Doravirine and Its Potential in the Treatment of HIV: An Evidence-Based Review of the Emerging Data

Alexander E Rock 1
Jeremy Lerner 2
Melissa E Badowski 1

1University of Illinois, College of Pharmacy, Department of Pharmacy Practice, Section of Infectious Diseases Pharmacotherapy, Chicago, IL 60612, USA; 2University of Illinois, College of Pharmacy, Department of Pharmacy Practice, Chicago, IL 60612, USA

Abstract: The utility of doravirine in the management of HIV-1 infection is approved for use in patients who are antiretroviral-naïve as well as patients who have achieved stable virologic suppression and are interested in replacing their current antiretroviral therapy. The role of doravirine continues to evolve as data emerges on the potential for new combination therapy with the investigational agent, islatravir, as well as a potential strategy to minimize post-marketing safety concerns with recommended first-line agents, such as integrase inhibitors. The goal of this review is to assess recent and emerging data on the non-nucleoside reverse transcriptase inhibitor, doravirine.

Keywords: doravirine, HIV-1, antiretroviral therapy, non-nucleoside reverse transcriptase inhibitor

Introduction

Doravirine, the most recent non-nucleoside reverse transcriptase inhibitor (NNRTI), is available as a co-formulated single-tablet regimen (STR) including lamivudine and tenofovir disoproxil fumarate (Delstrigo™) and independently distributed as Pifeltro™. Doravirine is approved as initial therapy as well as for those virologically suppressed on a stable antiretroviral (ARV) regimen. 1 This addition provides much needed variety to the NNRTI class with a daily medication that has less frequent side effects compared to other NNRTIs, maintains activity in the presence of common NNRTI resistance mutations (ie K103N, E138K, Y181C, etc.) and potentially provides a therapeutic alternative for individuals experiencing adverse events from other medication classes (ie integrase strand transfer inhibitors (INSTIs), protease inhibitors (PIs), etc.). Currently, the European AIDS Clinical Society (EACS) recommends the use of doravirine/lamivudine/tenofovir disoproxil fumarate STR for initial therapy in patients who are ARV-naïve whereas the United States Department of Health and Human Services (DHHS) reserves this combination for certain clinical situations. 2,3

With emerging data suggesting possible links to weight gain with both tenofovir alafenamide and the INSTI class there is a need for highly effective antiretroviral therapy (ART) with benign side effect profiles. 4–6 To date, there is minimal data addressing weight gain with doravirine. Recent cumulative data from three clinical trials (Phase 2, P007; Phase 3, DRIVE-FORWARD and DRIVE-AHEAD) concluded there may be slightly more weight gain associated
with doravirine than efavirenz but no significant weight gain when compared to boosted darunavir regimens in treatment-naïve people living with HIV (PLWH). The co-formulation of doravirine with lamivudine and tenofovir disoproxil fumarate may become more appealing because this STR may be an ideal alternative for patients suffering from ongoing issues related to weight gain or hypertriglyceridemia. One potential concern with the co-formulation of tenofovir disoproxil fumarate focuses on renal and bone health. Tenofovir disoproxil fumarate has a significant history associated with increased rates of renal injury, and decreased bone mineral density. Although these events are serious, fortunately their incidence is rare.

When highly treatment-experienced patients are unable to achieve virologic suppression it is not an uncommon practice to manage someone an INSTI with a boosted PI with or without a NRTI backbone, based on resistance patterns and tolerability. In circumstances where this boosted PI needs to be avoided, such as patients who are awaiting transplantation or individuals with drug interactions precluding the use of a boosted PI, doravirine becomes an attractive option. Although this occurs in clinical practice there has yet to be a randomized clinical trial performed to validate this approach.

Another potential role of doravirine is being evaluated in combination with lamivudine and islatravir, a nucleoside reverse transcriptase translocation inhibitor (NRTTI). With renewed interest in doravirine, the goal of this manuscript is to review current and emerging data on the role of doravirine in the management of HIV-1 infection.

