PLETAL®
(PLAY-tal)
(cilostazol) (sil-OS-tah-zol)
Tablets

CONTRAINDICATION
PLETAL is contraindicated in patients with congestive heart failure of any severity. Cilostazol and several of its metabolites are inhibitors of phosphodiesterase III. Several drugs with this pharmacologic effect have caused decreased survival compared to placebo in patients with class III-IV congestive heart failure.

DESCRIPTION
PLETAL (cilostazol) is a quinolinone derivative that inhibits cellular phosphodiesterase (more specific for phosphodiesterase III). The empirical formula of cilostazol is C_{20}H_{27}N_{5}O_{2}, and its molecular weight is 369.46. Cilostazol is 6-[4-(1-cyclohexyl-1H-tetrazol-5-yl)butoxy]-3,4-dihydro-2(1H)-quinolinone, CAS-73963-72-1. The structural formula is:

![Structural formula of cilostazol]

CILOSTAZOL

Cilostazol occurs as white to off-white crystals or as a crystalline powder that is slightly soluble in methanol and ethanol, and is practically insoluble in water, 0.1 N HCl, and 0.1 N NaOH.

PLETAL (cilostazol) tablets for oral administration are available in 50 mg triangular and 100 mg round, white debossed tablets. Each tablet, in addition to the active ingredient, contains the following inactive ingredients: carboxymethylcellulose calcium, corn starch, hydroxypropyl methylcellulose 2910, magnesium stearate, and microcrystalline cellulose.

CLINICAL PHARMACOLOGY
Mechanism of Action
The mechanism of the effects of PLETAL on the symptoms of intermittent claudication is not fully understood. PLETAL and several of its metabolites are cyclic AMP (cAMP) phosphodiesterase III inhibitors (PDE III inhibitors), inhibiting phosphodiesterase activity and suppressing cAMP degradation with a resultant increase in cAMP in platelets and blood vessels, leading to inhibition of platelet aggregation and vasodilation, respectively.

PLETAL reversibly inhibits platelet aggregation induced by a variety of stimuli, including thrombin, ADP, collagen, arachidonic acid, epinephrine, and shear stress. Effects on circulating plasma lipids have been examined in patients taking PLETAL. After 12 weeks, as compared to placebo, PLETAL 100 mg twice daily produced a reduction in triglycerides of 29.3 mg/dL (15%) and an increase in HDL-cholesterol of 4.0 mg/dL (± 10%).

Cilostazol affects both vascular beds and cardiovascular function. It produces non-homogeneous dilation of vascular beds, with greater dilation in femoral beds than in vertebral, carotid or superior mesenteric arteries. Renal arteries were not responsive to the effects of cilostazol.
Pharmacodynamics
In dogs or cynomolgous monkeys, cilostazol increased heart rate, myocardial contractile force, and coronary blood flow as well as ventricular automaticity, as would be expected for a PDE III inhibitor. Left ventricular contractility was increased at doses required to inhibit platelet aggregation. A-V conduction was accelerated. In humans, heart rate increased in a dose-proportional manner by a mean of 5.1 and 7.4 beats per minute in patients treated with 50 and 100 mg twice daily, respectively. In 264 patients evaluated with Holter monitors, numerically more cilostazol-treated patients had increases in ventricular premature beats and non-sustained ventricular tachycardia events than did placebo-treated patients; the increases were not dose-related.

Pharmacokinetics
PLETAL is absorbed after oral administration. A high fat meal increases absorption, with an approximately 90% increase in C_max and a 25% increase in AUC. Absolute bioavailability is not known. Cilostazol is extensively metabolized by hepatic cytochrome P-450 enzymes, mainly 3A4, and, to a lesser extent, 2C19, with metabolites largely excreted in urine. Two metabolites are active, with one metabolite appearing to account for at least 50% of the pharmacologic (PDE III inhibition) activity after administration of PLETAL.

Pharmacokinetics are approximately dose proportional. Cilostazol and its active metabolites have apparent elimination half-lives of about 11-13 hours. Cilostazol and its active metabolites accumulate about 2-fold with chronic administration and reach steady state blood levels within a few days. The pharmacokinetics of cilostazol and its two major active metabolites were similar in healthy subjects and patients with intermittent claudication due to peripheral arterial disease (PAD).

