Clinical Trial of Herbal Treatment Gene-Eden-VIR/Novirin in Oral Herpes

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Abstract

Background. Our previous articles showed that suppressive or preventive treatment with the herbal Gene-Eden-VIR/Novirin reduced the number and duration of genital herpes outbreaks with no adverse effects. These studies also revealed that the herbal Gene-Eden-VIR/Novirin is mostly superior to acyclovir, valacyclovir, and famciclovir drugs in genital herpes. This study tested the effect of Gene-Eden-VIR/Novirin in oral herpes (also called cold sores and fever blisters). Methods. The framework of the study was a retrospective chart review. The study included 68 participants. The participants took 1 to 4 capsules per day over a period of 2 to 36 months. The study included 2 Food and Drug Administration–recommended controls: baseline and a no-treatment. Results. Gene-Eden-VIR/Novirin was effective in 89.3% of participants. The treatment reduced the mean number of outbreaks per year from 6.0 and 3.6 in the control groups to 2.0 in the treatment group ($P < .0001$ and $P = .07$, respectively). Gene-Eden-VIR/Novirin reduced the mean duration of outbreaks from 9.8 and 5.8 days in the control groups to 3.2 days in the treatment group ($P < .0001$ and $P = .02$, respectively). There were no reports of adverse experiences. Gene-Eden-VIR/Novirin was compared to acyclovir and valacyclovir in 6 tests. In all tests, Gene-Eden-VIR/Novirin showed higher efficacy. Gene-Eden-VIR/Novirin also showed superior safety. Conclusions. This clinical study showed that suppressive or preventive treatment with the herbal Gene-Eden-VIR/Novirin reduced the number and duration of outbreaks in oral herpes without any adverse effects. The study also showed that the herbal Gene-Eden-VIR/Novirin had better clinical effects than acyclovir and valacyclovir, the leading drugs in the category. Based on these results, we recommend using the herbal Gene-Eden-VIR/Novirin as preventive treatment for oral herpes and, specifically, as an alternative to the acyclovir and valacyclovir drugs.

Keywords

HSV1, HSV2, acyclovir, valacyclovir, famciclovir, natural treatment, herbal treatment, oral herpes, cold sores, fever blisters, outbreaks, gene-Eden-VIR, Novirin

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The output of the sessions are the charts that we analyzed in this study. Clinical studies have shown that a 4-month suppressive treatment with ACV and VACV decreased the mean number of outbreaks by more than 50%, and increased the number of recurrence-free patients by more than 20%. Furthermore, suppressive treatment resulted in a longer time (24-72 days) to first recurrence.

Gene-Eden-VIR/Novirin is a patented herbal treatment. The Gene-Eden-VIR/Novirin formula includes 5 ingredients: a 100 mg extract of quercetin, a 150 mg extract of green tea, a 50 mg extract of cinnamon, a 25 mg extract of licorice, and 100 μg of selenium. According to microcompetition theory, latent viruses cause most major diseases. To reverse the effect of latent viruses on the host, a team of scientists developed the herbal Gene-Eden-VIR/Novirin. The treatment was introduced to the public at the end of 2009. Previous clinical studies conducted at the Center for the Biology of Chronic Disease tested the effect of Gene-Eden-VIR/Novirin on viruses. The studies showed that the herbal treatment has an antiviral effect. Since viruses are linked to fatigue, another clinical study tested the effect of Gene-Eden-VIR/Novirin on fatigue. The study reported that Gene-Eden-VIR/Novirin safely reduced the feeling of fatigue in individuals infected with a latent virus. In 2016, Polansky et al reported that Gene-Eden-VIR/Novirin safely decreased the number and duration of outbreaks in individuals suffering from genital herpes. The study also compared the clinical effects of the herbal Gene-Eden-VIR/Novirin to the leading drugs ACV, VACV, and FCV. The comparison showed that Gene-Eden-VIR/Novirin was better than these drugs.

