Childhood maltreatment and adult psychopathology: pathways to hypothalamic-pituitary-adrenal axis dysfunction

Maus-tratos na infância e psicopatologia do adulto: caminhos para a disfunção do eixo hipotálamo-pituitária-adrenal

Marcelo F. Mello,¹ Alvaro A. Faria,¹ Andrea F. Mello,¹ Linda L. Carpenter,² Audrey R. Tyrka,² Lawrence H. Price²

Abstract

Objective: The aim of this paper was to examine the relationship between childhood maltreatment and adult psychopathology, as reflected in hypothalamic-pituitary-adrenal axis dysfunction. Method: A selective review of the relevant literature was undertaken in order to identify key and illustrative research findings. Results: There is now a substantial body of preclinical and clinical evidence derived from a variety of experimental paradigms showing how early-life stress is related to hypothalamic-pituitary-adrenal axis function and psychological state in adulthood, and how that relationship can be modulated by other factors. Discussion: The risk for adult psychopathology and hypothalamic-pituitary-adrenal axis dysfunction is related to a complex interaction among multiple experiential factors, as well as to susceptibility genes that interact with those factors. Although acute hypothalamic-pituitary-adrenal axis responses to stress are generally adaptive, excessive responses can lead to deleterious effects. Early-life stress alters hypothalamic-pituitary-adrenal axis function and behavior, but the pattern of hypothalamic-pituitary-adrenal dysfunction and psychological outcome in adulthood reflect both the characteristics of the stressor and other modifying factors. Conclusion: Research to date has identified multiple determinants of the hypothalamic-pituitary-adrenal axis dysfunction seen in adults with a history of childhood maltreatment or other early-life stress. Further work is needed to establish whether hypothalamic-pituitary-adrenal axis abnormalities in this context can be used to develop risk endophenotypes for psychiatric and physical illnesses.

Descriptors: Cortisol; Risk factor; Psychopathology; Child abuse; Hypothalamus

Resumo

Objetivo: A meta deste artigo foi a de estudar as relações ente maus-tratos na infância e psicopatologia no adulto, como reflexo de uma disfunção do eixo hipotálamo-pituitária-adrenal. Método: Uma revisão seletiva da literatura relevante foi feita para identificar achados-chave e ilustrativos. Resultados: Existe atualmente um volume significativo de achados científicos pré-clínicos e clínicos derivados de paradigmas experimentais, que demonstram que o estresse precoce está relacionado à função do eixo hipotálamo-pituitária-adrenal e a estados psicológicos no indivíduo adulto, e como esta relação pode ser modulada por outros fatores. Discussão: O risco para o desenvolvimento de psicopatologia no adulto e disfunções do eixo hipotálamo-pituitária-adrenal está relacionado à complexa interação de múltiplos fatores vivenciais, assim como a genes que levam a uma susceptibilidade, que interagem com estes fatores. Embora as respostas agudas do eixo hipotálamo-pituitária-adrenal sejam geralmente adaptativas, as respostas excessivas podem levar a efeitos deletérios. O estresse precoce pode alterar a função do eixo hipotálamo-pituitária-adrenal assim como o comportamento, porém, o padrão da disfunção do eixo hipotálamo-pituitária-adrenal e a evolução psicológica na vida adulta refletem ambas as características do estressor e outros fatores modificadores. Conclusão: A pesquisa atual identificou múltiplos determinantes da disfunção do eixo hipotálamo-pituitária-adrenal encontrados em adultos com história de maus-tratos na infância ou outros estressores precoces. Trabalhos futuros são necessários para estabelecer se as anormalidades do eixo hipotálamo-pituitária-adrenal neste contexto podem ser usadas para o desenvolvimento de endofenótipos de risco para doenças físicas ou psiquiátricas.

Descritores: Cortisol; Fatores de risco; Psicopatologia; Maus-tratos infantis; Hipotálamo

Correspondence
Marcelo F. Mello
Rua Botucatu, 431
04023-061 São Paulo, SP, Brazil
Phone: (+55 11) 5082-2860
E-mail: mf-mello@uol.com.br

¹ Instituto PROVE (Programa de Atendimento e Pesquisa em Violência), Universidade Federal de São Paulo (UNIFESP), São Paulo (SP), Brazil

² Mood Disorders Research Program, Butler Hospital, Warren Alpert Medical School, Brown University, Providence (RI), USA

Introduction

For many years, empirical observations deriving from psychoanalytic, psychotherapeutic and general psychiatric clinical settings have suggested that childhood maltreatment has long-term consequences on mental health in adulthood. 1-3 Recently, more methodologically rigorous clinical and epidemiological studies have confirmed these earlier findings, while advances in basic and applied neurosciences have led to greater insights into possible mechanisms of pathogenesis.

Psychiatric consequences of childhood maltreatment

Childhood maltreatment is a major social problem. It is common and can result in serious physical injury or even death. Moreover, its psychological consequences can acutely affect a child's mental health well into adulthood.4 Four types of maltreatment are commonly recognized: physical abuse, sexual abuse, emotional (or psychological) abuse and neglect.

