HIGHLIGHTS OF PRESCRIBING INFORMATION

These highlights do not include all the information needed to use SELZENTRY safely and effectively. See full prescribing information for SELZENTRY.

SELZENTRY (maraviroc) Tablets Initial U.S. Approval: 2007

WARNING: HEPATOTOXICITY

- See full prescribing information for complete boxed warning
 Hepatotoxicity has been reported which may be preceded by severe rash or other features of a systemic allergic reaction (e.g., fever, eosinophilia, or elevated IgE).
- Immediately evaluate patients with signs or symptoms of hepatitis or allergic reaction. (5.1)

-----RECENT MAJOR CHANGES ------

Boxed Warning, Hepatotoxicity (5.1)	July 2011
Warnings and Precautions, Hepatotoxicity (5.1)	July 2011

-----INDICATIONS AND USAGE------

SELZENTRY is a CCR5 co-receptor antagonist indicated for combination antiretroviral treatment of adults infected with only CCR5-tropic HIV-1.

- In treatment-naive subjects, more subjects treated with SELZENTRY experienced virologic failure and developed lamivudine resistance compared with efavirenz. (12.4,14.3)
- Tropism testing with a highly sensitive tropism assay is required for the appropriate use of SELZENTRY. (1)

 DOSAGE	AND	ADMINIS	TRATIO	N

DOORGE AND ADMINIOTRATIO	
When given with potent CYP3A inhibitors (with or	150 mg
without potent CYP3A inducers) including PIs	twice daily
(except tipranavir/ritonavir), delavirdine (2, 7.1)	
With NRTIs, tipranavir/ritonavir, nevirapine,	300 mg
raltegravir, and other drugs that are not potent	twice daily
CYP3A inhibitors or CYP3A inducers (2, 7.1)	
With potent CYP3A inducers including efavirenz	600 mg
(without a potent CYP3A inhibitor) (2, 7.1)	twice daily
A more complete list of coadministered drugs is listed in	Dosage and

A more complete list of coadministered drugs is listed in *Dosage and Administration* (2).

Dose adjustment may be necessary in patients with renal impairment. (2.2)

-----CONTRAINDICATIONS ----

 SELZENTRY should not be used in patients with severe renal impairment or end-stage renal disease (ESRD) (CrCl <30 mL/min) who are taking potent CYP3A inhibitors or inducers. (4)

FULL PRESCRIBING INFORMATION: CONTENTS*

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------ WARNINGS AND PRECAUTIONS ------

- Hepatotoxicity accompanied by severe rash or systemic allergic reaction including potentially life-threatening events has been reported. Hepatic laboratory parameters including ALT, AST, and bilirubin should be obtained prior to starting SELZENTRY and at other time points during treatment as clinically indicated. If rash or symptoms or signs of hepatitis or allergic reaction develop, hepatic laboratory parameters should be monitored and discontinuation of treatment should be considered. Use caution when administering SELZENTRY to patients with pre-existing liver dysfunction or who are co-infected with viral hepatitis B or C. (5.1)
- More cardiovascular events including myocardial ischemia and/or infarction were observed in treatment-experienced subjects who received SELZENTRY. Use with caution in patients at increased risk of cardiovascular events. (5.2)
- If patients with severe renal impairment or end-stage renal disease (ESRD) receiving SELZENTRY (without concomitant CYP3A inducers or inhibitors) experience postural hypotension, the dose of SELZENTRY should be reduced from 300 mg twice daily to 150 mg twice daily. (5.2)

----- ADVERSE REACTIONS ------

The most common adverse events in treatment-experienced subjects (>8% incidence) which occurred at a higher frequency compared with placebo are upper respiratory tract infections, cough, pyrexia, rash, and dizziness. (6)

To report SUSPECTED ADVERSE REACTIONS, contact ViiV Healthcare at 1-877-844-8872 or FDA at 1-800-FDA-1088 or www.fda.gov/medwatch.

-----DRUG INTERACTIONS ------DRUG INTERACTIONS

- Coadministration with CYP3A inhibitors, including protease inhibitors (except tipranavir/ritonavir) and delavirdine, will increase the concentration of SELZENTRY. (7.1)
- Coadministration with CYP3A inducers, including efavirenz, may decrease the concentration of SELZENTRY. (7.1)

------USE IN SPECIFIC POPULATIONS------

- SELZENTRY should only be used in pregnant women if the potential benefit justifies the potential risk to the fetus. (8.1)
- There are no data available in pediatric patients; therefore, SELZENTRY should not be used in patients aged <16 years. (8.4)

See 17 for PATIENT COUNSELING INFORMATION and MEDICATION GUIDE.

Revised: 7/2011

- 8.5 Geriatric Use
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*Sections or subsections omitted from the full prescribing information are not listed.

8

1 FULL PRESCRIBING INFORMATION

2	WARNING: HEPATOTOXICITY
3	Hepatotoxicity has been reported with use of SELZENTRY. Severe rash or evidence
4	of a systemic allergic reaction (e.g., fever, eosinophilia, or elevated IgE) prior to the
5	development of hepatotoxicity may occur. Patients with signs or symptoms of hepatitis or
6	allergic reaction following use of SELZENTRY should be evaluated immediately [see
7	Warnings and Precautions (5.1)].
8	1 INDICATIONS AND USAGE
9	SELZENTRY, in combination with other antiretroviral agents, is indicated for adult
10	patients infected with only CCR5-tropic HIV-1.
11	This indication is based on analyses of plasma HIV-1 RNA levels in 2 controlled studies
12	of SELZENTRY in treatment-experienced subjects and one study in treatment-naive subjects.
13	Both studies in treatment-experienced subjects were conducted in clinically advanced, 3-class
14	antiretroviral-experienced (nucleoside reverse transcriptase inhibitor [NRTI], non-nucleoside
15	reverse transcriptase inhibitor [NNRTI], protease inhibitor [PI], or enfuvirtide) adults with
16	evidence of HIV-1 replication despite ongoing antiretroviral therapy.
17	The following points should be considered when initiating therapy with SELZENTRY:
18	• Adult patients infected with only CCR5-tropic HIV-1 should use SELZENTRY.
19	• Tropism testing must be conducted with a highly sensitive tropism assay that has
20	demonstrated the ability to identify patients appropriate for use of SELZENTRY. Outgrowth
21	of pre-existing low-level CXCR4- or dual/mixed-tropic HIV-1 not detected by tropism
22	testing at screening has been associated with virologic failure on SELZENTRY [see
23	Microbiology (12.4), Clinical Studies (14.3)].
24	• Use of SELZENTRY is not recommended in subjects with dual/mixed- or CXCR4-tropic
25	HIV-1 as efficacy was not demonstrated in a Phase 2 study of this patient group.
26	• The safety and efficacy of SELZENTRY have not been established in pediatric patients.
27	• In treatment-naive subjects, more subjects treated with SELZENTRY experienced virologic
28	failure and developed lamivudine resistance compared with efavirenz [see Microbiology
29	(12.4), Clinical Studies (14.3)].
30	2 DOSAGE AND ADMINISTRATION
31	2.1 Dose Recommendations for Patients With Normal Renal Function
32	The recommended dose of SELZENTRY differs based on concomitant medications due
33	to drug interactions (see Table 1). SELZENTRY can be taken with or without food.
34	SELZENTRY must be given in combination with other antiretroviral medications.
35	Table 1 gives the recommended dose adjustments [see Drug Interactions (7.1)].
36	

Table 1. Recommended Dosing Regimen

Concomitant Medications	Dose of SELZENTRY
Potent CYP3A inhibitors (with or without a potent CYP3A inducer)	Dose of SELECTRY
including:	
 protease inhibitors (except tipranavir/ritonavir) 	
 delavirdine 	150 mg twice daily
• ketoconazole, itraconazole, clarithromycin	
• other potent CYP3A inhibitors (e.g., nefazodone, telithromycin)	
Other concomitant medications, including tipranavir/ritonavir,	200 ma truica dailer
nevirapine, raltegravir, all NRTIs, and enfuvirtide	300 mg twice daily
Potent CYP3A inducers (without a potent CYP3A inhibitor)	
including:	
• efavirenz	600 mg twice daily
• rifampin	000 mg twice dany
• etravirine	
carbamazepine, phenobarbital, and phenytoin	

2.2 Dose Recommendations for Patients With Renal Impairment

- 40 Table 2 provides dosing recommendations for patients based on renal function and
- 41 concomitant medications.

Table 2. Recommended Dosing Regimens Based on Renal Function

	Dose of SELZENTRY Based on Renal Function				
					End-Stage
		Mild	Moderate		Renal Disease
	Normal	(CrCl >50	$(CrCl \ge 30)$	Severe	(ESRD) On
	(CrCl>80	and ≤ 80	and ≤ 50	(CrCl <30	Regular
Concomitant Medications ^a	mL/min)	mL/min)	mL/min)	mL/min)	Hemodialysis
Potent CYP3A inhibitors	150 mg	150 mg	150 mg	NR	NR
(with or without a CYP3A	twice daily	twice daily	twice daily		
inducer) ^a					
Other concomitant	300 mg	300 mg	300 mg	300 mg	300 mg twice
medications ^a	twice daily	twice daily	twice daily	twice	daily ^b
				daily ^b	
Potent CYP3A Inducers	600 mg	600 mg	600 mg	NR	NR
(without a potent CYP3A	twice daily	twice daily	twice daily		
inhibitor) ^a					

- NR = Not recommended.
- 45 ^a See Table 1 for the list of concomitant medications.

- ^b The dose of SELZENTRY should be reduced to 150 mg twice daily if there are any symptoms
- 47 of postural hypotension [see Warnings and Precautions (5.2)].

48 3 DOSAGE FORMS AND STRENGTHS

- 49 150-mg blue, oval, film-coated tablets debossed with "MVC 150" on one side and plain on
 50 the other.
- 51 300-mg blue, oval, film-coated tablets debossed with "MVC 300" on one side and plain on
 52 the other.

53 4 CONTRAINDICATIONS

54 SELZENTRY should not be used in patients with severe renal impairment or end-stage 55 renal disease (ESRD) (CrCl <30 mL/min) who are taking potent CYP3A inhibitors or inducers.

56 5 WARNINGS AND PRECAUTIONS

57 5.1 Hepatotoxicity

58 Hepatotoxicity with allergic features including life-threatening events has been reported 59 in clinical trials and postmarketing. Severe rash or evidence of systemic allergic reaction 60 including drug-related rash with fever, eosinophilia, elevated IgE, or other systemic symptoms 61 have been reported in conjunction with hepatotoxicity. These events occurred approximately 1 62 month after starting treatment. Among reported cases of hepatitis, some were observed in the 63 absence of allergic features or with no pre-existing hepatic disease.

Appropriate laboratory testing including ALS, AST, and bilirubin should be conducted prior to initiating therapy with SELZENTRY and at other time points during treatment as clinically indicated. Hepatic laboratory parameters should be obtained in any patient who develops rash, or signs or symptoms of hepatitis, or allergic reaction. Discontinuation of SELZENTRY should be considered in any patient with signs or symptoms of hepatitis, or with

69 increased liver transaminases combined with rash or other systemic symptoms.

70 Caution should be used when administering SELZENTRY to patients with pre-existing 71 liver dysfunction or who are coinfected with viral hepatitis B or C. The safety and efficacy of 72 SELZENTRY have not been specifically studied in patients with significant underlying liver 73 disorders. In studies of treatment-experienced HIV-infected subjects, approximately 6% of 74 subjects were co-infected with hepatitis B and approximately 6% were co-infected with hepatitis 75 C. Due to the small number of co-infected subjects studied, no conclusions can be drawn 76 regarding whether they are at an increased risk for hepatic adverse events with administration of 77 SELZENTRY.

