

Department of Pharmacology and Experimental Department of Pharmacology and Experimental Therapeutics Faculty Papers Therapeutics

5-14-2020

Twice-Daily Doravirine Overcomes the Interaction Effect from Once-Weekly Rifapentine and Isoniazid in Healthy Volunteers.

Edwin Lam Thomas Jefferson University

Joseph Schaefer Thomas Jefferson University

Richard Zheng Thomas Jefferson University

Tingting Zhan Thomas Jefferson University

Walter K. Kraft *Thomas Jefferson University* Follow this and additional works at: https://jdc.jefferson.edu/petfp

Part of the Medical Pharmacology Commons
<u>Let us know how access to this document benefits you</u>

Recommended Citation

Lam, Edwin; Schaefer, Joseph; Zheng, Richard; Zhan, Tingting; and Kraft, Walter K., "Twice-Daily Doravirine Overcomes the Interaction Effect from Once-Weekly Rifapentine and Isoniazid in Healthy Volunteers." (2020). *Department of Pharmacology and Experimental Therapeutics Faculty Papers*. Paper 120. https://jdc.jefferson.edu/petfp/120

This Article is brought to you for free and open access by the Jefferson Digital Commons. The Jefferson Digital Commons is a service of Thomas Jefferson University's Center for Teaching and Learning (CTL). The Commons is a showcase for Jefferson books and journals, peer-reviewed scholarly publications, unique historical collections from the University archives, and teaching tools. The Jefferson Digital Commons allows researchers and interested readers anywhere in the world to learn about and keep up to date with Jefferson scholarship. This article has been accepted for inclusion in Department of Pharmacology and Experimental Therapeutics Faculty Papers by an authorized administrator of the Jefferson Digital Commons. For more information, please contact: JeffersonDigitalCommons@jefferson.edu.

1	Title: Twice-Daily Doravirine Overcomes the Interaction Effect from Once-Weekly Rifapentine and
2	Isoniazid in Healthy Volunteers
3	
4 5	Authors: Edwin Lam ¹ , Joseph Schaefer ² , Richard Zheng ¹ , Tingting Zhan ¹ , & Walter K. Kraft ¹
6	Affiliations: Department of Pharmacology & Experimental Therapeutics ¹ and Sidney Kimmel Medical
7	College ² , Thomas Jefferson University, Philadelphia, PA, USA
8	
9	Corresponding Author:
10	Edwin Lam, PharmD
11	Department of Pharmacology & Experimental Therapeutics
12	Thomas Jefferson University
13	132 South 10 th Street, 1170 Main Building
14	Tel: 215-955-9076
15 16	Email: Edwin.lam@jefferson.edu
17	Conflicts of Interest: All authors declared no competing interests for this work.
18	
19	Funding: This work was supported by an Investigator Studies Program grant provided by Merck & Co,
20	Inc. [MISP58495]. Edwin Lam is supported by the National Institutes of Health institutional training grant
21	[T32GM008562].
22	
23	Keywords: Human immunodeficiency virus; tuberculosis, non-nucleoside reverse transcriptase inhibitor;
24	ritamycin; drug interaction; doravirine
25 26	
20	
28	
29	
30	
31	
32	
33	
34	
35	
36	
37	
38	
39	
40 //1	
42	
43	
44	
45	
46	
47	

48 Abstract:

49 Doravirine (DOR) is a non-nucleoside reverse transcriptase inhibitor indicated for the treatment of human 50 immunodeficiency virus-1 (HIV-1). Its use in combination with rifapentine (RPT), an anti-tuberculosis 51 antibiotic, may reduce the exposure of DOR compromising viral suppression in those living with HIV-1 co-52 infected with tuberculosis. We conducted a prospective, phase I, open label, two-period, fixed sequence, 53 drug interaction study to evaluate the effect of once-weekly RPT and isoniazid (INH) on the pharmacokinetics of DOR in healthy volunteers. DOR 100 mg was administered alone twice-daily for 4 54 55 days in period 1. In period 2, once-weekly RPT+INH was co-administered with multiple doses of DOR 100 mg twice-daily for study days 7, 14, and 21. Plasma was obtained for DOR pharmacokinetics when given 56 57 alone and co-administered with RPT+INH. Eleven healthy volunteers enrolled and completed the study. The geometric mean ratios and 90% confidence intervals for DOR AUC₀₋₁₂ and C₁₂ in the presence of 58 59 RPT+INH compared to DOR alone were 0.71 (0.60-0.85) and 0.69 (0.54-0.89), respectively. Although 60 exposures were moderately reduced in the presence of RPT+INH, trough DOR values were within the 61 concentration range associated with virological suppression. These results demonstrate that a modest 62 decrease in doravirine exposure would unlikely be clinically relevant in a virally suppressed patient coadministered once-weekly RPT+INH. 63 64 65 66

