Patients optimizing epilepsy management via an online community

The POEM Study

ABSTRACT

Objective: The study objective was to test whether engaging in an online patient community improves self-management and self-efficacy in veterans with epilepsy.

Methods: The study primary outcomes were validated questionnaires for self-management (Epilepsy Self-Management Scale [ESMS]) and self-efficacy (Epilepsy Self-Efficacy Scale [ESES]). Results were based on within-subject comparisons of pre- and postintervention survey responses of veterans with epilepsy engaging with the PatientsLikeMe platform for a period of at least 6 weeks. Analyses were based on both completer and intention-to-treat scenarios.

Results: Of 249 eligible participants enrolled, 92 individuals completed both surveys. Over 6 weeks, completers improved their epilepsy self-management (ESMS total score from 139.7 to 142.7, \( p = 0.02 \)) and epilepsy self-efficacy (ESES total score from 244.2 to 254.4, \( p = 0.02 \)) scores, with greatest impact on an information management subscale (ESMS–information management total score from 20.3 to 22.4, \( p < 0.001 \)). Results were similar in intention-to-treat analyses. Median number of logins, postings to forums, leaving profile comments, and sending private messages were more common in completers than noncompleters.

Conclusions: An internet-based psychosocial intervention was feasible to implement in the US veteran population and increased epilepsy self-management and self-efficacy scores. The greatest improvement was noted for information management behaviors. Patients with chronic conditions are increasingly encouraged to self-manage their condition, and digital communities have potential advantages, such as convenience, scalability to large populations, and building a community support network.

Classification of evidence: This study provides Class IV evidence that for patients with epilepsy, engaging in an online patient community improves self-management and self-efficacy.

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GLOSSARY

ESES = Epilepsy Self-Efficacy Scale; ESMS = Epilepsy Self-Management Scale; ICD-9 = International Classification of Diseases, Ninth Revision; ITT = intention-to-treat; PLM = PatientsLikeMe; POEM = Policy for Optimized Epilepsy Management; VISN = Veteran’s Integrated Service Network.

Many patients with epilepsy are not well-informed about their condition, and educational programs have been developed to increase health literacy and support self-management. The potential for digital educational programs has long been recognized; however, there remains limited evaluation of their effectiveness. One program, WebEase, used structured educational modules to promote enhanced patient self-efficacy on topics of medication use, sleep, and stress. However, these studies were limited by significant attrition, suggesting that highly structured educational interventions may not be optimally suited for some patients. An alternative approach to formal education is to simply connect patients with one another such that novice
patients can benefit from “expert patients” who can suggest practical advice on how to live well with their disease or have better conversations with health care professionals.\textsuperscript{13,14}

A patient-powered research network, PatientsLikeMe (PLM), provides an epilepsy-specific support system, as well as digital tools for tracking seizures, symptoms, medications, and comorbidities. An earlier study of the PLM platform suggested that patients experienced self-reported benefits including symptom management, medication adherence, and improved care, with the greatest predictor of benefits being the number of other patients with whom they shared a social connection.\textsuperscript{14}

However, this study was a cross-sectional retrospective survey without validated measures.

We sought to conduct a prospective study with a defined population of US veterans with epilepsy using validated survey measures of self-management and self-efficacy to determine the effectiveness of PLM. We hypothesized that an unstructured exposure to PLM would demonstrate statistically significant improvements in both self-management and self-efficacy.

**METHODS**

Primary research question. Does participation in an online patient community improve validated metrics of self-management and self-efficacy in veterans with epilepsy? This study provides Class IV evidence that 6 weeks of participation in a patient-driven, online community increases scoring on validated surveys for epilepsy self-management and self-efficacy. Total survey scores for both self-management ($p = 0.02$) and self-efficacy ($p = 0.02$) demonstrated significant improvements; a subscale for information management also demonstrated significant improvement ($p < 0.001$).

