Stress Hormones: Good or Bad for Posttraumatic Stress Disorder Risk?

Questions a new study in Biological Psychiatry

Philadelphia, PA, September 12, 2012 – Glucocorticoids, a group of hormones that includes cortisol, are considered stress hormones because their levels increase following stress. When their relationship to stress was first identified, it was shown that the release of cortisol prepared the body to cope with the physical demands of stress. Subsequently, high levels of cortisol were linked to depression and other stress-related disorders, giving rise to the hypothesis that high levels of cortisol on a long-term basis may impair the psychological capacity to cope with stress.

For this reason, drugs such as mifepristone that block glucocorticoid activity, called glucocorticoid receptor antagonists, have been tested as treatments for depression. But other recent data suggest that, in animal models and in humans, elevating glucocorticoid levels may reduce the development of posttraumatic stress disorder or PTSD.

This hypothesis is now supported by a new study in Biological Psychiatry. Using an animal model of PTSD, Rajnish Rao and colleagues demonstrate that elevated levels of glucocorticoids at the time of acute stress confers protection against anxiety-like behavior and the delayed enhancing effect of stress on synaptic connectivity in the basolateral amygdala.

“It seems, increasingly, that the ‘trauma’ in posttraumatic stress disorder is the impact of stress on brain structure and function,” commented Dr. John Krystal, Editor of Biological Psychiatry. “The study by Rao and colleagues provides evidence that glucocorticoids may have protective effects in their animal model that prevent from these changes in synaptic connectivity, potentially shedding light on protective effects of glucocorticoids described in relation to PTSD.”

Senior author Prof. Sumantra Chattarji from the National Centre for Biological Sciences in Bangalore, India, explained the reasoning behind their work: “First, this work was inspired by a puzzle - counterintuitive clinical reports - that individuals having lower levels of cortisol are more susceptible to developing PTSD and that cortisol treatment in turn reduces the cardinal symptoms of PTSD. Second, using a rodent model of acute stress, we were not only able to capture the essence of these clinical reports, but also identify a possible cellular mechanism in the amygdala, the emotional hub of the brain.”

Their results are consistent with clinical reports on the protective effects of glucocorticoids against the development of PTSD symptoms triggered by traumatic stress.

Two successive manipulations, both of which elevate corticosterone levels by themselves, together reset the number of synapses in the amygdala and restored anxiety behavior to normal levels in rats. Strikingly, these high and low numbers of synapses in the amygdala appear to be reliable predictors of high and low anxiety states respectively.

“With the increasing costs and suffering associated with PTSD victims, it is our hope that basic research of the kind reported in this study will help in developing new therapeutic strategies against this debilitating disorder,” concluded Chattarji.


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Notes for editors
Full text of the article is available to credentialed journalists upon request; contact Rhiannon Bugno at +1 214 648 0880 or Biol.Psych@utsouthwestern.edu. Journalists wishing to interview the authors may contact Sumantra Chattarji at +91 80 23666121 or shona@ncbs.res.in.

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