ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS
1. **NAME OF THE MEDICINAL PRODUCT**

Sycrest 5 mg sublingual tablets

2. **QUALITATIVE AND QUANTITATIVE COMPOSITION**

Each sublingual tablet contains 5 mg asenapine (as maleate).

For the full list of excipients, see section 6.1.

3. **PHARMACEUTICAL FORM**

Sublingual tablet
Round, white to off-white, sublingual tablets debossed with “5” on one side.

4. **CLINICAL PARTICULARS**

4.1 **Therapeutic indications**

Sycrest is indicated for the treatment of moderate to severe manic episodes associated with bipolar I disorder in adults.

4.2 **Posology and method of administration**

**Posology**

The recommended starting dose of Sycrest as monotherapy is 5 mg twice daily. One dose should be taken in the morning and one dose should be taken in the evening. The dose can be increased to 10 mg twice daily based on individual clinical response and tolerability. See section 5.1. For combination therapy a starting dose of 5 mg twice daily is recommended. Depending on the clinical response and tolerability in the individual patient, the dose can be increased to 10 mg twice daily.

**Special populations**

**Elderly**

Sycrest should be used with care in the elderly. Limited data on efficacy in patients 65 years of age and older are available. Available pharmacokinetic data are described in section 5.2.

**Renal impairment**

No dose adjustment is required for patients with renal impairment. There is no experience with asenapine in patients with severe renal impairment who have a creatinine clearance less than 15 mL/min.

**Hepatic impairment**

No dose adjustment is required for patients with mild hepatic impairment. The possibility of elevated asenapine plasma levels cannot be excluded in some patients with moderate hepatic impairment (Child-Pugh B) and caution is advised. In subjects with severe hepatic impairment (Child-Pugh C), a 7-fold increase in asenapine exposure was observed. Thus, Sycrest is not recommended in patients with severe hepatic impairment.

**Paediatric population**

A pharmacokinetic study and a short term efficacy and safety study were performed in a paediatric population (ages 10-17 years) with manic or mixed episodes associated with bipolar I disorder. Long
term safety in this population was explored in a 50-week, open-label, uncontrolled extension study. Currently available data are described in sections 4.8, 5.1 and 5.2 but no recommendation on a posology can be made.

Method of administration

The tablet should not be removed from the blister until ready to take it. Dry hands should be used when touching the tablet. The tablet should not be pushed through the tablet pack. The tablet pack should not be cut or torn. The coloured tab should be peeled back and the tablet should be removed gently. The tablet should not be crushed.

To ensure optimal absorption, the Sycrest sublingual tablet should be placed under the tongue and allowed to dissolve completely. The tablet will dissolve in saliva within seconds. Sycrest sublingual tablets should not be chewed or swallowed. Eating and drinking should be avoided for 10 minutes after administration.
When used in combination with other medicinal products, Sycrest should be taken last.

Treatment with Sycrest is not advised in patients who are unable to comply with this method of administration, as the bioavailability of asenapine when swallowed is low (<2% with an oral tablet formulation).

4.3 Contraindications

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

4.4 Special warnings and precautions for use

Elderly patients with dementia-related psychosis
Elderly patients with dementia-related psychosis treated with antipsychotic substances are at an increased risk of death.
Sycrest is not approved for the treatment of patients with dementia-related psychosis and is not recommended for use in this particular group of patients.

Neuroleptic malignant syndrome
Neuroleptic malignant syndrome (NMS), characterised by hyperthermia, muscle rigidity, autonomic instability, altered consciousness and elevated serum creatine phosphokinase levels, has been reported to occur with antipsychotics, including asenapine. Additional clinical signs may include myoglobinuria (rhabdomyolysis) and acute renal failure.
If a patient develops signs and symptoms indicative of NMS Sycrest must be discontinued.

Seizures
In clinical trials, cases of seizure were occasionally reported during treatment with asenapine. Therefore, Sycrest should be used with caution in patients who have a history of seizure disorder or have conditions associated with seizures.

Suicide
The possibility of a suicide attempt is inherent in psychotic illnesses and bipolar disorder and close supervision of high-risk patients should accompany treatment.

Orthostatic hypotension
Asenapine may induce orthostatic hypotension and syncope, especially early in treatment, probably reflecting its α1-adrenergic antagonist properties. Elderly patients are particularly at risk for experiencing orthostatic hypotension (see section 4.8). In clinical trials, cases of syncope were occasionally reported during treatment with Sycrest. Sycrest should be used with caution in elderly patients and in patients with known cardiovascular disease (e.g., heart failure, myocardial infarction
or ischemia, conduction abnormalities), cerebrovascular disease, or conditions that predispose the patient to hypotension (e.g., dehydration and hypovolemia).

**Tardive dyskinesia**

Medicinal products with dopamine receptor antagonistic properties have been associated with the induction of tardive dyskinesia characterised by rhythmical, involuntary movements, predominantly of the tongue and/or face. In clinical trials, cases of tardive dyskinesia were occasionally reported during treatment with asenapine. The onset of extrapyramidal symptoms is a risk factor for tardive dyskinesia. If signs and symptoms of tardive dyskinesia appear in a patient on Sycrest, discontinuation of treatment should be considered.

**Hyperprolactinaemia**

Increases in prolactin levels were observed in some patients with Sycrest. In clinical trials, there were few adverse reactions related to abnormal prolactin levels reported.

**QT interval**

Clinically relevant QT prolongation does not appear to be associated with asenapine. Caution should be exercised when Sycrest is prescribed in patients with known cardiovascular disease or family history of QT prolongation, and in concomitant use with other medicinal products thought to prolong the QT interval.

**Hyperglycaemia and diabetes mellitus**

Hyperglycaemia or exacerbation of pre-existing diabetes has occasionally been reported during treatment with asenapine. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes mellitus in patients with schizophrenia or bipolar disorder and the increasing incidence of diabetes mellitus in the general population. Appropriate clinical monitoring is advisable in diabetic patients and in patients with risk factors for the development of diabetes mellitus.

**Dysphagia**

Esophageal dysmotility and aspiration have been associated with antipsychotic treatment. Cases of dysphagia were occasionally reported in patients treated with Sycrest.

**Body temperature regulation**

Disruption of the body’s ability to reduce core body temperature has been attributed to antipsychotic medicines. From the clinical trials, it is concluded that clinically relevant body temperature dysregulation does not appear to be associated with asenapine. Appropriate care is advised when prescribing Sycrest for patients who will be experiencing conditions that may contribute to an elevation in core body temperature, e.g. exercising strenuously, exposure to extreme heat, receiving concomitant medicinal products with anticholinergic activity or being subject to dehydration.

**Patients with severe hepatic impairment**

Asenapine exposure is increased 7-fold in patients with severe hepatic impairment (Child-Pugh C). Therefore, Sycrest is not recommended in such patients.

**Parkinson’s disease and dementia with Lewy bodies**

Physicians should weigh the risks versus the benefits when prescribing Sycrest to patients with Parkinson’s disease or dementia with Lewy Bodies (DLB) since both groups may be at increased risk of neuroleptic malignant syndrome as well as having an increased sensitivity to antipsychotics. Manifestation of this increased sensitivity can include confusion, obtundation, postural instability with frequent falls, in addition to extrapyramidal symptoms.

**Falls**

Asenapine may cause adverse effects such as somnolence, orthostatic hypotension, dizziness and extrapyramidal symptoms, which may lead to falls and, consequently, fractures or other injuries. Patients at risk for fall should be evaluated prior to prescribing asenapine.
4.5 Interaction with other medicinal products and other forms of interaction

Given the primary effects of asenapine on the central nervous system (CNS) (see section 4.8), caution should be used when it is taken in combination with other centrally acting medicinal products. Patients should be advised to avoid alcohol while taking Sycrest.

Potential for other medicines to affect Sycrest
Asenapine is cleared primarily through direct glucuronidation by UGT1A4 and oxidative metabolism by cytochrome P450 isoenzymes (predominantly CYP1A2). The potential effects of inhibitors and an inducer of several of these enzyme pathways on asenapine pharmacokinetics were studied, specifically fluvoxamine (CYP1A2 inhibitor), paroxetine (CYP2D6 inhibitor), imipramine (CYP1A2/2C19/3A4 inhibitor), cimetidine (CYP3A4/2D6/1A2 inhibitor), carbamazepine (CYP3A4/1A2 inducer) and valproate (UGT inhibitor). Except for fluvoxamine, none of the interacting medicinal products resulted in clinically relevant alterations in asenapine pharmacokinetics. During combined administration with a single dose of asenapine 5 mg, fluvoxamine 25 mg twice daily resulted in a 29% increase in asenapine AUC. The full therapeutic dose of fluvoxamine would be expected to produce a greater increase in asenapine plasma concentrations. Therefore, co-administration of asenapine and fluvoxamine should be approached with caution.

Potential for Sycrest to affect other medicines
Because of its α1-adrenergic antagonism with potential for inducing orthostatic hypotension (see section 4.4), Sycrest may enhance the effects of certain antihypertensive agents.

Asenapine may antagonise the effect of levodopa and dopamine agonists. If this combination is deemed necessary, the lowest effective dose of each treatment should be prescribed.

In vitro studies indicate that asenapine weakly inhibits CYP2D6. Clinical drug interaction studies investigating the effects of CYP2D6 inhibition by asenapine showed the following results:

- Following co-administration of dextromethorphan and asenapine in healthy subjects, the ratio of dextrorphan/dextromethorphan (DX/DM) as a marker of CYP2D6 activity was measured. Indicative of CYP2D6 inhibition, treatment with asenapine 5 mg twice daily resulted in a fractional decrease in DX/DM ratio to 0.43. In the same study, treatment with paroxetine 20 mg daily decreased the DX/DM ratio to 0.032.

- In a separate study, co-administration of a single 75 mg dose of imipramine with a single 5 mg dose of asenapine did not affect the plasma concentrations of the metabolite desipramine (a CYP2D6 substrate).

- Co-administration of a single 20 mg dose of paroxetine (a CYP2D6 substrate and inhibitor) during treatment with 5 mg asenapine twice daily in 15 healthy male subjects resulted in an almost 2-fold increase in paroxetine exposure.

In vivo asenapine appears to be at most a weak inhibitor of CYP2D6. However, asenapine may enhance the inhibitory effects of paroxetine on its own metabolism. Therefore, Sycrest should be co-administered cautiously with medicinal products that are both substrates and inhibitors for CYP2D6.

