Title
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Permalink
https://escholarship.org/uc/item/3gb7625z

Journal
Peptides, 72

ISSN
01969781

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Publication Date
2015-10-01

DOI
10.1016/j.peptides.2015.03.020

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Peer reviewed
FETAL EXPOSURE TO PLACENTAL CORTICOTROPIC-RELEASING HORMONG (pCRH) PROGRAMS DEVELOPMENTAL TRAJECTORIES

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Abstract

The maternal endocrine stress system is profoundly altered during the course of human pregnancy. The human placenta expresses the genes for CRH as early as the seventh week of gestation and it is the exponential increase in placental CRH (pCRH) over the course of human gestation that is responsible for the greatest modification in the maternal stress system. The bi-directional placental release of hormones into the maternal and fetal compartments has profound influences for both. The influential Fetal Programming model predicted that early or fetal exposures to maternal signals of threat or adverse conditions have lifelong consequences for health outcomes. A basic assumption of this model was that developing organisms play a dynamic role in their own construction. Data are reviewed and new data are presented that elevated pCRH over the course of human gestation plays a fundamental role in the organization of the fetal nervous system, modifies birth phenotype (the timing of the onset of spontaneous labor and delivery), and influences developmental temperamental and metabolic trajectories. Evidence for sex differences and conserved function across species is presented. Finally, a model is presented that proposes several pathways that pCRH can program risk for health and disease.

Keywords
Corticotropic-releasing-hormone; fetal programming; Stress; Human pregnancy; Fetal development; Sex differences; Child development

PRECIS

It is difficult to know where to begin to describe my relationship with Abba, Mexico City? Philadelphia? Breckenridge? Atlanta? Cozumel? Utrecht? Chicago? or perhaps at the beginning, in New Orleans. I was drawn to the Deep South in the mid-1960’s by my desire to become involved in the Civil Rights movement. I had never travelled to the Deep South.
from the relative tranquility of my California home and, unlike today, there was a scarcity of information available about the culture and politics of various destinations, so my decision about where to go was somewhat random. I was in my mid-twenties and made the decision that graduate school would be the vehicle to allow me and my small family to make a move. When I arrived on the Louisiana State University (LSU) campus in Baton Rouge, there were fewer than 30 African-American students enrolled among the 15,000 plus students. Martin Luther King was assassinated during my tenure at LSU and that tragedy was the impetus for a fledgling civil right movement on campus. I was a founding member of the Martin Luther King Action Movement (MLKAM) on campus and in parallel I quickly co-opted the psychology national honorary society to begin a petition against the Viet Nam war. The MLKAM remained local and contentious (and dangerous for some) but the petition against the war in Viet Nam mysteriously became national and resulted in a late Sunday night knock on my door from an agent of the US government. The faculty was passively supportive of my activities but as I found out, labeled me a “trouble-maker.”

As part of my training, the graduate program assigned me in the summers to the New Orleans Veterans Administration (NOVA) hospital. During my second summer I attended a grand round presentation related to a patient I had been working with. The patient (let’s call him Nate) was a young, very smart African American man who had recently returned from Viet Nam. Nate had taken a job at a New Orleans gas station and when an impolite customer had called him “boy” Nate promptly knocked him to the ground. Because no “sane” African American in the 1960’s would strike a white man in the Deep South, Nate was admitted to the psychiatric ward. He was perfectly sane, smart and angry and refused to comply with some of the degrading demands of the staff (such as stripping and taking medications). During the grand rounds I was more than shocked to hear an eminent consultant seriously suggest that a lobotomy might be considered a reasonable treatment. When I regained my senses and realized that this was a serious proposal, I raised my hand in protest and claimed that I could work with Nate and perhaps we could re-evaluate his case later. Without going into detail, once I told Nate what was going to happen and how he could avoid it, he was discharged within a month. Because the label of “trouble-maker” had followed me to the NOVA, coupled with my “defiance” of authority in the Grand Rounds there was an effort to re-channel my energy. The director of the psychology service, who always seemed to be terrified by me, told me about a request made by a group of researchers in a “mysterious” building for someone interested in collaborating. None of the faculty were interested and the director thought I might be (not really, he just wanted to get rid of me).

