1 SUSTIVA<sup>®</sup>

# 2 (efavirenz) capsules and tablets

3

**Rx only** 

# 4 **DESCRIPTION**

SUSTIVA<sup>®</sup> (efavirenz) is a human immunodeficiency virus type 1 (HIV-1) specific, nonnucleoside, reverse transcriptase inhibitor (NNRTI).

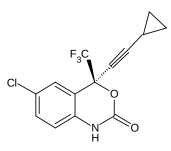
Capsules: SUSTIVA is available as capsules for oral administration containing either
50 mg, 100 mg, or 200 mg of efavirenz and the following inactive ingredients: lactose
monohydrate, magnesium stearate, sodium lauryl sulfate, and sodium starch glycolate.
The capsule shell contains the following inactive ingredients and dyes: gelatin, sodium
lauryl sulfate, titanium dioxide, and/or yellow iron oxide. The capsule shells may also
contain silicon dioxide. The capsules are printed with ink containing carmine 40 blue,
FD&C Blue No. 2, and titanium dioxide.

**Tablets:** SUSTIVA is available as film-coated tablets for oral administration containing 600 mg of efavirenz and the following inactive ingredients: croscarmellose sodium, hydroxypropyl cellulose, lactose monohydrate, magnesium stearate, microcrystalline cellulose, and sodium lauryl sulfate. The film coating contains Opadry<sup>®</sup> Yellow and Opadry<sup>®</sup> Clear. The tablets are polished with carnauba wax and printed with purple ink, Opacode<sup>®</sup> WB.

20 Efavirenz is chemically described as (S)-6-chloro-4-(cyclopropylethynyl)-1,4-dihydro-4-

21 (trifluoromethyl)-2H-3,1-benzoxazin-2-one.

22 Its empirical formula is  $C_{14}H_9ClF_3NO_2$  and its structural formula is:



23

- Efavirenz is a white to slightly pink crystalline powder with a molecular mass of 315.68.
- 25 It is practically insoluble in water ( $<10 \mu g/mL$ ).

#### 26 MICROBIOLOGY

#### 27 Mechanism of Action

Efavirenz (EFV) is a non-nucleoside reverse transcriptase inhibitor (NNRTI) of human immunodeficiency virus type 1 (HIV-1). EFV activity is mediated predominantly by noncompetitive inhibition of HIV-1 reverse transcriptase (RT). HIV-2 RT and human cellular DNA polymerases  $\alpha$ ,  $\beta$ ,  $\gamma$ , and  $\delta$  are not inhibited by EFV.

#### 32 Antiviral Activity In Vitro

33 The concentration of EFV inhibiting in vitro replication of wild-type laboratory adapted strains and clinical isolates by 90-95% (IC<sub>90-95</sub>) ranged from 1.7 to 25 nM in 34 35 lymphoblastoid cell lines, peripheral blood mononuclear cells (PBMCs), and 36 macrophage/monocyte cultures. EFV demonstrated antiviral activity against most non-37 clade B isolates (subtypes A, AE, AG, C, D, F, G, J, N), but had reduced antiviral activity 38 against group O viruses. EFV demonstrated additive antiviral activity without 39 cytotoxicity against HIV-1 in cell culture when combined with the NNRTIs delavirdine 40 (DLV) and nevirapine (NVP), NRTIs (abacavir, didanosine, emtricitabine, lamivudine 41 [LAM], stavudine, tenofovir, zalcitabine, zidovudine [ZDV]), PIs (amprenavir, indinavir 42 [IDV], lopinavir, nelfinavir, ritonavir, saquinavir), and the fusion inhibitor enfuvirtide. 43 EFV demonstrated additive to antagonistic antiviral activity in vitro with atazanavir. EFV 44 was not antagonistic with adefovir, used for the treatment of hepatitis B virus infection, 45 or ribavirin, used in combination with interferon for the treatment of hepatitis C virus 46 infection.

#### 47 **Resistance**

In vitro: HIV-1 isolates with reduced susceptibility to EFV (>380-fold increase in IC<sub>90</sub>
 value) emerged rapidly under *in vitro* selection. Genotypic characterization of these
 viruses identified mutations resulting in single amino acid substitutions L100I or V179D,
 double substitutions L100I/V108I, and triple substitutions L100I/V179D/ Y181C in RT.

52 **Clinical studies:** Clinical isolates with reduced susceptibility *in vitro* to EFV have been 53 obtained. One or more RT substitutions at amino acid positions 98, 100, 101, 103, 106, 54 108, 188, 190, 225, and 227 were observed in patients failing treatment with EFV in 55 combination with IDV, or with ZDV plus LAM. The mutation K103N was the most 56 frequently observed. Long-term resistance surveillance (average 52 weeks, range 4-106 57 weeks) analyzed 28 matching baseline and virologic failure isolates. Sixty-one percent 58 (17/28) of these failure isolates had decreased EFV susceptibility *in vitro* with a median 59 88-fold change in EFV susceptibility (IC<sub>50</sub> value) from reference. The most frequent 60 NNRTI mutation to develop in these patient isolates was K103N (54%). Other NNRTI 61 mutations that developed included L100I (7%), K101E/Q/R (14%), V108I (11%), 62 G190S/T/A (7%), P225H (18%), and M230I/L (11%).

#### 63 Cross-Resistance

64 Cross-resistance among NNRTIs has been observed. Clinical isolates previously 65 characterized as EFV-resistant were also phenotypically resistant *in vitro* to DLV and 66 NVP compared to baseline. DLV- and/or NVP-resistant clinical viral isolates with 67 NNRTI resistance-associated substitutions (A98G, L100I, K101E/P, K103N/S, V106A, 68 Y181X, Y188X, G190X, P225H, F227L, or M230L) showed reduced susceptibility to 69 EFV *in vitro*. Greater than 90% of NRTI-resistant clinical isolates tested *in vitro* retained 70 susceptibility to EFV.

## 71 CLINICAL PHARMACOLOGY

#### 72 Pharmacokinetics

73 **Absorption:** Peak efavirenz plasma concentrations of 1.6-9.1  $\mu$ M were attained by 74 5 hours following single oral doses of 100 mg to 1600 mg administered to uninfected

volunteers. Dose-related increases in C<sub>max</sub> and AUC were seen for doses up to 1600 mg;
the increases were less than proportional suggesting diminished absorption at higher
doses.

In HIV-infected patients at steady state, mean  $C_{max}$ , mean  $C_{min}$ , and mean AUC were dose proportional following 200-mg, 400-mg, and 600-mg daily doses. Time-to-peak plasma concentrations were approximately 3-5 hours and steady-state plasma concentrations were reached in 6-10 days. In 35 patients receiving SUSTIVA 600 mg once daily, steady-state  $C_{max}$  was 12.9  $\pm$  3.7  $\mu$ M (mean  $\pm$  SD), steady-state  $C_{min}$  was 5.6  $\pm$  3.2  $\mu$ M, and AUC was 184  $\pm$  73  $\mu$ M•h.

#### 84 Effect of Food on Oral Absorption:

85 *Capsules*—Administration of a single 600-mg dose of efavirenz capsules with a high-86 fat/high-caloric meal (894 kcal, 54 g fat, 54% calories from fat) or a reduced-fat/normal-87 caloric meal (440 kcal, 2 g fat, 4% calories from fat) was associated with a mean increase 88 of 22% and 17% in efavirenz AUC<sub> $\infty$ </sub> and a mean increase of 39% and 51% in efavirenz 89 C<sub>max</sub>, respectively, relative to the exposures achieved when given under fasted 90 conditions. (See **DOSAGE AND ADMINISTRATION** and **PRECAUTIONS:** 91 **Information for Patients**.)

92 *Tablets*—Administration of a single 600-mg efavirenz tablet with a high-fat/high-caloric 93 meal (approximately 1000 kcal, 500-600 kcal from fat) was associated with a 28% 94 increase in mean  $AUC_{\infty}$  of efavirenz and a 79% increase in mean  $C_{max}$  of efavirenz 95 relative to the exposures achieved under fasted conditions. (See **DOSAGE AND** 96 **ADMINISTRATION** and **PRECAUTIONS: Information for Patients**.)

97 **Distribution:** Efavirenz is highly bound (approximately 99.5-99.75%) to human plasma 98 proteins, predominantly albumin. In HIV-1 infected patients (n=9) who received 99 SUSTIVA 200 to 600 mg once daily for at least one month, cerebrospinal fluid 100 concentrations ranged from 0.26 to 1.19% (mean 0.69%) of the corresponding plasma 101 concentration. This proportion is approximately 3-fold higher than the non-protein-bound 102 (free) fraction of efavirenz in plasma.

103 **Metabolism:** Studies in humans and *in vitro* studies using human liver microsomes have 104 demonstrated that efavirenz is principally metabolized by the cytochrome P450 system to 105 hydroxylated metabolites with subsequent glucuronidation of these hydroxylated 106 metabolites. These metabolites are essentially inactive against HIV-1. The *in vitro* studies 107 suggest that CYP3A4 and CYP2B6 are the major isozymes responsible for efavirenz 108 metabolism.

109 Efavirenz has been shown to induce P450 enzymes, resulting in the induction of its own

110 metabolism. Multiple doses of 200-400 mg per day for 10 days resulted in a lower than

- 111 predicted extent of accumulation (22-42% lower) and a shorter terminal half-life of 40-55
- 112 hours (single dose half-life 52-76 hours).

**Elimination:** Efavirenz has a terminal half-life of 52-76 hours after single doses and 40-55 hours after multiple doses. A one-month mass balance/excretion study was conducted using 400 mg per day with a <sup>14</sup>C-labeled dose administered on Day 8. Approximately 14-34% of the radiolabel was recovered in the urine and 16-61% was recovered in the feces. Nearly all of the urinary excretion of the radiolabeled drug was in the form of metabolites. Efavirenz accounted for the majority of the total radioactivity measured in feces.

## 120 Special Populations

Hepatic Impairment: The pharmacokinetics of efavirenz have not been adequately
 studied in patients with hepatic impairment (see PRECAUTIONS: General).

123 Renal Impairment: The pharmacokinetics of efavirenz have not been studied in 124 patients with renal insufficiency; however, less than 1% of efavirenz is excreted 125 unchanged in the urine, so the impact of renal impairment on efavirenz elimination should 126 be minimal.

127 Gender and Race: The pharmacokinetics of efavirenz in patients appear to be similar128 between men and women and among the racial groups studied.

#### 129 Geriatric: see PRECAUTIONS: Geriatric Use

#### 130 Pediatrics: see PRECAUTIONS: Pediatric Use

#### Drug Interactions also CONTRAINDICATIONS (see and 131 **PRECAUTIONS:** Drug Interactions) 132

133 Efavirenz has been shown *in vivo* to cause hepatic enzyme induction, thus increasing the 134 biotransformation of some drugs metabolized by CYP3A4. In vitro studies have shown 135 that efavirenz inhibited P450 isozymes 2C9, 2C19, and 3A4 with K<sub>i</sub> values (8.5-17 µM) in the range of observed efavirenz plasma concentrations. In *in vitro* studies, efavirenz 136 137 did not inhibit CYP2E1 and inhibited CYP2D6 and CYP1A2 (K<sub>i</sub> values 82-160 µM) only 138 at concentrations well above those achieved clinically. The effects on CYP3A4 activity 139 are expected to be similar between 200-mg, 400-mg, and 600-mg doses of efavirenz. 140 Coadministration of efavirenz with drugs primarily metabolized by 2C9, 2C19, and 3A4 141 isozymes may result in altered plasma concentrations of the coadministered drug. Drugs which induce CYP3A4 activity would be expected to increase the clearance of efavirenz 142 143 resulting in lowered plasma concentrations.

