Short-course therapy for recurrent genital herpes and herpes labialis

Entering an era of greater convenience, better treatment adherence, and reduced cost

**Practice recommendations**
- Consider giving patients an oral antiviral (OAV) medication to self-administer when HSV prodromal symptoms occur.
- Patient-initiated, short-course, high-dose OAV treatment of recurrent HSV outbreaks may be as effective as the traditional, longer-course regimens.

**hit early, hit hard.** That expression arose during the evolution of treatment for human immunodeficiency virus (HIV).

Our review focuses on episodic treatment of acute recurrent HSV outbreaks for immunocompetent persons. We do not discuss suppressive therapy, which may be indicated for frequent or severe recurrences (6 or more per year) in immunocompromised patients, or as an adjunctive measure to reduce genital herpes transmission.

As we will describe in detail, the efficacy of the new short-course therapy is, at minimum, comparable to that seen with the older, longer-course trials of topical and oral antiviral therapy. In one head-to-head comparison, Leone et al compared a short-course regimen (3 days) of valacyclovir with 5 days of treatment; they found no difference in results.

If the efficacy of short-course treatment is the same as that of longer courses, the increased convenience and expected improvement in patient adherence with these new regimens argue strongly in their favor. (See Scope of the problem.)

**The strategy**
- Take advantage of a brief therapeutic window

The innate and acquired immune responses of chronically infected, immunocompetent persons rapidly limit cutaneous viral replication, thereby truncating the duration of recurrent HSV outbreaks. In both recurrent herpes labialis and genital herpes, HSV viral titers peak in the first 24 hours following lesion onset (Figure 1A). 

Herpes labialis lesion size and pain are also greatest in the first 24 hours. Most herpes labialis lesions progress from the vesicle stage to the ulcer/soft crust stage within 48 hours, with a hard crust forming by day 2 or 3 (Figure 1B).

With genital lesions, crust formation depends on whether the skin area is dry (3–4 days) or moist (8–9 days).

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The likely events are a burst of virus replication in the first 24 hours of outbreak that lyses basal keratinocytes in a discreet area of epidermis innervated by the infected neuron(s), followed by a vigorous immune response that curtails the infection and creates, in part, the clinical disease (erythema, swelling, vesiculation, and ulceration). The subsequent elements of the illness, which are the majority of the lesion course, are related to wound healing.

**Recognizing the window.** Given the brief period of viral replication and the rapid evolution of lesions, the therapeutic window for treating HSV outbreaks with antiviral drugs is both early and short, making it problematic to effectively treat HSV recurrences. Patients often have mature lesions by the time they consult a physician, rendering subsequent antiviral treatment less effective. However, before lesions appear, many patients experience prodromal symptoms such as pain, burning, or itching. These symptoms can be a prompt to start treatment early, thereby taking advantage of the transient therapeutic window.

If a patient is able to self-administer therapy when prodromal symptoms occur, there may be a greater benefit to treatment. Giving patients drugs for self-administration is therefore an important strategy in managing HSV recurrences.

Traditionally, patient-initiated episodic therapy for recurrent genital herpes and herpes labialis has involved multiple daily doses of topical or oral antiviral agents for 4 to 5 days. Studies of the pathogenesis of HSV recurrences, however, indicate—as said earlier—that the period of virus replication is early and brief, such that a shorter duration of treatment might be more appropriate and equally effective. Other recent clinical studies have indicated that patient-initiated, short-course, high-dose OAV treatment of recurrent HSV infections may be as effective as the traditional therapies in immunocompetent patients. In the section that follows, we examine and compare the results of these trials. (See The agents and how they work.)

**Clinical trials**

**Short-course, high-dose, patient-initiated episodic OAV therapy for recurrent genital herpes**

**Three-day vs 5-day valacyclovir therapy.** The efficacy of 3-day treatment with oral valacyclovir was compared with that of 5-day treatment in immunocompetent patients.
**Scope of the problem**

Herpes simplex virus (HSV) type 1 (HSV-1) or type 2 (HSV-2) results in periodic, recurrent outbreaks of skin lesions after first infection. Herpes labialis (fever blisters or cold sores) is usually caused by HSV-1, while genital herpes is usually caused by HSV-2. Patients with HSV-1 genital herpes typically have fewer recurrences than those with HSV-2 genital infection. The prevalence of HSV-1 and HSV-2 infection varies according to age, geography, gender, and population subgroup, such as people who exhibit high-risk sexual behavior. Approximately 45% of Americans are infected with HSV-1 by adolescence, and approximately 22% of all American adults are infected with HSV-2. The global prevalence of HSV is even greater: as many as 60% to 90% of older adults worldwide are seropositive for HSV-1, and as many as 30% are seropositive for HSV-2. HSV-2 seropositivity is more prevalent among women than men. Overall, the burden of recurrent genital herpes outbreaks can have a profound, negative impact on patient quality of life. The psychological impact of recurrent herpes labialis has not been thoroughly investigated, but an undefined burden is thought to exist, particularly in young patients with frequent or severe recurrences.

