Predicted antiviral activity of tenofovir versus abacavir in combination with a cytosine analogue and the integrase inhibitor dolutegravir in HIV-1-infected South African patients initiating or failing first-line ART

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Objectives: The WHO recently recommended the use of a new first-line ART containing dolutegravir. We investigated the efficacy of NRTI backbones (tenofovir or abacavir with a cytosine analogue) in low- and middle-income countries where there is significant prior exposure to antiretrovirals and drug resistance to NRTIs.

Methods: Within the treatment-as-prevention study in South Africa, we selected participants with available next-generation sequencing (NGS) data for the HIV-1 pol gene at trial entry; they were either ART initiators (n = 1193) or already established on ART (n = 94). NGS of the HIV-1 pol gene was carried out using MiSeq technology; reverse transcriptase drug resistance mutations (DRMs) were detected at 5% (DRM5%) and 20% (DRM20%) for all 1287 participants. Genotypic susceptibility was assessed using the Stanford HIVDB resistance interpretation algorithm.

Results: NRTI DRM20% and DRM5% were detected among 5/1193 (0.4%) and 9/1193 (0.8%) of ART initiators, respectively. There was tenofovir exposure in 73/94 (77.7%) of those established on ART, with full susceptibility to abacavir in 57/94 (60.6%) and 56/94 (60.6%) for DRM20% and DRM5%, respectively, while 67/94 (71.3%) and 64/94 (68.1%) were fully susceptible to tenofovir, respectively. The differences between tenofovir and abacavir were not statistically significant at the 20% or 5% variant level (P = 0.16 and 0.29, respectively). NGS detection of variants at the 5% level increased detection of K65R in both naive and treated groups. One of 607 integrase sequences carried a DRM20% (Q148R).

Conclusions: Dolutegravir with a cytosine analogue plus tenofovir or abacavir appears to have similar efficacy in South Africans naive to ART. NGS should be considered in HIV drug resistance surveillance.

Introduction

Countries such as South Africa are implementing a universal test-and-treat strategy recommended by the WHO, therefore increasing the number of people eligible for ART. This policy may lead to higher levels of acquired drug resistance and transmitted drug resistance and compromised ART efficacy in a proportion of patients. For this reason, a robust, cheap and well tolerated fixed-dose combination (FDC) first-line therapy, with a high genetic barrier, is highly desirable. Although NNRTI-based regimens were efficacious, the low genetic barrier to resistance has resulted in an increasing level of pretreatment drug resistance (PDR) to NNRTIs across low- and middle-income countries (LMICs), compounded by the use of thymidine analogues (TAS) prior to availability of tenofovir-based NRTI backbones. In response to rising PDR in LMICs, the WHO recommended in 2017 the use of a novel anti-retroviral, dolutegravir, a second-generation integrase strand transfer inhibitor (INSTI), in countries with a PDR level >10%.

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Cheaper to manufacture, dolutegravir appears to be a good candidate with a high genetic barrier to resistance based on first-line ART studies in northern countries with predominantly subtype B viruses.8–10 However, the NRTI backbone needs to be carefully chosen as pretreatment NRTI resistance in ART-naîve/prior-ART-exposed individuals may compromise the successful implementation of dolutegravir-based first-line therapy.11 Moreover, if its implementation is based on a public health approach, some patients already on efavirenz-based first-line ART may be switched to the new recommended first-line therapy, without prior monitoring of virological suppression status. This raises the very real possibility that patients with virological failure and extensive NRTI resistance mutations1 could be switched to dolutegravir-based ART.

So far, only the FDCs of abacavir/lamivudine/dolutegravir,12 and since April 2018 tenofovir/lamivudine/dolutegravir,13 have been manufactured and are available. In 2009, Sax et al.13 observed a shorter time to virological failure and to first adverse event in people starting ART with an abacavir/lamivudine backbone, compared with tenofovir/emtricitabine, in combination with either efavirenz or atazanavir. The FDC of tenofovir/emtricitabine/efavirenz has been available since 2010 and recommended as first-line therapy. Wide-scale use of the same NRTI backbone for previous NNRTI and future INSTI-based regimens has raised concerns regarding activity in those patients with PDR or ongoing viral failure at the time of switch from efavirenz to dolutegravir.

We investigated drug resistance to NRTIs, NNRTIs and INSTIs in the ANRS 12249 treatment-as-prevention (TasP) trial in ART initiators/initials, as well as in people failing their first-line ART enrolled in the trial, in order to understand whether abacavir- or tenofovir-based backbones would be more active.

