8-2-2013

Weight loss in persons with serious mental illness

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appropriateness of including our trial in the same group as the other randomized glutamine trials performed in the critical care setting, given differences in patient population, administration route, dosage, and approach to feeding.

Buijs et al. express several concerns that we believe are overstated. First, we attempted to optimize nutrition delivery through the application of evidence-informed strategies. Our success with feeding is probably consistent with the results in most intensive care units worldwide. The overall degree of imbalance in baseline characteristics is consistent with expected random variation and did not affect our conclusions. Our prespecified analysis plan, which followed best practices for factorial, randomized, controlled trials, did not adjust for the number of organ failures. Nevertheless, when we adjust for important baseline characteristics (including organ failures), the adjusted odds ratio of 28-day mortality for glutamine as compared with placebo remains consistent with our primary analysis, at 1.4 (95% confidence interval, 1.0 to 2.1; \( P = 0.05 \)). Finally, our secondary 6-month mortality results were based on survival analysis techniques that remain valid in the presence of random loss to follow-up. Besides, almost the entire treatment effect was observed within the first 30 days, when there was almost no loss to follow-up.

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Since publication of their article, the authors report no further potential conflict of interest.


DOI: 10.1056/NEJMc1306658

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**Weight Loss in Persons with Serious Mental Illness**

**TO THE EDITOR:** Daumit and colleagues (April 25 issue) describe the results of the Randomized Trial of Achieving Healthy Lifestyles in Psychiatric Rehabilitation (ACHIEVE), and they report that an intensive behavioral weight-loss intervention significantly reduced weight in adults with severe mental illness. This comprehensive intervention used existing rehabilitation-program staff in community-based psychiatric rehabilitation settings.

Unfortunately, the 2011 Centers for Medicare and Medicaid Services (CMS) policy for weight-loss counseling would not cover this effective intervention. The CMS policy covers only brief weight-loss counseling delivered by primary care physicians in primary care settings, not the comprehensive counseling by behavioral providers in a rehabilitation setting that Daumit and colleagues found effective. Current CMS policy will have little effect on health disparities because populations that are disproportionately affected by obesity often use the services of community-based organizations. Further, unlike the offices of most primary care doctors, community clinics employ behavioral counselors and typically have facilities (e.g., kitchens and recreation rooms) that are conducive to behavioral weight-loss programs.

The study by Daumit et al. and other community-based obesity trials provide support for the case for CMS to cover a wider range of settings and providers. This is not only consistent with the evidence but is also the most feasible approach to reducing weight in underserved populations.
TO THE EDITOR: Daumit et al. bring to light an important and often overlooked dynamic of treating patients with mental illness. Since one third of the U.S. population is classified as obese, and mentally ill patients are at increased risk for being overweight, it is critical that all health care professionals, specifically those working within the mental health field, integrate within the patient–physician relationship a discussion regarding healthy eating and lifestyle habits.

As mental health care providers struggle to do more with less funding, this study exemplifies what can be accomplished with relatively low overhead. The seamless integration of the ACHIEVE protocol into outpatient mental health centers and the use of the already established infrastructure within these centers increase the validity of this study.

Future studies of the effect of this behavioral intervention on coexisting conditions such as cardiovascular disease, diabetes, and dyslipidemia may reinforce the importance of addressing the benefits of a healthy lifestyle with patients who have mental illness.

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No potential conflict of interest relevant to this letter was reported.

THE AUTHORS REPLY: The implementation and dissemination of effective lifestyle interventions are critical to stem the epidemic of obesity and obesity-related conditions in persons with serious mental illness. However, incorporating weight-loss interventions into behavioral health settings will require financial and organizational resources that are unavailable in most community mental health settings. Community mental health programs are often under financial stress, and mental health, not physical health, is their first priority.

As Pagoto et al. state, the current CMS policy of providing reimbursement for weight counseling by primary care physicians would not be applicable to the ACHIEVE intervention, which was delivered by health coaches and staff. However, the federal government does provide two potential sources of funding for health promotion to organizations that serve persons with serious mental illness. The Patient Protection and Affordable Care Act (ACA) “health home” provision gives states additional federal funding for care management and the use of clinical information systems by Medicaid providers to enhance coordination of medical and behavioral health care. Eligible Medicaid recipients must have a serious and persistent mental health condition, two or more chronic mental or physical health conditions, or one mental or physical condition with a high risk of another. Primarily through ACA funding, the Substance Abuse and Mental Health Services Administration has awarded more than 60 grants to community-based agencies with the goal of building infrastructure to support integration of primary care services for persons with serious mental illness. These initiatives do not explicitly fund lifestyle interventions, although program resources could be used for incorporating health promotion into mental health care. Still, these two initiatives are modest in scope. We concur with Pagoto and colleagues that broader coverage of weight-management counseling is needed.

We also agree with Ackerman that incorporating healthy-lifestyle interventions into community mental health programs, particularly those such as psychiatric rehabilitation programs that mental health consumers attend frequently, is a wonderful but unrealized opportunity to leverage the existing infrastructure.

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DOI: 10.1056/NEJMc1306994
Weight-loss interventions could be incorporated into regular programming. In any case, implementing the ACHIEVE intervention into a broader array of community mental health settings would require resources to ensure the effective translation of our findings into practice.

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Since publication of their article, the authors report no further potential conflict of interest.


DOI: 10.1056/NEJMc1306994

Darbepoetin Alfa in Systolic Heart Failure

TO THE EDITOR: In their report on the Reduction of Events by Darbepoetin Alfa in Heart Failure (RED-HF) trial, Swedberg et al. (March 28 issue) conclude that the correction of anemia with the use of darbepoetin alfa in patients with heart failure does not reduce the rate of cardiovascular end points. This finding is in line with the results of the Trial to Reduce Cardiovascular Events with Aranesp Therapy (TREAT). However, it may be difficult to generalize such results to all forms of anemia in heart failure, especially to anemia caused by iron deficiency. As found by Toblli et al. and also shown in the Ferinject Assessment in Patients with Iron Deficiency and Chronic Heart Failure (FAIR-HF) trial, iron supplementation in patients with heart failure and iron deficiency reduces N-terminal pro–brain natriuretic peptide levels and improves symptoms and functional capacity. Since the FAIR-HF trial did not evaluate objective end points, the follow-up Iron in Congestive Heart Failure (ICHF) trial (ClinicalTrials.gov number, NCT01837082) will evaluate the effect of iron supplementation on left ventricular ejection fraction and renal function with the use of magnetic resonance imaging of the heart and radionuclide measurement.

It seems plausible to assume that taking into account the patient’s iron status in the therapeutic regimen may be superior to an approach focusing only on the hemoglobin level. Therefore, it is important that the authors stratify the primary end point according to the iron status at baseline, especially since iron deficiency was rigorously corrected during the trial.

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No potential conflict of interest relevant to this letter was reported.


DOI: 10.1056/NEJMc1306321

TO THE EDITOR: The RED-HF investigators found that darbepoetin alfa had no effect in patients with anemia and heart failure. But was the untreated group really untreated? Almost all the patients came into the program with a normal transferrin saturation of more than 20%; indeed, the median level was 24% (interquartile range, 19 to 31). This percentage is much above what is typically found in heart failure, in which the majority of patients have a transferrin saturation of less than 20%, suggesting iron deficiency. Thus, presumably the iron deficiency was corrected before they entered the study. In addition, once enrolled in the program, the great majority of patients received intravenous iron, oral iron,