I entered Abba’s laboratory with large vats of frogs. Frogs were everywhere. He explained how frog skins were used as a bioassay for Melanocyte-Stimulating-Hormone (MSH) and that although MSH was known to influence pigmentation in amphibians, new data published by the group at the University of Utrecht indicated that it may have effects in the mammalian brain. Rats treated with MSH appeared to learn a simple discrimination faster than rats treated with placebo. The evolutionary and teleological implications regarding what was conserved and why, was captivating and we discussed possible directions for a research project. He armed me with several papers to read and asked me to return with a specific idea to test in rats. He did not seem to care that I might have been a trouble-maker or concerned that I had never worked with animals before (I probably did not tell him). The
available literature at that time only examined the effects of MSH on tasks with aversive or stressful consequences so our first collaborative and published paper reported that MSH improved learning on an appetitive task (55). Many papers describing the effects of peptides on animal and human development have followed and currently I continue to examine the influence of gestational exposures to neurobiological signals of stress (including peptides and hormones) on the brain and behavior of children. The influence that Abba had on my personal and professional life cannot be overstated.

STRESS, HUMAN PREGNANCY AND FETAL EXPOSURE TO PLACENTAL CRH

Based on our initial and novel studies that described the long lasting (perhaps permanent) effects on the brain and behavior of rats exposed to stress peptides during very early-life (fetal or early infancy; 2, 6, 8, 53, 59, 60), a program of research (Figure 1) was initiated to examine of the effects of activation of the maternal endocrine stress axis during pregnancy on the human fetus. A fundamental assumption of our research program is that prenatal exposure to maternal HPA and placental hormones represent primary mechanisms underlying the effects of maternal psychological distress on subsequent infant and child development. From this program we have learned that during human pregnancy there is a complex relation between psychosocial and biological markers of prenatal stress. Both sources of stress have programming consequences for the human fetal nervous system, birth outcome and risk for subsequent health and disease (44–47).

The maternal endocrine stress (hypothalamic-pituitary-adrenal; HPA) system is profoundly altered during the course of human pregnancy. The pituitary gland doubles in size increasing by several fold the synthesis, and release of pituitary peptides into the maternal circulation. Production from target tissues, such as cortisol from the adrenal gland also increases two to four-fold over the course of pregnancy. But it is the growth and development of a new fetal organ, the placenta in primates that is primarily responsible for the profound changes in the maternal/fetal stress systems (45,47). The human placenta expresses the genes for CRH (hCRHmRNA) (as well as proopiomelanocortin, the precursor for ACTH and beta-endorphin (BE)) as early as the seventh week of gestation and it is the expotential (20 to 40-fold) increase in placental CRH (pCRH) over the course of human gestation that is responsible for the greatest modification in the maternal stress system (Figure 2).

Placental CRH is identical to hypothalamic CRH in structure, immunoreactivity and bioactivity (40, 63,64). However in contrast to the inhibitory influence (negative feedback) on the promoter region of the CRH gene in the hypothalamus, maternal stress signals from the adrenal glands (i.e., cortisol) activate the promoter region in the placenta and stimulate the expression of hCRHmRNA establishing a positive feedback loop that allows for the simultaneous increase of placental CRH (pCRH), ACTH and cortisol over the course of gestation (Figure 2). Thus, all of the stress related peptides/hormones levels rise as pregnancy advances, peaking during labor, and falling to basal or, in the case of pCRH, undetectable levels within 24 hours after delivery. The normative exponential increase of these stress signals, (especially pCRH) over the latter part of human gestation, (i) plays a fundamental role in the organization of the fetal nervous system (61), (ii) influences the

*Peptides. Author manuscript; available in PMC 2016 October 01.*
maturation of the fetal HPA axis and other systems, (iii) modifies birth phenotype (the timing of the onset of spontaneous labor and delivery) 35, 65, 66, 68, 69, 72 and (iv) regulates maternal adaptation during pregnancy (21, 46).

