

HHS Public Access

Neurotoxicol Teratol. Author manuscript; available in PMC 2022 November 01.

Published in final edited form as:

Author manuscript

Neurotoxicol Teratol. 2021; 88: 107033. doi:10.1016/j.ntt.2021.107033.

Co-occurrence of preconception maternal childhood adversity and opioid use during pregnancy: Implications for offspring brain development

Madeleine Allen^a, Nora K. Moog^b, Claudia Buss^{b,c}, Elizabeth Yen^f, Hanna Gustafsson^a, Elinor L. Sullivan^{a,d,e}, Alice M. Graham^a

^aDepartment of Psychiatry, Oregon Health & Science University, 3181 SW Sam Jackson Park Rd., Portland, OR, 97239, United States

^bCharité – Universitätsmedizin Berlin, corporate member of Freie Universität Berlin and Humboldt-Universität zu Berlin, Institute of Medical Psychology, Luisenstrasse 57, 10117 Berlin, Germany

^cDevelopment, Health and Disease Research Program, University of California, Irvine, 837 Health Sciences Drive, Irvine, California, 92697, United States

^dDivision of Neuroscience, Oregon National Primate Research Center, 505 NW 185th Ave., Beaverton, OR, 97006

^eDepartment of Behavioral Neuroscience, Oregon Health & Science University, 3181 SW Sam Jackson Park Rd., Portland, OR, 97239, United States

^fDepartment of Pediatrics, Tufts Medical Center, Boston, MA, 02111, United States.

Abstract

Understanding of the effects of *in utero* opioid exposure on neurodevelopment is a priority given the recent dramatic increase in opioid use among pregnant individuals. However, opioid abuse does not occur in isolation—pregnant individuals abusing opioids often have a significant history of adverse experiences in childhood, among other co-occurring factors. Understanding the specific pathways in which these frequently co-occurring factors may interact and cumulatively influence offspring brain development *in utero* represents a priority for future research in this area. We highlight maternal history of childhood adversity (CA) as one such co-occurring factor that is

Correspondence: grahaal@ohsu.edu (Alice M. Graham, PhD).

Financial disclosures for the manuscript entitled, "Co-occurrence of preconception childhood trauma history and opioid use during pregnancy: Implications for offspring brain development."

Ms. Allen reported no biomedical financial interests or potential conflicts of interest.

Dr. Moog reported no biomedical financial interests or potential conflicts of interest.

Dr. Buss reported no biomedical financial interests or potential conflicts of interest.

Dr. Yen reported no biomedical financial interests or potential conflicts of interest.

Dr. Gustaffson reported no biomedical financial interests or potential conflicts of interest.

Dr. Sullivan reported no biomedical financial interests or potential conflicts of interest.

Dr. Graham reported no biomedical financial interests or potential conflicts of interest.

Publisher's Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.

more prevalent among individuals using opioids during pregnancy and which is increasingly shown to affect offspring neurodevelopment through mechanisms beginning *in utero*. Despite the high incidence of CA history in pregnant individuals using opioids, we understand very little about the effects of comorbid prenatal opioid exposure and maternal CA history on fetal brain development. Here, we first provide an overview of current knowledge regarding effects of opioid exposure and maternal CA on offspring neurodevelopment that may occur during gestation. We then outline potential mechanistic pathways through which these factors might have interactive and cumulative influences on offspring neurodevelopment as a foundation for future research in this area.

Keywords

maternal childhood adversity; prenatal opioid exposure; *in ntero* exposure; opioid epidemic; brain development; maternal-placental fetal biology

1 - Introduction

The marked increase in opioid misuse in recent decades (Cicero et al., 2014; Substance Abuse and Mental Health Services Administration (US) and Office of the Surgeon General (US), 2018), including high rates of usage among pregnant individuals (Desai et al., 2014; Maeda et al., 2014; Terplan et al., 2010), has raised concern regarding effects of *in utero* opioid exposure on offspring brain development. (Abdel-Latif et al., 2013; Camden et al., 2021; Maeda et al., 2014). Advancing understanding of how in utero opioid exposure influences fetal brain development is of critical importance-the prenatal period is the most rapid period of brain development across the lifespan, and biological signaling mechanisms guiding embryonic and fetal brain development are highly sensitive to cues from the extrauterine environment (Entringer et al., 2015). Furthermore, alterations to brain development during gestation have implications for adaptive potential in response to the postnatal environment (Hartman and Belsky, 2018) and long-term risk for psychiatric disorders (O'Donnell and Meaney, 2016). While there is some empirical evidence in support of poorer physiological and neurodevelopmental outcomes soon after birth in infants exposed to opioids in utero versus non-exposed peers (Beckwith and Burke, 2015; de Cubas and Field, 1993; McGlone and Mactier, 2015; Moe, 2002; Nygaard et al., 2016, 2015; Ornoy et al., 2001), these observations have not been consistently replicated (Baldacchino et al., 2014), and long-term effects of prenatal opioid exposure on neurodevelopment remain poorly understood (Conradt et al., 2018).

A significant challenge in understanding the short- and long-term effects of prenatal opioid exposure lies in disentangling the potential influence of prenatal opioid exposure from the multiple, frequently co-occurring factors in the pre- and postnatal environment with potential to influence neurodevelopment. Previous literature in this area does not thoroughly account for the wide array of co-occurring factors prevalent in pregnant individuals using opioids (Conradt et al., 2019). Focusing on the potential mechanisms by which *in utero* opioid exposure and commonly co-occurring risk factors in the prenatal environment influence neurodevelopmental trajectories has potential to increase clarity into

the etiology of poor neurodevelopmental outcomes with implications for prevention and early intervention strategies.

Among pregnant individuals using opioids, there is an increased prevalence of exposure to childhood adversity (CA), which is a well-established risk factor for substance use (Brown and Shillington, 2017; Racine et al., 2020). Here we consider CA to include a range of negative experiences in childhood shown to have long-term consequences for psychiatric and other health outcomes. These include experiences involving threat (physical abuse, sexual abuse, exposure to domestic and community violence), deprivation (emotional or physical neglect), and instability (loss of a caregiver, caregiver transitions, family conflict and divorce) (Koss and Gunnar, 2018; McLaughlin et al., 2014). The number of CA exposures (also referred to as Adverse Childhood Experiences) proportionally increased the odds of opioid misuse in adulthood in a study of a large community sample while controlling for a variety of other sociodemographic factors (Merrick et al., 2020; Quinn et al., 2019). Additionally, childhood abuse and neglect were shown to increase the risk for adulthood prescription opioid misuse in a large nationally-representative sample (Austin and Shanahan, 2018). Among adults receiving treatment for opioid use disorder, CA is correlated with increased likelihood of relapse (Derefinko et al., 2019), younger age of opioid use initiation, more recent injection drug use, and increased likelihood of overdosing (Stein et al., 2017). The high co-occurrence of CA and opioid misuse is of particular relevance for advancing understanding of fetal brain development in the context of maternal opioid use given recent evidence for effects of maternal CA on maternal-placental-fetal (MPF) biology and fetal brain development (Buss et al., 2017; Hendrix et al., 2020; Lehrner and Yehuda, 2018; Moog et al., 2018, 2016a).

The current review aims to provide a foundation for examining the potential cumulative and interactive influences of maternal history of CA and opioid use during pregnancy on offspring neurodevelopment. We note that neonatal opioid withdrawal syndrome (NOWS) is a common focus both in clinical settings and in prior research on in utero opioid exposure (Bakhireva et al., 2019; Kelty and Preen, 2019; Kocherlakota, 2014). However, NOWS is not a primary focus of this manuscript given the lack of understanding of its role in neurodevelopment and long-term outcomes (Conradt et al., 2019). Further, NOWS diagnoses are variable, and a sizable portion of infants exposed to opioids in utero do not receive a NOWS diagnosis (Jones et al., 2018)-we aim to include all opioid-exposed neonates in this review to ensure a comprehensive examination of *in utero* opioid exposure. We first provide a brief review of current knowledge regarding the effects of prenatal opioid exposure and maternal CA on the developing brain *in utero*. We highlight research with potential to disentangle prenatal versus postnatal effects of prenatal opioid exposure and maternal CA, given that transmission of these risk factors may occur prenatally via alterations to MPF biology and epigenetic mechanisms, as well as through alterations to the postnatal environment. This literature is then contextualized with a discussion of the candidate mechanistic pathways by which opioids and sequelae related to maternal CA can influence offspring brain development *in utero*. Finally, we make recommendations for future research with the aim of increasing clarity regarding the implications of maternal opioid use during pregnancy on offspring neurodevelopment in the context of accompanying risk factors. We speak to the potential relevance of such research for eventually informing

public health messaging and the stigma related to opioid use during pregnancy, in addition to prevention and intervention efforts for individuals at risk for misusing opioids.

2 – Neurodevelopmental outcomes associated with *in utero* opioid exposure and preconception maternal CA

2.1 – Long-term neurodevelopmental outcomes associated with *in utero* opioid exposure and maternal preconception CA

Given the focus of this review on mechanistic pathways for the *in utero* effects of opioid exposure and maternal CA on human offspring neurodevelopment, clinical research examining the effects of both factors on neonatal human brain outcomes represent a primary interest due to enhancing capacity for distinguishing pre- versus postnatal effects. However, it is worth noting that research with animal models, which allows for experimental control to isolate effects of specific exposures during certain developmental windows, has identified long-term impacts of *in utero* opioid exposure and maternal CA on a range of outcomes beyond the neonatal period. While it is challenging to differentiate the role of the pre- versus post-natal environment in clinical studies investigating long-term neurodevelopmental outcomes of *in utero* opioid exposure and maternal CA, these studies suggest the potential for neurodevelopmental alterations related to these exposures to persist well beyond infancy into later childhood and adulthood. We therefore provide a brief review of literature examining long-term neurodevelopmental alterations in relation to these factors to contextualize the impetus for advancing understanding of the relevant mechanistic pathways leading to these alterations.

Pre-clinical studies suggest that long-term neurodevelopmental alterations in offspring exposed to opioids *in utero* include increased risk for behaviors analogous to symptoms of mood and anxiety disorders (Ahmadalipour et al., 2015; Hung et al., 2013; Wu et al., 2020), altered social and reward processing (Buisman-Pijlman et al., 2009b; Hol et al., 1996; Niesink et al., 1996; Vanderschuren et al., 1995; Vathy and Katay, 1992), cognitive differences (Ahmadalipour et al., 2015; Niu et al., 2009; Šlamberová et al., 2003; Wang et al., 2017; Wang and Han, 2009), in addition to increased seizure potential and alterations in endogenous opioid system and endocrinal stress-response functioning (Boggess and Risher, 2020; Byrnes and Vassoler, 2018). These studies primarily employ rodents (rats or mice), with the exception of one study which used chicks (Wang et al., 2017). Despite significant differences in brain morphology and the timing of neurodevelopment between rodents and humans (Ohmura and Kuniyoshi, 2017), there is some evidence to support potential translation to humans. This includes reports of emotional challenges and cognitive deficits in toddlers and school-aged children exposed to opioids during gestation (Levine and Woodward, 2018; Nelson et al., 2020; Nygaard et al., 2016, 2015; Yeoh et al., 2019), although these findings have not been consistently replicated (Bakhireva et al., 2019; Conradt et al., 2019).

Several factors require consideration in translating findings from animal models of prenatal opioid exposure. First, the role of the postnatal environment on offspring development can be challenging to model in animal studies. Animal models that include variation

in the postnatal environment indicate that the role of the postnatal environment plays a significant role in developmental trajectories of offspring exposed to opioids *in utero*. Some evidence suggests that effects of prenatal opioid exposure on offspring neurodevelopment may not persist in a beneficial postnatal environment—adult rodents exposed to opioids during pregnancy showed depressive-like behaviors that were prevented with postnatal environmental enrichment (Ahmadalipour et al., 2015) and exercise (Wu et al., 2017). These rodent models suggest an important modulatory role of the postnatal environment on neurodevelopment in offspring exposed to opioids *in utero*, although studies examining the role of the early postnatal environment on rodent development must be translated to human findings with caution. Rodents are born at the equivalent of about mid-gestation in human fetuses (Clancy et al., 2007), implying that early postnatal factors may have differential effects on brain development in rodents than in humans.