68

- 69
- 70
- ,
- 71

72 Introduction

73 Doravirine is a non-nucleoside reverse transcriptase inhibitor (NNRTI) used in combination with 74 other antiretrovirals (ARV) for the treatment of HIV-1 infection in adults. It is non-inferior to current standard of care ARV regimens with fewer adverse events.^{1, 2} The elimination of doravirine is primarily 75 76 through metabolism via cytochrome P450 (CYP) 3A mediated oxidation to an M9 metabolite.³ In vitro 77 studies demonstrate low potential of interference from the parent or metabolite on phase I and II metabolizing enzymes and drug transporters.³ Therefore, doravirine has a lower potential for drug-drug 78 79 interactions compared to other NNRTIs and does not impact the pharmacokinetics of other drugs. 80 Considering the predominant route of elimination is through CYP3A, co-administration with strong 81 inhibitors and inducers may alter the pharmacokinetic profile of doravirine. 82 The Centers for Disease Control and Prevention (CDC) currently recommends several latent 83 tuberculosis infection (LTBI) treatment options for persons with HIV and includes either once-daily 84 isoniazid for nine months, rifampin or rifabutin once-daily for four months, or once-weekly rifapentine 85 and isoniazid for three months.⁴ The rifamycin class of anti-tuberculosis agents are strong inducers of 86 CYP enzymes, including CYP3A, with rifabutin being the less potent inducer of the three. Co-87 administration with multiple-doses of rifampin significantly reduced doravirine trough concentrations by 97% with multiple-dose rifabutin similarly reducing troughs by 68%.^{5, 6} With the exception of rifabutin, 88 89 where twice-daily doses of doravirine are predicted to overcome the interaction⁶, the use of rifampin 90 and rifabutin together with once-daily doses of doravirine is contraindicated. A three month 91 rifapentine based regimen has comparable efficacy and safety compared to lengthy once-daily anti-92 tuberculosis regimens.⁷ The magnitude of drug-drug interaction between doravirine when co-93 administered with rifapentine and isoniazid has not been defined.

94 The objective of this study was to evaluate the effects of once-weekly rifapentine and isoniazid
95 on the steady-state pharmacokinetics of twice-daily doravirine, and to assess the safety and tolerability
96 of these co-administered drugs.

97 Methods

98 The study was approved by the Thomas Jefferson University Institutional Review Board and 99 registered on clinicaltrials.gov (NCT03886701). The study was conducted in accordance to Good Clinical 100 Practice standards and applicable federal and/or local regulatory requirements. All participants provided 101 written informed consent prior to beginning the study.

102 Study population

103 Healthy HIV and TB negative adult men or non-pregnant women between 18-60 years old with a 104 body mass index of 19-33 kg/m² weighing between 45-120 kg at screening were enrolled in the study. 105 Women of childbearing potential and their partners were required to use acceptable methods of 106 contraception during the time of the study and until four weeks after the last dose of drug. Women who 107 were postpartum for less than 12 months were excluded. Other exclusion criteria included any clinically 108 significant disease, current drug or alcohol abuse, known anaphylactic or systemic reactions to doravirine, 109 rifapentine, or isoniazid, and those who have received another study drug within four weeks or five half-110 lives (whichever occurring first).