The mean ± SEM plasma concentration-time profile at steady state after multiple dosing of PLETAL 100 mg twice daily is shown below:

Distribution:
Plasma Protein and Erythrocyte Binding
Cilostazol is 95 - 98% protein bound, predominantly to albumin. The mean percent binding for 3,4-dehydro-cilostazol is 97.4% and for 4'-trans-hydroxy-cilostazol is 66%. Mild hepatic impairment did not affect protein binding. The free fraction of cilostazol was 27% higher in subjects with renal impairment than in healthy subjects. The displacement of cilostazol from plasma proteins by erythromycin, quinidine, warfarin, and omeprazole was not clinically significant.

Reference ID: 3691430
Metabolism and Excretion:
Cilostazol is eliminated predominately by metabolism and subsequent urinary excretion of metabolites. Based on in vitro studies, the primary isoenzymes involved in cilostazol’s metabolism are CYP3A4 and, to a lesser extent, CYP2C19. The enzyme responsible for metabolism of 3,4-dehydro-cilostazol, the most active of the metabolites, is unknown.

Following oral administration of 100 mg radiolabeled cilostazol, 56% of the total radioactivity AUC in plasma was cilostazol, 15% was 3,4-dehydro-cilostazol (4-7 times as active as cilostazol), and 4% was 4’-trans-hydroxy-cilostazol (one fifth as active as cilostazol).

The primary route of elimination was via the urine (74%), with the remainder excreted in feces (20%). No measurable amount of unchanged cilostazol was excreted in the urine, and less than 2% of the dose was excreted as 3,4-dehydro-cilostazol. About 30% of the dose was excreted in urine as 4’-trans-hydroxy-cilostazol. The remainder was excreted as other metabolites, none of which exceeded 5%. There was no evidence of induction of hepatic microenzymes.

Special Populations:
Age and Gender:
The total and unbound oral clearances, adjusted for body weight, of cilostazol and its metabolites were not significantly different with respect to age and/or gender across a 50-to-80-year-old age range.

Smokers:
Population pharmacokinetic analysis suggests that smoking decreased cilostazol exposure by about 20%.

Hepatic Impairment:
The pharmacokinetics of cilostazol and its metabolites were similar in subjects with mild hepatic disease as compared to healthy subjects. Patients with moderate or severe hepatic impairment have not been studied.

Renal Impairment:
The total pharmacologic activity of cilostazol and its metabolites was similar in subjects with mild to moderate renal impairment and in healthy subjects. Severe renal impairment increases metabolite levels and alters protein binding of the parent. The expected pharmacologic activity, however, based on plasma concentrations and relative PDE III inhibiting potency of parent drug and metabolites, appeared little changed. Patients on dialysis have not been studied, but, is unlikely that cilostazol can be removed efficiently by dialysis because of its high protein binding (95 - 98%).

Pharmacokinetic and Pharmacodynamic Drug-Drug Interactions:
Cilostazol could have pharmacodynamic interactions with other inhibitors of platelet function and pharmacokinetic interactions because of effects of other drugs on its metabolism by CYP3A4 or CYP2C19. A reduced dose of PLETAL should be considered when taken concomitantly with CYP3A4 or CYP2C19 inhibitors. Cilostazol does not appear to inhibit CYP3A4 (see Pharmacokinetic and Pharmacodynamic Drug-Drug Interactions, Lovastatin).

Aspirin:
Short-term (≤4 days) co-administration of aspirin with PLETAL increased the inhibition of ADP-induced ex vivo platelet aggregation by 22% - 37% when compared to either aspirin or PLETAL alone. Short-term (≤4 days) co-administration of aspirin with PLETAL increased the inhibition of arachidonic acid-induced ex vivo platelet aggregation by 20% compared to PLETAL alone and by 48% compared to aspirin alone. However, short-term co-administration of aspirin with PLETAL had no clinically significant impact on PT, aPTT, or bleeding time compared to aspirin alone. Effects of long-term co-administration in the general population are unknown. In eight randomized, placebo-controlled, double-blind clinical trials, aspirin was coadministered with cilostazol to 201 patients. The most frequent doses and mean durations of aspirin therapy were 75-81 mg daily for 137 days.
(107 patients) and 325 mg daily for 54 days (85 patients). There was no apparent increase in frequency of hemorrhagic adverse effects in patients taking cilostazol and aspirin compared to patients taking placebo and equivalent doses of aspirin.

