PRODUCT MONOGRAPH
INCLUDING PATIENT MEDICATION INFORMATION

PrSITAVIG™
acyclovir mucoadhesive buccal tablets
Mucoadhesive Buccal Tablet 50 mg
Antiviral

Cipher Pharmaceuticals Inc.
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Mississauga, Ontario
L5N 8K4, Canada

Date of Preparation:
November 27, 2017

Submission Control No: 189933
PART I: HEALTH PROFESSIONAL INFORMATION

SUMMARY PRODUCT INFORMATION

<table>
<thead>
<tr>
<th>Route of Administration</th>
<th>Dosage Form / Strength</th>
<th>Clinically Relevant Nonmedicinal Ingredients</th>
</tr>
</thead>
<tbody>
<tr>
<td>Buccal</td>
<td>Mucoadhesive buccal tablet 50 mg</td>
<td>None</td>
</tr>
<tr>
<td></td>
<td></td>
<td>For a complete listing see Dosage Forms, Composition and Packaging section</td>
</tr>
</tbody>
</table>

INDICATIONS AND CLINICAL USE

SITAVIG™ (acyclovir mucoadhesive buccal tablets) is indicated for the treatment of recurrent herpes labialis (cold sores) in immunocompetent adults.

Geriatrics (≥ 65 years of age):
Clinical studies of SITAVIG™ did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects.

Pediatrics (<18 years of age):
Safety and effectiveness of SITAVIG™ in pediatric patients have not been established.

CONTRAINdications

- Patients with known hypersensitivity (e.g., anaphylaxis) to acyclovir, milk protein concentrate, or any other component of the product.
- For a complete listing, see the DOSAGE FORMS, COMPOSITION AND PACKAGING section of the product monograph.
WARNINGS AND PRECAUTIONS

General

SITAVIG™ is a single dose therapy for the treatment of recurrent herpes labialis. Therapy should be initiated at the onset of prodromal symptoms, and prior to the appearance of any signs of herpes labialis lesions. SITAVIG™ should be used in the upper gum region (canine fossa), for recurrent herpes labialis (cold sores) of the lips and around the mouth.

Special Populations

Pregnant Women: SITAVIG™ should be used during pregnancy only if the potential benefit outweighs the potential risk to the fetus. No studies with SITAVIG™ have been performed in pregnant women. Systemic exposure of acyclovir following buccal administration of SITAVIG™ is minimal.

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

Animal reproduction studies have not been conducted with SITAVIG. Acyclovir was not teratogenic in the mouse, rabbit or rat at exposures greatly in excess of human exposure.

Nursing Women: There is no experience with SITAVIG™ in nursing mothers. SITAVIG™ should be administered to a nursing mother with caution.

It is not known whether topically applied acyclovir is excreted in breast milk. Systemic exposure following buccal administration is minimal.

After oral (systemic) administration of acyclovir, concentrations have been documented in breast milk in 2 women and ranged from 0.6 to 4.1 times the corresponding plasma levels. These concentrations would potentially expose the nursing infant to a dose of acyclovir up to 0.3 mg/kg/day.

Geriatrics (≥ 65 years of age): Clinical studies of SITAVIG™ did not include sufficient numbers of subjects aged 65 and over to determine whether they respond differently from younger subjects.

Pediatrics (< 18 years of age): Safety and effectiveness of SITAVIG™ in pediatric patients have not been established.

Immunocompromised Patients: The safety of SITAVIG™ has not been studied in immunocompromised subjects.
ADVERSE REACTIONS

Adverse Drug Reaction Overview

The safety of SITAVIS™ was assessed in a Phase III clinical trial reflecting exposure to SITAVIS™ in 378 adult patients with recurrent herpes labialis.

The adverse drug reactions that occurred in ≥ 1% of patients included headache and application site pain. There was no discontinuation of SITAVIS™ due to adverse drug reactions. Most treatment related adverse events were mild or moderate in severity. There were no serious adverse drug reactions reported.