111 Study design and treatments

This was a phase I, prospective, open-label, two-period, fixed-sequence, drug-drug interaction study conducted in healthy volunteers. (**Figure 1**) Subjects received doravirine 100 mg twice-daily for four study days during the first period. In the second period, once-weekly weight based rifapentine and isoniazid and pyridoxine 50 mg was co-administered with doravirine 100 mg twice-daily. The doses of rifapentine were weight based and included 900 mg (≥50 kg) or 750 mg (32.1-49.9 kg) with a 900 mg maximum dose. Isoniazid was given as a 15 mg/kg dose and rounded up to the nearest 50 or 100 mg

118 with a maximum dose of 900 mg. The doses and dosing schedule for rifapentine, isoniazid, and

119 pyridoxine were selected based on the Centers for Disease Control and Prevention treatment regimens

120 for LTBI.⁴ Subjects received a light meal prior to each doravirine dose for study days 1-4. For the morning

of study days 7, 14, and 21, isoniazid and pyridoxine was dosed following an overnight fast with

doravirine and rifapentine dosed after the subject received a meal. For days 8-13 and 15-20, subjects

received a light meal before each dose of doravirine. All subjects received 8 ounces of water with each

124 dose.

125 Pharmacokinetic sampling and bioanalysis

126 Plasma samples for doravirine were collected at pre-dose (0 hour), 0.5, 1, 1.5, 2, 3, 6, 12, 24, 36, 127 48 and 72 hours post dose on study days 4-7 (period one) and study days 21-24 (period two). A pre-dose 128 sample was taken for all subjects prior to starting period 1 and doravirine dosing to ensure no previous 129 doses of doravirine were taken. A doravirine trough concentration was collected on study days 15, 16, 130 and 20 prior to the administration of the second doravirine dose. Approximately 3 mL of blood was 131 collected into K2-EDTA vials and inverted 8-10 times before being centrifuged at 3000 rpm for 10 132 minutes. Plasma was aliquoted into a cryotube and was stored at -20°C before analysis. Plasma 133 doravirine concentrations were determined using a validated ultra-performance liquid chromatography 134 tandem mass spectrometry with a lower limit of quantification of 1 ng/mL over a calibration range of 1-135 1000 ng/mL (developed and validated by Syneos Health Clinique, Quebec, QC, Canada). 136 Safety and tolerability 137 Safety and tolerability were assessed throughout the study and included monitoring for adverse events, 138 physical exams, vital signs, electrocardiograms, and laboratory safety tests (blood chemistry, 139 hematology, and urinalysis). Laboratory safety assessments were conducted in period 1 (study day 5) 140 and period 2 (study days 8, 15, and 22).

141 Pharmacokinetic and statistical analysis

143 analysis and performed on R version 3.6.1 with the PKNCA and ncappc packages. Doravirine 144 pharmacokinetic parameters included the area under the concentration-time curve during the dosing 145 interval (AUC₀₋₁₂), steady-state trough plasma concentration at the end of the dosing interval (C_{12}), 146 average steady-state plasma drug concentration during multiple dose administration (Cave), peak plasma 147 concentration at steady-state (C_{max}), terminal elimination half-life ($t_{1/2}$), apparent clearance at steady-148 state (CL_{ss}/F), and the accumulation ratio for both periods. 149 The parameters AUC₀₋₁₂, C₁₂, C_{avg}, and C_{max} were separately evaluated using a generalized estimating 150 equations (GEE) model with log-link with the primary predictor of period. Geometric means and 151 corresponding 95% confidence intervals were calculated for each treatment group. The 90% confidence 152 intervals were estimated from the GEE model for the geometric mean ratios for (doravirine + rifapentine 153 + isoniazid)/doravirine) alone. The statistical analysis was performed on R version 3.6.1 with the geepack 154 package.⁸ Sample size was calculated from the reported variability in doravirine trough concentration in 155 HIV-infected men (1540 nmol/L and 95% CI: 1110-2140 nmol/L).⁹ Using a significance level of 5% with a 156 two-sided paired t-test, a sample size of 11 provided greater than 80% power to detect a change of 50% 157 in doravirine trough concentrations. The magnitude of change was selected on the assumption that a 158 50% change in trough concentrations would be clinically significant.