Standard protocol approvals, registrations, and patient consents. Approval was obtained from the University of California San Francisco Committee for Human Research Institutional Review Board and the San Francisco VA Medical Center Research & Development Committee. All participating patients gave informed consent through an online process. Prospective participants entered the study through a Web site, which initially required an e-mail address and unique username. They were presented with an electronic version of the approved informed consent document; an electronic “signature” of approval was required to enter the study, and this action was date- and time-stamped. The trial was registered at www.clinicaltrials.gov (NCT01762215).

Procedures. Potential participants were primarily identified using administrative ICD-9 diagnostic code data for epilepsy and seizures within the Veteran’s Integrated Service Network 21 (VISN 21), a region covering the western United States and Pacific, and were contacted by mail and/or phone. Patients were also identified through VA partner hospitals in VISN 17 (Texas) and VISN 6 (Virginia and North Carolina). For the identified cohorts, patient addresses, telephone numbers, and e-mail addresses, if available, were obtained. Per VA research policies, patients were mailed a “Dear Patient” letter inviting them to participate and a prestamped “opt-out” card to decline participation. If the opt-out card was not received back within 3 weeks, investigators attempted to follow up by phone.

Additional participants were recruited through paper and digital advertisement. The Web site www.poemstudy.com provided a study description, participation requirements, and online informed consent process. Eligibility was self-reported using online screening questions, and respondents who were not a US veteran, diagnosed with epilepsy (seizure disorder), or 18 years of age or older were excluded from participation.

Following informed consent, participants created a PLM account and performed baseline assessments. Participants who completed a follow-up assessment around 6 weeks later were given a $50 Amazon gift card.

**Intervention.** In addition to existing epilepsy-specific tracking tools, forums, and educational materials, POEM (Policy for Optimized Epilepsy Management) study participants had the option of receiving a special “veteran” designation, being part of an exclusive veteran discussion forum, and obtaining specialized information about epilepsy care within the VHA Epilepsy Center of Excellence system. Participants engaged with the Web site functions as much or as little as desired during the 6-week study period; participants received newsletter e-mails from PLM but were not contacted by study investigators.

On the basis of a small pilot study ($n = 40$), patient-reported outcomes on perceived self-management skills were assessed using 2 previously validated surveys, which served as co-primary outcome measures: the Epilepsy Self-Efficacy Scale (ESES) and Epilepsy Self-Management Scale (ESMS).\textsuperscript{11,12} The ESES is a 33-item questionnaire that measures a patient’s confidence in managing their own care related to their epilepsy diagnosis and contains 3 subscales: medication management (e.g., compliance), seizure management (e.g., managing triggers), and general management (e.g., attending appointments). An example ESES item reads “I can always take my seizure medication when I am away from home.” The survey responses are graded on an 11-point Likert scale, with 0 = I cannot do, 5 = moderately sure I can do, and 10 = sure I can do (range 0–330). The ESMS is a 38-item scale that assesses a patient’s frequency of self-management behaviors and includes 5 subscales: medication (e.g., side effect management), information (e.g., tracking seizures in a diary), safety (e.g., avoiding hazards), seizure (e.g., planning self-care), and lifestyle (e.g., sleep, diet). An example ESMS item reads “I write down how often I have seizures and when they occur.” This questionnaire is graded on a 5-point Likert scale, with 1 = never do and 5 = always do (range 38–190). Higher scores in both instruments indicate better self-management capabilities.

Using data from the pilot study, we determined a target sample size of approximately 250 participants to adequately test the study hypothesis, assuming an approximate 6-week study completion rate of 40%.

Participants also completed a user satisfaction survey,\textsuperscript{14} which assessed benefits that the platform may provide. Site usage was continually monitored throughout the study, including number of participant logins, posts to a public forum, profile comments to another user, and private messages sent to another user.

