Valaciclovir versus aciclovir in patient initiated treatment of recurrent genital herpes: a randomised, double blind clinical trial


Objective: To compare the efficacy and safety of twice daily valaciclovir with five times daily aciclovir in the treatment of an episode of recurrent genital herpes simplex virus (HSV) infection in immunocompetent individuals.

Methods: 739 patients with a history of recurrent genital HSV infection received either oral valaciclovir (500 mg twice daily) or aciclovir (200 mg five times daily) for 5 days for treatment of their next recurrent episode in a controlled, randomised, double blind trial. Patients self initiated therapy at the first signs and/or symptoms of the HSV recurrence, then were assessed in clinic on five occasions over 7 days, and twice weekly thereafter until lesions had healed. Safety was evaluated through adverse experience reports and haematology and biochemistry monitoring.

Results: No significant differences were detected between valaciclovir and aciclovir for the primary endpoint, the duration of all signs and symptoms which included lesion healing and pain/discomfort. The hazard ratio [95% confidence interval] for valaciclovir vs aciclovir was 0.93 [0.79, 1.08]. Lesion healing time was similar in each treatment group [hazard ratio valaciclovir vs aciclovir 0.96 [0.80, 1.14]]. The odds ratio of valaciclovir vs aciclovir in preventing the development of vesicular/ulcerative lesions was 1.08 [0.82, 1.42]. Percentages of patients in whom all HSV cultures were negative were similar in the valaciclovir and aciclovir groups at 59% and 54% respectively; for patients having equal to or more than one positive culture result after treatment initiation, cessation of virus shedding was similarly rapid for the two treatments [hazard ratio 0.98 [0.75, 1.27]]. The safety profiles of valaciclovir and aciclovir were comparable with adverse experiences being infrequent and generally mild.

Conclusion: This study has demonstrated that valaciclovir 500 mg twice daily is equivalent in efficacy to aciclovir 200 mg five times daily as episodic treatment of recurrent genital HSV infection. Valaciclovir maintains the established efficacy and safety of aciclovir but offers a much more convenient twice daily dosing regimen.

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Keywords: valaciclovir; genital herpes; recurrent; herpes simplex virus

Introduction

Genital herpes simplex virus (HSV) infection continues to be a major public health problem throughout the world. Epidemiological surveys undertaken in both industrialised and developing countries indicate that the prevalence of HSV infection is rising rapidly, and HSV is now the most common cause of genital ulceration in many countries.1 The seroprevalence of HSV-2 is highest in Africa (30–40%) followed by the USA (13–40%) and Europe (7–16%).2 Studies show an annual acquisition rate of 2–4% for HSV-2 depending upon the population studied.3 Over the past decade, aciclovir (Zovirax) has become the therapy of choice for the management of recurrent genital HSV infection.4,5 The recommended oral regimen of 200 mg five times daily for five days has been shown to reduce lesion healing time and shorten the course of virus shedding when treatment is initiated within 24 hours of the first signs and/or symptoms of a recurrent HSV episode.6,7 Early treatment may prevent the development of vesicular or ulcerative lesions in some patients if treatment is initiated during the prodrome.8,9 Over 15 years of clinical experience has shown aciclovir to be an effective and well tolerated drug for the management of HSV infections.10 However, its limited oral bioavailability (10–20%) and short plasma elimination half life mean that frequent dosing is required. A formulation providing higher oral aciclovir bioavailability would therefore improve upon its clinical usefulness.11

Valaciclovir (Valtrex, Zelitrex) is the L-valyl ester of aciclovir and once absorbed, is rapidly and almost completely converted to aciclovir and the essential amino acid, L-valine after oral administration. The absolute bioavailability of aciclovir after oral valaciclovir administration is 54%, a three- to fivefold increase over oral aciclovir itself.12,13 Two controlled studies comparing valaciclovir with placebo and with aciclovir and placebo as episodic treatment for recurrent genital HSV infection have shown that both active therapies are effective in speeding resolution of signs and symptoms.14,15 The present study was undertaken to investigate whether the efficacy of valaciclovir twice daily could be maintained with the lower unit dose of 500 mg.
Patients and methods

STUDY DESIGN
This was a multicentre, randomised, double blind comparison of twice daily valaciclovir and five times daily aciclovir given for five days for the treatment of a single episode of recurrent genital herpes infection. For the purpose of this study, genital herpes was defined as a history of recurrent HSV infection involving genital, perianal, or closely related sites.