78 **5.2 Cardiovascular Events**

79 Use with caution in patients at increased risk for cardiovascular events. Eleven subjects

80 (1.3%) who received SELZENTRY had cardiovascular events including myocardial ischemia

- 81 and/or infarction during the Phase 3 studies in treatment-experienced studies (total exposure
- 82 609 patient-years [300 on SELZENTRY once daily + 309 on SELZENTRY twice daily]), while
- 83 no subjects who received placebo had such events (total exposure 111 patient-years). These

subjects generally had cardiac disease or cardiac risk factors prior to use of SELZENTRY, and
 the relative contribution of SELZENTRY to these events is not known.

In the Phase 2b/3 study in treatment-naive subjects, 3 subjects (0.8%) who received
SELZENTRY had events related to ischemic heart diseases and 5 subjects (1.4%) who received
efavirenz had such events (total exposure 506 and 508 patient-years for SELZENTRY and
efavirenz, respectively).

When SELZENTRY was administered to healthy volunteers at doses higher than the
recommended dose, symptomatic postural hypotension was seen at a greater frequency than in
placebo. However, when SELZENTRY was given at the recommended dose in HIV subjects in
Phase 3 studies, postural hypotension was seen at a rate similar to placebo (approximately 0.5%).
Caution should be used when administering SELZENTRY in patients with a history of postural
hypotension or on concomitant medication known to lower blood pressure.

96 Postural Hypotension in Patients With Renal Impairment: Patients with impaired 97 renal function may have cardiovascular co-morbidities and could be at increased risk of 98 cardiovascular adverse events triggered by postural hypotension. An increased risk of postural 99 hypotension may occur in patients with severe renal insufficiency or in those with ESRD due to 100 increased maraviroc exposure in some patients. SELZENTRY should be used in patients with 101 severe renal impairment or ESRD only if they are not receiving a concomitant potent CYP3A 102 inhibitor or inducer. However, the use of SELZENTRY in these patients should only be 103 considered when no alternative treatment options are available. If patients with severe renal 104 impairment or ESRD experience any symptoms of postural hypotension while taking 300 mg 105 twice daily, the dose should be reduced to 150 mg twice daily [see Dosage and Administration 106 (2.2)].

107 **5.3 Immune Reconstitution Syndrome**

108Immune reconstitution syndrome has been reported in patients treated with combination109antiretroviral therapy, including maraviroc. During the initial phase of combination antiretroviral110treatment, patients whose immune system responds may develop an inflammatory response to111indolent or residual opportunistic infections (such as infection with *Mycobacterium avium*,112cytomegalovirus, *Pneumocystis* jirovecii, *Mycobacterium* tuberculosis, or reactivation of *Herpes*113simplex and *Herpes* zoster), which may necessitate further evaluation and treatment.

114 **5.4 Potential Risk of Infection**

115 SELZENTRY antagonizes the CCR5 co-receptor located on some immune cells, and 116 therefore could potentially increase the risk of developing infections. The overall incidence and 117 severity of infection, as well as AIDS-defining category C infections, was comparable in the 118 treatment groups during the Phase 3 treatment-experienced studies of SELZENTRY. While there 119 was a higher rate of certain upper respiratory tract infections reported in the arm receiving 120 SELZENTRY compared with placebo (23% versus 13%), there was a lower rate of pneumonia 121 (2% vs 5%) reported in subjects receiving SELZENTRY. A higher incidence of Herpes virus 122 infections (11 per 100 patient-years) was also reported in the arm receiving SELZENTRY when 123 adjusted for exposure compared with placebo (8 per 100 patient-years).

124 In the Phase 2b/3 study in treatment-naive subjects, the incidence of AIDS-defining 125 Category C events when adjusted for exposure was 1.8 for SELZENTRY compared with 2.4 for 126 efavirenz per 100 patient-years of exposure. 127 Patients should be monitored closely for evidence of infections while receiving 128 SELZENTRY. 129 5.5 Potential Risk of Malignancy 130 While no increase in malignancy has been observed with SELZENTRY, due to this 131 drug's mechanism of action it could affect immune surveillance and lead to an increased risk of 132 malignancy. 133 The exposure-adjusted rate for malignancies per 100 patient-years of exposure in 134 treatment-experienced studies was 4.6 for SELZENTRY compared with 9.3 on placebo. In 135 treatment-naive subjects, the rates were 1.0 and 2.4 per 100 patient-years of exposure for 136 SELZENTRY and efavirenz, respectively. 137 Long-term follow-up is needed to more fully assess this risk. 138 6 ADVERSE REACTIONS 139 The following adverse reactions are discussed in other sections of the labeling: 140 Hepatotoxicity [see Boxed Warning, Warnings and Precautions (5.1)] 141 Cardiovascular events [see Warnings and Precautions (5.2)] • 142 **Clinical Trials Experience** 6.1 143 Because clinical trials are conducted under widely varying conditions, adverse reaction 144 rates observed in the clinical trials of a drug cannot be directly compared with rates in the 145 clinical trials of another drug and may not reflect the rates observed in practice. 146 Studies in Treatment-Experienced Subjects: The safety profile of SELZENTRY is 147 primarily based on 840 HIV-infected subjects who received at least 1 dose of SELZENTRY 148 during two Phase 3 trials. A total of 426 of these subjects received the indicated twice-daily 149 dosing regimen. 150 Assessment of treatment-emergent adverse events is based on the pooled data from 151 2 studies in subjects with CCR5-tropic HIV-1 (A4001027 and A4001028). The median duration 152 of therapy with SELZENTRY for subjects in these studies was 48 weeks, with the total exposure 153 on SELZENTRY twice daily at 309 patient-years versus 111 patient-years on placebo + 154 optimized background therapy (OBT). The population was 89% male and 84% white, with mean 155 age of 46 years (range: 17 to 75 years). Subjects received dose equivalents of 300 mg maraviroc 156 once or twice daily. 157 The most common adverse events reported with twice-daily therapy with SELZENTRY 158 with frequency rates higher than placebo, regardless of causality, were upper respiratory tract 159 infections, cough, pyrexia, rash, and dizziness. Additional adverse events that occurred with 160 once-daily dosing at a higher rate than both placebo and twice-daily dosing were diarrhea, 161 edema, influenza, esophageal candidiasis, sleep disorders, rhinitis, parasomnias, and urinary 162 abnormalities. In these 2 studies, the rate of discontinuation due to adverse events was 5% for

- 163 subjects who received SELZENTRY twice daily + OBT as well as those who received placebo +
- 164 OBT. Most of the adverse events reported were judged to be mild to moderate in severity. The

165 data described below occurred with twice-daily dosing of SELZENTRY.

- 166 The total number of subjects reporting infections were 233 (55%) and 84 (40%) in the 167 group receiving SELZENTRY twice daily and the placebo group, respectively. Correcting for
- 168 the longer duration of exposure on SELZENTRY compared with placebo, the exposure-adjusted
- 169 frequency (rate per 100 subject-years) of these events was 133 for both SELZENTRY twice
- 170 daily and placebo.
- 171 Dizziness or postural dizziness occurred in 8% of subjects on either SELZENTRY or 172 placebo, with 2 subjects (0.5%) on SELZENTRY permanently discontinuing therapy (1 due to 173 syncope, 1 due to orthostatic hypotension) versus 1 subject on placebo (0.5%) permanently 174 discontinuing therapy due to dizziness.
- 175 Treatment-emergent adverse events, regardless of causality, from A4001027 and
- 176 A4001028 are summarized in Table 3. Selected events occurring at $\geq 2\%$ of subjects and at a

177 numerically higher rate in subjects treated with SELZENTRY are included; events that occurred

- 178 at the same or higher rate on placebo are not displayed.
- 179

180 Table 3. Percentage of Subjects With Selected Treatment-Emergent Adverse Events (All

- 181 Causality) (≥2% on SELZENTRY and at a higher rate compared with placebo)
- 182 Studies A4001027 and A4001028 (Pooled Analysis, 48 Weeks)

	SEI	LZENTRY		
	Twice Daily ^a			Placebo
		Exposure-		Exposure-
		adjusted rate		adjusted rate
	N = 426	(per 100 pt-yrs)	N = 426	(per 100 pt-yrs)
	(%)	$PYE = 309^{b}$	(%)	$PYE = 111^{b}$
Eye Disorders				
Conjunctivitis	2	3	1	3
Ocular infections, inflammations,	2	3	1	2
and associated manifestations				
Gastrointestinal Disorders				
Constipation	6	9	3	6
General Disorders and				
Administration Site Conditions				
Pyrexia	13	20	9	17
Pain and discomfort	4	5	3	5
Infections and Infestations				
Upper respiratory tract infection	23	37	13	27
Herpes infection	8	11	4	8
Sinusitis	7	10	3	6

Bronchitis	7	9	5	9
Folliculitis	4	5	2	4
Pneumonia	2	3	5	10
Anogenital warts	2	3	1	3
Influenza	2	3	0.5	1
Otitis media	2	3	0.5	1
Metabolism and Nutrition Disorders				
Appetite disorders	8	11	7	13
Musculoskeletal and Connective				
Tissue Disorders				
Joint-related signs and symptoms	7	10	3	5
Muscle pains	3	4	0.5	1
Neoplasms Benign, Malignant, and				
Unspecified				
Skin neoplasms benign	3	4	1	3
Nervous System Disorders				
Dizziness/postural dizziness	9	13	8	17
Paresthesias and dysesthesias	5	7	3	6
Sensory abnormalities	4	6	1	3
Disturbances in consciousness	4	5	3	6
Peripheral neuropathies	4	5	3	6
Psychiatric Disorders				
Disturbances in initiating and	8	11	5	10
maintaining sleep				
Depressive disorders	4	6	3	5
Anxiety symptoms	4	5	3	7
Renal and Urinary Disorders				
Bladder and urethral symptoms	5	7	1	3
Urinary tract signs and symptoms	3	4	1	3
Respiratory, Thoracic, and				
Mediastinal Disorders				
Coughing and associated symptoms	14	21	5	10
Upper respiratory tract signs and	6	9	3	6
symptoms				
Nasal congestion and inflammations	4	6	3	5
Breathing abnormalities	4	5	2	5
Paranasal sinus disorders	3	4	0.5	1
Skin and Subcutaneous Tissue				
Disorders				
Rash	11	16	5	11

Apocrine and eccrine gland	5	7	4	7.5
disorders				
Pruritus	4	5	2	4
Lipodystrophies	3	5	0.5	1
Erythemas	2	3	1	2
Vascular Disorders				
Vascular hypertensive disorders	3	4	2	4

^a300-mg dose equivalent.

184 $^{b}PYE = Patient-years of exposure.$

185

Laboratory Abnormalities: Table 4 shows the treatment-emergent Grade 3-4 laboratory
 abnormalities that occurred in >2% of subjects receiving SELZENTRY.

188

189Table 4. Maximum Shift in Laboratory Test Values (Without Regard to Baseline)

- 190 Incidence ≥2% of Grade 3-4 Abnormalities (ACTG Criteria) Studies A4001027 and
- 191 **A4001028 (Pooled Analysis, 48 Weeks)**

		SELZENTRY	
		Twice Daily + OBT	Placebo + OBT
Laboratory Parameter		$(N = 421)^{a}$	$(N = 207)^{a}$
Preferred Term	Limit	%	%
Aspartate aminotransferase	>5.0x ULN	4.8	2.9
Alanine aminotransferase	>5.0x ULN	2.6	3.4
Total bilirubin	>5.0x ULN	5.5	5.3
Amylase	>2.0x ULN	5.7	5.8
Lipase	>2.0x ULN	4.9	6.3
Absolute neutrophil count	<750/mm ³	4.3	2.4

^aPercentages based on total subjects evaluated for each laboratory parameter.