Childhood maltreatment is associated with a diverse range of psychiatric consequences. In children and adolescents, it increases the risk of behavioral problems, including internalizing (anxiety, depression) and externalizing (aggression, acting out) behavior. 5-9 Maltreated children have a moderately increased risk of depression in adolescence and adulthood (adjusted odds ratios ranging from 1.3 to 2.4), which will partly mirror the family context in which the maltreatment occured. 5,7,8,10-13

Because depression is common, about a quarter to a third of maltreated children will meet the criteria for major depression by their late 20's,8,14,15 thus representing a substantial public health burden. For many of the affected individuals, the onset of depression begins in childhood, hence the importance of focusing on early intervention before the symptoms of depression appear in the abused and neglected children. Depression is commonly associated with neglect and physical and sexual abuse, with no clear evidence of more specific effects of any particular type of maltreatment. Some investigators have shown a dose-response relationship, with depression more likely occurring with harsh or severe physical abuse than with less severe forms of maltreatment.8,16 Consistent evidence suggests that both physical and sexual abuse are associated with a doubling of the risk of attempted suicide in young people who are followed up on into their late 20's. For physical and sexual abuse, these effects persist after adjustment for confounding family and individual variables^{8,17} but, for neglect, these effects are mainly explained by family context.11

Other psychiatric disorders in adulthood that are frequently associated with childhood maltreatment include personality disorders, 18,19 alcohol abuse and dependence, 11 eating disorders, 20 anxiety disorders,²¹ and posttraumatic stress disorder (PTSD).²²

Neurobiological consequences of stress

The term stress has been widely used to denote an individual's response to environmental or psychosocial conditions which require change. Allostasis and allostatic load are more useful concepts for understanding the neurobiological consequences of stress. Allostasis involves the maintenance of stability (i.e., of homeostasis). The concept of allostatic load was proposed to refer to the physiological degradation of the individual as a result of repeated cycles of allostasis, as well as to the inefficient turning-on or shutting-off of responses to stressors. 23,24 The concepts of allostasis and allostatic load invoke a cascade of cause and effect that begins with primary stress mediators, such as catecholamines and cortisol, and leads to primary effects and then to secondary and tertiary outcomes.

The brain acts as an integrative center coordinating the behavioral, neuroendocrine and other neurobiological responses of the individual to environmental challenges and internal physiological needs (e.g., thirst, hunger, sleep-wake cycle). There are considerable individual differences in coping with such challenges based upon interacting genetic, developmental and experiential factors. Moreover, there can be an amplifying effect of genetic predisposition and early developmental events, such as childhood maltreatment, thus predisposing certain individuals throughout their lives to over-react physiologically and behaviorally to events. The capacity to adapt and maintain homeostasis (i.e., allostasis) in the face of challenges and the ability to turn adaptive responses on and off efficiently are vital to survival; at the same time, excessive demand for such adaptive responses can lead to cumulative effects over long time intervals.23,24

These cumulative effects, i.e., the allostatic load caused by forced adaptations to various psychosocial challenges and adverse environments are manifested as a structural and functional deterioration of the organism. Among the many factors that contribute to allostatic load are genes, early experiences and early development, as well as learned behaviors reflecting life-style choices with respect to diet, exercise, smoking, and drinking. All these factors influence the reactivity of the systems that produce the physiological stress mediators. As a result, allostatic load partially reflects comes to reflect a genetically or developmentallyprogrammed inefficiency in handling not only unusual stressors, but also the normal challenges of daily life (e.g., thirst, hunger, need for sleep), as well as the adverse physiological consequences of an unhealthy diet, sedentariness, excessive alcohol or smoking. 23,24

The HPA axis and psychopathology

Hyperactivity of the hypothalamic-pituitary-adrenal (HPA) axis in major depression is suggested by a large body of research, including basal and provoked measurements of plasma HPA axis hormone concentrations, imaging of pituitary and adrenal gland volume, cerebrospinal fluid (CSF) levels of corticotrophin releasing hormone (CRH), and post-mortem measures of brain CRH receptor binding and CRH messenger ribonucleic acid (mRNA) levels. 25,26 Indeed, hyperactivity of the HPA axis as a state marker for major depression is one of the most prominent findings in psychoneuroendocrinology.²⁷ ²⁹ Hyperactive CRH neurons, manifested as CRH hypersecretion and impaired efficacy of the glucocorticoid-mediated feedback are considered to be reliable hallmarks of the disturbed neuroendocrine regulation associated with, and perhaps causally related to depressive disorders.30

Allostatic load leads to alterations in hippocampal neuroplasticity, which is partially regulated by the brain-derived neurotrophic factor (BDNF). Hippocampal neurons subjected to an increased allostatic load show reductions in dendritic arborization and in BDNF expression, which could be one of the factors mediating the dendritic effects. The reduction in BDNF is partly mediated by excessive glucocorticoids, a characteristic feature of HPA axis hyperactivity, which can interfere with the normal transcriptional mechanisms that control BDNF expression. Decreased neurogenesis and increased rate of neuronal death and atrophy are also mediated by excessive glucocorticoids.31 Clinical studies are consistent with these findings: depressed patients have a significant reduction in gray matter in the hippocampus, anterior cingulate cortex and dorsomedial prefrontal cortex compared to controls. The reduction is more pronounced in unremitted patients than in remitted patients.32

However, the relationship between HPA function and emotional disturbances involving mood and anxiety is not entirely unambiguous.

Moreover, HPA axis dysfunction is not restricted to depressive illness, but has been linked to several other psychiatric and functional somatic disorders. A key question is whether such dysfunction operates as a pre-existing pathogenic risk factor, whether it reflects the ongoing pathogenesis of the illness, or whether it represents some combination of both.

HPA axis function in individuals with a history of childhood maltreatment

1. Studies on the pituitary–adrenal and autonomic responses to psychosocial stress

A number of studies have used structured laboratory test settings to evaluate responses to psychosocial stress. Among the most widely employed is the Trier Social Stress Test (TSST), which consists of a public speaking task and a mental arithmetic task, both standardized and conducted in front of a panel of judges. The TSST has been shown to reliably induce HPA axis and sympathetic activation.³⁸ Blood and/or saliva samples for the determination of plasma or salivary adrenocorticotropin (ACTH) and cortisol levels, as well as heart rate measures are obtained before, during, and after the stress induction.

