Editors’ Note: Association of Prestroke Metformin Use, Stroke Severity, and Thrombolysis Outcome

In "Association of prestroke metformin use, stroke severity, and thrombolysis outcome," Westphal et al. reviewed data from the European Thrombolysis in Ischemic Stroke Patients collaboration and compared stroke severity (NIH Stroke Scale), 3-month functional outcome (modified Rankin score), and mortality in 757 patients with type 2 diabetes who received metformin before stroke and 1,162 patients with type 2 diabetes who did not receive metformin before stroke. The authors reported that patients on metformin had less severe strokes, better functional outcome, and lower mortality and concluded that these findings suggest metformin has a protective effect in this patient population. Although he found the findings interesting, Dasheiff believed that this conclusion was premature because (1) correlation does not imply causation, (2) the statistical analysis did not fully compensate for the myriad confounders, and (3) there is no mention of other diabetic medications such as insulin use, which, in addition to the HbA1c and blood glucose may reflect disease severity. Westphal et al. agreed that causality cannot be inferred but emphasized the strengths of their study including the large sample size and their use of propensity score matching. The authors suggested their findings may precipitate additional studies on the relationship between metformin and outcome after stroke and increased use of metformin in patients with diabetes and vascular risk factors.

Ariane Lewis, MD, and Steven Galetta, MD
Neurology® 2021;96:500. doi:10.1212/WNL.0000000000011565

Reader Response: Association of Prestroke Metformin Use, Stroke Severity, and Thrombolysis Outcome

Richard Dasheiff (Plano, TX)

When I first read the article by Westphal et al., I was excited about the possible positive effects of metformin on stroke. I note that a one-point difference in NIHSS is not clinically significant, a one-point change in modified Rankin Score is somewhat significant, and 13% vs 18% death rate is more clinically significant. However, further critical analysis challenges this interpretation. First, the authors conclude that "[t]his suggests a protective effect of MET [metformin]." Neurology reviewers and editors should admonish authors about inferring causality from inappropriately designed studies. In addition, as the authors report, "The number of confounders in this study was 19; together with the treatment group (MET+/MET−), 20 independent variables resulted." This can only partially be compensated by their propensity score matching and multiple imputation. I would point out several other problems. For example, missing values included international normalized ratio (INR) on admission (29.7%). How could this be if everyone got IV thrombolysis? Per the article by Scheitz et al. cited in reference 8, "Laboratory measures obtained on hospital admission include glucose (mmol/L), blood cell counts (platelets, hemoglobin, and leucocytes), international normalized ratio and partial thromboplastin time/ Not all centers have to provide data on all variables but have given a commitment to add missing variables retrospectively, if considered relevant to answer a specific research question."
So, another question is whether everyone in this study had an INR before IV thrombolysis? Finally, perhaps the most important factor is the severity of the subject’s type 2 diabetes. Although blood glucose and HbA1c are surrogates for the patient’s clinical state, there is no assessment of other diabetes comedications. Patients usually move along a spectrum of diet control, oral hypoglycemics, and insulin. The composition of the MET group could be weighted toward insulin dependence, which would make them, on average, treated longer for their diabetes and having a more severe illness. That alone might account for their worse clinical outcomes, and not be a protective effect of metformin.


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Author Response: Association of Prestroke Metformin Use, Stroke Severity, and Thrombolysis Outcome

Laura P. Westphal (Zurich), Ulrike Held (Zurich), Stefan Engelter (Basel, Switzerland), and Susanne Wegener (Zurich)