**Data analysis.** Baseline characteristics of the study population were summarized using standard descriptive statistics. Characteristics of completers and noncompleters were compared using t tests for continuous variables and χ² tests for categorical
variables. For all survey questions, participants were given the option of answering “not relevant” or “prefer not to answer,” and a variety of approaches were used to address missing data. First, analyses were performed using mean Likert scores for all items that were answered in a given scale, ignoring items that were not answered. Second, mean responses from items that were answered were used to calculate extrapolated full-scale and subscale scores. Third, sensitivity analyses were performed excluding participants who replied “not relevant” or “prefer not to answer” for more than 10% of the items in a given scale. Results were similar using all statistical approaches; therefore, only results using the extrapolated scores are reported in the results section. Our primary analytic approach was maximum likelihood random-effects regression with time as the independent variable. Both intention-to-treat (ITT) and completer analyses were performed. For ITT analyses, we took the conservative approach of assuming zero change for study participants who did not complete the follow-up survey. We decided a priori to use a \( p \) value of 0.05 for both of our co-primary outcomes, and to use Bonferroni-corrected \( p \) values of 0.01 for the 5 ESMS subscales and 0.0167 for the 3 ESES subscales. Finally, exploratory analyses were performed to identify baseline factors associated with change scores for each of our co-primary outcomes.

All analyses were performed using Stata/SE version 12.1 (StataCorp LP, College Station, TX).

RESULTS A total of 453 individuals registered, of whom 204 were excluded (14 opted out, 51 did not consent, 68 were not veterans, 22 did not have epilepsy, 28 did not complete the initial survey, and 21 were identified as duplicates or invalid) and 249 (55%) were enrolled (figure). Of 249 enrollees, 92 (37%) completed the follow-up survey and were included in completer analyses.

The mean age for the eligible participants was 50.2 years (SD 13.9), and approximately 80% were men (table 1). Most participants (75%) were non-Hispanic white, more than half were married, and approximately one-quarter had children (table 1). Nearly one-third reported their health status as fair or poor, and nearly two-thirds were not working for pay or were receiving disability income. Completers were slightly more likely than noncompleters to have received education beyond high school (83% vs 72%, \( p = 0.055 \)) although this was not statistically significant. There were no other demographic or baseline survey scoring differences between completers and noncompleters (table e-1 on the Neurology® Web site at Neurology.org).

We observed statistically significant improvement in both of our co-primary outcomes of the full ESMS (self-management) and ESES (self-efficacy) scores (table 2). ESMS total scores increased 1.1 points (\( p = 0.02 \)) in the ITT analyses and 2.9 points (\( p = 0.02 \)) in the completer analyses. Similarly, ESES total scores increased 3.6 points (\( p = 0.03 \)) in the ITT analyses and 10.2 points (\( p = 0.02 \)) in the completer analyses. The subscale with the greatest improvement was information management (ESMS-IM), which increased 0.9 points in the ITT analyses and 2.1 points in the completer analyses (both \( p < 0.001 \), table 3). The greatest ESES improvement was general...
Table 1 Change over time in co-primary outcomes (ESMS total and ESES total)*

<table>
<thead>
<tr>
<th>Method</th>
<th>No.</th>
<th>Preintervention, mean (SD)</th>
<th>Postintervention, mean (SD)</th>
<th>Change, mean (95% CI)</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>ESMS total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ITT</td>
<td>249</td>
<td>138.6 (15.9)</td>
<td>139.8 (15.6)</td>
<td>+1.2 (0.2, 2.1)</td>
<td>0.02*</td>
</tr>
<tr>
<td>Completer</td>
<td>92</td>
<td>139.8 (16.9)</td>
<td>142.7 (15.4)</td>
<td>+2.9 (0.4, 5.4)</td>
<td>0.02*</td>
</tr>
<tr>
<td>ESES total</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>ITT</td>
<td>248</td>
<td>237.3 (58.5)</td>
<td>240.9 (58.5)</td>
<td>+3.6 (0.3, 6.9)</td>
<td>0.03*</td>
</tr>
<tr>
<td>Completer</td>
<td>92</td>
<td>244.2 (46.8)</td>
<td>254.4 (44.7)</td>
<td>+10.2 (1.5, 19.0)</td>
<td>0.02*</td>
</tr>
</tbody>
</table>

Abbreviations: CI = confidence interval; ESES = Epilepsy Self Efficacy Scale; ESMS = Epilepsy Self-Management Scale; ITT = intention-to-treat.