In an important study, Heim et al. used the TSST to evaluate neuroendocrine and autonomic responses in four groups of women who had been carefully categorized according to the presence or absence of early-life abuse and current major depression, as follows: 1) with a history of early-life abuse and with current major depression; 2) with a history of early-life abuse and without current major depression; 3) without a history of early-life abuse and with current major depression; and 4) without a history of early-life abuse and without a history of psychiatric disorder (i.e., healthy controls).³⁹

Women with a history of childhood abuse, with or without current major depression, exhibited increased ACTH responses to stress compared with controls. The net ACTH response was more than 6-fold greater in abused women with current major depression than in controls. These women also demonstrated increased cortisol and heart rate responses to psychosocial stress. Abused women who were not currently depressed exhibited normal cortisol responses, despite their increased ACTH response, perhaps suggesting adrenal adaptation to central sensitization as a marker of resilience against depression after early stress. Depressed women without a history of abuse demonstrated normal neuroendocrine responses.

These findings, which are consistent with studies of early-life stress in laboratory animals, suggest that HPA axis and autonomic nervous system hyperreactivity, presumably due to CRH hypersecretion, may be a long-lasting consequence of childhood abuse in women and one which may constitute a risk factor for adult psychopathology.²⁷ Analyzing the same data using multiple regression techniques, Heim et al. showed that childhood maltreatment was the strongest predictor of ACTH responsiveness, followed by a number of abuse events, adulthood traumas and depression.⁴⁰ These findings indicate that a history of childhood abuse *per se* in women is related to increased HPA axis reactivity, which is further enhanced when additional trauma occurs in adulthood.²⁷

In contrast, a more recent study by Carpenter et al. found that men and women with a history of childhood maltreatment and no history of depression had *decreased* cortisol responses to the TSST.⁴¹ These findings are consistent with other recent studies showing attenuated cortisol responses to psychosocial or neurobiological challenge in adults with a history of maltreatment, as discussed further below.

2. Studies on the pituitary–adrenal and autonomic responses to pharmacological provocation tests

The CRH and ACTH stimulation tests, derived from classical neuroendocrinology, have also been used to explore HPA axis dysfunction related to psychopathology and associated risks factors. The former evaluates the responsiveness of the anterior pituitary to CRH, with diminished reactivity presumably reflecting downregulation of corticothrophs in response to chronically increased stimulation by the CRH-containing parvocellular neurons originating in the paraventricular nucleus (PVN) of the hypothalamic median eminence.⁴² The latter evaluates the responsiveness of the adrenal cortex to its major stimulating hormone, i.e., ACTH.⁴³

In their study of depressed and non-depressed women with and without a history of childhood maltreatment, Heim et al. administered both tests to the same subjects. 42,44 They found that abused women without depression exhibited increased ACTH responses to CRH, but both groups of depressed women (with and without childhood maltreatment) exhibited a blunted ACTH response to CRH, which is consistent with many previous studies of major depression. 45,46 Abused women without depression had a lower cortisol response than other groups after the ACTH stimulation test. 42 Laboratory studies on early-life stress using non-human primate models have demonstrated similar results. 47,48

Heim et al. hypothesized that their results could reflect both a sensitization of the pituitary and a counter-regulatory adaptation of the adrenal gland in abused women without current depression. As cortisol has inhibitory effects on the central CRH and noradrenergic systems, a relative decreased availability of cortisol, as a consequence of childhood trauma, might facilitate disinhibition of central stress responses. When subjected to even more stress, such women might then repeatedly hypersecrete CRH, eventually resulting in pituitary CRH receptor downregulation and symptoms of depression through CRH effects in extra-hypothalamic circuits.²⁷

However, caution is required when interpreting such findings, as underscored by Carpenter et al., who used multiple regression analysis in an effort to disentangle the effects of depression and early-life stress on CSF CRH in adults.⁴⁹ In their study of depressed and healthy control adults, perceived early-life stress was significantly correlated with concentrations of CSF CRH, but the presence of depression was not. However, the relationship between CSF CRH and early-life stress was complex: CSF CRH concentrations were negatively correlated (i.e., lower) with adversity in the perinatal and pre-teen years (ages 6-13 years), but positively correlated (i.e., higher) with stress in the pre-school years (ages 0-5 years). These findings were interpreted in light of considerable preclinical research in laboratory animals, demonstrating that timing of exposure to stressors in early life is critically important in determining their long-term effects on neurobiology and behavior.

3. Studies on glucocorticoid receptor (GR) sensitivity using pharmacological provocation tests

Basal hypocortisolism and hypersuppresion of cortisol after low doses of dexamethasone have been observed both in patients with PTSD and in women with a history of early-life stress and PTSD.^{36,50} These findings are believed to reflect increased GR sensitivity. The

combined dexamethasone/CRH (Dex/CRH) test was developed to allow for a more rigorous in vivo examination of GR function by increasing the sensitivity of the classical dexamethasone suppression test (DST).⁵¹ In the Dex/CRH test, CRH-induced escape of cortisol from suppression reflects impaired glucocorticoid-mediated feedback control of the HPA axis under conditions of increased hypothalamic drive.

Heim et al. used this approach to study HPA axis function in men with and without major depression and childhood abuse.52 Abused men demonstrated increased cortisol responses compared to non-abused men, regardless of the presence of major depression. Increased responses were associated with exposure to both sexual and physical abuse, and were correlated with the severity of abuse. Their results suggested that childhood maltreatment is associated with impaired glucocorticoid-mediated feedback control of the HPA axis under stimulated conditions.