PATIENTS
Male and female patients of at least 18 years of age were eligible to enter the study if they had experienced four or more recurrences of genital HSV reactivation in the previous 12 months. Those who had been previously receiving suppressive aciclovir had to have experienced a recurrence within three months of stopping therapy. Patients within the three months before enrolment into the study in order to be eligible. Patients were not eligible if they had significant hepatic or renal impairment, were pregnant or nursing mothers, had malabsorption syndrome, were immunocompromised, known to be HIV seropositive, or were receiving probenecid, other systemic antiviral medications, or immunomodulatory treatment.

Ethics committee approval was obtained at each study site before recruitment commenced and all patients gave full written informed consent.

STUDY PROCEDURES
Eligible patients were screened then randomised in equal numbers to one of two treatment arms—oral valaciclovir 500 mg twice daily or oral aciclovir 200 mg five times daily for five days. A blood sample was taken at the screening visit for baseline haematology and clinical chemistry, and to determine HSV-2 serological status if there was no record of a positive HSV culture during the study or a documented history of culture proved genital HSV infection in the patient’s usual medical record. Serological testing was by western blot at a single reference laboratory. Pregnancy tests were performed on all females of childbearing potential at screening and again on the first day of treatment (day 1).

Study medication was dispensed at screening. Patients randomised to valaciclovir also received placebo (dummy) aciclovir tablets; those randomised to aciclovir also received placebo valaciclovir tablets.

Patients self initiated treatment at the first signs or symptoms of their next recurrence. Signs and symptoms included mucocutaneous lesions and/or pain or discomfort. They attended the clinic for evaluation within 24 hours of starting medication. Clinical evaluation and staging of external lesions (macule/papule, vesicle/fissure, crust, or healed) was performed on days 1, 2, 3, 5, and 7. Patients attended the clinic twice weekly thereafter if lesions had not healed by day 7. Diaries were used throughout the treatment and follow up period for patients to record the date and time of prodrome/first signs of genital lesions, compliance with the dosing schedule, the patient’s own assessment of lesion healing, and the pain/discomfort level (no pain, mild, moderate, or severe pain). Swabs for HSV culture were taken at each clinic visit until all lesions had healed. Virus isolation was performed at a single reference laboratory in each city according to local protocol. Safety was assessed by adverse experience reporting at each visit and from haematology (haemoglobin, white blood cell, and platelet counts) and clinical chemistry (creatinine, alkaline phosphatase, and AST or ALT) evaluations on days 1 and 5.

Blood was drawn for determination of steady state plasma aciclovir concentrations by radioimmunoassay on days 2 and 5. Compliance was assessed by pill count of returned blister packs. All patients who did not start treatment were asked to return unused study medication.

EFFICACY ENDPOINTS
The primary efficacy endpoint on which the sample size calculation was based was length of episode, which was defined as the number of days from treatment initiation to complete resolution of all signs and symptoms. In the majority of patients, genital HSV lesions were expected to progress through the usual stages of vesiculation or ulceration, and scabbing before healing. Lesion healing time was therefore also considered a primary efficacy endpoint and was defined as the number of days between initiation of treatment and complete re-epithelialisation of all mucocutaneous lesions. Secondary endpoints were whether or not the HSV episode aborted (defined by the presence of symptoms such as pain/discomfort but failure of lesions to develop beyond the macule/papule stage), duration and severity of pain/discomfort, percentages of patients having no positive HSV culture after treatment initiation, and duration of virus shedding in those patients having at least one positive HSV culture.

SAMPLE SIZE CALCULATION
The minimum sample size of 250 treated patients per arm provided sufficient power to detect differences between treatments in the duration of all signs and symptoms, assuming proportional hazards functions. The sample size provided 80% power to detect hazard ratios between 0.76 and 1.31 at the 5% level of significance, assuming that 45% of the aciclovir group would have signs or symptoms at day five. This is equivalent to detecting an arithmetic reduction of 10% (45% to 35%) in the proportion of patients still having signs or symptoms at day 5. Smaller treatment differences were considered clinically unimportant; therefore, if the resulting 95% confidence intervals for the hazard ratio [95% CI] fell within these clinically unimportant ranges, the power of the study was such that equivalence could be concluded.

