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Suboptimal bonding impairs hormonal, epigenetic and neuronal development in preterm infants, but these impairments can be reversed

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INTRODUCTION
Over the past 20 years, the incidence of preterm birth has increased and stabilised at an estimated 12% in the USA and 4-8% in high-income countries in Europe (1). Both the mortality rate and the incidence of major disabilities, such as cerebral palsy and mental retardation, have decreased due to improvements in perinatal health care (2). However, the long-term morbidity of these infants is still a serious health concern (1). Minor morbidities such as attention deficits, academic underachievement and hyperkinetic disorders occur frequently (1,2). Marcos et al. stated that both major and minor morbidities are probably induced by the fact that a preterm infant is exposed to unnatural stimuli that it would not be exposed to in the uterus (1). Indeed, it has been widely established that unnatural stimuli such as painful procedures, impaired nutrition and medical conditions such as hypoxia–ischaemia and infection can disturb homeostasis and hence normal organ development (3). Development might also be influenced by another unnatural stimulus that is an inevitable consequence of premature birth and that is altered parent–infant bonding (4).

In the medical literature, developmental consequences of altered parent–infant bonding are overshadowed by the consequences of many other unnatural stimuli. This could be explained by the fact that there is no objectively measurable physiological parameter for the degree of bonding. This creates a challenge for directly addressing the research question about whether suboptimal bonding alters development. On the other hand, bonding-related perinatal aspects, or in other words elements of bonding, have been studied separately. For example, there is an abundance of literature on the influence of perinatal maternal separation available (5), as well as literature on the physiological effects of the amount of perinatal exposure to licking and grooming. Licking and grooming is the animal analogue of caressing (6). In this review, we try to answer the previously formulated research question about whether suboptimal bonding alters development as accu-

Key notes
- In addition to hypoxia–ischaemia, infection and malnutrition, suboptimal bonding is one of the many unnatural stimuli that preterm infants are exposed to.
- The physiological consequences of suboptimal bonding are less frequently addressed in the literature than those of other threatening unnatural stimuli.
- This review found that suboptimal bonding significantly impaired hormonal, epigenetic and neuronal development, but these impairments could be reversed by bonding interventions investigated at neonatal intensive care units.

Abbreviations
BDNF, Brain-derived neurotrophic factor; EEG, Electro-encephalogram; HPA, Hypothalamic–pituitary–adrenal; KC, Kangaroo care; mRNA, Messenger-RNA; NICU, Neonatal intensive care unit; NMDA, N-methyl-D-aspartate.
Impairments caused by suboptimal bonding

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RESULTS

We extracted 23 bonding-related perinatal factors from the literature, including the following: licking, grooming, presence or absence (separation), feeding routines, sensitivity, received care, tactile stimulation and body heat. However, many papers used more general terms to capture bonding-related perinatal factors, such as early life experiences, early life adversity, NICU stay or NICU admission.

NICU-related early life experiences that altered physiology

NICU-related early life experiences were reviewed in a booklet edited by Browne and White called Clinics in Perinatology – Foundations in Developmental Care (13). They stated that, in the light of the remarkable plasticity during early development, the significant modifications of sensory experience that come with preterm birth and a necessary NICU admission are likely to ‘have a range of effects on the normal course of development’. However, they also stated that ‘investigators are a long way from understanding the particulars’. Another key point made in the booklet was that, compared with their term-born peers, preterm children have a much less developed behavioural repertoire, because of the immaturity of their brains. Important differences between the brains of term infants and preterm infants were highlighted by Browne et al. For example, brain volumes and brain region volumes are significantly different. Furthermore, the protein brain-derived neurotrophic factor (BDNF) appears to be lower in preterm infants. BDNF is a protein that supports neuronal survival and encourages the growth of new neurons. It thereby stimulates brain development and brain plasticity (13). Additionally, a significantly reduced capacity for, and duration of, long-term depression neuroplasticity has been found in preterm infants (15). Long-term depression is an immediate reduction in the efficacy of neuronal synapses after activation of those synapses. It plays an important role in the activity-dependent pruning or reinforcing changes of neuronal development. This means that when neurons fire frequently, their pathways are reinforced, for instance, their myelin sheets will thicken. However, when firing in certain neuronal pathways is scarce, neuronal apoptosis will follow (15). Research needs to be extended in order to investigate whether long-term potentiation neuroplasticity, the opposite of long-term depression neuroplasticity, is also reduced (13). Either way, whether or not that is the case, the demonstrated physiological differences in preterm infants make them extremely vulnerable. The fact that the physiological differences we report here are only some of the many known physiological differences caused by being born prematurely makes them even more vulnerable (13).

This instant presence of physiological differences due to premature birth is relevant to the research question about whether suboptimal bonding alters development. Apparently, preterm infants are already more vulnerable prior to the exposure to any additional unnatural stimulus, including unnatural bonding experiences. This was both a limitation and a strength of this review. It was a limitation because it impeded the assessment of the isolated effects of exposure to unnatural stimuli. On the other hand, it was a strength, because it suggested that, due to their vulnerability, the effects of all unnatural stimuli, including suboptimal bonding, could have major effects and they should therefore receive clinical attention.