Plasma doravirine pharmacokinetic parameters were estimated using a non-compartmental

159

142

160 Results

161 Subject demographics

Eleven subjects (10 male and 1 female) were enrolled and completed the study with a mean (<u>+</u>SD) age of
46.4 (<u>+</u> 9.9) years old. Subjects were Black or African American (73%) or white (27%). Mean body
weights were 92.9 kg (range 75.5-109.4 kg) which allowed the maximum doses of rifapentine and

isoniazid (900 mg) for all subjects during the study. The mean (<u>+</u>SD) body mass index was 31.2 (<u>+</u>2.6)
 kg/m². All subjects were included in the PK and safety analysis.

167 **Doravirine plasma concentration time profile**

168 All pharmacokinetic data were included in the analysis and figures. All subjects had undetectable 169 doravirine concentrations in plasma at pre-dose prior to beginning study day 1. Two subjects had 170 undetectable doravirine concentrations in plasma at the 72-hour time point in the second period (study 171 day 24). Mean doravirine plasma concentration profiles alone and in combination with rifapentine and 172 isoniazid for all sampled points was plotted against time (figure 2). The mean doravirine plasma 173 concentration during the dosing interval for doravirine alone or in combination with rifapentine and 174 isoniazid is shown in **figure 3.** Co-administration with once-weekly rifapentine and isoniazid modestly reduced the steady-state AUC₀₋₁₂, C₁₂, C_{avg}, and C_{max} by 29%, 31%, 29%, and 25%, respectively. Figure 4 175 176 displays the individual and geometric mean ratios for rifapentine + isoniazid + doravirine/doravirine 177 alone AUC₀₋₁₂, C₁₂, C_{avg}, and C_{max}. Rifapentine co-administration reduced doravirine half-life by 58% while 178 increasing the steady-state total plasma clearance by 41% (table 1).

179 Safety and Tolerability

180 Doravirine in combination with rifapentine and isoniazid was generally well tolerated with no 181 serious adverse events. Adverse events were mild in intensity with 9 out of 11 subjects (82%) reporting 182 at least one adverse event. The most common adverse event throughout the study was intravenous 183 catheter site pain and redness (45.5%) where blood sampling occurred. Nausea and vomiting was the 184 most common reported adverse event (9%) during period 1 where doravirine was dosed alone. During 185 the second period where doravirine was co-administered with weekly rifapentine and isoniazid, one 186 subject (9%) reported dysuria following the second week of rifapentine and isoniazid dosing. The same 187 subject reported chills, headache, and a fever following the third week of dosing rifapentine and

isoniazid. These symptoms subsided and were resolved two days after the report adverse event. All

189 laboratory parameters were within normal limits during the course of the study.

190

191 Discussion

In patients infected with HIV and LTBI, the current therapeutic options include daily rifampin or rifabutin for 4 months or isoniazid daily for 9 months. Although both INH and rifamycin based regimens are similarly effective in the treatment of LTBI in patients with HIV, patients are more likely to complete shorter and convenient regimens. While rifapentine and isoniazid affords a shorter duration and dosing frequency for treatment than INH monotherapy, drug interaction studies are infrequent to evaluate this regimen co-administered with HIV antiretroviral therapies.

198 In this study, twice-daily doses of doravirine 100 mg dosed to steady-state was selected based 199 on nonparametric superposition predictions from a single dose drug interaction study with rifabutin.⁶ 200 Furthermore, doravirine was generally well tolerated across multiple doses of up to 750 mg with robust 201 antiviral activity at 200 mg once-daily in patients with HIV-1 with a terminal half-life of approximately 15 hours.^{9, 10} Therefore, a twice-daily dosing regimen for 4 days was selected in the first period for several 202 203 reasons: 1) given the safety profile of doravirine reported previously, a 100 mg twice-daily dose of 204 doravirine is expected to not be a safety concern, 2) it is expected that the 100 mg twice-daily doses of 205 doravirine should obtain the same level of virological efficacy as seen with patients dosed 200 mg once-206 daily, and 3) steady-state exposure is expected within 3 days of dosing allowing steady-state doravirine 207 concentrations to be sampled by the fourth study day in the first period. 208 Doravirine is contraindicated when co-administered with drugs that are strong CYP3A inducers. 209 The current study evaluated the pharmacokinetics of steady-state doravirine in the presence of