193

194Study in Treatment-Naive Subjects: Treatment-Emergent Adverse Events:195Treatment-emergent adverse events, regardless of causality, from Study A4001026, a196double-blind, comparative, controlled study in which 721 treatment-naive subjects received197SELZENTRY 300 mg twice daily (N = 360) or efavirenz (N = 361) in combination with198zidovudine/lamivudine for 96 weeks, are summarized in Table 5. Selected events occurring at199 $\geq 2\%$ of subjects and at a numerically higher rate in subjects treated with SELZENTRY are200included; events that occurred at the same or higher rate on efavirenz are not displayed.201

202 Table 5. Percentage of Subjects With Selected Treatment-Emergent Adverse Events (All

203 Causality) (≥2% on SELZENTRY and at a higher rate compared with efavirenz)

204 Study A4001026 (96 Weeks)

	SELZENTRY	Efavirenz
	300 mg Twice Daily +	600 mg Once Daily +
	Zidovudine/Lamivudine	Zidovudine/Lamivudine
	(N = 360)	(N = 361)
	%	%
Blood and Lymphatic System Disorders		
Anemias NEC	8	5
Neutropenias	4	3
Ear and Labyrinth Disorders		
Ear disorders NEC	3	2
Gastrointestinal Disorders		
Flatulence, bloating, and distention	10	7
Gastrointestinal atonic and hypomotility	9	5
disorders NEC		
Gastrointestinal signs and symptoms	3	2
NEC		
General Disorders and Administration		
Site Conditions		
Body temperature perception	3	1
Infections and Infestations		
Bronchitis	13	9
Herpes infection	7	6
Upper respiratory tract infection	32	30
Bacterial infections NEC	6	3
Herpes zoster/varicella	5	4
Lower respiratory tract and lung	3	2
infections		
Neisseria infections	3	0
Tinea infections	4	3
Viral infections NEC	3	2
Musculoskeletal and Connective Tissue		
Disorders		
Joint-related signs and symptoms	6	5
Nervous System Disorders		
Memory loss (excluding dementia)	3	1
Paresthesias and dysesthesias	4	3

Renal and Urinary Disorders		
Bladder and urethral symptoms	4	3
Reproductive System and Breast		
Disorders		
Erection and ejaculation conditions and	3	2
disorders		
Respiratory, Thoracic, and Mediastinal		
Disorders		
Upper respiratory tract signs and	9	5
symptoms		
Skin and Subcutaneous Disorders		
Acnes	3	2
Alopecias	2	1
Lipodystrophies	4	3
Nail and nail bed conditions (excluding	6	2
infections and infestations)		

205

206 Laboratory Abnormalities:

207

208Table 6. Maximum Shift in Laboratory Test Values (Without Regard to Baseline)

209 Incidence ≥2% of Grade 3-4 Abnormalities (ACTG Criteria) Study A4001026 (96 Weeks)

		SELZENTRY	Efavirenz
		300 mg Twice Daily +	600 mg Once Daily+
		Zidovudine/Lamivudine	Zidovudine/Lamivudine
Laboratory Parameter		$(N = 353)^{a}$	$(N = 350)^{a}$
Preferred Term	Limit	%	%
Aspartate aminotransferase	>5.0 x ULN	4.0	4.0
Alanine aminotransferase	>5.0 x ULN	3.9	4.0
Creatine kinase		3.9	4.8
Amylase	>2.0 x ULN	4.3	6.0
Absolute neutrophil count	<750/mm ³	5.7	4.9
Hemoglobin	<7.0 g/dL	2.9	2.3

 a N = Total number of subjects evaluable for laboratory abnormalities.

211 Percentages based on total subjects evaluated for each laboratory parameter. If the same subject

in a given treatment group had >1 occurrence of the same abnormality, only the most severe is

- counted.
- 214

215Less Common Adverse Events in Clinical Trials: The following adverse events216occurred in <2% of subjects treated with SELZENTRY. These events have been included</td>

- 217 because of their seriousness and either increased frequency on SELZENTRY or are potential
- 218 risks due to the mechanism of action. Events attributed to the patient's underlying HIV infection 219 are not listed.
- 220 Blood and Lymphatic System: Marrow depression and hypoplastic anemia.
- 221 *Cardiac Disorders:* Unstable angina, acute cardiac failure, coronary artery disease,
 222 coronary artery occlusion, myocardial infarction, myocardial ischemia.
- 223 *Hepatobiliary Disorders:* Hepatic cirrhosis, hepatic failure, cholestatic jaundice, 224 portal vein thrombosis, hypertransaminasemia, jaundice.
- 225 *Infections and Infestations:* Endocarditis, infective myositis, viral meningitis,
 226 pneumonia, treponema infections, septic shock, *Clostridium* difficile colitis, meningitis.
- 227 *Musculoskeletal and Connective Tissue Disorders:* Myositis, osteonecrosis,
- rhabdomyolysis, blood CK increased.
- Neoplasms Benign, Malignant, and Unspecified (Including Cysts and Polyps):
 Abdominal neoplasm, anal cancer, basal cell carcinoma, Bowen's disease, cholangiocarcinoma,
- 231 diffuse large B-cell lymphoma, lymphoma, metastases to liver, esophageal carcinoma,
- 232 nasopharyngeal carcinoma, squamous cell carcinoma, squamous cell carcinoma of skin, tongue
- 233 neoplasm (malignant stage unspecified), anaplastic large cell lymphomas T- and null-cell types,
- bile duct neoplasms malignant, endocrine neoplasms malignant and unspecified.
- 235 Nervous System Disorders: Cerebrovascular accident, convulsions and epilepsy,
 236 tremor (excluding congenital), facial palsy, hemianopia, loss of consciousness, visual field
 237 defect.
- 238 6.2 Postmarketing Experience
- The following events have been identified during post-approval use of SELZENTRY.
 Because these reactions are reported voluntarily from a population of unknown size, it is not
- 241 possible to estimate their frequency or establish a causal relationship to exposure to
- 242 SELZENTRY.
- 243
 - Skin and Subcutaneous Tissue Disorders: Stevens-Johnson syndrome.
- 244 7 DRUG INTERACTIONS

245 **7.1** Effect of Concomitant Drugs on the Pharmacokinetics of Maraviroc

Maraviroc is a substrate of CYP3A and Pgp and hence its pharmacokinetics are likely to be modulated by inhibitors and inducers of these enzymes/transporters. Therefore, a dose adjustment may be required when maraviroc is coadministered with those drugs [see Dosage and Administration (2)].

- Concomitant use of maraviroc and St. John's wort (*Hypericum perforatum*) or products
 containing St. John's wort is not recommended. Coadministration of maraviroc with St. John's
 wort is expected to substantially decrease maraviroc concentrations and may result in suboptimal
- 253 levels of maraviroc and lead to loss of virologic response and possible resistance to maraviroc.
- 254 For additional drug interaction information see *Clinical Pharmacology* (12.3).

255 8 USE IN SPECIFIC POPULATIONS

256 8.1 Pregnancy

257 <u>Pregnancy Category B:</u> The incidence of fetal variations and malformations was not 258 increased in embryofetal toxicity studies performed with maraviroc in rats at exposures (AUC) 259 approximately 20-fold higher and in rabbits at approximately 5-fold higher than human 260 exposures at the recommended daily dose (up to 1,000 mg/kg/day in rats and 75 mg/kg/day in 261 rabbits). During the pre- and postnatal development studies in the offspring, development of the 262 offspring, including fertility and reproductive performance, was not affected by the maternal 263 administration of maraviroc.

However, there are no adequate and well-controlled studies in pregnant women. Because animal reproduction studies are not always predictive of human response, SELZENTRY should be used during pregnancy only if clearly needed.

Antiretroviral Pregnancy Registry: To monitor maternal-fetal outcomes of pregnant
 women exposed to SELZENTRY and other antiretroviral agents, an Antiretroviral Pregnancy
 Registry has been established. Physicians are encouraged to register patients by calling 1-800 258-4263.

271 8.3 Nursing Mothers

- 272The Centers for Disease Control and Prevention recommend that HIV-infected273mothers not breastfeed their infants to avoid risking postnatal transmission of HIV
- infection. Studies in lactating rats indicate that maraviroc is extensively secreted into rat milk. It
 is not known whether maraviroc is secreted into human milk. Because of the potential for both
 HIV transmission and serious adverse reactions in nursing infants, mothers should be instructed
 not to breastfeed if they are receiving SELZENTRY.

278 8.4 Pediatric Use

- The pharmacokinetics, safety and efficacy of maraviroc in patients aged <16 years have not been established. Therefore, maraviroc should not be used in this patient population.
- 281 8.5 Geriatric Use

There were insufficient numbers of subjects aged 65 and over in the clinical studies to determine whether they respond differently from younger subjects. In general, caution should be exercised when administering SELZENTRY in elderly patients, also reflecting the greater frequency of decreased hepatic and renal function, of concomitant disease and other drug therapy.

287 8.6 Renal Impairment

- 288 Recommended doses of SELZENTRY for patients with impaired renal function
- 289 (CrCl ≤80 mL/min) are based on the results of a pharmacokinetic study conducted in healthy
- 290 subjects with various degrees of renal impairment. The pharmacokinetics of maraviroc in
- subjects with mild and moderate renal impairment was similar to that in subjects with normal
- renal function [see Clinical Pharmacology (12.3)]. A limited number of subjects with mild and
- 293 moderate renal impairment in the Phase 3 clinical trials (n = 131 and n = 12, respectively)
- received the same dose of SELZENTRY as that administered to subjects with normal renal

function. In these subjects there was no apparent difference in the adverse event profile formaraviroc compared with subjects with normal renal function.

If patients with severe renal impairment or ESRD not receiving a concomitant potent CYP3A inhibitor or inducer experience any symptoms of postural hypotension while taking SELZENTRY 300 mg twice daily, the dose should be reduced to 150 mg twice daily. No studies have been performed in subjects with severe renal impairment or ESRD co-treated with potent CYP3A inhibitors or inducers. Hence, no dose of SELZENTRY can be recommended, and SELZENTRY is contraindicated for these patients *[see Dosage and Administration (2.2)*,

303 *Contraindications (4), Warnings and Precautions (5.2), Clinical Pharmacology (12.3)].*

304 8.7 Hepatic Impairment

305 Maraviroc is principally metabolized by the liver; therefore, caution should be exercised 306 when administering this drug to patients with hepatic impairment, because maraviroc 307 concentrations may be increased. Maraviroc concentrations are higher when SELZENTRY 308 150 mg is administered with a potent CYP3A inhibitor compared with following administration 309 of 300 mg without a CYP3A inhibitor, so patients with moderate hepatic impairment who 310 receive SELZENTRY 150 mg with a potent CYP3A inhibitor should be monitored closely for 311 maraviroc-associated adverse events. Maraviroc has not been studied in subjects with severe 312 hepatic impairment [see Warnings and Precautions (5.1), Clinical Pharmacology (12.3)]. 313 8.8 Gender

Population pharmacokinetic analysis of pooled Phase 1/2a data indicated gender (female: n = 96, 23.2% of the total population) does not affect maraviroc concentrations. Dosage adjustment based on gender is not necessary.

317 **8.9 Race**

318Population pharmacokinetic analysis of pooled Phase 1/2a data indicated exposure was31926.5% higher in Asians (N = 95) as compared with non-Asians (n = 318). However, a study320designed to evaluate pharmacokinetic differences between Caucasians (n = 12) and Singaporeans321(n = 12) showed no difference between these 2 populations. No dose adjustment based on race is322needed.

323 10 OVERDOSAGE

The highest dose administered in clinical studies was 1,200 mg. The dose-limiting adverse event was postural hypotension, which was observed at 600 mg. While the recommended dose for SELZENTRY in patients receiving a CYP3A inducer without a CYP3A inhibitor is 600 mg twice daily, this dose is appropriate due to enhanced metabolism.