In her review, Grunau (3) also drew attention to effects of certain unnatural stimuli. She reported on the long-term effects of stress caused by a NICU admission, specifically procedural pain-related stress. Grunau stated that procedural pain and stress in preterm infants are associated with abnormal brain development in the NICU, above and beyond other clinical risk factors. Grunau was also part of the research team that found that greater neonatal pain and stress exposure, measured by the number of skin breaking procedures, was associated with slower body and head growth. This was associated with altered cortical grey matter on diffusion tensor imaging at term. Greater exposure to procedural pain-related stress was associated with reduced development of white matter and subcortical grey matter (16). Furthermore, Grunau pointed out a longitudinal study from Doesburg et al. In that study, cumulative neonatal pain exposure in infants that were born prematurely was associated with changes in background cortical rhythmicity on magnetoencephalography measured later in childhood. The spectral structure of cortical oscillations expressed in power ratios among oscillations in different frequency ranges was assessed. Corrected for potential confounders, the found alterations in such spontaneous brain oscillations remained negatively correlated with visual-perceptual abilities at school age (17).

Anand et al. also reviewed the literature on alterations in adult brains due to repetitive neonatal pain and stress experiences. They focused more on maternal separation stress and the mechanisms that caused the effects of pain and stress. They proposed a two-way mechanism: hypoactivation of N-methyl-D-aspartate (NMDA) receptors is caused by maternal separation and sensory isolation, which
leads to increased apoptosis in multiple areas of the immature brain, while at the same time hyperactivation of NMDA receptors in other brain areas is caused by an exposure to repetitive pain (18). According to Anand et al., the latter results in excitotoxic damage to developing neurons. NMDA receptors are a subtype receptor for the most dominant excitatory transmitter system in the brain, the glutamate system. Thereby, NMDA receptors play a central role in the activity-dependent pruning or reinforcing of neurons (18). The complexity of the mechanism by which pain causes neuronal damage is not fully understood, but the fact that repetitive neonatal pain is harmful, and that there is involvement of the hypothesised mechanisms, is supported by various later studies (19,20). Furthermore, recent electro-encephalogram (EEG) findings suggest that the preterm neonatal brain is more sensitive to pain and that preterm infants are not very capable of distinguishing tactile from nociceptive stimulation, creating even more complexity for the preterm infant’s caretakers (3). The infant’s immaturity in mimicry, day and night rhythm and behavioural repertoire also imposes on the infant’s readiness to enter the caregiving dialogue, while on the parental side fear and the hindrance of the tubes, patches and the incubator disturb the caregiving dialogue (13). Programmes designed to recognise infant cues and to provide supportive care have, therefore, developed over the past decade, such as the Newborn Individualised Developmental Care and Assessment Program (NIDCAP) mainly established by Als.

The concept of developmental care is a framework for NICU staff that ensures that each individual infant is as stable, well organised and competent as possible (14). The infant’s own physiological and behavioural expressions are professionally assessed and seen as a reliable guide for caregivers to act upon. The family is seen as the primary caregiver and thus stimulated to be maximally involved in the caregiving (14). In a Cochrane review that included 627 preterm infants, however, no significant evidence was found that NIDCAP improved long-term neurodevelopmental or short-term medical outcomes (21). This is remarkable, as there is converging evidence that preterm infants are more developmentally vulnerable to parent–infant interaction, suggesting that the parents’ behaviours play a key role in their child’s neurodevelopment (3). Maximum parental involvement, for instance, due to a programme such as NIDCAP, should compensate for adverse clinical exposure and compromised early brain development (3,13). In some of the trials included in the Cochrane review such compensating effects were indeed found. Whereas the primary outcomes of reduction of mortality and severe disabilities showed no significant differences, shorter hospital stays, growth benefits and improved scores on neurodevelopmental scales at nine months of corrected age were reported (21). Nonetheless, based on all of the results and its cost-effectiveness, Ohlsson et al. concluded that no clear benefits of NIDCAP could be identified. The implementation of NIDCAP in its present form was therefore not recommended. However, Ohlsson et al. did recommend maximum involvement of the parents in caregiving, as they believed in the benefits of this (22).

This was supported by a cohort study by Grunau et al. that demonstrated that greater positive maternal interaction during painful procedures buffered the relationship between neonatal procedural pain exposure and poorer focused attention in very preterm infants at eight months of corrected age. More maternal interaction was also protective against infants’ internalising anxiety or depressive behaviours at 18 months of corrected age (3). So perhaps, compared to the current standard care in NICUs, NIDCAP is not enriching enough to lead to significant results. Over the past decades, even without having a programme such as NIDCAP implemented, most NICUs started focusing on individualised caregiving and also on involving the parents (23). A more easily measurable factor for caregiving enrichment would be an increase in parents visiting the NICU. Only very few studies have looked at this. They identified infrequent maternal visits as a risk factor for later psychological development in preterm infants (24). The above implies that both bonding-related and non-bonding-related early life experiences affect physiological development in a context-dependent way. Painful stimuli, maternal separation and abuse can be excitotoxic for neuronal development, whereas adequate caregiving can be protective.