4.6 Fertility, pregnancy and lactation

Pregnancy
There are no adequate data from the use of Sycrest in pregnant women. Asenapine was not teratogenic in animal studies. Maternal and embryo toxic effects were found in animal studies (see section 5.3).
Newborn infants exposed to antipsychotics (including Sycrest) during the third trimester of pregnancy are at risk of adverse reactions including extrapyramidal and/or withdrawal symptoms that may vary in severity and duration following delivery. There have been reports of agitation, hypertonia, hypotonia, tremor, somnolence, respiratory distress, or feeding disorder in newborn infants. Consequently, newborn infants should be monitored carefully.

Sycrest should not be used during pregnancy unless the clinical condition of the woman requires treatment with asenapine and only if the potential benefit outweighs the potential risk to the foetus.

**Breast-feeding**
Asenapine was excreted in milk of rats during lactation. It is not known whether asenapine or its metabolites are excreted in human milk. Breast-feeding should be discontinued during treatment with Sycrest.

**Fertility**
No impairment of fertility has been observed in nonclinical studies (see section 5.3).

### 4.7 Effects on ability to drive and use machines

Asenapine may cause somnolence and sedation. Therefore, patients should be cautioned about driving and using machines until they are reasonably certain that Sycrest therapy does not affect them adversely.

### 4.8 Undesirable effects

**Summary of safety profile**
The most frequently reported adverse drug reactions (ADRs) associated with the use of asenapine in clinical trials were somnolence and anxiety. Serious hypersensitivity reactions have been reported. Other serious ADRs are discussed in more detail in section 4.4.

**Tabulated list of adverse reactions**
The incidences of the ADRs associated with asenapine therapy are tabulated below. The table is based on adverse reactions reported during clinical trials and/or post-marketing use. All ADRs are listed by system organ class and frequency; very common (≥ 1/10), common (≥ 1/100 to < 1/10), uncommon (≥ 1/1,000 to < 1/100), rare (≥ 1/10,000 to < 1/1,000) and not known (cannot be estimated from the available data). Within each frequency grouping, ADRs are presented in order of decreasing seriousness.

<table>
<thead>
<tr>
<th>System organ class</th>
<th>Very common</th>
<th>Common</th>
<th>Uncommon</th>
<th>Rare</th>
<th>Not known</th>
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<tbody>
<tr>
<td>Blood and lymphatic disorders</td>
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<td></td>
<td>Neutropenia</td>
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<td>Immune system disorders</td>
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<td>Allergic reactions</td>
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<td>Metabolism and nutrition disorders</td>
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<td></td>
<td>Hyperglycaemia</td>
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<tr>
<td>Psychiatric disorders</td>
<td>Anxiety</td>
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<tr>
<td>System organ class</td>
<td>Very common</td>
<td>Common</td>
<td>Uncommon</td>
<td>Rare</td>
<td>Not known</td>
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<tr>
<td>Nervous system disorders</td>
<td>Somnolence</td>
<td>Dystonia Akathisia Dyskinesia Parkinsonism Sedation Dizziness Dysgeusia</td>
<td>Syncope Seizure Extrapyramidal disorder Dysarthria Restless legs syndrome</td>
<td>Neuroleptic malignant syndrome</td>
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<td>Eye disorders</td>
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<td>Accommodation disorder</td>
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<td>Cardiac disorders</td>
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<td>Sinus bradycardia Bundle branch block Electrocardiogram QT prolonged Sinus tachycardia</td>
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<td>Vascular disorders</td>
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<td>Orthostatic hypotension Hypotension</td>
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<tr>
<td>Respiratory, thoracic and mediastinal disorders</td>
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<td>Pulmonary embolism</td>
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<tr>
<td>Gastrointestinal disorders</td>
<td>Hypoaesthesia oral Nausea Salivary hypersecretion</td>
<td>Swollen tongue Dysphagia Glossodynia Paraesthesia oral Oral mucosal lesions (ulcerations, blistering and inflammation)</td>
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<tr>
<td>Hepatobiliary disorders</td>
<td>Alanine aminotransferase increased</td>
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<tr>
<td>Injury, poisoning and procedural complications</td>
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<td></td>
<td>Falls*</td>
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<tr>
<td>Musculoskeletal and connective tissue disorders</td>
<td>Muscle rigidity</td>
<td>Rhabdomyolysis</td>
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<td>Pregnancy, puerperium and perinatal conditions</td>
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<td></td>
<td></td>
<td>Drug withdrawal syndrome neonatal (see 4.6)</td>
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<tr>
<td>System organ class</td>
<td>Very common</td>
<td>Common</td>
<td>Uncommon</td>
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<td>Reproductive system and breast disorders</td>
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<td>Sexual dysfunction</td>
<td>Gynaecomastia</td>
<td>Galactorrhoea</td>
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<tr>
<td>General disorders and administration site conditions</td>
<td>Fatigue</td>
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</table>

* See subsection “Falls” below

**Description of selected adverse reactions**

**Extrapyramidal Symptoms (EPS)**
In clinical trials, the incidence of extrapyramidal symptoms in asenapine-treated patients was higher than placebo (15.4 % vs 11.0 %).
From the short-term (6 weeks) schizophrenia trials there appears to be a dose-response relationship for akathisia in patients treated with asenapine, and for parkinsonism there was an increasing trend with higher doses.
Based on a small pharmacokinetic study, paediatric patients appeared to be more sensitive to dystonia with initial dosing with asenapine when a gradual up-titration schedule was not followed (see section 5.2). The incidence of dystonia in paediatric clinical trials using a gradual up-titration was similar to that seen in adult trials.

**Weight increase**
In the combined short-term and long-term schizophrenia and bipolar mania trials in adults, the mean change in body weight for asenapine was 0.8 kg. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at endpoint) in the short-term schizophrenia trials was 5.3 % for asenapine compared to 2.3 % for placebo. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at endpoint) in the short-term, flexible-dose bipolar mania trials was 6.5 % for asenapine compared to 0.6 % for placebo.
In a 3-week, placebo-controlled, randomized, fixed-dose efficacy and safety trial in paediatric patients 10 to 17 years of age with bipolar I disorder, the mean change from baseline to endpoint in weight for placebo and asenapine 2.5 mg, 5 mg, and 10 mg twice daily, was 0.48, 1.72, 1.62, and 1.44 kg, respectively. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at Day 21) was 14.1 % for asenapine 2.5 mg twice daily, 8.9 % for asenapine 5 mg twice daily, and 9.2 % for asenapine 10 mg twice daily, compared to 1.1 % for placebo. In the long-term extension trial (50 weeks), a total of 34.8 % of subjects experienced clinically significant weight increase (i.e., ≥ 7 % increase in body weight at endpoint). Overall mean (SD) weight gain at study endpoint was 3.5 (5.76) kg.

**Orthostatic hypotension**
The incidence of orthostatic hypotension in elderly subjects was 4.1 % compared to 0.3 % in the combined phase 2/3 trial population.

**Falls**
Falls may occur as a result of one or more adverse events such as the following: Somnolence, Orthostatic hypotension, Dizziness, Extrapyramidal symptoms.

**Hepatic enzymes**
Transient, asymptomatic elevations of hepatic transaminases, alanine transferase (ALT), aspartate transferase (AST) have been seen commonly, especially in early treatment.

**Other findings**
Cerebrovascular events have been reported in patients treated with asenapine but there is no evidence of any excess incidence over what is expected in adults between 18 and 65 years of age.
Asenapine has anaesthetic properties. Oral hypoaesthesia and oral paraesthesia may occur directly after administration and usually resolves within 1 hour.

There have been post-marketing reports of serious hypersensitivity reactions in patients treated with asenapine, including anaphylactic/anaphylactoid reactions, angioedema, swollen tongue and swollen throat (pharyngeal oedema).

**Paediatric population**

Asenapine is not indicated for the treatment of children and adolescent patients below 18 years (see section 4.2). The clinically relevant adverse experiences identified in the paediatric bipolar and schizophrenia trials were similar to those observed in adult bipolar and schizophrenia trials.

The most common adverse reactions (≥ 5 % and at least twice the rate of placebo) reported in paediatric patients with bipolar I disorder were somnolence, sedation, dizziness, dysgeusia, hypoaesthesia oral, paraesthesia oral, nausea, increased appetite, fatigue, and weight increased (see Weight increase above).

The most common adverse reactions (proportion of patients ≥ 5 % and at least twice placebo) reported in paediatric patients with schizophrenia were somnolence, sedation, akathisia, dizziness, and hypoaesthesia oral. There was a statistically significant higher incidence of patients with ≥ 7 % weight gain (from baseline to endpoint) compared to placebo (3.1 %) for Sycrest 2.5 mg twice daily (9.5 %) and Sycrest 5 mg twice daily (13.1 %).

**Reporting of suspected adverse reactions**

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

**4.9 Overdose**

Few cases of overdose were reported in the asenapine program. Reported estimated doses were between 15 and 400 mg. In most cases it was not clear if asenapine had been taken sublingually. Treatment-related adverse reactions included agitation and confusion, akathisia, orofacial dystonia, sedation, and asymptomatic ECG findings (bradycardia, supraventricular complexes, intraventricular conduction delay).

No specific information is available on the treatment of overdose with Sycrest. There is no specific antidote to Sycrest. The possibility of multiple medicinal product involvement should be considered. Cardiovascular monitoring is necessary to detect possible arrhythmias and management of overdose should concentrate on supportive therapy, maintaining an adequate airway oxygenation and ventilation, and management of symptoms. Hypotension and circulatory collapse should be treated with appropriate measures, such as intravenous fluids and/or sympathomimetic agents (epinephrine and dopamine should not be used, since beta stimulations may worsen hypotension in the setting of Sycrest-induced alpha blockade). In case of severe extrapyramidal symptoms, anticholinergic medicines should be administered. Close medical supervision and monitoring should continue until the patient recovers.

**5. PHARMACOLOGICAL PROPERTIES**

**5.1 Pharmacodynamic properties**

Pharmacotherapeutic group: Psycholeptics, antipsychotics, ATC code: N05AH05
Mechanism of action
The mechanism of action of asenapine is not fully understood. However, based on its receptor pharmacology, it is proposed that the efficacy of asenapine is mediated through a combination of antagonist activity at D2 and 5-HT2A receptors. Actions at other receptors e.g., 5-HT1A, 5-HT1B, 5-HT2C, 5-HT6, 5-HT7, D3, and α2-adrenergic receptors, may also contribute to the clinical effects of asenapine.

Pharmacodynamic effects
Asenapine exhibits high affinity for serotonin 5-HT1A, 5-HT1B, 5-HT2A, 5-HT2B, 5-HT2C, 5-HT5, 5-HT6, and 5-HT7 receptors, dopamine D2, D3, D4, and D1 receptors, α1 and α2-adrenergic receptors, and histamine H1 receptors, and moderate affinity for H2 receptors. In in vitro assays asenapine acts as an antagonist at these receptors. Asenapine has no appreciable affinity for muscarinic cholinergic receptors.