Evidence from clinical research also emphasizes the need to consider postnatal environmental influences on neurodevelopment in offspring exposed to opioids in utero. Compared to children without prenatal substance exposure, children exposed to opioids in *utero* are more likely to experience a range of adversities in the postnatal environment, including abuse, neglect, housing instability, low socioeconomic status (SES), poor nutrition, parental psychopathology, low access to healthcare, and caregiver disruptions (Conradt et al., 2018; Levine et al., 2021). While consideration of such co-occurring adversities is rare in studies examining the effects of prenatal opioid exposure on long-term neurodevelopment in humans (Conradt et al., 2018), those that attempt to control for factors such as low SES, maternal education level, and the quality of the home environment have found that the detrimental effects of prenatal opioid exposure on cognitive and psychomotor development were no longer evident (Hans and Jeremy, 2001; Messinger et al., 2004). A recent study by Levine et al. (2021) similarly reported that deficits in motor development, cognitive development, and emotional and behavioral dysregulation found in 2-year-old children with prenatal methadone exposure were mediated by gestational age at birth, and aspects of the postnatal environment, including, breastfeeding participation, maternal stress, and punitive parenting. They also found that the children exposed to methadone in utero demonstrated deficits in language development at 2 years old that did not remain significant after controlling for maternal education and polysubstance use during pregnancy (Levine et al., 2021). This work highlights the role of the postnatal environment in the developmental trajectories of offspring exposed to opioids in utero.

Similar to human studies of children exposed to opioids *in utero* (Boggess and Risher, 2020; Conradt et al., 2019; Kirkegaard et al., 2020; Skumlien et al., 2020; Winklbaur et al., 2009), offspring of mothers who experienced CA are at increased risk for a myriad of poor neurodevelopmental and physiological outcomes (Buss et al., 2017). In childhood, offspring with maternal CA exposure show increased physiological anxiety markers (Jovanovic et al., 2011), higher rates of obesity and smoking behaviors (Roberts et al., 2014), increased externalizing behaviors (Miranda et al., 2013; Myhre et al., 2014; Plant et al., 2017; Rijlaarsdam et al., 2014), increased negative emotionality (Bouvette-Turcot et al., 2015), and increased risk for autism (Roberts et al., 2013). While several of these studies identified postnatal mediators of the relationship between maternal CA and offspring outcomes (Collishaw et al., 2007; Miranda et al., 2013; Plant et al., 2017; Rijlaarsdam et al., 2014;

Roberts et al., 2014), others also indicated co-occurring factors in the prenatal environment that modulate the association between poorer long-term neurodevelopmental outcomes and maternal CA history, such as antenatal depression (Collishaw et al., 2007; Plant et al., 2017) and genetic variation (Bouvette-Turcot et al., 2015). Limited pre-clinical studies examining the effect of parental early life adversity on offspring neurodevelopment partially support findings from clinical studies, suggesting either maternal or paternal early life adversity may increase the risk for biological and behavioral stress phenotypes in mice and primate offspring (Cowan et al., 2016). These findings further emphasize the need to examine offspring neurodevelopmental alterations at birth in order to identify alterations related to prenatal opioid exposure and maternal CA that truly occur before exposure to the postnatal environment.

2.2 – Offspring neonatal brain outcomes associated with *in utero* opioid exposure and maternal preconception CA

2.2.1 – **Whole-brain outcome measures**—*In utero* opioid exposure and maternal CA have both been associated with altered neonatal head circumference, which is thought to be associated with smaller intracranial, or brain, volume (Cheong et al., 2008; Lindley et al., 1999), suggesting that these exposures may impede or alter fetal brain growth during gestation. Head circumference at birth is often utilized as a predictor for later neurodevelopment—neonates with head circumferences at the top or bottom 2% are at significantly greater risk for neurodevelopmental disorder diagnosis later in life (Wright and Emond, 2015).

Prenatal opioid exposure has been associated with reduced neonatal head circumference (Craig et al., 2020; Monnelly et al., 2018; Peterson et al., 2020; Towers et al., 2019; Visconti et al., 2015). Additionally, two studies directly examining intracranial volume using MRI techniques identified global volumetric reductions in brains of neonates exposed to opioids in utero compared to unexposed controls (Peterson et al., 2020) and previous literature documenting normative neonatal brain development (Yuan et al., 2014). However, consideration of co-occurring factors has been limited in these studies. Additionally, the type of opioid exposure is likely to influence fetal head growth. Methadone and buprenorphine are two opioid maintenance treatments that are currently the frontline treatments for pregnant individuals with opioid dependence (World Health Organization, 2014). Both drugs have affinity for the µ-opioid receptor (MOR), and reduce mortality rates and risk of relapse associated with opioid misuse due to their long half-life (methadone) and partial agonist (buprenorphine) properties (Bonhomme et al., 2012). Of these two opioid maintenance treatments, only methadone appears to be associated with smaller neonatal head circumference (Jones et al., 2014, 2010; O'Connor et al., 2019; Pritham et al., 2012; Zedler et al., 2016). Poly-substance use during pregnancy, which is very common among opioid-using pregnant individuals, may exacerbate reductions in offspring head circumference associated with prenatal opioid exposure-among prenatally opioid-exposed infants, additional exposure to tobacco has been associated with further reduced head circumference (Winklbaur et al., 2009). Limited evidence from clinical literature suggests that reduction in neonatal head circumference related to prenatal opioid exposure does not appear to be dose-dependent (Gray et al., 2010; O'Connor et al., 2019), although this will

need to be replicated in future studies. Smaller cerebral size at birth is associated with poorer intellectual and executive functioning outcomes, particularly when subsequent postnatal head growth does not catch up to peers (Aagaard et al., 2018; Bilder et al., 2013; Ferrer et al., 2019; Gale et al., 2006; Heinonen et al., 2008; Kirkegaard et al., 2020; Langridge et al., 2013; Wright and Emond, 2015).

Maternal CA has also been associated with alterations in global cerebral development at birth, however interpretation of findings is limited by the sparsity of research in this area. One study reported an association between maternal CA history and a higher cephalization index (ratio of head circumference to body weight) at birth when adjusting statistically for factors related to maternal demographics, maternal health, delivery, infant sex assigned at birth, and infant gestational age at delivery (Appleton et al., 2019). These findings are consistent with reports of a higher offspring cephalization index linked to earlier age of maternal menarche (Holdsworth and Appleton, 2020), given that exposure to CA is associated with earlier pubertal onset (Colich et al., 2020; Holdsworth and Appleton, 2020; Lei et al., 2018; Noll et al., 2017). Another group demonstrated positive correlations between maternal CA and offspring head circumference and weight at birth, while accounting for multiple maternal psychosocial, health, and nutritional factors (Apanasewicz-Grzegorczyk et al., 2020). Additionally, maternal CA has been associated with reduced neonatal cortical gray matter, which contributed to an overall smaller intracranial volume (Moog et al., 2018). These results persisted after adjusting for multiple potential confounds frequently associated with maternal history of CA, including SES, complications during pregnancy, obesity, recent exposure to interpersonal violence, stress throughout pregnancy and in the early postpartum period, and length of gestation (Moog et al., 2018).

Overall, there appears to be potential for both prenatal opioid exposure and maternal CA to influence fetal cerebral growth, suggesting potential for cumulative or interactive influences to be examined in future studies. In addition, future research would benefit from examining trajectories of postnatal intracranial growth to see if differences persist over time, with consideration of additional moderating factors (including environmental risks and weight at birth).

2.2.2 - Outcomes in large-scale brain networks and global connectivity-

Examining connectivity both with resting state functional MRI and diffusion tensor imaging is of great interest for understanding the long-term effects of fetal neurodevelopment, as key aspects of adult brain organization, such as small worldness (component of brain network organization consisting of close and highly interconnected nodes) and nacent forms of large-scale brain networks, are detectable during the neonatal period and are predictive of neurodevelopment throughout childhood (De Asis-Cruz et al., 2020; Graham et al., 2021; Schneider et al., 2004; Smyser et al., 2010; van den Heuvel et al., 2015). Some evidence suggests that prenatal opioid exposure is related to altered maturation of white matter fiber tracts in the neonatal brain. Monnelly et al. (2018) reported decreased fractional anisotropy (FA), a measure of white matter integrity and connection orientation, in the neonatal white matter skeleton, which represents the center of each white matter tract common to the sample. Additionally, Walhovd et al., (2012) found increased mean diffusivity (MD), indicative of reduced white matter integrity, in the superior longitudinal fasciculi of neonates

exposed to opioids during pregnancy. Furthermore, Merhar et al., (2019) found preliminary evidence of increased risk of white matter lesions and abnormalities in neonates prenatally exposed to opioids. In contrast, while controlling for gestational age, offspring sex, tobacco and alcohol use, maternal age, SES, race/ethnicity, depression, anxiety, and prenatal stress, a small cohort study found that frontal and parietal white matter in prenatally opioid-exposed neonates showed increased FA and reduced MD (Peterson et al., 2020). These studies have attempted to account for key pre- and post-natal environmental covariates, but maternal CA history has not been considered either as a covariate or moderator. Moreover, small sample sizes (ranging from 20 to 60 participants) and lack of replication in independent datasets decrease confidence in these findings, particularly in light of the expected small effect sizes and recent literature highlighting the lack of reproducibility in neuroimaging studies (Boekel et al., 2015; Kharabian Masouleh et al., 2019; Ks et al., 2013; Marek et al., 2020).

Utilizing resting state functional connectivity MRI has become increasingly popular in neonatal brain development research, as it takes advantages of brain activity at rest and can reveal neonatal functional brain network activity with potential implications for longterm neurodevelopment. Recent work by Salzwedel et al. (2020) employing resting state functional connectivity included a larger sample size relative to prior studies (n=133), although only 18 infants in the study had *in utero* opioid exposure. However, examining the potential link between alterations in neonatal brain connectivity and subsequent cognitive development, and employing multivariate modeling to account for co-occurring prenatal influences (including polysubstance exposure and maternal depression, although not maternal CA history) represent significant strengths of the study. The findings indicate an association between prenatal opioid exposure and alterations in neonatal resting state functional connectivity of the left middle frontal and right angular gyrus, as well as the cingulo-opercular network. These alterations in the neonatal brain were not associated with cognitive, language, or motor composite scores at 3 months of age. Thus, while this study suggests that prenatal opioid exposure may be associated with alterations in developing brain systems involved in cognition and executive functioning, it is unclear if these effects persist long after birth.

Maternal CA has not been examined in relation to offspring neonatal global brain connectivity. Alterations in region-specific connectivity (Hendrix et al., 2020) and global gray matter differences (Moog et al., 2018) in neonates exposed to maternal CA, in addition to functional connectivity differences in school-aged children with maternal CA (Zhang et al., 2021), suggest that the effects of maternal CA on neonatal functional connectivity may be an area of interest for future research.

2.2.3 - Region-specific volume and connectivity outcomes—Additional

findings from neonatal brain imaging reveal that prenatal opioid exposure is associated with volumetric alterations of multiple cortical and subcortical brain regions with potential implications for basic motor and sensory processing and integration, as well as higher-order cognitive and emotional processing. One pilot study demonstrated decreased volume of the basal ganglia and larger volume of the lateral ventricle in neonates exposed to buprenorphine or methadone during pregnancy (Yuan et al., 2014). Additionally, research employing mid-pregnancy ultrasounds (18-22 weeks gestation) identified increased thalamic diameters in

fetuses exposed to opioids in utero (Schulson et al., 2014). A pilot study utilized MRI in fetuses at approximately 33 weeks of gestation to identify reduced anteroposterior diameter of the cerebellar vermis in fetuses exposed to opioids in utero compared to unexposed fetuses (Radhakrishnan et al., 2021a). Peterson et al. (2020) also found alterations in brain volume across multiple cortical regions, with increases observed in the middle temporal and inferior frontal gyri, posterior cingulate cortex, and inferior medial prefrontal cortex (mPFC), and decreases in the middle frontal gyrus, orbitofrontal cortex, and some dorsal and lateral regions of the prefrontal cortex. A recent pilot study identified that prenatally opioidexposed neonates showed increased right amygdala-mPFC connectivity and decreased amygdala connectivity to other regions within the medial temporal lobe while controlling for maternal depression and infant sex (Radhakrishnan et al., 2021b). These findings indicate potential alterations in the cerebellum, multiple subcortical brain regions involved in sensory processing, movement, arousal, stress and emotion processing and reactivity, and distributed cortical brain regions, including those involved in higher level cognitive processes. However, limitations in sample size and a lack of replication in independent samples again hamper the generalizability of these findings.