Such a parental protection hypothesis is significantly supported by previous work from Sullivan’s research group. They addressed the question of why a child became attached to caregivers regardless of the quality of care they received. The key point that Sullivan stated was that the child’s brain was not an immature version of the adult brain. Its circuitry was designed to ensure attachment to caregivers. As proximity to a caregiver is necessary in order to survive, a child should therefore attach to any caregiver (25). In their studies, attachments to abusive caregivers resulted in dysregulation in neural networks, especially in disregulated amygdala function. The amygdala is a brain area within the limbic system, which is required for fear and avoidance learning in adult animals. By carefully manipulating the abuse-related attachment learning in rat pups, Sullivan et al. discovered that during a sensitive period early in life, the activation of the amygdala was blocked whenever there was maternal presence. Parental abuse did therefore not cause a fear and learn to avoid reaction when an infant’s mother was present. In their laboratory, pups formed attachments even when painful abuse was experienced. This infant experience resulted in later life depressive-like behaviour and altered amygdalas with suboptimal connectivity to the prefrontal cortex. Experiencing nonabusive painful procedures on the other hand did not result in amygdala changes, indicating that during attachment learning, pain was processed differently (25). More knowledge of this would be extremely valuable for NICUs, because even though painful experiences are inevitable, the negative effects of these might be minimised by high-quality parental caregiving. Moreover, the differences in physiological parameters due to prematurity itself are difficult to treat,
but optimising early life experiences might be easier. Indeed, noninvasive interventions targeted at parental care are relatively easy to implement, according to Walker et al. Moreover, these interventions might have a significant effect on the health outcome of the offspring, particularly in a vulnerable population of preterm babies (26).

Over the past few decades, researchers in the Braun laboratory have attempted to provide evidence for this parental protection theory. In one study, they used a maternal separation paradigm on the Octodon Degus trumpet-tailed rat species and investigated whether a mother’s voice during maternal separation could protect from separation-induced changes of brain function. Compared to the control group, the separation group showed an upregulation of NMDA receptor density in the prefrontal cortex, hippocampus and amygdala. In the third group, presentation of the maternal call during the separation period suppressed the NMDA receptor upregulation in all brain regions (27). In a 2014 review, Bock et al. (28) summarised their work, and the work of others, to provide an integrative view on this topic. They reported that ‘evidence is now accumulating that the specific and individual effects of early life experiences on the functional development of brain and behaviour emerge as a function of type, intensity, timing and the duration of the events’. In their studies, the effect of separation seemed to be delayed pruning or, in other words, a disturbance in the guidance for the neuronal cells to optimally specialise (28). However, the most frequently reported physiological disturbances in other studies examining early life experiences are those related to the development of the hypothalamic–pituitary–adrenal (HPA) axis, also known as the stress axis.

Mother–infant-related early life experiences and stress axis regulation

Due to the sudden transition from intra-uterine to extra-uterine life, early life is a very stressful period. Without a protection mechanism, the ongoing stress and the constantly hyperactive HPA axis would lead to chronic glucocorticoid exposure, which would cause cellular damage (26). A mechanism to protect organisms against this damage can, therefore, also be described as an adequate regulatory mechanism. However, the newborn infant’s own regulatory mechanisms are not sufficiently mature to accomplish this protection. The newborn infant needs coregulation for optimal protection and thus optimal development. Animal research has demonstrated that mothers are able to directly regulate the pup’s homeostasis, including his or her cortisol levels, through hidden regulators (29). These regulators are for instance sensory, motor, nutrient and thermal stimuli embedded in typical mother–infant interactions, such as licking, grooming and arched-back nursing. The parental influences on the infant’s physiology often cannot be seen directly, but become evident in periods of stress or separation for instance demonstrated by fluctuations in blood pressure or hormonal concentrations (9). That is because the infant’s stress regulatory system is, as previously stated, not yet mature enough to cope with separation or stress without parental support. The main axis within the elaborate stress regulatory system is the HPA axis, therefore also called the stress axis. Parent–infant interactions can thus exert an immediate coregulatory influence on the infant’s HPA axis (26,30). During the very stressful early life period, this coregulation results in suppressed cortisol activity, to protect the brain from otherwise constant glucocorticoid exposure. This period is called the stress hyporesponsive period.

During the stress hyporesponsive period, cortisol and the HPA axis are not the only systems that are affected. Other hormonal systems and, therefore, behavioural responses are affected in both the short-term and long-term. The immediate regulatory influence of typical mother–infant interactions is elaborate and necessary for optimal development (26,30). Inadequate mother–infant interaction thus impairs optimal development. Long episodes of maternal separation, lasting 180 minutes, changed corticotropin-releasing factor signalling pathways and glucocorticoid receptors involved in negative feedback over the HPA axis in adult rats. Separated rats exhibited visceral hypersensitivity and were more prone to stress-induced intestinal mucosal dysfunction (5). Additionally, Walker et al. reported that inflammatory reactions were reduced in pups reared by their mothers compared to artificially reared pups (26).

Does this mean that the more care a mother provides, the better her offspring? The short answer is no. For example, a maximal maternal stimulation protocol used by Walker et al. showed higher pain sensitivity when compared to less maternal stimulation. This suggests that this protocol was beyond the optimal degree of stimulation provided by the mother. Too much maternal stimulation provided to a compromised brain, such as a premature brain, might be detrimental (26). Indeed, not only the presence, but also the sensitivity and responsiveness from caregivers during the early life period appeared critical in maintaining the desirable, low cortisol concentrations (6). To define optimal care giving, the relationship between the rats’ HPA axis development and naturally occurring, individual differences in maternal care and thus maternal coregulation has frequently been evaluated (9). Liu et al. reported that the magnitude of the HPA response to stress in adult animals was strongly correlated with the degree of maternal licking and grooming of young pups (31). HPA axis alterations thus appeared context dependent. Indeed, both Rinaman et al. and Walker et al. reported that different lengths of maternal separation led to different HPA axis alterations (5,26). In older pups and humans, social companionship instead of separation has also been demonstrated to influence HPA responses, a process called social buffering (32). Similar to maternal presence, this also works through coregulation, enabled by receiving multiple sensory cues from the social companion. However, the very early life social buffering system, the stress hyporesponsive period, has unique features. It is adapted specifically to the mother–infant dyad (32). According to Rincón-Cortés et al., maternal care and stressors early in life jointly programme HPA axis responses and functioning in later life (30). The question is...
no longer whether early life experiences influence the HPA axis, but exactly how they do it (33).