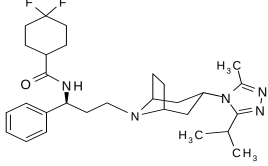
Prolongation of the QT interval was seen in dogs and monkeys at plasma concentrations 6 and 12 times, respectively, those expected in humans at the intended exposure of 300 mg equivalents twice daily. However, no significant QT prolongation was seen in the studies in treatment-experienced subjects with HIV using the recommended doses of maraviroc or in a specific pharmacokinetic study to evaluate the potential of maraviroc to prolong the QT interval

333 [see Clinical Pharmacology (12.3)].

- There is no specific antidote for overdose with maraviroc. Treatment of overdose should consist of general supportive measures including keeping the patient in a supine position, careful assessment of patient vital signs, blood pressure, and ECG.
- 337 If indicated, elimination of unabsorbed active maraviroc should be achieved by emesis or
- 338 gastric lavage. Administration of activated charcoal may also be used to aid in removal of
- 339 unabsorbed drug. Since maraviroc is moderately protein-bound, dialysis may be beneficial in
- 340 removal of this medicine.

341 **11 DESCRIPTION**

- SELZENTRY (maraviroc) is a selective, slowly reversible, small molecule antagonist of
 the interaction between human CCR5 and HIV-1 gp120. Blocking this interaction prevents
 CCR5-tropic HIV-1 entry into cells.
- 345 SELZENTRY is available as film-coated tablets for oral administration containing either
- 346 150 or 300 mg of maraviroc and the following inactive ingredients: dibasic calcium phosphate
- 347 (anhydrous), magnesium stearate, microcrystalline cellulose, and sodium starch glycolate. The
- 348 film coat (Opadry[®] II Blue [85G20583]) contains FD&C blue #2 aluminum lake, soya lecithin,
- 349 polyethylene glycol (macrogol 3350), polyvinyl alcohol, talc, and titanium dioxide.
- 350 Maraviroc is chemically described as 4,4-difluoro-*N*-{(1*S*)-3-[*exo*-3-(3-isopropyl-5-
- 351 methyl-4*H*-1,2,4-triazol-4-yl)-8-azabicyclo[3.2.1]oct-8-yl]-1-
- 352 phenylpropyl}cyclohexanecarboxamide.
- 353 The molecular formula is $C_{29}H_{41}F_2N_5O$ and the structural formula is:



354

Maraviroc is a white to pale-colored powder with a molecular weight of 513.67. It is highly soluble across the physiological pH range (pH 1.0 to 7.5).

357 12 CLINICAL PHARMACOLOGY

358 **12.1 Mechanism of Action**

359 Maraviroc is an antiviral drug [see Clinical Pharmacology (12.4)].

360 **12.2 Pharmacodynamics**

- 361 Exposure-Response Relationship in Treatment-Experienced Subjects: The
- 362 relationship between maraviroc, modeled plasma trough concentration (C_{min}) (1 to 9 samples per
- 363 patient taken on up to 7 visits), and virologic response was evaluated in
- 364 973 treatment-experienced HIV-1-infected subjects with varied optimized background
- antiretroviral regimens in Studies A4001027 and A4001028. The C_{min} , baseline viral load,

- 366 baseline CD4+ cell count, and overall sensitivity score (OSS) were found to be important
- 367 predictors of virologic success (defined as viral load <400 copies/mL at 24 weeks). Table 7
- illustrates the proportions of subjects with virologic success (%) within each C_{min} quartile for 368
- 369 150-mg twice-daily and 300-mg twice-daily groups.
- 370

				°8-° ~		
		150 mg	Twice Daily		300 mg	g Twice Daily
		(With CYF	P3A Inhibitors)	(Without CYP3A Inhibitors)		
		Median	% Subjects With		Median	% Subjects With
	n	C _{min}	Virologic Success	n	C _{min}	Virologic Success
Placebo	160	-	30.6	35	_	28.6
Q1	78	33	52.6	22	13	50.0
Q2	77	87	63.6	22	29	68.2
Q3	78	166	78.2	22	46	63.6
Q4	78	279	74.4	22	97	68.2

371 Table 7. Treatment-Experienced Subjects With Virologic Success by C_{min} Quartile (Q1-Q4)

372

- 373 Exposure-Response Relationship in Treatment-Naive Subjects: The relationship 374 between maraviroc, modeled plasma trough concentration (C_{min}) (1 to 12 samples per patient 375 taken on up to 8 visits), and virologic response was evaluated in 294 treatment-naive HIV-1-376 infected subjects receiving maraviroc 300 mg twice daily in combination with 377
- zidovudine/lamivudine in Study A4001026. Table 8 illustrates the proportion (%) of subjects 378 with virologic success <50 copies/mL at 48 weeks within each C_{min} quartile for the 300-mg
- 379 twice-daily dose.
- 380

381 Table 8. Treatment-Naive Subjects With Virologic Success by C_{min} Quartile (Q1-Q4)

		300 mg Twice Daily				
	n	Median C _{min}	% Subjects With Virologic Success			
Q1	75	23	57.3			
Q2	72	39	72.2			
Q3	73	56	74.0			
Q4	74	81	83.8			

382

383

Eighteen of 75 (24%) subjects in Q1 had no measurable maraviroc concentration on at 384 least one occasion vs. 1 of 73 and 1 of 74 in Q3 and Q4, respectively.

385 Effects on Electrocardiogram: A placebo-controlled, randomized, crossover study to 386 evaluate the effect on the QT interval of healthy male and female volunteers was conducted with 387 3 single oral doses of maraviroc and moxifloxacin. The placebo-adjusted mean maximum (upper 388 1-sided 95% CI) increases in QTc from baseline after 100, 300, and 900 mg of maraviroc were 389 -2 (0), -1 (1), and 1 (3) msec, respectively, and 13 (15) msec for moxifloxacin 400 mg. No

- 390 subject in any group had an increase in QTc of ≥60 msec from baseline. No subject experienced
- an interval exceeding the potentially clinically relevant threshold of 500 msec.
- 392 **12.3 Pharmacokinetics**
- 393

			AUC ₁₂	C _{max}	C_{min}
Patient Population	Maraviroc Dose	Ν	(ng.hr/mL)	(ng/mL)	(ng/mL)
Healthy volunteers (Phase 1)	300 mg twice daily	64	2,908	888	43.1
Asymptomatic HIV subjects (Phase 2a)	300 mg twice daily	8	2,550	618	33.6
Treatment-experienced	300 mg twice daily	94	1,513	266	37.2
HIV subjects (Phase 3) ^a	150 mg twice daily (+ CYP3A inhibitor)	375	2,463	332	101
Treatment-naive HIV subjects (Phase 2b/3) ^a	300 mg twice daily	344	1,865	287	60

394 **Table 9. Mean Maraviroc Pharmacokinetic Parameters**

^a The estimated exposure is lower compared with other studies possibly due to sparse sampling,
 food effect, compliance, and concomitant medications.

397

<u>Absorption:</u> Peak maraviroc plasma concentrations are attained 0.5 to 4 hours following
 single oral doses of 1 to 1,200 mg administered to uninfected volunteers. The pharmacokinetics
 of oral maraviroc are not dose proportional over the dose range.

401 The absolute bioavailability of a 100-mg dose is 23% and is predicted to be 33% at
402 300 mg. Maraviroc is a substrate for the efflux transporter P-glycoprotein.

403 <u>Effect of Food on Oral Absorption:</u> Coadministration of a 300-mg tablet with a 404 high-fat breakfast reduced maraviroc C_{max} and AUC by 33% in healthy volunteers. There were 405 no food restrictions in the studies that demonstrated the efficacy and safety of maraviroc [see 406 *Clinical Studies (14)*]. Therefore, maraviroc can be taken with or without food at the 407 recommended dose [see Dosage and Administration (2)].

408 <u>Distribution:</u> Maraviroc is bound (approximately 76%) to human plasma proteins, and 409 shows moderate affinity for albumin and alpha-1 acid glycoprotein. The volume of distribution 410 of maraviroc is approximately 194 L.

411 <u>Metabolism:</u> Studies in humans and in vitro studies using human liver microsomes and 412 expressed enzymes have demonstrated that maraviroc is principally metabolized by the

413 cytochrome P450 system to metabolites that are essentially inactive against HIV-1. In vitro

414 studies indicate that CYP3A is the major enzyme responsible for maraviroc metabolism. In vitro

- studies also indicate that polymorphic enzymes CYP2C9, CYP2D6, and CYP2C19 do not
- 416 contribute significantly to the metabolism of maraviroc.
- 417 Maraviroc is the major circulating component (~42% drug-related radioactivity)
- 418 following a single oral dose of $300 \text{ mg} [^{14}\text{C}]$ -maraviroc. The most significant circulating
- 419 metabolite in humans is a secondary amine (~22% radioactivity) formed by N-dealkylation. This

polar metabolite has no significant pharmacological activity. Other metabolites are products of
 mono-oxidation and are only minor components of plasma drug-related radioactivity.

422 <u>Excretion:</u> The terminal half-life of maraviroc following oral dosing to steady state in 423 healthy subjects was 14 to 18 hours. A mass balance/excretion study was conducted using a 424 single 300-mg dose of ¹⁴C-labeled maraviroc. Approximately 20% of the radiolabel was 425 recovered in the urine and 76% was recovered in the feces over 168 hours. Maraviroc was the 426 major component present in urine (mean of 8% dose) and feces (mean of 25% dose). The 427 remainder was excreted as metabolites.

428 Hepatic Impairment: Maraviroc is primarily metabolized and eliminated by the liver. A 429 study compared the pharmacokinetics of a single 300-mg dose of SELZENTRY in subjects with 430 mild (Child-Pugh Class A, n = 8), and moderate (Child-Pugh Class B, n = 8) hepatic impairment to pharmacokinetics in healthy subjects (n = 8). The mean C_{max} and AUC were 11% and 25% 431 432 higher, respectively, for subjects with mild hepatic impairment, and 32% and 46% higher, 433 respectively, for subjects with moderate hepatic impairment compared with subjects with normal 434 hepatic function. These changes do not warrant a dose adjustment. Maraviroc concentrations are 435 higher when SELZENTRY 150 mg is administered with a potent CYP3A inhibitor compared 436 with following administration of 300 mg without a CYP3A inhibitor, so patients with moderate 437 hepatic impairment who receive SELZENTRY 150 mg with a potent CYP3A inhibitor should be 438 monitored closely for maraviroc-associated adverse events. The pharmacokinetics of maraviroc 439 have not been studied in subjects with severe hepatic impairment *[see Warnings and Precautions*]

440 *(5.1)]*.