Clinical Trial Adverse Drug Reactions

Because clinical trials are conducted under very specific conditions the adverse reaction rates observed in the clinical trials may not reflect the rates observed in practice and should not be compared to the rates in the clinical trials of another drug. Adverse drug reaction information from clinical trials is useful for identifying drug-related adverse events and for approximating rates.

One randomized, double-blind, placebo controlled trial was conducted in patients with recurrent herpes labialis (cold sores), having at least 4 herpes labialis episodes in the previous year. In this trial, 378 herpes simplex virus (HSV) infected subjects used SITAVIS™ as a single dose, and 397 subjects used placebo. The incidence of adverse drug reactions was similar in the two groups.

Table 1 provides adverse drug reactions that were reported in ≥ 1% of patients treated with SITAVIS™.

Table 1 - Adverse Drug Reactions Reported in ≥ 1% of Patients in Phase 3 Trial

<table>
<thead>
<tr>
<th>Event</th>
<th>SITAVIS™ N = 378 (%)</th>
<th>Placebo N = 397 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nervous System Disorders</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Headache</td>
<td>4 (1.1)</td>
<td>9 (2.3)</td>
</tr>
<tr>
<td>General Disorders and Administration Site Conditions</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Application Site Pain</td>
<td>4 (1.1)</td>
<td>4 (1.0)</td>
</tr>
</tbody>
</table>

Less Common Clinical Trial Adverse Drug Reactions (<1%)

- **Blood and Lymphatic System Disorders**: Lymphadenopathy, thrombocytopenia.
- **Ear and Labyrinth Disorders**: Vertigo.
- **Gastrointestinal Disorders**: Abdominal pain upper, aphthous stomatitis, diarrhoea, dry mouth, gastrooesophageal reflux disease, gingival pain, lip dry, lip swelling, mouth ulceration, nausea, oral pain, stomach discomfort, stomatitis, vomiting.
**General Disorders and Administration Site Conditions:** Application site discomfort, application site erythema, application site irritation, application site paraesthesia, influenza like illness, mucosal erosion.

**Infections and Infestations:** Oral herpes.

**Investigations:** Alanine aminotransferase increased, aspartate aminotransferase increased, blood creatine increased, blood creatinine increased, gamma-glutamyltransferase increased, mean cell hemoglobin increased.

**Musculoskeletal and Connective Tissue Disorder:** Myalgia.

**Nervous System Disorders:** Dizziness, lethargy.

**Skin and Subcutaneous Tissue Disorders:** Erythema, heat rash, pruritus, rash, rash erythematous.

**Vascular Disorders:** Epistaxis, flushing.

**Post-Market Adverse Drug Reactions**

In addition to adverse events identified in clinical trials, the following post-marketing events have been included due to their seriousness, frequency of reporting, potential causal association with SITAVIG™, or a combination of these factors. Because they are reported spontaneously from a population of unknown size, estimates of incidence cannot be made.

**Allergic:** Swelling of tongue and lip, glossitis and cheilitis.

**Central Nervous System:** Nightmares, agitation and insomnia.

**Other:** Incorrect route of administration.

**DRUG INTERACTIONS**

**Overview**

No interaction studies have been performed with SITAVIG™. Acyclovir is primarily eliminated unchanged in the urine via active tubular secretion. Drugs administered concomitantly that compete with tubular secretion may increase acyclovir plasma concentrations. However, due to the low dose and minimal systemic absorption of SITAVIG™, systemic drug interactions are unlikely.

**Drug-Food Interactions**

There was no formal food effect study conducted with SITAVIG™; however, in clinical studies
patients were allowed to eat and drink while taking SITAVIG™.

**Drug-Laboratory Interactions**

Interactions of SITAVIG with laboratory tests have not been established.

**DOSAGE AND ADMINISTRATION**

**Dosing Considerations**

SITAVIG™ is a single dose therapy for the treatment of recurrent herpes labialis. Therapy should be initiated at the onset of prodromal symptoms, and prior to the appearance of any signs of herpes labialis lesions.