144 Drug interaction studies were performed with efavirenz and other drugs likely to be 145 coadministered or drugs commonly used as probes for pharmacokinetic interaction. The 146 effects of coadministration of efavirenz on the C<sub>max</sub>, AUC, and C<sub>min</sub> are summarized in Table 1 (effect of efavirenz on other drugs) and Table 2 (effect of other drugs on 147 148 efavirenz). For information regarding clinical recommendations see **PRECAUTIONS**:

149	Drug Interactions.	
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			Number	Coadministered Drug (mean % change)		
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)
Atazanavir	400 mg qd with a light meal d 1-20	600 mg qd with a light meal d 7-20	27	↓ 59% (49-67%)	↓ 74% (68-78%)	↓ 93% (90-95%)
	400 mg qd d 1-6, then 300 mg qd d 7-20 with ritonavir 100 mg qd and a light meal	600 mg qd 2 h after atazanavir and ritonavir d 7-20	13	↑ 14% <sup>a</sup> (↓ 17-↑ 58%)	↑ 39% <sup>a</sup> (2-88%)	↑ 48% <sup>a</sup> (24-76%)
Indinavir	1000 mg q8h x 10 days After morning dose	600 mg x 10 days	20	⇔p	↓ 33% <sup>b</sup> (26-39%)	↓ 39% <sup>b</sup> (24-51%)

 Table 1:
 Effect of Efavirenz on Coadministered Drug Plasma C<sub>max</sub>, AUC, and C<sub>min</sub>

				Coadministered Drug (mean % change)		
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)
	After afternoon dose			$\leftrightarrow^{\mathrm{b}}$	$\downarrow 37\%^{b}$	$\downarrow 52\%^{b}$
	After evening dose			$\downarrow 29\%^{b}$	(26-46%) ↓ 46% <sup>b</sup>	(47-57%) ↓ 57% <sup>b</sup>
				(11-43%)	(37-54%)	(50-63%)
Lopinavir/ ritonavir	400/100 mg q12h x 9 days	600 mg x 9 days	11,7 <sup>c</sup>	$\leftrightarrow^{d}$	↓ 19% <sup>d</sup>	$\downarrow 39\%^{d}$
NT 10	750 01	(00	10	A 210/	(↓ 36-↑ 3%)	(3-62%)
Nelfinavir	750 mg q8h x 7 days	600 mg x 7 days	10	↑ 21%	↑ 20%	$\leftrightarrow$
Metabolite	7 duys	/ days		(10-33%) ↓ 40%	(8-34%) ↓ 37%	1 420/
				-		↓ 43% (21,50%)
AG-1402				(30-48%)	(25-48%)	(21-59%)
Ritonavir	500 mg q12h x 8 days	600 mg x 10 days	11			
	After AM dose			↑ 24%	↑ 18%	↑ 42%
				(12-38%)	(6-33%)	(9-86%) <sup>e</sup>
	After PM dose			$\leftrightarrow$	$\leftrightarrow$	↑ 24% (3-50%) <sup>e</sup>
Saminavir	1200 mg q8h x	600 mg v	12	↓ 50%	↓ 62%	↓ 56%
Saquinavir SGC <sup>f</sup>	1200 mg q8n x 10 days	600 mg x 10 days	12	-		-
SGC	10 44 95	10 44 95		(28-66%)	(45-74%)	(16-77%)
Lamivudine	150 mg q12h x 14 days	600 mg x 14 days	9	$\leftrightarrow$	$\leftrightarrow$	↑ 265% (37-873%
Tenofovir <sup>g</sup>	300 mg qd	600 mg x 14 days	29	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Zidovudine	300 mg q12h x	600 mg x	9	$\leftrightarrow$	$\leftrightarrow$	↑ 225%
	14 days	14 days				(43-640%)
Azithromycin	600 mg single dose	400 mg x	14	↑ 22%	$\leftrightarrow$	NA
5	0 0	7 days		(4-42%)		
Clarithromycin	500 mg q12h x	400 mg x	11	↓ 26%	↓ 39%	↓ 53%
	7 days	7 days		(15-35%)	(30-46%)	(42-63%)
14-OH metabolite	-	-		(15 5570) ↑ 49%	(30 <del>4</del> 0/0) ↑ 34%	(42 0370) ↑ 26%
				(32-69%)	(18-53%)	(9-45%)
Fluconazole	200 mg x 7 days	400 mg x 7 days	10	↔	↔	↔
Itraconazole	200 mg q12h x	600 mg x	18	↓ 37%	↓ 39%	↓ 44%
	28 days	14 days		(20-51%)	(21-53%)	(27-58%)
Hydroxyitraconazole				↓ 35%	↓ 37%	↓ 43%
,,,				(12-52%)	(14-55%)	(18-60%)
Rifabutin	300 mg qd x	600 mg x	9	↓ 32%	↓ 38%	↓ 45%
i internet i	14 days	14 days	,	(15-46%)	(28-47%)	(31-56%)
Voriconazole	400 mg po q12h x 1 day then 200 mg po q12h x 8 days	400 mg x 9 days	NA	↓ 61% <sup>h</sup>	↓ 77% <sup>h</sup>	(51-5076) NA
	300 mg po q12h days 2-7	300 mg x 7 days	NA	$\downarrow$ 36% <sup>i</sup> (21-49%)	↓ 55% <sup>i</sup> (45-62%)	NA

# Table 1: Effect of Efavirenz on Coadministered Drug Plasma C<sub>max</sub>, AUC, and C<sub>min</sub>

	of Elavirenz on		Number	Coadministered Drug (mean % change)			
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)	
	400 mg po q12h days 2-7	300 mg x 7 days	NA	↑ 23% <sup>i</sup> (↓ 1-↑ 53%)	↓ 7% <sup>i</sup> (↓ 23-↑ 13%)	NA	
Atorvastatin	10 mg qd x 4 days	600 mg x 15 days	14	↓ 14% (1-26%)	↓ 43% (34-50%)	↓ 69% (49-81%)	
Total active (including metabolites)				↓ 15% (2-26%)	↓ 32% (21-41%)	↓ 48% (23-64%)	
Pravastatin	40 mg qd x 4 days	600 mg x 15 days	13	↓ 32% (↓ 59-↑ 12%)	↓ 44% (26-57%)	↓ 19% (0-35%)	
Simvastatin	40 mg qd x 4 days	600 mg x 15 days	14	↓ 72% (63-79%)	↓ 68% (62-73%)	↓ 45% (20-62%)	
Total active (including metabolites)				↓ 68% (55-78%)	↓ 60% (52-68%)	NA <sup>j</sup>	
Carbamazepine	200 mg qd x 3 days, 200 mg bid x 3 days, then 400 mg qd x 29 days	600 mg x 14 days	12	↓ 20% (15-24%)	↓ 27% (20-33%)	↓ 35% (24-44%)	
Epoxide metabolite				$\leftrightarrow$	$\leftrightarrow$	↓ 13% (↓ 30-↑ 7%)	
Cetirizine	10 mg single dose	600 mg x 10 days	11	↓ 24% (18-30%)	$\leftrightarrow$	NA	
Diltiazem	240 mg x 21 days	600 mg x 14 days	13	↓ 60% (50-68%)	↓ 69% (55-79%)	↓ 63% (44-75%)	
Desacetyl diltiazem				↓ 64% (57-69%)	↓ 75% (59-84%)	↓ 62% (44-75%)	
N-monodesmethyl diltiazem				↓ 28% (7-44%)	↓ 37% (17-52%)	↓ 37% (17-52%)	
Ethinyl estradiol	50 µg single dose	400 mg x 10 days	13	$\leftrightarrow$	↑ 37% (25-51%)	NA	
Lorazepam	2 mg single dose	600 mg x 10 days	12	↑ 16% (2-32%)	$\leftrightarrow$	NA	
Methadone	Stable maintenance 35-100 mg daily	600 mg x 14-21 days	11	↓ 45% (25-59%)	↓ 52% (33-66%)	NA	
Paroxetine	20 mg qd x 14 days	600 mg x 14 days	16	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$	
Sertraline	50 mg qd x 14 days	600 mg x 14 days	13	↓ 29% (15-40%)	↓ 39% (27-50%)	↓ 46% (31-58%)	

# Table 1: Effect of Efavirenz on Coadministered Drug Plasma C<sub>max</sub>, AUC, and C<sub>min</sub>

# Table 1: Effect of Efavirenz on Coadministered Drug Plasma C<sub>max</sub>, AUC, and C<sub>min</sub>

			Number	(	Coadministered Dr (mean % change)	0
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)
	cates decrease ↔	→ Indicates no change or	a mean incre	ase or decrease of	f<10%.	
a Compared with atazanavi	400 mg qd alone.					
b Comparator dose of indina	avir was 800 mg q	8h x 10 days.				
<sup>c</sup> Parallel-group design; n fo	or efavirenz + lopi	navir/ritonavir, n for lopi	navir/ritonav	ir alone.		
<sup>d</sup> Values are for lopinavir; t	he pharmacokineti	ics of ritonavir 100 mg q	12h are unaffe	ected by concurre	nt efavirenz.	
e 95% CI.						
<sup>f</sup> Soft Gelatin Capsule.						
<sup>g</sup> Tenofovir disoproxil fuma	ırate.					
<sup>h</sup> 90% CI not available.						

i Relative to steady-state administration of voriconazole (400 mg for 1 day, then 200 mg po q12h for 2 days).

j Not available because of insufficient data.

NA = not available.

#### 150

# Table 2: Effect of Coadministered Drug on Efavirenz Plasma $C_{max}, \mbox{AUC}, \mbox{and} C_{min}$

			Number		Efavirenz (mean % change)	
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)
Indinavir	800 mg q8h x 14 days	200 mg x 14 days	11	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Lopinavir/ ritonavir	400/100 mg q12h x 9 days	600 mg x 9 days	11,12 <sup>a</sup>	$\leftrightarrow$	↓ 16% (↓ 38-↑ 15%)	↓ 16% (↓ 42-↑ 20%)
Nelfinavir	750 mg q8h x 7 days	600 mg x 7 days	10	$ \begin{array}{c} \downarrow 12\% \\ (\downarrow 32-\uparrow 13\%)^{b} \end{array} $	↓ 12% (↓ 35-↑ 18%) <sup>b</sup>	↓ 21% (↓ 53-↑ 33%)
Ritonavir	500 mg q12h x 8 days	600 mg x 10 days	9	↑ 14% (4-26%)	↑ 21% (10-34%)	↑25% (7-46%) <sup>b</sup>
Saquinavir SGC <sup>c</sup>	1200 mg q8h x 10 days	600 mg x 10 days	13	↓ 13% (5-20%)	↓ 12% (4-19%)	$\begin{array}{c} \downarrow 14\% \\ (2-24\%)^{b} \end{array}$
Tenofovir <sup>d</sup>	300 mg qd	600 mg x 14 days	30	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Azithromycin	600 mg single dose	400 mg x 7 days	14	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Clarithromycin	500 mg q12h x 7 days	400 mg x 7 days	12	↑ 11% (3-19%)	$\leftrightarrow$	$\leftrightarrow$
Fluconazole	200 mg x 7 days	400 mg x 7 days	10	$\leftrightarrow$	↑ 16% (6-26%)	↑ 22% (5-41%)
Itraconazole	200 mg q12h x 14 days	600 mg x 28 days	16	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Rifabutin	300 mg qd x 14 days	600 mg x 14 days	11	$\leftrightarrow$	$\leftrightarrow$	↓ 12% (↓ 24-↑ 1%)
Rifampin	600 mg x 7 days	600 mg x 7 days	12	↓ 20% (11-28%)	↓ 26% (15-36%)	↓ 32% (15-46%)