*ITT analyses assume that those who did not complete the second survey experienced no change in scores. Mean change scores and p values based on maximum likelihood random-effects regression with time as the independent variable.

**Significance levels set a priori at p < 0.05 for both co-primary outcomes.

Table 2 Baseline characteristics of 249 veterans with epilepsy

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Mean ± SD or n (%)*</th>
</tr>
</thead>
<tbody>
<tr>
<td>Age, y</td>
<td>50.2 ± 13.9</td>
</tr>
<tr>
<td>Male</td>
<td>201 (80.7)</td>
</tr>
<tr>
<td>&gt;High school education</td>
<td>188 (75.8)</td>
</tr>
<tr>
<td>Race/ethnicity</td>
<td></td>
</tr>
<tr>
<td>Non-Hispanic white</td>
<td>187 (75.1)</td>
</tr>
<tr>
<td>Black/African American</td>
<td>20 (8.0)</td>
</tr>
<tr>
<td>Hispanic</td>
<td>17 (6.8)</td>
</tr>
<tr>
<td>Other</td>
<td>25 (10.0)</td>
</tr>
<tr>
<td>Marital status</td>
<td></td>
</tr>
<tr>
<td>Currently married</td>
<td>137 (56.2)</td>
</tr>
<tr>
<td>Widowed/divorced/separated</td>
<td>76 (31.1)</td>
</tr>
<tr>
<td>Never married</td>
<td>31 (12.7)</td>
</tr>
<tr>
<td>Have children</td>
<td>66 (26.5)</td>
</tr>
<tr>
<td>Work status</td>
<td></td>
</tr>
<tr>
<td>Full-time</td>
<td>47 (19.9)</td>
</tr>
<tr>
<td>Part-time</td>
<td>35 (14.8)</td>
</tr>
<tr>
<td>Not working for pay</td>
<td>154 (65.3)</td>
</tr>
<tr>
<td>Annual household income &lt;$25,000</td>
<td>77 (35.0)</td>
</tr>
<tr>
<td>Receive disability income</td>
<td>152 (63.6)</td>
</tr>
</tbody>
</table>

*Data missing as follows: education (n = 1), health status (n = 6), marital status (n = 5), work status (n = 13), household income (n = 29), and disability income (n = 10).

We found that veterans with epilepsy who enrolled in an online patient community reported statistically significant improvements in previously validated self-efficacy and self-management measures. The greatest improvement was seen in information management, which includes the use of a seizure diary, tracking side effects, and tools for medication adherence. This result potentially demonstrates that study participants believed that the availability of online tracking tools or social support mechanisms improved their self-management capabilities.

The study demographics were largely reflective of the general US veteran population. The mean age and sex were not unexpected, although we had anticipated that a younger cohort of patients, such as Veterans from Operation Iraqi Freedom and Operation Enduring Freedom, might be more attracted to the concept of a digital health platform. These results indicate that the platform was accepted and used by a broad cohort of patients. Furthermore, there was no demographic difference between study completers and noncompleters, indicating that the older patients who enrolled had similar experiences to the general US veteran population. The mean age and sex were not unexpected, although we had anticipated that a younger cohort of patients, such as Veterans from Operation Iraqi Freedom and Operation Enduring Freedom, might be more attracted to the concept of a digital health platform. These results indicate that the platform was accepted and used by a broad cohort of patients. Furthermore, there was no demographic difference between study completers and noncompleters, indicating that the older patients who enrolled had similar experiences to the general US veteran population.

DISCUSSION We found that veterans with epilepsy who enrolled in an online patient community reported statistically significant improvements in previously validated self-efficacy and self-management measures. The greatest improvement was seen in information management, which includes the use of a seizure diary, tracking side effects, and tools for medication adherence. This result potentially demonstrates that study participants believed that the availability of online tracking tools or social support mechanisms improved their self-management capabilities.