441 Renal Impairment: A study compared the pharmacokinetics of a single 300-mg dose of 442 SELZENTRY in subjects with severe renal impairment (CLcr <30 mL/min, n = 6) and ESRD 443 (n = 6) to healthy volunteers (n = 6). Geometric mean ratios for maraviroc C_{max} and AUC_{inf} were 444 2.4-fold and 3.2-fold higher, respectively, for subjects with severe renal impairment, and 1.7-fold 445 and 2.0-fold higher, respectively, for subjects with ESRD as compared with subjects with normal 446 renal function in this study. Hemodialysis had a minimal effect on maraviroc clearance and 447 exposure in subjects with ESRD. Exposures observed in subjects with severe renal impairment 448 and ESRD were within the range observed in previous 300-mg single-dose studies of 449 SELZENTRY in healthy volunteers with normal renal function. However, maraviroc exposures 450 in the subjects with normal renal function in this study were 50% lower than that observed in 451 previous studies. Based on the results of this study, no dose adjustment is recommended for 452 patients with renal impairment receiving SELZENTRY without a potent CYP3A inhibitor or 453 inducer. However, if patients with severe renal impairment or ESRD experience any symptoms 454 of postural hypotension while taking SELZENTRY 300 mg twice daily, their dose should be 455 reduced to 150 mg twice daily [see Dosage and Administration (2.2); Warnings and Precautions 456 (5.2)]. 457 In addition, the study compared the pharmacokinetics of multiple-dose SELZENTRY in

458 combination with saquinavir/ritonavir 1,000/100 mg twice daily (a potent CYP3A inhibitor 450 1000 1

459 combination) for 7 days in subjects with mild renal impairment (CLcr >50 and \leq 80 mL/min,

- 460 n = 6) and moderate renal impairment (CLcr \ge 30 and \le 50 mL/min, n = 6) to healthy volunteers
- 461 with normal renal function (n = 6). Subjects received 150 mg of SELZENTRY at different dose
- 462 frequencies (healthy volunteers every 12 hours; mild renal impairment every 24 hours;
- 463 moderate renal impairment every 48 hours). Compared with healthy volunteers (dosed every
- 464 12 hours), geometric mean ratios for maraviroc AUC_{tau}, C_{max} , and C_{min} were 50% higher, 20%
- 465 higher, and 43% lower, respectively, for subjects with mild renal impairment (dosed every
- 466 24 hours). Geometric mean ratios for maraviroc AUC_{tau} , C_{max} , and C_{min} were 16% higher, 29%
- lower, and 85% lower, respectively, for subjects with moderate renal impairment (dosed every
- 468 48 hours) compared with healthy volunteers (dosed every 12 hours). Based on the data from this
- study, no adjustment in dose is recommended for patients with mild or moderate renal
 impairment [see Dosage and Administration (2.2)].
- 471 <u>Effect of Concomitant Drugs on the Pharmacokinetics of Maraviroc:</u> Maraviroc is a
 472 substrate of CYP3A and Pgp and hence its pharmacokinetics are likely to be modulated by
 473 inhibitors and inducers of these enzymes/transporters. The CYP3A/Pgp inhibitors ketoconazole,
 474 lopinavir/ritonavir, ritonavir, darunavir/ritonavir, saquinavir/ritonavir, and atazanavir ± ritonavir
 475 all increased the C_{max} and AUC of maraviroc (see Table 10). The CYP3A inducers rifampin,
- 476 etravirine, and efavirenz decreased the C_{max} and AUC of maraviroc (see Table 10).
- 477 Tipranavir/ritonavir (net CYP3A inhibitor/Pgp inducer) did not affect the steady-state
 478 pharmacokinetics of maraviroc (see Table 10). Cotrimoxazole and tenofovir did not affect the
 479 pharmacokinetics of maraviroc.

480

			Ratio (90% CI) of Maraviroc Pharmacokinetic Parameters With/Without Coadministered Drug			
Coadministered Drug		Dose of		(No Effect = 1.00)		
and Dose	Ν	SELZENTRY	C _{min}	AUC _{tau}	C _{max}	
CYP3A and/or P-gp Inhi		1				
Ketoconazole	12	100 mg b.i.d.	3.75	5.00	3.38	
400 mg q.d.			(3.01, 4.69)	(3.98, 6.29)	(2.38, 4.78)	
Ritonavir	8	100 mg b.i.d.	4.55	2.61	1.28	
100 mg b.i.d.			(3.37, 6.13)	(1.92, 3.56)	(0.79, 2.09)	
Saquinavir (soft gel	11	100 mg b.i.d.	11.3	9.77	4.78	
capsules) /ritonavir 1,000 mg/100 mg b.i.d.			(8.96, 14.1)	(7.87, 12.14)	(3.41, 6.71)	
Lopinavir/ritonavir	11	300 mg b.i.d.	9.24	3.95	1.97	
400 mg/100 mg b.i.d.			(7.98, 10.7)	(3.43, 4.56)	(1.66, 2.34)	
Atazanavir	12	300 mg b.i.d.	4.19	3.57	2.09	
400 mg q.d.			(3.65, 4.80)	(3.30, 3.87)	(1.72, 2.55)	
Atazanavir/ritonavir	12	300 mg b.i.d.	6.67	4.88	2.67	
300 mg/100 mg q.d.			(5.78, 7.70)	(4.40, 5.41)	(2.32, 3.08)	
Darunavir/ritonavir	12	150 mg b.i.d.	8.00	4.05	2.29	
600 mg/100 mg b.i.d.		-	(6.35, 10.1)	2.94, 5.59	(1.46, 3.59)	
CYP3A and/or P-gp Indu	icers					
Efavirenz	12	100 mg b.i.d.	0.55	0.552	0.486	
600 mg q.d.			(0.43, 0.72)	(0.492, 0.620)	(0.377, 0.626)	
Efavirenz	12	200 mg b.i.d.	1.09	1.15	1.16	
600 mg q.d.		(+efavirenz):	(0.89, 1.35)	(0.98, 1.35)	(0.87, 1.55)	
		100 mg b.i.d.				
		(alone)				
Rifampicin	12	100 mg b.i.d.	0.22	0.368	0.335	
600 mg q.d.		-	(0.17, 0.28)	(0.328, 0.413)	(0.260, 0.431	
Rifampicin	12	200 mg b.i.d.	0.66	1.04	0.97	
600 mg q.d.		(+rifampicin):	(0.54, 0.82)	(0.89, 1.22)	(0.72, 1.29)	
		100 mg b.i.d.				
		(alone)				
Etravirine	14	300 mg b.i.d.	0.609	0.468	0.400	
200 mg b.i.d.			(0.525, 0.707)	(0.381, 0.576)	(0.282, 0.566)	
Nevirapine ^a	8	300 mg SD	-	1.01	1.54	
200 mg b.i.d.				(0.65, 1.55)	(0.94, 2.51)	
(+ lamivudine 150 mg b.i.d	1.,					
tenofovir 300 mg q.d.)						

481 Table 10. Effect of Coadministered Agents on the Pharmacokinetics of Maraviroc

Lopinavir/ritonavir +	11	300 mg b.i.d.	6.29	2.53	1.25
efavirenz 400 mg/100 mg		_	(4.72, 8.39)	(2.24, 2.87)	(1.01, 1.55)
b.i.d. + 600 mg q.d.					
Saquinavir(soft gel	11	100 mg b.i.d.	8.42	5.00	2.26
capsules) /ritonavir +			(6.46, 10.97)	(4.26, 5.87)	(1.64, 3.11)
efavirenz					
1,000 mg/100 mg b.i.d. +					
600 mg q.d.					
Darunavir/ritonavir +	10	150 mg b.i.d.	5.27	3.10	1.77
etravirine			(4.51, 6.15)	(2.57, 3.74)	(1.20, 2.60)
600 mg/100 mg b.i.d. +			(4.51, 0.15)	(2.57, 5.77)	
200 mg b.i.d.					
Tipranavir/ritonavir	12	150 mg b.i.d.	1.80	1.02	0.86
500 mg/200 mg b.i.d.			(1.55, 2.09)	(0.850, 1.23)	(0.61, 1.21)
Other					
Raltegravir	17	300 mg b.i.d.	0.90	0.86	0.79
400 mg b.i.d.			(0.85, 0.96)	(0.80, 0.92)	(0.67, 0.94)

482 ^aCompared with historical data.

- 483
- <u>Effect of Maraviroc on the Pharmacokinetics of Concomitant Drugs:</u> Maraviroc is
 unlikely to inhibit the metabolism of coadministered drugs metabolized by the following
 cytochrome P enzymes (CYP1A2, CYP2B6, CYP2C8, CYP2C9, CYP2C19, and CYP3A)
 because maraviroc did not inhibit activity of those enzymes at clinically relevant concentrations
 in vitro. Maraviroc does not induce CYP1A2 in vitro.
- 100 11 1

In vitro results indicate that maraviroc could inhibit P-glycoprotein in the gut and maythus affect bioavailability of certain drugs.

491Drug interaction studies were performed with maraviroc and other drugs likely to be492coadministered or commonly used as probes for pharmacokinetic interactions (see Table 6).

493 Maraviroc had no effect on the pharmacokinetics of zidovudine or lamivudine. Maraviroc

decreased the C_{min} and AUC of raltegravir by 27% and 37%, respectively, which is not clinically

significant. Maraviroc had no clinically relevant effect on the pharmacokinetics of midazolam,

496 the oral contraceptives ethinylestradiol and levonorgestrel, no effect on the urinary 6β -

- 497 hydroxycortisol/cortisol ratio, suggesting no induction of CYP3A in vivo. Maraviroc had no
- 498 effect on the debrisoquine metabolic ratio (MR) at 300 mg twice daily or less in vivo and did not
- 499 cause inhibition of CYP2D6 in vitro until concentrations >100 μ M. However, there was 234%
- 500 increase in debrisoquine MR on treatment compared with baseline at 600 mg once daily,
- 501 suggesting potential inhibition of CYP2D6 at higher dose.
- 502 **12.4 Microbiology**

503 <u>Mechanism of Action:</u> Maraviroc is a member of a therapeutic class called CCR5 co-504 receptor antagonists. Maraviroc selectively binds to the human chemokine receptor CCR5 505 present on the cell membrane, preventing the interaction of HIV-1 gp120 and CCR5 necessary for CCR5-tropic HIV-1 to enter cells. CXCR4-tropic and dual-tropic HIV-1 entry is not inhibitedby maraviroc.

508 <u>Antiviral Activity in Cell Culture:</u> Maraviroc inhibits the replication of CCR5-tropic 509 laboratory strains and primary isolates of HIV-1 in models of acute peripheral blood leukocyte 510 infection. The mean EC₅₀ value (50% effective concentration) for maraviroc against HIV-1 511 group M isolates (subtypes A to J and circulating recombinant form AE) and group O isolates

512 ranged from 0.1 to 4.5 nM (0.05 to 2.3 ng/mL) in cell culture.

513 When used with other antiretroviral agents in cell culture, the combination of maraviroc 514 was not antagonistic with NNRTIs (delavirdine, efavirenz, and nevirapine), NRTIs (abacavir, 515 didanosine, emtricitabine, lamivudine, stavudine, tenofovir, zalcitabine, and zidovudine), or 516 protease inhibitors (amprenavir, atazanavir, darunavir, indinavir, lopinavir, nelfinavir, ritonavir, 517 saquinavir, and tipranavir). Maraviroc was additive/synergistic with the HIV fusion inhibitor 518 enfuvirtide. Maraviroc was not active against CXCR4-tropic and dual-tropic viruses (EC₅₀ value

519 >10 μM). The antiviral activity of maraviroc against HIV-2 has not been evaluated.
 520 *Resistance in Cell Culture:* HIV-1 variants with reduced susceptibility to maraviroc
 521 have been selected in cell culture, following serial passage of 2 CCR5-tropic viruses (CC1/85

and RU570). The maraviroc-resistant viruses remained CCR5-tropic with no evidence of a

523 change from a CCR5-tropic virus to a CXCR4-using virus. Two amino acid residue substitutions

in the V3-loop region of the HIV-1 envelope glycoprotein (gp160), A316T, and I323V (HXB2

525 numbering), were shown to be necessary for the maraviroc-resistant phenotype in the HIV-1 526 isolate CC1/85. In the RU570 isolate a 3-amino acid residue deletion in the V3 loop, Δ QAI

(HXB2 positions 315 to 317), was associated with maraviroc resistance. The relevance of the
 specific gp120 mutations observed in maraviroc-resistant isolates selected in cell culture to

clinical maraviroc resistance is not known. Maraviroc-resistant viruses were characterized
 phenotypically by concentration-response curves that did not reach 100% inhibition in

531 phenotypic drug assays, rather than increases in EC_{50} values.

532 *Cross-Resistance in Cell Culture:* Maraviroc had antiviral activity against HIV-1 533 clinical isolates resistant to NNRTIs, NRTIs, PIs, and the fusion inhibitor enfuvirtide in cell 534 culture (EC_{50} values ranged from 0.7 to 8.9 nM (0.36 to 4.57 ng/mL). Maraviroc-resistant viruses 535 that emerged in cell culture remained susceptible to the enfuvirtide and the protease inhibitor 536 saquinavir.