STATISTICAL ANALYSIS
The data listing and statistical analyses were performed using Statistical Analysis Systems software version 6.07 (SAS Institute Inc,
Cary, NC, USA). The intent to treat analysis group was defined as all patients randomised who returned to the clinic for assessment of an HSV recurrence. This group was used for the analysis of the primary efficacy endpoint of episodic length and relevant secondary endpoints, as well as for exploratory analyses and the evaluation of safety. Patients in whom vesicular/ulcerative lesions were prevented were excluded from the analysis of lesion healing time. Patients for whom there was no positive HSV culture result during treatment or follow up were excluded from the analysis of duration of virus shedding.

The distributions of time to event endpoints (length of episode, lesion healing time, duration of pain/discomfort, and duration of viral shedding) were estimated using the Kaplan-Meier product limit method. With the exception of duration of viral shedding, all other times to event were calculated in hours, from signs or symptom onset as recorded in patient diaries. Cox’s proportional hazards models were used to estimate treatment differences. Formal hypothesis testing was performed for length of episode. The Cochran-Mantel-Haenszel test was used to investigate treatment differences in the percentages of patients with pain/discomfort on days three and seven and in those in whom vesicular/ulcerative lesions were prevented. Odds ratios were also derived for percentages of patients in whom lesions failed to progress to vesicles/ulcers. Since previous studies have identified sex as an important factor influencing efficacy,\(^8\,^9\) it was included as a covariate in the Cox’s proportional hazard models or logistic regression models as appropriate. Hodges-Lehmann estimates [95% CI] were used to assess any treatment differences or changes over time in haematology and biochemistry variables. All adverse experience data were tabulated and compared across the two treatment groups.

Additional exploratory analyses were conducted on episode length, lesion healing time, and percentages of patients in whom vesicular/ulcerative lesions were prevented. Age (years) and time from prodrome/first sign to initiation of therapy (hours) were fitted as continuous variables; prior use of suppressive aciclovir therapy (used \(v\) not used), and number of recurrences in the previous year \((\leq 8 \, \text{v} \geq 9)\) were also incorporated as covariates into the models in order to identify their impact on the clinical endpoints.

### Results

#### Patient characteristics

In all, 999 patients were randomised to treatment at 48 study sites in Europe and Australia. The intent to treat group consisted of 739 patients who returned for clinic assessment. Overall, 59% of treated patients did not have a documented positive culture for HSV at a genital site and required serological confirmation (table 1). In total, 378 patients were treated with valaciclovir and 361 received aciclovir. Those patients who did not return to the clinic were unlikely to have had a recurrence and therefore would not have commenced study drug treatment.

The demographic characteristics and HSV disease history of treated subjects at screening are shown in table 1 and were similar in the valaciclovir and aciclovir groups. The sex split was equal in both groups; ages ranged from 18 to 77 years (median 32 years). HSV disease history was similar in the valaciclovir and aciclovir groups although slightly more patients in the valaciclovir group had experienced \(\geq 9\) recurrences in the previous 12 months compared with the aciclovir group (37% \(v\) 29%). The median time from the initial HSV episode was approximately four years. Overall, 14%–17% had received suppressive aciclovir therapy in the previous year for a median time of four months. A positive HSV culture was recorded for 41% of patients before enrolment. The slight imbalance noted in previous HSV recurrence frequency necessitated routine inclusion of this factor as a covariate in the efficacy analyses. There were insufficient numbers of patients with proved HSV-1 infection (1%–2%) for separate analysis.

### Analysis of efficacy

#### Length of episode

No significant difference between valaciclovir and aciclovir was detected in length of episode, as evidenced by the hazard ratio [95% CI] of 0.93 [0.79, 1.08]. \(p = 0.34\) (table 2). The median episode duration for the valaciclovir and aciclovir groups was 4.7 and 4.6 days respectively (fig 1).