Methods

Study design and setting

The ANRS 12249 TasP trial was a cluster-randomized trial implemented in 22 clusters (2×11) in the Hlabisa sub-district in rural KwaZulu-Natal, South Africa, from March 2012 to June 2016. There were 1000 individuals ≥16 years residing in each cluster with an HIV prevalence of ~30%. The full trial protocol has been described previously; participants residing in the intervention clusters were offered ART after HIV diagnosis, regardless of their CD4 counts, whereas participants in control clusters were offered ART according to the prevailing South African guidelines (i.e. CD4 count <350 cells/mm³ in March 2012, then <500 cells/mm³ from January 2015).14

Study population

Participants attending the trial clinics were either ART naive at entry into the trial or already established on ART prescribed through the Hlabisa HIV treatment and care programme. All individuals were asked to complete study questionnaires and provide plasma samples at their first trial clinic visit. Plasma samples were used for viral load (VL) testing, using the Abbott RealTime HIV-1 m2000rt (Abbott Molecular Inc., Des Plaines, IL, USA), as well as for Sanger drug resistance testing in the Africa Health Research Institute diagnostic laboratory when clinically indicated.

Next-generation sequencing (NGS)

NGS was used to characterize HIV pol from participants’ plasma samples with a VL ≥1000 copies/mL, adapting a protocol that was previously described by Gall et al.15 Briefly, RNAs were extracted from 1 mL of plasma, using the QIAamp Viral RNA Mini Kit (Qiagen, Hilden, Germany) and were eluted in 60 μL of elution buffer. The near-full HIV genome was amplified with four subtype-C–specific primer pairs, generating four overlapping amplicons of 2.1, 2.3, 2.2 and 3.9 kb. DNA concentrations of amplicons were quantified with the Qubit dsDNA HS Assay kit (Invitrogen, Carlsbad, CA, USA). Diluted at 0.2 ng/μL each, amplicons were pooled equimolarly and prepared for the library using the Nextera XT DNA Library preparation and the Nextera XT DNA sample preparation index kits (Illumina, San Diego, CA, USA), following the manufacturer’s protocol. The runs comprised a total of 96 samples, including three controls: 1 negative sample, 1 inter-run sample and 1 intra-run sample. If amplification or sequencing failed for the pol region of the HIV genome, samples were re-amplified with primers that partially covered the pol gene, excluding the integrase, and were sequenced in a 386-sample run, with the same controls as the 96-sample run. The intra-run and inter-run controls were used to assess the reproducibility and accuracy of our method (n=29). The mean identity of the consensus sequences derived for each duplicate was 99.81% (SD=0.35%). The mean difference between the SNP frequencies, detected from 0.2% to 100%, was 1.4% (SD=2.5%); therefore, we set with confidence our detection level of minority variants at 5%.

Read assembly was performed using Geneious 10.0.6 software;16 briefly, reads between 100 and 300 bp were selected and those with Phred scores <30 were excluded. The sequences were trimmed up to 10 bp from 5’ and 30 bp at the 3’ end and mapped against a subtype C reference sequence (AF411967) annotated for WHO surveillance of drug resistance mutations (DRMs).17 Minority variants between 5% and 20% were included when they were also detected by a BaseSpace application, MiCall.18 DRMs in the reverse transcriptase and integrase were detected at 20% (DRM20%) and 5% (DRM50%) levels of detection.

DRM penalty scores and resistance interpretation were estimated using the Stanford HIV Drug Resistance database (https://hivdb.stanford.edu/) for the following drugs: abacavir, cytosine analogues (lamivudine and emtricitabine), zidovudine, tenofovir, efavirenz/nevirapine and dolutegravir. The mutation scores were classified as follows: 0–9, susceptible; 10–29, low-level resistant; and ≥30, resistant. According to the REGA algorithm V10.0.0, the genotypic susceptibility score (GSS) was calculated for the entire abacavir/lamivudine/dolutegravir or tenofovir/lamivudine/dolutegravir regimen; a GSS score ≥3 and ≥2 represents full susceptibility to the regimen amongst ART initiators and ART-exposed persons, respectively.19

Ethics

The trial was approved by the Biomedical Research Ethics Committee (BFC 104/11) at the University of KwaZulu-Natal and the Medicines Control Council of South Africa (ClinicalTrials.gov: NCT01509508; South African National Clinical Trials Register: DOH-27-0512-3974). All trial participants gave written consent or witnessed thumbprint informed consent prior to undertaking any study procedures.