537 *Clinical Resistance:* Virologic failure on maraviroc can result from genotypic and 538 phenotypic resistance to maraviroc, through outgrowth of undetected CXCR4-using virus present 539 before maraviroc treatment (see *Tropism* below), through resistance to background therapy drugs 540 (Table 11), or due to low exposure to maraviroc [see Clinical Pharmacology (12.2)].

541 Antiretroviral Treatment-Experienced Subjects (Studies A4001027 and 542 A4001028): Week 48 data from treatment-experienced subjects failing maraviroc-containing 543 regimens with CCR5-tropic virus (n = 58) have identified 22 viruses that had decreased 544 susceptibility to maraviroc characterized in phenotypic drug assays by concentration-response

- 545 curves that did not reach 100% inhibition. Additionally, CCR5-tropic virus from 2 of these 546 treatment-failure subjects had \geq 3-fold shifts in EC₅₀ values for maraviroc at the time of failure.
- 547 Fifteen of these viruses were sequenced in the gp120 encoding region and multiple amino
- 548 acid substitutions with unique patterns in the heterogeneous V3 loop region were detected.
- 549 Changes at either amino acid position 308 or 323 (HXB2 numbering) were seen in the V3 loop
- 549 in 7 of the subjects with decreased maraviroc susceptibility. Substitutions outside the V3 loop of
- 551 gp120 may also contribute to reduced susceptibility to maraviroc.
- 552 Antiretroviral Treatment-Naive Subjects (Study A4001026): Treatment-naive 553 subjects receiving SELZENTRY had more virologic failures and more treatment-emergent 554 resistance to the background regimen drugs compared with those receiving efavirenz (Table 11).
- 555
- 556 **Table 11. Development of Resistance to Maraviroc or Efavirenz and Background Drugs in**
- 557 Antiretroviral Treatment-Naive Trial A4001026 for Patients with CCR5-Tropic Virus at

	Maraviroc	Efavirenz	
Total N in dataset (as-treated)	273	241	
Total virologic failures (as-treated)	85(31%)	56 (23%)	
Evaluable virologic failures with post baseline genotypic and phenotypic data	73	43	
Lamivudine resistance	39 (53%)	13 (30%)	
Zidovudine resistance	2 (3%)	0	
Efavirenz resistance		23 (53%)	
• Phenotypic resistance to maraviroc ^a	19 (26 %)		

558 Screening Using Enhanced Sensitivity TROFILE[®] Assay

- ^a Includes subjects failing with CXCR4- or dual/mixed-tropism because these viruses are not
 intrinsically susceptible to maraviroc.
- 561

In an as-treated analysis of treatment-naive subjects at 96 weeks, 32 subjects failed a
 maraviroc-containing regimen with CCR5-tropic virus and had a tropism result at failure; 7 of
 these subjects had evidence of maraviroc phenotypic resistance defined as

565 concentration-response curves that did not reach 95% inhibition. One additional subject had a

- \geq 3-fold shift in the EC₅₀ value for maraviroc at the time of failure. A clonal analysis of the V3
- loop amino acid envelope sequences was performed from 6 of the 7 subjects. Changes in V3
- loop amino acid sequence differed between each of these different subjects, even for those
- infected with the same virus clade suggesting that that there are multiple diverse pathways to
- 570 maraviroc resistance. The subjects who failed with CCR5-tropic virus and without a detectable
- 571 maraviroc shift in susceptibility were not evaluated for genotypic resistance.
- 572 Of the 32 maraviroc virologic failures failing with CCR5-tropic virus, 20(63%) also had 573 genotypic and/or phenotypic resistance to background drugs in the regimen (lamivudine, 574 zidovudine).
- 575

Tropism: In both treatment-experienced and treatment-naive subjects, detection of

576 CXCR4-using virus prior to initiation of therapy has been associated with a reduced virologic 577 response to maraviroc.

578 Antiretroviral Treatment-Experienced Subjects: In the majority of cases, treatment 579 failure on maraviroc was associated with detection of CXCR4-using virus (i.e., CXCR4- or 580 dual/mixed-tropic) which was not detected by the tropism assay prior to treatment. 581 CXCR4-using virus was detected at failure in approximately 55% of subjects who failed 582 treatment on maraviroc by Week 48, as compared with 9% of subjects who experienced 583 treatment failure in the placebo arm. To investigate the likely origin of the on-treatment 584 CXCR4-using virus, a detailed clonal analysis was conducted on virus from 20 representative 585 subjects (16 subjects from the maraviroc arms and 4 subjects from the placebo arm) in whom 586 CXCR4-using virus was detected at treatment failure. From analysis of amino acid sequence 587 differences and phylogenetic data, it was determined that CXCR4-using virus in these subjects 588 emerged from a low level of pre-existing CXCR4-using virus not detected by the tropism assay 589 (which is population-based) prior to treatment rather than from a coreceptor switch from 590 CCR5-tropic virus to CXCR4-using virus resulting from mutation in the virus. 591 Detection of CXCR4-using virus prior to initiation of therapy has been associated with a 592 reduced virological response to maraviroc. Furthermore, subjects failing maraviroc twice daily at 593 Week 48 with CXCR4-using virus had a lower median increase in CD4+ cell counts from baseline (+41 cells/mm³) than those subjects failing with CCR5-tropic virus (+162 cells/mm³). 594 595 The median increase in CD4+ cell count in subjects failing in the placebo arm was +7 cells/mm³. 596 Antiretroviral Treatment-Naive Subjects: In a 96-week study of antiretroviral 597 treatment-naive subjects, 14% (12/85) who had CCR5-tropic virus at screening with an enhanced 598 sensitivity tropism assay (TROFILE) and failed therapy on maraviroc had CXCR4-using virus at 599 the time of treatment failure. A detailed clonal analysis was conducted in 2 previously 600 antiretroviral treatment-naive subjects enrolled in a Phase 2a monotherapy study who had 601 CXCR4-using virus detected after 10 days treatment with maraviroc. Consistent with the detailed 602 clonal analysis conducted in treatment-experienced subjects, the CXCR4-using variants appear 603 to emerge from outgrowth of a pre-existing undetected CXCR4-using virus. Screening with an 604 enhanced sensitivity tropism assay reduced the number of maraviroc virologic failures with 605 CXCR4- or dual/mixed-tropic virus at failure to 12 compared with 24 when screening with the 606 original tropism assay. All but one (11/12; 92%) of the maraviroc failures failing with CXCR4-607 or dual/mixed-tropic virus also had genotypic and phenotypic resistance to the background drug 608 lamivudine at failure and 33% (4/12) developed zidovudine-associated resistance substitutions. 609 Subjects who had CCR5-tropic virus at baseline and failed maraviroc therapy with

- 610 CXCR4-using virus had a median increase in CD4+ cell counts from baseline of +113 cells/mm³
- 611 while those subjects failing with CCR5-tropic virus had an increase of +135 cells/mm³. The
- 612 median increase in CD4+ cell count in subjects failing in the efavirenz arm was + 95 cells/mm³.
- 613 13 NONCLINICAL TOXICOLOGY

614 13.1 Carcinogenesis, Mutagenesis, Impairment of Fertility

615 <u>Carcinogenesis:</u> Long-term oral carcinogenicity studies of maraviroc were carried out

- 616 in rasH2 transgenic mice (6 months) and in rats for up to 96 weeks (females) and 104 weeks
- 617 (males). No drug-related increases in tumor incidence were found in mice at 1,500 mg/kg/day
- and in male and female rats at 900 mg/kg/day. The highest exposures in rats were approximately
- 619 11 times those observed in humans at the therapeutic dose of 300 mg twice daily for the
- 620 treatment of HIV-1 infection.
- 621 <u>Mutagenesis:</u> Maraviroc was not genotoxic in the reverse mutation bacterial test (Ames 622 test in *Salmonella* and *E. coli*), a chromosome aberration test in human lymphocytes, and rat 623 bone marrow micronucleus test.
- 624 <u>Impairment of Fertility:</u> Maraviroc did not impair mating or fertility of male or female 625 rats and did not affect sperm of treated male rats at approximately 20-fold higher exposures 626 (AUC) than in humans given the recommended 300-mg twice-daily dose.

627 14 CLINICAL STUDIES

The clinical efficacy and safety of SELZENTRY is derived from analyses of data from 3 ongoing studies in adult subjects infected with CCR5-tropic HIV-1: A4001027 and A4001028 in antiretroviral treatment-experienced adult subjects and A4001026 in treatment-naive subjects. These studies are supported by a 48-week study in antiretroviral treatment-experienced adult subjects infected with dual/mixed-tropic HIV-1, A4001029.

633 14.1 Studies in CCR5-Tropic, Treatment-Experienced Subjects

634 Studies A4001027 and A4001028 are ongoing, double-blind, randomized,

placebo-controlled, multicenter studies in subjects infected with CCR5-tropic HIV-1. Subjects
 were required to have an HIV-1 RNA of greater than 5,000 copies/mL despite at least 6 months

637 of prior therapy with at least 1 agent from 3 of the 4 antiretroviral drug classes (\geq 1 NRTI,

 ≥ 1 NNRTI, ≥ 2 PIs, and/or enfuvirtide) or documented resistance to at least 1 member of each

639 class. All subjects received an optimized background regimen consisting of 3 to 6 antiretroviral

- agents (excluding low-dose ritonavir) selected on the basis of the subject's prior treatment
- history and baseline genotypic and phenotypic viral resistance measurements. In addition to the

642 optimized background regimen, subjects were then randomized in a 2:2:1 ratio to SELZENTRY

643 300 mg once daily, SELZENTRY 300 mg twice daily, or placebo. Doses were adjusted based on
644 background therapy as described in *Dosing and Administration*, Table 1.

In the pooled analysis for A4001027 and A4001028, the demographics and baseline
characteristics of the treatment groups were comparable (Table 12). Of the 1,043 subjects with a
CCR5-tropism result at screening, 7.6% had a dual/mixed-tropism result at the baseline visit 4 to
6 weeks later. This illustrates the background change from CCR5- to dual/mixed-tropism result
over time in this treatment-experienced population, prior to a change in antiretroviral regimen or

administration of a CCR5 co-receptor antagonist.

651

652Table 12. Demographic and Baseline Characteristics of Subjects in Studies A4001027 and

653 **A4001028**

A4001020		
	SELZENTRY	
	Twice Daily	Placebo
	(N = 426)	(N = 209)
Age (years)		
Mean (range)	46.3 (21-73)	45.7 (29-72)
Sex		
Male	382 (89.7%)	185 (88.5%)
Female	44 (10.3%)	24 (11.5%)
Race		
White	363 (85.2%)	178 (85.2%)
Black	51 (12.0%)	26 (12.4%)
Other	12 (2.8%)	5 (2.4%)
Region		
U.S.	276 (64.8%)	135 (64.6%)
Non-U.S.	150 (35.2%)	74 (35.4%)
Subjects with previous enfuvirtide use	142 (33.3%)	62 (29.7%)
Subjects with enfuvirtide as part of OBT	182 (42.7%)	91 (43.5%)
Baseline plasma HIV-1 RNA (log ₁₀ copies/mL)	1 95 (2 06 6 99)	1 96 (2 46 7 07)
Mean (range)	4.85 (2.96-6.88)	4.86 (3.46-7.07)
Subjects with screening viral load \geq 100,000 copies/mL	179 (42.0%)	84 (40.2%)
Baseline CD4+ cell count (cells/mm ³)	167 (2-820)	171 (1-675)
Median (range)		
Subjects with baseline CD4+ cell count $\leq 200 \text{ cells/mm}^3$)	250 (58.7%)	118 (56.5%)
Subjects with Overall Susceptibility Score (OSS): ^a		
0	57 (13.4%)	35 (16.7%)
1	136 (31.9%)	44 (21.1%)
2	104 (24.4%)	59 (28.2%)
≥3	125 (29.3%)	66 (31.6%)
Subjects with enfuvirtide resistance mutations	90 (21.2%)	45 (21.5%)
Median number of resistance-associated: ^b		
PI mutations	10	10
NNRTI mutations	1	1
NRTI mutations	6	6

^a OSS - Sum of active drugs in OBT based on combined information from genotypic and

655 phenotypic testing.