In an attempt to answer that question, Strüber et al. proposed a two-pathway model in which they addressed more hormones than just the stress axis hormones. With this model, they provided an explanation for why both the HPA axis hyperfunction and hypofunction were demonstrated in animal and human studies that evaluated the later life effects of different early life experiences (33). In their critical review, Tang et al. also advocated a more complex reality than the more the better. Their theory was called maternal modulation (6). According to the authors, the role of the mother is multidimensional, including not only maternal behaviour quantity, but also the quality of the maternal behaviour as demonstrated by reliability and sensitivity and the mother’s capacity to regulate her own HPA function. Indeed, it is possible that infants who are fed by a stressed mother with a high cortisol concentration in her breastmilk have prematurely ended stress hyporesponsive periods (30). Interestingly, Tang et al. demonstrated, with results from animal and human studies, that the most influential factor for shaping key parameters of the infant’s regulatory system, such as the HPA axis and other hormonal systems, was the caregiving prior to, and immediately after, exposure to a stressful stimulus. When offspring are stressed, it is the reliability, not the quantity, of maternal care that determines whether or not stressful novelty exposure will have a positive impact on the plasticity of the HPA axis during later life (6). Tang et al. concluded that high-quality parenting following negative life events could enhance positive adaptation during subsequent negative life events. This could also be true for high-quality parenting following negative events during a NICU stay. From these results, it can be deduced that differences in maternal care and early life experiences set a cascade of physiological events in motion, impacting on the essence of human beings throughout the rest of their lives.

Mother–infant-related early life experiences altering genetics

Research has indeed demonstrated that the HPA axis is not the only physiological system that is altered by early life experiences and bonding-related experiences in particular. A growing body of neuroscience research has argued convincingly that an individual’s early life experiences, or early life environment, together with an individual’s genotype, influence his entire health and behavioural phenotype (34). Over the past few decades, an explanation for this has been discovered. Environmental factors can alter gene expression, without altering the genetic code itself. Instead, the frequency with which the genes are transcribed is altered. The transcription of the genome is regulated by intracellular molecules. The concentrations of these molecules are influenced by sensorial information about environmental factors. Together, those molecules and the genome itself are called the epigenome (35).

Epigenetic changes occur immediately after conception, providing a mechanism by which early caregiving environ-
alter oxytocin and vasopressin-regulated social behaviours (39). She incorporated seven different research paradigms in her review: maternal separation, early stress, single mothers, neonatal handling, communal nursing, high versus low licking and grooming and the manipulation 0 paradigm, a paradigm with no extra neonatal handling or manipulation, just weekly cage changes, while pups are transported in plastic cups. She concluded that all the paradigms had been shown to induce changes in the oxytocin and vasopressin systems. Therefore, those systems have shown a high level of plasticity as a function of the early life social environment. Currently, most authors deem that the question of whether or not early life experience has the ability to lead to long-term changes in oxytocin and vasopressin systems is undisputed. Instead, they consider the process and directionality by which experiences affect those and other systems (39,40). Oxytocin systems are highly connected to for instance serotonin, noradrenaline and dopamine pathways.

Dopamine pathways are important for experiencing rewards. When neurohormonal circuitry functions adequately, sociality is experienced as rewarding. Studies have indicated that early life experiences have long-term programming effects on these pathways as well. In these studies, the long-term effects were defined as effects still present in adulthood (42). The direction of the changes was again complicated, as the influences were brain region related and often gender dependent (42). In addition to the dopamine system, influences of maternal care and early life experiences on the reproductive system have also been studied. These studies were reviewed by Toufexis et al. Their review demonstrated that the reproductive system was mainly investigated by examining differences in the functioning of the hypothalamic–pituitary–gonadal axis. Within this axis, alterations in the expression of sex hormone receptors such as the oestrogen receptor-α were reported (43). This hormone regulates sexual behaviour and the sexual strategy used by females. By influencing the sexual behaviour of the offspring, the effects from differences in maternal care become transgenerational (43).

Camozzato et al. (44) used a different approach to assess the hypothalamic–pituitary–gonadal axis. They explored the development of a brain area important to the control of ovulation, the medial pre-optic area. Using a neonatal handling paradigm, they found an approximate reduction percentage of 50% in the number of neuronal cells in that area at postnatal day 11 and still at day 90. A compensatory neuronal cell size increase was not found. On the contrary, the neonatal handling procedure decreased cell size. This study strengthened the conclusion of an early life environmental effect on the hypothalamic–pituitary–gonadal axis by focusing on alterations in neuronal development.

Mother–infant-related early life experiences that alter neuronal development

Many studies have focused on altered neuronal development caused by different early life experiences. Some studies focused on specific brain areas, such as the medial pre-optic area. Other studies examined larger regions or even the entire brain, such as the study by Sarro et al. (45). They explored whether the cortical activity of rat pups was directly influenced by interactions with the mother. They recorded spontaneous neocortical potentials in freely behaving infant rats during natural interactions with their mother on postnatal days 12–19. Maternal absence increased cortical desynchrony. According to the authors, cortical desynchronisation was not necessarily negative, ‘it is an activity pattern conducive to neural plasticity and information transfer across brain regions’. Removing littermates induced further desynchronisation. The authors stated that ‘the results uncover that the mere presence of the mother provides an immediate impact on the state of cortical neural activity’. (45) The fact that the mere presence of a mother can be comforting to a child could be of value for families during a NICU stay. Furthermore, Sarro et al. found that maternal behaviours such as grooming and nurturing modulated infant cortical activity, sometimes inducing desynchronisation and sometimes reducing it. According to Sarro et al., there ‘may be a mechanism allowing variation in maternal care to create variability in brain development and behavioural outcome. The nature and duration of specific maternal behaviours, as well as the temporal patterning of interaction and absence, could induce robust individual differences in arousal and consolidation, as well as network maturation’.