We thank Dr. Dasheiff for his interest in our article.1 We agree that based on observational registry-based data, it is not straightforward to claim causality and impossible to account for all, including unknown, confounders. However, a thorough analysis based on propensity score–matched (PSM) sets of patients is mimicking a randomized experiment, and because the data set was very large, we were able to account for many known potential confounders simultaneously. It is common in real-life databases that there are some observations missing. In our study, we tackled this with a complex approach, in which we performed imputation and PSM repeatedly in multiple iterative steps. The observation that international normalized ratio (INR) values were incomplete is explained by the fact that standard operating procedures for intravenous thrombolysis (IVT) differed across centers. Some centers do not determine INR before IVT in patients with neither anticoagulation nor liver or coagulation disorder. We agree that duration of diabetes, nonpharmacological and other cotreatments may matter. This limitation was mentioned in the study. By accounting for the age difference between metformin (MET)-/MET+ groups with PSM, we intended to address this aspect. Our data may serve as an argument for patients and GPs to continue MET in patients with diabetes and vascular risk factors and spur planning of further trials that may prove the suggested protective effect of MET.


Copyright © 2021 American Academy of Neurology
Editors’ Note: Association of Guideline Publication and Delays to Treatment in Pediatric Status Epilepticus

In “Association of guideline publication and delays to treatment in pediatric status epilepticus,” Fernandez et al. reported that despite publication of evidence of delays in treatment of refractory status epilepticus (SE) in the Pediatric Status Epilepticus Research Group (pSERG) at the end of 2014, there was no difference in time to initiation of benzodiazepines, nonbenzodiazepine antiepileptic drugs (AEDs), and continuous infusions to patients with refractory SE in pSERG in the hospital between 2011–2014 and 2015–2019. The authors identified a number of proposed actions to overcome potential barriers to the timely initiation of AEDs. Albuja et al. commented that they improved the time to administration of second-line AEDs at their institution by creating a SE alert system to contact neurology, pharmacy, the rapid response team, and the bed manager simultaneously when an inpatient is suspected to be in SE. Amengual-Gual et al. applauded them for implementing this initiative. It is imperative to continue to identify quality improvement measures to identify SE and ensure that AEDs are given in a timely fashion to patients with SE because it becomes more resistant over time, which may lead to increased morbidity and mortality.1,2

Ariane Lewis, MD, and Steven Galetta, MD

Reader Response: Association of Guideline Publication and Delays to Treatment in Pediatric Status Epilepticus

Ana C. Albuja (Providence, RI), Meriem K. Bensalem-Owen (Lexington, KY), and Mauricio F. Villamar (Providence, RI)

Sánchez Fernández et al.1 reported that the publication of evidence on delays in treatment of pediatric refractory convulsive status epilepticus (rSE) was not associated with improvement in time to treatment of rSE (TTTSE). They propose interventions to reduce TTTSE. We would like to share a successful model that we implemented. In 2013, we identified delays in TTTSE in our institution.2 We saw no improvement in TTTSE after a dual intervention consisting of educational programs for healthcare professionals and the development of an electronic SE order set.3 This prompted us to develop an SE alert system that was modeled after our “code stroke” protocol.4 When an inpatient is suspected to have SE, the SE alert is activated by calling an operator who simultaneously pages the neurology and pharmacy house staff, the rapid response team, and the manager responsible for bed assignment. Use of this alert system led to considerable reduction in time to administration of second-line antiseizure medications (22.21 ± 3.44 minutes) compared with standard care (58.30 ± 6.72 minutes; p < 0.0001).4 Because many hospitals have alert protocols for management of acute stroke, SE alert systems could be replicated with local resources to facilitate timely management and coordinated care of SE in adults and children.
Author Response: Association of Guideline Publication and Delays to Treatment in Pediatric Status Epilepticus

Marta Amengual-Gual (Boston), Cristina Barcia Aguilar (Boston), and Tobias Loddenkemper (Boston)
Neurology® 2021;96:503. doi:10.1212/WNL.0000000000011560

