



Table 24. Outcomes of Randomized Treatment Through Week 48 (Study 88)

Table with 3 columns: Outcome, Logipinar and Rivinor + nevirapine + NRTIs (n=144), Logipinar and Rivinor Twice Daily (n=144). Rows include Response, Virologic failure, Relapse, etc.

Table 25. Outcomes of Randomized Treatment Through Week 48 (Study 89)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 26. Outcomes of Randomized Treatment Through Week 48 (Study 90)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 27. Outcomes of Randomized Treatment Through Week 48 (Study 91)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 28. Outcomes of Randomized Treatment Through Week 48 (Study 92)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 29. Outcomes of Randomized Treatment Through Week 48 (Study 93)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 30. Outcomes of Randomized Treatment Through Week 48 (Study 94)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 31. Outcomes of Randomized Treatment Through Week 48 (Study 95)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 32. Outcomes of Randomized Treatment Through Week 48 (Study 96)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 33. Outcomes of Randomized Treatment Through Week 48 (Study 97)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 34. Outcomes of Randomized Treatment Through Week 48 (Study 98)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 35. Outcomes of Randomized Treatment Through Week 48 (Study 99)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 36. Outcomes of Randomized Treatment Through Week 48 (Study 100)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 37. Outcomes of Randomized Treatment Through Week 48 (Study 101)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 38. Outcomes of Randomized Treatment Through Week 48 (Study 102)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 39. Outcomes of Randomized Treatment Through Week 48 (Study 103)

Table with 3 columns: Outcome, Logipinar and Rivinor + NRTIs (n=292), Logipinar and Rivinor Twice Daily (n=292). Rows include Virologic Success, Virologic failure, Relapse, etc.

Table 40. Pharmacokinetic Properties of Logipinar

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 41. Steady-State Pharmacokinetic Parameters of Logipinar, Mean ± SD

Table with columns: Parameter, Value. Rows include C_{max} (ng/mL), C_{trough} (ng/mL), AUC (ng·h/mL), etc.

Table 42. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 43. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 44. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

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Table 45. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

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Table 46. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 47. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 48. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 49. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 50. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 51. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 52. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 53. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 54. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

Table 55. Pharmacokinetic Properties of Logipinar in Patients with HIV-1 Infection

Table with columns: Parameter, Value. Rows include T_{1/2} (h), C_{max} (ng/mL), AUC (ng·h/mL), etc.

The yellow, 100 mg logipinar/25 mg rivinor tablets contain the following inactive ingredients: colloidal silicon dioxide, copovidone, sodium stearoyl fumarate, sorbitol monolaurate and opafolony which contains colloidal anhydrous silica, copovidone, sodium stearoyl fumarate, sorbitol monolaurate and opafolony which contains colloidal anhydrous silica, copovidone, hydroxypropyl cellulose, iron oxide yellow, polyethylene glycol, polyborate 80, talc and titanium dioxide.

12.1 Mechanism of Action
Logipinar and Rivinor is a fixed-dose combination of HIV-1 antiretroviral drugs [see Microbiology (12.4)] and Rivinor. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar, thereby providing increased plasma levels of logipinar.

12.2 Pharmacokinetics
The pharmacokinetic properties of logipinar are summarized in Table 13. The steady-state pharmacokinetic parameters of logipinar are summarized in Table 14. Under fed conditions, logipinar concentrations were similar following administration of logipinar and rivinor tablets to capsules with less pharmacokinetic variability. Under conditions (50% food, 25% from fat), logipinar concentrations were similar following administration of logipinar and rivinor capsules and oral solution.

12.3 Pharmacokinetics
The pharmacokinetic properties of rivinor are summarized in Table 15. The steady-state pharmacokinetic parameters of rivinor are summarized in Table 16. Under fed conditions, rivinor concentrations were similar following administration of logipinar and rivinor tablets to capsules with less pharmacokinetic variability. Under conditions (50% food, 25% from fat), rivinor concentrations were similar following administration of logipinar and rivinor capsules and oral solution.