Warfarin:
The cytochrome P-450 isoenzymes involved in the metabolism of R-warfarin are CYP3A4, CYP1A2, and CYP2C19, and in the metabolism of S-warfarin, CYP2C9. Cilostazol did not inhibit either the metabolism or the pharmacologic effects (PT, aPTT, bleeding time, or platelet aggregation) of R- and S-warfarin after a single 25-mg dose of warfarin. The effect of concomitant multiple dosing of warfarin and PLETAL on the pharmacokinetics and pharmacodynamics of both drugs is unknown.

Clopidogrel:
Multiple doses of clopidogrel do not significantly increase steady state plasma concentrations of cilostazol.

Inhibitors of CYP3A4:

**Strong Inhibitors of CYP3A4**
Administration of ketoconazole 400 mg with cilostazol 100 mg resulted in a 94% increase in the Cmax and a 117% increase in the AUC of cilostazol.

A dose reduction to 50 mg twice daily should be considered when administered with strong inhibitors of CYP3A4 (e.g. ketoconazole itraconazole, clarithromycin, telithromycin, nelfinavir, indinavir, ritonavir). (see DOSAGE AND ADMINISTRATION and PRECAUTIONS).

**Moderate Inhibitors of CYP3A4**

**Erythromycin**
Administration of erythromycin with cilostazol resulted in a 47% increase in the Cmax and a 72% increase in the AUC of cilostazol. The AUC of the 4'-trans-hydroxy-cilostazol metabolite was increased by 141%. (see DOSAGE AND ADMINISTRATION).

**Diltiazem**
Administration of diltiazem with cilostazol decreased the clearance of cilostazol by ~30%. Cilostazol Cmax increased ~30% and AUC increased ~40%. (see DOSAGE AND ADMINISTRATION).

**Grapefruit Juice**
Administration of a single dose of 100 mg cilostazol with 240 ml grapefruit juice (an inhibitor of intestinal CYP3A4) increased the Cmax of cilostazol by ~50%, but had no effect on AUC.

A dose reduction to 50 mg twice daily should be considered when administered with moderate CYP3A4 inhibitors (e.g., erythromycin, fluconazole, diltiazem, grapefruit juice) (see DOSAGE AND ADMINISTRATION and PRECAUTIONS).

Inhibitors of CYP2C19

**Omeprazole:**
Administration of omeprazole with 100 mg cilostazol did not significantly affect the metabolism of cilostazol, but the AUC of the 3,4-dehydro-cilostazol was increased by 69%.

A dose reduction to 50 mg twice daily should be considered when administered with CYP2C19 inhibitors (e.g., ticlopidine, fluconazole, omeprazole, fluoxetine) (see DOSAGE AND ADMINISTRATION and PRECAUTIONS).
Quinidine

Concomitant administration of quinidine with a single dose of cilostazol 100 mg did not alter cilostazol pharmacokinetics.

Lovastatin

The concomitant administration of lovastatin with cilostazol decreases cilostazol $C_{ss,max}$ and $AUC_{\tau}$ by 15%. There is also a decrease, although nonsignificant, in cilostazol metabolite concentrations. Co-administration of cilostazol with lovastatin increases lovastatin and $\beta$-hydroxi lovastatin $AUC$ approximately 70%. This is most likely clinically insignificant.

Effect of Cilostazol on CYP3A4

PLETAL does not appear to cause increased blood levels of drugs metabolized by CYP3A4, as it had no effect on lovastatin, a drug with metabolism very sensitive to CYP3A4 inhibition.