Clinical efficacy

Clinical efficacy in bipolar I disorder
The efficacy of asenapine in the treatment of a DSM-IV manic or mixed episode of bipolar I disorder with or without psychotic features was evaluated in two similarly designed 3-week, randomized, double-blind, flexible-dose, placebo- and active controlled (olanzapine) monotherapy trials involving 488 and 489 patients, respectively. All patients met the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) diagnostic criteria for bipolar I disorder, current episode manic (DSM-IV 296.4x), or mixed (DSM-IV 296.6x) and had a Young Mania Rating Scale (Y-MRS) score of ≥ 20 at screening and baseline. Patients with rapid cycling were excluded from these studies. Asenapine demonstrated superior efficacy to placebo in the reduction of manic symptoms over 3 weeks. Point estimates [95 % CI] for the change from baseline to endpoint in YMRS using LOCF analysis in the two studies were as follows:

-11.5 [-13.0, -10.0] for asenapine vs -7.8 [-10.0, -5.6] for placebo and

A statistically significant difference between asenapine and placebo was seen as early as day 2.

Patients from the two pivotal 3 week trials were studied for a further 9 weeks an extension trial. Maintenance of effect during the episode after 12 weeks of randomised treatment was demonstrated in this trial.

In one double-blind, fixed-dose, parallel-group, 3-week placebo controlled trial in subjects with bipolar I disorder experiencing an acute manic or mixed episode involving 367 patients of which 126 received placebo, 122 received asenapine 5 mg twice daily (BID), and 119 received asenapine 10 mg BID, the primary efficacy hypothesis was met. Both asenapine doses (5 mg BID and 10 mg BID) were superior to placebo and showed statistically significant improvement in change from baseline in Y-MRS total score at Day 21 compared with placebo. Based upon a LOCF analysis including all patients treated, the difference in least squares (LS) mean change from baseline to Day 21 in the Y-MRS total score between asenapine 5 mg BID and placebo was -3.1 points (95 % CI [-5.7, -0.5]; p-value = 0.0183). The difference in LS mean change from baseline to Day 21 in the Y-MRS total score between asenapine 10 mg BID and placebo was -3.0 points (95 % CI [-5.6, -0.4]; p-value = 0.0244). A statistically significant difference between asenapine and placebo was seen as early as day 2. In this short-term, fixed-dose controlled trial there was no evidence of added benefit with a 10 mg twice daily dose compared to 5 mg twice daily.

In a 12-week, placebo-controlled trial involving 326 patients with a manic or mixed episode of bipolar I disorder, with or without psychotic features, who were partially non-responsive to lithium or valproate monotherapy for 2 weeks at therapeutic serum levels, the addition of asenapine as adjunctive therapy resulted in superior efficacy to lithium or valproate monotherapy at week 3 (point estimates [95 % CI] for the change from baseline to endpoint in YMRS using LOCF analysis were -10.3 [-11.9, -8.8] for asenapine and -7.9 [-9.4, -6.4] for placebo) and at week 12 (-12.7 [-14.5, -10.9] for asenapine and -9.3 [-11.8, -7.6] for placebo) in the reduction of manic symptoms.
Paediatric population
Asenapine is not indicated for the treatment of children and adolescent patients below 18 years (see section 4.2).

The safety and efficacy of Sycrest was evaluated in 403 paediatric patients with bipolar I disorder who participated in a single, 3-week, placebo-controlled, double-blind trial, of whom 302 patients received Sycrest at fixed doses ranging from 2.5 mg to 10 mg twice daily. Study results showed statistically significant superiority for all three Sycrest doses in improving the Young Mania Rating Scale (YMRS) total score as measured by the change from baseline to Day 21, as compared with placebo. Long term efficacy could not be established in a 50-week, uncontrolled, open-label extension trial. The clinically relevant adverse reactions identified in the paediatric trials were generally similar to those observed in the adult trials. However, adverse effects of treatment on weight gain and on plasma lipid profile appeared to be greater than effects observed in the adult trials.

Efficacy of Sycrest was not demonstrated in an 8-week, placebo-controlled, double-blind, randomized, fixed-dose trial in 306 adolescent patients aged 12-17 years with schizophrenia at doses of 2.5 and 5 mg twice daily.

Paediatric studies with Sycrest were performed using flavoured sublingual tablets. The European Medicines Agency has deferred the obligation to submit the results of studies with Sycrest in one or more subsets of the paediatric population in bipolar I disorder (see section 4.2 for information on paediatric use).

5.2 Pharmacokinetic properties

Absorption
Following sublingual administration, asenapine is rapidly absorbed with peak plasma concentrations occurring within 0.5 to 1.5 hours. The absolute bioavailability of sublingual asenapine at 5 mg is 35 %. The absolute bioavailability of asenapine when swallowed is low (< 2 % with an oral tablet formulation). The intake of water several (2 or 5) minutes after asenapine administration resulted in decreased (19 % and 10 %, respectively) asenapine exposure. Therefore, eating and drinking should be avoided for 10 minutes after administration (see section 4.2).

Distribution
Asenapine is rapidly distributed and has a large volume of distribution (approximately 20-25 L/kg), indicating extensive extravascular distribution. Asenapine is highly bound (95 %) to plasma proteins, including albumin and α1-acid glycoprotein.

Biotransformation
Asenapine is extensively metabolized. Direct glucuronidation (mediated by UGT1A4) and cytochrome P450 (primarily CYP1A2, with contributions of 2D6 and 3A4) mediated oxidation and demethylation are the primary metabolic pathways for asenapine. In an in vivo study in humans with radio-labelled asenapine, the predominant drug-related entity in plasma was asenapine N’-glucuronide; others included N-desmethasenapine, N-desmethasenapine N-carbamoyl glucuronide, and unchanged asenapine in smaller amounts. Sycrest activity is primarily due to the parent compound.

Asenapine is a weak inhibitor of CYP2D6. Asenapine does not cause induction of CYP1A2 or CYP3A4 activities in cultured human hepatocytes. Co-administration of asenapine with known inhibitors, inducers or substrates of these metabolic pathways has been studied in a number of drug-drug interaction studies (see section 4.5).

Elimination
Asenapine is a high clearance compound, with a clearance after intravenous administration of 52 L/h. In a mass balance study, the majority of the radioactive dose was recovered in urine (about 50 %) and faeces (about 40 %), with only a small amount excreted in faeces (5-16 %) as unchanged compound.
Following an initial more rapid distribution phase, the terminal half-life of asenapine is approximately 24 h.

**Linearity/non-linearity**
Increasing the dose from 5 to 10 mg twice daily (a two-fold increase) results in less than linear (1.7 times) increases in both the extent of exposure and maximum concentration. The less than proportional increase of Cmax and AUC with dose may be attributed to limitations in the absorption capacity from the oral mucosa following sublingual administration. During twice-daily dosing, steady-state is attained within 3 days. Overall, steady-state asenapine pharmacokinetics are similar to single-dose pharmacokinetics.

**Pharmacokinetics in special populations**

**Hepatic impairment**
The pharmacokinetics of asenapine were similar among subjects with mild (Child-Pugh A) or moderate (Child-Pugh B) hepatic impairment and subjects with normal hepatic function. In subjects with severe hepatic impairment (Child-Pugh C), a 7-fold increase in asenapine exposure was observed (see section 4.2).

**Renal impairment**
The pharmacokinetics of asenapine following a single dose of 5 mg asenapine were similar among subjects with varying degrees of renal impairment and subjects with normal renal function. There is no experience with asenapine in severe renal impairment patients with a creatinine clearance less than 15 mL/min.

**Elderly**
In elderly patients (between 65 and 85 years of age), exposure to asenapine is approximately 30 % higher than in younger adults.

**Paediatric population (children and adolescents)**
In a PK study using unflavoured sublingual tablets, at the 5 mg twice daily dose level, asenapine pharmacokinetics in adolescent patients (12 to 17 years of age, inclusive) are similar to those observed in adults. In adolescents, the 10 mg twice daily dose did not result in increased exposure compared to 5 mg twice daily.
In a second PK study using flavoured sublingual tablets, the 10 mg twice daily dose in a paediatric population (10 to 17 years of age, inclusive) resulted in an approximate dose-proportional increase in asenapine exposure compared to 5 mg twice daily.

**Gender**
A population pharmacokinetic analysis indicated that there is no evidence of gender-related differences in the pharmacokinetics of asenapine.

**Race**
In a population pharmacokinetic analysis, no clinical relevant effects of race on the pharmacokinetics of asenapine were found.

**Smoking status**
A population pharmacokinetic analysis indicated that smoking, which induces CYP1A2, has no effect on the clearance of asenapine. In a dedicated study, concomitant smoking during administration of a single 5 mg sublingual dose had no effect on the pharmacokinetics of asenapine.

### 5.3 Preclinical safety data

Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology. Repeat-dose toxicity studies in rat and dog showed mainly dose-limiting pharmacological effects, such as sedation. Furthermore, prolactin-mediated effects on mammary
glands and oestrus cycle disturbances were observed. In dogs high oral doses resulted in hepatotoxicity that was not observed after chronic intravenous administration. Asenapine has some affinity to melanin-containing tissues. However, when tested in vitro it was devoid of phototoxicity. In addition, histopathological examination of the eyes from dogs treated chronically with asenapine did not reveal any signs of ocular toxicity, demonstrating the absence of a phototoxic hazard. Asenapine was not genotoxic in a battery of tests. In subcutaneous carcinogenicity studies in rats and mice, no increases in tumour incidences were observed. Effects in non-clinical studies were observed only at exposures considered sufficiently in excess of the maximum human exposure indicating little relevance to clinical use. Asenapine did not impair fertility in rats and was not teratogenic in rat and rabbit. Embryotoxicity was found in reproduction toxicology studies using rats and rabbits. Asenapine caused mild maternal toxicity and slight retardation of foetal skeletal development. Following oral administration to pregnant rabbits during the period of organogenesis, asenapine adversely affected body weight at the high dose of 15 mg kg⁻¹ twice daily. At this dose foetal body weight decreased. When asenapine was administered intravenously to pregnant rabbits, no signs of embryotoxicity were observed. In rats, embryofetal toxicity (increased post-implantation loss, decreased foetal weights, and delayed ossification) was observed following oral or intravenous administration during organogenesis or throughout gestation. Increased neonatal mortality was observed among the offspring of female rats treated during gestation and lactation. From a cross-fostering study it was concluded that asenapine induced peri- and postnatal losses are caused by impairment of the pups rather than altered nursing behaviour of the dams.

6. **PHARMACEUTICAL PARTICULARS**

6.1 **List of excipients**

Gelatin  
Mannitol (E421)

6.2 **Incompatibilities**

Not applicable.