12.4 Microbiology
Logipinar, an inhibitor of the HIV-1 protease, prevents cleavage of the viral Gag-Pol polyprotein, resulting in the production of immature, non-infectious viral particles.
Antiviral Activity
In the absence of human serum, the mean 50% effective concentration (EC₅₀) values of logipinar against five different HIV-1 genotypes (B, CRF01_AE, CRF02_AG, CRF07_BC, CRF08_OB) ranged from 10 to 100 nM (0.0001 to 0.001 μg/mL). In the presence of human serum, the mean EC₅₀ values of logipinar against these HIV-1 laboratory strains ranged from 0.5 to 10 nM (0.0005 to 0.01 μg/mL).
HIV-1 isolates with reduced susceptibility to logipinar have been selected in cell culture. The presence of rivinor does not appear to influence the selection of logipinar-resistant viruses in cell culture.

12.5 Resistance Information
Logipinar and Rivinor have been used in combination with other HIV-1 protease inhibitors. The antiviral activity in cell culture of logipinar against clinical isolates from patients previously treated with a single protease inhibitor was reduced compared to logipinar monotherapy. The presence of rivinor did not appear to influence the selection of logipinar-resistant viruses in cell culture.
Cross-resistance - Nonclinical Studies
Viruses with reduced susceptibility to logipinar have been observed among HIV-1 protease inhibitors. The antiviral activity in cell culture of logipinar against clinical isolates from patients previously treated with a single protease inhibitor was reduced compared to logipinar monotherapy.

12.6 Pregnancy
There is a pregnancy exposure registry that monitors pregnancy outcomes in women exposed to logipinar and rivinor during pregnancy. Physicians are encouraged to register patients by calling the Antiretroviral Pregnancy Registry at 1-800-233-2323.
Risk Summary
Available data from the Antiretroviral Pregnancy Registry show no difference in the risk of overall major birth defects compared to the background rate for major birth defects of 2.7% in the U.S. reference population of the Metropolitan Atlanta Congenital Defects Program (MACDP) [see Data]. The estimated background rate of major birth defects in live births was 2.5% (95% CI: 1.4%-3.9%) following first-trimester exposure to logipinar-containing regimens and 3.0% (95% CI: 2.4%-3.8%) following first-trimester exposure to rivinor-containing regimens. The presence of both drugs in live births was 2.5% (95% CI: 1.4%-3.9%) following first-trimester exposure to logipinar-containing regimens and 3.0% (95% CI: 2.4%-3.8%) following first-trimester exposure to rivinor-containing regimens. The presence of both drugs in live births was 2.5% (95% CI: 1.4%-3.9%) following first-trimester exposure to logipinar-containing regimens and 3.0% (95% CI: 2.4%-3.8%) following first-trimester exposure to rivinor-containing regimens. The presence of both drugs in live births was 2.5% (95% CI: 1.4%-3.9%) following first-trimester exposure to logipinar-containing regimens and 3.0% (95% CI: 2.4%-3.8%) following first-trimester exposure to rivinor-containing regimens.

12.7 Lactation
The Centers for Disease Control and Prevention recommend that HIV-1 infected mothers not breastfeed their infants to avoid risking potential transmission of HIV-1. Because of the potential for HIV transmission in HIV-infected infants, 2) developing viral resistance (in HIV-positive infants), and 3) adverse reactions in the breastfed infant, instruct mothers to breastfeed if they are receiving logipinar and rivinor.

12.8 Fertility and Reproductive Potential
Logipinar and Rivinor may reduce the efficacy of combined hormonal contraceptives. Advise patients using combined hormonal contraceptives to use an effective alternative contraceptive method or an additional barrier method of contraception [see Drug Interactions (7.3)].
12.9 Pediatric Use
The safety, efficacy, and pharmacokinetic properties of logipinar and rivinor in pediatric patients below the age of 14 years have not been established. Logipinar and Rivinor should not be administered once daily in pediatric patients.
An open-label, multicenter, randomized, parallel, controlled trial evaluating the pharmacokinetic profile, tolerability, safety, and efficacy of logipinar and Rivinor in HIV-1 infected children aged 6 to 14 years. The study included 143 children, 14 days after 6 months of age, and 6 months after 12 months of age. The study included 143 children, 14 days after 6 months of age, and 6 months after 12 months of age. The study included 143 children, 14 days after 6 months of age, and 6 months after 12 months of age.