This article reports the effects of suppressive or preventive treatment with Gene-Eden-VIR/Novirin on the number and duration of oral herpes outbreaks. The article also compares the effects of the herbal Gene-Eden-VIR/Novirin to those of ACV and VACV, the 2 leading drugs in the category.

Methods

Objective

The objective of the study was to measure the effects of suppressive or preventive treatment with the herbal Gene-Eden-VIR/Novirin on the frequency and duration of oral herpes outbreaks and to compare these effects to those of the leading drugs in the category.

Framework

The framework of the study was a retrospective chart review. Lilac Corp, the company that sells Gene-Eden-VIR/Novirin to the public, assists its customers in tracking the effects of the treatment on their health. To track these effects, Lilac Corp uses the Natural Origin Treatment Clinical Questionnaire (NotCiq). Professional interviewers collect the answers to the questionnaire over the phone in single sessions. The answers are considered patient-reported outcomes (PROs). The output of the sessions are the charts that we analyzed in this study.

Efficacy

The primary end points were the frequency, or number of outbreaks per period, and the duration of outbreaks, or the number of days passed between initiation of symptoms and signs and complete resolution of the symptoms. There were several secondary end points. Those included the number of participants who had over a 50% reduction in their recurrence rate, the percentage of patients who were recurrence-free, the time to a first recurrence in all participants, and the percentage of patients with prevented or delayed recurrences.

Ethical Consideration

The institutional review board that approved the study was Salus Institutional Review Board. Since the study was a retrospective chart review, the institutional review board approved a waiver of the requirement to obtain an informed consent from the participants under the exemption status of the federal regulations 45 CFR 46.101(b)(4).
Safety
We analyzed all reports of adverse events.

Population
We excluded all participants who used Gene-Eden-VIR/Novirin for other purposes, including those who used it as treatment for other diseases. We also excluded participants who were concurrently using antiviral drugs a suppressive treatment, specifically ACV or VACV. In the duration of outbreaks analysis, 2 additional participants were excluded due to concurrent episodic treatment with ACV. The final list of participants included 68 participants. All participants had at least one oral herpes outbreak per year.

Statistical Analysis
The analysis of the data in the study was based on the intent-to-treat population. We used the Kaplan-Meier product-limit method to calculate the time to first recurrence of an oral herpes outbreak. We evaluated the differences between the curves using the log-rank, Wilcoxon, and Tarone-Ware tests. We also calculated the difference between pretreatment and treatment (Δ), and calculated the statistical difference between the deltas. We performed the statistical analysis using a one-tail t test assuming unequal variances, or a single-factor analysis of variance. We considered a P ≤ .05 as statistically significant.

Results

Patient Demographic and Baseline Characteristics
The study included 68 participants. We divided the participants into 2 groups: a treatment group, which included 56 participants, and a no-treatment control group, which included 12 participants. Note that the 56 individuals in the treatment group also comprise the 56 individuals in the baseline or pretreatment control group (see Table 2). Table 1 summarizes the demographic and clinical characteristics of the pretreatment/treatment and no-treatment groups.

Efficacy
Primary Efficacy Endpoints. Out of the 56 participants in the pretreatment group, 89.3% reported a decrease, and 80.4% a greater than 50% decrease in the number (frequency) of outbreaks per year (Table 2). The mean number of outbreaks per year decreased from 3.58 and 5.96, in the no-treatment and pretreatment controls, respectively, to 2.02 in the treatment group (44% and 66%, P = .07 and P < .0001, respectively; Table 2).