Similar results have been reported for women with borderline personality disorder. Rinne et al. compared borderline personality disorder patients with and without sustained childhood abuse and comorbid PTSD or major depression with healthy control subjects. 53 The borderline patients who had been chronically abused had increased ACTH and cortisol responses to the Dex/CRH test; the presence of comorbid PTSD significantly attenuated the ACTH response.

Once again, however, specific factors relating to the nature of the abuse can markedly affect the results of such studies. Carpenter et al. recently reported their findings with the Dex/CRH test in a large sample of 230 adults without major Axis I disorders.54 Using a general linear models analysis, the authors tested for a large number of potentially confounding variables, including age, gender, education level, socioeconomic adversity in childhood, current depressive and anxiety symptoms, smoking, exogenous hormone use (for contraception or estrogen replacement) and types of maltreatment. They found that a history of self-reported childhood emotional abuse was independently and significantly associated with diminished cortisol response to the Dex/CRH test, whereas physical abuse, sexual abuse, emotional neglect and physical neglect had no significant independent effects.

Currently available studies indicate that much still remains to be learned about the different clinical factors that affect GR function in humans and how this process takes places. At the same time, recent preclinical research has shown, more clearly than ever, how GR function could be implicated in the pathogenesis of HPA axis dysfunction in clinical settings. For example, early adversity in rodents induces reduced expression of GR's at the epigenomic level by inducing DNA methylation at a promoter site of the GR gene. 55,56 It would be logical to expect that the impaired glucocorticoid effects resulting from this may contribute to increased stress reactivity and promote symptoms of depression.57

Long-term sequelae of early-life stress: moderating and mediating factors

As is evident from the preceding discussion, the ultimate effects of childhood maltreatment in terms of HPA axis function and psychiatric outcome in adulthood reflect a complex interplay of other moderating and mediating conditions. Elucidation of these factors and the nature of their relationships with adulthood outcomes is a major challenge.

For example, clinical experience and the preclinical animal literature suggest that early developmental events can have a major impact on subsequent psychological and neurobiological function.

We recently studied 42 premature infants born at 33.3 ± 1.9 weeks gestation of whom 15 were maintained in incubators for least 2 weeks and 27 were cared for with the kangaroo method, in which infant and mother consistently maintain skin-to-skin contact using specially-applied bandages (Mello et al., unpublished data). Groups did not differ with respect to gender, socioeconomic status, birth weight, gestational age or age at evaluation. We hypothesized that incubator care could serve as a naturalistic human model of maternal deprivation. Regression analysis showed that, at the age 6 months, infants who had been placed under incubator care/who had received care through an incubator had higher awakening salivary cortisol concentrations and were shorter compared to those receiving the kangaroo method. There were no differences in weight or cognitive and psychomotor development. These results are consistent with studies in laboratory animals showing that experimental early maternal deprivation increases HPA axis activity, which could in turn be a risk factor for psychiatric illness.

Similarly, late life events in an individual's personal history can also promote risk or resilience to the development of psychopathology. Tyrka et al., while studying ACTH and cortisol responses to the Dex/CRH test in healthy adults with or without a history of parental loss found that the former group had increased cortisol responses, an effect that was particularly robust in men.58 However, levels of parental care moderated the effect of loss: subjects with parental desertion and very low levels of care had attenuated cortisol responses. ACTH responses did not differ between groups.

Finally, it should be noted that moderating and mediating factors need not only be extrinsic events in an individual's history as they can also be intrinsic. Tyrka et al. used the TSST and the Dex/CRH test to evaluate the effects of temperament on HPA axis in healthy adults without significant psychopathology. 59,60 They found that in both psychosocial stress and pharmacological challenge tests, those with inhibited temperament (low novelty-seeking behavior) had higher cortisol responses compared to those with high noveltyseeking behavior.

The HPA axis and genetics

If HPA axis function can be affected by an individual characteristic such as temperament, which is, in itself, complexly reflective of both constitutional and early experiential factors, it should be obvious that genetic influences on the HPA axis are likely to be significant. Empirical demonstrations of such influences in humans have begun to appear. For example, Baghai et al. demonstrated that the angiotensin converting enzyme (ACE) insertion/deletion polymorphism significantly altered the cortisol response to the Dex/CRH test in patients with major depression, with insertion homozygotes showing higher cortisol responses than deletion homozygotes; differences were no longer evident after successful antidepressant treatment.61 Similarly, Wust et al. showed that glucocorticoid receptor gene polymorphisms determined the cortisol response to both provocative pharmacological testing and psychosocial stress, as measured by the TSST.62

Genetics and the long-term sequelae of early-life stress

Contemporary work on the interaction of genetics and stress has been profoundly influenced by the findings of Caspi et al., who studied a functional polymorphism in the promoter region of the serotonin transporter gene (5-HTTLPR) in a sample of 847 children who were followed prospectively from ages 3 to 26 years. 63 Individuals with one or two copies of the short allele of the polymorphism were more likely to develop depression and related symptoms in response to stressful life events than were individuals who were homozygous for the long allele. This study has been one of the most convincing empirical demonstrations to date of the gene-environment interaction in the pathogenesis of psychiatric illness. Kaufman et al. extended these findings to maltreated children, showing that low social support and the short allele of the 5-HTTLPR polymorphism interact with each other to incur in the greatest risk for depression.⁶⁴

Barr et al. addressed the interaction of genetics and early-life stress in modulating HPA axis function in a study of ACTH and cortisol responses to stress in rhesus macaque monkeys. ⁶⁵ They found that the genotype of the rh5-HTTLPR polymorphism (which is orthologous with the 5-HTTLPR polymorphism in humans) and rearing condition independently influenced HPA axis responses to stress. Moreover, rearing, rh5-HTTLPR genotype and sex interacted with one another: adverse rearing conditions and the rh5-HTTLPR short allele resulted in lower cortisol responses to stress in females, but not in males. More recently, Bradley et al. have extended this work to genes that directly control HPA axis function, demonstrating that reported childhood abuse interacts with single nucleotide polymorphisms (SNPs) in the CRH Type I receptor gene, thus allowing for the prediction of depressive symptoms in adulthood. ⁶⁶

Exciting preclinical studies have begun to elucidate the molecular basis of such findings, showing that environmental factors are actually capable of altering gene expression. This reflects recent advances in the field of *epigenetics*, the study of how differences in phenotype and gene expression can be caused by changes in chromosome or chromatin packaging (as opposed to by changes in the DNA sequence). There is emerging evidence that such changes can remain stable between cell divisions, and might show transgenerational inheritance.