6.3 **Shelf life**

3 years

6.4 **Special precautions for storage**

Store in the original package in order to protect from light and moisture. This medicinal product does not require any special temperature storage conditions.

6.5 **Nature and contents of container**

Peelable aluminium/aluminium blisters in cartons of 20, 60 or 100 sublingual tablets per carton. Not all pack sizes may be marketed.

6.6 **Special precautions for disposal**

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.
7. MARKETING AUTHORISATION HOLDER

N.V. Organon, Kloosterstraat 6, NL-5349 AB Oss, The Netherlands

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/640/001
EU/1/10/640/002
EU/1/10/640/003

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORISATION

Date of first authorisation: 01 September 2010
Date of latest renewal: 05 May 2015

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
1. NAME OF THE MEDICINAL PRODUCT

Sycrest 10 mg sublingual tablets

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each sublingual tablet contains 10 mg asenapine (as maleate).

For the full list of excipients, see section 6.1.

3. PHARMACEUTICAL FORM

Sublingual tablet
Round, white to off-white, sublingual tablets debossed with "10" on one side.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Sycrest is indicated for the treatment of moderate to severe manic episodes associated with bipolar I disorder in adults.

4.2 Posology and method of administration

Posology

The recommended starting dose of Sycrest as monotherapy is 5 mg twice daily. One dose should be taken in the morning and one dose should be taken in the evening. The dose can be increased to 10 mg twice daily based on individual clinical response and tolerability. See section 5.1. For combination therapy a starting dose of 5 mg twice daily is recommended. Depending on the clinical response and tolerability in the individual patient, the dose can be increased to 10 mg twice daily.

Special populations

Elderly
Sycrest should be used with care in the elderly. Limited data on efficacy in patients 65 years of age and older are available. Available pharmacokinetic data are described in section 5.2.

Renal impairment
No dose adjustment is required for patients with renal impairment. There is no experience with asenapine in patients with severe renal impairment who have a creatinine clearance less than 15 mL/min.

Hepatic impairment
No dose adjustment is required for patients with mild hepatic impairment. The possibility of elevated asenapine plasma levels cannot be excluded in some patients with moderate hepatic impairment (Child-Pugh B) and caution is advised. In subjects with severe hepatic impairment (Child-Pugh C), a 7-fold increase in asenapine exposure was observed. Thus, Sycrest is not recommended in patients with severe hepatic impairment.

Paediatric population
A pharmacokinetic study and a short term efficacy and safety study were performed in a paediatric population (ages 10-17 years) with manic or mixed episodes associated with bipolar I disorder. Long
term safety in this population was explored in a 50-week, open-label, uncontrolled extension study. Currently available data are described in sections 4.8, 5.1 and 5.2 but no recommendation on a posology can be made.

**Method of administration**

The tablet should not be removed from the blister until ready to take it. Dry hands should be used when touching the tablet. The tablet should not be pushed through the tablet pack. The tablet pack should not be cut or torn. The coloured tab should be peeled back and the tablet should be removed gently. The tablet should not be crushed.

To ensure optimal absorption, the Sycrest sublingual tablet should be placed under the tongue and allowed to dissolve completely. The tablet will dissolve in saliva within seconds. Sycrest sublingual tablets should not be chewed or swallowed. Eating and drinking should be avoided for 10 minutes after administration.

When used in combination with other medicinal products, Sycrest should be taken last.

Treatment with Sycrest is not advised in patients who are unable to comply with this method of administration, as the bioavailability of asenapine when swallowed is low (< 2 % with an oral tablet formulation).

**4.3 Contraindications**

Hypersensitivity to the active substance or to any of the excipients listed in section 6.1.

**4.4 Special warnings and precautions for use**

**Elderly patients with dementia-related psychosis**

Elderly patients with dementia-related psychosis treated with antipsychotic substances are at an increased risk of death.

Sycrest is not approved for the treatment of patients with dementia-related psychosis and is not recommended for use in this particular group of patients.

**Neuroleptic malignant syndrome**

Neuroleptic malignant syndrome (NMS), characterised by hyperthermia, muscle rigidity, autonomic instability, altered consciousness and elevated serum creatine phosphokinase levels, has been reported to occur with antipsychotics, including asenapine. Additional clinical signs may include myoglobinuria (rhabdomyolysis) and acute renal failure.

If a patient develops signs and symptoms indicative of NMS Sycrest must be discontinued.

**Seizures**

In clinical trials, cases of seizure were occasionally reported during treatment with asenapine. Therefore, Sycrest should be used with caution in patients who have a history of seizure disorder or have conditions associated with seizures.

**Suicide**

The possibility of a suicide attempt is inherent in psychotic illnesses and bipolar disorder and close supervision of high-risk patients should accompany treatment.

**Orthostatic hypotension**

Asenapine may induce orthostatic hypotension and syncope, especially early in treatment, probably reflecting its α1-adrenergic antagonist properties. Elderly patients are particularly at risk for experiencing orthostatic hypotension (see section 4.8). In clinical trials, cases of syncope were occasionally reported during treatment with Sycrest. Sycrest should be used with caution in elderly patients and in patients with known cardiovascular disease (e.g., heart failure, myocardial infarction
or ischemia, conduction abnormalities), cerebrovascular disease, or conditions that predispose the patient to hypotension (e.g., dehydration and hypovolemia).

**Tardive dyskinesia**
Medicinal products with dopamine receptor antagonistic properties have been associated with the induction of tardive dyskinesia characterised by rhythmical, involuntary movements, predominantly of the tongue and/or face. In clinical trials, cases of tardive dyskinesia were occasionally reported during treatment with asenapine. The onset of extrapyramidal symptoms is a risk factor for tardive dyskinesia. If signs and symptoms of tardive dyskinesia appear in a patient on Sycrest, discontinuation of treatment should be considered.

**Hyperprolactinaemia**
Increases in prolactin levels were observed in some patients with Sycrest. In clinical trials, there were few adverse reactions related to abnormal prolactin levels reported.

**QT interval**
Clinically relevant QT prolongation does not appear to be associated with asenapine. Caution should be exercised when Sycrest is prescribed in patients with known cardiovascular disease or family history of QT prolongation, and in concomitant use with other medicinal products thought to prolong the QT interval.

**Hyperglycaemia and diabetes mellitus**
Hyperglycaemia or exacerbation of pre-existing diabetes has occasionally been reported during treatment with asenapine. Assessment of the relationship between atypical antipsychotic use and glucose abnormalities is complicated by the possibility of an increased background risk of diabetes mellitus in patients with schizophrenia or bipolar disorder and the increasing incidence of diabetes mellitus in the general population. Appropriate clinical monitoring is advisable in diabetic patients and in patients with risk factors for the development of diabetes mellitus.

**Dysphagia**
Esophageal dysmotility and aspiration have been associated with antipsychotic treatment. Cases of dysphagia were occasionally reported in patients treated with Sycrest.

**Body temperature regulation**
Disruption of the body’s ability to reduce core body temperature has been attributed to antipsychotic medicines. From the clinical trials, it is concluded that clinically relevant body temperature dysregulation does not appear to be associated with asenapine. Appropriate care is advised when prescribing Sycrest for patients who will be experiencing conditions that may contribute to an elevation in core body temperature, e.g. exercising strenuously, exposure to extreme heat, receiving concomitant medicinal products with anticholinergic activity or being subject to dehydration.

**Patients with severe hepatic impairment**
Asenapine exposure is increased 7-fold in patients with severe hepatic impairment (Child-Pugh C). Therefore, Sycrest is not recommended in such patients.

**Parkinson’s disease and dementia with Lewy bodies**
Physicians should weigh the risks versus the benefits when prescribing Sycrest to patients with Parkinson’s disease or dementia with Lewy Bodies (DLB) since both groups may be at increased risk of neuroleptic malignant syndrome as well as having an increased sensitivity to antipsychotics. Manifestation of this increased sensitivity can include confusion, obtundation, postural instability with frequent falls, in addition to extrapyramidal symptoms.

**Falls**
Asenapine may cause adverse effects such as somnolence, orthostatic hypotension, dizziness and extrapyramidal symptoms, which may lead to falls and, consequently, fractures or other injuries. Patients at risk for fall should be evaluated prior to prescribing asenapine.
4.5 Interaction with other medicinal products and other forms of interaction

Given the primary effects of asenapine on the central nervous system (CNS) (see section 4.8), caution should be used when it is taken in combination with other centrally acting medicinal products. Patients should be advised to avoid alcohol while taking Sycrest.

Potential for other medicines to affect Sycrest
Asenapine is cleared primarily through direct glucuronidation by UGT1A4 and oxidative metabolism by cytochrome P450 isoenzymes (predominantly CYP1A2). The potential effects of inhibitors and an inducer of several of these enzyme pathways on asenapine pharmacokinetics were studied, specifically fluvoxamine (CYP1A2 inhibitor), paroxetine (CYP2D6 inhibitor), imipramine (CYP1A2/2C19/3A4 inhibitor), cimetidine (CYP3A4/2D6/1A2 inhibitor), carbamazepine (CYP3A4/1A2 inducer) and valproate (UGT inhibitor). Except for fluvoxamine, none of the interacting medicinal products resulted in clinically relevant alterations in asenapine pharmacokinetics.

During combined administration with a single dose of asenapine 5 mg, fluvoxamine 25 mg twice daily resulted in a 29% increase in asenapine AUC. The full therapeutic dose of fluvoxamine would be expected to produce a greater increase in asenapine plasma concentrations. Therefore, co-administration of asenapine and fluvoxamine should be approached with caution.

Potential for Sycrest to affect other medicines
Because of its α1-adrenergic antagonism with potential for inducing orthostatic hypotension (see section 4.4), Sycrest may enhance the effects of certain antihypertensive agents.

Asenapine may antagonise the effect of levodopa and dopamine agonists. If this combination is deemed necessary, the lowest effective dose of each treatment should be prescribed.

In vitro studies indicate that asenapine weakly inhibits CYP2D6. Clinical drug interaction studies investigating the effects of CYP2D6 inhibition by asenapine showed the following results:

- Following co-administration of dextromethorphan and asenapine in healthy subjects, the ratio of dextrorphan/dextromethorphan (DX/DM) as a marker of CYP2D6 activity was measured. Indicative of CYP2D6 inhibition, treatment with asenapine 5 mg twice daily resulted in a fractional decrease in DX/DM ratio to 0.43. In the same study, treatment with paroxetine 20 mg daily decreased the DX/DM ratio to 0.032.

- In a separate study, co-administration of a single 75 mg dose of imipramine with a single 5 mg dose of asenapine did not affect the plasma concentrations of the metabolite desipramine (a CYP2D6 substrate).