#### Lesion healing

In those patients with lesions that progressed to vesicles/ulcers, complete re-epithelialisation occurred as rapidly with valaciclovir as aciclovir. The hazard ratio [95% CI] for valaciclovir \(v\) aciclovir was 0.96 [0.80, 1.14] (table

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**Table 1**  Demographic characteristics and HSV disease history

<table>
<thead>
<tr>
<th></th>
<th>Valaciclovir ((n = 378))</th>
<th>Aciclovir ((n = 361))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Median age (years)</td>
<td>32 (18–77)</td>
<td>33 (20–74)</td>
</tr>
<tr>
<td>Sex ((%) male/female</td>
<td>50/50</td>
<td>49/51</td>
</tr>
<tr>
<td>Median time from first genital HSV episode (years)</td>
<td>3.66</td>
<td>4.15</td>
</tr>
<tr>
<td>No of recurrences in previous year (%)</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>1–3</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>4–8</td>
<td>60</td>
<td>68</td>
</tr>
<tr>
<td>(\geq 9)</td>
<td>37</td>
<td>29</td>
</tr>
<tr>
<td>Use of suppressive aciclovir in previous year</td>
<td>17%</td>
<td>14%</td>
</tr>
<tr>
<td>Median time on suppressive therapy (years)</td>
<td>0.34</td>
<td>0.34</td>
</tr>
<tr>
<td>HSV culture before or at screening</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HSV-2</td>
<td>29.9%</td>
<td>25.5%</td>
</tr>
<tr>
<td>HSV-1</td>
<td>0.8%</td>
<td>1.7%</td>
</tr>
<tr>
<td>Not typed</td>
<td>11.6%</td>
<td>12.7%</td>
</tr>
<tr>
<td>HSV-2 infection confirmed by serology*</td>
<td>57.7%</td>
<td>60.1%</td>
</tr>
</tbody>
</table>

*Performed on patients without documented positive HSV culture at genital site.*
Table 2  Valaciclovir versus aciclovir: time to event endpoints

<table>
<thead>
<tr>
<th></th>
<th>Valaciclovir v aciclovir</th>
<th>Hazard ratio (95% CI)</th>
<th>Median time to event (days)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length of episode*</td>
<td>0.93 (0.79, 1.08) $p = 0.34$</td>
<td>4.7 ± 4.6</td>
<td></td>
</tr>
<tr>
<td>Lesion healing†</td>
<td>0.96 (0.80, 1.14)</td>
<td>4.4 ± 4.5</td>
<td></td>
</tr>
<tr>
<td>Duration of pain/discomfort‡</td>
<td>overall</td>
<td>0.91 (0.78, 1.06)</td>
<td>2 ± 2</td>
</tr>
<tr>
<td></td>
<td>males</td>
<td>0.89 (0.71, 1.12)</td>
<td>2.5 ± 2.0</td>
</tr>
<tr>
<td></td>
<td>females</td>
<td>0.97 (0.77, 1.22)</td>
<td>3.0 ± 3.0</td>
</tr>
<tr>
<td>Duration of virus shedding‡</td>
<td>overall</td>
<td>0.98 (0.75, 1.27)</td>
<td>2 ± 2</td>
</tr>
<tr>
<td></td>
<td>males</td>
<td>0.93 (0.65, 1.33)</td>
<td>2 ± 2</td>
</tr>
<tr>
<td></td>
<td>females</td>
<td>0.95 (0.61, 1.47)</td>
<td>2 ± 2</td>
</tr>
</tbody>
</table>

*Time from treatment initiation to resolution of all signs and symptoms (n = 710).
†Defined as loss of crusts and re-epithelialisation, includes only patients who developed vesicular/ulcerative lesions (n = 559).
‡All patients (n = 712).
§Proportional hazards assumption not satisfied for sex, therefore separate analyses of males and females was necessary.

Figure 1  Kaplan-Meier plot of percentages of patients with signs or symptoms of a genital herpes recurrence following treatment with valaciclovir or aciclovir.

Figure 2  Kaplan-Meier plot of percentages of patients with cutaneous lesions of a genital herpes recurrence not yet healed following treatment with valaciclovir or aciclovir.