Methodology
A literature search current through 27 March 2020 was conducted using Embase (n=162), Medline via PubMed (n=89), and Medline via Ovid (n=77) with the search terms utilized being doravirine OR mk-1439. The same terms were used to review abstract results from the Conference on Retroviruses and Opportunistic Infections (CROI) (n=13), International AIDS Society Conference on HIV Science (IAS) (n=6), European AIDS Clinical Society (EACS) (n=7), International Workshop of Clinical Pharmacology on HIV and Hepatitis Therapy (n=5), and IDWeek (n=6). Searches of these databases were limited to the English language and adult patients but were not limited beyond these criteria. The above results were compared for duplicates and resulted in n=119 unique published papers and n=37 conference abstracts (Figure 1).

Results
The greatest amount of data existing on doravirine is derived from the DRIVE trials along with Phase II dose finding trials (Table 1). These trials outline efficacy and safety data for doravirine and are the basis for its approval by the US Food and Drug Administration. Long-term efficacy data is available up to 96 weeks in the DRIVE-FORWARD trials and DRIVE-AHEAD trials. These studies assessed treatment-naïve patients as well as...
### Table 1  Doravirine Clinical Trials

| Title | Study Agents | Study Design | Outcomes (VL <50 Copies/mL) | Emergent Resistance | Adverse Events (AE) (%) |
|-------|--------------|--------------|------------------------------|---------------------|------------------------|
| **Treatment-Naïve** | | | **Week 96** | **PDVF** | **Week 96** |
| DRIVE-FORWARD Week 96 (2020) Week 48 (2018) | ● DOR (100 mg daily) + 2 NRTIs (n=383) ● DRV/r (800 mg/100 mg daily) + 2 NRTIs (n=383) | ● Phase 3 randomised controlled, double blind, non-inferiority study performed at 125 clinical centres in 15 countries ● December 1, 2014 – October 20, 2015 ● Age ≥ 18 years ● HIV-1 RNA ≥ 1000 copies/mL ● No known resistance | W96 DOR: 73% (277/383) DRV/r: 66% (248/383) Non-inferiority demonstrated | PDVF DOR: 9% (n=34) DRV/r: 11% (n=43) | DOR: 17 Diarrhea Nausea Headache Discontinuation due to AE |
| | | | | n=1 (V106I, H221Y, F227C, M184V) | W96: 17 W48: 5 |
| | | | | n=1 (V106A, P225T/H, V118I, M184I) | W96: 12 W48: 7 |
| | | | | DRV/r: 11% (n=43) | W96: 15 W48: 6 |
| | | | | n=1 phenotypic resistance to FTC | Discontinuation due to AE |
| | | | | | W48: 2 (n=9) W48: 2 (n=6) |
| Gatell JM (2019) | ● DOR (100 mg) + TDF/FTC ● EFV (600 mg) + TDF/FTC | ● Phase Ib randomized, open-label, double-blind, dose-finding, multi-center study performed in 11 countries | W48 DOR: 84% (312/383) DRV/r: 80% (306/383) Non-inferiority demonstrated | PDVF DOR: 5% (n=19) DRV/r: 6% (n=24) | W96: 2 (n=9) W48: 2 (n=6) |
| | | | | | W96: 4 (n=14) W48: 3 (n=12) |
| | | | | | CNS AEs Discontinuation due to AE |
| | | | | | W24: 26.9 W 24: 47.2* |
| | | | | | W48: 2.8 5.6 |