In addition,
- SITAVIG™ should not be crushed, chewed, sucked or swallowed.
- Food and drink can be taken normally when SITAVIG™ is in place.
- Avoid any situations which may interfere with adhesion of the tablet such as chewing gum, touching or pressing the tablet after placement, wearing upper denture, and brushing teeth. If the teeth need to be cleaned while the tablet is in place, rinse the mouth gently.
- Drink plenty of liquids in the case of dry mouth.

**Recommended Dose and Dosage Adjustment**

One SITAVIG™ tablet should be applied as a single dose to the upper gum region (canine fossa). Once applied, the tablet stays in position and should be left in place until it dissolves.

**Administration**

SITAVIG™ should be applied with a dry finger immediately after taking it out of the blister. The rounded side of the tablet should be placed to the upper gum just above the incisor tooth (canine fossa) and held in place with a slight pressure over the upper lip for 30 seconds to ensure adhesion. For comfort the rounded side should be placed to the upper gum with the flat side facing the inside of the lip.

- If SITAVIG™ does not adhere or falls off within the first 6 hours, the same tablet should be repositioned immediately. If the tablet cannot be repositioned, a new tablet should be placed.
- If SITAVIG™ is swallowed within the first 6 hours, the patient should drink a glass of water and a new tablet should be applied.
- SITAVIG™ does not need to be reapplied if the tablet falls out or is swallowed after the first 6 hours.
OVERDOSAGE

Acyclovir absorption and systemic exposure following application of SITAVIG™ are minimal. Overdose is therefore unlikely.

Symptomatic and supportive care is the basis for management.

For management of a suspected drug overdose, contact your regional Poison Control Centre.

ACTION AND CLINICAL PHARMACOLOGY

Mechanism of Action

Acyclovir is an antiviral drug. It is a synthetic purine nucleoside with inhibitory activity against herpes simplex virus type 1 (HSV-1) and type 2 (HSV-2). Acyclovir is phosphorylated intracellularly by the viral encoded thymidine kinase (TK) of HSV into acyclovir monophosphate, a nucleotide analogue. The monophosphate is further converted into diphosphate by cellular guanylate kinase and into triphosphate by a number of cellular enzymes. In a biochemical reaction, acyclovir triphosphate inhibits replication of herpes viral DNA by competing with nucleotides for binding to the viral DNA polymerase and by incorporation into and termination of the growing viral DNA chain. The cellular thymidine kinase of normal, uninfected cells does not use acyclovir effectively as a substrate, hence toxicity to mammalian host cells is low.

Pharmacokinetics

Table 2 - Summary of Acyclovir Buccal Tablet’s Pharmacokinetic Parameters in Saliva Following Application of a Single SITAVIG™ 50 mg Tablet in Healthy Volunteers (N = 12)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Single dose mean ± SD</th>
<th>(Min – Max)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cmax (ng/mL)</td>
<td>440 ± 241</td>
<td>(149 – 959)</td>
</tr>
<tr>
<td>Tmax (h)</td>
<td>7.95 ± 4.08</td>
<td>(3.07 – 18.05)</td>
</tr>
<tr>
<td>AUC0-24h (h·ng/mL)</td>
<td>2900 ± 2400</td>
<td>(849 – 9450)</td>
</tr>
</tbody>
</table>

Absorption: SITAVIG™ is applied topically to the gum and releases acyclovir as the buccal tablet gradually dissolves. As shown in Table 2, single dose application of SITAVIG containing 50 mg of acyclovir to the buccal mucosa in 12 healthy volunteers provided mean maximum salivary concentrations of 440 μg/mL 8 hours after application of the tablet. The single local application of SITAVIG provided rapid (<30 min), high (≥IC50) and sustained (≥ 24 hours) acyclovir concentrations in saliva and in labial mucosa, the replication site of HSV-1.