			Number		Efavirenz (mean % change)	
Coadministered Drug	Dose	Efavirenz Dose	of Subjects	C <sub>max</sub> (90% CI)	AUC (90% CI)	C <sub>min</sub> (90%CI)
Voriconazole	400 mg po q12h x 1 day then 200 mg po q12h x 8 days	400 mg x 9 days	NA	↑ 38% <sup>e</sup>	↑44% <sup>e</sup>	NA
	300 mg po q12h days 2-7	300 mg x 7 days	NA	↓ 14% <sup>f</sup> (7-21%)	$\leftrightarrow^{\mathrm{f}}$	NA
	400 mg po q12h days 2-7	300 mg x 7 days	NA	$\overset{f}{\leftrightarrow}$	↑17% <sup>f</sup> (6-29%)	NA
Atorvastatin	10 mg qd x 4 days	600 mg x 15 days	14	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Pravastatin	40 mg qd x 4 days	600 mg x 15 days	11	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Simvastatin	40 mg qd x 4 days	600 mg x 15 days	14	↓ 12% (↓ 28-↑ 8%)	$\leftrightarrow$	↓ 12% (↓ 25-↑ 3%)
Aluminum hydroxide 400 mg magnesium hydroxide 400 mg, plus simethicone 40 mg	30 mL single dose	400 mg single dose	17	$\leftrightarrow$	$\leftrightarrow$	NA
Carbamazepine	200 mg qd x 3 days, 200 mg bid x 3 days, then 400 mg qd x 15 days	600 mg x 35 days	14	↓ 21% (15-26%)	↓ 36% (32-40%)	↓ 47% (41-53%)
Cetirizine	10 mg single dose	600 mg x 10 days	11	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Diltiazem	240 mg x 14 days	600 mg x 28 days	12	↑ 16% (6-26%)	↑ 11% (5-18%)	↑ 13% (1-26%)
Ethinyl estradiol	50 µg single dose	400 mg x 10 days	13	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Famotidine	40 mg single dose	400 mg single dose	17	$\leftrightarrow$	$\leftrightarrow$	NA
Paroxetine	20 mg qd x 14 days	600 mg x 14 days	12	$\leftrightarrow$	$\leftrightarrow$	$\leftrightarrow$
Sertraline	50 mg qd x 14 days	600 mg x 14 days	13	↑ 11% (6-16%)	$\leftrightarrow$	$\leftrightarrow$

#### Effect of Coadministered Drug on Efavirenz Plasma C<sub>max</sub>, AUC, and Table 2: Cmin

<sup>b</sup> 95% CI.

c Soft Gelatin Capsule.

d Tenofovir disoproxil fumarate.

e 90% CI not available.

f Relative to steady-state administration of efavirenz (600 mg once daily for 9 days).

NA = not available.

151

# 152 INDICATIONS AND USAGE

153 SUSTIVA (efavirenz) in combination with other antiretroviral agents is indicated for the 154 treatment of HIV-1 infection. This indication is based on two clinical trials of at least one 155 year duration that demonstrated prolonged suppression of HIV RNA.

# 156 **Description of Studies**

157 Study 006, a randomized, open-label trial, compared SUSTIVA (600 mg once daily) + zidovudine (ZDV, 300 mg q12h) + lamivudine (LAM, 150 mg q12h) or SUSTIVA 158 159 (600 mg once daily) + indinavir (IDV, 1000 mg q8h) with indinavir (800 mg q8h) + 160 zidovudine (300 mg q12h) + lamivudine (150 mg q12h). Twelve hundred sixty-six 161 patients (mean age 36.5 years [range 18-81], 60% Caucasian, 83% male) were enrolled. All patients were efavirenz-, lamivudine-, NNRTI-, and PI-naive at study entry. The 162 median baseline CD4+ cell count was 320 cells/mm<sup>3</sup> and the median baseline HIV-1 163 RNA level was 4.8 log<sub>10</sub> copies/mL. Treatment outcomes with standard assay (assay 164 165 limit 400 copies/mL) through 48 and 168 weeks are shown in Table 3. Plasma HIV RNA levels were quantified with standard (assay limit 400 copies/mL) and ultrasensitive (assay 166 limit 50 copies/mL) versions of the AMPLICOR HIV-1 MONITOR<sup>®</sup> assay. During the 167 168 study, version 1.5 of the assay was introduced in Europe to enhance detection of nonclade B virus. 169

		A + ZDV AM	SUSTIV	'A + IDV	IDV + ZI	DV + LAM
	n=	422	n=	429	n=	415
Outcome	Week 48	Week 168	Week 48	Week 168	Week 48	Week 168
Responder <sup>a</sup>	69%	48%	57%	40%	50%	29%
Virologic failure <sup>b</sup>	6%	12%	15%	20%	13%	19%
Discontinued for adverse events	7%	8%	6%	8%	16%	20%
Discontinued for other reasons <sup>c</sup>	17%	31%	22%	32%	21%	32%
CD4+ cell count (cells/mm <sup>3</sup> )						
Observed subjects (n)	(279)	(205)	(256)	(158)	(228)	(129)
Mean change from baseline	190	329	191	319	180	329

Table 3: Outcomes of Randomized Treatment Through 48 and 168 Weeks, Study 006

170 <sup>a</sup> Patients achieved and maintained confirmed HIV-1 RNA <400 copies/mL through Week 48 or Week</li>
 168.

<sup>b</sup> Includes patients who rebounded, patients who were on study at Week 48 and failed to achieve
 confirmed HIV-1 RNA <400 copies/mL at time of discontinuation, and patients who discontinued due to</li>
 lack of efficacy.

<sup>c</sup> Includes consent withdrawn, lost to follow-up, noncompliance, never treated, missing data, protocol
 violation, death, and other reasons. Patients with HIV-1 RNA levels <400 copies/mL who chose not to</li>
 continue in the voluntary extension phases of the study were censored at date of last dose of study
 medication.

For patients treated with SUSTIVA + zidovudine + lamivudine, SUSTIVA + indinavir, or indinavir + zidovudine + lamivudine, the percentage of responders with HIV-1 RNA <50 copies/mL was 65%, 50%, and 45%, respectively, through 48 weeks, and 43%, 31%, and 23%, respectively, through 168 weeks. A Kaplan-Meier analysis of time to loss of virologic response (HIV RNA <400 copies/mL) suggests that both the trends of virologic response and differences in response continue through 4 years.

185 ACTG 364 is a randomized, double-blind, placebo-controlled, 48-week study in NRTI-186 experienced patients who had completed two prior ACTG studies. One-hundred ninety-187 six patients (mean age 41 years [range 18-76], 74% Caucasian, 88% male) received 188 NRTIs in combination with SUSTIVA (efavirenz) (600 mg once daily), or nelfinavir (NFV, 750 mg TID), or SUSTIVA (600 mg once daily) + nelfinavir in a randomized, 189 double-blinded manner. The mean baseline CD4+ cell count was 389 cells/mm<sup>3</sup> and 190 mean baseline HIV-1 RNA level was 8130 copies/mL. Upon entry into the study, all 191 192 patients were assigned a new open-label NRTI regimen, which was dependent on their

193 previous NRTI treatment experience. There was no significant difference in the mean 194 CD4+ cell count among treatment groups; the overall mean increase was approximately 195 100 cells at 48 weeks among patients who continued on study regimens. Treatment 196 outcomes are shown in Table 4. Plasma HIV RNA levels were quantified with the 197 AMPLICOR HIV-1 MONITOR<sup>®</sup> assay using a lower limit of quantification of 500 198 copies/mL.

Outcome	SUSTIVA + NFV + NRTIs n=65	SUSTIVA + NRTIs n=65	NFV + NRTIs n=66
HIV-1 RNA <500 copies/mL <sup>a</sup>	71%	63%	41%
HIV-1 RNA ≥500 copies/mL <sup>b</sup>	17%	34%	54%
CDC Category C Event	2%	0%	0%
Discontinuations for adverse events <sup>c</sup>	3%	3%	5%
Discontinuations for other reasons <sup>d</sup>	8%	0%	0%

Table 4: Outcomes of Randomized Treatment Through 48 Weeks, Study ACTG 364\*

199 \* For some patients, Week 56 data were used to confirm the status at Week 48.

<sup>a</sup> Subjects achieved virologic response (two consecutive viral loads <500 copies/mL) and maintained it through Week 48.</li>

<sup>b</sup> Includes viral rebound and failure to achieve confirmed <500 copies/mL by Week 48.

203 <sup>c</sup> See ADVERSE REACTIONS for a safety profile of these regimens.

204 <sup>d</sup> Includes loss to follow-up, consent withdrawn, noncompliance.

205 A Kaplan-Meier analysis of time to treatment failure through 72 weeks demonstrates a

206 longer duration of virologic suppression (HIV RNA <500 copies/mL) in the SUSTIVA-

207 containing treatment arms.

#### 208 **CONTRAINDICATIONS**

209 SUSTIVA (efavirenz) is contraindicated in patients with clinically significant 210 hypersensitivity to any of its components.

211 SUSTIVA should not be administered concurrently with astemizole, bepridil, cisapride,

212 midazolam, pimozide, triazolam, or ergot derivatives because competition for CYP3A4

213 by efavirenz could result in inhibition of metabolism of these drugs and create the

214 potential for serious and/or life-threatening adverse events (eg, cardiac arrhythmias,

215 prolonged sedation, or respiratory depression). SUSTIVA should not be administered 216 concurrently with standard doses of voriconazole because SUSTIVA significantly

- 217 decreases voriconazole plasma concentrations. Adjusted doses of voriconazole and
- 218 efavirenz may be administered concomitantly (see **CLINICAL PHARMACOLOGY**,
- 219 Tables 1 and 2; **PRECAUTIONS: Drug Interactions**, Table 5; and **DOSAGE AND**
- 220 ADMINISTRATION: Dosage Adjustment).

#### 221 WARNINGS

# ALERT: Find out about medicines that should NOT be taken with SUSTIVA. This statement is also included on the product's bottle labels. (See CONTRAINDICATIONS and PRECAUTIONS: Drug Interactions.)

SUSTIVA must not be used as a single agent to treat HIV-1 infection or added on as a sole agent to a failing regimen. As with all other non-nucleoside reverse transcriptase inhibitors, resistant virus emerges rapidly when efavirenz is administered as monotherapy. The choice of new antiretroviral agents to be used in combination with efavirenz should take into consideration the potential for viral cross-resistance.

230 **Psychiatric Symptoms:** Serious psychiatric adverse experiences have been reported 231 in patients treated with SUSTIVA. In controlled trials of 1008 patients treated with 232 regimens containing SUSTIVA for a mean of 2.1 years and 635 patients treated with 233 control regimens for a mean of 1.5 years, the frequency of specific serious psychiatric 234 events among patients who received SUSTIVA or control regimens, respectively, were: 235 severe depression (2.4%, 0.9%), suicidal ideation (0.7%, 0.3%), nonfatal suicide attempts 236 (0.5%, 0), aggressive behavior (0.4%, 0.5%), paranoid reactions (0.4%, 0.3%), and manic 237 reactions (0.2%, 0.3%). When psychiatric symptoms similar to those noted above were 238 combined and evaluated as a group in a multifactorial analysis of data from Study 006, 239 treatment with efavirenz was associated with an increase in the occurrence of these 240 selected psychiatric symptoms. Other factors associated with an increase in the 241 occurrence of these psychiatric symptoms were history of injection drug use, psychiatric 242 history, and receipt of psychiatric medication at study entry; similar associations were 243 observed in both the SUSTIVA and control treatment groups. In Study 006, onset of new 244 serious psychiatric symptoms occurred throughout the study for both SUSTIVA-treated 245 and control-treated patients. One percent of SUSTIVA-treated patients discontinued or 246 interrupted treatment because of one or more of these selected psychiatric symptoms.

There have also been occasional postmarketing reports of death by suicide, delusions, and psychosis-like behavior, although a causal relationship to the use of SUSTIVA cannot be determined from these reports. Patients with serious psychiatric adverse experiences should seek immediate medical evaluation to assess the possibility that the symptoms may be related to the use of SUSTIVA, and if so, to determine whether the risks of continued therapy outweigh the benefits (see **ADVERSE REACTIONS**).