The literature examining maternal CA in relation to specific brain volumes and connectivity in the neonatal period is even more limited. However, one recent study observed an association between maternal childhood emotional neglect and stronger offspring neonatal functional connectivity between the amygdala and dorsal anterior cingulate cortex, as well as between the amygdala and ventromedial prefrontal cortex (Hendrix et al., 2020). Increasing sample sizes, testing for replication in independent data sets, and considering potential cumulative and interactive effects of maternal CA and opioid use, among other co-occurring factors, represent important next steps in advancing understanding of whether coordinated functioning of the amygdala with regions of the prefrontal cortex, and other early neural phenotypes, play a role in mediating effects of these prenatal exposures on subsequent development.

2.2.4 – Summary of neonatal brain outcomes associated with *in utero* opioid exposure and maternal preconception CA—The findings to date indicate that both maternal CA and opioid use during pregnancy have potential to influence offspring brain development *in utero*. Prenatal opioid exposure appears to be associated with several neurodevelopmental alterations at birth, including smaller cerebral size, white matter abnormalities, alterations to functional connectivity networks, and volumetric changes to various brain regions. However, there are substantial limitations to existing literature examining the neurodevelopmental effects of prenatal opioid exposure during the fetal and neonatal period, including small sample sizes and lack of replication of findings, making it particularly difficult to draw conclusions about specific functional connectivity or volume changes. A significant challenge in studying the effects of prenatal opioid exposure in humans is how varying dosages, timings, frequencies, and specific opioid compounds will alter fetal neurodevelopment, given the substantial variability of each of these factors in the population of pregnant individuals using opioids. Furthermore, we continue to lack understanding of long-term neurodevelopmental consequences of prenatal opioid exposure in humans, and our review did not identify evidence to support the commonly held belief

that alterations at birth associated with prenatal opioid exposure will persist into later childhood and adulthood.

Additionally, studies examining potential neonatal brain alterations associated with maternal CA are currently very rare. To our knowledge, only four studies examine how maternal CA influences offspring brain outcomes at birth: two studies indicating a higher cephalization index (Apanasewicz-Grzegorczyk et al., 2020; Appleton et al., 2019), one reporting smaller gray matter and overall intracranial volumes (Moog et al., 2018), and one showing altered frontoamygdala connectivity (Hendrix et al., 2020). While understanding the specific effects of maternal CA on fetal neurodevelopment will be challenging without replication of this work, the current evidence does suggest that maternal CA alters offspring neurodevelopmental trajectories during gestation, and this is a compelling area for further research. Beyond increasing sample sizes to increase likely reproducibility of findings, future investigations into the effects of both prenatal opioid exposure and maternal CA on offspring neurodevelopment would benefit from study designs which carefully balance major potential confounds and provide sufficient statistical power to model cumulative and interactive influences.

3 - Pathways by which in utero opioid exposure and maternal

preconception CA influence fetal brain development

3.1 - Endogenous opioid system

The endogenous opioid system is a complex neuromodulatory system that consists of several families of peptides and receptors that influence a wide range of behavioral and biological processes, including pain, reward processing, stress responsivity, cell survival, respiratory depression, ionic homeostasis, digestion, euphoria, cardiovascular health, and sedation (Fricker et al., 2020; Shenoy and Lui, 2020). Opioid peptides, including enkephalins, endorphins, dynorphins, nociceptin and endomorphins, have distinct effects depending on the regional and developmental context, as well as their differential affinities for the opioid receptors, including the μ -opioid receptor (MOR), δ -opioid receptor (DOR), κ -opioid receptor (KOR), nociceptin receptor (NOR), and zeta opioid receptor (ZOR) (Dhaliwal and Gupta, 2021; Fricker et al., 2020). The endogenous opioid system is distributed throughout the body, although it is particularly active in the central and peripheral nervous systems (Fricker et al., 2020).

During the prenatal period, the endogenous opioid system is thought to play a unique modulatory role in fetal brain development. Rodent studies suggest that developing neural cells begin to produce opioid receptors and opioid peptides early in gestation (Farid et al., 2008; Hauser and Knapp, 2018), while limited studies in humans verify endogenous opioid system activity by 11 (Tripathi et al., 2008) and 20 (Kinney et al., 2008; Magnan and Tiberi, 1989; Wang et al., 2006) weeks of gestation. Endogenous opioids modulate a range of early fetal neurodevelopmental processes involved with neuronal and glial maturation, and some opioid peptide activity appears to be reserved for developmental processes alone (Farid et al., 2008; Hauser and Knapp, 2018). While there are exceptions, opioid receptor activation tends to suppress fetal brain growth via inhibition of neuronal and glial proliferation

and differentiation, in addition to increased neuronal cell death, although specific activity differs by brain region and type of opioid receptor involved (Hauser and Knapp, 2018). One significant exception is the association of endogenous opioid activity with growth of oligodendrocytes, glial cells responsible for myelination. MOR activation during the prenatal period leads to increased mitosis of immature oligodendrocytes, which do not yet produce myelin (Knapp et al., 1998; Knapp and Hauser, 1996), while KOR agonists promote embryonic myelin production and mature oligodendrocyte differentiation and proliferation (Knapp et al., 2009, 2001; Mei et al., 2016), at the cost of neuron and astrocyte genesis (Hahn et al., 2010).

Opioid drugs (also referred to as exogenous opioids) with addiction potential take effect by mimicking endogenous opioid peptides and acting on opioid receptors throughout the body, both prenatally via placental transfer (Hauser and Knapp, 2018) and postnatally (Davis and Pasternak, 2005). When used during pregnancy, exogenous opioids travel across the placenta in significant amounts, reaching drug equilibrium between the pregnant individual and the fetus (Gerdin and Lindberg, 1990; Griffiths and Campbell, 2015). While developing fetal brain cells often appear to transiently express opioid receptors at different concentrations throughout gestation (Hauser and Knapp, 2018), some evidence suggests that certain neural cells are more sensitive to exogenous opioids during parts of the prenatal period than in adulthood—rat fetal neurons have been found to bind to methadone at a rate 2-14 times higher than in adults (Pertschuk et al., 1977). The potential for exogenous opioids to exaggerate the differential endogenous opioid activity on the development of various neural cell types, including promotion of oligodendrocyte growth and inhibition of neuron and astrocyte growth, suggests a potential pathway for reduced overall head circumference (Craig et al., 2020; Monnelly et al., 2018; Peterson et al., 2020; Towers et al., 2019; Visconti et al., 2015) and intracranial volume (Peterson et al., 2020; Yuan et al., 2014), in addition to altered white matter structure in opioid-exposed neonates (Merhar et al., 2019; Monnelly et al., 2018; Peterson et al., 2020; Walhovd et al., 2012).

Additionally, animal models suggest that prenatal opioid exposure may have specific programing effects on the rapidly developing fetal endogenous opioid system (Byrnes and Vassoler, 2018). The effects of prenatal opioid exposure on postnatal endogenous opioid system functioning appear to vary depending on the brain region of interest, postnatal age, and hormonal factors (Byrnes and Vassoler, 2018). Overall, the majority of literature in this area suggests that rodent offspring chronically exposed to exogenous opioids during fetal development show reduced opioid receptor binding across specific brain regions in the early postnatal period, but increased receptor binding in the same regions in adulthood (Byrnes and Vassoler, 2018). However, this does not appear to be consistent across brain regions—adult animals prenatally exposed to morphine showed reduced MOR binding in the bilateral amygdala (Šlamberová et al., 2005) and the medial preoptic area (Vathy et al., 2003). Prenatal opioid exposure may also be associated with increased endogenous opioid release in brain regions important for reward processing (substantia nigra, piriform cortex, and septum) in adulthood (Buisman-Pijlman et al., 2009a). Additionally, whole-brain analyses in rats exposed to opioids in utero showed increased MOR expression and binding during the neonatal period, but not in adulthood (Bhat et al., 2006), further suggesting

that opioid receptor alterations induced by prenatal opioid exposure appear to vary across developmental periods.

Direct CA exposure has also been shown to permanently alter the endogenous opioid system, which may in turn alter offspring development during pregnancy (Vazquez et al., 2005). To our knowledge, there is only one study that examined the intergenerational effects of adversity on offspring endogenous opioid system functioning—non-stressed male offspring of female rats exposed to chronic stress (beginning post-weaning through adulthood) showed significantly decreased spinal cord MOR gene expression compared to non-stressed male rats without parental stress exposure (Hormozi et al., 2018). Spinal cord MOR gene expression in male offspring exposed to only paternal stress or both maternal and paternal stress did not significantly differ from unexposed male rats (Hormozi et al., 2018). These findings verify that maternal chronic stress, including stress over the course of development, does influence offspring endogenous opioid system functioning.

Direct exposure to early adversity in rodents is associated with alterations in mRNA expression of brain opioid receptors in a time-, region- and sex-specific manner (Nakamoto et al., 2020). In mice exposed to maternal separation and social isolation, an animal model of CA, expression of KOR, MOR and DOR mRNA in the periaqueductal gray area was reduced, but KOR mRNA in the amygdala was significantly increased (Nakamoto et al., 2020). Chronic lifetime stress was associated with decreased MOR mRNA in the spinal cord of adult male rats (Hormozi et al., 2018). Additionally, chronic stress in mice is associated with increased dynorphin release and subsequent increased KOR activation in the basolateral amygdala, nucleus accumbens (NAc), dorsal raphe, and hippocampus (Land et al., 2008). Increased KOR activation of serotonergic neurons in the dorsal raphe nucleus projecting to the NAc appear to mediate the aversive stress response (Land et al., 2009). A rat model of early life adversity reported alterations to KOR and dynorphin activity in the lateral habenula, a brain region associated with reward- and aversion-related learning and depression (Simmons et al., 2020). They reported that juvenile, adolescent, and adult rats exposed to early life adversity exhibited increased dynorphin levels and significantly decreased KOR mRNA expression in the lateral habenula compared to rats unexposed to early life adversity. Karkhanis et al. (2016) observed that compared to unexposed controls, adult rats exposed to chronic early life stress demonstrated differences in KOR and dynorphin activity in the NAc, including decreased dynorphin levels, increased KOR agonist-mediated inhibition of dopamine, and increased dopamine levels in response to a KOR antagonist. Chang et al. (2019) observed reductions in KOR and MOR mRNA in the NAc of neonatal female rats recently exposed to predator odor, but increased MOR and DOR mRNA in juvenile females exposed to predator odor during the neonatal period. In humans, postmortem brain tissue of individuals with a history of abuse who died by suicide revealed that KOR expression was significantly decreased in the anterior insula compared to controls—an effect that was not observed in suicide victims without CA exposure and that was associated with decreased DNA methylation in an intron of the KOR gene (Lovallo et al., 2018; Lutz et al., 2018).

Findings from both animal and human literature indicate that direct CA exposure appears to alter endogenous opioid system functioning, particularly KOR and dynorphin functioning

in brain regions associated with reward processing. If carried forward into pregnancy, these alterations have implications for fetal neurodevelopment via the mechanisms previously discussed. Moreover, among pregnant individuals using opioids, these alterations may interact with the dysregulating effects of exogenous opioids on the endogenous opioid system and thereby exacerbate effects on the developing fetal brain. Future research will need to examine how CA-induced alterations to endogenous opioid system functioning may alter offspring neurodevelopment during pregnancy, although preliminary evidence suggests that MOR gene expression in the spinal cord may be downregulated in offspring with maternal chronic lifetime stress exposure.

Alterations to the maternal endogenous opioid system related to preconception CA have potential to subsequently alter maternal hypothalamic-pituitary-adrenal (HPA) axis functioning (Areda et al., 2005; Bilkei-Gorzo et al., 2008; Brunton, 2019; Hale et al., 2003; Jaschke et al., 2021; Kudryavtseva et al., 2004; Marinelli et al., 2004; Yamamoto et al., 2003). In adults assigned female at birth (AFAB)¹ without CA exposure, administration of an opioid receptor antagonist, naltrexone, typically causes a strong increase in HPA axis activity; however this activation was suppressed in adults AFAB exposed to CA, suggesting that endogenous opioid modulation of the HPA axis is reduced in CA-exposed individuals AFAB (Lovallo et al., 2018). During pregnancy, the neurosteroid allopregnanolone, a metabolite of progesterone that increases in concentration during pregnancy, potentiates endogenous opioid inhibition of HPA axis reactivity (Brunton et al., 2009; Kammerer et al., 2002; Russell et al., 2008). Individuals that experienced CA show an exaggerated blunting of the HPA axis during pregnancy compared to pregnant individuals without preconception CA (Morrison et al., 2017), which can be mimicked in non-pregnant mice when they are administered allopregnanolone (Morrison et al., 2020). This provides further evidence that maternal CA alters endogenous opioid modulation of the HPA axis during pregnancy. Alterations in HPA axis functioning during pregnancy in turn have significant implications for fetal neurodevelopment, which is discussed further below.