Table 3  Valaciclovir versus aciclovir for recurrent genital herpes: prevention of lesion development, pain, and virus shedding proportions

<table>
<thead>
<tr>
<th></th>
<th>Valaciclovir</th>
<th>Aciclovir</th>
<th>Valaciclovir v aciclovir Relative risk (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage in whom vesicular/ulcerative lesions were prevented</td>
<td>22</td>
<td>21</td>
<td>1.08 [0.82, 1.42]</td>
</tr>
<tr>
<td>overall</td>
<td>20</td>
<td>18</td>
<td>1.26 [0.82, 1.93]</td>
</tr>
<tr>
<td>males</td>
<td>43</td>
<td>42</td>
<td>0.96 [0.67, 1.37]</td>
</tr>
<tr>
<td>females</td>
<td>11</td>
<td>11</td>
<td>0.70† [0.40, 1.23]</td>
</tr>
<tr>
<td>Percentage with pain on day 3*</td>
<td>50</td>
<td>52</td>
<td>0.87† [0.55, 1.38]</td>
</tr>
<tr>
<td>none</td>
<td>43</td>
<td>42</td>
<td>1.08 [0.67, 1.73]</td>
</tr>
<tr>
<td>mild</td>
<td>13</td>
<td>11</td>
<td>1.08 [0.58, 2.04]</td>
</tr>
<tr>
<td>moderate/severe</td>
<td>11</td>
<td>11</td>
<td>1.08 [0.58, 2.04]</td>
</tr>
<tr>
<td>Percentage with pain on day 7‡</td>
<td>59</td>
<td>54</td>
<td>ND</td>
</tr>
<tr>
<td>none</td>
<td>50</td>
<td>52</td>
<td>0.87† [0.55, 1.38]</td>
</tr>
<tr>
<td>mild</td>
<td>4</td>
<td>2</td>
<td>1.08 [0.67, 1.73]</td>
</tr>
<tr>
<td>moderate/severe</td>
<td>1</td>
<td>1</td>
<td>1.08 [0.58, 2.04]</td>
</tr>
<tr>
<td>Percentage with no positive HSV culture</td>
<td>59</td>
<td>54</td>
<td>ND</td>
</tr>
</tbody>
</table>

*Pain scores missing for some patients.
†Cochran–Mantel-Haenszel statistic.
ND = not determined.

2). Median healing times were 4.4 and 4.5 days for the valaciclovir and aciclovir groups respectively (fig 2).

Prevention of vesicular/ulcerative lesions

The percentages of patients in the valaciclovir and aciclovir groups in whom vesicular/ulcerative lesions were prevented was similar, at 22% and 21% respectively, and did not differ with sex (table 3). The odds ratio [95% CI] for the chance of vesicular lesions being prevented with valaciclovir treatment compared with aciclovir was 1.08 [0.82, 1.42].

Pain/discomfort

No differences between valaciclovir and aciclovir were evident in the percentages of patients experiencing no/mild pain or moderate/severe pain on day 3 or day 7 (table 3). Resolution of pain/discomfort was achieved as rapidly with valaciclovir as aciclovir in all patients and in males and females when analysed separately. Hazard ratios [95% CI] indicated no treatment differences (table 2).

Virus shedding

In patients with at least one swab for HSV culture, results were all negative in similar proportions in each treatment group, at 59% for valaciclovir and 54% for aciclovir (table 3). For those in whom at least one culture was positive during treatment or follow up, termination of virus shedding was as rapid in each group, as evidenced by the hazard ratio [95% CI] for valaciclovir v aciclovir of 0.97 [0.75, 1.26], with no differences being apparent between the sexes (table 2).

Exploratory analyses

Results of the exploratory analyses suggest that episode resolution was 20% faster in those patients who had previously received suppressive aciclovir therapy in the year before enrolment compared with those who had not (hazard ratio [95% CI] 0.80 [0.63, 1.01] p = 0.06). Earlier treatment initiation accelerated episode resolution (hazard ratio for the difference [95% CI] 0.99 [0.98, 1.00] p = 0.01) and lesion healing (hazard ratio for the difference [95% CI] 0.99 [0.98, 1.00] p = 0.03). Hence, in patients starting treatment six hours after first signs/symptoms, episode resolution and lesion healing were 15% and 13% faster, respectively, than in those starting treatment 24 hours after first signs/symptoms.