253 **Nervous System Symptoms:** Fifty-three percent of patients receiving SUSTIVA in 254 controlled trials reported central nervous system symptoms compared to 25% of patients 255 receiving control regimens. These symptoms included, but were not limited to, dizziness 256 (28.1%), insomnia (16.3%), impaired concentration (8.3%), somnolence (7.0%), 257 abnormal dreams (6.2%), and hallucinations (1.2%). These symptoms were severe in 258 2.0% of patients, and 2.1% of patients discontinued therapy as a result. These symptoms 259 usually begin during the first or second day of therapy and generally resolve after the first 260 2-4 weeks of therapy. After 4 weeks of therapy, the prevalence of nervous system 261 symptoms of at least moderate severity ranged from 5% to 9% in patients treated with 262 regimens containing SUSTIVA and from 3% to 5% in patients treated with a control 263 regimen. Patients should be informed that these common symptoms were likely to 264 improve with continued therapy and were not predictive of subsequent onset of the less 265 frequent psychiatric symptoms (see WARNINGS: Psychiatric Symptoms). Dosing at 266 bedtime may improve the tolerability of these nervous system symptoms (see ADVERSE 267 **REACTIONS and DOSAGE AND ADMINISTRATION**).

Analysis of long-term data from Study 006 (median follow-up 180 weeks, 102 weeks, and 76 weeks for patients treated with SUSTIVA + zidovudine + lamivudine, SUSTIVA + indinavir, and indinavir + zidovudine + lamivudine, respectively) showed that, beyond 24 weeks of therapy, the incidences of new-onset nervous system symptoms among SUSTIVA-treated patients were generally similar to those in the indinavir-containing control arm.

Patients receiving SUSTIVA should be alerted to the potential for additive central
nervous system effects when SUSTIVA is used concomitantly with alcohol or
psychoactive drugs.

Patients who experience central nervous system symptoms such as dizziness, impaired
concentration, and/or drowsiness should avoid potentially hazardous tasks such as driving
or operating machinery.

280 Drug Interactions: Concomitant use of SUSTIVA and St. John's wort (Hypericum 281 John's wort-containing products is *perforatum*) or St. not recommended. 282 Coadministration of non-nucleoside reverse transcriptase inhibitors (NNRTIs), including 283 SUSTIVA, with St. John's wort is expected to substantially decrease NNRTI 284 concentrations and may result in suboptimal levels of efavirenz and lead to loss of 285 virologic response and possible resistance to efavirenz or to the class of NNRTIs.

286 Reproductive Risk Potential: Pregnancy Category D. Efavirenz may cause fetal 287 harm when administered during the first trimester to a pregnant woman. Pregnancy 288 should be avoided in women receiving SUSTIVA. Barrier contraception should always 289 be used in combination with other methods of contraception (eg, oral or other hormonal 290 contraceptives). Women of childbearing potential should undergo pregnancy testing 291 before initiation of SUSTIVA. If this drug is used during the first trimester of pregnancy, 292 or if the patient becomes pregnant while taking this drug, the patient should be apprised 293 of the potential harm to the fetus.

294 There are no adequate and well-controlled studies in pregnant women. SUSTIVA should 295 be used during pregnancy only if the potential benefit justifies the potential risk to the 296 fetus, such as in pregnant women without other therapeutic options. As of July 2005, the 297 Antiretroviral Pregnancy Registry has received prospective reports of 282 pregnancies 298 exposed to efavirenz-containing regimens, nearly all of which were first-trimester 299 exposures (277 pregnancies). Birth defects occurred in 5 of 228 live births (first-trimester 300 exposure) and 1 of 14 live births (second/third-trimester exposure). None of these 301 prospectively reported defects were neural tube defects. However, there have been four 302 retrospective reports of findings consistent with neural tube defects, including 303 meningomyelocele. All mothers were exposed to efavirenz-containing regimens in the 304 first trimester. Although a causal relationship of these events to the use of SUSTIVA has 305 not been established, similar defects have been observed in preclinical studies of 306 efavirenz.

307 Malformations have been observed in 3 of 20 fetuses/infants from efavirenz-treated 308 cynomolgus monkeys (versus 0 of 20 concomitant controls) in a developmental toxicity

309 study. The pregnant monkeys were dosed throughout pregnancy (postcoital days 20-150) 310 with efavirenz 60 mg/kg daily, a dose which resulted in plasma drug concentrations 311 similar to those in humans given 600 mg/day of SUSTIVA. Anencephaly and unilateral 312 anophthalmia were observed in one fetus, microophthalmia was observed in another 313 fetus, and cleft palate was observed in a third fetus. Efavirenz crosses the placenta in 314 cynomolgus monkeys and produces fetal blood concentrations similar to maternal blood concentrations. Efavirenz has been shown to cross the placenta in rats and rabbits and 315 316 produces fetal blood concentrations of efavirenz similar to maternal concentrations. An 317 increase in fetal resorptions was observed in rats at efavirenz doses that produced peak 318 plasma concentrations and AUC values in female rats equivalent to or lower than those 319 achieved in humans given 600 mg once daily of SUSTIVA. Efavirenz produced no 320 reproductive toxicities when given to pregnant rabbits at doses that produced peak plasma 321 concentrations similar to and AUC values approximately half of those achieved in 322 humans given 600 mg once daily of SUSTIVA.

Antiretroviral Pregnancy Registry: To monitor fetal outcomes of pregnant women
exposed to SUSTIVA, an Antiretroviral Pregnancy Registry has been established.
Physicians are encouraged to register patients by calling (800) 258-4263.

#### 326 **PRECAUTIONS**

#### 327 General

Skin Rash: In controlled clinical trials, 26% (266/1008) of patients treated with 600 mg 328 329 SUSTIVA experienced new-onset skin rash compared with 17% (111/635) of patients 330 treated in control groups. Rash associated with blistering, moist desquamation, or 331 ulceration occurred in 0.9% (9/1008) of patients treated with SUSTIVA. The incidence of 332 Grade 4 rash (eg, erythema multiforme, Stevens-Johnson syndrome) in patients treated 333 with SUSTIVA in all studies and expanded access was 0.1%. The median time to onset 334 of rash in adults was 11 days and the median duration, 16 days. The discontinuation rate 335 for rash in clinical trials was 1.7% (17/1008). SUSTIVA should be discontinued in patients developing severe rash associated with blistering, desquamation, mucosal 336 337 involvement, or fever. Appropriate antihistamines and/or corticosteroids may improve 338 the tolerability and hasten the resolution of rash.

Rash was reported in 26 of 57 pediatric patients (46%) treated with SUSTIVA capsules. One pediatric patient experienced Grade 3 rash (confluent rash with fever), and two patients had Grade 4 rash (erythema multiforme). The median time to onset of rash in pediatric patients was 8 days. Prophylaxis with appropriate antihistamines prior to initiating therapy with SUSTIVA in pediatric patients should be considered (see **ADVERSE REACTIONS**).

Liver Enzymes: In patients with known or suspected history of hepatitis B or C infection and in patients treated with other medications associated with liver toxicity, monitoring of liver enzymes is recommended. In patients with persistent elevations of serum transaminases to greater than five times the upper limit of the normal range, the benefit of continued therapy with SUSTIVA needs to be weighed against the unknown risks of significant liver toxicity (see ADVERSE REACTIONS: Laboratory Abnormalities).

Because of the extensive cytochrome P450-mediated metabolism of efavirenz and limited clinical experience in patients with hepatic impairment, caution should be exercised in administering SUSTIVA to these patients.

355 Convulsions: Convulsions have been observed infrequently in patients receiving 356 efavirenz, generally in the presence of known medical history of seizures. Patients who 357 are receiving concomitant anticonvulsant medications primarily metabolized by the liver, 358 such as phenytoin and phenobarbital, may require periodic monitoring of plasma levels 359 (see PRECAUTIONS: Drug Interactions). Caution must be taken in any patient with a 360 history of seizures.

361 *Animal toxicology*: Nonsustained convulsions were observed in 6 of 20 monkeys 362 receiving efavirenz at doses yielding plasma AUC values 4- to 13-fold greater than those 363 in humans given the recommended dose.

364 Cholesterol: Monitoring of cholesterol and triglycerides should be considered in
365 patients treated with SUSTIVA (see ADVERSE REACTIONS).

Fat Redistribution: Redistribution/accumulation of body fat including central obesity,
 dorsocervical fat enlargement (buffalo hump), peripheral wasting, facial wasting, breast
 enlargement, and "cushingoid appearance" have been observed in patients receiving

antiretroviral therapy. The mechanism and long-term consequences of these events arecurrently unknown. A causal relationship has not been established.

Immune Reconstitution Syndrome: Immune reconstitution syndrome has been reported in patients treated with combination antiretroviral therapy, including SUSTIVA. During the initial phase of combination antiretroviral treatment, patients whose immune system responds may develop an inflammatory response to indolent or residual opportunistic infections (such as *Mycobacterium avium* infection, cytomegalovirus, *Pneumocystis jiroveci* pneumonia [PCP], or tuberculosis), which may necessitate further evaluation and treatment.

#### 378 Information for Patients

A statement to patients and healthcare providers is included on the product's bottle
labels: ALERT: Find out about medicines that should NOT be taken with
SUSTIVA. A Patient Package Insert (PPI) for SUSTIVA is available for patient
information.

Patients should be informed that SUSTIVA is not a cure for HIV-1 infection and that they may continue to develop opportunistic infections and other complications associated with HIV-1 disease. Patients should be told that there are currently no data demonstrating that SUSTIVA therapy can reduce the risk of transmitting HIV to others through sexual contact or blood contamination.

Patients should be advised to take SUSTIVA every day as prescribed. SUSTIVA must always be used in combination with other antiretroviral drugs. Patients should be advised to take SUSTIVA on an empty stomach, preferably at bedtime. Taking SUSTIVA with food increases efavirenz concentrations and may increase the frequency of adverse events. Dosing at bedtime may improve the tolerability of nervous system symptoms (see **ADVERSE REACTIONS** and **DOSAGE AND ADMINISTRATION**). Patients should remain under the care of a physician while taking SUSTIVA.

Patients should be informed that central nervous system symptoms including dizziness, insomnia, impaired concentration, drowsiness, and abnormal dreams are commonly reported during the first weeks of therapy with SUSTIVA. Dosing at bedtime may improve the tolerability of these symptoms, and these symptoms are likely to improve

with continued therapy. Patients should be alerted to the potential for additive central
nervous system effects when SUSTIVA is used concomitantly with alcohol or
psychoactive drugs. Patients should be instructed that if they experience these symptoms
they should avoid potentially hazardous tasks such as driving or operating machinery (see
WARNINGS: Nervous System Symptoms). In clinical trials, patients who develop
central nervous system symptoms were not more likely to subsequently develop
psychiatric symptoms (see WARNINGS: Psychiatric Symptoms).

406 Patients should also be informed that serious psychiatric symptoms including severe 407 depression, suicide attempts, aggressive behavior, delusions, paranoia, and psychosis-like 408 symptoms have also been infrequently reported in patients receiving SUSTIVA. Patients 409 should be informed that if they experience severe psychiatric adverse experiences they 410 should seek immediate medical evaluation to assess the possibility that the symptoms may be related to the use of SUSTIVA, and if so, to determine whether discontinuation of 411 412 SUSTIVA may be required. Patients should also inform their physician of any history of 413 mental illness or substance abuse (see WARNINGS: Psychiatric Symptoms).

414 Patients should be informed that another common side effect is rash. These rashes usually 415 go away without any change in treatment. In a small number of patients, rash may be 416 serious. Patients should be advised that they should contact their physician promptly if 417 they develop a rash.

418 Women receiving SUSTIVA should be instructed to avoid pregnancy (see WARNINGS: 419 **Reproductive Risk Potential**). A reliable form of barrier contraception should always be 420 used in combination with other methods of contraception, including oral or other 421 hormonal contraception, because the effects of efavirenz on hormonal contraceptives are 422 not fully characterized. Women should be advised to notify their physician if they 423 become pregnant while taking SUSTIVA. If this drug is used during the first trimester of 424 pregnancy, or if the patient becomes pregnant while taking this drug, she should be 425 apprised of the potential harm to the fetus.

SUSTIVA may interact with some drugs; therefore, patients should be advised to report
to their doctor the use of any other prescription, nonprescription medication, or herbal
products, particularly St. John's wort.