In a study on human subjects, Hane et al. (46) confirmed altered cortical activity due to differences in caregiving. They scored mothers according to variations in their maternal caregiving behaviour when their infants were nine months old. A correlation between these scores and infant EEG at three years was then investigated as an indicator of stress reactivity in human infants. When compared to high-quality maternal caregiving, low-quality maternal caregiving led to an increase in stress reactivity determined by right frontal EEG asymmetry.

In their booklet, Browne et al. described that the establishment of adequate bonding through such high-quality caregiving has profound effects on animals’ emotions and responses to stress, accompanied by important changes in their central levels of BDNF. As previously mentioned, BDNF is a regulator of neuronal development. Increased BDNF promotes and accelerates central nervous system maturation, possibly leading to more adaptive coping styles and reduced vulnerability to psychopathology (13). Browne et al. stated that ‘researchers have therefore sought for ways to increase cerebral BDNF endogenously or exogenously in humans’. They concluded that the most feasible option to increase BDNF, as well as other substances that stimulate development, would be endogenously. Interventions that enrich the early life environment by focusing more on parent–infant bonding could as a result cause an endogenous increase in BDNF, as well as an increase in other substances important to development (13).

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Human studies that assessed caregiving or interventions that increased bonding

The most common example of a caregiving method that increases the focus on bonding in NICUs is Kangaroo care (KC). KC is a method of creating skin-to-skin contact by holding a baby in a supine position on a naked caregiver's chest when the baby is wearing nothing but a diaper (47). A longitudinal study by Feldman et al. was set up to examine the effect of KC on preterm infants, parent–infant interaction and child development (4). Infants who received KC for 14 consecutive days were matched to control infants, with both groups containing 73 infants. They were all assessed at three time intervals during the first half year and seven other time intervals during the first decade. During the first half year, family interaction and home environments were more adapted in the KC group. Mothers reported less depression and infants scored higher on the Bayley Mental Developmental Index and the Psychomotor Developmental Index. Feldman et al. speculated that KC had both a direct impact on the infants' neurophysiological regulation as well as an indirect impact by improving parental mood, perceptions and interactive behaviour (4). During the infants' first decade, KC increased maternal attachment behaviour and infants' autonomic functioning as measured by respiratory sinus arrhythmia. Furthermore, child cognitive development and executive functions were enhanced. By 10 years of age, KC children showed attenuated stress responses, more mature respiratory sinus arrhythmia patterns, better organised sleep and better cognitive control. Respiratory sinus arrhythmia patterns and maternal behaviour were dynamically inter-related over time, demonstrating coregulation (4). Such positive effects of KC were reinforced in a Cochrane review on Kangaroo mother care. This term is used to describe KC within a broader caregiving principle that adds almost exclusive breastfeeding and attempted early discharge from the hospital to daily, preferably long, KC sessions (47).

Other interventions aimed at increasing bonding described in the literature were, for instance, maternal voice recordings or maternal biological sounds (48) music therapy (49) or a breathing bear (50). Lahav et al. performed a number of studies on using maternal sounds in the NICU, all with promising effects. For example, in one study increased cardiorespiratory stability, with a reduced frequency of apnoea and bradycardia, was reported (48). Research, therefore, appears to support the use of familiar or positive sounds in NICUs. This was also confirmed by a recent meta-analysis on music therapy that concluded that evidence-based NICU music therapy was highly beneficial with an overall large significant effect size. Benefits were greatest with live music therapy, such as mothers singing (49). Voice vibrations and the rhythmicity of a maternal voice, as well as maternal breathing patterns and heart beats, are features that healthy, term born neonates normally rely upon to support their regulatory development (48). Access to such audible features could thus be of even more importance to preterm infants, who, as previously stated, are exposed to many unnatural stimuli and have a far more vulnerable physique (10). Breathing is a serious challenge for many preterm infants due to lung immaturity (1). Extra coregulatory support for breathing using audible biological sounds or music with vital rhythms could thus be very beneficial (48).

Thoman et al. used breathing sounds in a device suitable for preterm and full-term infants, called the breathing bear. This is a teddy bear designed to permit entrainment of the infant's breathing capacity and related neural functions. It breathes quietly to ensure that the intervention is not imposed on the infant. In a study with irritable full-term infants, the device was placed in the crib and the infant could control whether, when and how long to make contact with it and experience its rhythmic body motion. Compared to a control group of irritable infants without exposure to the breathing bear, positive effects on contact behaviour, temperament, maternal stress and maternal depression were reported (50). Superior effects to those of the breathing bear could presumably be caused by sleeping in the same bed as a sibling or parent.