ANALYSIS OF SAFETY

Adverse experiences reported during treat-

Table 4  Adverse experiences reported by $\geq 2\%$ of patients

<table>
<thead>
<tr>
<th>Adverse experience</th>
<th>Valaciclovir n = 378 (%)</th>
<th>Aciclovir n = 361 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Headache</td>
<td>14</td>
<td>9</td>
</tr>
<tr>
<td>Nausea</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Diarrhoea</td>
<td>2</td>
<td>5</td>
</tr>
<tr>
<td>Abdominal pain</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Asthenia</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Dizziness</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Insomnia</td>
<td>2</td>
<td>1</td>
</tr>
</tbody>
</table>
ment and follow up, irrespective of association, which occurred in >2% of subjects, are listed in table 4. The most frequent were headache and nausea; no major differences between treatments were evident. Headache was considered possibly attributable to valaciclovir or aciclovir in 10% and 7% of patients, with nausea thought possibly attributable in 8% and 7%, respectively. Three adverse experiences in three patients were considered serious, but only one, a patient receiving aciclovir who suffered abdominal pain, was thought possibly attributable to the drug. Treatment was discontinued prematurely in five patients (valaciclovir three, aciclovir two) as a result of adverse experiences.

There were no clinically important changes from screening in any haematology or clinical chemistry variable with valaciclovir or aciclovir treatment or differences between groups during treatment or between values from samples taken on day one and day five. Mean serum creatinine values (SD) on day one were 87·9 (14·7) and 87·2 (13·2) µM for the valaciclovir and aciclovir groups, respectively. On day five, respective mean values were 87·9 (14·1) and 87·5 (13·8) µM, indicating no meaningful differences between the groups or with treatment.

**PLASMA ACICLOVIR LEVEL MONITORING**

Estimated mean peak aciclovir concentrations for patients randomised to valaciclovir and aciclovir were 12·3 µM (2·8 µg/ml) and 2·9 µM (0·65 µg/ml) respectively. Corresponding estimates of the areas under the aciclovir plasma concentration time curves over 24 hours were 99·5 µM/h (22·4 µg/ml) and 49·3 µM/h (11·1 h/µg/ml). Thus, the 500 mg valaciclovir regimen increased the systemic exposure to aciclovir by twofold over the standard oral aciclovir regimen. Since a 500 mg dose of valaciclovir contains approximately 347 mg aciclovir, the bioavailability of aciclovir from valaciclovir 500 mg twice daily was 2·9 times greater than that from aciclovir 200 mg five times daily in this study.

**Discussion**

This study has found that valaciclovir at a dose of 500 mg twice daily for five days and aciclovir at the standard dose of 200 mg five times daily also for five days are equally effective in patient initiated treatment of recurrent genital herpes. Moreover, this clinical equivalence is achieved without apparent compromise to the excellent safety profile established for aciclovir. This was one of two large trials conducted to examine a 500 mg twice daily regimen of valaciclovir, after an earlier placebo controlled study of similar design demonstrated that 1000 mg was as effective as the standard aciclovir regimen in shortening all signs and symptoms of a recurrence. The companion study has shown 500 mg and 1000 mg unit doses of valaciclovir taken twice daily to be equivalent for the same efficacy endpoints and both to be superior to placebo. From this and the results of the present study, it may be concluded that valaciclovir 500 mg twice daily achieves the same clinical efficacy in treating genital herpes recurrences as the standard oral aciclovir regimen.

The five times a day dosing regimen for aciclovir for recurrent genital herpes has been supported by several clinical studies and is currently recommended by the US Centers for Disease Control and Prevention. An effective twice daily treatment regimen for recurrent genital herpes may confer benefits in addition to the greater convenience and improved compliance usually attributed to simplified dosing schedules. There is a degree of general community awareness of the indications for aciclovir (Zovirax). Consequently, many patients with anxiety about the potential social consequences of their diagnosis, including fear of detection, will appreciate a regimen that allows all treatment to be taken (and stored) in the privacy of the home.

Simulated plasma aciclovir concentration vs time profiles indicated that the 500 mg twice daily valaciclovir regimen would increase systemic aciclovir exposure by 1·5 to 2-fold compared with that following oral aciclovir. Plasma aciclovir level monitoring in this study in young adults with genital herpes, a population of similar age to that in the formal phase 1 study, has confirmed that at steady state, 500 mg twice daily valaciclovir does result in the predicted doubling of aciclovir exposure.