The human placenta integrates numerous sources of maternal stress signals and responds with a dose dependent release of CRH. Detection by the fetal/placental unit of stress signals from the maternal environment (for instance, cortisol) “informs” the fetus that there may be a threat to survival. The placental/fetal unit responds to this by synthesizing and releasing CRH (positive feedback) and activating a cascade of consequences. Depending on the timing during gestation and the severity of the stress the myometria are activated resulting in abbreviated gestation and early fetal escape from a hostile environment. In parallel the fetus incorporates the bidirectional information to adjust its developmental trajectory and modifies its nervous system to ensure survival in a potentially hostile postpartum environment (48). Survival under these conditions can be associated with compromised growth, reproductive success, motor, cognitive and emotional function.

Moreover, it is believed that high levels of placental CRH circulating in the maternal bloodstream down-regulates maternal corticotrophs blunting the communication between the hypothalamus and pituitary gland and affecting her response to the environment (21). This not only explains the decrease in maternal responses to stress as pregnancy advances but also has been proposed as a mechanism for postpartum depression (22).

The bi-directional placental release of hormones into the maternal and fetal compartments is a powerful route of communication between the fetus and mother with long term and profound influences for both. The Fetal Programming or Developmental Origins of Health and Disease Models (1) were proposed to explain the persisting influences associated with prenatal exposure to stress. This influential model predicted that early or fetal exposures to maternal signals of threat or adverse conditions have lifelong consequences for health outcomes. A basic assumption of this model was that developing organisms play a dynamic role in their own construction. (44, 46, 47). Signals from the maternal stress systems are continuous sources of information to the fetus over the course of gestation. As described above, these systems are undergoing massive but orderly changes as pregnancy advances. Deviations in these systems from normative patterns can be signals to the fetus that there is a threat to the host (mother). Depending on the severity and the timing of deviant (stressful) maternal signals the sequence of neural development can be disrupted resulting in programmed consequences for brain structure and behavior. Fetal exposure to deviant elevations of stress and placental hormones early in pregnancy have been associated with less optimal neurodevelopment, more irritable and fearful temperament, larger volumes in limbic areas of the brain and slower behavioral recovery from pain (5, 11, 12). Fetal exposure to increases in these hormones occurring very late in gestation has been associated in some instances with enhanced cognitive development (56). These influences occur against the background of the timing and sequence of fetal organ development.

The fetal period in the life cycle is unmatched by any other in growth and development, and it is the stage in the human life span that is most vulnerable to both organizing and disorganizing (programming) influences. Fetal organs develop from progenitor stem cells at
precise times and in a specific sequence from conception to maturity, so the timing of maternal signals is a critical factor in determining the structure of the neurodevelopmental program. Disruption in the timing or sequence of organ development can result in tissue remodeling producing smaller organs or altered organ morphology. Remodeled tissue modifies the function and physiological capacity of the organ throughout the lifespan and is a fundamental assumption of how fetal exposures influence health and disease.

The human fetal nervous system is a primary target for these circulating programming influences because it is undergoing dramatic growth over a prolonged period of time. For instance, radial neuronal cell migration begins in the human brain around 42 days GA (71) and by 16 weeks GA, form the subplate zone. Concurrently, cells accumulating in the outer cerebral wall form the cortical plate which will become the cerebral cortex. By gestational week 20, axons form synapses with the cortical plate and there is an exponential increase in cortical thickness (22) and by gestational week 24, cortical circuits are organized (31). The human fetal brain is forming secondary and tertiary gyri, and exhibiting neuronal differentiation, dendritic arborization, axonal elongation, synapse formation and collateralization, and myelination by gestational week 28. Near term the fetal human brain contains billions of neurons and is 40% greater in number than in the adult (25). The rate of synaptogenesis reaches an astonishing peak so that at gestational week 34 through 24 months postpartum, there is an increase of 40,000 synapses per minute (32).