CLINICAL STUDIES

The ability of PLETAL to improve walking distance in patients with stable intermittent claudication was studied in eight, randomized, placebo-controlled, double-blind trials of 12 to 24 weeks’ duration involving 2,274 patients using dosages of 50 mg twice daily ($n=303$), 100 mg twice daily ($n=998$), and placebo ($n=973$). Efficacy was determined primarily by the change in maximal walking distance from baseline (compared to change on placebo) on one of several standardized exercise treadmill tests.

Compared to patients treated with placebo, patients treated with PLETAL 50 or 100 mg twice daily experienced statistically significant improvements in walking distances both for the distance before the onset of claudication pain and the distance before exercise-limiting symptoms supervened (maximal walking distance). The effect of PLETAL on walking distance was seen as early as the first on-therapy observation point of two or four weeks.

The following figure depicts the percent mean improvement in maximal walking distance, at study end for each of the eight studies.

![Percent Mean Improvement in Maximal Walking Distance at Study End for the Eight Randomized, Double-Blind, Placebo-Controlled Clinical Trials](chart.png)
Across the eight clinical trials, the range of improvement in maximal walking distance in patients treated with PLETAL 100 mg twice daily, expressed as the percent mean change from baseline, was 28% to 100%.

The corresponding changes in the placebo group were –10% to 41%.

The Walking Impairment Questionnaire, which was administered in six of the eight clinical trials, assesses the impact of a therapeutic intervention on walking ability. In a pooled analysis of the six trials, patients treated with either PLETAL 100 mg twice daily or 50 mg twice daily reported improvements in their walking speed and walking distance as compared to placebo. Improvements in walking performance were seen in the various subpopulations evaluated, including those defined by gender, smoking status, diabetes mellitus, duration of peripheral artery disease, age, and concomitant use of beta blockers or calcium channel blockers. PLETAL has not been studied in patients with rapidly progressing claudication or in patients with leg pain at rest, ischemic leg ulcers, or gangrene. Its long-term effects on limb preservation and hospitalization have not been evaluated.

A randomized, double-blind, placebo-controlled Phase IV study was conducted to assess the long-term effects of cilostazol, with respect to mortality and safety, in 1,439 patients with intermittent claudication and no heart failure. The trial stopped early due to enrollment difficulties and a lower than expected overall death rate. With respect to mortality, the observed 36-month Kaplan-Meier event rate for deaths on study drug with a median time on study drug of 18 months was 5.6% (95% CI of 2.8 to 8.4 %) on cilostazol and 6.8% (95% CI of 1.9 to 11.5 %) on placebo. These data appear to be sufficient to exclude a 75% increase in the risk of mortality on cilostazol, which is the \textit{a priori} study hypothesis.

**INDICATIONS AND USAGE**
lf PLETAL is indicated for the reduction of symptoms of intermittent claudication, as indicated by an increased walking distance.

**CONTRAINDICATIONS**
PLETAL is contraindicated in patients with congestive heart failure of any severity. Cilostazol and several of its metabolites are inhibitors of phosphodiesterase III. Several drugs with this pharmacologic effect have caused decreased survival compared to placebo in patients with class III-IV congestive heart failure.

PLETAL is contraindicated in patients with haemostatic disorders or active pathologic bleeding, such as bleeding peptic ulcer and intracranial bleeding. PLETAL inhibits platelet aggregation in a reversible manner.

PLETAL is contraindicated in patients with known or suspected hypersensitivity to any of its components.

**PRECAUTIONS**

*Hematologic:*
Rare cases have been reported of thrombocytopenia or leukopenia progressing to agranulocytosis when PLETAL was not immediately discontinued. The agranulocytosis, however, was reversible on discontinuation of PLETAL. Monitor platelets and white blood cell counts periodically.

*Cardiovascular:*
Based on its mechanism of action, cilostazol may induce tachycardia, palpitation, tachyarrhythmia and/or hypotension. The increase in heart rate associated with cilostazol is approximately 5 to 7 bpm. Patients with a history of ischemic heart disease may be at risk for exacerbations of angina pectoris or myocardial infarction.