In the Phase 3 study, the levels of acyclovir in saliva were measured within 24 hours of SITAVIG™ application in 56 patients with recurrent herpes labialis (mean value 88.1
micrograms per mL) and were within the range of those observed in the pharmacokinetic study in healthy volunteers.

In healthy volunteers, the median duration of buccal adhesion was 14 hours following application of a single SITAVIG™ 50 mg tablet.

**Distribution:** Plasma concentrations of acyclovir were measured in 12 healthy volunteers after a single-dose application of SITAVIG™ 50 mg buccal tablet. Acyclovir concentrations had a delayed appearance (undetectable at 5 hours) and were below the concentrations required for antiviral activity (range: 17.5 to 55.3 ng/mL).

**Metabolism:** Acyclovir is metabolized to 9-[(carboxymethoxy)methyl]guanine (CMMG) and 8-hydroxy-acyclovir (8-OH-ACV) by oxidation and hydroxylation.

**Excretion:** Acyclovir is primarily excreted unchanged by the kidneys.

**STORAGE AND STABILITY**

SITAVIG™ should be stored at 20 to 25°C. Protect from moisture. The shelf life is 36 months.

Keep out of reach and sight of children.

**DOSAGE FORMS, COMPOSITION AND PACKAGING**

SITAVIG™ contains 50 mg of acyclovir and the following inactive ingredients: colloidal silicon dioxide, hypromellose, magnesium stearate, microcrystalline cellulose, milk protein concentrate, povidone and sodium lauryl sulfate.

SITAVIG™ is supplied as a round white to slightly yellow tablet, with a rounded side and a flat side, debossed with an “AL21” on the flat side. SITAVIG™ tablets are packaged into boxes of 2 blisters, each blister containing 1 tablet.
PART II: SCIENTIFIC INFORMATION

PHARMACEUTICAL INFORMATION

Drug Substance

Proper name: acyclovir; aciclovir

Chemical name: 6H-Purin-6-one, 2-amino-1,9-dihydro-9-[(2-hydroxyethoxy)methyl]-9-[(2-Hydroxyethoxy)methyl]guanine

Molecular formula and molecular mass: C_8H_11N_5O_3, 225.2 g/mol

Structural formula:

![Structural formula of acyclovir]

Physicochemical properties: Acyclovir drug substance is a white or almost white crystalline powder. Freely soluble in dimethylsulfoxide, slightly soluble in dimethylformamide, very slightly soluble in methanol and water.

CLINICAL TRIALS

The efficacy and safety of SITAVIG™ (acyclovir mucoadhesive buccal tablets) was evaluated in a randomized, double-blind, placebo-controlled, patient-initiated, multicenter trial. SITAVIG™ was administered as a single dose (n = 378) versus placebo (n = 397) in patients with recurrent herpes labialis (cold sores). All patients had at least 4 herpes episodes in the previous year of whom 68.4% had ≥ 5 episodes. The mean duration of previous herpes episodes was 8.6 (± 3.1) days in the SITAVIG™ group and 8.3 (± 3.0) days in the placebo group.

Patients were instructed to initiate treatment within one hour after the onset of prodromal symptoms and before the appearance of any signs of herpes labialis lesions by applying the tablet to the buccal mucosa in the canine fossa. If the tablet was detached within the first 6 hours, subjects were instructed to reapply a tablet.

Table 3 provides a summary of the Patient Demographics for the Clinical Trial in Recurrent Herpes Labialis.
Study demographics and trial design

Table 3 - Summary of Patient Demographics for Clinical Trial in Recurrent Herpes Labialis

<table>
<thead>
<tr>
<th>Study #</th>
<th>Trial Design</th>
<th>Dosage, Route of Administration and Duration</th>
<th>Study Subjects – ITT Population (n=771)</th>
<th>Mean Age (Range)</th>
<th>Gender M/F</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA2005/21/02</td>
<td>Multicentre, multinational, randomized, double-blind</td>
<td>SITAVIG™ 50 mg Placebo Single-dose mucoadhesive tablet applied to the gum</td>
<td>376 395</td>
<td>40 (18 – 80) 42 (18 – 73)</td>
<td>118/258 124/271</td>
</tr>
</tbody>
</table>