PREGNATAL EXPOSURE TO pCRH AND FETAL DEVELOPMENT

Our program of research has been focused on the influence of fetal exposure to maternal psychobiological markers of stress on fetal development. As discussed above, we particularly have been interested in variations in the concentrations of pCRH (i) because it is a major stress peptide with direct effects on the nervous system and (ii) because deviations in pCRH provides objective evidence that the fetus has been exposed, and is responding, to maternal signals of stress. Studies of human fetal heart rate in response to stimulation provide a model for assessing neurological integrity before there are postpartum influences on development such as socialization and parenting (62). In a prospective longitudinal study, we assessed fetal nervous system maturation at 26, 31 and 37 weeks gestation by monitoring fetal heart rate (FHR) responses to vibroacoustic stimulation (VAS) in 191 maternal/fetal dyads (4). The VAS is used routinely to awake a sleeping fetus and consists of a tone and vibration from a device placed on the mother’s abdomen. Reliable startle responses were not detected in all fetuses until 31 weeks gestational age. There was, however, evidence that some fetuses did respond at 26 weeks, so we considered that there may be individual differences in response patterns that could be explained by exposures to placental markers of stress. To examine this we (8) evaluated in 138 maternal/fetal dyads the association between startle responses at 26 weeks and pCRH levels collected at 15, 20 and 26 weeks of gestation. Elevated placental CRH levels at 15 weeks of gestation, but not later points, predicted (r (137)=.24, p<.01) smaller peak FHR responses to the VAS (FIGURE 3).

Figure 4 shows that fetal exposure to the lowest levels of CRH at 15 weeks gestation is associated with a clear response and recovery pattern to the VAS at 26 weeks gestation. Fetal exposure to the highest level of CRH is associated with failure to respond. These

*Peptides. Author manuscript; available in PMC 2016 October 01.*
findings suggested that fetal exposure to the lowest concentrations of pCRH (optimal, low stress exposure) early in gestation reflected greater fetal maturity and accelerated neurological development. In an earlier study we had assessed effects of fetal exposure to pCRH on fetal memory and attention by presenting a series of repeated VAS, interrupted by a novel VAS. In this model, improved memory and attention is reflected in the extent to which the novel stimulation disrupts the pattern of habituation. We reported that lower placental CRH during the third trimester was associated with an improved ability to habituate to repeated presentations of the same VAS stimulus and to identify a novel stimulus (61).

We repeated the habituation experiment by presenting pure tone stimulation to the fetus delivered by speakers attached to the mother’s abdomen. Following previously reported methods (52,62) we compared the slopes of the FHR response to the last four stimuli of the initial trials (trials 12–15, slope2) with the first four trials after the novel, dishabituating stimulus (trials 17–20, slope 3) (Figure 5). It is apparent that fetal responses at 36 weeks are characterized by a large initial FHR response followed by a progressive decrease in response (habituation). At 30 weeks fetuses show sensitization (progressive FHR response to subsequent stimulation) and there is no evidence of a response at 26 weeks GA. Moreover, at 36 weeks there is a fetal heart rate response following the novel stimulus (tone sixteen) and then a trend toward dishabituation. We found that dishabituation increases with age in both groups however fetuses exposed to low concentrations of pCRH show a greater dishabituation response (better learning) than fetuses exposed to high levels of CRH ($F_{(1,130)}= 5.3$, $p= 0.02$). Based on the first four tones, we divided the groups who decrease (habituators) and those who increase (sensitizers) heart rate over these four trials. There was a trend suggesting that fetuses classified as habituators at 36 weeks gestation were exposed to lower CRH concentrations at 15 weeks gestation than the sensitizing fetuses (Figure 6). These findings suggested that exposure to CRH early in pregnancy influenced fetal maturation as measured by startle and habituation: high concentrations “retard” maturation and low levels accelerate it.