*Strong or Moderate CYP3A4 and CYP2C19 inhibitors*
Cilostazol plasma concentrations and overall pharmacological activity are increased when cilostazol is administered with strong or moderate CYP3A4 inhibitors (e.g. ketoconazole, itraconazole, erythromycin, diltiazem) and strong CYP2C19 inhibitors (e.g. ticlopidine, fluconazole, omeprazole) (see \textit{Pharmacokinetic and Pharmacodynamic Drug-Drug Interactions}). Dose reduction to 50 mg twice daily should be considered (see \textit{DOSAGE AND ADMINISTRATION}).
**Information for Patients**

Please refer to the patient package insert. Patients should be advised:

- to read the patient package insert for PLETAL carefully before starting therapy and to reread it each time therapy is renewed in case the information has changed.
- to take PLETAL at least one-half hour before or two hours after food.
- that the beneficial effects of PLETAL on the symptoms of intermittent claudication may not be immediate. Although the patient may experience benefit in 2 to 4 weeks after initiation of therapy, treatment for up to 12 weeks may be required before a beneficial effect is experienced.

**Hepatic Impairment**

Patients with moderate or severe hepatic impairment have not been studied in clinical trials. Special caution is advised when PLETAL is used in such patients.

**Renal Impairment**

Patients on dialysis have not been studied, but, it is unlikely that cilostazol can be removed efficiently by dialysis because of its high protein binding (95-98%).

Special caution is advised when PLETAL is used in patients with severe renal impairment: estimated creatinine clearance < 25 ml/min.

**Cardiovascular Toxicity**

Repeated oral administration of cilostazol to dogs (30 or more mg/kg/day for 52 weeks, 150 or more mg/kg/day for 13 weeks, and 450 mg/kg/day for 2 weeks), produced cardiovascular lesions that included endocardial haemorrhage, hemosiderin deposition and fibrosis in the left ventricle, haemorrhage in the right atrial wall, haemorrhage and necrosis of the smooth muscle in the wall of the coronary artery, intimal thickening of the coronary artery, and coronary arteritis and periarteritis. At the lowest dose associated with cardiovascular lesions in the 52-week study, systemic exposure (AUC) to unbound cilostazol was less than that seen in humans at the maximum recommended human dose (MRHD) of 100 mg twice daily. Similar lesions have been reported in dogs following the administration of other positive inotropic agents (including PDE III inhibitors) and/or vasodilating agents. No cardiovascular lesions were seen in rats following 5 or 13 weeks of administration of cilostazol at doses up to 1500 mg/kg/day. At this dose, systemic exposures (AUCs) to unbound cilostazol were only about 1.5 and 5 times (male and female rats, respectively) the exposure seen in humans at the MRHD. Cardiovascular lesions were also not seen in rats following 52 weeks of administration of cilostazol at doses up to 150 mg/kg/day. At this dose, systemic exposures (AUCs) to unbound cilostazol were about 0.5 and 5 times (male and female rats, respectively) the exposure in humans at the MRHD. In female rats, cilostazol AUCs were similar at 150 and 1500 mg/kg/day. Cardiovascular lesions were also not observed in monkeys after oral administration of cilostazol for 13 weeks at doses up to 1800 mg/kg/day. While this dose of cilostazol produced pharmacologic effects in monkeys, plasma cilostazol levels were less than those seen in humans given the MRHD, and those seen in dogs given doses associated with cardiovascular lesions.

**Carcinogenesis, Mutagenesis, Impairment of Fertility**

Dietary administration of cilostazol to male and female rats and mice for up to 104 weeks, at doses up to 500 mg/kg/day in rats and 1000 mg/kg/day in mice, revealed no evidence of carcinogenic potential. The maximum doses administered in both rat and mouse studies were, on a systemic exposure basis, less than the human exposure at the MRHD of the drug. Cilostazol tested negative in bacterial gene mutation, bacterial DNA repair, mammalian cell gene mutation, and mouse in vivo bone marrow chromosomal aberration assays. It was, however, associated with a significant increase in chromosomal aberrations in the in vitro Chinese Hamster Ovary Cell assay. Cilostazol did not affect fertility or mating performance of male and female rats at doses as high as 1000 mg/kg/day. At this dose, systemic exposures (AUCs) to unbound cilostazol were less than 1.5 times in males, and about 5 times in females, the exposure in humans at the MRHD.
Pregnancy