Study results

The mean and median durations of the recurrent herpes labialis episode (ITT population, n = 771) were approximately half a day shorter in patients treated with SITAVIG™ compared with patients treated with placebo.
DETAILED PHARMACOLOGY

Pharmacokinetics/Pharmacodynamics

A single-center, randomized, cross-over, pharmacokinetic/pharmacodynamics study was conducted in 12 healthy subjects to compare the pharmacokinetic parameters and tolerability of a single dose of SITAVIG™ in plasma, saliva and labial mucosa to those of a single acyclovir 200 mg oral (systemic) tablet. SITAVIG™ was applied in the morning to the upper gum as a single dose. The comparator, acyclovir 200 mg oral tablet, was administered only once in the morning.

Acyclovir saliva, mucosa and plasma concentrations, duration of SITAVIG™ adhesion, local and general tolerability were assessed.

The single local application of SITAVIG™ provided rapid (<30 min), high (≥IC50) and sustained (≥ 24 hours) acyclovir concentrations in saliva and in labial mucosa, markedly over the IC50 (22.5 ng/mL) and those obtained after a single administration of acyclovir 200 mg tablet (11,700-fold higher for saliva and at least 4-fold higher for labial mucosa). In contrast, plasma concentrations were lower than those observed with acyclovir 200 mg oral tablet with a relative bioavailability corrected by the dose of 49% for SITAVIG™.

The detection of very high acyclovir concentrations in saliva and labial mucosa persisting several hours after SITAVIG™ dislodgment or complete erosion, and the higher acyclovir concentrations in labial mucosa than in saliva support the assumption of acyclovir storage in mucosa.

SITAVIG™ provides low (below the IC50) plasma concentrations and high (over the IC50) concentrations in saliva and labial mucosa. Rapid, high and sustained concentrations of acyclovir at the replication site of HSV-1, allow for a prolonged inhibition of the virus and thereby circumvent the poor intracellular half-life of acyclovir triphosphate.

MICROBIOLOGY

Antiviral Activity

The quantitative relationship between the cell culture susceptibility of herpes viruses to antivirals and the clinical response to therapy has not been established in humans, and virus sensitivity testing has not been standardized. Sensitivity testing results, expressed as the concentration of drug required to inhibit by 50% the growth of virus in cell culture (EC50), vary greatly depending upon a number of factors. Using plaque-reduction assays on Vero cells, the median EC50 value of acyclovir against clinical herpes virus isolates (subjects receiving placebo) was 1.3 μM (range: < 0.56 to 3.3 μM).
Drug Resistance

Resistance of HSV to acyclovir can result from qualitative and quantitative changes in the viral TK and/or DNA polymerase. Clinical isolates of HSV with reduced susceptibility to acyclovir have been recovered from immunocompromised subjects, especially with advanced HIV infection. While most of the acyclovir-resistant mutant isolates from immunocompromised subjects thus far have been found to be TK-deficient, other mutant isolates involving the viral TK gene (TK partial and TK altered) or DNA polymerase have been identified. TK-negative mutants may cause severe disease in infants and immunocompromised adults.

The possibility of viral resistance to acyclovir should be considered in immunocompromised subjects who show poor clinical response during therapy.

TOXICOLOGY

Acute Studies

Single doses of acyclovir by intravenous, oral and intraperitoneal routes, were administered to rats, and no deaths were observed at the maximum practical dose volume. The oral median lethal dose (LD$_{50}$) of acyclovir in rats was greater than 20,000 mg/kg. The intravenous LD$_{50}$ was greater than 600 mg/kg, and the intraperitoneal LD$_{50}$ values in male and female rats were 1,305 and 1,210 mg/kg, respectively.