656 ^b Resistance mutations based on IAS guidelines.¹

657

The Week 48 results for the pooled Studies A4001027 and A4001028 are shown in

659 Table 13.

660

661 Table 13. Outcomes of Randomized Treatment at Week 48

662 Studies A4001027 and A4001028

	SELZENTRY		
	Twice Daily	Placebo	Mean
Outcome	(N = 426)	(N = 209)	Difference
Mean change from Baseline to Week 48 in	-1.84	-0.78	-1.05
HIV-1 RNA (log ₁₀ copies/mL)			
<400 copies/mL at Week 48	239 (56%)	47 (22%)	34%
<50 copies/mL at Week 48	194 (46%)	35 (17%)	29%
Discontinuations			
Insufficient clinical response	97 (23%)	113 (54%)	
Adverse events	19 (4%)	11 (5%)	
Other	27 (6%)	18 (9%)	
Subjects with treatment-emergent CDC Category C events	22 (5%)	16 (8%)	
Deaths (during study or within 28 days of last dose)	9 (2%) ^a	1 (0.5%)	

^a One additional subject died while receiving open-label therapy with SELZENTRY subsequent
 to discontinuing double-blind placebo due to insufficient response.

665

666 After 48 weeks of therapy, the proportions of subjects with HIV-1 RNA <400 copies/mL 667 receiving SELZENTRY compared with placebo were 56% and 22%, respectively. The mean 668 changes in plasma HIV-1 RNA from baseline to Week 48 were $-1.84 \log_{10}$ copies/mL for 669 subjects receiving SELZENTRY + OBT compared with $-0.78 \log_{10}$ copies/mL for subjects 670 receiving OBT only. The mean increase in CD4+ cell counts was higher on SELZENTRY twice 671 daily + OBT (124 cells/mm³) than on placebo + OBT (60 cells/mm³).

672 14.2 Study in Dual/Mixed-Tropic, Treatment-Experienced Subjects

673 Study A4001029 was an exploratory, randomized, double-blind, multicenter trial to 674 determine the safety and efficacy of SELZENTRY in subjects infected with dual/mixed 675 coreceptor tropic HIV-1. The inclusion/exclusion criteria were similar to those for Studies 676 A4001027 and A4001028 above and the subjects were randomized in a 1:1:1 ratio to 677 SELZENTRY once daily, SELZENTRY twice daily, or placebo. No increased risk of infection 678 or HIV disease progression was observed in the subjects who received SELZENTRY. Use of 679 SELZENTRY was not associated with a significant decrease in HIV-1 RNA compared with 680 placebo in these subjects and no adverse effect on CD4+ cell count was noted.

681 **14.3 Study in Treatment-Naive Subjects**

682 Study A4001026 is an ongoing, randomized, double-blind, multicenter study in subjects 683 infected with CCR5-tropic HIV-1 classified by the original TROFILE tropism assay. Subjects 684 were required to have plasma HIV-1 RNA \geq 2,000 copies/mL and could not have: 1) previously 685 received any antiretroviral therapy for >14 days, 2) an active or recent opportunistic infection or 686 a suspected primary HIV-1 infection, or 3) phenotypic or genotypic resistance to zidovudine, 687 lamivudine, or efavirenz. Subjects were randomized in a 1:1:1 ratio to SELZENTRY 300 mg 688 once daily, SELZENTRY 300 mg twice daily, or efavirenz 600 mg once daily, each in 689 combination with zidovudine/lamivudine. The efficacy and safety of SELZENTRY are based on 690 the comparison of SELZENTRY twice daily versus efavirenz. In a pre-planned interim analysis 691 at 16 weeks, SELZENTRY 300 mg once daily failed to meet the pre-specified criteria for 692 demonstrating non-inferiority and was discontinued.

The demographic and baseline characteristics of the maraviroc and efavirenz treatment
 groups were comparable (Table 14). Subjects were stratified by screening HIV-1 RNA levels and
 by geographic region. The median CD4+ cell counts and mean HIV-1 RNA at baseline were

- 696 similar for both treatment groups.
- 697

SELZENTRY	Efavirenz
300 mg Twice Daily +	600 mg Once Daily +
Zidovudine/Lamivudine	Zidovudine/Lamivudine
(N = 360)	(N = 361)
36.7	37.4
20-69	18-77
104 (29)	102 (28)
204 (57)	198 (55)
123 (34)	133 (37)
6 (2)	5 (1)
27 (8)	25 (7)
241 (5-1,422)	254 (8-1,053)
4.9 (3-7)	4.9 (3-7)
	300 mg Twice Daily + Zidovudine/Lamivudine (N = 360) 36.7 20-69 104 (29) 204 (57) 123 (34) 6 (2) 27 (8) 241 (5-1,422)

698Table 14. Demographic and Baseline Characteristics of Subjects in Study A4001026

699

700 The treatment outcomes at 96 weeks for Study A4001026 are shown in Table 15. 701 Treatment outcomes are based on reanalysis of the screening samples using a more sensitive 702 tropism assay, Enhanced sensitivity TROFILE HIV tropism assay, which became available after 703 the Week 48 analysis, approximately 15% of the subjects identified as CCR5-tropic in the 704 original analysis had dual/mixed- or CXCR4-tropic virus. Screening with enhanced sensitivity 705 version of the TROFILE tropism assay reduced the number of maraviroc virologic failures with 706 CXCR4- or dual/mixed-tropic virus at failure to 12 compared with 24 when screening with the 707 original TROFILE HIV tropism assay.

708

709	Table 15: Study Outcome (Snapshot) at	Week 96 Using Enhanced	Sensitivity Assay ^a				
		SELZENTRY	Efavirenz				
		300 mg Twice Daily +	600 mg Once Daily +				
		Zidovudine/Lamivudine	Zidovudine/Lamivudine				
		N = 311	N = 303				
	Outcome at Week 96 ^b	n (%)	n (%)				
	Virologic Responders:	199 (64)	195 (64)				
	(HIV-1 RNA <400 copies/mL)						
	Virologic Failure:						
	Non-sustained HIV-1 RNA	39 (13)	22 (7)				
	suppression						
	HIV-1 RNA never suppressed	9 (3)	1 (<1)				
	Virologic Responders:	183 (59)	190 (63)				
	(HIV-1 RNA <50 copies/mL)						
	Virologic Failure:						
	• Non-sustained HIV-1 RNA	43 (14)	25 (8)				
	suppression						
	• HIV-1 RNA never suppressed	21 (7)	3 (1)				
	Discontinuations due to:						
	Adverse Events	19 (6)	47 (16)				
	• Death	2(1)	2(1)				
	• Other ^c	43 (14)	36 (12)				
710	^a The total number of subjects (Ns) in Tabl		-				
711	virus in the reanalysis of screening sampl	•					
712	reanalysis reclassified approximately 15%		-				
713	or CXCR4-tropic virus. These numbers are different than those presented in Table 14 because						
714	the numbers in Table 14 reflect the subject	ets with CCR5-tropic virus	according to the original				
715	tropism assay.						
716	^b Week 48 results: Virologic responders (<	400): 228/311 (73%) in SE	LZENTRY, 219/303				
717	(72%) in efavirenz;						

7(

Virologic responders (<50): 213/311 (69 %) in SELZENTRY, 207/303 (68%) in efavirenz. 718 ^c Other reasons for discontinuation include lost to follow-up, withdrawn, protocol violation, and 719

720

other.

721

The median increase from baseline in CD4+ cell counts at Week 96 was 184 cells/mm³ 722 for the arm receiving SELZENTRY compared with 155 cells/mm³ for the efavirenz arm. 723

724 **15 REFERENCES**

- 1. IAS-USA Drug Resistance Mutations Figures.
- 726 http://www.iasusa.org/pub/topics/2006/issue3/125.pdf

727 16 HOW SUPPLIED/STORAGE AND HANDLING

- 728 SELZENTRY film-coated tablets are available as follows:
- 150- and 300-mg tablets are blue, biconvex, oval, film-coated tablets debossed with "MVC 150"
- 730 or "MVC 300" on one side and plain on the other.
- 731 Bottle packs 150-mg tablets: 60 tablets (NDC 49702-223-18).
- Bottle packs 300-mg tablets: 60 tablets (NDC 49702-224-18).
- 733 SELZENTRY film-coated tablets should be stored at 25° C (77°F); excursions permitted
- between 15°C and 30°C (59°F-86°F) [see USP Controlled Room Temperature].
 Shelf life is 24 months.

736 17 PATIENT COUNSELING INFORMATION

- 737 *See Medication Guide.*
- Patients should be informed that liver problems including life-threatening cases have
 been reported with SELZENTRY. Patients should be informed that if they develop signs or
- 740 symptoms of hepatitis or allergic reaction following use of SELZENTRY (rash, skin, or eyes
- 741 look yellow; dark urine; vomiting; abdominal pain), they should stop SELZENTRY and seek
- medical evaluation immediately. Patients should understand that laboratory tests for liver
- real enzymes and bilirubin will be ordered prior to starting SELZENTRY, at other times during
- treatment, and if they develop severe rash or signs and symptoms of hepatitis or an allergic
- reaction on treatment [see Warnings and Precautions (5.1)].
- Patients should be informed that SELZENTRY is not a cure for HIV infection and
 patients may still develop illnesses associated with HIV infection, including opportunistic
 infections. The use of SELZENTRY has not been shown to reduce the risk of transmission of
- 749 HIV to others through sexual contact, sharing needles, or blood contamination.
- 750 Patients should be advised that it is important to:
- remain under the care of a physician when using SELZENTRY;
- take SELZENTRY every day as prescribed and in combination with other antiretroviral drugs;
- report to their physician the use of any other prescription or nonprescription medication or
 herbal products;
- inform their physician if they are pregnant, plan to become pregnant or become pregnant
 while taking SELZENTRY;
- not change the dose or dosing schedule of SELZENTRY or any antiretroviral medication
 without consulting their physician.
- Patients should be advised that it is important to take all their anti-HIV medicines asprescribed and at the same time(s) each day.

- Patients should be advised that when their supply of SELZENTRY starts to run low, theyshould ask their doctor or pharmacist for a refill.
- Patients should be advised that if they forget to take a dose, they should take the next dose of SELZENTRY as soon as possible and then take their next scheduled dose at its regular time. If it is less than 6 hours before their next scheduled dose, they should not take the missed dose and should instead wait and take the next dose at the regular time.
- Caution should be used when administering SELZENTRY in patients with a history of
 postural hypotension or on concomitant medication known to lower blood pressure. Patients
 should be advised that if they experience dizziness while taking SELZENTRY, they should
 avoid driving or operating machinery.
- 772
- 773 TROFILE[®] is a registered trademark of Monogram Biosciences, Inc.
- 774

775 Manufactured for:



776	Healthcare
777	ViiV Healthcare
778	Research Triangle Park, NC 27709
779	
780	by:
781	Pfizer Manufacturing Deutschland GmbH
782	Freiburg, Germany
783	
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785	
786	July 2011
787	SEL: 4PI
788	
789	PHARMACIST-DETACH HERE AND GIVE MEDICATION GUIDE TO PATIENT
790	
791	MEDICATION GUIDE
792	SELZENTRY [®] (sell-ZEN-tree) Tablets
793	(maraviroc)
794	
795	Read the Medication Guide that comes with SELZENTRY before you start taking it
796	and each time you get a refill. There may be new information. This information
797	does not take the place of talking with your healthcare provider about your medical
798	condition or treatment.