Out of 56 participants in the pretreatment group, 91% reported a decrease in the duration of outbreaks. The mean duration of outbreaks decreased from 5.83 days and 9.78 days,

Table 1. Demographic and Clinical Characteristics of the Participants.

|                      | Pretreatment and Treatment Groups | No-Treatment Group |
|----------------------|----------------------------------|-------------------|
| Age, average (years) | 49                               | 46                |
| Age, n (%)           |                                  |                   |
| 20-40                | 15 (26.8%)                       | 2 (16.7%)         |
| 41-50                | 14 (25.0%)                       | 5 (41.7%)         |
| 51-60                | 16 (28.6%)                       | 4 (33.3%)         |
| 61-80                | 11 (19.6%)                       | 1 (8.3%)          |
| Gender, n (%)        |                                  |                   |
| Male                 | 33 (58.9%)                       | 3 (25%)           |
| Female               | 23 (41.1%)                       | 9 (75%)           |
| Race, n (%)          |                                  |                   |
| African American     | 8 (14.3%)                        | 4 (33.3%)         |
| Caucasian            | 36 (64.3%)                       | 8 (66.7%)         |
| Hispanic             | 6 (10.7%)                        | 0 (0%)            |
| Other                | 6 (10.7%)                        | 0 (0%)            |
| Years since diagnosis by physician | 0.5-50 (range), 9.0 (mean), 4.5 (median) | 1-40 (range), 8.5 (mean), 2.5 (median) |
| Years since initial episode | 1-40 (range), 18.7 (mean), 19 (median) | 1-40 (range), 13.5 (mean), 12 (median) |
| Percentage diagnosed by physician | 67.9% | 58.3% |
| Percentage that had a laboratory test to confirm diagnosis (out of those diagnosed by a physician) | 65.8% | 57.1% |
| Symptoms of infection, n (%) |                                  |                   |
| Oral blisters/ulcers | 49 (87.5%)                       | 9 (75.0%)         |
| Local pain           | 38 (67.9%)                       | 9 (75.0%)         |
| General discomfort   | 25 (44.6%)                       | 6 (50.0%)         |
| Light sensitivity    | 13 (23.2%)                       | 2 (16.7%)         |
| Flu-like symptoms    | 20 (35.7%)                       | 7 (58.3%)         |
| Duration of treatment (months), n (%) |                                  |                   |
| 2-5                  | 21 (37.5%)                       | N/A               |
| 6-18                 | 24 (42.9%)                       | N/A               |
| >18                  | 11 (19.6%)                       | N/A               |
in the no-treatment and pre-treatment control groups, respectively, to 3.21 days in the treatment group ($P = .02$ and $P < .0001$, respectively; Table 2).

We tested the internal consistency of the participants’ answers using a second set of questions. On frequency of symptoms, the participants were asked, “How often did the symptoms appear before (after) taking Gene-Eden-VIR/Novirin?” using a scale from 1 to 7, where 1 is “very often” and 7 is “not at all.” The mean scores were 3.05 and 5.57 for the “before” and “after” taking Gene-Eden-VIR/Novirin questions, respectively ($P < .0001$). On duration of symptoms, the participants were asked, “How long did your symptoms last, before (after) taking Gene-Eden-VIR/Novirin?” using a scale from 1 to 7, where 1 is “very long time,” and 7 is “didn’t have symptoms.” The mean scores were 3.00 and 5.73 for the “before” and “after” taking Gene-Eden-VIR/Novirin questions, respectively ($P < .0001$). These results show that the participants provide consistent answers.

To test for a possible difference between current and past users of Gene-Eden-VIR/Novirin, we compared the answers of these groups. The decrease in number of days per episode in current and past users was 6.71 and 6.28, respectively ($P = .36$). The decrease in number of episodes per year in current and past users was 4.32 and 3.17, respectively ($P = .15$). No statistical difference was found between the answers provided by participants in the 2 groups.

We also tested the effect of duration of treatment. We divided the treatment group into 3 subgroups according to their duration of treatment: 2 to 5 months ($N = 21$), 6 to 18 months ($N = 24$), >18 months ($N = 11$). The number of outbreaks per year for the 3 subgroups before treatment were 5.77, 4.57, and 8.27, for the 2 to 7 months, 8 to 18 months, and over 18 months, respectively. There was no statistically significant difference between these values ($P = .15$). The results showed that there is a statistically significant difference in the delta values between the 3 subgroups (3.05, 3.25, and 7.18, respectively, $F[2, 53] = 5.43$, $P = .007$; Table 3). Longer duration of treatment was associated with a larger decrease in the frequency of symptoms. These results show a larger effect of treatment.