The results of this study have demonstrated both the clinical and statistical equivalence in efficacy of valaciclovir and the standard oral aciclovir regimen on the primary endpoint. We believe that the study was adequately protected against type 2 error; a 10% difference in the number of patients with signs or symptoms at day 5 could have been detected with 80% power. Although not statistically powered for hypothesis testing on all the other efficacy endpoints, analyses consistently indicated in each case no evidence for any meaningful differences between treatment groups. Of particular clinical importance are the results showing that vesicular lesions were prevented in similar percentages of patients and that in 59% and 54% of valaciclovir and aciclovir recipients, respectively, no HSV culture was positive once treatment had begun.

The primary efficacy endpoint, length of episode, encompasses all clinical signs and symptoms, specifically the more traditional and easily quantifiable resolution of pain, itching, or other discomfort. The intent to treat population, on which the primary time to event analysis of length of episode was based, thus comprises two subpopulations defined according to clinical outcome—those in whom clinically evident skin lesions progress through all stages of healing and those whose initial symptoms indicated a pending recurrence but which subsequently fails to develop beyond the papule stage (aborted episode). Lesions that fail to progress have been recognised in several earlier clinical trials of aciclovir and represent the most desirable clinical outcome.

Patient initiated treatment is designed to
Valaciclovir versus aciclovir in patient-initiated treatment of recurrent genital herpes: a randomised, double-blind clinical trial

Valaciclovir confers all the clinical benefits of aciclovir but with the convenience and security of a twice daily dosing regimen and without apparent compromise to aciclovir’s excellent safety profile.

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The International Valaciclovir HSV Study Group

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**Gram’s stain**

**Historical vignette**

Hans Christian Joachim Gram was born in Copenhagen in 1853, the son of a professor of law. He qualified in medicine in 1878, and after holding junior posts he joined a course in microbiology. His tutor formed a high opinion of him, and asked Carl Friedländer if he could study in his laboratory in Berlin. He arrived there in 1883, and almost at once embarked on an investigation into the problem of identifying bacteria in stained tissue sections. The techniques in use at this time made this difficult because cell nuclei, fibrin, and bacteria stained with equal intensity. Gram found that if he applied a solution of iodine and potassium iodide to sections already stained with gentian violet, then immersed them in absolute alcohol, he could decolourize the tissue cells without changing the deep blue colour of many bacteria. Subsequently, he found it useful to counterstain the tissue with Bismarck or aniline brown.

Three weeks after his arrival in Berlin, Gram wrote to his tutor in Copenhagen: “I have had the good luck to find what seems to be a very good method of staining cocci while the tissue and cell nuclei remain unstained. Dr F is delighted with the method.” Friedländer soon mentioned Gram’s method in a paper on the micrococi of pneumonia, then Gram himself published a detailed account of it. By this time, he had found that it could be applied to smears as well as tissues, so that it was possible to distinguish bacteria which retained their blue colour after the iodine/alcohol treatment (“Gram positive”) from those which were decolourized (“Gram negative”). He ended his paper with these words: “Hopefully, the method will prove useful in the hands of other investigators.” Having said this, he did no further work on the technique, leaving its refinement to others. Having been in Berlin for only five months, he returned to Copenhagen, where in due course he became professor of pharmacology and senior physician at Frederik Hospital. He died in 1938 at the age of 85 years.

Neisser first identified the gonococcus in 1879 by staining smears of pus from the male urethra with methyl violet, and it was soon realised that it stained equally well with other basic aniline dyes such as gentian violet and methylene blue. Gram did not try his technique on the gonococcus himself, but in 1886 Roux wrote: “One can always recognise the true nature [of gonococci] if, after establishing their presence with gentian violet alone, the liquid of Gram and alcohol are successively added. If the cocci completely disappear, they are indeed those of Neisser.” Counterstaining with Bismarck brown revealed the organisms more clearly, but later workers found that a better contrast was obtained with carbol fuchsin or methyl red.

Gram staining for the diagnosis of gonorrhoea was widely used by the end of the 19th century although some venereologists still preferred a single stain, particularly methylene blue. A few physicians—Harrison was one—advocated staining by methylene blue to confirm the result of a Gram stain, but in modern times strict protocols have made such precautions unnecessary. Gram’s is the commonest staining procedure in venereology, and indeed in the whole of clinical microbiology. In later life he was amused that he was best known for a single discovery that he had made, almost by chance, as a young man. He said he had been lucky, but luck often favours those who deserve it.

J D Oriel
Valaciclovir versus aciclovir in patient initiated treatment of recurrent genital herpes: a randomised, double blind clinical trial.

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