 $20 \ of \ 45$ 

- Patients should be informed that redistribution or accumulation of body fat may occur inpatients receiving antiretroviral therapy and that the cause and long-term health effects of
- 431 these conditions are not known at this time.

# 432 Drug Interactions (see also CONTRAINDICATIONS and 433 CLINICAL PHARMACOLOGY: Drug Interactions)

Efavirenz has been shown *in vivo* to induce CYP3A4. Other compounds that are substrates of CYP3A4 may have decreased plasma concentrations when coadministered with SUSTIVA. *In vitro* studies have demonstrated that efavirenz inhibits 2C9, 2C19, and 3A4 isozymes in the range of observed efavirenz plasma concentrations. Coadministration of efavirenz with drugs primarily metabolized by these isozymes may result in altered plasma concentrations of the coadministered drug. Therefore, appropriate dose adjustments may be necessary for these drugs.

441 Drugs which induce CYP3A4 activity (eg, phenobarbital, rifampin, rifabutin) would be 442 expected to increase the clearance of efavirenz resulting in lowered plasma 443 concentrations. Drug interactions with SUSTIVA are summarized in Tables 5 and 6. The 444 tables include potentially significant interactions, but are not all inclusive.

Drug Class: Drug Name	<b>Clinical Comment</b>
Antifungal: voriconazole	CONTRAINDICATED at standard doses. SUSTIVA significantly decreases voriconazole plasma concentrations, and coadministration may decrease the therapeutic effectiveness of voriconazole. Also, voriconazole significantly increases SUSTIVA plasma concentrations, which may increase the risk of SUSTIVA- associated side effects. When voriconazole is coadministered with SUSTIVA, voriconazole maintenance dose should be increased to 400 mg every 12 hours and SUSTIVA dose should be decreased to 300 mg once daily using the capsule formulation. SUSTIVA tablets should not be broken. (See CLINICAL PHARMACOLOGY, Tables 1 and 2; CONTRAINDICATIONS; and DOSAGE AND ADMINISTRATION: Dosage Adjustment.)
Antihistamine: astemizole	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as cardiac arrhythmias.
Antimigraine: ergot derivatives (dihydroergotamine, ergonovine, ergotamine, methylergonovine)	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as acute ergot toxicity characterized by peripheral vasospasm and ischemia of the extremities and other tissues.
Benzodiazepines: midazolam, triazolam	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as prolonged or increased sedation or respiratory depression.
Calcium channel blocker: bepridil	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as cardiac arrhythmias.
GI motility agent: cisapride	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as cardiac arrhythmias.
Neuroleptic: pimozide	CONTRAINDICATED due to potential for serious and/or life-threatening reactions such as cardiac arrhythmias.
St. John's wort (Hypericum perforatum)	Expected to substantially decrease plasma levels of efavirenz; has not been studied in combination with SUSTIVA.

Table 5:Drugs That Are Contraindicated or Not Recommended for Use With<br/>SUSTIVA

445

Concomitant Drug Class: Drug Name	Effect on Concentration of SUSTIVA or Concomitant Drug	Clinical Comment
Antiretroviral agents		
Protease inhibitor: Amprenavir	↓ amprenavir	SUSTIVA has the potential to decrease serum concentrations of amprenavir.
Protease inhibitor: Fosamprenavir calcium	↓ amprenavir	Fosamprenavir (unboosted): Appropriate doses of the combinations with respect to safety and efficacy have not been established.
		Fosamprenavir/ritonavir: An additional 100 mg/day (300 mg total) of ritonavir is recommended when SUSTIVA is administered with fosamprenavir/ritonavir once daily. No change in the ritonavir dose is required when SUSTIVA is administered with fosamprenavir plus ritonavir twice daily.
Protease inhibitor: Atazanavir	↓ atazanavir <sup>a</sup>	When coadministered with SUSTIVA in treatment-naive patients, the recommended dose of atazanavir is 300 mg with ritonavir 100 mg and SUSTIVA 600 mg (all once daily). Dosing recommendations for SUSTIVA and atazanavir in treatment-experienced patients have not been established.
Protease inhibitor: Indinavir	↓ indinavir <sup>a</sup>	The optimal dose of indinavir, when given in combination with SUSTIVA, is not known. Increasing the indinavir dose to 1000 mg every 8 hours does not compensate for the increased indinavir metabolism due to SUSTIVA. When indinavir at an increased dose (1000 mg every 8 hours) was given with SUSTIVA (600 mg once daily), the indinavir AUC and C <sub>min</sub> were decreased on average by 33-46% and 39-57%, respectively, compared to when indinavir (800 mg every 8 hours) was given alone.
Protease inhibitor: Lopinavir/ritonavir	$\downarrow$ lopinavir <sup>a</sup>	A dose increase of lopinavir/ritonavir to 533/133 mg (4 capsules or 6.5 mL) twice daily taken with food is recommended when used in combination with SUSTIVA
Protease inhibitor: Ritonavir	↑ ritonavir <sup>a</sup> ↑ efavirenz <sup>a</sup>	When ritonavir 500 mg q12h was coadministered with SUSTIVA 600 mg once daily, the combination was associated with a higher frequency of adverse clinical experiences (eg, dizziness, nausea, paresthesia) and laboratory abnormalities (elevated liver enzymes). Monitoring of liver enzymes is recommended when SUSTIVA is used in combination with ritonavir.

# Table 6:Established<sup>a</sup> and Other Potentially Significant<sup>b</sup> Drug Interactions:<br/>Alteration in Dose or Regimen May Be Recommended Based on Drug<br/>Interaction Studies or Predicted Interaction

446

Concomitant Drug Class: Drug Name	Effect on Concentration of SUSTIVA or Concomitant Drug	Clinical Comment
Protease inhibitor: Saquinavir	↓ saquinavir <sup>a</sup>	Should not be used as sole protease inhibitor in combination with SUSTIVA.
Other agents		
Anticoagulant: Warfarin	↑ or ↓ warfarin	Plasma concentrations and effects potentially increased or decreased by SUSTIVA.
Anticonvulsants: Carbamazepine	↓ carbamazepine <sup>a</sup> ↓ efavirenz <sup>a</sup>	There are insufficient data to make a dose recommendation for efavirenz. Alternative anticonvulsant treatment should be used.
Phenytoin Phenobarbital	↓ anticonvulsant ↓ efavirenz	Potential for reduction in anticonvulsant and/or efavirenz plasma levels; periodic monitoring of anticonvulsant plasma levels should be conducted.
Antidepressant: Sertraline	$\downarrow$ sertraline <sup>a</sup>	Increased in sertraline dose should be guided by clinical response.
Antifungals:		
Itraconazole	↓ itraconazole <sup>a</sup> ↓ hydroxyitraconazole <sup>a</sup>	Since no dose recommendation for itraconazole can be made, alternative antifungal treatment should be considered.
Ketoconazole	↓ ketoconazole	Drug interaction studies with SUSTIVA and ketoconazole have not been conducted. SUSTIVA has the potential to decrease plasma concentrations of ketoconazole. (See Table 5 for guidance on coadministration with adjusted doses of voriconazole.)
Anti-infective: Clarithromycin	↓ clarithromycin <sup>a</sup> ↑ 14-OH metabolite <sup>a</sup>	Plasma concentrations decreased by SUSTIVA; clinical significance unknown. In uninfected volunteers, 46% developed rash while receiving SUSTIVA and clarithromycin. No dose adjustment of SUSTIVA is recommended when given with clarithromycin. Alternatives to clarithromycin, such as azithromycin, should be considered (see <b>Other Drugs</b> , following table). Other macrolide antibiotics, such as erythromycin, have not been studied in combination with SUSTIVA.
Antimycobacterial: Rifabutin	↓ rifabutin <sup>a</sup>	Increase daily dose of rifabutin by 50%. Consider doubling the rifabutin dose in regimens where rifabutin is given 2 or 3 times a week.
Antimycobacterial: Rifampin	$\downarrow$ efavirenz <sup>a</sup>	Clinical significance of reduced efavirenz concentrations is unknown. Dosing recommendations for concomitant use of SUSTIVA and rifampin have not been established.

# Table 6:Established<sup>a</sup> and Other Potentially Significant<sup>b</sup> Drug Interactions:<br/>Alteration in Dose or Regimen May Be Recommended Based on Drug<br/>Interaction Studies or Predicted Interaction

Concomitant Drug Class: Drug Name	Effect on Concentration of SUSTIVA or Concomitant Drug	Clinical Comment
Calcium channel blockers: Diltiazem	↓ diltiazem <sup>a</sup> ↓ desacetyl diltiazem <sup>a</sup> ↓ N-monodesmethyl diltiazem <sup>a</sup>	Diltiazem dose adjustments should be guided by clinical response (refer to the complete prescribing information for diltiazem). No dose adjustment of efavirenz is necessary when administered with diltiazem.
Others (eg, felodipine, nicardipine, nifedipine, verapamil)	↓ calcium channel blocker	No data are available on the potential interactions of efavirenz with other calcium channel blockers that are substrates of the CYP3A4 enzyme. The potential exists for reduction in plasma concentrations of the calcium channel blocker. Dose adjustments should be guided by clinical response (refer to the complete prescribing information for the calcium channel blocker).
HMG-CoA reductase inhibitors: Atorvastatin Pravastatin Simvastatin	↓ atorvastatin <sup>a</sup> ↓ pravastatin <sup>a</sup> ↓ simvastatin <sup>a</sup>	Plasma concentrations of atorvastatin, pravastatin, and simvastatin decreased. Consult the complete prescribing information for the HMG-CoA reductase inhibitor for guidance on individualizing the dose.
Narcotic analgesic: Methadone	↓ methadone <sup>a</sup>	Coadministration in HIV-infected individuals with a history of injection drug use resulted in decreased plasma levels of methadone and signs of opiate withdrawal. Methadone dose was increased by a mean of 22% to alleviate withdrawal symptoms. Patients should be monitored for signs of withdrawal and their methadone dose increased as required to alleviate withdrawal symptoms.
Oral contraceptive: Ethinyl estradiol	↑ ethinyl estradiol <sup>a</sup>	Plasma concentrations increased by SUSTIVA; clinical significance unknown. The potential interaction of efavirenz with oral contraceptives has not been fully characterized. A reliable method of barrier contraception should be used in addition to oral contraceptives.

# Table 6:Established<sup>a</sup> and Other Potentially Significant<sup>b</sup> Drug Interactions:<br/>Alteration in Dose or Regimen May Be Recommended Based on Drug<br/>Interaction Studies or Predicted Interaction

<sup>a</sup> See **CLINICAL PHARMACOLOGY**, Tables 1 and 2 for magnitude of established interactions.

<sup>b</sup> This table is not all-inclusive.

447

448 Other Drugs: Based on the results of drug interaction studies (see Tables 1 and 2), no 449 dosage adjustment is recommended when SUSTIVA (efavirenz) is given with the 450 following: aluminum/magnesium hydroxide antacids, azithromycin, cetirizine, 451 famotidine, fluconazole, lamivudine, lorazepam, nelfinavir, paroxetine, tenofovir 452 disoproxil fumarate, and zidovudine.

453 Specific drug interaction studies have not been performed with SUSTIVA and NRTIS 454 other than lamivudine and zidovudine. Clinically significant interactions would not be 455 expected since the NRTIs are metabolized via a different route than efavirenz and would 456 be unlikely to compete for the same metabolic enzymes and elimination pathways.