**Diagnosis**

Thirty-eight participants were diagnosed by a physician and 18 used self-diagnosis. We tested the differences between these groups. Table 4 presents the results. All participants in the 2 groups reported a statistically significant improvement in the number, duration, severity, interference in daily life, and the level of pain. However, those diagnosed by a physician were younger, had more episodes, experienced more severe and painful symptoms, and reported a greater interference in their daily life (borderline significance) during the pretreatment period. In contrast, the participants in the 2 groups report similar duration of episodes during this period. The results also showed that there is no statistically significant difference in the treatment period between the 2 groups in all aspects except the interference in daily life (borderline significance). These results are consistent with those reported in the literature on help-seeking behavior, which shows a positive relationship between the level of pain and disability and the likelihood of seeking professional help.

**Secondary Efficacy Endpoints.** We used the Kaplan-Meier product-limit method to calculate the time to first recurrence. The number of days to a first recurrence, or mean survival time, was 106.5 and 175.91 during the pretreatment and no-treatment periods, respectively, and 364.6 days during the treatment period ($P < .0001$ and $P = .01$ for the pretreatment and no-treatment controls, respectively; Figure 1). These results show that treatment with the herbal Gene-Eden-VIR/Novirin increased the time to a first recurrence.

No participants in both the pretreatment and no-treatment controls were recurrence-free (Table 2). Following treatment with the herbal Gene-Eden-VIR/Novirin, 46.4% were recurrence-free ($P < .0001$). Also, the herbal Gene-Eden-VIR/Novirin treatment prevented or delayed 89% of the

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**Table 2.** Summary of the Efficacy Endpoints in the Current Study. All Participants Had at Least One Outbreak per Year.

| Treatment   | Mean Number of Outbreaks per Year | Time to First Recurrence (Days) | % Recurrence-Free | % With Decrease in Recurrence | % With >50% Decrease | Mean Duration of Outbreak (Days) |
|-------------|----------------------------------|---------------------------------|------------------|-------------------------------|----------------------|-------------------------------|
| No-Tx control | 12                               | 3.58                            | 175.9            | 0.0%                          | —                    | 5.83                          |
| Pre-Tx control | 56                              | 5.96                            | 106.5            | 0.0%                          | 89.3%                | 9.78                          |
| Tx           | 56                               | 2.02 ($P = .07$, no-Tx)         | 364.6 ($P < .0001$, pre-Tx) | 46.4% ($P < .0001$, both controls) | 80.4% | 3.2 ($P = .02$, no-Tx) ($P < .0001$, pre-Tx) |

Abbreviation: Tx, treatment.

*Using Kaplan-Meier test.

**Table 3.** Duration of Treatment Effect in Those Taking Gene-Eden-VIR/Novirin for 2 to 5 Months Versus Those Taking Gene-Eden-VIR/Novirin for 6 to 18 Months and More Than 18 Months.

| Duration of Treatment | Delta (Change in Number of Outbreaks) | $P$ Value | $F$ Value |
|-----------------------|---------------------------------------|-----------|-----------|
| 2-5 months ($n = 21$) | 3.05                                  | $P < .01$ | $F(2, 53) = 5.43$ |
| 6-18 months ($n = 24$) | 3.25                                  |           |           |
| >18 months ($n = 11$) | 7.18                                  |           |           |

**Table 4.** Summary of the Efficacy Endpoints in the Current Study. All Participants Had at Least One Outbreak per Year.

| Treatment   | Mean Number of Outbreaks per Year | Time to First Recurrence (Days) | % Recurrence-Free | % With Decrease in Recurrence | % With >50% Decrease | Mean Duration of Outbreak (Days) |
|-------------|----------------------------------|---------------------------------|------------------|-------------------------------|----------------------|-------------------------------|
| No-Tx control | 12                               | 3.58                            | 175.9            | 0.0%                          | —                    | 5.83                          |
| Pre-Tx control | 56                              | 5.96                            | 106.5            | 0.0%                          | 89.3%                | 9.78                          |
| Tx           | 56                               | 2.02 ($P = .07$, no-Tx)         | 364.6 ($P < .0001$, pre-Tx) | 46.4% ($P < .0001$, both controls) | 80.4% | 3.2 ($P = .02$, no-Tx) ($P < .0001$, pre-Tx) |

Abbreviation: Tx, treatment.

