Complete and Partial Responses of the TEMPI Syndrome to Bortezomib

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is one of the very few to have been published. This lack of data on the subject is extraordinary considering the number of fatalities involved. There is an urgent need to study the epidemiologic characteristics related to drowning and the nature of rescue efforts and the management of care after such disasters.

One of the greatest diagnostic challenges is the sudden, unexpected death of a swimmer with no significant medical history and in whom a conventional autopsy is negative. Cardiac channelopathies, particularly those that produce the long-QT syndrome, have been implicated, but they are considered rare. Nevertheless, such a diagnosis can be of considerable medicolegal and preventive importance. Tester and Ackerman and their colleagues have published studies that have gone a long way toward demonstrating that the cardiac channelopathies are not as rare as originally thought. Their letter is a reminder of the importance of awareness of the problem. The reason why certain mutations are associated with sudden death during swimming, rather than other forms of exercise, remains a mystery. As molecular autopsies become more widely available and, it is hoped, cheaper to perform, we may find some answers. In the meantime, those at risk should receive advice regarding supervised swimming and appropriate cardiac assessment and preventive interventions.

Joost J.L.M. Bierens, M.D., Ph.D. Society to Rescue People from Drowning Amsterdam, the Netherlands
Anthony J. Handley, M.D. Royal Life Saving Society UK Broom, United Kingdom
James P. Orlowski, M.D. Florida Hospital Tampa, FL

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


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TO THE EDITOR: We recently described the TEMPI syndrome (Aug. 4, 2011), which is characterized by the pentad of telangiectasias, elevated erythropoietin level and erythrocytosis, monoclonal gammopathy, perinephric fluid collections, and intrapulmonary shunting. One of the patients (Patient 2) had a dramatic response to treatment with the proteasome inhibitor bortezomib, and we hypothesized that the paraprotein may play a role in the pathophysiology of the TEMPI syndrome.

Patient 2, a 48-year-old woman, received a total of eight cycles of intravenous bortezomib (four doses of 1.3 mg per square meter of body-surface area per cycle). Her telangiectasias disappeared (Fig. 1, Panels A through D), her perinephric fluid collections disappeared (Fig. 1E and 1F), and her serum levels of erythropoietin decreased from 6400 mIU per milliliter to 19 mIU per milliliter. Levels of IgG kappa paraprotein became undetectable. Before treatment, she required a wheelchair and continuous supplemental oxygen; since the completion of treatment, her intrapulmonary shunting has resolved and she has recently resumed jogging. She remains in complete remission 13 months after receiving her last dose of bortezomib.

Patient 3, a 55-year-old woman, received six cycles of intravenous bortezomib. Her telangiectasias resolved, her serum erythropoietin level normalized from a peak of 507 mIU per milliliter, and her partial pressure of oxygen in arterial blood while breathing ambient air improved from 44 mm Hg to 70 mm Hg. Production of perinephric fluid, which drained into her abdomen after surgical fenestration of the renal capsule, decreased, as indicated by a decreasing requirement for large-volume paracentesis. However, after 4 months off treatment, levels of IgG kappa paraprotein began to increase, as did her serum erythropoietin level. Retreatment with bortezomib has been difficult because of the development of severe pulmonary hypertension.

Complete and Partial Responses of the TEMPI Syndrome to Bortezomib
Patient 1, a 52-year-old man, has received 9 weekly cycles of intravenous bortezomib. His serum level of erythropoietin has decreased from 5500 mIU per milliliter to 2500 mIU per milliliter; treatment is ongoing. The response to bortezomib of another patient with four features of the TEMPI syndrome (there was no intrapulmonary shunting), for whom there was limited follow-up, was recently described in the Journal.²

The efficacy of bortezomib treatment, as well
as the completely reversible nature of the symptoms, suggests that the abnormal plasma-cell clone and monoclonal gammopathy are the likely cause of the TEMPI syndrome. Efforts to identify the antigenic target of the paraprotein are under way. We suspect that there exist other patients with the TEMPI syndrome — as well as patients with other disorders — whose symptoms might be explained by a plasma-cell dyscrasia or underlying monoclonal gammopathy. We welcome any reader insights into this unusual syndrome.

Wilfried Schroyens, M.D., Ph.D.
Antwerp University Hospital
Antwerp, Belgium

Casey O’Connell, M.D.
University of Southern California
Los Angeles, CA

David B. Sykes, M.D., Ph.D.
Massachusetts General Hospital
Boston, MA
dbsykes@partners.org

Disclosure forms provided by the authors are available with the full text of this letter at NEJM.org.


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CORRECTIONS

Clinical and Biomarker Changes in Dominantly Inherited Alzheimer’s Disease (published Online First at NEJM.org on July 11, 2012; DOI: 10.1056/NEJMoa1202753). In Table 2 (page 5), the parenthetical unit of measure for Aβ42 in the CSF should have been pg/ml, rather than mg/ml. We regret the error. The article is correct at NEJM.org.

A 12-Month Phase 3 Study of Pasireotide in Cushing’s Disease (March 8, 2012;366:914-24). In the legend for Figure 1 (page 918), the first sentence should have ended, “50 patients had a substantial reduction (either normalization or ≥50% reduction from baseline) in urinary free cortisol level at month 6,” rather than “61 patients had a reduction of at least 50% in urinary free cortisol levels. . . .” In the first paragraph of the Discussion (page 922), the first sentence should have read, “This randomized, double-blind trial showed that 50 of 103 patients had a substantial reduction (either normalization or ≥50% reduction from baseline) in the urinary free cortisol levels at month 6 . . . ,” rather than “. . . 61 of 103 patients had a substantial reduction (≥50%) in the urinary free cortisol level. . . .” Also, several names were misspelled on page 2 of the Supplementary Appendix. The article is correct and the Supplementary Appendix has been replaced at NEJM.org.

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