Cobedding was investigated in a Cochrane review (51). Only five studies could be included in that review, all of which were small and reported their outcomes differently. There was, therefore, insufficient evidence to make recommendations for future practice. No significant differences in weight gain, apnoea, bradycardia, desaturation, infection incidence, length of hospital stay or parental perceptions were found. Cobedded twins appeared to spend more time crying, but they also appeared to spend more time in quiet sleep. Morgan et al. (52) examined whether infants should be cobedding with their mothers. In a within-subject design, they measured heart rate variability to assess autonomic function in 16, two-day-old full-term healthy neonates sleeping in skin-to-skin contact with their mothers and sleeping alone prior to discharge. All neonates were admitted to the hospital because they were born by Caesarean section. Both sleeping methods were measured for one hour. Results show a 176% increase in autonomic activity and a 86% decrease in quiet sleep duration during maternal separation compared to skin-to-skin contact. The separation thus appeared to increase stress.

FLacking et al. (23) reviewed the concept of closeness and separation in neonatal intensive care units more generally than the cobedding concept. They too highlighted the need for acknowledging the importance and impact of both physical and emotional closeness between the preterm infant and parent in the NICU. Emotional closeness referred to parental feelings of being emotionally connected, feelings of love, warmth and affection. In their review, the evidence that illustrated the importance of closeness came from stress and cortisol studies. Close physical contact between a parent and a preterm infant decreased the infant's cortisol levels and pain responses (35). In addition, an increase in the amount of physical contact and family-centred care synchronised cortisol variation between the preterm infant and his or her mother.
It appeared that sharing the same environment increased the concordance between the HPA axis reactivity of the preterm infants and their mothers, reflected by the cortisol levels in their saliva (53). In addition, various interventions that have supported the abilities of parents to observe and interpret their infant’s behaviour have all been associated with improved cognition years later (4,54). Flacking et al. thus concluded that parent-infant closeness should be an absolute priority within NICUs (23).

The importance of closeness was also described in the Postnatal maternal care, tactile stimulation and brain development section in the review by Samra et al. Concerning that, they reported that preterm infants were vulnerable to programmed changes if their care environments were less than optimal. Furthermore, they reported on evidence linking prematurity and family-centred care to epigenetic modifications. They concluded that more research is needed to understand these variables and their moderating or mediating relationships to each other (55). An attempt to address this was made by Milgrom et al. and Welch et al. in their studies, which examined caregiving environments by investigating different interventions that supported parents.

With their early sensitivity training programme, Milgrom et al. (56) reached a significantly enhanced white matter maturation and connectivity on magnetic resonance imaging in prematurely born infants when they were 38–40 weeks of age. Absolute volumes of cerebral tissue types revealed no differences. In this randomised, controlled trial, 45 women with infants younger than 30 weeks of gestation were involved. The intervention (n = 22) consisted of 10 sessions of parents working with therapists to learn to recognise signs of infant stress, shutdown mechanisms, alert available behaviour, quality of motor behaviours, facial expressions and posture or muscle tone. In comparison with NIDCAP, this intervention had relatively low costs due to less nurse training and nurse time assessing the infant. The parents assessed the preterm infant, which had possible postdischarge benefits as well. In addition to the increased white matter maturation, other medical outcomes demonstrated a general improvement for children in the intervention group as well. For example, fewer days on oxygen and a decreased length in hospital stay were reported. So, even though not all of these improvements were statistically significant, the findings of this study concurred with recent evidence that the quality of early life experiences influences physiological development and especially brain development. The results provided a platform for consideration of larger randomised trials investigating this type of interventions in preterm infants (56).

Welch et al. conducted such a large, randomised controlled trial that investigated an intervention with a different approach. The intervention developed and studied by the authors was called the Family Nurture Intervention. In their research, they assessed the impact of that intervention on EEG activity in 134 preterm infants born between 26–34 weeks of gestation (57). The distinguishing feature of this intervention was that the staff facilitated communication on affect and physiological coregulation between the mother and her infant. The intervention started early and continued over the full course of hospitalisation. It entailed mother and infant scent cloth exchange, sustained touch, vocal soothing, eye contact, wrapped or skin-to-skin holding and family-based support sessions (58). Family Nurture Intervention infants showed robust increases in frontal brain activity on EEG, which other investigators found predictive of better neurobehavioural outcomes. Effects were significant in both quiet and active sleep, regardless of gender, gestational age or birth weight. Significant group-by-age effects in other brain regions were found as well (57). Additional studies assessing this intervention demonstrated not only improved maternal caregiving behaviour (59), but also significantly improved scores across multiple domains of neurodevelopment, social relatedness and attention problems in the intervention group compared to the control group receiving standard care (60). These results suggest widespread changes in developmental trajectories mediated by the Family Nurture Intervention, an intervention focused entirely on enhanced coregulation, or in other words, enhanced bonding.

**DISCUSSION**

Recent findings suggest that early life conditions turn out as either harmful or protective, depending on the organism’s later life environmental context. This implies that early life conditions may prepare the infant for their future life through glucocorticoid programming and phenotypic plasticity with the goal to match with future environmental demands (10,39,40). This concept has led to the hypothesis that a mismatch between early and later life conditions can enhance vulnerability to disease. For example, although the offspring of low maternal care dams seemingly displayed lower cognitive performance under basal conditions when compared to the offspring of high maternal care dams, a different outcome was described when the rats were tested under stressful conditions, that is a situation reminiscent of their early life environment (28,33). These findings suggest that what used to be perceived as a maladaptive phenotype may turn out to represent an adaptation to the environmental context that prevailed during development, which appropriately predicts and matches with the environment later in life. This is a phenomenon that has adaptive values in particular contexts, but can be maladaptive when mismatched and persistent (34,37,40).