Pregnancy Category C: In a rat developmental toxicity study, oral administration of 1000 mg cilostazol/kg/day was associated with decreased fetal weights, and increased incidences of cardiovascular, renal, and skeletal anomalies (ventricular septal, aortic arch and subclavian artery abnormalities, renal pelvic dilation, 14th rib, and retarded ossification). At this dose, systemic exposure to unbound cilostazol in nonpregnant rats was about 5 times the exposure in humans given the MRHD. Increased incidences of ventricular septal defect and retarded ossification were also noted at 150 mg/kg/day (5 times the MRHD on a systemic exposure basis). In a rabbit developmental toxicity study, an increased incidence of retardation of ossification of the sternum was seen at doses as low as 150 mg/kg/day. In nonpregnant rabbits given 150 mg/kg/day, exposure to unbound cilostazol was considerably lower than that seen in humans given the MRHD, and exposure to 3,4-dehydro-cilostazol was barely detectable. When cilostazol was administered to rats during late pregnancy and lactation, an increased incidence of stillborn and decreased birth weights of offspring was seen at doses of 150 mg/kg/day (5 times the MRHD on a systemic exposure basis).

There are no adequate and well-controlled studies in pregnant women.

Nursing Mothers
Transfer of cilostazol into milk has been reported in experimental animals (rats). Because of the potential risk to nursing infants, a decision should be made to discontinue nursing or to discontinue PLETAL.

Pediatric Use
The safety and effectiveness of PLETAL in pediatric patients have not been established.

Geriatric Use
Of the total number of subjects (n = 2274) in clinical studies of PLETAL, 56 percent were 65-years-old and over, while 16 percent were 75-years-old and over. No overall differences in safety or effectiveness were observed between these subjects and younger subjects, and other reported clinical experience has not identified differences in responses between the elderly and younger patients, but greater sensitivity of some older individuals cannot be ruled out. Pharmacokinetic studies have not disclosed any age-related effects on the absorption, distribution, metabolism, and elimination of cilostazol and its metabolites.

ADVERSE REACTIONS
Adverse events were assessed in eight placebo-controlled clinical trials involving 2274 patients exposed to either 50 or 100 mg twice daily PLETAL (n=1301) or placebo (n=973), with a median treatment duration of 127 days for patients on PLETAL and 134 days for patients on placebo.

The only adverse event resulting in discontinuation of therapy in ≥ 3% of patients treated with PLETAL 50 or 100 mg twice daily was headache, which occurred with an incidence of 1.3%, 3.5%, and 0.3% in patients treated with PLETAL 50 mg twice daily, 100 mg twice daily, or placebo, respectively. Other frequent causes of discontinuation included palpitation and diarrhea, both 1.1% for cilostazol (all doses) versus 0.1% for placebo.

The most commonly reported adverse events, occurring in ≥ 2% of patients treated with PLETAL 50 or 100 mg twice daily, and with a frequency higher than placebo, are shown in the table (below).

Other events seen with an incidence of ≥ 2%, but occurring in the placebo group at least as frequently as in the 100 mg twice daily group were: asthenia, hypertension, vomiting, leg cramps, hypesthesia, paresthesia, dyspnea, rash, hematuria, urinary tract infection, flu syndrome, angina pectoris, arthritis, and bronchitis.
Most Commonly Reported AEs (Incidence ≥ 2%) in Patients on PLETAL (PLT) 50 mg twice daily or 100 mg twice daily and Occurring at a Rate in the 100 mg twice daily Group Higher Than in Patients on Placebo