*Using Kaplan-Meier test.
Table 4. Diagnosis by a Physician Versus Self-Diagnosis.

|                          | Self-Diagnosis | Diagnosis by Physician | Statistics (Between Groups) |
|--------------------------|----------------|------------------------|----------------------------|
| Mean age at diagnosis (years) | 42             | 25                     | $P < .0001$                |
| Mean number of episodes  | 4.28 (pretreatment) | 6.76 (pretreatment) | $P = .01$                 |
|                          | 1.33 (treatment)  | 2.34 (treatment)       | $P = .12$                 |
| Mean duration of episodes| 9.83 (pretreatment) | 9.75 (pretreatment) | $P = .48$                 |
|                          | 4.28 (treatment)  | 2.70 (treatment)       | $P = .11$                 |
|                         | ($P = .0008$)    | ($P < .0001$)          |                           |
| Severity of symptoms; 1 is “very bad” and 7 is “not bad at all” | 3.89 (pretreatment) | 2.89 (pretreatment) | $P = .02$                 |
|                          | 6.17 (treatment)  | 5.89 (treatment)       | $P = .93$                 |
|                         | ($P < .0001$)    | ($P < .0001$)          |                           |
| Interference in daily life; 1 is “interfered all the time” and 7 is “did not interfere” | 3.70 (pretreatment) | 2.84 (pretreatment) | $P = .09$                 |
|                          | 6.50 (treatment)  | 5.80 (treatment)       | $P = .06$                 |
|                         | ($P = .0001$)    | ($P < .0001$)          |                           |
| Pain level; 1 is “very painful” and 7 is “not at all” | 3.56 (pretreatment) | 2.75 (pretreatment) | $P = .03$                 |
|                          | 5.89 (treatment)  | 6.03 (treatment)       | $P = .35$                 |
|                         | ($P < .0001$)    | ($P < .0001$)          |                           |

Figure 1. Kaplan-Meier plots of time to first oral herpes recurrence in the current study from participants with a history of at least 1 recurrence per year compared to (A) treatment versus pretreatment groups, (B) treatment versus no-treatment groups, (C) percentage of participants who have a certain number of outbreaks per year, treatment versus pretreatment groups.
recurrences, and 45 of 56 (80%) participants had a greater than 50% decrease in the recurrence rate.

Safety. There were no reports of adverse experiences following treatment with the herbal Gene-Eden-VIR/Novirin.

Gene-Eden-VIR/Novirin Versus Leading Drugs

The next section compares the herbal Gene-Eden-VIR/Novirin with ACV and VACV, the 2 leading drugs in the category. This study did not include what the FDA calls “concurrent” groups of ACV or VACV. According to the FDA guidelines, concurrent groups are test or control groups that “are chosen from the same population and treated concurrently.” Instead, we compared the effect of Gene-Eden-VIR/Novirin to “external” ACV and VACV groups.

We started creating the external ACV and VACV groups by searching PubMed (MEDLINE) for articles that reported the results of clinical studies that tested suppressive treatment with ACV and VACV in oral herpes. We used the following keywords: “herpes,” “labialis,” “acyclovir,” “valacyclovir,” and “valaciclovir.” We limited the search by the article type “randomized controlled trial,” and the English language. There were no date restrictions. We included all articles that tested the effect of oral treatment on the frequency of oral herpes outbreaks in immunocompetent adult men and nonpregnant women. The final list included 2 articles, a crossover study that tested the efficacy of a 4-month use of oral ACV in preventing HSL, and a placebo-controlled study that tested the effects of VACV 500 mg once daily for 16 weeks in preventing HSL.

