Approved by the FDA on October 11, 2019, asenapine transdermal system (Secuado, manufactured by Hisamitsu Pharmaceutical Co., Inc. and distributed by Noven Therapeutics) is the first “patch” approved by the FDA for the treatment of adults with schizophrenia (Table 1).1-3 Asenapine is a second-generation antipsychotic that was previously available only as a sublingual formulation (Saphris, Allergan [now AbbVie] under license from Merck Sharp & Dohme B.V.).4-7 Asenapine’s reformulation potentially simplifies using this antipsychotic by reducing the dosing frequency from twice daily (recommended for the sublingual tablet) to once daily (recommended for the patch). The food and drink restrictions posed by the sublingual formulation are eliminated. Also avoided are dysgeusia (bad taste) and hypoesthesia of the tongue (numbing). Asenapine transdermal system offers a different method to manage schizophrenia, and this novel method of administration through the skin is worth considering.8

The asenapine transdermal system is available in 3 patch sizes: 20, 30, and 40 cm², which deliver 3.8, 5.7, and 7.6 mg/24 hours of asenapine, respectively.3 Based on the average exposure (area under the plasma concentration curve [AUC]) of asenapine, 3.8 mg/24 hours corresponds to 5 mg twice daily of sublingual asenapine, and 7.6 mg/24 hours corresponds to 10 mg twice daily of sublingual asenapine.5 The “in-between” dose strength of 5.7 mg/24 hours would correspond to exposure to a total of 15 mg/d of sublingual asenapine. The recommended starting dose for asenapine transdermal system is 3.8 mg/24 hours. The dosage may be increased to

| Table 1 |
| Fast facts about asenapine transdermal system |

| Brand name: Secuado |
| Class: Second-generation antipsychotic, transdermal formulation |
| Indication: Adults with schizophrenia |
| Approval date: October 11, 2019 |
| Availability date: March 2020 |
| Manufacturer: Hisamitsu Pharmaceutical Co., Inc. (distributed by Noven Therapeutics) |
| Dosing forms: Three patch sizes, 20, 30, and 40 cm², delivering 3.8, 5.7, and 7.6 mg/24 hours of asenapine, respectively |
| Recommended dosage: Recommended starting dose is 3.8 mg/24 hours; the dosage may be increased to 5.7 mg/24 hours or 7.6 mg/24 hours, as needed, after 1 week |

Asenapine transdermal system for schizophrenia

Leslie Citrome, MD, MPH

This transdermal formulation provides sustained delivery of asenapine via a once-daily patch

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Disclosure
No writing assistance or external financial support was utilized in the production of this article. Dr. Citrome has been a consultant for Noven Pharmaceuticals since 2011 on the development of antipsychotic transdermal formulations, and on the development of the asenapine transdermal system since 2014. In the past 5 years Dr. Citrome has engaged in collaborative research with, or received consulting or speaking fees from, AbbVie, Acadia, Alexza, Alkermes, Allergan, Astellas, AstraZeneca, Avanir, Axsome, BioXcel, Boehringer Ingelheim, Bristol-Myers Squibb, Cadent Therapeutics, Eisai, Eli Lilly, Forum, Genentech, Impel, Indivior, Intra-Cellular Therapies, Janssen, Jazz, Karuna, Lundbeck, Lyne, Meiji, Merck, Medivation, Mylan, Neurocrine, Novartis, Noven, Osmotica, Otsuka, Pfizer, Reckitt Benckiser, Relmada, Reviva, Sage, Shire, Sunovion, Takeda, Teva, Valeant, and Vanda.

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5.7 mg/24 hours or 7.6 mg/24 hours, as needed, after 1 week. The safety of doses above 7.6 mg/24 hours has not been evaluated in clinical studies. Asenapine transdermal system is applied once daily and should be worn for 24 hours only, with only 1 patch at any time. Application sites include the upper arm, upper back, abdomen, and hip. A different application site of clean, dry, intact skin should be selected each time a new patch is applied. Although showering is permitted, the use of asenapine transdermal system during swimming or taking a bath has not been evaluated. Of note, prolonged application of heat over an asenapine transdermal system increases plasma concentrations of asenapine, and thus application of external heat sources (eg, heating pads) over the patch should be avoided.