(Continued)
| Title            | Study Agents                  | Study Design                                                                 | Outcomes (VL <50 Copies/mL)                                                                 | Emergent Resistance | Adverse Events (AE) (%) |
|------------------|-------------------------------|------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------|---------------------|-------------------------|
| DRIVE-AHEAD Week 96 (2019) | ● DOR/3TC/TDF (n=364)         | ● Phase 3, open-label, parallel-group, randomized, non-inferiority switch study performed at 126 centers  | W96: DOR 77.5% (282/364) EFV 73.6% (268/364) Non-inferiority demonstrated              | PDVF                | DOR                     |
|                  | ● EFV/FTC/TDF (n=364)         | ● Age ≥ 18 years                                                             |                                                                                           |                     | EFV                     |
|                  |                               | ● HIV-1 RNA ≥ 1000 copies/mL                                                  |                                                                                           |                     | Dizziness               |
|                  |                               | ● No known resistance to study drugs                                          |                                                                                           |                     | W96: 10.2               |
|                  |                               | ● CrCL ≥ 50 mL/min                                                           |                                                                                           |                     | W48: 8.8                |
|                  |                               | W96: 77.5% (282/364) EFV 73.6% (268/364) Non-inferiority demonstrated        |                                                                                           | PDVF: DOR 6% (n=22) | W96: 14                 |
|                  |                               | W48: 84.3% (307/364) EFV 80.8% (294/364) Non-inferiority demonstrated        |                                                                                           |                     | W48: 12.1               |
|                  |                               | W96: 77.5% (282/364) EFV 73.6% (268/364) Non-inferiority demonstrated        |                                                                                           |                     | 25.2%                   |
|                  |                               | W48: 84.3% (307/364) EFV 80.8% (294/364) Non-inferiority demonstrated        |                                                                                           | PDVF: 3.8% (n=14)  | W96: 8.5                |
|                  |                               | W48: 84.3% (307/364) EFV 80.8% (294/364) Non-inferiority demonstrated        |                                                                                           |                     | W48: 8.2                |
|                  |                               | W96: 100% (8/8) W48: 100% (8/8)                                              |                                                                                           |                     | W96: 3 (NR)             |
|                  |                               | *Targeted enrollment not achieved                                            |                                                                                           |                     | W48: 3 (n=11)           |
|                  |                               | *2 subjects not included in final analysis                                   |                                                                                           |                     | W48: 7 (n=24)           |
|                  |                               | ● GI disorders (40%)                                                         |                                                                                           |                     |                         |
|                  |                               | ● Fatigue (20%)                                                              |                                                                                           |                     |                         |
|                  |                               | ● Nervous system disorders (20%)                                             |                                                                                           |                     |                         |
|                  |                               | ● Insomnia (10%)                                                            |                                                                                           |                     |                         |
| DRIVE-BEYOND     | ● DOR/3TC/TDF (n=10)          | ● Phase 2, multicenter, open-label, single arm study                        | W96: 100% (8/8) W48: 100% (8/8)                                                           |                     |                         |
|                  | ● K103N (n=8)                 | ● Age ≥ 18 years                                                             |                                                                                           |                     |                         |
|                  | ● G190A (n=2)                 | ● HIV-1 RNA ≥ 1000 copies/mL                                                  |                                                                                           |                     |                         |
|                  |                               | ● CD4 count ≥ 100 cells/mm³                                                   |                                                                                           |                     |                         |
|                  |                               | ● Single NNRTI mutation (K103N, Y181C, or G190A)                              |                                                                                           |                     |                         |
|                  |                               | ● No known resistance to study drugs                                          |                                                                                           |                     |                         |
|                  |                               | W96: 100% (8/8) W48: 100% (8/8)                                              |                                                                                           |                     |                         |
|                  |                               | *Targeted enrollment not achieved                                            |                                                                                           |                     |                         |
|                  |                               | *2 subjects not included in final analysis                                   |                                                                                           |                     |                         |
|                  |                               | ● n=1 with G190A PDVF                                                        |                                                                                           |                     |                         |
|                  |                               | ● n=1 with K103N LTFU                                                        |                                                                                           |                     |                         |
|                  |                               | PDVF: 0/8 in final analysis                                                  |                                                                                           |                     |                         |
### Treatment-Experienced

| DRIVE-SHIFT  |  |  |  |  |  |
|--------------|---|---|---|---|---|
| ● Immediate switch group (ISG) to DOR/3TC/TDF (n=447) | ● Phase 3, open-label, randomized, active-controlled, non-inferiority study performed at 122 sites in Europe, North America, Latin America, and Asia | ● Virologically-suppressed ≥ 6 months on 2 NRTIs + boosted PI, EVG, or NNRTI | ● June 17, 2015 – February 10, 2017 | ● Age ≥ 18 years | ● No history of virologic failure |
| ● Delayed switch group (DSG) (continued for first 24 weeks) to DOR/3TC/TDF (n=223) |  |  |  |  |  |