In their review of the role of epigenetics in psychiatric disorders, Tsankova et al. illustrated the significance of such findings by providing the example of BDNF. This protein has been extensively implicated in the pathophysiology of depression and the mechanism of action of antidepressant treatments, and recent work has focused on the regulation of its gene, bdnf.67 Studies using an animal model of depression (chronic defeat) have shown that, in a controlled environment with no stress, bdnf chromatin demonstrates moderate histone H3 acetylation and no histone H3-K27 dimethylation. In this state, histone deacetylase 5 (HDAC5) represses unnecessary activation of BDNF and maintains chromatin balance. The stress of chronic defeat induces the dimethylation of histone H3-K27, resulting in a "closed" chromatin state at bdnf promoters and repression of bdnf transcript expression. Imipramine treatment after defeat stress down-regulates *Hdac5* gene expression and increases H3 acetylation, "reopening" the repressed chromatin state and reactivating transcriptional activity of the bdnf gene. In other words, an experiential factor (chronic defeat stress) alters BDNF function by physically modifying its gene, an effect which can, in turn, be reversed by antidepressant treatment.

Determinants of HPA axis function following early-life stress

To do a review, it is essential to recognize that, although the effects of early-life stress on the HPA axis are now well documented, such effects cannot be thought of in simplistic or bivariate terms. Rather, the understanding of these effects requires that a broad range of factors that can exert their influence via the stressor, the HPA axis, or the interaction of the two be systematically considered.

For example, above, we have considered evidence that *timing of the stressor* may be of importance, where attenuated HPA function appears following preschool stressors and enhanced function following preteen stress. ^{68,69} The *nature of the stressor* is relevant, with some studies suggesting increased HPA reactivity following early-life physical or sexual abuse or parental loss, but diminished reactivity following emotional abuse. At the same time, these effects can be *moderated or mediated by other factors* such as parental desertion or neglectful care (decreasing reactivity), inhibited temperament (increasing reactivity) or advancing age (decreasing reactivity). *Comorbid psychiatric conditions, recent stressors,* and *genetic vulnerability* also clearly impact HPA axis responsivity.

These factors are all, of course, extrinsic to the HPA axis itself. Divergent findings across and even within studies have underscored the fact that the way in which the "HPA axis function" is evaluated depends on which *specific parameters* are assessed. Such parameters include the *tissue* being assayed (e.g., brain, CSF, blood, saliva, lymphocytes), the *biochemical substance* being measured (e.g., mRNA, receptors, CRH, ACTH, cortisol), the *experimental method* employed (e.g., basal vs. provocative [e.g., TSST, Dex/CRH]), and the *index* used for analysis (e.g., peak change, recovery, area-under-the-curve [AUC]).

Conclusion

Risk for adult psychopathology is related to a complex interaction of multiple experiential factors: prenatal, perinatal, past environmental, and current environmental. Specific genes confer susceptibility to certain kinds of experiential factors that can lead to or modulate HPA axis dysfunction and psychopathological/ psychopathology risk. HPA axis response to stress can be thought of as a mirror of the organism's response to stress: acute responses are generally adaptive, but excessive or prolonged responses can lead to deleterious effects. These deleterious effects, which may not be clinically obvious, have the potential to serve as endophenotypic markers of susceptibility to disease. Early-life stress in rodents, nonhuman primates and humans (including childhood maltreatment) alters HPA axis function and behavior, but the specific pattern of HPA dysfunction and probably the nature of the adult psychopathology are reflective of both the characteristics of the stressor and an array of modifying factors.

Disclosures

Writing group member	Employment	Research grant ¹	Other research grant or medical continuous education ²	Speaker's honoraria	Ownership interest	Consultant/ Advisory Board	Other ³
Marcelo F. Mello	UNIFESP	FAPESP** CNPq**	Eli Lilly Janssen Cilag Servier Boeringher Inghelheim Bristol Myers Squibb	Eli Lilly* Roche* Pfizer*	-	-	Libbs* Torrent* Lundbeck*
Linda L. Carpenter	Warren Alpert Medical School	NIMH** US Department of the Interior** US Department of Defense**	AstraZeneca* Spears*	Cyberonics**	-	Cyberonics Neuronetics* Novartis* Wyeth*	-
Audrey R. Tyrka	Warren Alpert Medical School	NIMH** US Department of the Interior** US Department of Defense**	-	-	-	-	-
Alvaro A. Faria	UNIFESP	-	-	-	-	-	-
Andréa Feijó Mello	UNIFESP	CNPq*	Servier**	AstraZeneca**	-	-	Libbs*
Lawrence H. Price	Warren Alpert Medical School	NIMH** US Department of the Interior** US Department of Defense**	-	Jazz Pharmaceutical**	-	Gerson Lehrman** Wiley** Springer** Lundbeck**	-

^{*}Modest

For more information, see Instructions for authors.