Efficacy. Table 5 presents a comparison of the effects of the herbal Gene-Eden-VIR/Novirin and 400 mg of the ACV drug on time to first recurrence, percent reduction in mean number of recurrences, percent reduction in the number of individuals with recurrences, and the mean duration of outbreaks. Table 6 presents a comparison of the effect of Gene-Eden-VIR/Novirin and VACV 500 mg on time to first recurrence, number of recurrences per month, and percent of participants who are recurrence-free. All 7 measures showed that the herbal Gene-Eden-VIR/Novirin is superior to the ACV and VACV drugs in decreasing the number and duration of oral herpes outbreaks (Table 7).

Safety. Rooney et al reported adverse events in one patient who dropped out of the study due to headache and nausea. Baker et al observed 22 events (33% of patients) in the VACV group. The most common adverse event was headache, reported 5 times among 3 patients taking VACV.

In addition, several articles reported noncommon adverse events associated with these 2 drugs. Yavuz et al, as an example, describes a case of a 78-year-old woman, who had a normal baseline renal function, and no contributing possible nephrotoxic factors. This woman developed irreversible renal dysfunction after oral treatment with ACV. Becker et al
Table 7. Efficacy of GEV/N Versus ACV and VACV.

| End Point                          | Superiority of Efficacy | Source          |
|-----------------------------------|-------------------------|-----------------|
| Mean number of recurrences per month | GEV/N > VACV             | Baker et al³    |
| Percentage recurrence-free        | GEV/N > VACV             | Baker et al³    |
| Time to first recurrence          | GEV/N > ACV              | Rooney et al¹⁶  |
| Time to first recurrence          | GEV/N > VACV             | Baker et al³    |
| Percent reduction in recurrence   | GEV/N > ACV              | Rooney et al¹⁶  |
| Percent reduction in those with recurrences | GEV/N > ACV              | Rooney et al¹⁶  |
| Mean duration of lesion           | GEV/N > ACV              | Rooney et al¹⁶  |

Abbreviations: GEV/NV, herbal Gene-Eden-VIR/Novirin; ACV, acyclovir; VACV, valacyclovir.

leaves another case of a patient who developed rapidly progressive acute renal failure with concomitant changes in mental status after treatment with high-dose parenteral ACV.³⁰

Another serious, yet uncommon side effect of treatment with ACV is neurotoxicity that may lead to hallucinations, confusion, seizures, and obtundation.³¹

Le Cleach et al tested the effectiveness and safety of the 3 oral antiviral drugs, acyclovir, famciclovir, and valacyclovir, using a meta-analysis of 26 clinical studies.³² They found that the number of withdrawals due to harms was mentioned in only 8 studies, that is, 31% of the studies. In these studies, there were 14 withdrawals due to harms in the placebo or no-treatment groups, and 31 in the antiviral groups. In addition, only 4 studies, that is, 16% of the studies, mentioned safety data in the form of the total number of adverse events. In total, the 4 studies reported 331 adverse events in 561 (59%) participants in the antiviral treatment groups compared with 115 adverse events in 291 (40%) participants in the placebo or no-treatment groups. Lam et al analyzed a cohort of 76,269 patients who were treated with acyclovir, or valacyclovir, and 84,646 who were treated with famciclovir.³³ The results showed that 0.27% of the patients treated with ACV or VACV were hospitalized with acute kidney injury.

In our study, the participants reported no adverse events following treatment with the herbal Gene-Eden-VIR/Novirin.

Discussion

This study showed that suppressive or preventive treatment with the herbal Gene-Eden-VIR/Novirin reduced the frequency and duration of oral herpes outbreaks. The study also showed the existence of a duration of treatment effect, that is, longer treatment was associated with fewer outbreaks. Finally, the results showed that Gene-Eden-VIR/Novirin is safe.