***W48***:
- ISG: 90.8% (406/447)
- DSG: NA W24
- ISG: 93.7% (419/447)
- DSG: 94.6% (211/223)

Non-inferiority demonstrated

### Investigational Agents

| Molina JM (2019) |  |  |  |  |
|------------------|---|---|---|---|
| ● ISL 0.25mg, 0.75mg, or 2.25mg + DOR + 3TC | ● Phase 2, randomized, double-blind, dose-ranging trial |  |  |  |
| ● DOR/3TC/TDF | ● Adults |  |  |  |
|  | ● Treatment-naive |  |  |  |
|  | ● HIV-1 RNA ≥ 1000 copies/mL |  |  |  |
|  | ● CD4 count ≥ 200 cells/mm³ |  |  |  |
|  | ● No ARV resistance |  |  |  |
|  | ● No active HCV or HBV coinfection |  |  |  |

***W48***:
- ISL 0.25 mg: 2.9% (26/28)
- ISL 0.75 mg: 9.1% (27/29)
- ISL 2.25 mg: 85.7% (24/28)
- DOR/3TC/TDF: 92.9% (26/28)

### Note

Statistical significance (*p*<0.05) indicated by (*).

**Abbreviations:** 3TC, lamivudine; CNS, central nervous system; CrCl, creatinine clearance; DOR, doravirine; DRV/r, ritonavir boosted darunavir; EFV, efavirenz; EVG, elvitegravir; FTC, emtricitabine; HBV, hepatitis B virus; HCV, hepatitis C virus; ISL, islatravir; LTFU, lost to follow-up; NR, not reported; NNRTI, non-nucleoside reverse transcriptase inhibitor; NRTI, nucleoside reverse transcriptase inhibitor; PDVF, protocol defined virologic failure; PI, protease inhibitor; TDF, tenofovir disoproxil fumarate; VL, viral load; W, week.
those switching from stable ART. The DRIVE trials in treatment-naive only directly compare doravirine to efavirenz/emtricitabine/tenofovir disoproxil fumarate and boosted darunavir. The DRIVE-Shift trial did include patients that switched to a doravirine regimen from either a boosted PI, elvitegravir, or NNRTI-based regimen, however, it did not stratify the results based on these characteristics. Across all of the DRIVE studies there was a consistent trend of doravirine being better tolerated than its comparator, the only exception being the DRIVE-FORWARD trial where the incidence of headache was higher at both 48 and 96 weeks for the doravirine arm compared to boosted darunavir.

DRIVE-FORWARD was a phase 3, double blind, non-inferiority trial, comparing treatment-naive patients whom were started in a 1:1 ratio on either doravirine or darunavir boosted with ritonavir with investigator selected NRTI backbone therapy. The mean effect on low-density lipoprotein (LDL-C), non-high density lipoprotein (HDL-C), HDL, total cholesterol (TC), and triglycerides (TG) at 96 weeks was (−0.4 mg/dL, −0.5 mg/dL, 4.5 mg/dL, 4.1 mg/dL, and −1.1 mg/dL) and (14.0 mg/dL, 17.7 mg/dL, 4.2 mg/dL, 21.9 mg/dL, and 22.5 mg/dL), for doravirine and darunavir boosted with ritonavir regimens, respectively. A significant difference was observed with a 95% confidence interval (CI) for all parameters of the boosted darunavir arm as well as HDL and cholesterol in the doravirine arm.

DRIVE-AHEAD was a phase 3, double blind, non-inferiority trial comparing treatment-naive patients whom were started in a 1:1 ratio on either doravirine/lamivudine/tenofovir disoproxil fumarate or efavirenz/emtricitabine/tenofovir disoproxil fumarate. At 0, 48, and 96 weeks, measured lipids were associated with a positive effect in the doravirine group. The mean effect on LDL and non-HDL at 96 weeks was (−0.62 mg/dL and −2.14 mg/dL) and (10.78 mg/dL and 14.95 mg/dL) for the doravirine and efavirenz arm, respectively. Fasting LDL-C and non-HDL-C increased in the efavirenz group but not in the doravirine group, yet the change in total cholesterol/HDL-C ratio was similar.