Before lesions appear, many patients have prodromal symptoms—pain, burning, or itching—that enable you to start treatment early.

adults with a history of ≥4 episodes of recurrent genital herpes and confirmed HSV infection. Eighty-four participants were randomized to receive 500 mg twice daily valacyclovir for 3 days (and placebo for the remaining 2 days) or 500 mg twice daily for 5 days, and were required to self-administer therapy no later than 24 hours after the onset of symptoms.

The primary endpoint was time to lesion healing (defined as the number of days from initiation of therapy to lesion reepithelialization). Secondary endpoints were pain duration, episode duration (defined as time from initiation of therapy to resolution of all symptoms) and percentage of patients with aborted lesions.

The 3-day valacyclovir treatment exhibited similar time to lesion healing, length of episode, and percentage of patients with aborted lesions as the 5-day treatment (Table 1), suggesting equal efficacy. Duration of pain was also similar (data not shown). Adverse events were similar for both treatment groups, with the most common being headache (10%), nausea (4%), and diarrhea (4%, 5-day treatment vs 2%, 3-day treatment).

**Placebo-controlled trial of 2-day acyclovir therapy.** Wald and coworkers examined the effect of a shorter treatment regimen of acyclovir (2 days) on recurrent genital herpes. Eighty-four immunocompetent HSV-2–infected patients with a history of ≥3 recurrences in the previous 12 months were randomized to receive either 2 days of 800 mg 3 times daily acyclovir or matching placebo. Patients were asked to take their medication no later than 12 hours after the first sign or symptom of an episode.

Efficacy endpoints were time to lesion healing, episode duration, and percentage of patients with aborted lesions. Short-course acyclovir therapy was shown to decrease time to healing (P=.001) and episode duration (P<.001) by 2 days compared with placebo (Table 1). Short-course acyclovir therapy also increased the percentage of patients with aborted lesions compared with placebo (27% vs 11%; P=.029 (Table 1). Adverse events were not recorded in this analysis.

**Placebo-controlled trial of single-day famciclovir therapy.** Aoki and colleagues performed a randomized, double-blind, patient-initiated, placebo-controlled trial to assess the efficacy and safety of patient-initiated, single-day famciclovir 1000 mg twice daily in immunocompetent adults with recurrent genital herpes. The 329 patients in the study were instructed to...
Short-course therapy for recurrent genital herpes

**TABLE 1**

<table>
<thead>
<tr>
<th>DRUG</th>
<th>TREATMENT DURATION</th>
<th>TREATMENT DOSE</th>
<th>CONTROL</th>
<th>MEDIAN TIME (DAYS) TO LESION HEALING (TREATMENT VS CONTROL)</th>
<th>MEDIAN EPISODE DURATION (DAYS) (TREATMENT VS CONTROL)</th>
<th>PATIENTS WITH ABORTED EPISODES (%) (TREATMENT VS CONTROL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valacyclovir(^a)</td>
<td>3 days</td>
<td>500 mg 2x daily</td>
<td>Valacyclovir 500 mg 2x/day for 5 days</td>
<td>4.4 vs 4.7 (P=NS)</td>
<td>4.3 vs 4.4 (P=NS)</td>
<td>25 vs 27 (P=NS)</td>
</tr>
<tr>
<td>Acyclovir(^b)</td>
<td>2 days</td>
<td>800 mg 3x daily</td>
<td>Placebo</td>
<td>4.0 vs 6.0 (P=.001)</td>
<td>4.0 vs 6.0 (P=.001)</td>
<td>27 vs 11 (P=0.029)</td>
</tr>
<tr>
<td>Famiclovir(^c)</td>
<td>1 day</td>
<td>1000 mg 2x daily</td>
<td>Placebo</td>
<td>4.3 vs 6.1 (P&lt;.001)</td>
<td>3.5 vs 5.0 (P&lt;.001)</td>
<td>23 vs 13 (P=.003)</td>
</tr>
</tbody>
</table>

