ATTACHMENT TO CAREGIVERS AND TYPE 1 DIABETES IN CHILDREN
Navezanost na starše in sladkorna bolezen tipa 1 pri otrocih

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Received/Prispelo: Oct 28, 2014
Accepted/Sprejeto: Dec 1, 2014

ABSTRACT
Keywords: attachment, children, type 1 diabetes, etiology, stress reactivity

Attachment is a behavioral and physiological system, which enables individual's dynamic adaptation to its environment. Attachment develops in close interaction between an infant and his/her mother, plays an important role in the development of the infant's brain, and influences the quality of interpersonal relationships throughout life. Security of attachment is believed to influence individual response to stress, exposing insecurely organized individuals to deregulated autonomic nervous system and exaggerated hypothalamic-pituitary-adrenal activity, which, in turn, produces increased and prolonged exposure to stress-hormones. Such stress responses may have considerable implications for the development of diverse health-risk conditions, such as insulin resistance and hyperlipidemia, shown by numerous studies. Although the mechanisms are not yet fully understood, there is compelling evidence highlighting the role of psychological stress in the development of type 1 diabetes (T1D). One of the possible contributing factors for the development of T1D may be the influence of attachment security on individual stress reactivity. Thus, the suggestion is that insecurely attached individuals are more prone to experience increased and prolonged influence of stress hormones and other mechanisms causing pancreatic beta-cell destruction. The present paper opens with a short overview of the field of attachment in children, the principal attachment classifications and their historic development, describes the influence of attachment security on individual stress-reactivity and the role of the latter in the development of T1D. Following is a review of recent literature on the attachment in patients with T1D with a conclusion of a proposed role of attachment organization in the etiology of T1D.

IZVLEČEK
Ključne besede: navezanost, otroci, starši, sladkorna bolezen tipa 1, etiologija, stresna reaktivnost

Navezanost je vedenjski in fiziološki sistem, ki posamezniku omogoča dinamično prilagajanje na okolje. Navezanost se razvija pri sovplivu med dojenčkom in materjo, igra pomembno vlogo pri razvoju otrokovih možgan in vpliva na kvaliteto posameznikovih socialnih odnosov vse življenje. Varnost ali oblika navezanosti vpliva na posameznikov odziv na stres (stresna reaktivnost). Tako pride pri negotovo navezanih posameznikih do slabše reguliranega avtonomnega živčnega sistema in preizkušene reaktivnosti hipotalama-hipoftizna-suprarenalne osi, zaradi česar so ti v življenju pogostejše in dalj časa izpostavljeni delovanju stresnih hormonov. Tovrsten odziv na stres pa ima pomembno vlogo pri razvoju inzulinske rezistence, hiperlipidemije in drugih stanj, ki predstavljajo tveganje za zdravje. Čeprav natančni mehanizmi še niso znani, je vedno več dokazov, da psihološki stres pomembno prispева k razvoju sladkorne bolezni tipa 1 (SBT1). Eden od mehanizmov razvoja te bolezni bi lahko bil tudi vpliv oblike navezanosti na posameznikovo stresno reaktivnost. Tako so lahko negotovo navezani posamezniki pogosteje, dlje in v večji meri izpostavljeni delovanju stresnih hormonov, ki skupaj z drugimi dejavniki povzročajo uničenje beta celic trebušne slinavke. Ta prispevek prikaže najprej kratak pregled področja navezanosti pri otrocih, glavne oblike navezanosti in njihov zgodovinski razvoj, orisuje vpliv oblike navezanosti na posameznikovo stresno reaktivnost in vpliv te reaktivnosti na razvoj SBT1. Zaključujo se s predlogom o vlogi oblike navezanosti pri razvoju SBT1 pri otrocih.