210 rifapentine and isoniazid. Once-weekly doses of rifapentine and isoniazid moderately reduced doravirine

steady AUC₀₋₁₂ and C₁₂ following twice-daily doses of 100 mg doravirine in healthy volunteers. This

212 reduction in exposure is reflected by the increase in doravirine steady-state clearance (8.4 L/hour versus 213 5.9 L/hour) and a shortened half-life (15.2 hours versus 6.4 hours) in the presence of rifapentine and 214 isoniazid. Rifapentine is a potent inducer of CYP450 metabolizing enzyme specifically impacting CYP3A4, CYP2C8, and CYP2C9 isoenzymes.¹¹ The potency of induction is 45% greater than rifabutin, with rifampin 215 216 being the most potent of the anti-tuberculosis rifamycins.¹² Therefore, this reduction in exposure with 217 subsequent increase in doravirine clearance was expected as seen in previous drug interaction studies, 218 where rifampin and rifabutin co-administration significantly reduced doravirine trough values following once-daily dosing.^{5, 6} This study reinforces the time-dependent change in the metabolic induction 219 220 capacity of once-weekly rifapentine. As seen in figure 5, mean doravirine C₁₂ concentrations reached a 221 nadir approximately 2 days following rifapentine dosing (study day 14).

222 In phase II studies doravirine doses of 25 mg and 200 mg have comparable virological efficacy in 223 patients with HIV-1 with geometric mean C_{24} values at 107 ng/mL (77-149 ng/mL).⁹ While there is no 224 established therapeutic range that correlates with clinically sustained virological suppression and 225 efficacy, an NNRTI class-specific steady-state concentration 6-fold above the in vitro 50% inhibitory concentration (IC₅₀) for wild type HIV would result in approximately 99% maximal viral reduction.¹⁵ In 226 227 the case of doravirine, the in vitro IC_{50} is 5.1 ng/mL which results in a pharmacokinetic target of 31.5 ng/mL.¹⁶ Although doravirine mean steady-state trough values at the second week of rifapentine dosing 228 229 were below the observed C_{24} in pivotal trials (figure 5), concentrations still remained 7.6-fold above the 230 values associated with maximal viral load reduction. Moreover, the nadir trough values seen at the 231 second week of rifapentine and isoniazid dosing were >50% of the values following a 25 mg dose in the 232 patient population where the antiviral activity was comparable to the higher dose levels. Based on the 233 strong correlation of doravirine trough values in the exposure-response relationship, trough 234 concentrations seen in this study are within the ranges associated with a >80% proportion of individuals 235 achieving HIV-1 RNA copies of <50 copies/mL.¹⁷

236 It should be noted that a true nadir cannot be defined given the absence of trough collections 237 on study days 17-19. Using the elimination rate and distribution volume of doravirine in the presence of 238 rifapentine observed in this study, trough values were predicted for study days 16-19 (figure 6). The 239 mean plasma trough concentrations at 12 and 24 hours was predicted to be 376.4 ng/mL suggesting 240 that concentrations of doravirine can be sustained above the IC_{50} during a once weekly course of 241 rifapentine. Although we observed a persistent reduction of up to 82% in doravirine C_{12} six days 242 following the last dose of rifapentine, the time-dependent metabolic induction is nonetheless similar to 243 reports in literature which observed up to 2-4 days of maximal induction following rifapentine administration.^{13, 14} 244

245 There were minimal adverse events observed in this study. Compared to previous reports using this regimen in drug interaction studies with other HIV-1 antiretrovirals,¹⁴ co-administration of 246 247 doravirine with once-weekly rifapentine and isoniazid at maximum doses was well tolerated. The most 248 common adverse event reported by 45.5% of subjects was intravenous catheter site pain and redness 249 which was unrelated to the study drugs. Only one female subject reported flu-like symptoms which 250 included fever, chills, and headache after the second dose of rifapentine, isoniazid, and doravirine. This 251 was anticipated, as a high incidence of flu-like symptoms have been reported following high dose rifapetine and isoniazid, particularly in older aged white women.¹⁸ Furthermore, safety lab value trends 252 253 were within the normal ranges throughout the entirety of the study.