Lesion healing time measures the duration of a subset of severe or classical herpetic outbreaks, characterized by the formation of vesicles, ulcers, or crusts (also papules in some studies\(^a\),\(^b\)). The endpoint is lesional reepithelialization/loss of crust. Episodes where there were only prodromal symptoms, erythema, and/or papule formation (or only symptoms and/or erythema in some studies) were considered “aborted” or prevented lesions. The occurrence of these favorable episode outcomes is described as a percentage of all episodes. Episode duration, sometimes called healing time of all lesions or time to return to normal skin, is the time to resolution of all episodes, regardless of lesion severity. The definition of normal skin varies among the different studies.

NS = not significant.

self-initiate therapy within 6 hours of the onset of prodromal symptoms or genital herpes lesions, and were asked to return to the clinic no later than 24 hours after initiation of therapy. Patients were followed until their lesions healed or for up to 14 days.

The primary endpoint was time to lesion healing of nonaborted lesions. Secondary endpoints were time to healing of all lesions (aborted and nonaborted), time to resolution of pain and other symptoms, and the percentage of patients who did not progress to a full outbreak.

Single-day treatment with famciclovir shortened the time to healing of nonaborted genital herpes lesions by approximately 2 days (P<.001), and the time to healing of all lesions by 1.5 days (P<.001) compared with placebo, and increased the percentage of patients who did not progress to a full outbreak (23% vs 13%; P=.003) (**TABLE 1**). Famiclovir also reduced the time to resolution of all symptoms by approximately 2 days (P<.001) (data not shown).

Adverse events were mild to moderate; the most common in the famciclovir and placebo groups, respectively, were headache (13.5% vs 5.4%), nausea (2.5% vs 3.6%), and diarrhea (4.9% vs 1.2%).

**FAST TRACK**

Studies indicate that the period of HSV replication is early and brief; a short time of treatment might be more appropriate and effective.
TABLE 2

Short-course, patient-initiated OAV therapy is effective against recurrent herpes labialis

<table>
<thead>
<tr>
<th>DRUG</th>
<th>TREATMENT DURATION</th>
<th>TREATMENT DOSE</th>
<th>COMPARATOR REGIMEN</th>
<th>CONTROL</th>
<th>MEDIAN TIME (DAYS) TO LESION HEALING (TREATMENT VS COMPARER VS CONTROL)*</th>
<th>MEDIAN EPISODE DURATION (DAYS) (TREATMENT VS COMPARER VS CONTROL)*</th>
<th>PATIENTS WITH ABORTED LESIONS (% (TREATMENT VS COMPARER VS CONTROL)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Valacyclovir†</td>
<td>1 day</td>
<td>2000 mg 2x daily</td>
<td>Valacyclovir 2000 mg 2x daily x 1 day, 1000 mg 2x daily for a 2nd day</td>
<td>Placebo</td>
<td>Study 1 4.3 vs 4.3 vs 5.1</td>
<td>Study 1 4.0 vs 4.5 vs 5.0</td>
<td>Study 1 44 vs 46 vs 38</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Study 2 4.8 vs 4.6 vs 5.4</td>
<td>Study 2 5.0 vs 5.5</td>
<td>Study 2 43 vs 43 vs 35</td>
</tr>
<tr>
<td>Famciclovir‡</td>
<td>1 dose</td>
<td>1500 mg</td>
<td>Famciclovir 750 mg 2x daily for 1 day</td>
<td>Placebo</td>
<td>4.4 vs 4.0 vs 6.2</td>
<td>4.5 vs 5.7 vs 7.0</td>
<td>33 vs 29 vs 34</td>
</tr>
</tbody>
</table>

Lesion healing time measures the duration of a subset of severe or classical herpetic outbreaks, characterized by the formation of vesicles, ulcers, or crusts (also papules in some studies28-29). The endpoint is lesion reepithelialization/loss of crust. Episodes where there were only prodromal symptoms, erythema, and/or papule formation (or only symptoms and/or erythema in some studies28-29) were considered "aborted" or prevented lesions. The occurrence of these favorable episode outcomes is described as a percentage of all episodes. Episode duration, sometimes called healing time of all lesions or time to return to normal skin, is the time to resolution of all episodes, regardless of lesion severity. The definition of normal skin varies among the different studies.