More evidence for this theory comes from considering the correlations between the reproductive system and human parental programming over an evolutionary time span. Under high-risk conditions, when the probability of survival is low, the optimal strategy is to maximise the number of offspring through accelerated mating. In contrast, a more propitious environment favours greater investment in individual offspring at the cost of mating (43). Thus, the adaptive maturation of neural and endocrine systems requires environmental input to optimise their...
functions. The outcome is somewhere on the continuum between resilience and psychopathology. It depends on the nature of the input, the balance between stimulant as well as challenging input, the caregiving during the challenges, the respective genetic predispositions and on the social context in later life, which all depend on each other as well (28). Due to this complexity, the directions of changes are sometimes inconsistent and unexpected and even small differences can affect the direction of change. Additionally, changes can even be different within one brain region (39).

This review was not intended to explain those varying, context-dependent directions of alterations. It meant to raise clinicians’ awareness of the omnipresence of changes caused by differences in bonding-related early life experiences, by providing a large, convincing amount of varying examples. Unfortunately, due to that variety, the actual summing up of findings, which would have resulted in a more number-oriented review, appeared impossible. Another limitation was the fact that the majority of this review consisted of animal studies. However, there is now strong evidence that human and animal parenting share many subcortical, neural and neurochemical mechanisms (8–10,55). Furthermore, albeit not always significant, numerous positive effects of interventions that aim to optimise bonding in human studies have been reported, in contrast to not one negative effect. Improving bonding should therefore be prioritised (10,13,60), particularly in high incidence, vulnerable populations such as preterm infants (1).

CONCLUSION
This review emphasises the importance of bonding by providing sound evidence for the fact that differences in bonding-related early life experiences and NICU-related early life experiences have a significant impact on physiological development in various ways. That impact is highly context dependent and frequently gender dependent as well. Furthermore, the impact can be reversed by improving the caregiving context and by optimising parent–infant bonding. The attention for adequate bonding in NICUs should therefore be increased.

CONFLICT OF INTEREST
None declared.

ACKNOWLEDGEMENTS
None.

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28. Bockm
28. Bockm

APPENDIX: METHODS

A MEDLINE and EMBASE search with broad search terms was undertaken through May–October 2014, to extract bonding-related perinatal factors from the available literature as a first step. There were four sorts of search terms; for the population (neonates, infants, etc.), for the determinant (bonding, caregiving, etc.) and for the changing (chang*, improv*, develop*) of the physiology with an emphasis on brain/neuronal physiology (brain, neuron*, EEG, physiology, organ, homeostas*, etc.; see Table A1). These search terms were decided upon in advance, based on their frequent usage in previously read, relevant articles. A thesaurus was used to find additional relevant terms. All matching titles were assessed and either included or excluded, according to predetermined, broad criteria (see Table A2). After this screening process, the included abstracts and their full texts were read in order to include or exclude articles according to new, more specific inclusion and exclusion criteria (that were also predetermined). For these criteria, see Table A4. Some articles were redistributed. In addition, reference lists of identified articles and related reviews were hand searched. For the final flowchart of all included articles per topic, see Figure A1.

The subsequent writing stage consisted of three more steps. First, an extensive, non-selective work containing all results was undertaken through May–October 2014, to extract bonding-related perinatal factors from the available literature as a first step. There were four sorts of search terms; for the population (neonates, infants, etc.), for the determinant (bonding, caregiving, etc.) and for the changing (chang*, improv*, develop*) of the physiology with an emphasis on brain/neuronal physiology (brain, neuron*, EEG, physiology, organ, homeostas*, etc.; see Table A1). These search terms were decided upon in advance, based on their frequent usage in previously read, relevant articles. A thesaurus was used to find additional relevant terms. All matching titles were assessed and either included or excluded, according to predetermined, broad criteria (see Table A2). During the assessment of all the titles, terms used for bonding-related perinatal factors and frequently occurring outcome parameters were extracted. They were written down to find a structured answer to the question of what does suboptimal bonding do to physiological development. With the written down terms, an associative and categorised framework was constructed (see Table A3), before reading the abstracts and full texts.

Successively, abstracts were screened to either subdivide them into their categories (preliminary inclusion) or to exclude them. The same broad inclusion and exclusion criteria that were previously used to screen the titles with were used for this abstract screening process (Table A2). After this screening process, the included abstracts and their full texts were read in order to include or exclude articles according to new, more specific inclusion and exclusion criteria (that were also predetermined). For these criteria, see Table A4. Some articles were redistributed. In addition, reference lists of identified articles and related reviews were hand searched. For the final flowchart of all included articles per topic, see Figure A1.

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| **Table A1** Search terms. At least one synonym of each term present in title/abstract. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|
| Term 1 | Term 2 | Term 3 | Term 4 |
| Develop* | Maternal care | Premature | Physiologic* |
| Improv* | Paternal care | Newborn* | Organ |
| Growth | Parental care | Childhood | Homeostas* |
| Chang* | Standard care | Infant* | Brain |
| Differentiat* | Caregiv* | Preterm | Neuron* |
| Matur* | Nurtur* | Perinatal | Hippocamp* |
| Bonding | Bonding | Postnatal | Axon* |
| Attachment | ‘Mother infant’ | Neonat* | EEG |
| ‘Mother infant’ | Offspring | MRI | Dendrit* |

*Means all possible extensions of the given word are included.

