (testing was negative) and poor viral fitness of the patient’s resistant viral strain. Another explanation that was not explored is the possibility for drug-genie interactions. NFV is metabolized to its active metabolite nelfinavir hydroxy-t-butylamide (M8) primarily by cytochrome P450 (CYP) isoenzyme 2C19 [13]. Had the patient remained on NFV, we would have considered both pharmacogenetics testing and therapeutic drug monitoring to evaluate the patient’s CYP2C19 expression. A poor metabolizer phenotype could lead to significantly higher NFV plasma drug concentrations that could have facilitated his viral suppression even in the setting of high-level NFV resistance. Given the patient had many DRMs, a phenotype resistance test may have also been helpful to determine the true sensitivities of the patient’s HIV-1 virus to his current ART and might have helped confirm the replicative capacity of the patient’s HIV-1 virus [1]. However, obtaining results from a phenotype are unlikely since the patient had low-level viremia.

To our knowledge, there have not been any other documented cases of HIV patients maintaining viral suppression with multidrug resistance to their current ART. Ultimately, the patient was successfully simplified to active ART with DRV/COBI monotherapy, maintaining an undetectable viral load on this new regimen, and reporting a much improved quality of life due to considerably decreased pill burden, decreased frequency of administration, and improved tolerability. We learned from this patient that ART is highly individualized and that using HIV-1 proviral DNA genotypes can be advantageous in guiding treatment simplification for treatment experienced patients who are on complex ART regimens, have virologic suppression and/or low-level viremia and have limited information regarding previous resistance testing.

References