#### 457 Carcinogenesis, Mutagenesis, and Impairment of Fertility

458 Long-term carcinogenicity studies in mice and rats were carried out with efavirenz. Mice 459 were dosed with 0, 25, 75, 150, or 300 mg/kg/day for 2 years. Incidences of 460 hepatocellular adenomas and carcinomas and pulmonary alveolar/bronchiolar adenomas 461 were increased above background in females. No increases in tumor incidence above 462 background were seen in males. In studies in which rats were administered efavirenz at 463 doses of 0, 25, 50, or 100 mg/kg/day for 2 years, no increases in tumor incidence above 464 background were observed. The systemic exposure (based on AUCs) in mice was 465 approximately 1.7-fold that in humans receiving the 600-mg/day dose. The exposure in rats was lower than that in humans. The mechanism of the carcinogenic potential is 466 467 unknown. However, in genetic toxicology assays, efavirenz showed no evidence of mutagenic or clastogenic activity in a battery of in vitro and in vivo studies. These 468 469 included bacterial mutation assays in S. typhimurium and E. coli, mammalian mutation 470 assays in Chinese hamster ovary cells, chromosome aberration assays in human 471 peripheral blood lymphocytes or Chinese hamster ovary cells, and an in vivo mouse bone 472 marrow micronucleus assay. Given the lack of genotoxic activity of efavirenz, the 473 relevance to humans of neoplasms in efavirenz-treated mice is not known.

Efavirenz did not impair mating or fertility of male or female rats, and did not affect sperm of treated male rats. The reproductive performance of offspring born to female rats given efavirenz was not affected. As a result of the rapid clearance of efavirenz in rats, systemic drug exposures achieved in these studies were equivalent to or below those achieved in humans given therapeutic doses of efavirenz.

#### 479 **Pregnancy**

480 **Pregnancy Category D:** See WARNINGS: Reproductive Risk Potential.

#### 481 Nursing Mothers

482 The Centers for Disease Control and Prevention recommend that HIV-infected 483 mothers not breast-feed their infants to avoid risking postnatal transmission of HIV. 484 Although it is not known if efavirenz is secreted in human milk, efavirenz is secreted into 485 the milk of lactating rats. Because of the potential for HIV transmission and the potential 486 for serious adverse effects in nursing infants, mothers should be instructed not to 487 breast-feed if they are receiving SUSTIVA.

#### 488 **Pediatric Use**

489 ACTG 382 is an ongoing, open-label study in 57 NRTI-experienced pediatric patients to 490 characterize the safety, pharmacokinetics, and antiviral activity of SUSTIVA in 491 combination with nelfinavir (20-30 mg/kg TID) and NRTIs. Mean age was 8 years (range 492 3-16). SUSTIVA has not been studied in pediatric patients below 3 years of age or who 493 weigh less than 13 kg. At 48 weeks, the type and frequency of adverse experiences was 494 generally similar to that of adult patients with the exception of a higher incidence of rash, 495 which was reported in 46% (26/57) of pediatric patients compared to 26% of adults, and a 496 higher frequency of Grade 3 or 4 rash reported in 5% (3/57) of pediatric patients 497 compared to 0.9% of adults (see ADVERSE REACTIONS, Table 8).

The starting dose of SUSTIVA was 600 mg once daily adjusted to body size, based on weight, targeting AUC levels in the range of 190-380  $\mu$ M•h. The pharmacokinetics of efavirenz in pediatric patients were similar to the pharmacokinetics in adults who received 600-mg daily doses of SUSTIVA. In 48 pediatric patients receiving the equivalent of a 600-mg dose of SUSTIVA, steady-state C<sub>max</sub> was 14.2 ± 5.8  $\mu$ M (mean ± SD), steady-state C<sub>min</sub> was 5.6 ± 4.1  $\mu$ M, and AUC was 218 ± 104  $\mu$ M•h.

#### 504 Geriatric Use

505 Clinical studies of SUSTIVA did not include sufficient numbers of subjects aged 65 506 years and over to determine whether they respond differently from younger subjects. In 507 general, dose selection for an elderly patient should be cautious, reflecting the greater 508 frequency of decreased hepatic, renal, or cardiac function and of concomitant disease or 509 other therapy.

# 510 ADVERSE REACTIONS

511 The most significant adverse events observed in patients treated with SUSTIVA are 512 nervous system symptoms, psychiatric symptoms, and rash. Unless otherwise specified, 513 the analyses described below included 1008 patients treated with regimens containing 514 SUSTIVA and 635 patients treated with a control regimen in controlled trials.

515 **Nervous System Symptoms:** Fifty-three percent of patients receiving SUSTIVA 516 reported central nervous system symptoms (see WARNINGS: Nervous System 517 **Symptoms**). Table 7 lists the frequency of the symptoms of different degrees of severity 518 and gives the discontinuation rates in clinical trials for one or more of the following 519 nervous system symptoms: dizziness, insomnia, impaired concentration, somnolence, 520 abnormal dreaming, euphoria, confusion, agitation, amnesia, hallucinations, stupor, 521 abnormal thinking, and depersonalization. The frequencies of specific central and 522 peripheral nervous system symptoms are provided in Table 9.

Percent of Patients with:	SUSTIVA 600 mg Once Daily (n=1008) %	Control Groups (n=635) %
Symptoms of any severity	52.7	24.6
Mild symptoms <sup>c</sup>	33.3	15.6
Moderate symptoms <sup>d</sup>	17.4	7.7
Severe symptoms <sup>e</sup>	2.0	1.3
Treatment discontinuation as a result of symptoms	2.1	1.1

 Table 7: Percent of Patients with One or More Selected Nervous System Symptoms<sup>a,b</sup>

523 <sup>a</sup> Includes events reported regardless of causality.

524 <sup>b</sup> Data from Study 006 and three Phase 2/3 studies.

- 525 <sup>c</sup> "Mild" = Symptoms which do not interfere with patient's daily activities.
- 526 <sup>d</sup> "Moderate" = Symptoms which may interfere with daily activities.

527 <sup>e</sup> "Severe" = Events which interrupt patient's usual daily activities.

**Psychiatric Symptoms:** Serious psychiatric adverse experiences have been reported in 528 529 patients treated with SUSTIVA. In controlled trials, the frequency of specific serious 530 psychiatric symptoms among patients who received SUSTIVA or control regimens, respectively, were severe depression (2.4%, 0.9%), suicidal ideation (0.7%, 0.3%), nonfatal 531 532 suicide attempts (0.5%, 0), aggressive behavior (0.4%, 0.5%), paranoid reactions (0.4%, 533 0.3%), and manic reactions (0.2%, 0.3%) (see WARNINGS: Psychiatric Symptoms). 534 Additional psychiatric symptoms observed at a frequency of >2% among patients treated 535 with SUSTIVA or control regimens, respectively, in controlled clinical trials were depression (19%, 16%), anxiety (13%, 9%), and nervousness (7%, 2%). 536

Skin Rash: Rashes are usually mild-to-moderate maculopapular skin eruptions that 537 538 occur within the first 2 weeks of initiating therapy with SUSTIVA. In most patients, rash 539 resolves with continuing SUSTIVA therapy within one month. SUSTIVA can be 540 reinitiated in patients interrupting therapy because of rash. Use of appropriate 541 antihistamines and/or corticosteroids may be considered when SUSTIVA is restarted. 542 SUSTIVA should be discontinued in patients developing severe rash associated with 543 blistering, desquamation, mucosal involvement, or fever. The frequency of rash by NCI 544 grade and the discontinuation rates as a result of rash are provided in Table 8.

Percent of Patients with:	Description of Rash Grade <sup>C</sup>	SUSTIVA 600 mg Once Daily Adults (n=1008) %	SUSTIVA Pediatric Patients (n=57) %	Control Groups Adults (n= 635) %
Rash of any grade		26.3	45.6	17.5
Grade 1 rash	Erythema, pruritus	10.7	8.8	9.8
Grade 2 rash	Diffuse maculopapular rash, dry desquamation	14.7	31.6	7.4
Grade 3 rash	Vesiculation, moist desquamation, ulceration	0.8	1.8	0.3
Grade 4 rash	Erythema multiforme, Stevens-Johnson syndrome, toxic epidermal necrolysis, necrosis requiring surgery, exfoliative dermatitis	0.1	3.5	0.0
Treatment discontinuation as a result of rash	—	1.7	8.8	0.3

## Table 8: Percent of Patients with Treatment-Emergent Rash<sup>a,b</sup>

<sup>a</sup> Includes events reported regardless of causality.

546 <sup>b</sup> Data from Study 006 and three Phase 2/3 studies.

547 <sup>c</sup> NCI Grading System.

548 As seen in Table 8, rash is more common in pediatric patients and more often of higher 549 grade (ie, more severe) (see **PRECAUTIONS: General**).

550 Experience with SUSTIVA (efavirenz) in patients who discontinued other antiretroviral 551 agents of the NNRTI class is limited. Nineteen patients who discontinued nevirapine 552 because of rash have been treated with SUSTIVA. Nine of these patients developed mild-553 to-moderate rash while receiving therapy with SUSTIVA, and two of these patients 554 discontinued because of rash.

Pancreatitis has been reported, although a causal relationship with efavirenz has not been
 established. Asymptomatic increases in serum amylase levels were observed in a

significantly higher number of patients treated with efavirenz 600 mg than in control

- 558 patients (see **ADVERSE REACTIONS: Laboratory Abnormalities**).
- 559 Selected clinical adverse experiences of moderate or severe intensity observed in  $\geq 2\%$  of
- 560 SUSTIVA-treated patients in two controlled clinical trials are presented in Table 9.

 $30 \ of \ 45$ 

Adverse Events		Study 006 NRTI-, and l itor-Naive Pat		Study ACTG 364 NRTI-experienced, NNRTI- and Protease Inhibitor-Naive Patients			
	SUSTIVA <sup>b</sup> + ZDV/LAM (n=412)	SUSTIVA <sup>b</sup> + Indinavir (n=415)	Indinavir + ZDV/LAM (n=401)	SUSTIVA <sup>b</sup> + Nelfinavir + NRTIs (n=64)	SUSTIVA <sup>b</sup> + NRTIs (n=65)	Nelfinavir + NRTIs (n=66)	
	180 weeks <sup>c</sup>	102 weeks <sup>c</sup>	76 weeks <sup>c</sup>	71.1 weeks <sup>c</sup>	70.9 weeks <sup>c</sup>	62.7 weeks <sup>c</sup>	
Body as a Whole							
Fatigue	8%	5%	9%	0	2%	3%	
Pain	1%	2%	8%	13%	6%	17%	
<b>Central and Peripheral</b>	Nervous Syste	m					
Dizziness	9%	9%	2%	2%	6%	6%	
Headache	8%	5%	3%	5%	2%	3%	
Insomnia	7%	7%	2%	0	0	2%	
Concentration impaired	5%	3%	<1%	0	0	0	
Abnormal dreams	3%	1%	0				
Somnolence	2%	2%	<1%	0	0	0	
Anorexia	1%	<1%	<1%	0	2%	2%	
Gastrointestinal							
Nausea	10%	6%	24%	3%	2%	2%	
Vomiting	6%	3%	14%				
Diarrhea	3%	5%	6%	14%	3%	9%	
Dyspepsia	4%	4%	6%	0	0	2%	
Abdominal pain	2%	2%	5%	3%	3%	3%	
Psychiatric							
Anxiety	2%	4%	<1%	_			
Depression	5%	4%	<1%	3%	0	5%	
Nervousness	2%	2%	0	2%	0	2%	
Skin & Appendages							
Rash	11%	16%	5%	9%	5%	9%	
Pruritus	<1%	1%	1%	9%	5%	9%	

#### Table 9: Selected Treatment-Emergent<sup>a</sup> Adverse Events of Moderate or Severe Intensity Reported in ≥2% of SUSTIVA-Treated Patients in Studies 006 and ACTG 364

<sup>a</sup> Includes adverse events at least possibly related to study drug or of unknown relationship for Study 006.
 Includes all adverse events regardless of relationship to study drug for Study ACTG 364.

563 <sup>b</sup> SUSTIVA provided as 600 mg once daily.

564 <sup>c</sup> Median duration of treatment.

565 — = Not Specified.