References

- Cohen P, Brown J, Smaile E. Child abuse and neglect and the development of mental disorders in the general population. *Dev Psychopathol*. 2001;13(4):981-99.
- Allen B. An analysis of the impact of diverse forms of childhood psychological maltreatment on emotional adjustment in early adulthood. Child Maltreat. 2008;13(3):307-12.
- Collishaw S, Pickles A, Messer J, Rutter M, Shearer C, Maughan B. Resilience to adult psychopathology following childhood maltreatment: evidence from a community sample. *Child Abuse* Negl. 2007;31(3):211-29.
- Gilbert R, Widom CS, Browne K, Fergusson D, Webb E, Janson S. Burden and consequences of child maltreatment in high-income countries. *Lancet*. 2009;373(9657):68-81.
- Lansford JE, Dodge KA, Pettit GS, Bates JE, Crozier J, Kaplow J. A 12year prospective study of the long-term effects of early child physical maltreatment on psychological, behavioral, and academic problems in adolescence. *Arch Pediatr Adolesc Med*. 2002;156(8):824-30.
- Manly JT, Kim JE, Rogosch FA, Cicchetti D. Dimensions of child maltreatment and children's adjustment: contributions of developmental timing and subtype. Dev Psychopathol. 2001;13(4):759-82.
- Thornberry TP, Ireland TO, Smith CA. The importance of timing: the varying impact of childhood and adolescent maltreatment on multiple problem outcomes. *Dev Psychopathol*. 2001;13(4):957-79.
- Fergusson DM, Boden JM, Horwood LJ. Exposure to childhood sexual and physical abuse and adjustment in early adulthood. *Child Abuse* Negl. 2008;32(6):607-19.
- Herrenkohl EC, Herrenkohl RC, Rupert LJ, Egolf BP, Lutz JG. Risk factors for behavioral dysfunction: the relative impact of maltreatment, SES, physical health problems, cognitive ability, and quality of parentchild interaction. *Child Abuse Negl*. 1995;19(2):191-203.
- Banyard VL, Williams LM, Siegel JA. The long-term mental health consequences of child sexual abuse: an exploratory study of the impact of multiple traumas in a sample of women. *J Trauma Stress*. 2001:14(4):697-715.

- Brown J, Cohen P, Johnson JG, Smailes EM. Childhood abuse and neglect: specificity of effects on adolescent and young adult depression and suicidality. J Am Acad Child Adolesc Psychiatry. 1999;38(12):1490-6.
- Johnson JG, Cohen P, Smailes EM, Skodol AE, Brown J, Oldham JM. Childhood verbal abuse and risk for personality disorders during adolescence and early adulthood. *Compr Psychiatry*. 2001;42(1): 16-23.
- Noll JG, Trickett PK, Susman EJ, Putnam FW. Sleep disturbances and childhood sexual abuse. J Pediatr Psychol. 2006;31(5):469-80.
- Widom CS, White HR, Czaja SJ, Marmorstein NR. Long-term effects of child abuse and neglect on alcohol use and excessive drinking in middle adulthood. J Stud Alcohol Drugs. 2007;68(3):317-26.
- Widom CS, DuMont K, Czaja SJ. A prospective investigation of major depressive disorder and comorbidity in abused and neglected children grown up. Arch Gen Psychiatry. 2007;64(1):49-56.
- Andrews GC, Slade J, Issakidis T, Swanston H. Child sexual abuse. Comparative quantification of health risks. Geneva: WHO; 2004.
- Widom C. Childhood victimization: early adversity and subsequent psychopathology, in Adversity, stress, and psychopathology. In: Dohrenwend B, editor. New York: Oxford University Press; 1998. p.81-95.
- Tarquis N. Neurobiological hypothesis relating to connections between psychopathy and childhood maltreatment. *Encephale*. 2006;32(3 Pt 1):377-84.
- Tyrka AR, Wyche MC, Kelly MM, Price LH, Carpenter LL. Childhood maltreatment and adult personality disorder symptoms: influence of maltreatment type. *Psychiatry Res.* 2009;165(3):281-7.
- **20.** Smyth JM, Heron KE, Wonderlich SA, Crosby RD, Thompson KM. The influence of reported trauma and adverse events on eating disturbance in young adults. *Int J Eat Disord*. 2008;41(3):195-202.
- 21. Stein MB, Schork NJ, Gelernter J. Gene-by-environment (serotonin transporter and childhood maltreatment) interaction for anxiety sensitivity, an intermediate phenotype for anxiety disorders. *Neuropsychopharmacology*. 2008;33(2):312-9.

^{**} Significant

^{***} Significant. Amounts given to the author's and institution or to a colleague for research in which the author has participation, not directly to the author.

Note: UNIFESP = Universidade Federal de São Paulo; FAPESP = Fundação de Amparo à Pesquisa do Estado de São Paulo; CNPq = Conselho Nacional de Desenvolvimento Científico e Tecnológico; NIMH = National Institute of Mental Health.