799	
800	What is the most important information I should know about SELZENTRY?
801	
802	Serious side effects have occurred with SELZENTRY, including liver
803	problems (liver toxicity). An allergic reaction may happen before liver problems
804	occur. Stop taking SELZENTRY and call your healthcare provider right away if you
805	get any of the following symptoms:
806	 an itchy rash on your body (allergic reaction)
807	 yellowing of your skin or whites of your eyes (jaundice)
808	 dark (tea-colored) urine
809	vomiting
810	 upper right stomach area (abdominal) pain
811	
812	What is SELZENTRY?
813	SELZENTRY is an anti-HIV medicine called a CCR5 antagonist. HIV (Human
814	Immunodeficiency Virus) is the virus that causes AIDS (Acquired Immune
815	Deficiency Syndrome).
816	
817	SELZENTRY is used with other anti-HIV medicines in adults with CCR5-tropic HIV-1
818	infection.
819	
820	Use of SELZENTRY is not recommended in people with dual/mixed or CXCR4-tropic
821	HIV-1.
822	
823	SELZENTRY will not cure HIV infection.
824	People taking SELZENTRY may still develop infections, including opportunistic
825	infections or other conditions that happen with HIV infection.
826	• It is very important that you stay under the care of your healthcare provider
827	during treatment with SELZENTRY.
828	• The long-term effects of SELZENTRY are not known at this time.
829	
830	SELZENTRY has not been studied in children less than 16 years of age.
831	
832	Does SELZENTRY lower the risk of passing HIV to other people?
833	No. CELZENTOV does not lower the risk of possing LUV to other possi-
834 825	No, SELZENTRY does not lower the risk of passing HIV to other people
835	through sexual contact, sharing needles, or being exposed to your blood.
836	Continue to practice safer sex.

837 838 839 840 841 842 843	 Use latex or polyurethane condoms or other barrier methods to lower the chance of sexual contact with any body fluids. This includes semen from a man, vaginal secretions from a woman, or blood. Never re-use or share needles. Ask your healthcare provider if you have any questions about safer sex or how to prevent passing HIV to other people.
844	How does SELZENTRY work?
845	HIV enters cells in your blood by attaching itself to structures on the surface of the
846	cell called receptors. SELZENTRY blocks a specific receptor called CCR5 that
847	CCR5-tropic HIV-1 uses to enter CD4 or T-cells in your blood. Your healthcare
848	provider will do a blood test to see if you have been infected with CCR5-tropic
849	HIV-1 before prescribing SELZENTRY for you.
850	
851 852 853 854	 When used with other anti-HIV medicines, SELZENTRY may: reduce the amount of HIV in your blood. This is called "viral load". increase the number of white blood cells called T (CD4) cells.
855	SELZENTRY does not work in all people with CCR5-tropic HIV-1 infection.
856	
857	Who should not take SELZENTRY?
858	People with severe kidney problems or who are on hemodialysis and are taking
858 859	People with severe kidney problems or who are on hemodialysis and are taking certain other medications should not take SELZENTRY. Talk to your healthcare
859	certain other medications should not take SELZENTRY. Talk to your healthcare
859 860	certain other medications should not take SELZENTRY. Talk to your healthcare
859 860 861	certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY?
859 860 861 862 863 864	certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you:
859 860 861 862 863 864 865	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C.
859 860 861 862 863 864 865 866	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems.
859 860 861 862 863 864 865 866 867	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems.
859 860 861 862 863 864 865 866 866 867 868	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure.
 859 860 861 862 863 864 865 866 867 868 869 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition.
 859 860 861 862 863 864 865 866 867 868 869 870 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition. are pregnant or plan to become pregnant. It is not known if SELZENTRY may
 859 860 861 862 863 864 865 866 867 868 869 870 871 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition. are pregnant or plan to become pregnant. It is not known if SELZENTRY may harm your unborn baby.
 859 860 861 862 863 864 865 866 867 868 869 870 871 872 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition. are pregnant or plan to become pregnant. It is not known if SELZENTRY may harm your unborn baby. Antiretroviral Pregnancy Registry. There is a pregnancy registry for women
 859 860 861 862 863 864 865 866 867 868 869 870 871 872 873 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition. are pregnant or plan to become pregnant. It is not known if SELZENTRY may harm your unborn baby. Antiretroviral Pregnancy Registry. There is a pregnancy registry for women who take antiviral medicines during pregnancy. The purpose of the registry is to
 859 860 861 862 863 864 865 866 867 868 869 870 871 872 	 certain other medications should not take SELZENTRY. Talk to your healthcare provider before taking this medicine if you have kidney problems. What should I tell my healthcare provider before taking SELZENTRY? Before you take SELZENTRY, tell your healthcare provider if you: have liver problems including a history of hepatitis B or C. have heart problems. have kidney problems. have low blood pressure or take medicines to lower blood pressure. have any other medical condition. are pregnant or plan to become pregnant. It is not known if SELZENTRY may harm your unborn baby. Antiretroviral Pregnancy Registry. There is a pregnancy registry for women

- 876 • are breastfeeding or plan to breastfeed. It is recommended that HIV-positive 877 women should not breastfeed their babies. This is because of the chance of passing HIV to your baby. You should not breastfeed if you are taking 878 879 SELZENTRY because the risk to your baby is unknown. Talk with your healthcare 880 provider about the best way to feed your baby.
- 881

882 Tell your healthcare provider about all the medicines you take, including 883 prescription and non-prescription medicines, vitamins, and herbal supplements. 884 Certain other medicines may affect the levels of SELZENTRY in your blood. Your 885 healthcare provider may need to change your dose of SELZENTRY when you take it 886 with certain medicines.

887

888 The levels of SELZENTRY in your blood may change and your healthcare provider 889 may need to adjust your dose of SELZENTRY when taking any of the following 890 medications together with SELZENTRY:

- 891
- 892 - darunavir (Prezista[®])
- 893 - lopinavir/ritonavir (Kaletra[®], Norvir[®])
- 894 - atazanavir (Reyataz[®])
- 895 - saguinavir (Invirase[®])
- 896 - nelfinavir (Viracept[®])
- 897 - indinavir (Crixivan[®])
- 898 - fosamprenavir (Lexiva[®])
- 899 - etravirine (Intelence[®])
- carbamezepine (Tegretol[®]) 900
- 901 - phenytoin (Dilantin[®])
- Ritonavir (Norvir[®]) 902
- 903

904 Do not take products that contain St. John's Wort (hypericum perforatum). 905 St. John's Wort may lower the levels of SELZENTRY in your blood so that it

906 will not work to treat your CCR5-tropic HIV infection.

- 907
- 908 Know the medicines you take. Keep a list of your medicines. Show the list to 909 your healthcare provider and pharmacist when you get a new medicine.
- 910
- 911 How should I take SELZENTRY?
- 912
- 913 Take SELZENTRY exactly as prescribed by your healthcare provider.
- 914 SELZENTRY comes in 150-mg and 300-mg tablets. Your healthcare provider will
- 915 prescribe the dose that is right for you.

- delavirdine (Rescriptor[®])
- ketoconazole (Nizoral[®])
- itraconazole (Sporanox[®])
- clarithromycin (Biaxin[®])
- nefazodone (Serzone^{®)}
- telithromycin (Ketek[®])
- efavirenz (Sustiva[®], Atripla[®])
- rifampin (Rifadin[®], Rifater[®])
- phenobarbital (Luminal[®])

916	•	Take SELZENTRY 2 times a day.
917	•	Swallow SELZENTRY tablets whole. Do not chew the tablets.
918	•	Take SELZENTRY tablets with or without food.
919	•	Always take SELZENTRY with other anti-HIV drugs as prescribed by your
920		healthcare provider.
921		
922	Do	o not change your dose or stop taking SELZENTRY or your other anti-HIV
923	m	edicines without first talking with your healthcare provider.
924		
925	•	If you take too much SELZENTRY, call your healthcare provider or the poison
926		control center right away.
927	•	If you forget to take SELZENTRY, take the next dose of SELZENTRY as soon as
928		possible and then take your next scheduled dose at its regular time. If it is less
929		than 6 hours before your next dose, do not take the missed dose. Wait and take
930		the next dose at the regular time. Do not take a double dose to make up for a
931		missed dose.
932	•	It is very important to take all your anti-HIV medicines as prescribed. This can
933		help your medicines work better. It also lowers the chance that your medicines
934		will stop working to fight HIV (drug resistance).
935	•	When your SELZENTRY supply starts to run low, ask your healthcare provider or
936		pharmacist for a refill. This is very important because the amount of virus in
937		your blood may increase and SELZENTRY could stop working if it is stopped for
938		even a short period of time.
939		
940	W	hat are the possible side effects of SELZENTRY?
941		
942	Th	ere have been serious side effects when SELZENTRY has been given with
943	ot	her anti-HIV drugs including:
944	•	Liver problems. See "What is the most important information I should know
945		about SELZENTRY?"
946	•	Heart problems including heart attack.
947	•	Low blood pressure when standing up (postural hypotension). Low blood
948		pressure when standing up can cause dizziness or fainting. Do not drive a car or
949		operate heavy machinery if you have dizziness while taking SELZENTRY.
950	•	Changes in your immune system. A condition called Immune Reconstitution
951		Syndrome can happen when you start taking HIV medicines. Your immune
952		system may get stronger and could begin to fight infections that have been
953		hidden in your body such as pneumonia, herpes virus, or tuberculosis. Tell your
954		healthcare provider if you develop new symptoms after starting your HIV
955		medicines.

956 957	• Possible chance of infection or cancer. SELZENTRY affects other immune system cells and therefore may possibly increase your chance for getting other
958 959	infections or cancer.
960	The most common side effects of SELZENTRY include colds, cough, fever,
961	rash, and dizziness.
962	
963	Tell your healthcare provider about any side effect that bothers you or does not go
964	away.
965	
966	These are not all of the side effects with SELZENTRY. For more information, ask
967	your healthcare provider or pharmacist.
968	
969	Call your doctor for medical advice about side effects. You may report side effects
970	to FDA at 1-800-FDA-1088.
971	
972	How should I store SELZENTRY?
973	 Store SELZENTRY tablets at room temperature from 59°F to 86°F (15°C to 20%C)
974 075	30°C).
975	• Safely throw away medicine that is out of date or that you no longer need.
976 977	Keep SELZENTRY and all medicines out of the reach of children.
978	Reep SELZENTRY and an medicines out of the reach of children.
979	General information about SELZENTRY
980	Medicines are sometimes prescribed for conditions that are not mentioned in
981	Medication Guides. Do not use SELZENTRY for a condition for which it was not
982	prescribed. Do not give SELZENTRY to other people, even if they have the same
983	symptoms you have. It may harm them.
984	
985	This Medication Guide summarizes the most important information about
986	SELZENTRY. If you would like more information, talk with your healthcare provider.
987	You can ask your healthcare provider or pharmacist for more information about
988	SELZENTRY that is written for health professionals.
989	For more information, go to www.selzentry.com.
990	
991	What are the ingredients in SELZENTRY?
992	Active ingredient: maraviroc
993	Inactive ingredients: microcrystalline cellulose, dibasic calcium phosphate
994	(anhydrous), sodium starch glycolate, magnesium stearate

- **Film-coat:** FD&C blue #2 aluminum lake, soya lecithin, polyethylene glycol
- 996 (macrogol 3350), polyvinyl alcohol, talc, and titanium dioxide
- 997
- 998 The brands listed are the trademarks or registered marks of their respective owners
- 999 and are not trademarks of ViiV Healthcare. The makers of these brands are not
- 1000 affiliated with and do not endorse ViiV Healthcare or its products.
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- 1002 This Medication Guide has been approved by the US Food and Drug Administration.
- 1003
- 1004
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