DRIVE-SHIFT was a switch study assessing two groups (baseline regimen vs doravirine/lamivudine/tenofovir disoproxil fumarate) before and after a switch where all patients ultimately received doravirine/lamivudine/tenofovir disoproxil fumarate. No weight gain or abnormal laboratory results related to changes in lipid profiles led to discontinuation of therapy. This study demonstrated superior lipid profiles for doravirine/lamivudine/tenofovir disoproxil fumarate as compared with boosted PI therapy. Statistically significant differences in both LDL and non-HDL at week 24 (p < 0.0001) were observed. However, the population effect of this may be limited, as only 1.6% of patients (n=3/184) in the baseline regimen group had a grade 3 laboratory abnormality value or greater (defined as LDL ≥190 mg/dL) at the end of the 24-week pre-switch time period. These three individuals did return to <190 mg/dL after 24 weeks on doravirine/lamivudine/tenofovir disoproxil fumarate.

Taken together, the DRIVE trials demonstrate that doravirine has proven efficacy over long periods of time while maintaining an acceptable adverse effect profile that may distinguish it from other regimens. One limitation noticed across all of the DRIVE trials is that the majority of participants were male, and there was a lack of data representing women who were pregnant.

**Emerging Literature and Discussion**

**Resistance**

Additional doravirine resistance data is beginning to emerge. One analysis evaluated 9764 sequences of ARV-naive samples in France, Greece, and Italy for the development of doravirine resistance-associated mutations (RAM). At least one doravirine RAM was identified in 1.4% (n = 137) of this population with most frequent mutations being V108I (n = 62; 0.6%), Y188L (n = 18; 0.2%), Y318F (n = 23; 0.2%), and H221Y (n = 10; 0.8%) compared to other NNRTIs (13.5%) and NRTIs (20.6%).

However, it is important to keep in mind that these tests were performed prior to doravirine being widely available and the mutations associated with doravirine resistance may not have been identified at the time. Additionally, this was only a genotypic assessment and does not demonstrate how all of these viruses would have phenotypically responded to doravirine. The same group went on to assess 9,199 sequences of which 381 belonged to treatment-experienced NNRTI failing patients. This subset, according to the Stanford algorithm, demonstrated a 42% (n=160) prevalence of doravirine resistance.

An algorithm to predict transmitted drug resistance (TDR) in treatment-naive patients in Spain demonstrated infrequent TDR associated with doravirine (1.8%) and PIs (0.8%) compared to other NNRTIs (13.5%) and NRTIs (3.0%). Unfortunately, TDR to INSTIs was not assessed in this analysis. Another study evaluated 6,893 NNRTI-experienced patients in Italy and identified intermediate
doravirine resistance (ie V106A) in 12.7% and high-level
doravirine resistance in 6.1% with the most common
mutation being Y188L. Previous treatment with efavirenz
or etravirine was associated with high-level doravirine
resistance while rifampicin was associated with a lower
probability of high-level doravirine resistance. A study
conducted by Ayitewala et al, assessed Ugandan patients
between 2006 and 2017 who failed therapy after a year of
efavirenz-based treatment and found 78.5% (n=143) of
these patients demonstrated high levels of resistance to
efavirenz. When this group was analyzed for cross-
resistance to the other NNRTIs, only 25.2% (n=36) were
susceptible to etravirine and rifampicin and 15.4% (n=22)
were susceptible to doravirine.

Pharmacokinetic/Pharmacodynamic
An excellent review performed by Boyle and colleagues
addresses the general pharmacodynamic, pharmacokinetic,
and drug interaction profile of doravirine. Since its pub-
lication in August 2019, a phase one study evaluated oral
granules as an alternative formulation for the single tablet
of combination doravirine/tenofovir disoproxil fumarate/
lamivudine. Equivalence of each component administered
was evaluated as granules as compared to their tablet
equivalent when taken alone and in combination with
either apple sauce or vanilla pudding. The absorption of
this formulation was determined to be equivalent to oral
tables. Although this formulation is not currently avail-
able, it would be an important option for patients unable to
tolerate or take oral tablets.

Another study assessed serum doravirine concentra-
tions after 72 hours of treatment cessation. This study,
importantly, demonstrated that at 72 hours after a single
dose of doravirine, 100% of the participants had an aver-
age trough concentration of 39 ng/mL (range: 33–61 ng/
ML) which was above the goal of 23 ng/mL set during
drug development. This is significant because it demon-
strates the forgiveness that standard doravirine dosing
would have in response to occasional non-compliance.