- Co-administration of a single 20 mg dose of paroxetine (a CYP2D6 substrate and inhibitor) during treatment with 5 mg asenapine twice daily in 15 healthy male subjects resulted in an almost 2-fold increase in paroxetine exposure.

In vivo asenapine appears to be at most a weak inhibitor of CYP2D6. However, asenapine may enhance the inhibitory effects of paroxetine on its own metabolism. Therefore, Sycrest should be co-administered cautiously with medicinal products that are both substrates and inhibitors for CYP2D6.

4.6 Fertility, pregnancy and lactation

Pregnancy
There are no adequate data from the use of Sycrest in pregnant women. Asenapine was not teratogenic in animal studies. Maternal and embryo toxic effects were found in animal studies (see section 5.3).
Newborn infants exposed to antipsychotics (including Sycrest) during the third trimester of pregnancy are at risk of adverse reactions including extrapyramidal and/or withdrawal symptoms that may vary in severity and duration following delivery. There have been reports of agitation, hypertonia, hypotonia, tremor, somnolence, respiratory distress, or feeding disorder in newborn infants. Consequently, newborn infants should be monitored carefully.

Sycrest should not be used during pregnancy unless the clinical condition of the woman requires treatment with asenapine and only if the potential benefit outweighs the potential risk to the foetus.

**Breast-feeding**
Asenapine was excreted in milk of rats during lactation. It is not known whether asenapine or its metabolites are excreted in human milk. Breast-feeding should be discontinued during treatment with Sycrest.

**Fertility**
No impairment of fertility has been observed in nonclinical studies (see section 5.3).

### 4.7 Effects on ability to drive and use machines

Asenapine may cause somnolence and sedation. Therefore, patients should be cautioned about driving and using machines until they are reasonably certain that Sycrest therapy does not affect them adversely.

### 4.8 Undesirable effects

**Summary of safety profile**
The most frequently reported adverse drug reactions (ADRs) associated with the use of asenapine in clinical trials were somnolence and anxiety. Serious hypersensitivity reactions have been reported. Other serious ADRs are discussed in more detail in section 4.4.

**Tabulated list of adverse reactions**
The incidences of the ADRs associated with asenapine therapy are tabulated below. The table is based on adverse reactions reported during clinical trials and/or post-marketing use. All ADRs are listed by system organ class and frequency; very common (≥ 1/10), common (≥ 1/100 to < 1/10), uncommon (≥ 1/1,000 to < 1/100), rare (≥ 1/10,000 to < 1/1,000) and not known (cannot be estimated from the available data). Within each frequency grouping, ADRs are presented in order of decreasing seriousness.

<table>
<thead>
<tr>
<th>System organ class</th>
<th>Very common</th>
<th>Common</th>
<th>Uncommon</th>
<th>Rare</th>
<th>Not known</th>
</tr>
</thead>
<tbody>
<tr>
<td>Blood and lymphatic disorders</td>
<td></td>
<td></td>
<td></td>
<td>Neutropenia</td>
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<tr>
<td>Immune system disorders</td>
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<td></td>
<td>Allergic reactions</td>
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<tr>
<td>Metabolism and nutrition disorders</td>
<td>Weight increased</td>
<td>Increased appetite</td>
<td>Hyperglycaemia</td>
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<tr>
<td>Psychiatric disorders</td>
<td>Anxiety</td>
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<tr>
<td>System organ class</td>
<td>Very common</td>
<td>Common</td>
<td>Uncommon</td>
<td>Rare</td>
<td>Not known</td>
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<tr>
<td>Nervous system disorders</td>
<td>Somnolence</td>
<td>Dystonia Akathisia Dyskinesia Parkinsonism</td>
<td>Syncope Seizure Extrapiramidal disorder</td>
<td>Dysarthria Restless legs syndrome</td>
<td>Neuroleptic malignant syndrome</td>
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<td></td>
<td></td>
<td>Sedation Dizziness</td>
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<td></td>
<td></td>
<td>Dizziness Dysgeusia</td>
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<tr>
<td>Eye disorders</td>
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<td>Accommodation disorder</td>
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<tr>
<td>Cardiac disorders</td>
<td></td>
<td></td>
<td>Sinus bradycardia Bundle branch block</td>
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<td>Electrocardiogram QT prolonged Sinus</td>
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<td>tachycardia</td>
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<tr>
<td>Vascular disorders</td>
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<td></td>
<td>Orthostatic hypotension</td>
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<td></td>
<td></td>
<td></td>
<td>Hypotension</td>
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<tr>
<td>Respiratory, thoracic and mediastinal</td>
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<td></td>
<td>Pulmonary embolism</td>
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<td>disorders</td>
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<tr>
<td>Gastrointestinal disorders</td>
<td></td>
<td>Hypoaesthesia oral Nausea Salivary hypersecretion</td>
<td>Swollen tongue Dysphagia Glossodynia Paraesthesia oral Oral mucosal lesions (ulcerations, blistering and inflammation)</td>
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<tr>
<td>Hepatobiliary disorders</td>
<td></td>
<td>Alanine aminotransferases increased</td>
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<tr>
<td>Injury, poisoning and procedural</td>
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<td></td>
<td></td>
<td>Falls*</td>
<td></td>
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<tr>
<td>complications</td>
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<tr>
<td>Musculoskeletal and connective tissue</td>
<td></td>
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<td>Muscle rigidity</td>
<td>Rhabdomyolysis</td>
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<td>disorders</td>
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<tr>
<td>Pregnancy, puerperium and perinatal</td>
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<td></td>
<td>Drug withdrawal syndrome neonatal (see 4.6)</td>
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<tr>
<td>conditions</td>
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</table>
System organ class | Very common | Common | Uncommon | Rare | Not known |
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<tbody>
<tr>
<td>Reproductive system and breast disorders</td>
<td>Sexual dysfunction</td>
<td>Amenorrhoea</td>
<td>Gynaecomastia</td>
<td>Galactorrhoea</td>
<td></td>
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<tr>
<td>General disorders and administration site conditions</td>
<td>Fatigue</td>
<td></td>
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</tbody>
</table>

* See subsection “Falls” below

**Description of selected adverse reactions**

* **Extrapyramidal Symptoms (EPS)**

In clinical trials, the incidence of extrapyramidal symptoms in asenapine-treated patients was higher than placebo (15.4 % vs 11.0 %).

From the short-term (6 weeks) schizophrenia trials there appears to be a dose-response relationship for akathisia in patients treated with asenapine, and for parkinsonism there was an increasing trend with higher doses.

Based on a small pharmacokinetic study, paediatric patients appeared to be more sensitive to dystonia with initial dosing with asenapine when a gradual up-titration schedule was not followed (see section 5.2). The incidence of dystonia in paediatric clinical trials using a gradual up-titration was similar to that seen in adult trials.

* **Weight increase**

In the combined short-term and long-term schizophrenia and bipolar mania trials in adults, the mean change in body weight for asenapine was 0.8 kg. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at endpoint) in the short-term schizophrenia trials was 5.3 % for asenapine compared to 2.3 % for placebo. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at endpoint) in the short-term, flexible-dose bipolar mania trials was 6.5 % for asenapine compared to 0.6 % for placebo.

In a 3-week, placebo-controlled, randomized, fixed-dose efficacy and safety trial in paediatric patients 10 to 17 years of age with bipolar I disorder, the mean change from baseline to endpoint in weight for placebo and asenapine 2.5 mg, 5 mg, and 10 mg twice daily, was 0.48, 1.72, 1.62, and 1.44 kg, respectively. The proportion of subjects with clinically significant weight gain (≥ 7 % weight gain from baseline at Day 21) was 14.1 % for asenapine 2.5 mg twice daily, 8.9 % for asenapine 5 mg twice daily, and 9.2 % for asenapine 10 mg twice daily, compared to 1.1 % for placebo. In the long-term extension trial (50 weeks), a total of 34.8 % of subjects experienced clinically significant weight increase (i.e., ≥ 7 % increase in body weight at endpoint). Overall mean (SD) weight gain at study endpoint was 3.5 (5.76) kg.

* **Orthostatic hypotension**

The incidence of orthostatic hypotension in elderly subjects was 4.1 % compared to 0.3 % in the combined phase 2/3 trial population.

* **Falls**

Falls may occur as a result of one or more adverse events such as the following: Somnolence, Orthostatic hypotension, Dizziness, Extrapyramidal symptoms.

* **Hepatic enzymes**

Transient, asymptomatic elevations of hepatic transaminases, alanine transferase (ALT), aspartate transferase (AST) have been seen commonly, especially in early treatment.

* **Other findings**

Cerebrovascular events have been reported in patients treated with asenapine but there is no evidence of any excess incidence over what is expected in adults between 18 and 65 years of age.
Asenapine has anaesthetic properties. Oral hypoaesthesia and oral paraesthesia may occur directly after administration and usually resolves within 1 hour.

There have been post-marketing reports of serious hypersensitivity reactions in patients treated with asenapine, including anaphylactic/anaphylactoid reactions, angioedema, swollen tongue and swollen throat (pharyngeal oedema).

**Paediatric population**

Asenapine is not indicated for the treatment of children and adolescent patients below 18 years (see section 4.2). The clinically relevant adverse experiences identified in the paediatric bipolar and schizophrenia trials were similar to those observed in adult bipolar and schizophrenia trials.

The most common adverse reactions (≥ 5 % and at least twice the rate of placebo) reported in paediatric patients with bipolar I disorder were somnolence, sedation, dizziness, dysgeusia, hypoaesthesia oral, paraesthesia oral, nausea, increased appetite, fatigue, and weight increased (see Weight increase above).

The most common adverse reactions (proportion of patients ≥ 5 % and at least twice placebo) reported in paediatric patients with schizophrenia were somnolence, sedation, akathisia, dizziness, and hypoaesthesia oral. There was a statistically significant higher incidence of patients with ≥ 7 % weight gain (from baseline to endpoint) compared to placebo (3.1 %) for Sycrest 2.5 mg twice daily (9.5 %) and Sycrest 5 mg twice daily (13.1 %).

**Reporting of suspected adverse reactions**

Reporting suspected adverse reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse reactions via the national reporting system listed in Appendix V.

**4.9 Overdose**

Few cases of overdose were reported in the asenapine program. Reported estimated doses were between 15 and 400 mg. In most cases it was not clear if asenapine had been taken sublingually. Treatment-related adverse reactions included agitation and confusion, akathisia, orofacial dystonia, sedation, and asymptomatic ECG findings (bradycardia, supraventricular complexes, intraventricular conduction delay).

No specific information is available on the treatment of overdose with Sycrest. There is no specific antidote to Sycrest. The possibility of multiple medicinal product involvement should be considered.