254 Several limitations should be noted. The study did not analyze the primary metabolite, M9, 255 which is a direct result of CYP-mediated oxidation. In the presence of an inducer such as rifapentine and 256 twice-daily dosing of doravirine, the exposure of M9 is expected to increase. The impact of the M9 257 metabolite on safety is unclear, as M9 is present as only 13% of parent dose, does not accumulate with 258 repeated dosing, and does not have activity against HIV reverse transcriptase.¹⁹ The study also enrolled 259 mostly male participants (10 male vs. 1 female). Although there was a gender imbalance in this study,

gender does not impact the pharmacokinetics of doravirine.¹⁷ Lastly, doravirine trough concentrations
were not collected for study days 17-19 during period two of the study. As such a true nadir cannot be
confirmed with certainty during that period where doravirine was co-administered with rifapentine and
isoniazid.

In summary, once-weekly oral rifapentine and isoniazid moderately reduced the AUC₀₋₁₂ and C₁₂ of twice-daily 100 mg doravirine by 29% and 31%, respectively. This reduction, however, was within the trough values associated with virological efficacy seen in pivotal clinical studies. As a result, doravirine 100 mg administered twice-daily may be considered to mitigate the drug interaction effect of rifapentine where the modest reduction in doravirine exposure is unlikely to be clinically relevant in a

269 virally suppressed patient.

270

271 Study Highlights

272 What is the current knowledge of the topic?

273 Co-infection with latent tuberculosis in persons living with Human Immunodeficiency Virus-1 (HIV-1) is

- 274 common. Rifapentine is an anti-tuberculosis antibiotic available as once-weekly treatment for latent
- tuberculosis and is a potent inducer of cytochrome (CYP) 3A metabolic enzyme. There are limited
- studies that evaluate rifapentine in the presence with antiretrovirals, including doravirine, a novel non-
- 277 nucleoside reverse transcriptase inhibitor metabolized by CYP3A.

278 What question did this study address?

- 279 This study evaluated the pharmacokinetics of twice-daily doravirine when co-administered with once-
- 280 weekly rifapentine and isoniazid.

281 What does this study add to our knowledge?

- 282 Following rifapentine-mediated CYP3A induction, doravirine steady-state trough concentrations
- 283 declined in a time-dependent manner. Steady-state clearance of doravirine increased with a >50%

284	reduction in	plasma half-life	following	g co-administration	with rifapentine	and isoniazid. Des	pite this
-----	--------------	------------------	-----------	---------------------	------------------	--------------------	-----------

reduction, doravirine exposure is in a range that is likely to maintain viral suppression.

286 How might this change clinical pharmacology or translational science?

- 287 Doravirine 100 mg given twice-daily can mitigate the interactive effects of once-weekly rifapentine. Co-
- administration was generally well tolerated with a modest decrease in doravirine exposure unlikely to
- 289 be clinically relevant. This dosing regimen may offer an alternative to virally suppressed patients on
- antiretroviral therapy co-infected with latent tuberculosis and considering a rifapentine based regimen.
- 291

292 Acknowledgements

- 293 We thank the volunteers, nurses, and staff of the Clinical Research Unit at Thomas Jefferson University
- 294 for their contributions to this research.
- 295

296 Author Contributions

- 297 E.L. designed the wrote the manuscript, research, and analyzed the data; J.S. and T.Z. contributed to
- analytical tools and analyzed the data; R.Z. and W.K.K. performed the research.