FETAL EXPOSURE TO pCRH AND BIRTH PHENOTYPE

Biological markers of stress consistently have been associated with adverse birth outcomes. The general findings among methodologically sound studies are that women reporting elevated levels of psychosocial stress during pregnancy are at significant risk for adverse birth outcomes (19, 33). Within the stress pathway, pCRH is most strongly linked to gestational length. Placental CRH has been characterized as controlling a “placental clock” that determines or alters the timing of onset of parturition (35, 65). Elevated levels and steeper trajectories of pCRH over the course of gestation initiate a cascade of events resulting in myometrial activation and in extreme cases preterm birth (35, 67). Because, it is the trajectory of pCRH production (i.e., the rate of acceleration), rather than the absolute hormone concentration that best predicts preterm birth it suggests that target cells are highly responsive to relative changes in pCRH concentrations (67).

In a study published in Peptides to examine the association between birth outcome and a panel of biological stress markers, maternal levels of B-endorphin, ACTH, cortisol and
pCRH were assessed at regular intervals from 15 to 36 weeks gestation in 203 pregnant women (49). Consistent with previous studies, pCRH levels in women destined to deliver preterm (before 37 weeks) had faster rates of increase and significantly higher levels of pCRH confined to the beginning of the early third trimester than women who subsequently delivered at term. Of the other maternal measures, only cortisol, as early as 15 weeks gestation, was elevated in women delivering preterm. Models that accounted for the independent and shared variance of pCRH and cortisol indicated that only pCRH between 26 and 31 weeks gestation predicted gestational length (Figure 7). However, we found that the best predictor of elevated pCRH at 31 weeks was elevated maternal cortisol at 15 weeks. The findings from this study indicated that a plausible stress-related endocrine signal, elevated cortisol from the mother very early in pregnancy, predicted the precocious rise in CRH leading to an abbreviated gestation. The pattern of findings supported the argument that the effect of elevated cortisol early in pregnancy reflected priming or programming effects on the eventual fetal/placental CRH response.

In addition to the regulation of gestational length, pCRH also plays a key role in the regulation of fetal maturation with consequences for birth phenotype. Neonatal evaluations of neuromuscular and physical characteristics of the newborn that develop over the course of gestation were done within 24 hours of birth to assess developmental maturation. In our study of 158 newborns within 24 hours after birth, fetal exposure to increased levels of maternal cortisol at 15 and at 19 weeks gestation and increased levels of pCRH at 31 weeks gestation were associated with significant decreases in newborn physical and neuromuscular maturation (20). These effects remained significant after adjusting for length of gestation.

FETAL EXPOSURE TO pCRH AND INFANT DEVELOPMENT

Research from our laboratory as well as from several international laboratories provides support for the influence of fetal exposures on developmental trajectories. It has become clear that one of the most consistent findings is that fetal exposure to psychological and biological stress signals is associated with more reactive physiological and behavioral responses to challenge during infancy perhaps reflecting more fearful temperament (11,15,16,38,73). In a sample of 247 mothers and their full term infants, we previously had examined the possible influence of the stress hormone pathway on infant temperament. We reported that elevated maternal cortisol at 30 gestational weeks (12) and elevated concentrations of pCRH at 25 gestational weeks each were significantly associated with greater maternal report of infant negative reactivity (11). Importantly these findings were independent of birth outcome, sociodemographic factors and postnatal maternal psychological measures. These novel findings were the first reports in infants/children of programming consequences of prenatal exposure to pCRH and suggested a biological pathway for the effects of maternal stress on the behavior of infants.

In addition to the effects on behavior and the central nervous system, there is evidence that prenatal maternal stress signals influence fetal growth, child obesity, and metabolic risk. In a recent study of 246 women and their healthy children, we (70) evaluated the association between fetal exposure to pCRH (from 15 to 37 weeks gestation) and child body mass index (BMI) in infants from 3 to 24 months of age. First, after adjusting for length of gestation,
elevated pCRH at 30 weeks gestation was associated with both BMI and weight at birth. Second, four profiles of childhood growth were identified during early childhood development. One of the growth profiles was characterized by early small body size followed by rapid catch-up growth. This pattern of catch-up growth has been associated with increased risk for obesity and was associated with elevated pCRH at 30 gestation. These data suggested that fetal exposure to pCRH contributes to programming of metabolic risk.