3.2 – HPA axis

The HPA axis directs the body's physiological response to acute and chronic stress through the sequential release of corticotropin-releasing hormone (CRH), adrenocorticotropic hormone (ACTH), and glucocorticoids (cortisol in humans). During pregnancy, maternal cortisol levels stimulate placental corticotropin releasing hormone (pCRH) (Rehman et al., 2007; Sandman et al., 2006), which acts on the fetal adrenal gland to stimulate cortisol synthesis *in utero* (Sandman et al., 2012). Maternal cortisol also passes through the placenta, particularly in adverse contexts (Benediktsson et al., 1997). Cortisol plays an obligatory role in fetal neurogenesis, gliogenesis, synaptogenesis, and growth of axons and dendrites, creating ample opportunity for alterations in cortisol levels during pregnancy to exert an influence on fetal neurodevelopment (Matthews, 2000).

¹The terminology "assigned female at birth" (AFAB) and "assigned male at birth" (AMAB) are utilized in this review to acknowledge that human sex is a composite of many traits (i.e. chromosomes, genes, hormones, sex organs, and sex characteristics) that vary between individuals and may not align with sex assigned at birth (de Vries and Södersten, 2009; Keevil et al., 2017; Montañez, 2017). References may not have used these terms in their studies.

Neurotoxicol Teratol. Author manuscript; available in PMC 2022 November 01.

There has been increasing interest in the co-occurrence of opioid use and dysregulated HPA axis in pregnant individuals, as both factors may influence offspring development through overlapping and interacting pathways (Lester and Padbury, 2009; Pastor et al., 2017). Some evidence suggests that opioid use during pregnancy may increase glucocorticoid release—pregnant rats administered daily morphine had significantly elevated glucocorticoid levels compared to pregnant controls (Kazemi et al., 2011). A recent pilot study found that higher levels of hair cortisol concentrations in pregnant individuals using opioids were associated with less severe withdrawal symptoms in offspring at birth (Wachman et al., 2020). Given that chronic stress and extensive opioid use have been associated with blunted HPA axis activity, the authors hypothesized that lower maternal cortisol levels in this sample may correspond to increased chronic stress and adversity history, which may contribute to worse opioid withdrawal symptoms at birth (de Vries et al., 2020; Wachman et al., 2020; Zhou et al., 2010; Zhou and Leri, 2016). Additionally, if exogenous opioid use exaggerates endogenous opioid inhibition of the HPA axis during pregnancy, higher rates of opioid use would likely correspond to both decreased cortisol and worse offspring withdrawal symptoms at birth. Thus, while the evidence to date is relatively mixed, it suggests that opioid use during pregnancy impacts the maternal HPA axis with implications for programming the fetal brain and HPA-axis. Future studies will be needed to advance understanding of this topic.

CA is well-known to produce long-term alterations in endocrine stress physiology, including greater HPA axis reactivity as well as hypocortisolism, and current literature suggests that HPA alterations related to preconception CA carry forward into pregnancy (Heim et al., 2019). Pregnant individuals with CA history have been found to have lower baseline levels of cortisol immediately after waking, an elevated cortisol awakening response (the rapid increase in cortisol levels occurring shortly after waking), and a flattened diurnal slope (slope of decreasing cortisol levels throughout the day) (Bublitz et al., 2014; Bublitz and Stroud, 2012a; Shea et al., 2007; Thomas et al., 2018; Thomas-Argyriou et al., 2020), as well as increased concentrations of cortisol in hair during mid- to late-pregnancy compared to pregnant individuals without a history of CA (Schreier et al., 2015a; Swales et al., 2018). Additionally, childhood sexual abuse may have more pronounced effects on HPA axis alterations than other adversities—pregnant individuals with a history of childhood sexual abuse showed increasing cortisol awakening responses throughout pregnancy when compared to pregnant individuals with histories of non-sexual childhood abuse and neglect, but diurnal slope did not significantly differ between groups (Bublitz and Stroud, 2012a). Maternal exposure to CA has also been associated with a steeper increase of pCRH during the third trimester of pregnancy (Moog et al., 2016b; Steine et al., 2020).

HPA axis functioning alterations during pregnancy frequently observed in relation to CA and opioid use are likely to mediate offspring developmental alterations. Elevated cortisol levels early in pregnancy are associated with a greater increase in pCRH during the third trimester of pregnancy (Sandman et al., 2006), and high concentrations of pCRH during the third trimester of pregnancy are associated with preterm birth and a more difficult infant temperament (Davis et al., 2005; Wadhwa et al., 2004). Furthermore, infants exposed to elevated maternal cortisol in late pregnancy show increased behavioral challenges and negative temperament beginning at 1 week old (de Weerth et al., 2003) and increased parent-

reported infant negative reactivity at 2 months of age (Davis et al., 2007). Interestingly, elevated maternal cortisol levels early in gestation were associated with slower and poorer offspring cognitive development throughout the first year of life, while elevated maternal cortisol levels late in gestation were associated with accelerated and more advanced cognitive development over the first year of life (Davis and Sandman, 2010). In one recent study, a flatter diurnal slope during the first and second trimesters of pregnancy was predictive of internalizing behavior in children AFAB and externalizing behavior in children AMAB at 4 years of age (Thomas-Argyriou et al., 2020). Additionally, higher average cortisol awakening response in individuals at any point in pregnancy mediated the association of maternal CA history with offspring internalizing, but not externalizing, problems at 4 years of age. Alterations to HPA axis activity that are typically observed in pregnant individuals with CA histories, particularly elevated cortisol levels and high late-gestation pCRH, appear to alter offspring developmental trajectories. Some evidence suggests that elevated cortisol during early-, mid-, and late-gestation differentially alters offspring development, but further research in this area will be needed to clarify these findings.

While it is challenging to definitively predict how maternal CA and opioid use during pregnancy may interact to alter maternal HPA axis functioning and subsequent offspring outcomes, both factors do appear to independently and uniquely alter maternal HPA axis activity. Given that the maternal HPA axis plays a prominent role in offspring neurodevelopment during pregnancy (Matthews, 2000), it will be important to investigate how prenatal opioid exposure may differentially alter fetal brain development in combination with frequently co-occurring factors, such as maternal CA, through alterations to maternal HPA axis fetal programming.

3.3 - Inflammation

Both opioid use and CA have repeatedly been shown to influence immune system functioning, and specifically to contribute to a pro-inflammatory phenotype in adulthood (Baumeister et al., 2016; Buchanan et al., 2010; Hutchinson et al., 2011; Lacagnina et al., 2017; Wang et al., 2012; Zhang et al., 2020), which appears to persist during pregnancy (Boeck et al., 2016; Moog et al., 2016a). Emerging preclinical data have shown nonneuronal actions of opioids on glial cells that activate pro-inflammatory cascades, including: elevated tumor necrosis factor alpha (TNF- α), interferon gamma (IFN- γ), interleukin 1 beta (IL1- β), interleukin 6 (IL-6), chemokine ligand 4 (CCL4), chemokine ligand 16 (CCL16) (Buchanan et al., 2010; Hutchinson et al., 2007; Wang et al., 2012; Zhang et al., 2020). Furthermore, heightened systemic inflammation secondary to exogenous opioids may create a feedback loop with neuroinflammation leading to increased opioid-seeking behavior. Specifically, opioid-induced activation of glial cells appears to enhance the analgesic and rewarding effects of opioids, and contribute to opioid tolerance (Arezoomandan et al., 2016; Bachtell et al., 2015; Hutchinson et al., 2012; Narita et al., 2006; Song and Zhao, 2001; Zhang et al., 2012).

In pregnant individuals, CA is associated with higher IL-6 and of CRP concentrations (Finy and Christian, 2018; Mitchell et al., 2018). These associations are partially moderated

by nutritional factors (Hantsoo et al., 2019a; McCormack et al., 2020) and depressive symptoms (McCormack et al., 2020; Walsh et al., 2016) and may be mediated by prepregnancy body-mass index (BMI) (Finy and Christian, 2018; Mitchell et al., 2018). A small study in pregnant individuals with gestational diabetes observed elevated IL-15 levels in association with a history of CA (Bublitz et al., 2017).

The potential for opioid use and maternal CA to independently lead to heightened maternal inflammation both preconception and during pregnancy has significant implications for programming offspring neurodevelopment. A host of animal and increasingly human literature have demonstrated the effects of heightened maternal inflammation during pregnancy on the developing fetal brain (Graham et al., 2018; Hantsoo et al., 2019b; Jantzie et al., 2020; Rasmussen et al., 2019; Rudolph et al., 2018; Yoon et al., 1997), infant and toddler socioemotional and cognitive development (Graham et al., 2018; Gustafsson et al., 2018; Rudolph et al., 2018), and subsequent risk for higher rates of neuropsychiatric disorders including schizophrenia, autism, ADHD, and obsessive-compulsive disorder (Gustafsson et al., 2020; Hantsoo et al., 2019b). While direct transfer of pro-inflammatory cytokines across the placenta seems to be limited (Aaltonen et al., 2005), maternal immune activation may indirectly increase cytokine concentrations in the fetal compartment via placental cytokine production (Ashdown et al., 2006; Urakubo et al., 2001). Importantly, the same pro-inflammatory cytokines activated by exogenous opioids (IL-6 and TNF- α) and maternal CA (IL-6 and CRP) have been identified in studies linking maternal inflammation during pregnancy to offspring neurobehavioral development (Coelho et al., 2014; Graham et al., 2018; Gustafsson et al., 2018; Rudolph et al., 2018).

Beyond pathways involving maternal inflammation during pregnancy, exogenous opioid transfer across the placenta and the fetal blood-brain-barrier may influence fetal neuroinflammation through similar pathways as adults after the initial development of opioid receptors at 9-10 weeks of gestation (Byrnes and Vassoler, 2018; Farid et al., 2008; Gerdin and Lindberg, 1990; Griffiths and Campbell, 2015). In rodents, prenatal methadone exposure has been associated with heightened levels of systemic inflammation (TNF- α , IL-1 β , IL-6, and chemokine CXC ligand 1 (CXCL1)) in pups at postnatal day 10 (roughly 40-42 weeks of postconceptional age in human neurodevelopment), but the majority of these markers returned to baseline at postnatal day 21 (roughly 9 months old in human neurodevelopment) (Jantzie et al., 2020). However, heightened inflammation specifically in 21-day-old offspring brains prenatally exposed to methadone was evidenced by elevated cytokine levels (TNF-a, IL-6, toll-like receptor 4 (TLR4), and myeloid differentiation primary response protein (Myd88)), reduced glial cell branching, and differences in structural tract formation (Jantzie et al., 2020). While further research is needed in animal models and humans to better understand the pathways through which opioid use may influence maternal inflammation, MPF biology and ultimately offspring brain and immune system functioning, findings to date indicate multiple potential pathways for such effects.

Overall, these findings indicate the potential for a history of CA combined with both pre-pregnancy and pregnancy opioid use to result in a significantly heightened inflammatory state during pregnancy (Table 1). Although evidence of inflammatory changes in fetuses with maternal CA and exposed to opioids *in utero* is limited, findings of heightened

maternal inflammation during pregnancy and heightened offspring inflammation postnatally may imply heightened offspring inflammation during gestation as well. If fetal offspring exhibit increased inflammation during gestation, neurodevelopment *in utero* may be altered, including increased oligodendrocyte proliferation and differentiation (Filipovic and Zecevic, 2008). Given the strong evidence for effects of heightened inflammation during pregnancy on offspring neurodevelopment and risk for psychiatric disorders, future research investigating the cumulative and interactive effects of maternal CA and opioid use on maternal inflammation during pregnancy represents a priority for future research.