300 References

- Molina, J.M., *et al.* Doravirine versus ritonavir-boosted darunavir in antiretroviral-naive adults
 with HIV-1 (DRIVE-FORWARD): 48-week results of a randomised, double-blind, phase 3, non inferiority trial. *Lancet HIV.* 5, e211-e220 (2018).
- Orkin, C., et al. Doravirine/Lamivudine/Tenofovir Disoproxil Fumarate is Non-inferior to
 Efavirenz/Emtricitabine/Tenofovir Disoproxil Fumarate in Treatment-naive Adults With Human
 Immunodeficiency Virus-1 Infection: Week 48 Results of the DRIVE-AHEAD Trial. *Clin Infect Dis.* 535-544 (2019).
- Sanchez, R.I., *et al.* Characterisation of the absorption, distribution, metabolism, excretion and mass balance of doravirine, a non-nucleoside reverse transcriptase inhibitor in humans.
 Xenobiotica. 49, 422-432 (2019).
- 3114.Sterling, T.R., et al. Three months of rifapentine and isoniazid for latent tuberculosis infection. N312Engl J Med. 365, 2155-2166 (2011).
- 3135.Yee, K.L., et al. The Effect of Single and Multiple Doses of Rifampin on the Pharmacokinetics of314Doravirine in Healthy Subjects. Clin Drug Investig. **37**, 659-667 (2017).
- Khalilieh, S.G., et al. Multiple Doses of Rifabutin Reduce Exposure of Doravirine in Healthy
 Subjects. J Clin Pharmacol. (2018).
- Sterling, T.R., *et al.* Three months of weekly rifapentine and isoniazid for treatment of
 Mycobacterium tuberculosis infection in HIV-coinfected persons. *AIDS.* **30**, 1607-1615 (2016).
 Højsgaard, S., Halekoh, U. & Yan, J. The R Package geepack for Generalized Estimating Equations.
- 320 2005. 15, 11 (2005).
 321 9. Schurmann, D., et al. A randomized, double-blind, placebo-controlled, short-term monotherapy
- study of doravirine in treatment-naive HIV-infected individuals. *AIDS*. **30**, 57-63 (2016).
 10. Anderson, M.S., *et al.* Safety, tolerability and pharmacokinetics of doravirine, a novel HIV nonnucleoside reverse transcriptase inhibitor, after single and multiple doses in healthy subjects. *Antivir Ther.* **20**, 397-405 (2015).
- 326 11. Burman, W.J., Gallicano, K. & Peloquin, C. Comparative pharmacokinetics and 327 pharmacodynamics of the rifamycin antibacterials. *Clin Pharmacokinet*. **40**, 327-341 (2001).
- Regazzi, M., Carvalho, A.C., Villani, P. & Matteelli, A. Treatment optimization in patients coinfected with HIV and Mycobacterium tuberculosis infections: focus on drug-drug interactions with rifamycins. *Clin Pharmacokinet*. **53**, 489-507 (2014).
- 13. Keung, A., et al. Enzyme induction observed in healthy volunteers after repeated administration
 of rifapentine and its lack of effect on steady-state rifapentine pharmacokinetics: part I. Int J
 Tuberc Lung Dis. 3, 426-436 (1999).
- Brooks, K.M., *et al.* Cytokine-Mediated Systemic Adverse Drug Reactions in a Drug-Drug
 Interaction Study of Dolutegravir With Once-Weekly Isoniazid and Rifapentine. *Clin Infect Dis.* **67**, 193-201 (2018).
- 33715.Xu, Y., et al. Characterizing Class-Specific Exposure-Viral Load Suppression Response of HIV338Antiretrovirals Using A Model-Based Meta-Analysis. Clin Transl Sci. 9, 192-200 (2016).
- Lai, M.T., *et al.* In vitro characterization of MK-1439, a novel HIV-1 nonnucleoside reverse
 transcriptase inhibitor. *Antimicrob Agents Chemother.* 58, 1652-1663 (2014).
- Yee, K.L., Ouerdani, A., Claussen, A., de Greef, R. & Wenning, L. Population Pharmacokinetics of
 Doravirine and Exposure-Response Analysis in Individuals with HIV-1. *Antimicrob Agents Chemother.* 63, (2019).

- Sterling, T.R., *et al.* Flu-like and Other Systemic Drug Reactions Among Persons Receiving Weekly
 Rifapentine Plus Isoniazid or Daily Isoniazid for Treatment of Latent Tuberculosis Infection in the
 PREVENT Tuberculosis Study. *Clin Infect Dis.* **61**, 527-535 (2015).
- 347 19. Sanchez, R.I., *et al.* Characterisation of the absorption, distribution, metabolism, excretion and
 348 mass balance of doravirine, a non-nucleoside reverse transcriptase inhibitor in humans.
 349 Xenobiotica. 1-11 (2018).
- 350

Figure and Table Legends

Table 1. Steady-state doravirine pharmacokinetic parameters^a and summary statistics following twice-

daily doses of doravirine 100 mg alone or twice-daily doravirine 100 mg co-administered with once-

weekly rifapentine and isoniazid.