**How it works**

Product labeling notes that asenapine is an atypical antipsychotic, and that its efficacy in schizophrenia could be mediated through a combination of antagonist activity at dopamine D2 and serotonin 5-HT2A receptors. The pharmacodynamic profile of asenapine is complex and receptor-binding assays performed using cloned human serotonin, norepinephrine, dopamine, histamine, and muscarinic receptors demonstrated picomolar affinity (extremely high) for 5-HT2C and 5-HT2A receptors, subnanomolar affinity (very high) for 5-HT7, 5-HT2B, 5-HT6, and D3 receptors, and nanomolar affinity (high) for D2 receptors, as well as histamine H1, D4, a1-adrenergic, a2-adrenergic, D1, 5-HT5, 5-HT1A, 5-HT1B, and histamine H2 receptors. Activity of asenapine is that of antagonism at these receptors. Asenapine

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**Clinical Point**

Asenapine transdermal system should be applied once daily and worn for 24 hours only.
has no appreciable affinity for muscarinic cholinergic receptors.

The asenapine receptor-binding “fingerprint” differs from that of other antipsychotics. Some of these receptor affinities are of special interest in terms of potential efficacy for pro-cognitive effects and amelioration of abnormal mood. In terms of tolerability, a relative absence of affinity to muscarinic receptors would predict a low risk for anticholinergic adverse effects, but antagonism at histamine H1 and at a1-adrenergic receptors, either alone or in combination, may cause sedation, and blockade of H1 receptors would also predict weight gain. Antagonism of a1-adrenergic receptors can be associated with orthostatic hypotension and neurally mediated reflex bradycardia.

**Clinical pharmacokinetics**

Three open-label, randomized, phase 1 studies were conducted to assess the relative bioavailability of asenapine transdermal system vs sublingual asenapine. These included single- and multiple-dose studies and clinical trials that examined the effects of different application sites and ethnic groups, and the effect of external heat on medication absorption. Studies were conducted in healthy individuals, except for the multiple-dose study, which was performed in adults with schizophrenia. The AUC for asenapine transdermal system was within the range of that of equivalent doses of sublingual asenapine, but peak exposure (maximum concentration) was significantly lower. As already noted, the AUC of the asenapine patch for 3.8 mg/24 hours and 7.6 mg/24 hours corresponds to sublingual asenapine 5 mg and 10 mg twice daily, respectively. Maximum asenapine concentrations are typically reached between 12 and 24 hours, with sustained concentrations during the 24-hour wear time. On average, approximately 60% of the available asenapine is released from the transdermal system over 24 hours. Steady-state plasma concentrations for asenapine transdermal system were achieved approximately 72 hours after the first application and, in contrast to sublingual asenapine, the peak-trough fluctuations were small (peak-to-trough ratio is 1.5 for asenapine transdermal system compared with >3 for sublingual asenapine). Dose-proportionality at steady state was evident for asenapine transdermal system. This is in contrast to sublingual asenapine, where exposure increases 1.7-fold with a 2-fold increase in dose. Following patch removal, the apparent elimination half-life

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**Table 2**

<table>
<thead>
<tr>
<th>Definition of response</th>
<th>Placebo N = 203</th>
<th>3.8 mg/24 hours N = 201</th>
<th>7.6 mg/24 hours N = 203</th>
</tr>
</thead>
<tbody>
<tr>
<td>≥30% improvement from baseline Positive and Negative Syndrome Scale total score</td>
<td>18.7</td>
<td>30.8</td>
<td>29.6</td>
</tr>
<tr>
<td>Clinical Global Impression– Improvement score of 1 (very much improved) or 2 (much improved)</td>
<td>34.0</td>
<td>49.8</td>
<td>43.3</td>
</tr>
</tbody>
</table>

*As calculated from the study report (reference 11, Supplementary Table 3). The 95% CI is provided if significant; otherwise ‘ns’ indicates ‘not significant’. CI: confidence interval; NNT: number needed to treat; ns: not significant.