3.4 - Oxidative stress

Given that oxidative stress and pro-inflammatory processes are highly interconnected mechanisms of pathology, it is not surprising that alterations in mitochondrial biology and increased oxidative stress markers have been observed in individuals exposed to CA and prenatal opioid exposure. Oxidative stress is a term used to describe a pronounced imbalance in oxidation-reduction homeostasis beyond normal redox signaling (Sies et al., 2017). Under typical physiological conditions, the mitochondrion produces a small amount of reactive oxygen species (ROS) which are counteracted by enzymatic defense mechanisms. Mitochondrial damage (e.g. fragmentation) may lead to an imbalance in ROS production and antioxidant defense capacity resulting in further cell-damaging oxidative stress (Gyllenhammer et al., 2020; Hoffmann and Spengler, 2018). The deleterious consequences of oxidative stress may include an increase in nucleic acid mutations, amino acid and protein damage, endoplasmic reticulum (ER) stress, and cell death, as well as numerous pathologies including cardiovascular disease, cancer, neurodegenerative disease, inflammatory disease, and viral infections (Sies et al., 2017). Individuals using opioids demonstrate evidence of oxidative stress (Awadalla and Salah-Eldin, 2016; Fan et al., 2015; Faria et al., 2016; Zhuo et al., 2012), likely resulting from mitochondrial impairment (Cunha-Oliveira et al., 2008; Faria et al., 2016; Mohamed et al., 2015; Zhuo et al., 2012).

Preclinical studies show a direct link between prenatal opioid exposure and elevated markers of oxidative stress in offspring brains throughout development (Aboulhoda and Hassan, 2018; Guzmán et al., 2006; Hung et al., 2013). Similarly, animals administered opioids were found to have increased ROS production, oxidative stress, and mitochondrial activity and density (Cunha-Oliveira et al., 2008; Faria et al., 2016; Mehdizadeh et al., 2017; Mohamed et al., 2015; Zhuo et al., 2012). Human adults AFAB with direct CA exposure showed increased ROS production, oxidative stress, and mitochondrial activity and density outside of the perinatal period (Boeck et al., 2016).

Similar signs of elevated oxidative stress were observed in association with preconception CA. The amount of mitochondrial DNA (mtDNA) in peripheral blood mononuclear cells or buccal cells is an indicator of the quality or health of mitochondria when considered with mitochondrial functional capacity, which has implications for oxidative stress (Picard et al., 2018). In two studies, mtDNA levels were observed to be increased in association with CA (Cai et al., 2015; Tyrka et al., 2016), but in one of these the alterations in mtDNA were contingent on presence of depressive state (Cai et al., 2015), and another study did not replicate these findings (Cai et al., 2020). During pregnancy, maternal stress and particularly

lifetime stress have been associated with reduced placental mtDNA content (Brunst et al., 2017) and differential expression of protein-coding mitochondrial genes in the placenta, which, in turn was associated with a more difficult infant temperament (Lambertini et al., 2015). However, the amount of mtDNA alone may not be a good indicator of mitochondrial quality or health without any information on mitochondrial functional capacity (Picard et al., 2018). Boeck et al. (2016) investigated mitochondrial functioning in individuals AFAB exposed to CA and observed a dose-response association with higher ROS production, higher oxidative stress and increased mitochondrial activity. Extending on this work, the same group demonstrated increased mitochondrial activity and density in individuals with CA shortly after parturition compared to controls (Gumpp et al., 2020), an association that was also observed 3 months postpartum, however, only in participants with high concentrations of cortisol (Boeck et al., 2018).

Animal studies demonstrate multiple deleterious brain outcomes associated with prenatal exposure to stress-induced oxidative stress, including cognitive impairment, dopamine D1 receptor dysfunction (D1DR), dysregulated N-Methyl-D-aspartate (NMDA) synaptic currents, neural apoptosis (especially in the hippocampus), and impaired long-term potentiation in CA1 (Cao et al., 2014; Giussani et al., 2012; Lu et al., 2013; Wang et al., 2014). Maternal oxidative stress appears to be indirectly associated with fetal oxidative stress through the reduction of placental perfusion and intrauterine increases in glucocorticoids and cytokines (Rakers et al., 2017). The increases in fetal oxidative stress, inflammation, and HPA axis activity all appear to contribute to the increased chances of offspring neurodevelopmental impairments associated with maternal oxidative stress and mitochondrial dysfunction (Buss, 2021; Graham et al., 2019, 2018). In addition to indirect effects of maternal mitochondrial biology on the developing fetus via alterations in placental function, stress hormone concentration, or pro-inflammatory processes, maternal mitochondria are physically passed from the oocyte to the zygote and thus directly influence offspring mitochondrial biology which may confer long-term effects on health and disease risk (Gyllenhammer et al., 2020). Thus elevations in oxidative stress during pregnancy, to which both maternal CA and opioid use may contribute, represent an important potential pathway for influencing offspring neurodevelopment.

3.5 – Epigenetics

Prenatal opioid exposure and maternal CA both have potential to influence offspring brain development via epigenetic mechanisms. Epigenetics refers to environmentally-induced alterations to gene expression through modification of DNA methylation and histone tails, chromatin structure, non-coding RNAs (i.e. microRNAs), and transposable elements (Jirtle and Skinner, 2007; Murrell et al., 2005; Slotkin and Martienssen, 2007, 2007; Wolffe and Matzke, 1999). While parental epigenetic marks are almost fully erased after fertilization (Seisenberger et al., 2012), some gene loci survive this methylation reprogramming, introducing the possibility of intergenerational and transgenerational transmission of epigenetic changes (Anway et al., 2005; Branco et al., 2016; Lane et al., 2003; Morgan et al., 1999; Radford, 2018; Rakyan et al., 2003; Sanchez-Delgado et al., 2016; Smallwood et al., 2011; Smith et al., 2012). As previously discussed, gestation represents a critical window in which developmental trajectories are more susceptible to changes in response to

environmental conditions through multiple pathways, including via epigenetic modifications (Jirtle and Skinner, 2007). Here we first discuss direct epigenetic alterations, which may occur in the fetus during gestation in response to biological cues from an MPF environment affected by opioid exposure and maternal CA. Second, we examine potential intergenerational inheritance of parental epigenetic changes in response to opioid use and preconception CA that may survive zygotic reprogramming and persist in offspring.

In utero opioid exposure is associated with offspring epigenetic modifications with implications for withdrawal symptoms shortly after birth (NOWS) and long-term development. Maternal exposure to morphine during the preconception, prenatal, and lactation periods in rats was associated with reduced hippocampal synaptic plasticity in rat offspring, with potential implications for offspring learning and memory abilities (Sarkaki et al., 2008). In humans, opioid-exposed neonates have increased methylation of adenosine triphosphate (ATP)-binding cassette sub-family B member 1 (ABCB1), cytochrome P450 family 2 subfamily D member 6 (CYP2D6), and the MOR gene in comparison to opioid-naïve neonates (McLaughlin et al., 2017). These genes are important for basic cellular and neurological function, in addition to the metabolism of opioids and other substances (Gaedigk, 2013; Hodges et al., 2011; Valentino and Volkow, 2018). Additionally, hypermethylation patterns on the MOR gene of human neonates exposed to opioids in utero were associated with greater severity of withdrawal symptoms at birth (Wachman et al., 2014), but this finding has not been consistently replicated (McLaughlin et al., 2017). These alterations reported in offspring with prenatal opioid exposure do demonstrate epigenetic modifications in this population. However, the exact mechanisms of these epigenetic alterations remain unclear. Additionally, the role of co-occurring prenatal environmental influences continues to complicate clinical studies in this area and will need to be considered in future research.

Telomeres, DNA-protein complexes which prevent chromosomal damage and maintain genomic stability (Blackburn, 2005), are of particular interest for tracking intrauterine epigenetic alterations and intergenerational epigenetic inheritance associated with offspring neurodevelopment. Telomere length is epigenetically altered throughout the lifespan in response to aging and environmental exposures, and shorter telomere length is associated with multiple psychiatric disorders (Lindqvist et al., 2015) and other health conditions (Zhu et al., 2011). The epigenetic regulation of fetal telomere length appears to begin *in utero* with input from stress-sensitive oxidative, immune, endocrine, and metabolic pathways in the MPF environment, and shorter telomere length at birth appears to increase risk for long-term adverse outcomes (Entringer et al., 2018). These findings suggest that both prenatal opioid exposure and maternal CA have potential to alter offspring telomere length *in utero* through several of the MPF pathways previously discussed.

Furthermore, epigenetically-altered parental telomere length appears to program zygote telomere length through both parental germ lines, suggesting that preconception parental exposures may alter offspring telomere length through intergenerational epigenetic inheritance (Bauch et al., 2019; Delgado et al., 2019; Factor-Litvak et al., 2016; Olsson et al., 2011). CA in particular is associated with shorter telomere length throughout the lifespan (Blaze et al., 2015; Kiecolt-Glaser et al., 2011; Li et al., 2017; Ridout et al., 2018), which

may be transmitted to offspring. Telomere length was shorter in 4-, 12-, and 18-month-old infants with maternal CA history, which corresponded to offspring externalizing problems at 18 months while controlling for prenatal stress and maternal depression (Esteves et al., 2019). While prenatal opioid exposure has not been examined in association with offspring telomere length, one study found that heroin use was associated with shorter telomere length in adults while controlling for psychiatric and physical comorbidities, stressful event exposures, age, sex, and smoking (Yang et al., 2013). This suggests a potential pathway through which preconception opioid use may alter offspring telomere length through intergenerational epigenetic inheritance. Future research will be needed to directly examine this pathway.

The majority of other studies investigating epigenetic inheritance through the germ line have been conducted in paternal germ cells. Limited evidence suggests that both adulthood opioid use and CA history may epigenetically alter human sperm (Chorbov et al., 2011; Roberts et al., 2018), but it is unclear if these alterations would survive post-fertilization methylation reprogramming. Additionally, several studies demonstrate that stress and fear may initiate epigenetic alterations to paternal germ cells in mice that are associated with offspring behavioral and physiological alterations (Dias and Ressler, 2014; Gapp et al., 2020; Rodgers et al., 2015, 2013).

While there is increasing evidence for true epigenetic inheritance via the paternal germ line, to our knowledge there is no study to date directly showing the inheritance of epigenetic marks of parental opioid use or CA via the maternal germ line. However, some animal studies provide indirect evidence for an epigenetic contribution to intergenerational effects of preconception maternal opioid use, demonstrating that offspring epigenetic alterations were associated with preconception maternal opioid administration (Byrnes et al., 2013; Vassoler et al., 2016). Several studies also indirectly suggest that epigenetic sequelae associated with maternal CA may be transmitted through oocyte alterations. Female rats that underwent chronic unpredictable stress in adulthood showed an increase in corticotropin releasing factor type 1 (CRF1) mRNA in the frontal cortex as well as in mature oocytes. The effects on brain CRF1 expression persisted into the next generation and were associated with behavioral abnormalities (Zaidan et al., 2013).

There does appear to be evidence that both prenatal opioid exposure and maternal (and paternal) CA have implications for epigenetic modifications in offspring, but much of the current evidence is not able to identify mechanisms for these modifications. Additionally, many of these findings lack a direct connection between epigenetic pathways of prenatal opioid exposure and maternal CA. However, epigenetic pathways of intergenerational CA appear to interact with other mechanisms influenced by prenatal opioid exposure, such as differential HPA axis functioning (Yehuda et al., 2014; Zaidan et al., 2013), highlighting the likelihood of complex, interactive effects with potential to influence offspring neurodevelopment. See Table 1 for a summary of findings related to pathways reviewed in Section 3.

4 – Conclusions and future directions

A substantial amount of research has been dedicated to understanding how *in utero* opioid exposure influences neurodevelopment due to the considerable increase in opioid use in recent decades. Our interpretation of the literature to date indicates some subtle alterations evident soon after birth following *in utero* opioid exposure, which may confer vulnerability to mood and anxiety disorders, differential social and reward processing, and learning and memory impairments. However, there is limited evidence that these behavioral and cognitive differences persist into early childhood or adulthood given that many of the findings appear to be completely or partially mitigated by differences in the postnatal environment (Ahmadalipour et al., 2015; Hartman and Belsky, 2018; O'Donnell and Meaney, 2016; Salzwedel et al., 2020). These findings are of particular interest because recent research in the area of prenatal programming has suggested that prenatal adversity does not program neurodevelopmental disorders, rather exposure to poorer circumstances during gestation may alter susceptibility to the influences of the postnatal environment, for better or for worse (Hartman and Belsky, 2018; O'Donnell and Meaney, 2016). However, much of our understanding comes from well-controlled animal research, which has acknowledged limitations, including cross-species differences in drug metabolism and gestational and neurodevelopmental timing, and challenges in approximating the multiple co-occurring risk factors typically accompanying opioid use during pregnancy in humans (Byrnes and Vassoler, 2018). Research in humans addressing effects of *in utero* opioid exposure on offspring neurodevelopment is limited due to small sample sizes and the challenges of addressing myriad commonly co-occurring pre- and postnatal factors with significant potential to influence offspring neurodevelopment.