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It was not unexpected that study completers used the platform more throughout the study period, although primarily in a smaller group of users who were highly engaged. These observations are consistent with the previously described trend in online platforms that attract a small cohort of “super-users,” while average participants engage less frequently, in a more passive fashion.15 About half of our participants agreed that PLM gave them more or better control over their condition and helped them understand their seizures, similar to results seen in an earlier study in a nonveteran population.14 Although we did not formally assess the potential for harm, no patient complained of any adverse effects. Site usage was entirely voluntary throughout the study.

<table>
<thead>
<tr>
<th>Table 3</th>
<th>Change over time in ESMS subscales</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td>No.</td>
</tr>
<tr>
<td>ESMS-IM</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESMS-LM</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESMS-MM</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESMS-SM</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESMS-SEM</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
</tbody>
</table>

Abbreviations: CI = confidence interval; ESMS = Epilepsy Self-Management Scale; IM = information management; ITT = intention-to-treat; LM = lifestyle management; MM = medication management; SEM = seizure management; SM = safety management.

Mean change scores and p values based on maximum likelihood random-effects regression with time as the independent variable.

* Statistical significance for ESMS subscale scores was set at p < 0.01 a priori to account for multiple comparisons.

<table>
<thead>
<tr>
<th>Table 4</th>
<th>Change over time in ESES subscales</th>
</tr>
</thead>
<tbody>
<tr>
<td>Method</td>
<td>No.</td>
</tr>
<tr>
<td>ESES-MED</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESES-GEN</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
<tr>
<td>ESES-SEIZ</td>
<td>ITT</td>
</tr>
<tr>
<td></td>
<td>Completer</td>
</tr>
</tbody>
</table>

Abbreviations: CI = confidence interval; ESES = Epilepsy Self-Efficacy Scale; GEN = general self-efficacy; ITT = intention-to-treat; MED = medication self-efficacy; SEIZ = seizure management self-efficacy.

Mean change scores and p values based on maximum likelihood random-effects regression with time as the independent variable.

* Statistical significance for ESES subscale scores was set at 0.0167 a priori to account for multiple comparisons.
Although our exploratory analyses did not demonstrate a statistical association between the improved outcomes and engagement on the social elements of the platform, we speculate that these features are worthy of further evaluation. The value of traditional “offline” social support networks for patient self-management has been demonstrated in earlier studies for epilepsy and other chronic conditions. Psycho-social interventions that improve self-management have previously been demonstrated to improve the health of the chronically ill. Furthermore, social interactions have also been directly correlated with improved levels of self-efficacy in the epilepsy population. Because this study was powered for detecting changes in the primary outcome measures, we believe that it would be inappropriate to conclude that social interactions on the forum did not affect the improvements. In fact, this work supports further hypothesis-generation and dedicated research on the specific impact of a social network “dose effect” on health-related outcomes.

For both the total ESMS and ESES scores, the overall improvements were modest, with the ESMS significance being driven primarily by the information management subscale (table 3) and the ESES more by the general self-efficacy subscale (table 4). Information management is more aligned with the features of the PLM platform (tools for condition tracking), and general self-efficacy (overall confidence in health management) is more consistent with the expected benefit of participating in PLM. We also note that our results share similar magnitudes of change when compared with other, more directive and resource-intensive education and disease-management platforms that have utilized the ESMS and ESES measures.

With evidence of improved self-management capabilities, it is reasonable to hypothesize that this type of online intervention could positively affect other metrics of epilepsy health, such as health care utilization, morbidity, and even epilepsy-related mortality. For our study, we believed it was premature to examine these outcomes since a foundation of validated research did not yet exist. Based on our results, we think that it would be compelling to explore more objective metrics of epilepsy-related health, such as emergency services utilization, frequency of interactions with epilepsy care teams, and medication adherence. Indirect effects of improved psychosocial support, such as work productivity, absences from work or school, and measures of social integration, might also be interesting measures to investigate.