- Grassi-Oliveira R, Stein LM. Childhood maltreatment associated with PTSD and emotional distress in low-income adults: the burden of neglect. *Child Abuse Negl.* 2008;32(12):1089-94.
- McEwen BS. Protective and damaging effects of stress mediators. N Engl J Med. 1998;338(3):171-9.
- 24. McEwen BS, Stellar E. Stress and the individual. Mechanisms leading to disease. *Arch Intern Med*. 1993;153(18):2093-101.
- Nemeroff CB. New vistas in neuropeptide research in neuropsychiatry: focus on corticotropin-releasing factor. Neuropsychopharmacology. 1992;6(2):69-75.
- Owens MJ, Nemeroff CB. Physiology and pharmacology of corticotropinreleasing factor. *Pharmacol Rev.* 1991;43(4):425-73.
- Heim C, Newport DJ, Mletzko T, Miller AH, Nemeroff CB. The link between childhood trauma and depression: insights from HPA axis studies in humans. *Psychoneuroendocrinology*. 2008;33(6):693-710.
- Arborelius L, Owens MJ, Plotsky PM, Nemeroff CB. The role of corticotropin-releasing factor in depression and anxiety disorders. J Endocrinol. 1999;160(1):1-12.
- Nestler EJ, Barrot M, DiLeone RJ, Eisch AJ, Gold SJ, Monteggia LM. Neurobiology of depression. *Neuron*. 2002;34(1):13-25.
- Mello AA, Mello MF, Carpenter LL, Price LH. Update on stress and depression: the role of the hypothalamic-pituitary-adrenal (HPA) axis. Rev Bras Psiguiatr. 2003;25(4):231-8.
- Berton O, Nestler EJ. New approaches to antidepressant drug discovery: beyond monoamines. Nat Rev Neurosci. 2006;7(2):137-51
- **32.** FrodI TS, Koutsouleris N, Bottlender R, Born C, Jager M, Scupin I, Reiser M, Maller HJ, Meisenzahl EM. Depression-related variation in brain morphology over 3 years: effects of stress? *Arch Gen Psychiatry*. 2008;65(10):1156-65.
- **33.** Posener JA, DeBattista C, Williams GH, Chmura Kraemer H, Kalehzan BM, Schatzberg AF. 24-Hour monitoring of cortisol and corticotropin secretion in psychotic and nonpsychotic major depression. *Arch Gen Psychiatry*. 2000;57(8):755-60.
- **34.** Oquendo MA, Echavarria G, Galfalvy HC, Grunebaum MF, Burke A, Barrera A, Cooper TB, Malone KM, John Mann J. Lower cortisol levels in depressed patients with comorbid post-traumatic stress disorder. *Neuropsychopharmacology*. 2003;28(3):591-8.
- **35.** Anisman H, Ravindran AV, Griffiths J, Merali Z.. Endocrine and cytokine correlates of major depression and dysthymia with typical or atypical features. *Mol Psychiatry*. 1999;4(2):182-8.
- Yehuda R, Yang RK, Buchsbaum MS, Golier JA. Alterations in cortisol negative feedback inhibition as examined using the ACTH response to cortisol administration in PTSD. *Psychoneuroendocrinology*. 2006;31(4):447-51.
- Ruiz J, Barbosa Neto J, Schoedl AF, Mello MF. Psychoneuroendocrinology of posttraumatic stress disorder. Rev Bras Psiquiatr. 2007;29(Supl1):7-12.
- **38.** Kirschbaum C, Pirke KM, Hellhammer DH. The 'Trier Social Stress Test'--a tool for investigating psychobiological stress responses in a laboratory setting. *Neuropsychobiology*. 1993;28(1-2):76-81.
- **39.** Heim C, Newport DJ, Heit S, Graham YP, Wilcox M, Bonsall R, Miller AH, Nemeroff CB. Pituitary-adrenal and autonomic responses to stress in women after sexual and physical abuse in childhood. *JAMA*. 2000;284(5):592-7.
- 40. Heim C, Newport DJ, Wagner D, Wilcox MM, Miller AH, Nemeroff CB. The role of early adverse experience and adulthood stress in the prediction of neuroendocrine stress reactivity in women: a multiple regression analysis. *Depress Anxiety*. 2002;15(3):117-25.
- Carpenter LL, Carvalho JP, Tyrka AR, Wier LM, Mello AF, Mello MF, Anderson GM, Wilkinson CW, Price LH. Decreased adrenocorticotropic hormone and cortisol responses to stress in healthy adults reporting significant childhood maltreatment. *Biol Psychiatry*. 2007;62(10):1080-7.
- **42**. Heim C, Newport DJ, Bonsall R, Miller AH, Nemeroff CB. Altered pituitary-adrenal axis responses to provocative challenge tests in adult surivors of childhood abuse. *Am J Psychiatry*. 2001;158(4): 575-81.
- **43**. Heim C, Ehlert U. Pharmakologische Provokationstests zur Einschatzung der neuroendokrinen Funktion. In: Kirschbaum C, Hellhammer DH, editor. *Enzyklopa die der Psychologie*. Hogrefe: Gottingen; 1999. p.307–59.