**Clinical Point**

Maximum asenapine concentrations are reached in 12 to 24 hours, with sustained concentrations during the 24-hour wear time.
is approximately 30 hours. The pharmacokinetics of the patch did not vary with regards to the application site (upper arm, upper back, abdomen, or hip area), and the pharmacokinetic profile was similar across the ethnic groups that participated in the study. Direct exposure to external heat did increase both the rate and extent of absorption, so external heat sources should be avoided.

**Efficacy**

The efficacy profile for asenapine transdermal system would be expected to mirror that for sublingual asenapine. In addition to data supporting the use of asenapine as administered sublingually, a phase 3 study specifically assessed efficacy and safety of asenapine transdermal system in adults with schizophrenia. This study was conducted in the United States and 4 other countries at a total of 59 study sites, and 616 patients with acutely exacerbated schizophrenia were enrolled. After a 3- to 14-day screening/single-blind run-in washout period, participants entered a 6-week inpatient double-blind period. Randomization was 1:1:1 to asenapine transdermal system 3.8 mg/24 hours, 7.6 mg/24 hours, or a placebo patch. Each of the patch doses demonstrated significant improvement vs placebo at Week 6 for the primary (change in Positive and Negative Syndrome Scale [PANSS] total score) and key secondary (change in Clinical Global Impression–Severity of Illness) endpoints. Response at endpoint, as defined by a ≥30% improvement from baseline PANSS total score, or by a Clinical Global Impression–Improvement score of 1 (very much improved) or 2 (much improved), was also assessed. For either definition of response, both doses of asenapine transdermal system were superior to placebo, with number needed to treat (NNT) values <10 for the 3.8 mg/24 hours dose (Table 2, page 41). These effect sizes are similar to what is known about sublingual asenapine as determined in a meta-analysis performed by the manufacturer and using individual patient data.

### Table 3

<table>
<thead>
<tr>
<th>Adverse reactions as noted in the highlights of prescribing information for asenapine transdermal system&lt;sup&gt;a&lt;/sup&gt;</th>
<th>Incidence (%)</th>
<th>Incidence (%)</th>
<th>NNH (95% CI)</th>
<th>Incidence (%)</th>
<th>Incidence (%)</th>
<th>NNH (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extrapyramidal disorder</td>
<td>2</td>
<td>8</td>
<td>19 (11-86)</td>
<td>13</td>
<td>10 (7-19)</td>
<td></td>
</tr>
<tr>
<td>Application site reaction</td>
<td>4</td>
<td>15</td>
<td>10 (7-20)</td>
<td>14</td>
<td>11 (7-24)</td>
<td></td>
</tr>
<tr>
<td>Weight gain</td>
<td>2</td>
<td>4</td>
<td>51 (ns)</td>
<td>6</td>
<td>26 (13-495)</td>
<td></td>
</tr>
<tr>
<td>Adverse reactions as noted in the highlights of prescribing information for sublingual asenapine&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Incidence (%)</td>
<td>Incidence (%)</td>
<td>NNH (95% CI)</td>
<td>Incidence (%)</td>
<td>Incidence (%)</td>
<td>NNH (95% CI)</td>
</tr>
<tr>
<td>Extrapyramidal disorder</td>
<td>2</td>
<td>4</td>
<td>67 (ns)</td>
<td>4</td>
<td>51 (ns)</td>
<td></td>
</tr>
<tr>
<td>Oral hypoesthesia</td>
<td>Not applicable</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Somnolence</td>
<td>1.5</td>
<td>4</td>
<td>34 (ns)</td>
<td>3</td>
<td>51 (ns)</td>
<td></td>
</tr>
</tbody>
</table>

<sup>a</sup>As calculated from the available 6-week clinical trials. Lower NNH values denote more frequently encountered adverse reactions. The 95% CI is provided if significant; otherwise ‘ns’ is noted for ‘not significant.’ Data for asenapine transdermal system taken from the product label, drug approval package, and study report, and for sublingual asenapine from the product label and adapted with permission from the author’s prior work. Incidence values <1% are estimated to be 0.5%

CI: confidence interval; NNH: number needed to harm; ns: not significant
The most common adverse reactions are extrapyramidal disorder, application site reactions, and weight gain.