The study had several limitations including self-reported validation of diagnosis, although most participants were recruited through VA databases and automated detection methods were used to flag suspect registrations. The outcome measures were also self-reported, although they have previously been validated and used in multiple epilepsy studies including controlled trials. Future research into whether clinical outcomes correlate with self-reported behaviors following PLM intervention would be valuable.

The time period of 6 weeks may have been too brief to accurately gauge long-term changes in self-management behaviors. We chose this time period based on the feasibility of engaging patients in the pilot study and on similar studies that used a 6-week trial period. We aimed for a reasonable balance of exposure, considering that some patients might primarily benefit from an initial online experience while others might establish a longer-term connection through the platform.

We also note that the rate of second survey completion only reached 37%, despite the use of an incentive for full study completion; this might represent an enriched sample of those who benefited from the platform. However, this completion rate was not unexpected based on similar results from a small pilot study conducted in 2012; in fact, the sample size targets for the full protocol were based on an approximate completion rate of 40%. Furthermore, we note that other research studies and reviews in this domain have previously reported similar completion rates over the same time frame, highlighting the expected attrition that occurs with digital interventions. We have not yet conducted quantitative assessments of the reasons for this attrition, but that exercise would be an appropriate next course. We attempted to statistically control for attrition by assuming no benefit in those who did not complete the second survey for the ITT analysis.

Finally, we acknowledge that we did not include a control group in this investigation because any patient assigned to a “wait list” could access PLM independently outside of the study mechanism, thus exposing the control group to the intervention. Despite these limitations, we contend that this study provides foundational evidence for the potential benefits of online patient-driven communities and self-management platforms for epilepsy. Traditional medications and other technological interventions will remain essential elements of epilepsy treatment, but there is a growing recognition of the importance of psychosocial support, education, and self-management tools for patients. Despite this, many patients with epilepsy are not provided with adequate access or resources under our current health care ecosystem; as Mittan asserts, “the greatest problem among psychosocial treatment and educational programs was that only a handful of these interventions were ever put into general use.” He suggests that the economics of reimbursement and the professional time required to
conductor these interventions impedes widespread adoption. As a cost-effective and more time-efficient alternative, online networks and management platforms offer more ubiquitous accessibility and a larger support system than traditional methods.

If patient engagement is, as some claim, the “blockbuster pill of the 21st century,”*22 how will we support its development and implementation? If digital communities and online self-management tools do have the potential to improve clinical outcomes, then we have a shared responsibility to rethink our current health care ecosystem, focusing on new methods for incentivizing interventions that are proven to benefit patients.

AUTHOR CONTRIBUTIONS
John D. Hixson: corresponding author, study concept, design, and overall supervision. Deborah Barnes: author, data analysis and interpretation. Karen Parks: author, critical revision of the manuscript for content. Tracy Duquin: author, study concept, design, and critical revision of the manuscript. Stephanie Van Wicks: author, study coordination, data analysis and interpretation. Arianne Graham: author, study coordination, design, and data analysis. Paul Wicks: author, study concept, design, data analysis and interpretation.

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REFERENCES

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This Week’s Neurology® Podcast

Early EEG contributes to multimodal outcome prediction of postanoxic coma (see p. 137)

This podcast begins and closes with Dr. Robert Gross, Editor-in-Chief, briefly discussing highlighted articles from the July 14, 2015, issue of Neurology. In the second segment, Dr. Andy Schomer talks with Dr. Jeannette Hofmeijer about her paper on early EEG contributing to multimodal outcome prediction of postanoxic coma. Dr. Sarah Wesley reads the e-Pearl of the week about Susac syndrome. In the next part of the podcast, Dr. Alberto Espay focuses his interview with Dr. Daniel Lowenstein on his H. Houston Merritt Lecture at the AAN Annual Meeting on the topic of advances in the treatments for epilepsy.

Disclosures can be found at Neurology.org.

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Patients optimizing epilepsy management via an online community: The POEM Study
John D. Hixson, Deborah Barnes, Karen Parko, et al.

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