Investigational Formulations
Another emerging role for the use of doravirine is being
evaluated with the use of the NRTTI, islatravir. In a dose
finding study (Table 1), various doses of islatravir were
paired with lamivudine and doravirine. Participants who
achieved viral suppression (<50 copies/mL) at week 20 or
later stopped lamivudine at their next visit and continued
with doravirine and islatravir. No differences in weight
change or body mass index were observed from baseline
and were similar between the varying doses of islatravir
combined with doravirine and doravirine/lamivudine/teno-
fovir disoproxil fumarate. A significantly lower impact
was observed on hip bone mineral density (BMD) with
islatravir combined with doravirine when compared to
doravirine/lamivudine/tenofovir disoproxil fumarate yet
changes in spine BMD were similar between the
groups. The two-drug combination of doravirine
100 mg and islatravir 0.75 mg is being considered in
treatment-naive, virologically suppressed, and highly treat-
ment-experienced populations.

Clinical Implications for Doravirine
The role of doravirine continues to evolve in clinical
practice as new data and potential combinations emerge
along with concerns of post-marketing safety data asso-
ciated with other drug classes. Although doravirine is
approved as initial therapy in patients who are ARV-
naïve, it is currently recommended to obtain a genotype
prior to the initiation of ART. In patients who are treat-
ment-naïve and no TDR is identified, the STR of doravir-
ine/lamivudine/tenofovir disoproxil fumarate may be an
appropriate option since baseline resistance to doravirine
is rather low but baseline genotypes must also consider the
presence of NRTI mutations. The development of doravir-
ine RAMs while on therapy is another concern in the
setting of medication non-adherence along with previous
use of efavirenz and etravirine. Although doravirine may
have a higher genetic barrier to resistance compared to
first-generation NNRTIs, second-generation INSTIs
(dolutegravir and bictegravir) and PIs are more forgiving
in settings of medication non-adherence and are associated
with higher genetic barriers to resistance. Furthermore,
development of V106A/M or Y188L with the use of
doravirine can precipitate the development of cross-
resistance across the NNRTI class.

In patients experiencing side effects from other ARVs,
such as weight gain with INSTIs or with tenofovir alafer-
namide, the role of doravirine may emerge as a potential
solution to reduce weight gain as well as long-term meta-
bolic consequences. Although not directly compared to
INSTIs, data on the effect of weight gain with doravirine
use was only recently presented. While no difference in
weight gain from baseline was observed between doravir-
ine (W48 = 1 kg; W96 = 1.5 kg) and boosted darunavir
(W48 = 0.59 kg; W96 = 0.7 kg), a significant difference
was observed between doravirine and efavirenz (W48 =
0 kg; W96 = 1 kg) based on 48 (p<0.001) and 96 (p=0.020) week data. In addition, DRIVE-AHEAD and DRIVE-FORWARD demonstrated better lipid profiles when doravirine was compared to efavirenz or boosted-darunavir. Additionally, DRIVE-SHIFT also demonstrated better lipid outcomes associated with doravirine, data on baseline regimens (ie boosted darunavir or boosted elvitegravir) were not separated to assess the degree of difference. A main limitation in this study is the lack of a true comparison to recommended INSTIs, dolutegravir and bictegravir.