Cardiovascular monitoring is necessary to detect possible arrhythmias and management of overdose should concentrate on supportive therapy, maintaining an adequate airway oxygenation and ventilation, and management of symptoms. Hypotension and circulatory collapse should be treated with appropriate measures, such as intravenous fluids and/or sympathomimetic agents (epinephrine and dopamine should not be used, since beta stimulations may worsen hypotension in the setting of Sycrest-induced alpha blockade). In case of severe extrapyramidal symptoms, anticholinergic medicines should be administered. Close medical supervision and monitoring should continue until the patient recovers.

**5. PHARMACOLOGICAL PROPERTIES**

**5.1 Pharmacodynamic properties**

Pharmacotherapeutic group: Psycholeptics, antipsychotics, ATC code: N05AH05
Mechanism of action
The mechanism of action of asenapine is not fully understood. However, based on its receptor pharmacology, it is proposed that the efficacy of asenapine is mediated through a combination of antagonist activity at D2 and 5-HT2A receptors. Actions at other receptors e.g., 5-HT1A, 5-HT1B, 5-HT2C, 5-HT6, 5-HT7, D3, and α2-adrenergic receptors, may also contribute to the clinical effects of asenapine.

Pharmacodynamic effects
Asenapine exhibits high affinity for serotonin 5-HT1A, 5-HT1B, 5-HT2A, 5-HT2B, 5-HT2C, 5-HT5, 5-HT6, and 5-HT7 receptors, dopamine D2, D3, D4, and D1 receptors), α1 and α2-adrenergic receptors, and histamine H1 receptors, and moderate affinity for H2 receptors. In in vitro assays asenapine acts as an antagonist at these receptors. Asenapine has no appreciable affinity for muscarinic cholinergic receptors.

Clinical efficacy

Clinical efficacy in bipolar I disorder
The efficacy of asenapine in the treatment of a DSM-IV manic or mixed episode of bipolar I disorder with or without psychotic features was evaluated in two similarly designed 3-week, randomized, double-blind, flexible-dose, placebo- and active controlled (olanzapine) monotherapy trials involving 488 and 489 patients, respectively. All patients met the Diagnostic and Statistical Manual of Mental Disorders, 4th Edition (DSM-IV) diagnostic criteria for bipolar I disorder, current episode manic (DSM-IV 296.4x), or mixed (DSM-IV 296.6x) and had a Young Mania Rating Scale (Y-MRS) score of ≥ 20 at screening and baseline. Patients with rapid cycling were excluded from these studies. Asenapine demonstrated superior efficacy to placebo in the reduction of manic symptoms over 3 weeks. Point estimates [95 % CI] for the change from baseline to endpoint in YMRS using LOCF analysis in the two studies were as follows:
-11.5 [-13.0, -10.0] for asenapine vs -7.8 [-10.0, -5.6] for placebo and
A statistically significant difference between asenapine and placebo was seen as early as day 2.

Patients from the two pivotal 3 week trials were studied for a further 9 weeks an extension trial. Maintenance of effect during the episode after 12 weeks of randomised treatment was demonstrated in this trial.

In one double-blind, fixed-dose, parallel-group, 3-week placebo controlled trial in subjects with bipolar I disorder experiencing an acute manic or mixed episode involving 367 patients of which 126 received placebo, 122 received asenapine 5 mg twice daily (BID), and 119 received asenapine 10 mg BID, the primary efficacy hypothesis was met. Both asenapine doses (5 mg BID and 10 mg BID) were superior to placebo and showed statistically significant improvement in change from baseline in Y-MRS total score at Day 21 compared with placebo. Based upon a LOCF analysis including all patients treated, the difference in least squares (LS) mean change from baseline to Day 21 in the Y-MRS total score between asenapine 5 mg BID and placebo was -3.1 points (95 % CI [-5.7, -0.5]; p-value = 0.0183). The difference in LS mean change from baseline to Day 21 in the Y-MRS total score between asenapine 10 mg BID and placebo was -3.0 points (95 % CI [-5.6, -0.4]; p-value = 0.0244). A statistically significant difference between asenapine and placebo was seen as early as day 2. In this short-term, fixed-dose controlled trial there was no evidence of added benefit with a 10 mg twice daily dose compared to 5 mg twice daily.

In a 12-week, placebo-controlled trial involving 326 patients with a manic or mixed episode of bipolar I disorder, with or without psychotic features, who were partially non-responsive to lithium or valproate monotherapy for 2 weeks at therapeutic serum levels, the addition of asenapine as adjunctive therapy resulted in superior efficacy to lithium or valproate monotherapy at week 3 (point estimates [95 % CI] for the change from baseline to endpoint in YMRS using LOCF analysis were -10.3 [-11.9, -8.8] for asenapine and -7.9 [-9.4, -6.4] for placebo) and at week 12 (-12.7 [-14.5, -10.9] for asenapine and -9.3 [-11.8, -7.6] for placebo) in the reduction of manic symptoms.
**Paediatric population**
Asenapine is not indicated for the treatment of children and adolescent patients below 18 years (see section 4.2).

The safety and efficacy of Sycrest was evaluated in 403 paediatric patients with bipolar I disorder who participated in a single, 3-week, placebo-controlled, double-blind trial, of whom 302 patients received Sycrest at fixed doses ranging from 2.5 mg to 10 mg twice daily. Study results showed statistically significant superiority for all three Sycrest doses in improving the Young Mania Rating Scale (YMRS) total score as measured by the change from baseline to Day 21, as compared with placebo. Long term efficacy could not be established in a 50-week, uncontrolled, open-label extension trial. The clinically relevant adverse reactions identified in the paediatric trials were generally similar to those observed in the adult trials. However, adverse effects of treatment on weight gain and on plasma lipid profile appeared to be greater than effects observed in the adult trials.

Efficacy of Sycrest was not demonstrated in an 8-week, placebo-controlled, double-blind, randomized, fixed-dose trial in 306 adolescent patients aged 12-17 years with schizophrenia at doses of 2.5 and 5 mg twice daily.

Paediatric studies with Sycrest were performed using flavoured sublingual tablets. The European Medicines Agency has deferred the obligation to submit the results of studies with Sycrest in one or more subsets of the paediatric population in bipolar I disorder (see section 4.2 for information on paediatric use).

5.2 **Pharmacokinetic properties**

**Absorption**
Following sublingual administration, asenapine is rapidly absorbed with peak plasma concentrations occurring within 0.5 to 1.5 hours. The absolute bioavailability of sublingual asenapine at 5 mg is 35 %. The absolute bioavailability of asenapine when swallowed is low (< 2 % with an oral tablet formulation). The intake of water several (2 or 5) minutes after asenapine administration resulted in decreased (19 % and 10 %, respectively) asenapine exposure. Therefore, eating and drinking should be avoided for 10 minutes after administration (see section 4.2).

**Distribution**
Asenapine is rapidly distributed and has a large volume of distribution (approximately 20-25 L/kg), indicating extensive extravascular distribution. Asenapine is highly bound (95 %) to plasma proteins, including albumin and α1-acid glycoprotein.

**Biotransformation**
Asenapine is extensively metabolized. Direct glucuronidation (mediated by UGT1A4) and cytochrome P450 (primarily CYP1A2, with contributions of 2D6 and 3A4) mediated oxidation and demethylation are the primary metabolic pathways for asenapine. In an *in vivo* study in humans with radio-labelled asenapine, the predominant drug-related entity in plasma was asenapine N’-glucuronide; others included N-desmethylasenapine, N-desmethylasenapine N-carbamoyl glucuronide, and unchanged asenapine in smaller amounts. Sycrest activity is primarily due to the parent compound.
Asenapine is a weak inhibitor of CYP2D6. Asenapine does not cause induction of CYP1A2 or CYP3A4 activities in cultured human hepatocytes. Co-administration of asenapine with known inhibitors, inducers or substrates of these metabolic pathways has been studied in a number of drug-drug interaction studies (see section 4.5).

**Elimination**
Asenapine is a high clearance compound, with a clearance after intravenous administration of 52 L/h. In a mass balance study, the majority of the radioactive dose was recovered in urine (about 50 %) and faeces (about 40 %), with only a small amount excreted in faeces (5-16 %) as unchanged compound.
Following an initial more rapid distribution phase, the terminal half-life of asenapine is approximately 24 h.

**Linearity/non-linearity**
Increasing the dose from 5 to 10 mg twice daily (a two-fold increase) results in less than linear (1.7 times) increases in both the extent of exposure and maximum concentration. The less than proportional increase of Cmax and AUC with dose may be attributed to limitations in the absorption capacity from the oral mucosa following sublingual administration. During twice-daily dosing, steady-state is attained within 3 days. Overall, steady-state asenapine pharmacokinetics are similar to single-dose pharmacokinetics.

**Pharmacokinetics in special populations**

**Hepatic impairment**
The pharmacokinetics of asenapine were similar among subjects with mild (Child-Pugh A) or moderate (Child-Pugh B) hepatic impairment and subjects with normal hepatic function. In subjects with severe hepatic impairment (Child-Pugh C), a 7-fold increase in asenapine exposure was observed (see section 4.2).

**Renal impairment**
The pharmacokinetics of asenapine following a single dose of 5 mg asenapine were similar among subjects with varying degrees of renal impairment and subjects with normal renal function. There is no experience with asenapine in severe renal impairment patients with a creatinine clearance less than 15 mL/min.

**Elderly**
In elderly patients (between 65 and 85 years of age), exposure to asenapine is approximately 30 % higher than in younger adults.

**Paediatric population (children and adolescents)**
In a PK study using unflavoured sublingual tablets, at the 5 mg twice daily dose level, asenapine pharmacokinetics in adolescent patients (12 to 17 years of age, inclusive) are similar to those observed in adults. In adolescents, the 10 mg twice daily dose did not result in increased exposure compared to 5 mg twice daily.
In a second PK study using flavoured sublingual tablets, the 10 mg twice daily dose in a paediatric population (10 to 17 years of age, inclusive) resulted in an approximate dose-proportional increase in asenapine exposure compared to 5 mg twice daily.

**Gender**
A population pharmacokinetic analysis indicated that there is no evidence of gender-related differences in the pharmacokinetics of asenapine.

**Race**
In a population pharmacokinetic analysis, no clinical relevant effects of race on the pharmacokinetics of asenapine were found.

**Smoking status**
A population pharmacokinetic analysis indicated that smoking, which induces CYP1A2, has no effect on the clearance of asenapine. In a dedicated study, concomitant smoking during administration of a single 5 mg sublingual dose had no effect on the pharmacokinetics of asenapine.