In addition, the study showed that preventive treatment with Gene-Eden-VIR/Novirin is more effective in decreasing the number and duration of oral herpes outbreaks, and is safer than ACV and VACV. In other words, Gene-Eden-VIR/Novirin produced better clinical results than ACV and VACV, the 2 leading drugs in the category.

Our study has some methodological advantages. The reviewed ACV and VACV clinical studies had a single duration of treatment. Specifically, both Rooney et al and Baker et al tested their patients after 4 months of treatment. In contrast, our study included a wide range of durations, from 2 months to 36 months, with an average of 10.3 months. Moreover, 42.9% of the participants in our study used the Gene-Eden-VIR/Novirin between 6 and 18 months, and 19.6% for 18 months and over, which is a much longer duration of treatment. Moreover, since we used a range of durations, rather than a single duration, we were able to test for a duration of treatment effect.

Another methodological advantage is using 2 types of questions in gathering information about the outbreaks. The first type of questions asked the participants to count the number of outbreaks per year, and to count the number of days an outbreak lasted. The second type asked the participants to rate the frequency and duration of their outbreaks on a scale from 1 to 7. By comparing the answers to these questions, we were able to test for consistency in the participants’ reports.

Placebo controls are regarded as the gold standard in medical research. However, this is a retrospective chart review, and therefore does not include a placebo control. Instead, this study included 2 other controls recommended by the FDA: a no-treatment concurrent control and a baseline control, a type of external control.²⁸

The scientific method argues in favor of randomization, large sample sizes, independent verification by different laboratories, and so on, to even out unique characteristics found in any specific study, that is, to minimize the effect of confounding factors. This method considers the same result, observed under dissimilar settings, as reliable. This study showed that the effect of Gene-Eden-VIR/Novirin is significant when compared with a variety of untreated groups with dissimilar characteristics created by independent scientists. Therefore, we conclude that the observed effect is, most likely, real, that is, not an artifact of our specific treatment population, or its matching specific controls, and, therefore, with a strong external validity.

This study uses as data PROs. See a discussion on PROs in our previous study.²¹ A possible limitation of PROs is the subjective report of symptoms. To address this issue, we compared the participants’ reports of their symptoms with those reported in the literature. The comparison clearly showed that the symptoms reported by the participants, and those reported in the literature, overlap.

One possible confounding factor in this study is the possible relationship between current use of the product and positive results. To test for this confounding factor, we compared the results reported by current and past users. We found no statistically significant difference between the 2 groups. Therefore, we can conclude that this possible confounding factor did not bias the results in this study.

The participants in this study were either diagnosed by a physician or self-diagnosed. First, studies showed that self-reports in conditions such as herpes zoster, oral opportunistic
infections among HIV patients, fractures, cataract, and macular degeneration and physical capacity produced accurate diagnostic information. Second, the differences we observed between the 2 groups are consistent with those reported in the literature on help-seeking behavior. For instance, a study showed that health care–seeking men, with genital ulcer disease, were older, had more ulceration episodes in the past year, and were more likely to test seropositive for HSV-2. Another study reported that patients who delayed seeking health care assistance were younger, came from a lower socioeconomic state, had milder symptoms, and experienced less interference with their daily activities. Other studies showed that an increase in pain and disability increased the likelihood of seeking professional help. These findings are consistent with those reported in our study.

Conclusion
In summary, this study showed that suppressive or preventative treatment with the herbal Gene-Eden-VIR/Novirin safely reduced the frequency and duration of oral herpes outbreaks. The study also showed that the herbal Gene-Eden-VIR/Novirin is more effective and safer than ACV and VACV, the leading drugs in the category. Finally, as far as we know, this study is the first clinical study that shows that a natural treatment is better than the leading drugs in a major drug category.

Author Contributions
All authors contributed equally to this article.

Declaration of Conflicting Interests
The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

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Ethical Approval
The institutional review board that approved the study was Salus Institutional Review Board. Since the study was a retrospective chart review, the institutional review board approved a waiver of the requirement to obtain an informed consent from the participants under the exemption status of the federal regulations 45 CFR 46.101(b)(4).

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