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1 INTRODUCTION

John Bowlby conceptualized attachment as an evolutionary behavioral system of comparable importance to the systems guiding feeding and reproductive behaviors (1). Contemporary theories describe it as a behavioral and physiological system with a biological basis in the orbitofrontal system of the right hemisphere and its cortical and subcortical connections (2). The infant or child dynamically adapts to its environment to regulate safety and survival by ensuring proximity of the caregiver or attachment figure in times of distress, using attachment behaviors that increase in their complexity with age. These include constant knowledge of the attachment figure’s (an older person, most commonly the mother) whereabouts, using sounds to make her return when gone too far or fleeing to her if the distress or perceived danger rises to a level that is too high to endure on his own. Infant’s attachment behavioral system is adapted to a complementary behavioral system of the attachment figure, namely, the care-giving system. This results in a behavioral interplay between the infant and the caregiver, in which the infant leaves ‘the secure base’ provided by the caregiver to explore, and returns back to ‘the safe-haven’ when in distress (tired, ill, hungry, alarmed or just too far) to be calmed down and ready to leave again. Over time, the repeated interactions between the caregiver and the infant are internalized by the infant and captured into implicit memory as ‘internal working models’ (2). These early models are thought to guide expectations and behaviors in a largely unconscious way, and are thought to remain relatively stable across the lifespan, provided the family environment and the wider ecology remain stable as well (3).

2 PATTERNS OF ATTACHMENT

According to Bowlby’s and Piaget’s theory, and in line with the current knowledge on brain development, the attachment to one principal (mostly the mother) and one or more secondary attachment figures is formed between the age of 7 months and 2 years, when the object’s constancy (the knowledge that mother exists even when not present or within sight) has developed (2, 4, 5). Attachment continues to develop throughout childhood, depending on the stability or changes in the family environment (3, 4, 6). Bowlby’s early observations of the striking behaviours of children subjected to separation from, and loss of, their attachment figures were further supported by Ainsworth’s observational studies of individual differences in the quality of the interactions between infants and their mothers. What emerged was the critical importance of how sensitive and responsive mothers responded to their infants’ attachment bids. These early observations paved the way for the development of a structured laboratory assessment, the Strange Situation Procedure, as a way of accessing children’s internal working models of attachment relationships (6). These were thought to be reflected in their behavior in response to a series of separations from, and reunions with, their caregivers. Three organized attachment strategies were identified and, subsequently, Main and Solomon added a fourth attachment category, capturing attachment disorganization (7). These were:

1. Secure attachment: infant shows signs of missing mother during her absence, greets her actively, seeks comfort, settles and returns to play. The mother’s behavior at home was tender, careful holding with contingent face-to-face pacing of interactions, and with sensitivity to the infant’s signals in the first year of life.
2. Insecure resistant/ambivalent attachment: infant appears preoccupied with mother throughout procedure, is either markedly angry or markedly passive, alternately seeking and resisting the mother, fails to settle or return to play on reunion and continues to focus on the mother and cry. The mother’s behavior was inept in holding, noncontingent in face-to-face interaction and unpredictable, but not rejecting.
3. Insecure avoidant: infant focuses on the toys, does not cry during separation, actively avoids and ignores mother on reunion, moves away, turns away, leans away when picked up. The mothers of these infants rejected attachment behavior and were particularly averse to tactual contact.
4. Insecure disorganized/disoriented: infant displays disorganized or disoriented behaviors in the parent’s presence (freezing all movements, rocking on hands and feet, hands in air, rise and then fall prone at parent’s entrance). Mostly infants whose parents behaved towards them in abusive manner (7).

Attachment to caregivers develops disregarding the way the child is treated by them, even when the child is maltreated (8). Attachment behavior, although in a different form, becomes activated in adults as well, in the way of monitoring the availability of attachment figures (most commonly romantic partners, parents or close friends) and seeking them as ‘stronger and wiser’ in times of stress (9). The patterns of attachment are manifested in different ways across the life span, using different, but related, methodologies. The focus shifted from the study of individual differences in behavior to the study of internal working models as reflected in narratives (8).