When evaluating doravirine for use in renal dysfunction, if administered as an individual agent, dosage adjustments are not required regardless of creatinine clearance. Yet, a major concern with co-formulating doravirine with lami-vudine and tenofovir disoproxil fumarate is renal and bone safety. Although bone and renal parameters were not specifically assessed in the DRIVE trials, 96-week data obtained from the DRIVE-FORWARD study was not associated with significant differences between Grade 2 or 3 adverse events when comparing doravirine or ritonavir-boosted darunavir with investigator selected NRTI backbone (tenofovir disoproxil fumarate/emtricitabine or abacavir/lamivudine). Unfortunately, data comparing backbone therapy was not described in order to more clearly evaluate changes in kidney function. The DRIVE-AHEAD study also did not specifically evaluate bone and renal parameters but did note fractures and drug-related adverse events occurred in less than 1% of the treatment group. Grade 3 adverse events related to changes in creatine were identified in 1.9% of the doravirine/lamivudine/tenofovir disoproxil fumarate group and 0.8% of the efavirenz/emtricitabine/tenofovir disoproxil fumarate group [95% CI 1.1 (−0.7, 3.2)] based on 48-week supplementary data. Updated 96-week data demonstrated a rise in serum creatinine by greater than 1.3–1.8 x Upper Limit of Normal (ULN) or increase of greater than 0.3 mg/dL above baseline (doravirine = 3%; efavirenz = 2%) or increase of ≥1.5 x above baseline (doravirine = 3%; efavirenz = 2%) was not significantly different between groups. Finally, of 670 patients enrolled in the DRIVE-SHIFT study, only 2 individuals experienced moderate (n=1) or serious (n=1) adverse renal events which resolved upon cessation of doravirine-based therapy. Post-marketing data should continue to evaluate the development of adverse renal effects in patients taking doravirine/lamivudine/tenofovir disoproxil fumarate.

One potential advantage of co-formulating doravirine with tenofovir disoproxil fumarate as opposed to tenofovir alafenamide may be the concern for alopecia which was recently described in a case series of 6 African American women. Five out of 6 women reported improvement of alopecia 1–5 months upon cessation of tenofovir alafenamide therapy. Although this cause–effect relationship is in a rather small number of individuals, additional studies should evaluate this potential adverse effect in greater detail.

Lack of food requirements make doravirine an attractive option when compared to rilpivirine, etravirine, efavirenz, PIs, and elvitegravir/cobicistat. In addition, while the DRIVE trials give a baseline level of efficacy, other evidence has shown that doravirine can be an effective NNRTI that has a place in modern HIV therapy. A common reason to switch between therapies has often been due to drug interactions. The lack of certain drug interactions with doravirine can potentially distinguish it from other therapies. For example, doravirine was not associated with any clinically significant interactions between pantoprazole or an aluminum-magnesium based antacid. Although the true number of patients living with HIV and requiring antacid therapy is not well defined, one study distributed a validated gastroesophageal reflux disease (GERD) questionnaire between two HIV clinics in Canada to 120 patients to assess the presence of GERD. Eighty-five patients were included in the analysis where 68 (81%) met criteria for diagnosis of GERD and 65 (95.6%) were taking medications, the majority being histamine 2 receptor antagonists (H2RA), to alleviate symptoms. While doravirine does not interact with H2RA or proton pump inhibitors, rilpivirine and atazanavir do have significant interactions.

Another common drug interaction to be concerned with is polyvalent cations and the widely used INSTIs. A retrospective analysis performed in Delaware, USA, identified 152 (42%) clinic patients as receiving a polyvalent cation and an INSTI. Forty-six patients (13%) went on to develop virologic failure with the use of either dolutegravir (15%), elvitegravir (11%), or raltegravir (18%), carrying an average 2.3 times risk for virologic failure when compared to those not reporting polyvalent cation use [95% CI 1.2–4.4]. These interaction profiles can potentially distinguish doravirine from other currently available agents.

An important drug interaction that warrants discussion is between doravirine and the rifamycin drug class. While the administration of rifampin in combination with doravirine is not recommended, as concentrations of doravirine
will be substantially decreased, rifabutin can be considered as long as doravirine is dosed twice daily.34,35

While the ideal candidate for doravirine STR must have a creatinine clearance of $\geq50$ mL/min, it can be administered irrespective of CD4 count or HIV-1 viral load. Doravirine, as a single agent, can be administered regardless of creatinine clearance and may be beneficial in patients who have susceptible virus to doravirine but documented resistance to other ARVs. Caution should be observed in patients with underlying NNRTI exposure and resistance since the durability of doravirine may be compromised. Both formulations of doravirine may be considered for patients experiencing side effects or drug interactions to other ARVs. While the role of doravirine continues to evolve, the focus of future studies should center on long-term safety and efficacy compared to current recommended regimens and standard of care as well as the potential combination with new agents being evaluated.

**Disclosure**

The authors report no conflicts of interest in this work.

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