12.10 Geriatric Use
Clinical studies of logipinar and rivinor did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently to general population. Caution should be exercised when administering the drug to the elderly and monitoring of logipinar and rivinor in elderly patients reflecting the greater frequency of decreased hepatic, renal, or cardiac function, and of concomitant disease or other drug therapy.
12.11 Hepatic Impairment
Logipinar and Rivinor is principally metabolized by the liver; therefore, caution should be exercised when administering the drug to patients with hepatic impairment because logipinar concentrations may be increased [see Warnings and Precautions (5.4) and Clinical Pharmacology (12.3)].
12.12 OVERDOSAGE
Overdoses with logipinar and rivinor oral solution have been reported. One of these reports described fatal cardiac shock in a 2.1 kg infant who received 100 mg of logipinar and rivinor oral solution (500 mg/25 mg) over a 14-day period. Symptoms were consistent with the recommended dosage of logipinar and rivinor oral solution. The patient was intubated and received mechanical ventilation. The patient was intubated and received mechanical ventilation. The patient was intubated and received mechanical ventilation.

12.13 Description
Logipinar and Rivinor Tablets, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
Logipinar USP is chemically designated as 1-[5-(1R)-1-(4S)-2,3-dihydro-1H-benzofuran-2-yl]-4-hydroxy-2-methyl-1H-imidazole-5-carboxamide (logipinar) and 2,2,6,6-tetramethyl-1-piperidine-1-oxide (rivinor). Logipinar USP is a white to off-white powder. It is highly soluble in methanol, ethanol and in isopropyl alcohol. Logipinar USP has the following structural formula:
Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.14 Description
Logipinar and Rivinor Capsules, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.15 Description
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12.16 Description
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.17 Description
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.18 Description
Logipinar and Rivinor Oral Solution, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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12.19 Description
Logipinar and Rivinor Tablets, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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12.20 Description
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.21 Description
Logipinar and Rivinor Oral Solution, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.22 Description
Logipinar and Rivinor Tablets, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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12.23 Description
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12.24 Description
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.25 Description
Logipinar and Rivinor Tablets, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.26 Description
Logipinar and Rivinor Capsules, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
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12.27 Description
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Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.28 Description
Logipinar and Rivinor Tablets, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
Logipinar USP is chemically designated as 1-[5-(1R)-1-(4S)-2,3-dihydro-1H-benzofuran-2-yl]-4-hydroxy-2-methyl-1H-imidazole-5-carboxamide (logipinar) and 2,2,6,6-tetramethyl-1-piperidine-1-oxide (rivinor). Logipinar USP is a white to off-white powder. It is highly soluble in methanol, ethanol and in isopropyl alcohol. Logipinar USP has the following structural formula:
Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.29 Description
Logipinar and Rivinor Capsules, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
Logipinar USP is chemically designated as 1-[5-(1R)-1-(4S)-2,3-dihydro-1H-benzofuran-2-yl]-4-hydroxy-2-methyl-1H-imidazole-5-carboxamide (logipinar) and 2,2,6,6-tetramethyl-1-piperidine-1-oxide (rivinor). Logipinar USP is a white to off-white powder. It is highly soluble in methanol, ethanol and in isopropyl alcohol. Logipinar USP has the following structural formula:
Cc1nc2c(nc1)C(=O)N2C3=CC=C(C=C3)C4=CC=CC=C4C5=CC=CC=C5

12.30 Description
Logipinar and Rivinor Oral Solution, USP is a co-formulation of logipinar USP and Rivinor USP. Logipinar USP is an inhibitor of the HIV-1 protease. As co-administered with logipinar and Rivinor, Rivinor inhibits the CYP3A-mediated metabolism of logipinar USP, thereby providing increased plasma levels of logipinar USP.
Logipinar USP is chemically designated as 1-[5-(1R)-1-(4S)-2,3-dihydro-1H-benzofuran-2-yl]-4-hydroxy-2-methyl-1H-imidazole-5-carboxamide (logipinar) and 2,2,6,6-tetramethyl-1-piperidine-1-oxide (rivinor). Logipinar USP is a white to off-white powder. It is highly soluble in methanol, ethanol and in isoprop