3 ATTACHMENT SECURITY AND STRESS REACTIVITY

Research findings drawn from diverse perspectives converge in showing that an individual’s behavioral and
physiological response to a specific stressor is consistent throughout time and comparable in different species (10). The responses depend on the perception of the stressor as well as on the individual’s stress reactivity and coping. Individual differences in stress response can be observed in different domains, namely: physiology, cognitive function, subjective experience and behavior (10). Within the physiological domain, changes responding to stressful experience can be divided into cardiovascular responses (indicated by blood pressure and heart rate), driven by autonomic nervous system activity, and metabolic responses produced by the output of the glucocorticoid hormone cortisol from the adrenal cortex, driven by hypothalamic-pituitary-adrenal (HPA) axis activity (10). Although physiological responses serve to meet metabolic demands posed by the stressor, prolonged exposure to the HPA axis hyperactivity may lead to insulin resistance, hyperlipidemia and other conditions with increased risk for health disturbance (11). Cognitive responses to stress include systems important in learning, memory as well as attentional processes (12). Within the subjective domain, the perception of the stressor in humans is believed to be determined within the orbitofrontal system of the right hemisphere (2), the most common emotional experience being fear and apprehension (13). Experiences that pose a risk are thought to activate the individual’s attachment system and various physiological and behavioral mechanisms that are all directed towards homeostatically regulating the stress response, resulting in the deactivation of the attachment system (14). For example, the presence of a large dog will induce the feeling of fear in a small child and result in the child fleeing towards his/her mother, aiming to be lifted up away from the danger, held and comforted until the danger subsides. When this occurs and the child finds a secure base in the attachment figure, his/her physiological functions (increased heart rate, shiver) and the feeling of fear return to normal, and the child is able to continue with play.

Secure attachment is believed to provide resilience in the face of stress, ‘which is expressed in the capacity to flexibly regulate emotional states via autoregulation and interactive regulation’ (2), using internalized coping mechanisms and interaction with the attachment figure (2). Studies investigating stress reactivity in humans and animals have shown secure attachment to result in adaptive hypoactivity of the HPA axis, resulting in lower levels of stress hormones’ release under the influence of psychological stress (15). Conversely, unresponsive or neglectful parenting and child maltreatment have been associated with blunted early morning cortisol levels, no diurnal decrease in cortisol and an exaggerated cortisol response to stressful situations, although the mechanisms behind are complex (16).

Various studies have shown that securely attached children, under stressful circumstances, maintain low levels of cortisol when in the presence of attachment figure, while insecurely attached (especially disorganized) children’s cortisol responses were high (17-20).

4 THE ROLE OF STRESS IN THE DEVELOPMENT OF TYPE 1 DIABETES

The etiology of type 1 diabetes (T1D) has been extensively studied, with some clear findings (21-24), but many questions have yet to be addressed (25). The mechanisms implied in the etiology of T1D are viral infection, seasonality, rapid growth, psychological stress, and many others, yielding conclusions of multi-factor origins of the disorder (21-24).

One of the proposed hypotheses on etiology of T1D assumed the destruction of pancreatic beta-cells was a result of functional overload of the cells, caused by overfeeding, accelerated growth at puberty, low physical activity as well as psychological stress (with elevations in cortisol, other stress hormones and autonomic nervous system imbalance), which all resulted in elevated needs for insulin production (24).

There is evidence for the role of diabetes-related autoimmunity in the destruction of pancreatic beta-cells and development of T1D. However, not all subjects with diabetes-specific autoantibodies develop clinical syndrome, and not all patients with T1D produce such antibodies. Sepa et al. conducted a wide prospective population-based study on more than 5000 newborns, with a follow-up of 1845 subjects after 2.5 years. Their results showed associations of serious life-events (such as divorce or death in the family) with diabetes-specific autoantibody status of babies without clinical diabetes. The association remained in the follow up sample, where 12.5% of children exposed to divorce developed autoimmunity, as compared to 3.8% of the children without the experience of divorce (26).