Examining candidate mechanistic pathways by which opioids and commonly co-occurring factors may influence offspring brain development represents an important direction for future research in this area. We highlight maternal CA history as a common yet understudied potential influence on offspring neurodevelopment in the context of maternal opioid use during pregnancy. Our review identifies multiple overlapping mechanistic pathways for the influence of maternal opioid use during pregnancy and maternal CA history on offspring neurodevelopment. These are aspects of MPF biology with evidence supporting sensitivity to both exogenous opioids and maternal CA history, and potential for programming fetal neurodevelopmental processes. The identified mechanistic pathways include the endogenous opioid system, the HPA axis, the immune system, epigenetics, and oxidative stress (Table 1). We also note that this review is not exhaustive, and other shared candidate mechanisms for effects of prenatal opioid exposure and maternal CA history likely include metabolic and other endocrine pathways (Buss et al., 2017; de Vries et al., 2020). The existence of these overlapping mechanistic pathways has important implications for research and policy.

From a research perspective, the existence of multiple shared mechanistic pathways for effects of *in utero* opioid exposure and maternal CA history on neurodevelopment suggests strong potential for cumulative and interactive influences, which call into question the utility and meaning of research focusing exclusively on effects of opioid exposure on neurodevelopment. A more fruitful approach will likely involve assessment of opioid use during pregnancy along with maternal CA history and other historical, environmental, and

demographic factors. Examination of these factors in relation to candidate shared biological mechanisms for effects on fetal neurodevelopment represents an important first step in this research. Such work will require interdisciplinary expertise to facilitate assessment of maternal substance use, CA history, psychological and physical health, environment and demographics, as well as MPF biology. A second critical step in this work will involve use of neuroimaging tools that can be used to assess brain structure and function shortly after birth, including structural and functional MRI and electroencephalograms, to minimize confounding effects of postnatal environmental influences on neurodevelopment. More generally, but also particularly important in the case of multivariate analyses and neuroimaging research, larger sample sizes (up to several thousands (Marek et al., 2020)) will be needed to identify reproducible findings. Furthermore, longitudinal studies assessing a wide variety of potentially co-occurring factors during pregnancy that follow offspring into childhood and beyond will be important for better understanding the roles of co-occurring risk factors and predictive pathways to offspring outcomes in childhood.

Implications for future directions include the need to facilitate and support collaborative science, changing public policy for individuals using opioids, and improving the treatment and prevention of opioid use in this population. Bringing together the necessary resources and expertise to recruit large samples of high risk, frequently stigmatized populations, while thoroughly assessing complex environmental, psychological, and biological systems, will be important next steps in this area of research. The heterogeneity of factors affecting offspring brain development in pregnant individuals using opioids, and the difficulty of disentangling these factors with the current scientific literature, also has implications for public policy, treatment, and prevention in this population. Many individuals are still penalized for using opioids during pregnancy and face high levels of societal pressure and stigma (Krans and Patrick, 2016; Patrick et al., 2017). Further, we acknowledge that the opioid crisis has received increasing resources, attention, public sympathy, and decriminalization as its demographics have shifted toward a primarily white population (Cicero et al., 2014; Hansen et al., 2020; Santoro and Santoro, 2018). White individuals are overrepresented in the opioid-misusing population because of two primary factors: 1) opioid prescriptions have fueled increasing nationwide opioid use in recent decades (Volkow and Blanco, 2021), and 2) Black, Indigenous, and People of Color (BIPOC) are less likely to receive opioid prescriptions due to racial biases among prescribers and reduced access to healthcare (Hansen et al., 2020; Om, 2018; Santoro and Santoro, 2018). The opioid epidemic is a great public health concern deserving of the resources it has been given; however, we acknowledge that many other current and historical public health crises primarily affecting marginalized populations in the United States have not received appropriate public and legislative support (Hardeman et al., 2018; Montoya-Barthelemy et al., 2020; Tester, 2017).

This review highlights findings indicating that individuals with CA experience long-lasting consequences that not only increase their risk of opioid abuse (Austin and Shanahan, 2018; Derefinko et al., 2019; Merrick et al., 2020; Savulich et al., 2017), but have implications for offspring neurodevelopment, and potential to exacerbate effects of *in utero* opioid exposure through shared mechanistic pathways. Further research in this area has potential to inform policy focused on ameliorating the negative sequelae of the opioid epidemic for the next generation by elucidating the biological programming potential of factors

co-occurring with opioid use, which may frequently be conceptualized as less relevant for offspring neurodevelopment. Such work has important implications for determining the extent to which resources will be devoted to making evidence-based trauma treatment readily available and increasing accessibility of trauma-informed treatment for pregnant individuals using opioids (SAMHSA, 2016). Long-term goals include increasing community resources and access to appropriate care, while supporting the most vulnerable members of our population.

Acknowledgements

Support for this work was provided by NIDA T32-DA007262 (Allen), R00 MH111805 (Graham), R34DA050291 (Graham & Fair), R34DA050291-S2 (Graham & Fair), P50 DA048756 (Mackiewicz Seghete), UG3 OD023349 (O'Connor, Buss, Miller, Simhan & Wadhwa), R01MH105538 (Wadhwa, Buss & Fair)

References

- Aagaard K, Bach CC, Henriksen TB, Larsen RT, Matthiesen NB, 2018. Head circumference at birth and childhood developmental disorders in a nationwide cohort in Denmark. Paediatr. Perinat. Epidemiol 32, 458–466. doi:10.1111/ppe.12479 [PubMed: 29882976]
- Aaltonen R, Heikkinen T, Hakala K, Laine K, Alanen A, 2005. Transfer of Proinflammatory Cytokines Across Term Placenta. Obstet. Gynecol 106, 802–807. doi:10.1097/01.AOG.0000178750.84837.ed [PubMed: 16199639]
- Abdel-Latif ME, Oei J, Craig F, Lui K, 2013. Profile of infants born to drug-using mothers A state-wide audit. J. Paediatr. Child Health 49, E80–E86. doi:10.1111/j.1440-1754.2012.02471.x [PubMed: 22530812]
- Aboulhoda BE, Hassan SS, 2018. Effect of prenatal tramadol on postnatal cerebellar development: Role of oxidative stress. J. Chem. Neuroanat 94, 102–118. doi:10.1016/j.jchemneu.2018.10.002 [PubMed: 30342117]
- Ahmadalipour A, Sadeghzadeh J, Vafaei AA, Bandegi AR, Mohammadkhani R, Rashidy-Pour A, 2015. Effects of environmental enrichment on behavioral deficits and alterations in hippocampal BDNF induced by prenatal exposure to morphine in juvenile rats. Neuroscience 305, 372–383. doi:10.1016/j.neuroscience.2015.08.015 [PubMed: 26272536]
- Anway MD, Cupp AS, Uzumcu M, Skinner MK, 2005. Epigenetic transgenerational actions of endocrine disruptors and male fertility. Science 308, 1466–1469. doi:10.1126/science.1108190 [PubMed: 15933200]
- Apanasewicz-Grzegorczyk A, Groth D, Scheffler C, Hermanussen M, Piosek M, Wychowaniec P, Babiszewska M, Barbarska O, Ziomkiewicz A, 2020. Traumatized women's infants are bigger than children of mothers without traumas. Anthropol. Anz 77. doi: 10.1127/anthranz/2020/1285
- Appleton AA, Kiley K, Holdsworth EA, Schell LM, 2019. Social Support During Pregnancy Modifies the Association Between Maternal Adverse Childhood Experiences and Infant Birth Size. Matern. Child Health J 23, 408–415. doi:10.1007/s10995-018-02706-z [PubMed: 30627949]
- Areda T, Kõks S, Philips M-A, Vasar E, Karis A, Asser T, 2005. Alterations in opioid system of the rat brain after cat odor exposure. Neurosci. Lett 377, 136–139. doi:10.1016/j.neulet.2004.11.083 [PubMed: 15740852]
- Arezoomandan R, Khodagholi F, Haghsparast A, 2016. Administration of the glial condition medium in the nucleus accumbens prolong maintenance and intensify reinstatement of morphine-seeking behavior. Neurochem Res 41, 855–68. [PubMed: 26547198]
- Ashdown H, Dumont Y, Ng M, Poole S, Boksa P, Luheshi GN, 2006. The role of cytokines in mediating effects of prenatal infection on the fetus: implications for schizophrenia. Mol. Psychiatry 11, 47–55. doi:10.1038/sj.mp.4001748 [PubMed: 16189509]
- Austin AE, Shanahan ME, 2018. Association of childhood abuse and neglect with prescription opioid misuse: Examination of mediation by adolescent depressive symptoms and pain. Child. Youth Serv. Rev 86, 84–93. doi:10.1016/j.childyouth.2018.01.023

- Awadalla EA, Salah-Eldin A-E, 2016. Molecular and histological changes in cerebral cortex and lung tissues under the effect of tramadol treatment. Biomed. Pharmacother 82, 269–280. doi:10.1016/ j.biopha.2016.04.024 [PubMed: 27470363]
- Bachtell R, Hutchinson MR, Wang X, Rice KC, Maier SF, Watkins LR, 2015. Targeting the toll of drug abuse: the translational potential for toll-like receptor 4. CNS Neurol Diord Drug Targets 14, 692–9.
- Bakhireva LN, Holbrook BD, Shrestha S, Leyva Y, Ashley M, Cano S, Lowe J, Stephen JM, Leeman L, 2019. Association between prenatal opioid exposure, neonatal opioid withdrawal syndrome, and neurodevelopmental and behavioral outcomes at 5–8 months of age. Early Hum. Dev 128, 69–76. doi:10.1016/j.earlhumdev.2018.10.010 [PubMed: 30554024]
- Baldacchino A, Arbuckle K, Petrie DJ, McCowan C, 2014. Neurobehavioral consequences of chronic intrauterine opioid exposure in infants and preschool children: a systematic review and metaanalysis. BMC Psychiatry 14, 104. doi:10.1186/1471-244X-14-104 [PubMed: 24708875]
- Bauch C, Boonekamp JJ, Korsten P, Mulder E, Verhulst S, 2019. Epigenetic inheritance of telomere length in wild birds. PLOS Genet. 15, e1007827. doi:10.1371/journal.pgen.1007827 [PubMed: 30763308]
- Baumeister D, Akhtar R, Ciufolini S, Pariante CM, Mondelli V, 2016. Childhood trauma and adulthood inflammation: a meta-analysis of peripheral C-reactive protein, interleukin-6 and tumour necrosis factor-a. Mol. Psychiatry 21, 642–649. doi:10.1038/mp.2015.67 [PubMed: 26033244]
- Beckwith AM, Burke SA, 2015. Identification of early developmental deficits in infants with prenatal heroin, methadone, and other opioid exposure. Clin. Pediatr. (Phila.) 54, 328–335. doi:10.1177/0009922814549545 [PubMed: 25189695]
- Benediktsson R, Calder AA, Edwards CR, Seckl JR, 1997. Placental 11 beta-hydroxysteroid dehydrogenase: a key regulator of fetal glucocorticoid exposure. Clin. Endocrinol. (Oxf.) 46, 161–166. doi:10.1046/j.1365-2265.1997.1230939.x [PubMed: 9135697]
- Bhat R, Chari G, Rao R, 2006. Effects of prenatal cocaine, morphine, or both on postnatal opioid (μ) receptor development. Life Sci. 78, 1478–1482. doi:10.1016/j.lfs.2005.07.023 [PubMed: 16242731]
- Bilder DA, Pinborough-Zimmerman J, Bakian AV, Miller JS, Dorius JT, Nangle B, McMahon WM, 2013. Prenatal and Perinatal Factors Associated with Intellectual Disability. Am. J. Intellect. Dev. Disabil 118, 156–176. doi:10.1352/1944-7558-118.2.156 [PubMed: 23464612]
- Bilkei-Gorzo A, Racz I, Michel K, Mauer D, Zimmer Anne, Klingmüller D, Zimmer Andreas, 2008. Control of hormonal stress reactivity by the endogenous opioid system. Psychoneuroendocrinology 33, 425–436. doi:10.1016/j.psyneuen.2007.12.010 [PubMed: 18280051]
- Blackburn EH, 2005. Telomeres and telomerase: their mechanisms of action and the effects of altering their functions. FEBS Lett. 579, 859–862. doi:10.1016/j.febslet.2004.11.036 [PubMed: 15680963]
- Blaze J, Asok A, Roth TL, 2015. The long-term impact of adverse caregiving environments on epigenetic modifications and telomeres. Front. Behav. Neurosci 9. doi:10.3389/fnbeh.2015.00079
- Boeck C, Gumpp AM, Calzia E, Radermacher P, Waller C, Karabatsiakis A, Kolassa I-T, 2018. The association between cortisol, oxytocin, and immune cell mitochondrial oxygen consumption in postpartum women with childhood maltreatment. Psychoneuroendocrinology 96, 69–77. doi:10.1016/j.psyneuen.2018.05.040 [PubMed: 29908404]
- Boeck C, Koenig AM, Schury K, Geiger ML, Karabatsiakis A, Wilker S, Waller C, Gündel H, Fegert JM, Calzia E, Kolassa I-T, 2016. Inflammation in adult women with a history of child maltreatment: The involvement of mitochondrial alterations and oxidative stress. Mitochondrion 30, 197–207. doi:10.1016/j.mito.2016.08.006 [PubMed: 27530300]
- Boekel W, Wagenmakers E-J, Belay L, Verhagen J, Brown S, Forstmann BU, 2015. A purely confirmatory replication study of structural brain-behavior correlations. Cortex J. Devoted Study Nerv. Syst. Behav 66, 115–133. doi:10.1016/j.cortex.2014.11.019
- Boggess T, Risher WC, 2020. Clinical and basic research investigations into the long-term effects of prenatal opioid exposure on brain development. J. Neurosci. Res doi:10.1002/jnr.24642