**SEX DIFFERENCES AND FETAL PROGRAMMING**

Under certain conditions secondary sex ratio (number of males to females born) is associated with, and perhaps programmed by, preconceptional and maternal/fetal exposures to signals of stress and adversity. Compared with females, more males (i) are born preterm (10), (ii) have poorer neonatal and infant health outcomes (39), (iii) have higher risk for motor and cognitive outcomes, and (iv) are less likely to survive in intensive care. It is believed that the differential mortality, “programming” of morbidity and developmental impairments in males occurs early in gestation (or even preconceptionally) so that a new reproductive cycle can be initiated and that parental resources can be directed to another pregnancy (75). For example, sex differences have been observed in mammalian animal models as early as meiosis (24). There is further evidence that within weeks of implantation the female placenta is more responsive than the male placenta to changes in stress signals including detection and response to maternal glucocorticoid concentration (9). In a recent review (50) we addressed sex differences by reviewing and reanalyzing data from previously published findings. We reasoned that if biological sex itself was influenced by early environmental exposures, then the influence of intrauterine exposures on males and females would be programmed differently.

As reviewed above, fetal exposure to cortisol early in gestation and pCRH at 31 weeks gestation was associated with delayed neuromuscular and motor maturation (20). Supplementary analyses indicated that these effects were significant only among males. Male fetuses exposed to elevated levels of cortisol early in pregnancy and CRH late in pregnancy exhibited delayed physical and neuromuscular maturation in very early infancy.

We previously had reported (and reviewed above) that fetal exposure to elevated pCRH at 25 gestational weeks in women was a risk factor for fearful or reactive temperament during infancy (11). New analyses illustrated that these effects were observed only among females. Specifically, elevated placental CRH at 25 gestational weeks was significantly associated with more fearful temperament and higher levels of distress behavior among female infants, but not male infants.

**CONCLUSION**

Figure 8 summarizes several pathways by which fetal exposure to stress generally, and pCRH specifically, can influence development and program risk for health and disease. Studies conducted (45, 47, 51) and underway in our program have been, and continue to be, designed to examine these possible pathways. One promising pathway is related to neuronal loss. Convincing evidence in animal models indicates that early life exposure to stress,
specifically including CRH, decreases dendritic spine density in areas of brain related to
cognitive and emotional regulation (7, 34). There is new evidence that fetal exposure to
biological and psychological markers of stress is associated with patterns of cortical and
subcortical development in children (5, 43) that may reflect dendritic arborization. We have
found that the patterns of nervous system remodeling in children mediate the association
between fetal exposure to adversity and subsequent behavioral trajectories and risk for
impairment. It will be a challenge to follow these findings to basic mechanisms of neuronal
loss.

In our initial study of peptide effects on behavior, indeed in the primary rationale for
examining the behavioral properties of MSH, we assumed that there must be conserved
functions of peptides across species. MSH and its fragments promotes escape behavior of
the amphibian by altering pigmentation (27) and in the mammal, including humans, by
increasing attention (26–30, 41,42, 53–59, 74). In both cases awareness of the environment
is essential suggesting a unifying evolutionary purpose.

In studies of early life exposures to CRH there is strong evidence for conserved function
across species that result in programming consequences for later life history outcomes. For
instance, surveillance and response systems have evolved and are conserved so that many
species including the desert dwelling Western Spadefoot tadpole can detect threats to
survival and adjust their developmental trajectory (3). In rapidly evaporating pools of desert
water with life-threatening consequences an elevation of CRH in the median eminence of
the tadpole precipitates metamorphosis (17,18). If the CRH response is blocked during
environmental desiccation, the rate of development is arrested and the tadpole’s survival is
compromised. There are long-term (life history) consequences for the surviving toad
exposed to this stress because it is smaller and is at a disadvantage against normally
developing toads in foraging for food and reproducing. As reviewed above, the human fetus
exposed to elevated pCRH may escape the inhospitable maternal host with abbreviated
gestation and/or suffer with temperamental or behavioral impairment. Just as the toad may
struggle to survive after early adversity, so will the developing child who has been exposed
to maternal signals of adversity including pCRH.