<table>
<thead>
<tr>
<th>Adverse Events (AEs) by Body System</th>
<th>PLT 50 mg twice daily (N=303)</th>
<th>PLT 100 mg twice daily (N=998)</th>
<th>Placebo (N=973)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BODY AS A WHOLE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>4</td>
<td>5</td>
<td>3</td>
</tr>
<tr>
<td>Back pain</td>
<td>6</td>
<td>7</td>
<td>6</td>
</tr>
<tr>
<td>Headache</td>
<td>27</td>
<td>34</td>
<td>14</td>
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<tr>
<td>Infection</td>
<td>14</td>
<td>10</td>
<td>8</td>
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<tr>
<td>CARDIOVASCULAR</td>
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<tr>
<td>Palpitation</td>
<td>5</td>
<td>10</td>
<td>1</td>
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<tr>
<td>Tachycardia</td>
<td>4</td>
<td>4</td>
<td>1</td>
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<tr>
<td>DIGESTIVE</td>
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<tr>
<td>Abnormal stools</td>
<td>12</td>
<td>15</td>
<td>4</td>
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<tr>
<td>Diarrhea</td>
<td>12</td>
<td>19</td>
<td>7</td>
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<tr>
<td>Dyspepsia</td>
<td>6</td>
<td>6</td>
<td>4</td>
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<tr>
<td>Flatulence</td>
<td>2</td>
<td>3</td>
<td>2</td>
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<tr>
<td>Nausea</td>
<td>6</td>
<td>7</td>
<td>6</td>
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<tr>
<td>METABOLIC &amp; NUTRITIONAL</td>
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<tr>
<td>Peripheral edema</td>
<td>9</td>
<td>7</td>
<td>4</td>
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<tr>
<td>MUSCULO-SKELETAL</td>
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<tr>
<td>Myalgia</td>
<td>2</td>
<td>3</td>
<td>2</td>
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<tr>
<td>NERVOUS</td>
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<td></td>
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<tr>
<td>Dizziness</td>
<td>9</td>
<td>10</td>
<td>6</td>
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<tr>
<td>Vertigo</td>
<td>3</td>
<td>1</td>
<td>1</td>
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<tr>
<td>RESPIRATORY</td>
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<tr>
<td>Cough increased</td>
<td>3</td>
<td>4</td>
<td>3</td>
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<tr>
<td>Pharyngitis</td>
<td>7</td>
<td>10</td>
<td>7</td>
</tr>
<tr>
<td>Rhinitis</td>
<td>12</td>
<td>7</td>
<td>5</td>
</tr>
</tbody>
</table>

Less frequent adverse events (<2%) that were experienced by patients exposed to PLETAL 50 mg twice daily or 100 mg twice daily in the eight controlled clinical trials and that occurred at a frequency in the 100 mg twice daily group greater than in the placebo group, regardless of suspected drug relationship, are listed below.

**Body as a whole:** Chills, face edema, fever, generalized edema, malaise, neck rigidity, pelvic pain, retroperitoneal haemorrhage.

**Cardiovascular:** Atrial fibrillation, atrial flutter, cerebral infarct, cerebral ischemia, congestive heart failure, heart arrest, haemorrhage, hypotension, myocardial infarction, myocardial ischemia, nodal arrhythmia, postural hypotension, supraventricular tachycardia, syncope, varicose vein, vasodilation, ventricular extrasystoles, ventricular tachycardia.

**Digestive:** Anorexia, cholelithiasis, colitis, duodenal ulcer, duodenitis, esophageal haemorrhage, esophagitis, increased GGT, gastritis, gastroenteritis, gum haemorrhage, hematemeses, melena, peptic ulcer, periodontal abscess, rectal haemorrhage, stomach ulcer, tongue edema.

**Endocrine:** Diabetes mellitus.

**Hemic and Lymphatic:** Anemia, ecchymosis, iron deficiency anemia, polycythemia, purpura.

**Metabolic and Nutritional:** Increased creatinine, gout, hyperlipemia, hyperuricemia.

**Muscloskeletal:** Arthralgia, bone pain, bursitis.

**Nervous:** Anxiety, insomnia, neuralgia.
Respiratory: Asthma, epistaxis, hemoptysis, pneumonia, sinusitis.
Skin and Appendages: Dry skin, furunculosis, skin hypertrophy, urticaria.
Special Senses: Amblyopia, blindness, conjunctivitis, diplopia, ear pain, eye haemorrhage, retinal haemorrhage, tinnitus.
Urogenital: Albuminuria, cystitis, urinary frequency, vaginal haemorrhage, vaginitis.