- Bonhomme J, Shim RS, Gooden R, Tyus D, Rust G, 2012. Opioid Addiction and Abuse in Primary Care Practice: A Comparison of Methadone and Buprenorphine as Treatment Options. J. Natl. Med. Assoc 104, 342–350. [PubMed: 23092049]
- Bouvette-Turcot A-A, Fleming AS, Wazana A, Sokolowski MB, Gaudreau H, Gonzalez A, Deslauriers J, Kennedy JL, Steiner M, Meaney MJ, MAVAN Research Team, 2015. Maternal childhood adversity and child temperament: an association moderated by child 5-HTTLPR genotype. Genes Brain Behav. 14, 229–237. doi:10.1111/gbb.12205 [PubMed: 25688466]
- Branco MR, King M, Perez-Garcia V, Bogutz AB, Caley M, Fineberg E, Lefebvre L, Cook SJ, Dean W, Hemberger M, Reik W, 2016. Maternal DNA Methylation Regulates Early Trophoblast Development. Dev. Cell 36, 152–163. doi:10.1016/j.devcel.2015.12.027 [PubMed: 26812015]
- Brown SM, Shillington AM, 2017. Childhood adversity and the risk of substance use and delinquency: The role of protective adult relationships. Child Abuse Negl. 63, 211–221. doi:10.1016/ j.chiabu.2016.11.006 [PubMed: 27884507]
- Brunst KJ, Sanchez Guerra M, Gennings C, Hacker M, Jara C, Bosquet Enlow M, Wright RO, Baccarelli A, Wright RJ, 2017. Maternal Lifetime Stress and Prenatal Psychological Functioning and Decreased Placental Mitochondrial DNA Copy Number in the PRISM Study. Am. J. Epidemiol 186, 1227–1236. doi:10.1093/aje/kwx183 [PubMed: 28595325]
- Brunton PJ, 2019. Endogenous opioid signalling in the brain during pregnancy and lactation. Cell Tissue Res. 375, 69–83. doi:10.1007/s00441-018-2948-1 [PubMed: 30415283]
- Brunton PJ, McKay AJ, Och dalski T, Piastowska A, R bas E, Lachowicz A, Russell JA, 2009. Central Opioid Inhibition of Neuroendocrine Stress Responses in Pregnancy in the Rat Is Induced by the Neurosteroid Allopregnanolone. J. Neurosci 29, 6449–6460 doi:10.1523/ JNEUROSCI.0708-09.2009 [PubMed: 19458216]
- Bublitz M, De La Monte S, Martin S, Larson L, Bourjeily G, 2017. Childhood maltreatment and inflammation among pregnant women with gestational diabetes mellitus: A pilot study. Obstet. Med 10, 120–124. doi:10.1177/1753495X17701320 [PubMed: 29051779]
- Bublitz MH, Parade S, Stroud LR, 2014. The effects of childhood sexual abuse on cortisol trajectories in pregnancy are moderated by current family functioning. Biol. Psychol 103, 152– 157. doi:10.1016/j.biopsycho.2014.08.014 [PubMed: 25220484]
- Bublitz MH, Stroud LR, 2012a. Childhood sexual abuse is associated with cortisol awakening response over pregnancy: preliminary findings. Psychoneuroendocrinology 37, 1425–1430. doi:10.1016/ j.psyneuen.2012.01.009 [PubMed: 22341730]
- Bublitz MH, Stroud LR, 2012b. Childhood sexual abuse is associated with cortisol awakening response over pregnancy: Preliminary findings. Psychoneuroendocrinology 37, 1425–1430. doi:10.1016/ j.psyneuen.2012.01.009 [PubMed: 22341730]
- Buchanan MM, Hutchinson M, Watkin LR, Yin H, 2010. Toll-like receptor 4 in CNS pathologies. J Neurochem 114, 13–27. [PubMed: 20402965]
- Buisman-Pijlman FTA, Gerrits M. a. F.M., Van Ree JM, 2009a. Increased opioid release in specific brain areas in animals exposed to prenatal morphine and emotional stress later in life. Neuroscience 159, 405–413. doi:10.1016/j.neuroscience.2008.11.010 [PubMed: 19138727]
- Buisman-Pijlman FTA, Gerrits MAFM, Van Ree JM, 2009b. Increased opioid release in specific brain areas in animals exposed to prenatal morphine and emotional stress later in life. Neuroscience 159, 405–413. doi:10.1016/j.neuroscience.2008.11.010 [PubMed: 19138727]
- Buss C, 2021. Maternal oxidative stress during pregnancy and offspring neurodevelopment. Brain. Behav. Immun 93, 6–7. doi:10.1016/j.bbi.2021.01.007 [PubMed: 33454300]
- Buss C, Entringer S, Moog NK, Toepfer P, Fair DA, Simhan HN, Heim CM, Wadhwa PD, 2017. Intergenerational Transmission of Maternal Childhood Maltreatment Exposure: Implications for Fetal Brain Development. J. Am. Acad. Child Adolesc. Psychiatry 56, 373–382. doi:10.1016/ j.jaac.2017.03.001 [PubMed: 28433086]
- Byrnes EM, Vassoler FM, 2018. Modeling prenatal opioid exposure in animals: Current findings and future directions. Front. Neuroendocrinol 51, 1–13. doi:10.1016/j.yfrne.2017.09.001 [PubMed: 28965857]

- Byrnes JJ, Johnson NL, Carini LM, Byrnes EM, 2013. Multigenerational effects of adolescent morphine exposure on dopamine D2 receptor function. Psychopharmacology (Berl.) 227, 263– 272. doi:10.1007/s00213-012-2960-1 [PubMed: 23314440]
- Cai N, Chang S, Li Yihan, Li Q, Hu Jingchu, Liang J, Song L, Kretzschmar W, Gan X, Nicod J, Rivera M, Deng H, Du B, Li Keqing, Sang W, Gao J, Gao S, Ha B, Ho H-Y, Hu C, Hu Jian, Hu Z, Huang G, Jiang G, Jiang T, Jin W, Li G, Li Kan, Li Yi, Li Yingrui, Li Youhui, Lin Y-T, Liu L, Liu T, Liu Ying, Liu Yuan, Lu Y, Lv L, Meng H, Qian P, Sang H, Shen J, Shi J, Sun J, Tao M, Wang Gang, Wang Guangbiao, Wang Jian, Wang L, Wang Xueyi, Wang Xumei, Yang H, Yang L, Yin Y, Zhang J, Zhang K, Sun N, Zhang W, Zhang X, Zhang Z, Zhong H, Breen G, Wang Jun, Marchini J, Chen Y, Xu Q, Xu X, Mott R, Huang G-J, Kendler K, Flint J, 2015. Molecular Signatures of Major Depression. Curr. Biol 25, 1146–1156. doi:10.1016/j.cub.2015.03.008 [PubMed: 25913401]
- Cai N, F ašková M, Kone ná K, Fojtová M, Fajkus J, Coomber E, Watt S, Soranzo N, Preiss M, Rektor I, 2020. No Evidence of Persistence or Inheritance of Mitochondrial DNA Copy Number in Holocaust Survivors and Their Descendants. Front. Genet 11, 87. doi:10.3389/fgene.2020.00087 [PubMed: 32211017]
- Camden A, Ray JG, To T, Gomes T, Bai L, Guttmann A, 2021. Identification of Prenatal Opioid Exposure Within Health Administrative Databases. Pediatrics 147. doi:10.1542/peds.2020-018507
- Cao K, Zheng A, Xu J, Li H, Liu Jing, Peng Y, Long J, Zou X, Li Y, Chen C, Liu Jiankang, Feng Z, 2014. AMPK activation prevents prenatal stress-induced cognitive impairment: Modulation of mitochondrial content and oxidative stress. Free Radic. Biol. Med 75, 156–166. doi:10.1016/j.freeradbiomed.2014.07.029 [PubMed: 25091899]
- Chang L, Kigar SL, Ho JH, Cuarenta A, Gunderson HC, Baldo BA, Bakshi VP, Auger AP, 2019. Early life stress alters opioid receptor mRNA levels within the nucleus accumbens in a sex-dependent manner. Brain Res. 1710, 102–108. doi:10.1016/j.brainres.2018.12.040 [PubMed: 30594547]
- Cheong JLY, Hunt RW, Anderson PJ, Howard K, Thompson DK, Wang HX, Bear MJ, Inder TE, Doyle LW, 2008. Head Growth in Preterm Infants: Correlation With Magnetic Resonance Imaging and Neurodevelopmental Outcome. Pediatrics 121, e1534–e1540. doi:10.1542/peds.2007-2671 [PubMed: 18519457]
- Chiou L-C, Yeh G-C, Fan S-H, How C-H, Chuang K-C, Tao P-L, 2003. Prenatal morphine exposure decreases analgesia but not K+ channel activation. Neuroreport 14, 239–242. doi:10.1097/00001756-200302100-00016 [PubMed: 12598737]
- Chorbov VM, Todorov AA, Lynskey MT, Cicero TJ, 2011. Elevated levels of DNA methylation at the OPRM1 promoter in blood and sperm from male opioid addicts. J. Opioid Manag 7, 258–264. [PubMed: 21957825]
- Cicero TJ, Ellis MS, Surratt HL, Kurtz SP, 2014. The Changing Face of Heroin Use in the United States: A Retrospective Analysis of the Past 50 Years. JAMA Psychiatry 71, 821–826. doi:10.1001/jamapsychiatry.2014.366 [PubMed: 24871348]
- Clancy B, Kersh B, Hyde J, Darlington RB, Anand KJS, Finlay BL, 2007. Web-based method for translating neurodevelopment from laboratory species to humans. Neuroinformatics 5, 79–94. doi:10.1385/NI:5:1:79 [PubMed: 17426354]
- Coelho R, Viola TW, Walss-Bass C, Brietzke E, Grassi-Oliveira R, 2014. Childhood maltreatment and inflammatory markers: a systematic review. Acta Psychiatr. Scand 129, 180–192. doi:10.1111/ acps.12217 [PubMed: 24205846]
- Colich NL, Platt JM, Keyes KM, Sumner JA, Allen NB, McLaughlin KA, 2020. Earlier age at menarche as a transdiagnostic mechanism linking childhood trauma with multiple forms of psychopathology in adolescent girls. Psychol. Med 50, 1090–1098. doi:10.1017/ S0033291719000953 [PubMed: 31020943]
- Collishaw S, Dunn J, O'Connor TG, Golding J, THE AVON LONGITUDINAL STUDY OF PARENTS AND CHILDREN STUDY TEAM, 2007. Maternal childhood abuse and offspring adjustment over time. Dev. Psychopathol 19. doi:10.1017/S0954579407070186
- Conradt E, Crowell SE, Lester BM, 2018. Early life stress and environmental influences on the neurodevelopment of children with prenatal opioid exposure. Neurobiol. Stress 9, 48–54. doi:10.1016/j.ynstr.2018.08.005 [PubMed: 30151420]
- Conradt E, Flannery T, Aschner JL, Annett RD, Croen LA, Duarte CS, Friedman AM, Guille C, Hedderson MM, Hofheimer JA, Jones MR, Ladd-Acosta C, McGrath M, Moreland A, Neiderhiser