Veronique Mead elaborated on a hypothesis, which included the role of stressful life-events in the critical period of brain development, namely, experience-dependent maturation. She argues that ‘disruptions in early bonding and attachment, including adverse events, such as traumatic stress, are capable of causing: (1) long-term imbalances in autonomic regulatory function and (2) relative dominance of sympathetic or parasympathetic activity.’ The proposed mechanisms would again program an individual’s autonomic nervous system and HPA axis, and expose the individual to higher insulin demands, as well as influence the immune system (27).
Taking all these into account, Ludvigsson proposed a unifying theory wherein various genetic, environmental and co-regulating factors were included as explanatory factors. In the proposed ‘beta-cell stress hypothesis,’ functional overload of the beta-cells as a result of multiple factors resulted in overexpression of specific antigens on the cells which mediated their destruction (28).

5 ATTACHMENT SECURITY AND DIABETES

There is only a scarcity of data on the attachment to caregivers or adults in close relationships in patients with diabetes. Ciechanowski et al. found dismissing attachment in the setting of poor patient-provider relationship to be associated with poorer treatment adherence in patients with type 1 and 2 diabetes. The study was performed using self-report measures of attachment security with 367 subjects (29). The same group then performed a cross-sectional study using self-report measures with 4095 adult primary care patients with diabetes type 1 or 2. Their results confirmed previous findings by showing that patients with dismissing attachment cooperated less well in their treatment than secure or preoccupied patients (lower levels of exercise, food care, diet and adherence to medications and higher rates of smoking), however, the patients with preoccupied attachment style were least likely to have worse metabolic control (not the securely attached). They concluded that insecure dismissing attachment is associated with worse diabetes outcomes, while secure and insecure preoccupied attachment predisposes to better outcomes (30).

Colton et al. studied disturbed eating behaviors in 106 girls with T1D in a one-year follow-up study using Children’s Eating Disorder Examination Interview. They concluded that a more disturbed attachment to mother was one of the predictors of a new onset of eating disturbances (31).

Rosenberg et al. performed a pilot study using self-report questionnaires on 31 families of adolescents with T1D, hypothesizing that attachment security to parents would be associated with metabolic control in adolescents. They were able to show only that maternal perceptions of more secure adolescents’ attachment, not adolescents’ reports, were associated with better glycemic control, although they concluded that the mechanism of the association was unclear (32).

Sepa et al. proposed that psychological stress could, via hormonal mechanisms, increase insulin resistance and trigger diabetes-related autoimmunity (33). They hypothesized that infants with diabetes-specific autoantibodies were more likely to have insecurely attached mothers than their antibody-negative peers. They interviewed 18 mothers of infants who were antibody-positive and 32 mothers of antibody-negative infants, using the Adult Attachment Interview (34).

Although failing to reach statistical significance, the proportion of children with insecurely attached mothers was substantially larger in the antibody-positive group (33% compared to 19% and 33% compared to 20% for two types of diabetes-specific autoantibodies) (33).

6 CONCLUSIONS

Although not yet fully understood, growing evidence points to the role of psychological stress in the development of T1D. One of the possible mechanisms could be mediated through the child’s attachment security and the influence of the attachment organization on stress reactivity, namely, the autonomous nervous system and the HPA axis reactivity. It is argued that insecurely attached, especially disorganized, children would be subjected to higher levels of stress hormones during exposure to stressful life-events. Insecure attachment and increased physiological stress reactivity would represent one of the factors contributing to the beta-cell overload, and, through the process of autoimmunity, peripheral tissue insulin resistance or some other as yet unknown process result in the destruction of beta-cells and clinical diabetes.

CONFLICTS OF INTEREST

The authors declare that no conflicts of interest exist.

FUNDING

None.

ETHICAL APPROVAL

Not required.

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