**5.3 Preclinical safety data**
Non-clinical data reveal no special hazard for humans based on conventional studies of safety pharmacology. Repeat-dose toxicity studies in rat and dog showed mainly dose-limiting pharmacological effects, such as sedation. Furthermore, prolactin-mediated effects on mammary
glands and oestrus cycle disturbances were observed. In dogs high oral doses resulted in hepatotoxicity that was not observed after chronic intravenous administration. Asenapine has some affinity to melanin-containing tissues. However, when tested in vitro it was devoid of phototoxicity. In addition, histopathological examination of the eyes from dogs treated chronically with asenapine did not reveal any signs of ocular toxicity, demonstrating the absence of a phototoxic hazard. Asenapine was not genotoxic in a battery of tests. In subcutaneous carcinogenicity studies in rats and mice, no increases in tumour incidences were observed. Effects in non-clinical studies were observed only at exposures considered sufficiently in excess of the maximum human exposure indicating little relevance to clinical use. Asenapine did not impair fertility in rats and was not teratogenic in rat and rabbit. Embryotoxicity was found in reproduction toxicology studies using rats and rabbits. Asenapine caused mild maternal toxicity and slight retardation of foetal skeletal development. Following oral administration to pregnant rabbits during the period of organogenesis, asenapine adversely affected body weight at the high dose of 15 mg.kg⁻¹ twice daily. At this dose foetal body weight decreased. When asenapine was administered intravenously to pregnant rabbits, no signs of embryotoxicity were observed. In rats, embryofoetal toxicity (increased post-implantation loss, decreased foetal weights, and delayed ossification) was observed following oral or intravenous administration during organogenesis or throughout gestation. Increased neonatal mortality was observed among the offspring of female rats treated during gestation and lactation. From a cross-fostering study it was concluded that asenapine induced peri- and postnatal losses are caused by impairment of the pups rather than altered nursing behaviour of the dams.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Gelatin
Mannitol (E421)

6.2 Incompatibilities

Not applicable.

6.3 Shelf life

3 years

6.4 Special precautions for storage

Store in the original package in order to protect from light and moisture. This medicinal product does not require any special temperature storage conditions.

6.5 Nature and contents of container

Peelable aluminium/aluminium blisters in cartons of 20, 60 or 100 sublingual tablets per carton. Not all pack sizes may be marketed.

6.6 Special precautions for disposal

Any unused medicinal product or waste material should be disposed of in accordance with local requirements.
7. MARKETING AUTHORISATION HOLDER

N.V. Organon, Kloosterstraat 6, NL-5349 AB Oss, The Netherlands

8. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/640/004
EU/1/10/640/005
EU/1/10/640/006

9. DATE OF FIRST AUTHORISATION/RENEWAL OF THE AUTHORIZATION

Date of first authorisation: 01 September 2010
Date of latest renewal: 05 May 2015

10. DATE OF REVISION OF THE TEXT

Detailed information on this medicinal product is available on the website of the European Medicines Agency http://www.ema.europa.eu.
ANNEX II

A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT
A. MANUFACTURER(S) RESPONSIBLE FOR BATCH RELEASE

Name and address of the manufacturer(s) responsible for batch release

Organon (Ireland) Ltd.
Drynam Road, Swords, Co. Dublin
Ireland

Schering-Plough Labo N.V.
Industriepark 30
B-2220 Heist-op-den-Berg, Belgium

The printed package leaflet of the medicinal product must state the name and address of the manufacturer responsible for the release of the concerned batch

B. CONDITIONS OR RESTRICTIONS REGARDING SUPPLY AND USE

Medicinal product subject to medical prescription.

C. OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORIZATION

- Periodic Safety Update Reports

The requirements for submission of periodic safety update reports for this medicinal product are set out in the list of Union reference dates (EURD list) provided for under Article 107c(7) of Directive 2001/83/EC and any subsequent updates published on the European medicines web-portal.

D. CONDITIONS OR RESTRICTIONS WITH REGARD TO THE SAFE AND EFFECTIVE USE OF THE MEDICINAL PRODUCT

- Risk Management Plan (RMP)

The MAH shall perform the required pharmacovigilance activities and interventions detailed in the agreed RMP presented in Module 1.8.2 of the Marketing Authorisation and any agreed subsequent updates of the RMP.

An updated RMP should be submitted:
- At the request of the European Medicines Agency;
- Whenever the risk management system is modified, especially as the result of new information being received that may lead to a significant change to the benefit/risk profile or as the result of an important (pharmacovigilance or risk minimisation) milestone being reached

If the dates for submission of a PSUR and the update of a RMP coincide, they can be submitted at the same time.
ANNEX III

LABELLING AND PACKAGE LEAFLET
A. LABELLING
PARTICULARS TO APPEAR ON THE OUTER PACKAGING

OUTER CARTON (5 mg)

1. NAME OF THE MEDICINAL PRODUCT

Sycrest 5 mg sublingual tablets
asenapine

2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each sublingual tablet contains 5 mg asenapine (as maleate).

3. LIST OF EXCIPIENTS

4. PHARMACEUTICAL FORM AND CONTENTS

20 sublingual tablets
60 sublingual tablets
100 sublingual tablets

5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.
Sublingual use.
Peelable blister. Do not crush, chew or swallow.
Keep the tablet under your tongue until it dissolves.
Do not eat or drink for 10 minutes after taking the tablet.

6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

7. OTHER SPECIAL WARNING(S), IF NECESSARY

8. EXPIRY DATE

EXP

9. SPECIAL STORAGE CONDITIONS

Store in the original package in order to protect from light and moisture.
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

N.V. Organon
Kloosterstraat 6
NL- 5349 AB Oss
The Netherlands

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/640/001 20 sublingual tablets
EU/1/10/640/002 60 sublingual tablets
EU/1/10/640/003 100 sublingual tablets

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

sycrest 5 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC:
SN:
NN:
<table>
<thead>
<tr>
<th>MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS</th>
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<tbody>
<tr>
<td><strong>BLISTER (5 mg)</strong></td>
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<tr>
<td><strong>1. NAME OF THE MEDICINAL PRODUCT</strong></td>
</tr>
<tr>
<td>Sycrest 5 mg sublingual tablets</td>
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<tr>
<td>asenapine</td>
</tr>
<tr>
<td><strong>2. NAME OF THE MARKETING AUTHORISATION HOLDER</strong></td>
</tr>
<tr>
<td>N.V. Organon</td>
</tr>
<tr>
<td><strong>3. EXPIRY DATE</strong></td>
</tr>
<tr>
<td>EXP</td>
</tr>
<tr>
<td><strong>4. BATCH NUMBER</strong></td>
</tr>
<tr>
<td>Lot</td>
</tr>
<tr>
<td><strong>5. OTHER</strong></td>
</tr>
</tbody>
</table>
### 1. NAME OF THE MEDICINAL PRODUCT

Sycrest 10 mg sublingual tablets  
assenapine

### 2. STATEMENT OF ACTIVE SUBSTANCE(S)

Each sublingual tablet contains 10 mg asenapine (as maleate).

### 3. LIST OF EXCIPIENTS

### 4. PHARMACEUTICAL FORM AND CONTENTS

<table>
<thead>
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<th>Number of Tablets</th>
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<tr>
<td>20 sublingual tablets</td>
</tr>
<tr>
<td>60 sublingual tablets</td>
</tr>
<tr>
<td>100 sublingual tablets</td>
</tr>
</tbody>
</table>

### 5. METHOD AND ROUTE(S) OF ADMINISTRATION

Read the package leaflet before use.  
Sublingual use.  
Peelable blister. Do not crush, chew or swallow.  
Keep the tablet under your tongue until it dissolves.  
Do not eat or drink for 10 minutes after taking the tablet.

### 6. SPECIAL WARNING THAT THE MEDICINAL PRODUCT MUST BE STORED OUT OF THE SIGHT AND REACH OF CHILDREN

Keep out of the sight and reach of children.

### 7. OTHER SPECIAL WARNING(S), IF NECESSARY

### 8. EXPIRY DATE

EXP

### 9. SPECIAL STORAGE CONDITIONS

Store in the original package in order to protect from light and moisture.
10. SPECIAL PRECAUTIONS FOR DISPOSAL OF UNUSED MEDICINAL PRODUCTS OR WASTE MATERIALS DERIVED FROM SUCH MEDICINAL PRODUCTS, IF APPROPRIATE

11. NAME AND ADDRESS OF THE MARKETING AUTHORISATION HOLDER

N.V. Organon
Kloosterstraat 6
NL- 5349 AB Oss
The Netherlands

12. MARKETING AUTHORISATION NUMBER(S)

EU/1/10/640/004 20 sublingual tablets
EU/1/10/640/005 60 sublingual tablets
EU/1/10/640/006 100 sublingual tablets

13. BATCH NUMBER

Lot

14. GENERAL CLASSIFICATION FOR SUPPLY

15. INSTRUCTIONS ON USE

16. INFORMATION IN BRAILLE

sycrest 10 mg

17. UNIQUE IDENTIFIER – 2D BARCODE

2D barcode carrying the unique identifier included.

18. UNIQUE IDENTIFIER – HUMAN READABLE DATA

PC:
SN:
NN:
## MINIMUM PARTICULARS TO APPEAR ON BLISTERS OR STRIPS

### BLISTER (10 mg)

<table>
<thead>
<tr>
<th>1. NAME OF THE MEDICINAL PRODUCT</th>
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<tbody>
<tr>
<td>Sycrest 10 mg sublingual tablets</td>
</tr>
<tr>
<td>asenapine</td>
</tr>
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</table>

<table>
<thead>
<tr>
<th>2. NAME OF THE MARKETING AUTHORISATION HOLDER</th>
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<td>N.V. Organon</td>
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<tr>
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</table>

<table>
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<tr>
<th>4. BATCH NUMBER</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lot</td>
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</table>

<table>
<thead>
<tr>
<th>5. OTHER</th>
</tr>
</thead>
</table>
B. PACKAGE LEAFLET
Package leaflet: Information for the patient

Sycrest 5 mg sublingual tablets
Sycrest 10 mg sublingual tablets
asenapine

Read all of this leaflet carefully before you start taking this medicine because it contains important information for you.
- Keep this leaflet. You may need to read it again.
- If you have any further questions, ask your doctor, pharmacist or nurse.
- This medicine has been prescribed for you only. Do not pass it on to others. It may harm them, even if their signs of illness are the same as yours.
- If you get any side effects, talk to your doctor, pharmacist or nurse. This includes any possible side effects not listed in this leaflet. See section 4.