Overall tolerability and safety

The systemic safety and tolerability profile for asenapine transdermal system would be expected to be similar to that for sublingual asenapine, unless there are adverse events that are related to high peak plasma concentrations or large differences between peak and trough plasma concentrations. Non-systemic local application site adverse events would, of course, differ between sublingual vs transdermal administration.

Use of asenapine transdermal system avoids the dysgeusia and oral hypoesthesia that can be observed with sublingual asenapine; however, dermal effects need to be considered (see Dermal safety, page 44). The most commonly observed adverse reactions (incidence ≥5% and at least twice that for placebo) for asenapine transdermal system are extrapyramidal disorder, application site reaction, and weight gain. For sublingual asenapine for adults with schizophrenia, the list includes akathisia, oral hypoesthesia, and somnolence. These adverse events can be further described using the metric of number needed to harm (NNH) as shown in Table 3 (page 42). Of note, extrapyramidal disorder and weight gain appear to be dose-related for asenapine transdermal system. Akathisia appears to be dose-related for sublingual asenapine but not for asenapine transdermal system. Somnolence appears to be associated with sublingual asenapine but not necessarily with asenapine transdermal system.

For sublingual asenapine, the additional indications (bipolar I disorder as acute monotherapy treatment of manic or mixed episodes in adults and pediatric patients age 10 to 17, adjunctive treatment to lithium or valproate in adults, and maintenance monotherapy treatment in adults) have varying commonly encountered adverse reactions.

Both transdermal asenapine system and sublingual asenapine are contraindicated in patients with severe hepatic impairment (Child-Pugh C) and those with known hypersensitivity to asenapine or to any components in the formulation. Both formulations carry similar
warnings in their prescribing information regarding increased mortality in older patients with dementia-related psychosis, cerebrovascular adverse reactions in older patients with dementia-related psychosis, neuroleptic malignant syndrome, tardive dyskinesia, metabolic changes, orthostatic hypotension, leukopenia (and neutropenia and agranulocytosis), QT prolongation, seizures, and potential for cognitive and motor impairment.

Adverse events leading to discontinuation of study treatment in the asenapine transdermal system pivotal trial occurred in 4.9%, 7.8%, and 6.8% of participants in the 3.8 mg/24 hour, 7.6 mg/24 hour, and placebo groups, respectively.

Dermal safety
In the pivotal efficacy study, the incidence of adverse events at patch application sites was higher in the active groups vs placebo (Table 3, page 42). The most frequently reported patch application site reactions were erythema and pruritus, occurring in approximately 10% and 4% in the active treatment arms vs 1.5% and 1.9% for placebo, respectively. With the exception of 1 adverse event of severe application site erythema during Week 2 (participant received 7.6 mg/24 hour, erythema resolved without intervention, and the patient continued the study), all other patch application site events were mild or moderate in severity. Rates of discontinuation due to application site reactions or skin disorders were ≤0.5% across all groups. In the pharmacokinetic studies, no patches were removed because of unacceptable irritation.

Why Rx?
Asenapine transdermal system is the first antipsychotic “patch” FDA-approved for the treatment of adults with schizophrenia. Asenapine has been available since 2009 as a sublingual formulation administered twice daily. The pharmacokinetic profile of the once-daily transdermal system demonstrates dose-proportional kinetics and sustained delivery of asenapine with a low peak-to-trough plasma level ratio. Three dosage strengths (3.8, 5.7, and 7.6 mg/24 hours) are available, corresponding to blood levels attained with sublingual asenapine exposures of 10, 15, and 20 mg/d, respectively. Application sites are rotated daily and include the upper arms, upper back, abdomen, or hip. Dysgeusia and hypohesthesia of the tongue are avoided with the use of the patch, and there are no food or drink restrictions. Attention will be needed in case of dermal reactions, similar to that observed with other medication patches.

Bottom Line
The asenapine transdermal drug delivery system appears to be efficacious and reasonably well tolerated. The treatment of schizophrenia is complex and requires individualized choices in order to optimize outcomes. A patch may be the preferred formulation for selected patients, and caregivers will have the ability to visually check if the medication is being used.
References


Clinical Point

Dysgeusia and hypoesthesia of the tongue are avoided with the use of the patch.