We thank Drs. Albuja, Bensalem-Owen, and Villamar for their comment on our study1 and for their crucial contribution, highlighting the developments toward improving status epilepticus treatment in this pilot study in 19 adult patients during inpatient care at a single center2 and our publication. We are entirely aligned with these essential suggestions and applaud the authors for their crucial contribution. Similarly—and simultaneously to the authors’ accomplishment—pediatric investigators were able to improve inpatient treatment times in children at a single center.3 Most ongoing delays continue to occur in patients with intermittent refractory status epilepticus—related to difficulties in assessment whether convulsive status epilepticus persists—and in the outpatient setting.4 We are actively working on measures that can identify ongoing seizures—not only at a single center but also for larger populations in multicenter settings and in the outpatient setting.5 We agree that—among other approaches such as implementing novel technologies and quality improvement techniques—learning from previous stroke treatment efforts will play an important role.6 We are grateful for the chance to add this outlook to our current article and congratulate Drs. Albuja, Bensalem-Owen, and Villamar on this innovative milestone toward our shared goal of implementing improved status epilepticus care for our patients.


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Read Capitol Hill Report: Be Informed, Get Engaged

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Author disclosures are available upon request (journal@neurology.org).
In the article “Evaluation of Multiple Sclerosis Disability Outcome Measures Using Pooled Clinical Trial Data” by M. Goldman et al., the text contained some discrepancies within the analyses of the MSOAC data set included in the paper. The authors found that the discrepancies were present because of a coding error that produced inaccurate results in the Kaplan-Meier results. The authors regret the errors. The text below, provided by the authors, describes the corrected material:

From the Authors:

In our publication in Neurology, we analyzed a pooled dataset comprising 12,776 clinical trial participants that had been assembled by the Multiple Sclerosis Outcome Assessments Consortium (MSOAC) to evaluate 4 performance tests as proposed components of a multidimensional test battery. We reported measurement properties; construct, convergent, and known group validity; and longitudinal performance of the Timed 25-Foot Walk (T25FW), 9-Hole Peg Test (9HPT), Low Contrast Letter Acuity (LCLA), and Symbol Digit Modalities Test (SDMT) individually and when combined into a multidimensional test battery compared to the Expanded Disability Status Scale (EDSS) and Short-Form-36 Physical Component Summary. The placebo arm data in the MSOAC database were made publicly available to support research by investigators in the MS field. Following publication of our paper, a colleague, using the publicly available placebo data from MSOAC, contacted us after finding seemingly different results than reported in our paper. To better understand the potential discrepancy in results, we returned to our analysis. Subsequently, errors in statistical program (Statistical Analysis System, SAS) coding for some of our analyses were discovered. These errors resulted in incorrect progression rates for the T25FW and 9HPT, as well as the composite measures of progression based on any 1 or any 2 measures, which included T25FW and 9HPT. In addition, while the graph of progression by LCLA was correct, in the text, we incorrectly gave results for LCLA progression (and its kappa coefficient for agreement with EDSS) using a 20% threshold, instead of the 7-point threshold stated. Herein, we report corrected results.

<table>
<thead>
<tr>
<th>Performance test</th>
<th>Kaplan-Meier estimates of the percentage (95% CI) of trial participants with ≥1 confirmed disability progression over 24 mo</th>
<th>Corrected</th>
<th>EDSS$^e,f$</th>
</tr>
</thead>
<tbody>
<tr>
<td>T25FW$^a$</td>
<td>6.5</td>
<td>23.8 (22.8–24.8)</td>
<td>20.2 (19.2–21.1)</td>
</tr>
<tr>
<td>9HPT$^b$</td>
<td>2.9</td>
<td>8.6 (8.0–9.3)</td>
<td>20.2 (19.2–21.1)</td>
</tr>
<tr>
<td>LCLA$^c$</td>
<td>13.1</td>
<td>13.8 (12.7–14.9)</td>
<td>16.1 (15.1–17.2)</td>
</tr>
<tr>
<td>SDMT$^d$</td>
<td>15.0f</td>
<td>15.0f (13.4–16.7)</td>
<td>14.5 (12.9–16.2)</td>
</tr>
<tr>
<td>Worsening on ≥1 test</td>
<td>Not previously reported</td>
<td>41.5 (38.9–44.1)</td>
<td>11.5 (9.9–13.3)</td>
</tr>
<tr>
<td>Worsening on ≥2 tests</td>
<td>Not previously reported</td>
<td>12.9 (11.2–14.8)</td>
<td>11.5 (9.9–13.3)</td>
</tr>
</tbody>
</table>