566 ZDV = zidovudine, LAM=lamivudine.

567 Clinical adverse experiences observed in  $\geq 10\%$  of 57 pediatric patients aged 3 to 16 years

568 who received SUSTIVA capsules, nelfinavir, and one or more NRTIs were: rash (46%),

- 569 diarrhea/loose stools (39%), fever (21%), cough (16%), dizziness/lightheaded/fainting
- 570 (16%), ache/pain/discomfort (14%), nausea/vomiting (12%), and headache (11%). The
- 571 incidence of nervous system symptoms was 18% (10/57). One patient experienced Grade
- 572 3 rash, two patients had Grade 4 rash, and five patients (9%) discontinued because of rash
- 573 (see also **PRECAUTIONS: Skin Rash** and **Pediatric Use**).

#### 574 **Postmarketing Experience**

575 Body as a Whole: allergic reactions, asthenia, redistribution/accumulation of body fat

#### 576 (see **PRECAUTIONS: Fat Redistribution**)

- 577 Central and Peripheral Nervous System: abnormal coordination, ataxia, convulsions,
- 578 hypoesthesia, paresthesia, neuropathy, tremor
- 579 Endocrine: gynecomastia
- 580 *Gastrointestinal:* constipation, malabsorption
- 581 *Cardiovascular:* flushing, palpitations
- 582 *Liver and Biliary System:* hepatic enzyme increase, hepatic failure, hepatitis
- 583 Metabolic and Nutritional: hypercholesterolemia, hypertriglyceridemia
- 584 *Musculoskeletal:* arthralgia, myalgia, myopathy
- 585 *Psychiatric:* aggressive reactions, agitation, delusions, emotional lability, mania, 586 neurosis, paranoia, psychosis, suicide
- 587 Respiratory: dyspnea
- 588 Skin and Appendages: erythema multiforme, nail disorders, photoallergic dermatitis, skin
- 589 discoloration, Stevens-Johnson syndrome
- 590 Special Senses: abnormal vision, tinnitus

#### 591 Laboratory Abnormalities

- 592 Selected Grade 3-4 laboratory abnormalities reported in ≥2% of SUSTIVA-treated
- 593 patients in two clinical trials are presented in Table 10.

		Study 006 LAM-, NNRTI-, and Protease Inhibitor-Naive Patients			Study ACTG 364 NRTI-experienced, NNRTI- and Protease Inhibitor-Naive Patients		
Variable	Limit	SUSTIVA <sup>a</sup> + ZDV/LAM (n=412) 180 weeks <sup>b</sup>	a + Indinavir (n=415) 102 weeks	Indinavir + ZDV/LAM (n=401) 76 weeks	a + Nelfinavir + NRTIs (n=64) 71.1 weeks	SUSTIVA <sup>a</sup> + NRTIs (n=65) 70.9 weeks <sup>b</sup>	Nelfinavir + NRTIs (n=66) 62.7 weeks
Chemistry							
ALT	>5 x ULN	5%	8%	5%	2%	6%	3%
AST	>5 x ULN	5%	6%	5%	6%	8%	8%
GGT <sup>c</sup>	>5 x ULN	8%	7%	3%	5%	0	5%
Amylase	>2 x ULN	4%	4%	1%	0	6%	2%
Glucose	>250 mg/dL	3%	3%	3%	5%	2%	3%
d Triglycerides	$\geq$ 751 mg/dL	9%	6%	6%	11%	8%	17%
Hematology							
Neutrophils	<750/mm <sup>3</sup>	10%	3%	5%	2%	3%	2%

Table 10: Selected Grade 3-4 Laboratory Abnormalities Reported in ≥2% of SUSTIVA-Treated Patients in Studies 006 and ACTG 364

<sup>a</sup> SUSTIVA provided as 600 mg once daily.

<sup>b</sup> Median duration of treatment.

<sup>c</sup> Isolated elevations of GGT in patients receiving SUSTIVA may reflect enzyme induction not associated with liver toxicity.

# <sup>d</sup> Nonfasting.

ZDV = zidovudine, LAM = lamivudine, ULN = Upper limit of normal, ALT = alanine aminotransferase, AST = aspartate aminotransferase, GGT = gamma-glutamyltransferase.

594 Liver function tests should be monitored in patients with a history of hepatitis B and/or C.

595 In the long-term data set from Study 006, 137 patients treated with SUSTIVA-containing

regimens (median duration of therapy, 68 weeks) and 84 treated with a control regimen

- 597 (median duration, 56 weeks) were seropositive at screening for hepatitis B (surface
- 598 antigen positive) and/or C (hepatitis C antibody positive). Among these co-infected

patients, elevations in AST to greater than five times ULN developed in 13% of patients in the SUSTIVA arms and 7% of those in the control arm, and elevations in ALT to greater than five times ULN developed in 20% of patients in the SUSTIVA arms and 7% of patients in the control arm. Among co-infected patients, 3% of those treated with SUSTIVA-containing regimens and 2% in the control arm discontinued from the study because of liver or biliary system disorders (see **PRECAUTIONS: General**).

605 Lipids: Increases from baseline in total cholesterol of 10-20% have been observed in some uninfected volunteers receiving SUSTIVA. In patients treated with SUSTIVA + 606 607 zidovudine + lamivudine, increases from baseline in nonfasting total cholesterol and HDL of approximately 20% and 25%, respectively, were observed. In patients treated 608 609 with SUSTIVA + indinavir, increases from baseline in nonfasting cholesterol and HDL 610 of approximately 40% and 35%, respectively, were observed. Nonfasting total cholesterol levels >240 mg/dL and >300 mg/dL were reported in 34% and 9%, respectively, of 611 612 patients treated with SUSTIVA + zidovudine + lamivudine; 54% and 20%, respectively, 613 of patients treated with SUSTIVA + indinavir; and 28% and 4%, respectively, of patients treated with indinavir + zidovudine + lamivudine. The effects of SUSTIVA on 614 615 triglycerides and LDL were not well characterized since samples were taken from nonfasting patients. The clinical significance of these findings is unknown (see 616 617 **PRECAUTIONS:** General).

618 *Cannabinoid Test Interaction*: Efavirenz does not bind to cannabinoid receptors. False-619 positive urine cannabinoid test results have been observed in non-HIV-infected 620 volunteers receiving SUSTIVA when the Microgenics CEDIA<sup>®</sup> DAU Multi-Level THC 621 assay was used for screening. Negative results were obtained when more specific 622 confirmatory testing was performed with gas chromatography/mass spectrometry.

623 Of the three assays analyzed (Microgenics CEDIA DAU Multi-Level THC assay, 624 Cannabinoid Enzyme Immunoassay [Diagnostic Reagents, Inc], and AxSYM<sup>®</sup> 625 Cannabinoid Assay), only the Microgenics CEDIA DAU Multi-Level THC assay showed 626 false-positive results. The other two assays provided true-negative results. The effects of 627 SUSTIVA on cannabinoid screening tests other than these three are unknown. The 628 manufacturers of cannabinoid assays should be contacted for additional information 629 regarding the use of their assays with patients receiving efavirenz.

#### 630 **OVERDOSAGE**

631 Some patients accidentally taking 600 mg twice daily have reported increased nervous632 system symptoms. One patient experienced involuntary muscle contractions.

Treatment of overdose with SUSTIVA (efavirenz) should consist of general supportive
measures, including monitoring of vital signs and observation of the patient's clinical
status. Administration of activated charcoal may be used to aid removal of unabsorbed
drug. There is no specific antidote for overdose with SUSTIVA. Since efavirenz is highly

637 protein bound, dialysis is unlikely to significantly remove the drug from blood.

## 638 **DOSAGE AND ADMINISTRATION**

#### 639 Adults

The recommended dosage of SUSTIVA (efavirenz) is 600 mg orally, once daily, in 640 641 combination with a protease inhibitor and/or nucleoside analogue reverse transcriptase 642 inhibitors (NRTIs). It is recommended that SUSTIVA be taken on an empty stomach, 643 preferably at bedtime. The increased efavirenz concentrations observed following 644 administration of SUSTIVA with food may lead to an increase in frequency of adverse 645 events (see CLINICAL PHARMACOLOGY: Effect of Food on Oral Absorption). 646 Dosing at bedtime may improve the tolerability of nervous system symptoms (see 647 WARNINGS: Nervous System Symptoms, PRECAUTIONS: Information for 648 Patients, and ADVERSE REACTIONS).

649 Concomitant Antiretroviral Therapy: SUSTIVA must be given in combination with
650 other antiretroviral medications (see CLINICAL PHARMACOLOGY: Drug
651 Interactions and PRECAUTIONS: Drug Interactions and INDICATIONS AND
652 USAGE).

**Dosage Adjustment:** If SUSTIVA is coadministered with voriconazole, the voriconazole maintenance dose should be increased to 400 mg every 12 hours and the SUSTIVA dose should be decreased to 300 mg once daily using the capsule formulation (three 100-mg capsules or one 200-mg and one 100-mg capsule). SUSTIVA tablets should not be broken. (See CLINICAL PHARMACOLOGY, Tables 1 and 2; **CONTRAINDICATIONS**; and **PRECAUTIONS: Drug Interactions**).

#### 659 **Pediatric Patients**

660 It is recommended that SUSTIVA be taken on an empty stomach, preferably at bedtime.

Table 11 describes the recommended dose of SUSTIVA for pediatric patients 3 years of

age or older and weighing between 10 and 40 kg. The recommended dosage of

663 SUSTIVA for pediatric patients weighing greater than 40 kg is 600 mg, once daily.

Body V	Body Weight		
kg	lbs	- SUSTIVA Dose (mg)	
10 to <15	22 to <33	200	
15 to <20	33 to <44	250	
20 to <25	44 to <55	300	
25 to <32.5	55 to <71.5	350	
32.5 to <40	71.5 to <88	400	
≥40	≥88	600	

Table 11: Pediatric Dose to be Administered Once Daily

#### 664 HOW SUPPLIED

#### 665 **Capsules**

666 SUSTIVA<sup>®</sup> (efavirenz) capsules are available as follows:

667 *Capsules 200 mg* are gold color, reverse printed with "SUSTIVA" on the body and 668 imprinted "200 mg" on the cap.

- 669 Bottles of 90 NDC 0056-0474-92
- 670 *Capsules 100 mg* are white, reverse printed with "SUSTIVA" on the body and imprinted 671 "100 mg" on the cap.
- 672 Bottles of 30 NDC 0056-0473-30
- 673 *Capsules 50 mg* are gold color and white, printed with "SUSTIVA" on the gold color cap
- and reverse printed "50 mg" on the white body.

675 Bottles of 30 NDC 0056-0470-30

#### 676 **Tablets**

- 677 SUSTIVA (efavirenz) tablets are available as follows:
- 678 *Tablets 600 mg* are yellow, capsular-shaped, film-coated tablets, with "SUSTIVA" 679 printed on both sides.
- 680 Bottles of 30 NDC 0056-0510-30
- 681 SUSTIVA capsules and SUSTIVA tablets should be stored at 25° C (77° F); excursions
- 682 permitted to 15°–30° C (59°–86° F) [see USP Controlled Room Temperature].
- 683 Distributed by:
- 684 Bristol-Myers Squibb Company
- 685 Princeton, NJ 08543 USA
- 686 SUSTIVA<sup>®</sup> is a registered trademark of Bristol-Myers Squibb Pharma Company.
- 687 Other brands listed are the trademarks of their respective owners and are not trademarks
- 688 of Bristol-Myers Squibb Company.
- 689
- 690 © Bristol-Myers Squibb Company 2007
- 691
- 692 Printed in USA
- 693 XX-XXXXXX-XX XXXXXXXX Revised \_\_\_\_\_
- 694

# 695 **PATIENT INFORMATION**

# 696 **SUSTIVA**<sup>®</sup> (sus-TEE-vah)

# 697 [efavirenz (eh-FAH-vih-rehnz)]

- 698 capsules and tablets
- 699

#### 700 ALERT: Find out about medicines that should NOT be taken with SUSTIVA.

Please also read the section "MEDICINES YOU SHOULD NOT TAKE WITH
SUSTIVA."

Read this information before you start taking SUSTIVA. Read it again each time you refill your prescription, in case there is any new information. This leaflet provides a summary about SUSTIVA and does not include everything there is to know about your medicine. This information is not meant to take the place of talking with your doctor.