Mice were also treated with single doses of acyclovir by intravenous, oral and intraperitoneal routes, and no deaths were observed at the maximum practical dose volume. The oral LD$_{50}$ of acyclovir was greater than 10,000 mg/kg. The intravenous LD$_{50}$ was 405 mg/kg, for male mice and greater than 600 mg/kg for male rats and the intraperitoneal LD$_{50}$ values in male and female rats were 1,454 and 999 mg/kg, respectively.

Repeat-Dose Studies

Acyclovir has been studied in repeat-dose toxicity studies in mice (1-month oral administration), rats (21-days intravenous administration) and Beagle dogs (1-month intravenous and 12-months oral administration).

In mice, there were no deaths and body weight gains and food consumption were comparable for control and treated mice. No signs of accumulation of acyclovir in plasma were apparent.

Treatment-related changes in rats were limited to the kidney and occurred at doses of 20 mg/kg/day and higher. These changes included increases in water consumption, urine output, blood urea nitrogen and absolute and relative kidney weights. The renal damage observed was reversible and affected parameters returned to normal after a 15-day post-dose recovery period. The No Observed Effect Level (NOEL) in rats was determined to be 10 mg/kg/day.

In studies of intravenous administration in dogs, high toxicity was revealed by 200 mg/kg/day (all dogs died by Day 8) and 100 mg/kg/day (5/8 dogs died between Day 21 and Day 31). Water
intake and urine output generally increased for dogs treated at dose levels of 40 mg/kg/day or higher. The increase in urine output was partially responsible for the clinical apparent dehydration, and was the sign of renal injury (decreased ability to concentrate urine). Microscopic alterations were only observed in the dogs receiving 50 mg/kg/day.

Evaluation of plasma levels in dogs administered 100 and 200 mg/kg/day revealed very high levels in the range of 45 to 254 μg/mL (200 to 1127 μM), whereas the 50 mg/kg/day animals had plasma levels in the 22.5 to 45 μg/mL (100 to 200 μM) range. All dogs given 50 mg/kg/day survived, and clinical signs of toxicity were limited. Dogs treated at 40 mg/kg/day had a slight increase in values for water intake and produced an increased volume of urine having decreased specific gravity and decreased osmolarity. There were no signs of toxicity observed in dogs given 20 mg/kg/day.

In studies of oral administration in dogs, the dose levels were decreased to 30 and 60 mg/kg/day administered three times a day after Day 15, due to acute gastrointestinal toxicity at 45 and 150 mg/kg/day. No toxic effects were observed at the revised dose levels during the remainder of the study. Acyclovir up to 60 mg/kg/day was well tolerated by beagle dogs. Signs of tenderness in the forepaws and breaking and loosening of nails were observed in the 45 and 150 mg/kg/day groups.

**Local Tolerance**

The local tolerance effect of SITAVIG™ in hamsters was evaluated using the cheek pouch technique. Seven days after application of SITAVIG™ into the hamster cheek pouch, there were no significant effects. The dosage used in this tolerance study is the human dosage that corresponds to a single dose of 500 mg/kg of SITAVIG™ in a hamster.

**Carcinogenicity**

Acyclovir was tested for carcinogenic potential in lifetime bioassays in rats and mice treated once daily at doses of 0, 50, 150 and 450 mg/kg administered by gavage. All rats and mice were observed for changes in physical condition, toxicosis, body weight, food consumption, and for death. All rats and mice found dead were necropsied. The study was considered as terminated once mortality decreased a group size to approximately 20% of the number of animals of that sex present in the test group when the study started. This was achieved after 110 weeks for male rats and 122 weeks for female rats, and 126 weeks for male mice and 111 weeks for female mice.