Results

Drug resistance cohort description

Of the 1287 participants with available NGS data for the reverse transcriptase gene, 1193 were classified as ART initiators (their ART status was naive (n=1054), their ART status was unknown (n=106) or they were previously exposed to a prevention of mother-to-child HIV transmission (PMTCT) regimen (n=33)), while 94 participants were currently exposed to ART (Table 1). Integrate NGS data were available for 524 ART initiators and 83 ART-exposed participants.
Among participants already on ART at trial entry, all were on an NNRTI-based regimen; 23 were on a TA backbone (zidovudine or stavudine) for a median time of 57.3 months (IQR 38.7–85.8), while 71 were on a tenofovir backbone for a median time of 22.4 months (IQR 11.6–37.2). Among people on a tenofovir-based regimen, 19 were previously on a stavudine-based regimen, and were found among nine participants who were previously or currently treated with TAs, with the exception of two participants who were on a tenofovir- and stavudine-based regimen, respectively. TAMs represented 14.4% of the DRMs (20/141; 14.2%), while the TAM-1 pathway (M41L and L210W) was represented by the TAM-2 mutations in addition to two mutations from the TAM-1 pathway (M41L and L210W). Notably, two additional TAMs were detected in one participant (n = 2), including Q148R and G163R.

**Genotypic antiretroviral susceptibility**

When resistance was detected at a 20% variant level, all ART initiators with available consensus sequence covering the entire integrase (n = 524) were fully susceptible to dolutegravir. Only 5 participants out of 1193 (0.4%) had low- to high-level NRTI resistance, while 115 had resistance to NNRTIs (9.6%) (Figure 2). At a variant level of 5%, more participants had RTI resistance detected among ART initiators; 9/1193 participants had NRTI resistance assessed, double the number of NRTI mutations were found (n = 22), mostly represented by K65R (n = 4/22, 18.2%) and the TAMs D67N and K219EQ (Figure 1b). Nine additional integrase DRM5% were found, mostly L74M (n = 5).

Among ART-exposed participants, a total of 217 DRM20% were detected and were equally distributed between the NRTI (104/217, 47.9%) and the NNRTI (113/217, 52.1%) classes. Fifteen participants had integrase DRM20% (15/83, 18.1%), including Q148R conferring low-level resistance to dolutegravir, and some accessory integrate mutations were also found: L74I/M (n = 114, 80.0%), E157Q (n = 35, 27.3%). The K65R mutation was the most prevalent (n = 35, 34.6%), followed by K65Q (n = 20, 19.2%) (Figure 1c). K65R was detected in 18/72 (25.0%) and 2/17 (11.8%) participants who were on a tenofovir- and stavudine-based regimen, respectively. TAMs represented 14.4% of the DRMs (n = 15/104); they belonged exclusively to the TAM-2 pathway and were found among nine participants who were previously or currently treated with TAs, with the exception of two participants who initiated ART with a tenofovir-based regimen.

Twenty-two more NRTI DRM5% were detected, mostly represented by the TAM-2 mutations in addition to two mutations from the TAM-1 pathway (M41L and L210W). Notably, two additional K65R mutations were detected at the 5% as compared with the 20% variant threshold (Figure 1d). Three additional participants had integrase DRM5%.
While 145/1193 (12.2%) had resistance to NNRTIs (Figure 2). Whether DRM20% or DRM5% were considered, 99.2% of ART initiators had full susceptibility (GSS to both dolutegravir-based regimens. There was no significant difference between the different ART-initiator sub-groups (ART naive, ART status unknown and previously exposed to a PMTCT regimen).

Among ART-exposed participants, only one had low-level resistance to dolutegravir (Q148R) among participants with available consensus sequence covering the protease (n = 83). This individual had no NRTI mutations. Regarding resistance to NRTI20%, 57/94 (60.6%) were fully susceptible to abacavir and cytosine analogues, while 67/94 (71.3%) were fully susceptible to tenofovir (P = 0.16; Figure 2). Accounting for DRM5%, 66.0% and 69.1% of participants were predicted to be fully susceptible (GSS to abacavir- and tenofovir-based dolutegravir regimens, respectively.

Full susceptibility to zidovudine was found among 92/94 (97.9%). Susceptibility to NNRTIs was compromised for 54/94

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**Figure 1.** DRMs among ART initiators (a and b) and ART-exposed participants (c and d), at 20% (a and c) and 5% (b and d) levels of detection.
(57.4%). When resistance was detected at 5%, full susceptibility to abacavir and cytosine analogues was found for 56/94 (59.6%), while 64/94 (68.1%) were fully susceptible to tenofovir ($P = 0.29$). Full susceptibility to zidovudine was found among 88/94 (93.6%). Susceptibility to NNRTIs was compromised for 59/94 (62.8%).

**Discussion**

In countries hosting large ART programmes with limited treatment options, it is important to evaluate whether the large-scale implementation of a new regimen will be effective. Following recent WHO guidelines, the South Africa will soon roll out a dolutegravir-based regimen as first-line ART. Our study aimed to predict genotypic susceptibility to this new regimen by analysing drug resistance over a large population with diverse ART exposure profiles. We found that a tenofovir + emtricitabine/lamivudine NRTI backbone had similar predicted efficacy to one based on abacavir in place of tenofovir.