## 707 What is SUSTIVA?

SUSTIVA is a medicine used in combination with other medicines to help treat infection
with Human Immunodeficiency Virus type 1 (HIV-1), the virus that causes AIDS
(acquired immune deficiency syndrome). SUSTIVA is a type of anti-HIV drug called a
"non-nucleoside reverse transcriptase inhibitor" (NNRTI). NNRTIS are not used in the
treatment of Human Immunodeficiency Virus type 2 (HIV-2) infection.

SUSTIVA works by lowering the amount of HIV-1 in the blood (viral load). SUSTIVA must be taken with other anti-HIV medicines. When taken with other anti-HIV medicines, SUSTIVA has been shown to reduce viral load and increase the number of CD4+ cells, a type of immune cell in blood. SUSTIVA may not have these effects in every patient.

- 718 SUSTIVA does not cure HIV or AIDS. People taking SUSTIVA may still develop other
- 719 infections and complications. Therefore, it is very important that you stay under the care
- of your doctor.
- 721 SUSTIVA has not been shown to reduce the risk of passing HIV to others. Therefore,
- continue to practice safe sex, and do not use or share dirty needles.

## 723 What are the possible side effects of SUSTIVA?

Serious psychiatric problems. A small number of patients experience severe depression, strange thoughts, or angry behavior while taking SUSTIVA. Some patients have thoughts of suicide and a few have actually committed suicide. These problems tend to occur more often in patients who have had mental illness. Contact your doctor right away if you think you are having these psychiatric symptoms, so your doctor can decide if you should continue to take SUSTIVA (efavirenz).

730 **Common side effects.** Many patients have dizziness, trouble sleeping, drowsiness, 731 trouble concentrating, and/or unusual dreams during treatment with SUSTIVA. These 732 side effects may be reduced if you take SUSTIVA at bedtime on an empty stomach. They 733 also tend to go away after you have taken the medicine for a few weeks. If you have these 734 common side effects, such as dizziness, it does not mean that you will also have serious 735 psychiatric problems, such as severe depression, strange thoughts, or angry behavior. Tell 736 your doctor right away if any of these side effects continue or if they bother you. It is 737 possible that these symptoms may be more severe if SUSTIVA is used with alcohol or 738 mood altering (street) drugs.

If you are dizzy, have trouble concentrating, or are drowsy, avoid activities that may bedangerous, such as driving or operating machinery.

Rash is common. Rashes usually go away without any change in treatment. In a small number of patients, rash may be serious. If you develop a rash, call your doctor right away. Rash may be a serious problem in some children. Tell your child's doctor right away if you notice rash or any other side effects while your child is taking SUSTIVA.

745 Other common side effects include tiredness, upset stomach, vomiting, and diarrhea.

746 Changes in body fat. Changes in body fat develop in some patients taking anti-HIV 747 medicine. These changes may include an increased amount of fat in the upper back and 748 neck ("buffalo hump"), in the breasts, and around the trunk. Loss of fat from the legs, 749 arms, and face may also happen. The cause and long-term health effects of these fat 750 changes are not known.

- 751 Tell your doctor or healthcare provider if you notice any side effects while taking752 SUSTIVA.
- Contact your doctor before stopping SUSTIVA because of side effects or for any otherreason.
- 755 This is not a complete list of side effects possible with SUSTIVA. Ask your doctor or
- pharmacist for a more complete list of side effects of SUSTIVA and all the medicinesyou will take.

# 758 How should I take SUSTIVA?

#### 759 General Information

- You should take SUSTIVA on an empty stomach, preferably at bedtime.
- Swallow SUSTIVA with water.
- Taking SUSTIVA with food increases the amount of medicine in your body, which
   may increase the frequency of side effects.
- Taking SUSTIVA at bedtime may make some side effects less bothersome.
- SUSTIVA must be taken in combination with other anti-HIV medicines. If you take
   only SUSTIVA, the medicine may stop working.
- Do not miss a dose of SUSTIVA. If you forget to take SUSTIVA, take the missed dose right away, unless it is almost time for your next dose. Do not double the next dose. Carry on with your regular dosing schedule. If you need help in planning the best times to take your medicine, ask your doctor or pharmacist.
- Take the exact amount of SUSTIVA your doctor prescribes. Never change the dose
   on your own. Do not stop this medicine unless your doctor tells you to stop.
- If you believe you took more than the prescribed amount of SUSTIVA, contact your
   local Poison Control Center or emergency room right away.

- Tell your doctor if you start any new medicine or change how you take old ones.
  Your doses may need adjustment.
- When your SUSTIVA supply starts to run low, get more from your doctor or
   pharmacy. This is very important because the amount of virus in your blood may
   increase if the medicine is stopped for even a short time. The virus may develop
   resistance to SUSTIVA and become harder to treat.
- Your doctor may want to do blood tests to check for certain side effects while you take SUSTIVA (efavirenz).

#### 783 Capsules

The dose of SUSTIVA capsules for adults is 600 mg (three 200-mg capsules, taken together) once a day by mouth. The dose of SUSTIVA for children may be lower (see Can children take SUSTIVA?).

#### 787 Tablets

- The dose of SUSTIVA tablets for adults is 600 mg (one tablet) once a day by mouth.
- 789

#### 790 Can children take SUSTIVA?

Yes, children who are able to swallow capsules can take SUSTIVA. Rash may be a serious problem in some children. Tell your child's doctor right away if you notice rash or any other side effects while your child is taking SUSTIVA. The dose of SUSTIVA for children may be lower than the dose for adults. Capsules containing lower doses of SUSTIVA are available. Your child's doctor will determine the right dose based on your child's weight.

## 797 Who should not take SUSTIVA?

Do not take SUSTIVA if you are allergic to the active ingredient, efavirenz, or to any
 of the inactive ingredients. Your doctor and pharmacist have a list of the inactive
 ingredients.

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# 801 What should I avoid while taking SUSTIVA?

Women taking SUSTIVA should not become pregnant. Serious birth defects have
 been seen in the offspring of animals and women treated with SUSTIVA during
 pregnancy. It is not known whether SUSTIVA caused these defects. Tell your doctor
 right away if you are pregnant. Also talk with your doctor if you want to become
 pregnant.

- Women should not rely only on hormone-based birth control, such as pills, injections, or implants, because SUSTIVA may make these contraceptives ineffective. Women must use a reliable form of barrier contraception, such as a condom or diaphragm, even if they also use other methods of birth control.
- **Do not breast-feed if you are taking SUSTIVA**. The Centers for Disease Control and Prevention recommend that mothers with HIV not breast-feed because they can pass the HIV through their milk to the baby. Also, SUSTIVA may pass through breast milk and cause serious harm to the baby. Talk with your doctor if you are breast-feeding. You may need to stop breast-feeding or use a different medicine.
- Taking SUSTIVA with alcohol or other medicines causing similar side effects as
   SUSTIVA, such as drowsiness, may increase those side effects.
- Do not take any other medicines without checking with your doctor. These medicines
   include prescription and nonprescription medicines and herbal products, especially
   St. John's wort.

#### 821 Before using SUSTIVA, tell your doctor if you

- have problems with your liver or have hepatitis. Your doctor may want to do tests
  to check your liver while you take SUSTIVA.
- have ever had mental illness or are using drugs or alcohol.
- have ever had seizures or are taking medicine for seizures [for example, Dilantin<sup>®</sup>
- 826 (phenytoin), Tegretol<sup>®</sup> (carbamazepine), or phenobarbital]. Your doctor may want to
- 827 check drug levels in your blood from time to time.

# 828 What important information should I know about taking other 829 medicines with SUSTIVA?

830 SUSTIVA may change the effect of other medicines, including ones for HIV, and
831 cause serious side effects. Your doctor may change your other medicines or change
832 their doses. Other medicines, including herbal products, may affect SUSTIVA. For this
833 reason, it is very important to:

- let all your doctors and pharmacists know that you take SUSTIVA.
- tell your doctors and pharmacists about all medicines you take. This includes those
   you buy over-the-counter and herbal or natural remedies.

Bring all your prescription and nonprescription medicines as well as any herbal remedies
that you are taking when you see a doctor, or make a list of their names, how much you
take, and how often you take them. This will give your doctor a complete picture of the
medicines you use. Then he or she can decide the best approach for your situation.

Taking SUSTIVA with St. John's wort (*Hypericum perforatum*), an herbal product sold as a dietary supplement, or products containing St. John's wort is not recommended. Talk with your doctor if you are taking or are planning to take St. John's wort. Taking St. John's wort may decrease SUSTIVA levels and lead to increased viral load and possible resistance to SUSTIVA or cross-resistance to other anti-HIV drugs.

# 846 MEDICINES YOU SHOULD NOT TAKE WITH SUSTIVA

The following medicines may cause serious and life-threatening side effects when takenwith SUSTIVA. You should not take any of these medicines while taking SUSTIVA:

- Hismanal<sup>®</sup> (astemizole)
- 850 Vascor<sup>®</sup> (bepridil)
- 851 Propulsid<sup>®</sup> (cisapride)
- 852 Versed<sup>®</sup> (midazolam)
- 853 Orap<sup>®</sup> (pimozide)
- Halcion<sup>®</sup> (triazolam)
- Ergot medications (for example, Wigraine<sup>®</sup> and Cafergot<sup>®</sup>)

- The following medicine should not be taken with SUSTIVA since it may lose its effect or may increase the chance of having side effects from SUSTIVA:
- Vfend<sup>®</sup> (voriconazole). Some doses of voriconazole can be taken at the same time as
  a lower dose of SUSTIVA, but you must check with your doctor first.

# 860 The following medicines may need to be replaced with another medicine when taken861 with SUSTIVA:

- 862 Fortovase<sup>®</sup>, Invirase<sup>®</sup> (saquinavir)
- Biaxin<sup>®</sup> (clarithromycin)
- Carbatrol<sup>®</sup>, Tegretol<sup>®</sup> (carbamazepine)
- Sporanox<sup>®</sup> (itraconazole)

# 866 The following medicines may require a change in the dose of either SUSTIVA or the 867 other medicine:

- Calcium channel blockers such as Cardizem<sup>®</sup> or Tiazac<sup>®</sup> (diltiazem), Covera HS<sup>®</sup> or Isoptin SR<sup>®</sup> (verapamil), and others.
- The cholesterol-lowering medicines Lipitor<sup>®</sup> (atorvastatin), PRAVACHOL<sup>®</sup>
   (pravastatin), and Zocor<sup>®</sup> (simvastatin).
- 872 Crixivan<sup>®</sup> (indinavir)
- Kaletra<sup>®</sup> (lopinavir/ritonavir)
- Methadone
- Mycobutin<sup>®</sup> (rifabutin)
- REYATAZ<sup>®</sup> (atazanavir sulfate). If you are taking SUSTIVA and REYATAZ, you should also be taking Norvir<sup>®</sup> (ritonavir).
- Rifadin<sup>®</sup> (rifampin) or the rifampin-containing medicines Rifamate<sup>®</sup> and Rifater<sup>®</sup>.
- 879 Zoloft<sup>®</sup> (sertraline)

## 880 These are not all the medicines that may cause problems if you take SUSTIVA. Be

sure to tell your doctor about all medicines that you take.

#### 882 General advice about SUSTIVA:

883 Medicines are sometimes prescribed for conditions that are not mentioned in patient 884 information leaflets. Do not use SUSTIVA for a condition for which it was not 885 prescribed. Do not give SUSTIVA to other people, even if they have the same 886 symptoms you have. It may harm them.

887 Keep SUSTIVA at room temperature  $(77^{\circ} \text{ F})$  in the bottle given to you by your 888 pharmacist. The temperature can range from 59° to 86° F.

889 Keep SUSTIVA out of the reach of children.

890

This leaflet summarizes the most important information about SUSTIVA. If you would like more information, talk with your doctor. You can ask your pharmacist or doctor for the full prescribing information about SUSTIVA, or you can visit the SUSTIVA website at *http://www.sustiva.com* or call 1-800-321-1335.

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