In rats, no treatment-related alteration in body weight of food consumption was observed. Ophthalmoscopic examination did not reveal any adverse effects related to the chronic administration of acyclovir. The effect of acyclovir on survival was assessed and revealed that acyclovir neither increased nor decreased the survival of the rats. The effect of treatment on the overall incidences of benign tumour bearing animals (TBAs), malignant TBAs, and total TBAs did not reveal any significant difference between the control group and the treated groups. None of the rats had unusual neoplasms, and there were no more benign or malignant neoplasms in treated rats at all dose levels than in the control group. Acyclovir was not carcinogenic as tested in rats.
In male mice, acyclovir neither increased nor decreased survival. However, in the 150 mg/kg and 450 mg/kg groups, female mice survived significantly longer than the control female mice group. The effect of treatment on the overall incidences of benign tumour bearing animals (TBAs), malignant TBAs, and total TBAs did not reveal any significant difference between the control group and the treated groups. None of the mice had unusual neoplasms, and chronic treatment with acyclovir did not produce non-neoplastic lesions or alter the incidence or severity of common spontaneous disease processes in mice. Acyclovir was not carcinogenic as tested in mice.

**Reproduction**

Animal reproduction studies have not been conducted with SITAVIG™. Acyclovir was not teratogenic in the rat, rabbit and mouse at exposures greatly in excess of human exposure.

Rats and rabbits were administered 0, 12, 25 and 50 mg/kg/day by subcutaneous injection from Day 6 through Day 15 of gestation in rats and Day 6 through Day 18 of gestation in rabbits. There were no signs of fetotoxicity or maternal toxicity in either species.

Fertility studies were conducted in mice administered 0, 50, 150 and 450 mg/kg/day. The two generation reproduction-fertility test in mice demonstrated that acyclovir did not interfere with fertility, breeding, conception, fetal development, parturition or lactation. Pert- and postnatal developmental indices were normal for all mice.

**REFERENCES**


READ THIS FOR SAFE AND EFFECTIVE USE OF YOUR MEDICINE
PATIENT MEDICATION INFORMATION

PrSITAVIG™
Acyclovir mucoadhesive buccal tablets

Read this carefully before you start taking SITAVIG™ and each time you refill your medication. This leaflet is a summary and will not tell you everything about this drug. Talk to your healthcare professional about your medical condition and treatment and ask if there is any new information about SITAVIG™.

What is SITAVIG™ used for?
- SITAVIG™ is a medication that is applied once to treat outbreaks of cold sores (herpes labialis) in adults.

How does SITAVIG™ work?
SITAVIG™ works by:
- Sticking to the gum, when placed on the upper gum by the incisor tooth (the tooth just to the right or left of your two front teeth).
- Over about 14 hours, the tablet dissolves.
- SITAVIG™ then:
  - Prevents the virus from multiplying.
  - This reduces the infection and symptoms such as burning, tingling, redness and itching.

What are the ingredients in SITAVIG™?
Medicinal ingredients: acyclovir

Non-medicinal ingredients: colloidal silicon dioxide, hypromellose, magnesium stearate, microcrystalline cellulose, milk protein concentrate, povidone, sodium lauryl sulfate.

SITAVIG™ comes in the following dosage form:
Mucoadhesive buccal tablet, 50 mg

Do not use SITAVIG™ if you:
- are allergic to acyclovir, milk protein concentrate, or any other ingredient of the product (see What are the ingredients of SITAVIG™?).

To help avoid side effects and ensure proper use, talk to your healthcare professional before you take SITAVIG™. Talk about any health conditions or problems you may have, including if you:
- have a weak immune system (become sick very easily).
- are pregnant or plan to become pregnant.
- are breastfeeding or plan to breastfeed.
- are under the age of 18 years.
Tell your healthcare professional about all the medicines you take, including any drugs, vitamins, minerals, natural supplements or alternative medicines.

The following may interact with SITAVIG™:
Other medicines are unlikely to affect how SITAVIG™ works or increase the chance of side effects. Drug interaction studies have not been done with SITAVIG™.