| **Table A2** Predetermined criteria for title inclusion or exclusion. |
|-----------------|-----------------|
| **Inclusion criteria** | **Exclusion criteria** |
| All childhood experiences | Article not in English, Dutch, French or German |
| All physiological outcomes | Explicitly about children with syndromal diseases only |
| | Explicitly comparing only socio-economical influences (e.g. poverty, malnutrition) |
| | Explicitly concerning mothers with psychiatric diseases only |

| **Table A3** Associative framework dividing all the articles into seven categories after inclusion. |
|-----------------|-----------------|
| **Terms for early life bonding-related factors** | **Physiological outcome parameters** |
| Parental licking and grooming | Dendritic spine density |
| Parental presence/separation | Oxytocin |
| Environmental enrichment | Cortisol |
| Kangaroo Mother Care | Altered amygdala appearance |
| Institutionalisation | Dopamine |
| Attachment style | HPA axis reactivity |
| Received caregiving | Stress-axis functionality |
| Bonding | Receptors |
| Feeding routines | Synapses |
| Parental sensitivity | Hormonal concentrations |
| Tactile stimulation | Methylation |
| Body heat | Epigenetic changes |
| Early life experience | Epigenetic modulations |
| Early life adversity | Hippocampal changes |
| Abuse | ...many more... |
| Neonatal handling | Physiological outcome parameters |
| Still-face paradigm | divided into five categories |
| Nurturing intervention | | |
| Comfort(ing) | Stress-axis parameters |
| Comfortin device | Genetic parameters |
| (High) quality caregiving | Hormonal parameters |
| Developmental care | Neuronal parameters |
| NICU standards/NICU care | All outcomes |

Seven categories into which all included articles are divided

1. All factors concerning parental care altering stress–axis parameters
2. All factors concerning parental care altering genetic parameters
3. All factors concerning parental care altering hormonal parameters
4. All factors concerning parental care altering neuronal parameters
5. Caregiving in institutions or hospitals altering all physiological outcomes
6. Enriched environments altering all physiological outcomes
7. Bonding increasing interventions altering all physiological outcomes
Table A4  Predetermined criteria for full-text inclusion and exclusion.

<table>
<thead>
<tr>
<th>Cat.</th>
<th>Inclusion criteria per category [AND]</th>
<th>Universal exclusion criteria, incl. validity crit. [OR]</th>
</tr>
</thead>
</table>
| 1    | • Comparing differences in parental care  
• At least one altered stress-axis measure  
  (incl. altered symp./parasymp. gut activity or altered vital parameters indic. stress)                                                                                                                                                                                                                                           | • Either one of the exclusion crit. from Table A2  
• About non-caregiving-related early life stress only  
• Use of I.Q. as the only outcome (= not physiological)  
• Physiological outcomes only apparent under extreme, controlled (laboratory) circumstances  
• Use of substance administration  
• Reversed research designs; investigating abnormal populations and their controls retrospectively  
• Animal research that was not within-litter nor randomised  
• Inappropriate statistical analysis  
• Human studies uncorrected for baseline differences  
• Double use of research population  
• Case report  
• Double article  
• Full text nowhere available e.g. conference abstract  
• Article not peer-reviewed                                                                                                                                         |
| 2    | • Comparing differences in parental care  
• At least one altered genetic outcome                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 3    | • Comparing differences in parental care  
• At least one altered hormonal outcome  
  [except for ’only cortisol’ outcomes (= cat. 1)]                                                                                                                                                                                                                                                                            |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 4    | • Comparing differences in parental care  
• At least one altered neuronal outcome                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 5    | • Caregiving in institutions or hospitals  
• Making a comparison (either to controls or with subgroups)  
• At least one altered physiological outcome                                                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 6    | • Use of an environmental enrichment paradigm  
• Making a comparison (either to controls or with subgroups)  
• At least one altered physiological outcome                                                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
| 7    | • Any bonding increasing intervention (compared to controls)  
• At least one altered physiological outcome                                                                                                                                                                                                                                                                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |

Cat, Category; incl, Including; crit, Criteria; symp./parasymp, Sympathetic/parasympathetic; indic, Indicating; I.Q, Intelligence quotient.

Table A5  Final filtering step; criteria to narrow down the results to be used in this review.

<table>
<thead>
<tr>
<th>Inclusion criteria</th>
<th>Exclusion criteria</th>
</tr>
</thead>
<tbody>
<tr>
<td>Clinically relevant</td>
<td>Not importantly contributing to theoretical background</td>
</tr>
<tr>
<td>Directly linked to the conclusion</td>
<td>Expected/similar finding to those previously described</td>
</tr>
<tr>
<td></td>
<td>Too difficult to translate to human research</td>
</tr>
</tbody>
</table>

findings from all included articles per topic was created. Then, the general conclusion was written, based on undisputed issues only. This conclusion was now used as the final measurement for information to be valued with, to minimise the selection bias while narrowing down the article. Significance to that conclusion, as decided upon with some final predetermined criteria (Table A5), led to the final inclusion, exclusion and reordering of the findings. Various illustrating results that were exemplifying but did not independently add value to the conclusion were thus excluded.

During the revision phase, the main search was repeated and an extra hand search was executed in order to be as up to date, as explicit and as complete as possible.
Fig. A1 Flow chart literature search. Art, articles; # art/cat, current number of articles per category; c.o. IN, cross over ingoing, from another category into this category; L.A.S., later automatic searches, automatically done by PubMed and EMBASE; H.S., handsearch, articles included by reference guided handsearches; tot. IN, total amount of articles included; c.o. OUT, cross-over Outgoing, from this category into another one; Excl acc. to tab 4, excluded according to Table A4; F. Incl, Final number of articles included.