Acknowledgments

Portions of the research reported here was supported by National Institute of Health grants NS-41298, HD-51852,
HD-28413 HD-40967, HD-50662, HD-65823 and Conte Center award MH-96889.

Particular gratitude is expressed to the families (mothers and children) who have continued in our longitudinal
studies. I am grateful for the expert contributions of students, post-doctoral fellows and collaborators, most of
whom are referenced in this manuscript. Special appreciation is extended to long-time and current collaborators,
Laura M. Glynn, PhD and Elysia Poggi Davis, PhD. Of course none of the research reported here and the many
other research projects that have made my career and life a special blessing would be (or have been) possible if it
were not for the mentoring and life-long support of Abba Kastin.

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Peptides. Author manuscript; available in PMC 2016 October 01.


*Peptides.* Author manuscript; available in PMC 2016 October 01.


Peptides. Author manuscript; available in PMC 2016 October 01.
HIGHLIGHTS

Placental CRH influences human fetal heart rate
Placental CRH organizes human neurobehavioral development
Elevated placental CRH is associated with preterm birth
Elevated placental CRH is associated with fearful temperament in children
Fetal exposure to CRH remolds the nervous system
Conservation of function across species
Figure 1.
The assessment protocol for research program. Pregnant women and their fetuses are initially assessed at ~14 weeks gestation. Participants are followed in current studies until they are adolescents.
Figure 2.
During the course of human pregnancy all stress hormones rise. Placental CRH (pCRH) which is detected in maternal circulation rises exponentially over the course of gestation. In contrast to the negative feedback regulation, during pregnancy maternal cortisol increases the production of CRH from the placenta. The effects of maternal cortisol on the fetus are modulated by the presence of a placental enzyme 11βHSD2 which oxidizes it into an inactive form, cortisone. Activity of this enzyme increases as pregnancy advances, and then drops near term so that maternal cortisol is available to promote maturation of the fetal lungs, central nervous system as well as other organ systems.
Figure 3.
Scatterplot that shows fetal exposure to elevated pCRH at 15 weeks gestation is associated with dampened heart rate responses to an external stimulus.
Figure 4.
Changes (delta) in fetal heart rate associated with low to high levels of pCRH. The greatest change in response (heart rate acceleration above baseline) was observed in conditions of exposure to low levels of pCRH. No response was observed at median-high or high levels of pCRH.
Specific age-related patterns in peak fetal heart rate changes to tone stimulation. The oldest fetuses (36 weeks gestation) have the largest response followed by a progressive decrease in response (habituation). Slopes of the last four stimuli of the initial trials (trials 12–15, slope 2) were compared with the first four trials after the novel, dishabituating stimulus (trials 17–20, slope 3) providing support for more pronounced dishabituation in the oldest fetuses.

**Figure 5.**
Specific age-related patterns in peak fetal heart rate changes to tone stimulation. The oldest fetuses (36 weeks gestation) have the largest response followed by a progressive decrease in response (habituation). Slopes of the last four stimuli of the initial trials (trials 12–15, slope 2) were compared with the first four trials after the novel, dishabituating stimulus (trials 17–20, slope 3) providing support for more pronounced dishabituation in the oldest fetuses.
Figure 6.
Slopes of the FHR response to the first four tones at 36 weeks of gestation (slope 1), were divided into those who decreased (habitua tors) and those who increased (sensitzers). There was a trend that fetuses classified as habitua tors were exposed to lower CRH concentrations at 15 weeks gestation than the sensitizing fetuses.
Figure 7.
The percentage of women delivering preterm (before 37 weeks gestation) was greatest who were among the highest decile (90th %tile) in third trimester CRH.
Figure 8.
Model that describes our research program to assess plausible pathways between fetal exposure to adversity (including pCRH) and developmental outcomes.