Table 1 Sensitivity to Change of Performance Tests and EDSS

Abbreviations: $^a$ 20% threshold. $^b$ 20% threshold. $^c$ Seven-point threshold: the previous paper had erroneously reported results for a 20% threshold. $^d$ Four-point threshold. $^e$ Baseline score 0: 1.5-point increase, baseline score 1.0–5.5: 1-point increase, baseline score ≥6.0: 0.5-point increase. $^f$ The study populations available for each comparison differed, leading to differing proportions with 3-mo confirmed worsening on EDSS.

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To compare the sensitivity to change of the performance measures with EDSS, time from baseline to 3-month confirmed worsening over 24 months was analyzed (table 1). We previously reported that the progression rates were lower for T25FW and 9HPT compared to EDSS. In the corrected analyses, the progression rate remained somewhat lower for 9HPT compared to EDSS, while progression rates for T25FW, LCLA, and SDMT were similar to or higher than the EDSS (results for LCLA and SDMT were the same as before, as the programming error affected only T25FW and 9HPT). When the performance tests were combined into a multidimensional outcome measure, the proportion of participants with confirmed worsening on any 1 performance test was substantially greater than the proportion with confirmed worsening on EDSS. When confirmed worsening on 2 performance tests was required, sensitivity to disability progression was similar to that of EDSS. Association of the progression events defined by the performance tests was better correlated with those defined by the EDSS in the corrected analysis, though remained modest (table 2).

Based on the previously reported analyses, we concluded that the results supported the use of the T25FW, 9HPT, LCLA, and SDMT as study outcome measures, both individually or combined into a multidimensional test battery. These corrected results demonstrate better sensitivity to change for the T25FW and 9HPT than previously reported, and further support our original conclusion.

Reference

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Table 2 Associations of 3-Month Confirmed Progression Events Defined by Performance Tests and by Expanded Disability Status Scale

<table>
<thead>
<tr>
<th>Comparison</th>
<th>Cohen’s κ (95% CI) Previously reported</th>
<th>Corrected</th>
</tr>
</thead>
<tbody>
<tr>
<td>T25FW vs EDSS</td>
<td>0.02 (−0.00 to 0.03)</td>
<td>0.37 (0.35 to 0.39)</td>
</tr>
<tr>
<td>9HPT vs EDSS</td>
<td>0.00 (−0.01 to 0.01)</td>
<td>0.25 (0.22 to 0.27)</td>
</tr>
<tr>
<td>LCLA vs EDSS</td>
<td>0.11 (0.08 to 0.14)</td>
<td>0.10 (0.07 to 0.13)</td>
</tr>
<tr>
<td>SDMT vs EDSS</td>
<td>−0.02 (−0.06 to 0.02)</td>
<td>−0.02 (−0.06 to 0.02)</td>
</tr>
</tbody>
</table>

Abbreviations: 9HPT = 9-Hole Peg Test; CI = confidence interval; EDSS = Expanded Disability Status Scale; LCLA = Low-Contrast Letter Acuity; SDMT = Symbol Digit Modalities Test; T25FW = Timed 25-Foot Walk.

a 20% threshold.
b 20% threshold.
c Seven-point threshold (results had previously been given for a 20% threshold).
d Four-point threshold.
e Baseline score 0: 1.5-point increase, baseline score 1.0–5.5: 1-point increase, baseline score ≥6.0: 0.5-point increase.
f The previously reported association between progression events defined by 20% worsening on LCLA and EDSS was correct. For consistency, the association between progression events defined by 7-point worsening on LCLA and EDSS is reported here.