Post-Marketing Experience
The following adverse drug reactions (ADRs) to PLETAL have been reported worldwide since the launch of PLETAL in the US.

- Blood and lymphatic system disorders:
  - Agranulocytosis, aplastic anemia, granulocytopenia, pancytopenia, thrombocytopenia, leukopenia, bleeding tendency
- Cardiac disorders:
  - Torsades de pointes, QTc prolongation (torsades de pointes and QTc prolongation occurred in patients with cardiac disorders, e.g. complete atrioventricular block, cardiac failure and bradyarrhythmia, when treated with cilostazol. Cilostazol was used “off label” due to its positive chronotropic action)
- Gastrointestinal disorders:
  - Gastrointestinal haemorrhage
- General disorders and administration site conditions:
  - Pain, chest pain, hot flushes
- Hepatobiliary disorders:
  - Hepatic dysfunction/abnormal liver function tests, jaundice
- Injury, poisoning and procedural complications:
  - Extradural haematoma and subdural haematoma
- Investigations:
  - Blood glucose increased, blood uric acid increased, platelet count decreased, white blood cell count decreased, increase in BUN (blood urea increased), blood pressure increase
- Nervous system disorders:
  - Intracranial haemorrhage, cerebral haemorrhage, cerebrovascular accident
- Respiratory, thoracic and mediastinal disorders:
  - Pulmonary haemorrhage, interstitial pneumonia
- Skin and subcutaneous tissue disorders:
  - Haemorrhage subcutaneous, pruritus, skin eruptions including Stevens-Johnson syndrome, skin drug eruption (dermatitis medicamentosa)
- Vascular disorders:
  - Subacute thrombosis (these cases of subacute thrombosis occurred in patients treated with aspirin and “off label” use of cilostazol for prevention of thrombotic complication after coronary stenting)

OVERDOSAGE
Information on acute overdosage with PLETAL in humans is limited. The signs and symptoms of an acute overdose can be anticipated to be those of excessive pharmacologic effect: severe headache, diarrhea, hypotension, tachycardia, and possibly cardiac arrhythmias. The patient should be carefully observed and given supportive treatment. Since cilostazol is highly protein-bound, it is unlikely that it can be efficiently removed by hemodialysis or peritoneal dialysis. The oral LD₃₀ of cilostazol is >5.0 g/kg in mice and rats and >2.0 g/kg in dogs.

DOSAGE AND ADMINISTRATION
The recommended dosage of PLETAL is 100 mg twice daily taken at least half an hour before or two hours after breakfast and dinner.

A dose reduction to 50 mg twice daily should be considered during co-administration with strong or moderate CYP3A4 inhibitors (e.g., ketoconazole, itraconazole, erythromycin and diltiazem) and during co-administration
with CYP2C19 inhibitors (e.g., ticlopidine, fluconazole and omeprazole. (see PRECAUTIONS AND Pharmacokinetic and Pharmacodynamic Drug-Drug Interactions) twice daily

Patients may respond as early as 2 to 4 weeks after the initiation of therapy, but treatment for up to 12 weeks may be needed before a beneficial effect is experienced. If symptoms are unimproved after 3 months, discontinue Pletal.

Discontinuation of Therapy

The available data suggest that the dosage of PLETAL can be reduced or discontinued without rebound (i.e., platelet hyperaggregability).

HOW SUPPLIED

PLETAL is supplied as 50 mg and 100 mg tablets. The 50 mg tablets are white, triangular, debossed with PLETAL 50, and provided in bottles of 60 tablets (NDC #59148-003-16). The 100 mg tablets are white, round, debossed with PLETAL 100, and provided in bottles of 60 tablets (NDC #59148-002-16).

Rx ONLY.

STORAGE

Store PLETAL tablets at 25℃ (77°F); excursions permitted to 15-30℃ (59-86°F) [See USP Controlled Room Temperature].

Manufactured for

OTSUKA AMERICA PHARMACEUTICAL, INC.
Rockville, MD 20850

Manufactured by

OTSUKA PHARMACEUTICAL CO., LTD.
Tokushima 771-0182, Japan

02US15L-XXXX
January 2015
U.S. Patent No. 4,277,479