* All of the healing time and episode duration values for the active treatment arms in both studies differed statistically significantly from placebo, except for famciclovir 750 mg twice daily for 1 day.
† None of the frequencies of aborted lesions in the active treatment arms in either study differed statistically significantly from placebo.

within 24 hours of initiation of therapy.

The primary endpoint in study 1 was clinician-observed duration of all herpes labialis lesions and the secondary endpoint was the percentage of subjects who had herpes labialis lesions that did not progress beyond the papule stage. In study 2, the endpoints were reversed: the primary endpoint was the percentage of patients with lesions that did not progress to full outbreak compared with placebo, but these differences were not statistically significant. The results with 2 days of valacyclovir treatment were similar. Adverse events were similar between the treatment groups and the placebo group.

Placebo-controlled trial of single-dose and single-day famciclovir therapy. Spruance and coworkers assessed patient-initiated famciclovir 1500 mg (single-dose) and 750 mg twice daily (single-day) in immunocompetent adults with recurrent cold sores.30 Subjects (N=1376) were at least 18 years of age and had experienced ≥3 episodes of cold sores over the previous 12 months. Subjects were instructed to administer 1500 mg (single-dose), 750 mg twice daily (single-day), or matching placebo within 1 hour of the onset of prodromal symptoms and before the onset of lesions, and were asked to return
to the clinic within 24 hours of initiating medication.

The primary endpoint was time to healing of primary vesicular lesions. Secondary endpoints included time to healing of all vesicular lesions (primary and secondary [secondary lesions are defined as lesions that developed in addition to and on 1 or more days after primary lesions and that were located at least 1 cm from primary lesions]), time to return to normal skin for all lesions (defined as loss of crust, swelling, and dry flaking), duration of lesion tenderness and pain, and proportion of patients with aborted lesions.

There was a statistically significant decrease in time to healing of primary vesicular lesions by approximately 2 days with both famciclovir treatments compared with placebo, with no significant differences seen in healing between the famciclovir arms (data not shown).

However, only single-dose famciclovir had a statistically significant decrease in the duration of lesion tenderness and pain and the time to return to normal skin compared with placebo (data not shown). No difference was noted between the famciclovir arms in the percentage of patients with aborted lesions compared with placebo. Adverse events in both famciclovir groups were similar to those in the placebo group.

DISCLOSURE
Dr Spruance has received research funding from, been a scientific consultant for, and served on speaker’s bureaus for GlaxoSmithKline and Novartis. Dr Aoki has received funds from GlaxoSmithKline and Novartis for participation in clinical trials and as a member of their Advisory Boards. Dr Tyring has received consultancies, honoraria, and grants from and served on speaker’s bureaus for GlaxoSmithKline and Novartis. Dr Stanberry has received consultancies from GlaxoSmithKline and Novartis. Dr Whitley belongs to speaker’s bureaus for GlaxoSmithKline and Novartis, received grants from the NIH, and is a consultant for Gilead Sciences and Achillion. Dr Hamed is an employee of Novartis.

TOPICAL ANTIVIRAL DRUG FORMULATIONS WERE THE FIRST TREATMENTS APPROVED FOR RECURRENT HSV-1 AND HSV-2 OUTBREAKS, BUT THESE WERE ONLY MARGINALLY EFFICACIOUS. ORALLY-ADMINISTERED ANTIVIRAL AGENTS APPEAR TO BE MORE EFFECTIVE, POSSIBLY BECAUSE OF BETTER DELIVERY OF THE DRUG TO THE SITE OF INFECTION. THREE ORAL ANTIVIRAL AGENTS (OAVs) ARE CURRENTLY APPROVED FOR THE TREATMENT OF RECURRENT GENITAL HERPES: ACYCLOVIR, AN ACYCLIC NUCLEOSIDE ANALOG; VALACYCLOVIR, THE PRODRUG OF ACYCLOVIR; AND FAMCICLOVIR, THE PRODRUG OF PENCICLOVIR, ANOTHER ACYCLIC NUCLEOSIDE ANALOG. ONE OAV (VALACYCLOVIR) IS CURRENTLY APPROVED FOR THE TREATMENT OF HERPES LABIALIS IN IMMUNOCOMPETENT PATIENTS. The prodrugs of acyclovir and penciclovir, valacyclovir and famciclovir, respectively, were synthesized to provide high oral bioavailability and thus permit less frequent administration and potentially greater efficacy compared to the parent compounds.
REFERENCES

FAST TRACK
One day and even a single dose of an oral antiviral decreased lesion healing time and duration of herpes labialis episodes by up to 2 days compared with placebo