We did not find any major mutations associated with high-level dolutegravir resistance irrespective of whether individuals were ART initiators or ART exposed. We did observe Q148R, which is associated with high-level dolutegravir resistance when accompanied by the G140S/A mutations. Indeed, INSTIs are used only for ART initiators and ART exposed. Our data suggest that NGS may not provide greater detection of drug resistance over Sanger sequencing under current conditions in ART-naive individuals.

Among ART initiators, including patients who were ART naive, previously exposed to a PMTCT regimen or for whom the ART status could not be clarified, nearly all were fully susceptible to both abacavir and tenofovir, with NRTI resistance found in just under 1% of the population, at both 20% and 5% levels of detection. In this specific population, both abacavir and tenofovir would be fully effective and the level of NNRTI resistance found in our population (>10%) confirmed the need to move towards INSTI-based regimens. Similarly, dolutegravir regimens with either abacavir or tenofovir were predicted to be fully active in 99% of ART-naive participants. Our data suggest that NGS may not provide greater detection of drug resistance over Sanger sequencing under current conditions in ART-naive individuals.

Among patients on ART, or those previously exposed to ART, full susceptibility to abacavir and cytosine analogues was found for 56/94 (59.6%), while 64/94 (68.1%) were fully susceptible to tenofovir using the 5% threshold. When accounting for DRM5%, full susceptibility to both dolutegravir regimens was not different and below 70%. Significant additional mutations, including K65R, were observed at the 5% level, but not the 20% level, suggesting NGS might be useful in treatment-experienced patients.

The higher, but non-statistically significant, level of resistance to abacavir compared with tenofovir in the context of use of an FDC containing tenofovir/emtricitabine/efavirenz in this population might be explained by the high prevalence of the M184V mutation, selected early in regimen failure by cytosine analogues, which can also confer partial resistance to abacavir. Moreover, the detection of TAMs, likely resulting from prior use of TAs, can enhance the resistance to abacavir in the presence of M184V whilst susceptibility to tenofovir might be increased. We found a low level of resistance to zidovudine among both ART initiators and
ART-exposed participants despite the detection of TAMs; this is explained by the fact that most of the TAMs were associated with M184V and/or K65R, mutations known to increase susceptibility to zidovudine.39 Therefore, these findings confirm the appropriate use of zidovudine for second-line PI-based therapy, after first-line virological failure. Finally, while the tenofovir-selected mutation K65R confers significant resistance to tenofovir and abacavir,40 its combination with M184V actually increases the level of resistance to abacavir, but decreases resistance to tenofovir.34

The preferable choice of using tenofovir as a backbone can be guided by clinical benefits: tenofovir has potent anti-HBV activity,41 and given the lack of testing for hepatitis B in most LMICs, where HIV and HBV prevalence is high, tenofovir is the preferred option, over abacavir. Secondly, HLA testing for HLA B*5701 is recommended where abacavir use is being considered due to the possibility of hypersensitivity reactions with this allele.42 Although one study found abacavir to be safe in African children regardless of HLA status,43 no such study has been conducted in adults. Abacavir use is preferred over tenofovir in children due to bone mineral density considerations,44 and therefore based on our data abacavir would be a suitable alternative to tenofovir if similar patterns of HIV drug resistance are found in ART-treated children about to initiate dolutegravir-based ART. In adults, tenofovir is contraindicated in moderate to severe kidney disease and in patients with high fracture risk. In an ageing population these conditions are more likely to occur. Although abacavir has been associated with increased risk of cardiovascular disease in retrospective cohorts,45 it is unclear whether this is relevant in black African populations.

This study was limited by a modest number of treated patients relative to naive patients. As expected the vast majority of naive participants were susceptible to both NRTIs and INSTIs. We used predicted activities based on genotypes and rule-based algorithms. Recent data from second-line studies have called into question the utility of such algorithms given better responses to second-line ART in those with more NRTI resistance.46 However, in the setting of second-line NRTI resistance is likely a surrogate of adherence and a boosted PI is likely sufficient to suppress virus in the majority of those with high adherence. Therefore, virological outcome studies are clearly needed to answer the question of impact of resistance to components of dolutegravir-containing ART.

We did not explore integrase resistance outside the integrase gene. Finally, we only genotyped those treated individuals with VL >1000 copies/mL. Although the WHO has defined virological failure as two consecutive VL counts of >1000 copies/mL, it is important to investigate drug resistance and outcomes among patients on ART with VL between 50 and 1000 copies/mL, as these individuals will also likely be switched to new dolutegravir first-line treatment without drug resistance testing.

In conclusion, our data suggest that in LMICs tenofovir and abacavir are predicted to have comparable effectiveness in combination with dolutegravir in treatment-naïve individuals.

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References