What is in this leaflet
1. What Sycrest is and what it is used for
2. What you need to know before you take Sycrest
3. How to take Sycrest
4. Possible side effects
5. How to store Sycrest
6. Contents of the pack and other information

1. What Sycrest is and what it is used for

Sycrest contains the active substance asenapine. This medicine belongs to a group of medicines called antipsychotics. Sycrest is used to treat moderate to severe manic episodes associated with bipolar I disorder in adults. Antipsychotic medicines affect the chemicals that allow communication between nerve cells (neurotransmitters). Illnesses that affect the brain, such as bipolar I disorder, may be due to certain chemicals in the brain, such as dopamine and serotonin, being out of balance and these imbalances may cause some of the symptoms you may be experiencing. Exactly how this medicine works is unknown, however, it is believed to adjust the balance of these chemicals.

Manic episodes associated with bipolar I disorder is a condition with symptoms such as feeling “high”, having excessive amounts of energy, needing much less sleep than usual, talking very quickly with racing ideas and sometimes severe irritability.

2. What you need to know before you take Sycrest

Do not take Sycrest
If you are allergic to asenapine or any of the other ingredients of this medicine (listed in section 6).

Warnings and precautions
Talk to your doctor, pharmacist or nurse before taking Sycrest.

Sycrest has not been studied in elderly patients with dementia. However, elderly patients with dementia, who are treated with other similar types of medicine, may have an increased risk of stroke or death. Sycrest is not approved for the treatment of elderly patients with dementia and is not recommended for use in this particular group of patients.

Sycrest may cause low blood pressure. In the early stages of treatment, some people may faint, especially when getting up from a lying or sitting position. This will usually pass on its own but if it does not, tell your doctor. Your dose may need to be adjusted.
Asenapine may cause sleepiness, sudden drops in blood pressure when you stand up, dizziness and changes in your ability to move and balance, which may lead to falls and, consequently, fractures or other injuries. Patients at risk for fall should be evaluated prior to prescribing asenapine.

**Tell your doctor immediately if you experience**
- involuntary rhythmic movements of the tongue, mouth and face. Withdrawal of Sycrest may be needed.
- fever, severe muscle stiffness, sweating or a lowered level of consciousness (a disorder called “neuroleptic malignant syndrome”). Immediate medical treatment may be needed.

Check with your doctor or pharmacist before taking Sycrest:
- if you have ever been diagnosed with a condition whose symptoms include high body temperature and muscle stiffness (also known as neuroleptic malignant syndrome).
- if you have ever experienced abnormal movements of the tongue or face (tardive dyskinesia). You should be aware that both of these conditions may be caused by this type of medicine.
- if you have a heart disease or a treatment for heart disease that makes you prone to low blood pressure
- if you are diabetic or prone to diabetes
- if you have Parkinson’s disease or dementia
- if you have epilepsy (seizures)
- if you experience any difficulty in swallowing (dysphagia)
- if you have severe liver problems. If you do, you should not take Sycrest
- if you have difficulty controlling core body temperature
- if you have thoughts of suicide
- if you have abnormally high levels of prolactin in the blood (hyperprolactinaemia)

Be sure to tell your doctor if you meet any of these conditions as he/she may want to adjust your dose or monitor you for a while. Also contact your doctor immediately if any of these conditions develops or worsens while using Sycrest.

**Children and adolescents**
Sycrest is not recommended for use in patients below the age of 18 years.

**Other medicines and Sycrest**
Tell your doctor or pharmacist if you are taking, have recently taken or might take any other medicines. Some medicines may reduce or increase the effect of Sycrest.

If you are taking other medicines, Sycrest should be taken last.

You should tell your doctor if you are taking antidepressant medicines (specifically fluvoxamine, paroxetine or fluoxetine), as it may be necessary to change your Sycrest or antidepressant medicine dose.

You should tell your doctor if you are taking medicines for Parkinson’s disease (such as levodopa), as this medicine may make them less effective.

Since Sycrest works primarily in the brain, interference from other medicines (or alcohol) that work in the brain could occur due to an additive effect on brain function.

Since Sycrest can lower blood pressure, care should be taken when Sycrest is taken with other medicines that lower blood pressure.

**Sycrest with food, drink and alcohol**
Do not eat or drink for 10 minutes after taking this medicine.
You should avoid drinking alcohol when taking this medicine.

**Pregnancy and breast-feeding**
If you are pregnant or breast-feeding, think you may be pregnant or are planning to have a baby, ask your doctor or pharmacist for advice before taking this medicine.

Do not take Sycrest while you are pregnant, unless your doctor tells you so. If you are taking this medicine and you become pregnant or you plan to get pregnant, ask your doctor as soon as possible whether you may continue taking Sycrest.

The following symptoms may occur in newborn babies, of mothers that have used Sycrest in the last trimester (last three months of their pregnancy): shaking, muscle stiffness and/or weakness, sleepiness, agitation, breathing problems, and difficulty in feeding. If your baby develops any of these symptoms you may need to contact your doctor.

Do not breast-feed when taking Sycrest.

**Driving and using machines**
Sycrest may cause sleepiness or sedation. Therefore, make sure your concentration and alertness are not affected before you drive or operate machinery.

3. **How to take Sycrest**

Always take this medicine exactly as your doctor or pharmacist has told you. Check with your doctor or pharmacist if you are not sure.

The recommended dose is a sublingual tablet of 5 mg or 10 mg two times a day. One dose should be taken in the morning and one dose should be taken in the evening.

**Instructions for use**
Sycrest is for sublingual use.
Sycrest is not advised if you are unable to take the tablet as described below. If you are unable to take this medicine as is described below, the treatment may not be effective for you.
- Do not remove a sublingual tablet from the blister until ready to take it.
- Use dry hands when touching the tablet.
- Do not push the tablet through the blister. Do not cut or tear the blister.
- Peel back the coloured tab (Figure 1).
- Gently remove the tablet (Figure 2). Do not crush the tablet.
- To ensure optimal absorption, place the tablet under the tongue and wait until it dissolves completely (Figure 3). The tablet will dissolve in saliva within seconds.
- Do not swallow or chew on the tablet.
- Do not eat or drink for 10 minutes after taking the tablet.

Figure 1  Figure 2  Figure 3
If you take more Sycrest than you should
If you take too much Sycrest, contact a doctor straight away. Take the medicine pack with you. In case of overdose you may feel sleepy or tired, or have abnormal body movements, problems with standing and walking, feel dizzy due to low blood pressure and feel agitated and confused.

If you forget to take Sycrest
Do not take a double dose to make up for a forgotten dose. If you miss one dose, take your next dose as usual. If you miss two or more doses, contact your doctor or pharmacist.

If you stop taking Sycrest
If you stop taking Sycrest, you will lose the effects of this medicine. You should not stop taking this medicine, unless your doctor tells you as your symptoms may return.

If you have any further questions on the use of this medicine, ask your doctor or pharmacist.

4. Possible side effects

Like all medicines, this medicine can cause side effects, although not everybody gets them.

Serious side effects have been reported with this medicine. Seek medical attention immediately if you have any of the following symptoms:
- allergic reactions (These usually involve a combination of effects such as difficulty in breathing or swallowing, swollen face, lips, tongue or throat, skin rash, itching and increased heart rate.)
- sudden increase in body temperature, with sweating, fast heartbeat, severe muscle stiffness, confusion and fluctuating blood pressure which may lead to coma
- convulsions, fits or seizures
- fainting
- falls which may occur as a result of one or more adverse events such as: sleepiness, sudden drops in blood pressure when you stand up, dizziness and changes in your ability to move and balance.

Tell your doctor right away if you have:
- signs of increased blood sugar levels such as excessive thirst, hunger or urination, weakness or onset of worsening of diabetes
- worm-like movements of the tongue, or other uncontrolled movements of the tongue, mouth, cheeks, or jaw which may progress to arms and legs

Other side effects reported with this medicine include:

**Very common side effects** (may affect more than 1 in 10 people)
- anxiety
- sleepiness

**Common side effects** (may affect up to 1 in 10 people)
- weight gain
- increased appetite
- slow or sustained muscle contractions
- restlessness
- involuntary muscle contractions
- slow movements, tremor
- sedation
- dizziness
- nausea
- change in taste
- numb feeling of the tongue or in the mouth
- increased saliva (drooling)
- muscle tightness
- fatigue
- increase in the level of liver proteins

**Uncommon side effects** (may affect up to 1 in 100 people)
- abnormal muscle movements: a collection of symptoms known as extrapyramidal symptoms (EPS) which may include one or more of the following: abnormal movements of muscles, tongue, or jaw, slow or sustained muscle contractions, muscle spasms, tremor (shaking), abnormal movements of the eyes, involuntary muscle contractions, slow movements, or restlessness
- unpleasant sensations in the legs (also called restless legs syndrome)
- speech problems
- abnormal slow or fast heartbeat
- middle heart block
- abnormal electrocardiogram (prolongation of the QT interval)
- low blood pressure upon standing
- low blood pressure
- tingling of the tongue or in the mouth
- swollen or painful tongue
- difficulty in swallowing
- ulcers, soreness, redness, swelling, and blisters within the mouth
- sexual dysfunction
- lack of regular menstrual periods

**Rare side effects** (may affect up to 1 in 1,000 people)
- changes in the levels of white blood cells
- difficulties in focusing with the eyes
- blood clots in blood vessels to the lungs causing chest pain and difficulty in breathing
- muscle disease presenting as unexplained aches and pains
- male breast enlargement
- leakage of milk or fluid from the breast

**Reporting of side effects**
If you get any side effects, talk to your doctor, pharmacist or nurse. This includes any possible side effects not listed in this leaflet. You can also report side effects directly via the national reporting system listed in Appendix V. By reporting side effects you can help provide more information on the safety of this medicine.

5. **How to store Sycrest**

Keep this medicine out of the sight and reach of children.

Do not use this medicine after the expiry date which is stated on the blister and on the carton. The expiry date refers to the last day of that month.

Store this medicine in the original package in order to protect from light and moisture.

This medicinal product does not require any special temperature storage conditions.

Do not throw away any medicines via wastewater or household waste. Ask your pharmacist how to throw away medicines you no longer use. These measures will help protect the environment.
6. Contents of the pack and other information

What Sycrest contains

- The active substance is asenapine.
- Each Sycrest 5 mg sublingual tablet contains 5 mg asenapine.
- Each Sycrest 10 mg sublingual tablet contains 10 mg asenapine.
- The exact amount is shown on your Sycrest tablet pack.
- The other ingredients are gelatin and mannitol (E421).

What Sycrest looks like and contents of the pack

The 5 mg sublingual tablets are round white to off-white tablets marked with “5” on one side.
The 10 mg sublingual tablets are round white to off-white tablets marked with “10” on one side.

The sublingual tablets are provided in peelable blisters containing 10 tablets each. Packs may contain 20, 60 or 100 tablets.
Not all pack sizes may be marketed.

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Detailed information on this medicine is available on the European Medicines Agency web site: http://www.ema.europa.eu.