- **44.** Heim C, Nemeroff CB. The role of childhood trauma in the neurobiology of mood and anxiety disorders: preclinical and clinical studies. *Biol Psychiatry*. 2001;49(12):1023-39.
- **45.** Gold PW, Chrousos G, Kellner C, Post R, Roy A, Augerinos P, Schulte H, Oldfield E, Loriaux DL. Psychiatric implications of basic and clinical studies with corticotropin-releasing factor. *Am J Psychiatry*. 1984;141(5):619-27.
- Holsboer F, Antonijevic I, Murck H, Kuenzel H, Steiger A. Blunted corticotropin and normal cortisol response to human corticotropin-releasing factor in depression. N Engl J Med. 1984;311(17):1127.
- 47. Coplan JD, Andrews MW, Rosenblum LA, Owens MJ, Friedman S, Gorman JM, Nemeroff CB. Persistent elevations of cerebrospinal fluid concentrations of corticotropin-releasing factor in adult nonhuman primates exposed to early-life stressors: implications for the pathophysiology of mood and anxiety disorders. *Proc Natl Acad Sci U S A*. 1996;93(4):1619-23.
- **48.** Dettling AC, Feldon J, Pryce CR. Early deprivation and behavioral and physiological responses to social separation/novelty in the marmoset. *Pharmacol Biochem Behav*. 2002;73(1):259-69.
- Carpenter LL, Tyrka AR, McDougle CJ, Malison RT, Owens MJ, Nemeroff CB, Price LH. Cerebrospinal fluid corticotropin-releasing factor and perceived early-life stress in depressed patients and healthy control subjects. *Neuropsychopharmacology*. 2004;29(4): 777-84.
- **50.** Newport DJ, Heim C, Bonsall R, Miller AH, Nemeroff CB. Pituitary-adrenal responses to standard and low-dose dexamethasone suppression tests in adult survivors of child abuse. *Biol Psychiatry*. 2004;55(1):10-20.
- 51. Heuser I, Yassouridis A, Holsboer F. The combined dexamethasone/ CRH test: a refined laboratory test for psychiatric disorders. *J Psychiatr Res.* 1994;28(4):341-56.
- **52.** Heim C, Mletzko T, Purselle D, Musselman DL, Nemeroff CB. The dexamethasone/corticotropin-releasing factor test in men with major depression: role of childhood trauma. *Biol Psychiatry*. 2008;63(4):398-405.
- 53. Rinne T, de Kloet ER, Wouters L, Goekoop JG, DeRijk RH, van den Brink W. Hyperresponsiveness of hypothalamic-pituitary-adrenal axis to combined dexamethasone/corticotropin-releasing hormone challenge in female borderline personality disorder subjects with a history of sustained childhood abuse. *Biol Psychiatry*. 2002;52(11):1102-12.
- Carpenter LL, Tyrka AR, Ross NS, Khoury L, Anderson GM, Price LH. Effect of childhood emotional abuse and age on cortisol responsivity in adulthood. *Biol Psychiatry*. 2009;66(1):69-75.
- Weaver IC, Cervoni N, Champagne FA, D'Alessio AC, Sharma S, Seckl JR, Dymor S, Szyf M, Meaney MJ. Epigenetic programming by maternal behavior. *Nat Neurosci*. 2004;7(8):847-54.
- Meaney MJ, Szyf M. Environmental programming of stress responses through DNA methylation: life at the interface between a dynamic environment and a fixed genome. *Dialogues Clin Neurosci*. 2005;7(2):103-23.
- **57.** Pariante CM, Thomas SA, Lovestone S, Makoff A, Kerwin RW. Do antidepressants regulate how cortisol affects the brain? *Psychoneuroendocrinology*. 2004;29(4):423-47.
- Tyrka AR, Wier L, Price LH, Ross N, Anderson GM, Wilkinson CW, Carpenter LL. Childhood parental loss and adult hypothalamicpituitary-adrenal function. *Biol Psychiatry*. 2008;63(12): 1147-54.
- Tyrka AR, Mello AF, Mello MF, Gagne GG, Grover KE, Anderson GM, Price LH, Carpenter LL. Temperament and hypothalamic-pituitaryadrenal axis function in healthy adults. *Psychoneuroendocrinology*. 2006;31(9):1036-45.
- **60.** Tyrka AR, Wier LM, Anderson GM, Wilkinson CW, Price LH, Carpenter LL. Temperament and response to the Trier Social Stress Test. *Acta Psychiatr Scand*. 2007;115(5):395-402.
- 61. Baghai TC, Schule C, Zwanzger P, Minov C, Zill P, Ella R, Eser D, Oezer S, Bondy B, Rupprecht R. Hypothalamic-pituitary-adrenocortical axis dysregulation in patients with major depression is influenced by the insertion/deletion polymorphism in the angiotensin I-converting enzyme gene. *Neurosci Lett.* 2002;328(3):299-303.
- Wust S, Van Rossum EF, Federenko IS, Koper JW, Kumsta R, Hellhammer DH. Common polymorphisms in the glucocorticoid

- receptor gene are associated with adrenocortical responses to psychosocial stress. J Clin Endocrinol Metab. 2004;89(2): 565-73.
- 63. Caspi A, Sugden K, Moffitt TE, Taylor A, Craig IW, Harrington H, McClay J, Mill J, Martin J, Braithwaite A, Poulton R. Influence of life stress on depression: moderation by a polymorphism in the 5-HTT gene. Science. 2003;301(5631):386-9.
- Kaufman J, Yang BZ, Douglas-Palumberi H, Houshyar S, Lipschitz D, Krystal JH, Gelernter J. Social supports and serotonin transporter gene moderate depression in maltreated children. Proc Natl Acad Sci U S A. 2004;101(49):17316-21.
- Barr CS, Newman TK, Schwandt M, Shannon C, Dvoskin RL, Lindell SG, Taubman J, Thompson B, Champoux M, Lesch KP, Goldman D, Suomi SJ, Higley JD. Sexual dichotomy of an interaction between early adversity and the serotonin transporter gene promoter variant in rhesus macagues. Proc Natl Acad Sci U S A. 2004;101(33): 12358-63.
- Bradley RG, Binder EB, Epstein MP, Tang Y, Nair HP, Liu W, Gillespie CF, Berg T, Evces M, Newport DJ, Stowe ZN, Heim CM, Nemeroff CB, Schwartz A, Cubells JF, Ressler KJ. Influence of child abuse on adult depression: moderation by the corticotropin-releasing hormone receptor gene. Arch Gen Psychiatry. 2008;65(2):190-200.
- Tsankova N, Renthal W, Kumar A, Nestler EJ. Epigenetic regulation in psychiatric disorders. Nat Rev Neurosci. 2007;8(5):355-67.
- Maercker A, Michael T, Fehm L, Becker ES, Margraf J. Age of traumatisation as a predictor of post-traumatic stress disorder or major depression in young women. Br J Psychiatry. 2004;184:482-7.
- Tyrka AR, Wier LM, Anderson GM, Wilkinson CW, Price LH, Carpenter LL. Temperament and cortisol response to the DEX/CRH test in healthy adults. Neuropsychopharmacology. 2004;29(Suppl):S213.