How to use SITAVIG™:
• Use SITAVIG™ exactly as your healthcare professional tells you.
• Do not crush, suck, chew or swallow the tablets.
• Use SITAVIG™:
  • as soon as possible for best results.
  • at first signs of a cold sore, such as burning, itching, tingling, redness.
  • before a cold sore appears.
• Apply the rounded side of SITAVIG™ to the upper gum, just above the incisor tooth.
• Do not apply SITAVIG™ to the lip or cheek.
• If SITAVIG™ does not stick to the gum or falls off:
  • before 6 hours, then:
    • reapply, or
    • if this does not stick, apply a new tablet.
  • after 6 hours, then do not reapply or apply a new tablet.
• If you swallow SITAVIG™:
  • before 6 hours, then drink a glass of water and apply a new tablet.
  • after 6 hours, then do not apply a new tablet.
• While using SITAVIG™:
  • you may eat and drink.
  • drink more liquids if your mouth becomes dry.
  • do not chew gum, wear dentures, brush teeth, or do any similar activity which may move the tablet.
  • if the teeth need to be cleaned while you are using the tablet, rinse the mouth gently.

How to apply SITAVIG™:

Step 1: Before you apply SITAVIG™, find the area on your upper gum, just above either the left or the right incisor. The incisor tooth is the tooth just to the right or left of your two front teeth (See Figure A). This is where you should apply SITAVIG™.
Step 2: Carefully PEEL back the cover of the blister pack. Take the SITAVIG™ tablet out of the blister pack. When removed from the blister pack, SITAVIG™ must be used right away. SITAVIG™ is round on one side and flat on the other side (See Figure B).

Step 3: Place the flat side of SITAVIG™ on your dry fingertip. Apply the round side of SITAVIG™ to your upper gum (See Figure C). The flat side will be facing the inside of your lip.

Step 4: Hold SITAVIG™ in place by pressing slightly with your finger on the outside of your upper lip, over the area where SITAVIG is placed, for 30 seconds. This will help SITAVIG™ stick to your gum (See Figure D).
Step 5: Leave the SITAVIG™ tablet in place until it dissolves, which can take about 14 hours.

Usual adult dose: 
SITAVIG™ is a single dose therapy. This means that only one tablet is applied to the upper gum, just once.

One SITAVIG™ tablet should be applied as a single dose to the area of the upper gum.

Overdose:

If you think you have taken too much SITAVIG™, contact your healthcare professional, hospital emergency department or regional Poison Control Centre immediately, even if there are no symptoms.

What are possible side effects from using SITAVIG™?
These are not all the possible side effects you may feel when taking SITAVIG™. If you experience any side effects not listed here, contact your healthcare professional. Please also see Warnings and Precautions.

- Headache
- Pain, discomfort, redness or tingling at or near where tablet was applied
- Lip dryness or swelling
- Mouth dryness, pain or ulcer
- Acid reflux disease (condition where stomach acid comes back up into your esophagus (the muscular tube that connects your throat and stomach))
- Nosebleeds
- Dizziness
- Tiredness
- Pain or discomfort in the abdomen
- Diarrhoea
- Nausea
- Vomiting
- Flu-like illness
- Muscle pain
- Redness, flushing, rash or itching

### Reporting Side Effects

You can report any suspected side effects associated with the use of health products to Health Canada by:

- Visiting the Web page on Adverse Reaction Reporting (http://www.hc-sc.gc.ca/dhp- mps/medeff/report-declaration/index-eng.php) for information on how to report online, by mail or by fax; or
- Calling toll-free at 1-866-234-2345.

*NOTE: Contact your healthcare professional if you need information about how to manage your side effects. The Canada Vigilance Program does not provide medical advice.*

### Storage:

Store at 20 - 25°C. Protect from moisture.

Keep out of reach and sight of children.

### If you want more information about SITAVIG™:

- Talk to your healthcare professional
- Find the full Product Monograph that is prepared for healthcare professionals and includes this Patient Medication Information by visiting the Health Canada website (www.healthcanada.gc.ca); the manufacturer’s website www.cipherpharma.com, or by calling 1-888-361-7207.

This leaflet was prepared by Cipher Pharmaceuticals Inc.

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