	Doravirine +	Doravirine	Doravirine + rifapentine /
	rifapentine		Doravirine
Parameter	Geometric mea	Geometric mean ratio	
		(90% CI)	
AUC ₀₋₁₂ (hr x ug/mL)	12.3 (10.4-14.3)	17.3 (14.9-20.0)	0.71 (0.60-0.85)
C ₁₂ (ug/mL)	0.9 (0.7-1.0)	1.2 (1.0-1.4)	0.69 (0.54-0.89)
C _{avg} (ug/mL) ^b	1.0 (0.8-1.2)	1.4 (1.2-1.7)	0.71 (0.60-0.85)
C _{max} (ug/mL)	1.3 (1.1-1.5)	1.7 (1.5-2.0)	0.75 (0.63-0.88)
t _{1/2} (hr) ^c	6.4 (17.0)	15.2 (19.4)	
CL/F (L/hr) ^c	8.4 (26.1)	5.9 (24.0)	
Accumulation ratio	1.2-1.6	1.8-3.2	

^{a.} Parameters with exposures are expressed as ug/mL.

^{b.} The average steady-state plasma concentration during multiple-dose administration was computed as AUC₀₋₁₂/dosing interval.

^{c.} Values for $t_{1/2}$ and CL/F are expressed as the geometric mean (%CV).

Figure 1. Study schematic. B₆ = vitamin B₆ (pyridoxine), BID = Twice-daily, DOR = doravirine, INH = isoniazid, PK = pharmacokinetics, RPT = rifapentine. Arrows indicate blood for PK or safety laboratory.



Figure 2. Mean (<u>+</u> SD) doravirine plasma concentration-time profiles following twice-daily doses of

doravirine 100 mg alone or together with once-weekly rifapentine and isoniazid. The inset represents
 the concentration-time profile plotted on a log-linear scale.



- **Figure 3.** Mean (<u>+</u> SD) doravirine plasma concentration-time profiles during the 12 hour dosing interval
- following twice-daily doses of doravrine 100 mg alone or together with once-weekly rifapentine and
- 412 isoniazid. Sampling at time 0 and 12 hours were taken prior to the second dose of doravirine. The inset
- 413 represents the concentration-time profile plotted on a log-linear scale.



432 Figure 4. Individual ratios (Doravirine/doravirine + rifapentine), geometric mean ratios, and

433 corresponding 90% confidence intervals for doravirine exposure parameters. AUC₀₋₁₂ = Area under the

- 434 concentration time during the 12 hour dosing interval, C_{12} = plasma concentration at the end of the
- dosing interval prior to the second dose, C_{avg} = average steady-state plasma concentration during the
- 436 dosing interval, C_{max} = maximum steady-state concentration



Figure 5. Trend of doravirine C_{12} concentrations throughout the study study days across the two study periods. The values represent the geometric mean C₁₂ concentration reported for that study day. Day 5 was used as a reference to calculate the percent change in C_{12} concentrations for subsequent days. The green down arrows indicate rifapentine, isoniaizd, and pyridoxine co-administered with the morning dose of doravirine 100 mg. The horizontal red line indicates the steady-state pharmacokinetic target 6-fold above the IC₅₀ associated with 99% viral reduction. The horizoantal blue line represents the steady-state C24 (% coefficient of variance) values observed in pivotal studies following once-daily doses of doravirine 100 mg. B6 = pyridoxine, C_{12} = observed trough concentration prior to the second dose for a twice-daily regimen, C_{24} = observed trough concentration prior to the second dose for a once-daily regimen, INH = Isoniazid, PK = pharmacokinetic, RPT = rifapentine.



Figure 6. Predicted mean doravirine plasma concentrations at study days 16-19. The gray shaded regions

represent the standard error of the mean. Predictions were based on doravirine pharmacokinetic
 parameters estimated following co-administration of rifapentine on day 21.