JM, Nguyen RHN, Posner J, Ross JL, Savitz DA, Ondersma SJ, Lester BM, 2019. Prenatal Opioid Exposure: Neurodevelopmental Consequences and Future Research Priorities. Pediatrics 144. doi:10.1542/peds.2019-0128

- Cowan CSM, Callaghan BL, Kan JM, Richardson R, 2016. The lasting impact of early-life adversity on individuals and their descendants: potential mechanisms and hope for intervention. Genes Brain Behav. 15, 155–168. doi:10.1111/gbb.12263 [PubMed: 26482536]
- Craig A, Wolfe I, Bell R, Lucas F, Gabrielson S, Cox D, 2020. The effect of antenatal opioid exposure on orbitofrontal head circumference at birth, a retrospective cohort study. Costas T Lambrew Res. Retreat 2020.
- Cunha-Oliveira T, Rego AC, Oliveira CR, 2008. Cellular and molecular mechanisms involved in the neurotoxicity of opioid and psychostimulant drugs. Brain Res. Rev 58, 192–208. doi:10.1016/ j.brainresrev.2008.03.002 [PubMed: 18440072]
- Davis EP, Glynn LM, Dunkel Schetter C, Hobel C, Chicz-Demet A, Sandman CA, 2005. Corticotropin-releasing hormone during pregnancy is associated with infant temperament. Dev. Neurosci 27, 299–305. doi:10.1159/000086709 [PubMed: 16137987]
- Davis EP, Glynn LM, Schetter CD, Hobel C, Chicz-demet A, Sandman CA, 2007. Prenatal Exposure to Maternal Depression and Cortisol Influences Infant Temperament. J. Am. Acad. Child Adolesc. Psychiatry 46, 737–746. doi:10.1097/chi.0b013e318047b775 [PubMed: 17513986]
- Davis EP, Sandman CA, 2010. The Timing of Prenatal Exposure to Maternal Cortisol and Psychosocial Stress Is Associated With Human Infant Cognitive Development. Child Dev. 81, 131–148. doi:10.1111/j.1467-8624.2009.01385.x [PubMed: 20331658]
- Davis M, Pasternak G, 2005. Opioid receptors and opioid pharmacodynamics, in: Opioids in Cancer Pain 2. pp. 29–38. doi:10.1093/med/9780199236640.003.0002
- De Asis-Cruz J, Krishnamurthy D, Zhao L, Kapse K, Vezina G, Andescavage N, Quistorff J, Lopez C, Limperopoulos C, 2020. Association of Prenatal Maternal Anxiety With Fetal Regional Brain Connectivity. JAMA Netw. Open 3, e2022349–e2022349. doi:10.1001/jamanetworkopen.2020.22349 [PubMed: 33284334]
- de Cubas MM, Field T, 1993. Children of methadone-dependent women: developmental outcomes. Am. J. Orthopsychiatry 63, 266–276. doi:10.1037/h0079429 [PubMed: 7683453]
- de Vries F, Bruin M, Lobatto DJ, Dekkers OM, Schoones JW, van Furth WR, Pereira AM, Karavitaki N, Biermasz NR, Zamanipoor Najafabadi AH, 2020. Opioids and Their Endocrine Effects: A Systematic Review and Meta-analysis. J. Clin. Endocrinol. Metab 105, 1020–1029. doi:10.1210/clinem/dgz022
- de Vries GJ, Södersten P, 2009. Sex differences in the brain: The relation between structure and function. Horm. Behav., 50th Anniversary of the Publication of Phoenix, Goy, Gerall & Young 1959: Organizational Effects of Hormones 55, 589–596. doi:10.1016/j.yhbeh.2009.03.012
- de Weerth C, Hees Y, Buitelaar JK, 2003. Prenatal maternal cortisol levels and infant behavior during the first 5 months. Early Hum. Dev 74, 139–151. doi:10.1016/s0378-3782(03)00088-4 [PubMed: 14580753]
- Delgado DA, Zhang C, Gleason K, Demanelis K, Chen LS, Gao J, Roy S, Shinkle J, Sabarinathan M, Argos M, Tong L, Ahmed A, Islam T, Rakibuz-Zaman M, Sarwar G, Shahriar H, Rahman M, Yunus M, Doherty JA, Jasmine F, Kibriya MG, Ahsan H, Pierce BL, 2019. The contribution of parent-to-offspring transmission of telomeres to the heritability of telomere length in humans. Hum. Genet 138, 49–60. doi:10.1007/s00439-018-1964-2 [PubMed: 30536049]
- Derefinko KJ, Salgado García FI, Talley KM, Bursac Z, Johnson KC, Murphy JG, McDevitt-Murphy ME, Andrasik F, Sumrok DD, 2019. Adverse childhood experiences predict opioid relapse during treatment among rural adults. Addict. Behav 96, 171–174. doi:10.1016/j.addbeh.2019.05.008 [PubMed: 31102882]
- Desai RJ, Hernandez-Diaz S, Bateman BT, Huybrechts KF, 2014. Increase in Prescription Opioid Use During Pregnancy Among Medicaid-Enrolled Women. Obstet. Gynecol 123, 997–1002. doi:10.1097/AOG.00000000000208 [PubMed: 24785852]
- Dhaliwal A, Gupta M, 2021. Physiology, Opioid Receptor, in: StatPearls. StatPearls Publishing, Treasure Island (FL).

- Dias BG, Ressler KJ, 2014. Parental olfactory experience influences behavior and neural structure in subsequent generations. Nat. Neurosci 17, 89–96. doi:10.1038/nn.3594 [PubMed: 24292232]
- Entringer S, Buss C, Wadhwa PD, 2015. Prenatal stress, development, health and disease risk: A psychobiological perspective—2015 Curt Richter Award Paper. Psychoneuroendocrinology 62, 366–375. doi:10.1016/j.psyneuen.2015.08.019 [PubMed: 26372770]
- Entringer S, de Punder K, Buss C, Wadhwa PD, 2018. The fetal programming of telomere biology hypothesis: an update. Philos. Trans. R. Soc. B Biol. Sci 373, 20170151. doi:10.1098/rstb.2017.0151
- Esteves KC, Jones CW, Wade M, Callerame K, Smith AK, Theall KP, Drury SS, 2019. Adverse Childhood Experiences: Implications for Offspring Telomere Length and Psychopathology. Am. J. Psychiatry 177, 47–57. doi:10.1176/appi.ajp.2019.18030335 [PubMed: 31509004]
- Factor-Litvak P, Susser E, Kezios K, McKeague I, Kark JD, Hoffman M, Kimura M, Wapner R, Aviv A, 2016. Leukocyte Telomere Length in Newborns: Implications for the Role of Telomeres in Human Disease. Pediatrics 137. doi:10.1542/peds.2015-3927
- Fan R, Schrott LM, Snelling S, Ndi J, Arnold T, Korneeva NL, 2015. Chronic oxycodone induces integrated stress response in rat brain. BMC Neurosci. 16, 58. doi:10.1186/s12868-015-0197-8 [PubMed: 26377394]
- Faria J, Barbosa J, Queirós O, Moreira R, Carvalho F, Dinis-Oliveira RJ, 2016. Comparative study of the neurotoxicological effects of tramadol and tapentadol in SH-SY5Y cells. Toxicology 359–360, 1–10. doi:10.1016/j.tox.2016.06.010
- Farid WO, Dunlop SA, Tait RJ, Hulse GK, 2008. The Effects of Maternally Administered Methadone, Buprenorphine and Naltrexone on Offspring: Review of Human and Animal Data. Curr. Neuropharmacol 6, 125–150. doi:10.2174/157015908784533842 [PubMed: 19305793]
- Ferrer M, García-Esteban R, Iñiguez C, Costa O, Fernández-Somoano A, Rodríguez-Delhi C, Ibarluzea J, Lertxundi A, Tonne C, Sunyer J, Julvez J, 2019. Head circumference and child ADHD symptoms and cognitive functioning: results from a large population-based cohort study. Eur. Child Adolesc. Psychiatry 28, 377–388. doi:10.1007/s00787-018-1202-4 [PubMed: 30027417]
- Filipovic R, Zecevic N, 2008. The effect of CXCL1 on human fetal oligodendrocyte progenitor cells. Glia 56, 1–15. doi:10.1002/glia.20582 [PubMed: 17910053]
- Finy MS, Christian LM, 2018. Pathways linking childhood abuse history and current socioeconomic status to inflammation during pregnancy. Brain. Behav. Immun 74, 231–240. doi:10.1016/ j.bbi.2018.09.012 [PubMed: 30217532]
- Fricker LD, Margolis EB, Gomes I, Devi LA, 2020. Five Decades of Research on Opioid Peptides: Current Knowledge and Unanswered Questions. Mol. Pharmacol 98, 96–108. doi:10.1124/ mol.120.119388 [PubMed: 32487735]
- Gaedigk A, 2013. Complexities of *CYP2D6* gene analysis and interpretation. Int. Rev. Psychiatry 25, 534–553. doi:10.3109/09540261.2013.825581 [PubMed: 24151800]
- Gale CR, O'Callaghan FJ, Bredow M, Martyn CN, Team, the A.L.S. of P. and C.S., 2006. The Influence of Head Growth in Fetal Life, Infancy, and Childhood on Intelligence at the Ages of 4 and 8 Years. Pediatrics 118, 1486–1492. doi:10.1542/peds.2005-2629 [PubMed: 17015539]
- Gapp K, van Steenwyk G, Germain PL, Matsushima W, Rudolph KLM, Manuella F, Roszkowski M, Vernaz G, Ghosh T, Pelczar P, Mansuy IM, Miska EA, 2020. Alterations in sperm long RNA contribute to the epigenetic inheritance of the effects of postnatal trauma. Mol. Psychiatry 25, 2162–2174. doi:10.1038/s41380-018-0271-6 [PubMed: 30374190]
- Gerdin E, Lindberg RA, 1990. Transplacental transfer of morphine in man. J. Perinat. Med 18, 305–312. doi:10.1515/jpme.1990.18.4.305 [PubMed: 2262875]
- Giussani DA, Camm EJ, Niu Y, Richter HG, Blanco CE, Gottschalk R, Blake EZ, Horder KA, Thakor AS, Hansell JA, Kane AD, Wooding FBP, Cross CM, Herrera EA, 2012. Developmental Programming of Cardiovascular Dysfunction by Prenatal Hypoxia and Oxidative Stress. PLOS ONE 7, e31017. doi:10.1371/journal.pone.0031017 [PubMed: 22348036]
- Graham AM, Marr M, Buss C, Sullivan EL, Fair DA, 2021. Understanding Vulnerability and Adaptation in Early Brain Development using Network Neuroscience. Trends Neurosci. 44, 276– 288. doi:10.1016/j.tins.2021.01.008 [PubMed: 33663814]

- Graham AM, Rasmussen JM, Entringer S, Ben Ward E, Rudolph MD, Gilmore JH, Styner M, Wadhwa PD, Fair DA, Buss C, 2019. Maternal Cortisol Concentrations During Pregnancy and Sex-Specific Associations With Neonatal Amygdala Connectivity and Emerging Internalizing Behaviors. Biol. Psychiatry 85, 172–181. doi:10.1016/j.biopsych.2018.06.023 [PubMed: 30122286]
- Graham AM, Rasmussen JM, Rudolph MD, Heim CM, Gilmore JH, Styner M, Potkin SG, Entringer S, Wadhwa PD, Fair DA, Buss C, 2018. Maternal Systemic Interleukin-6 During Pregnancy Is Associated With Newborn Amygdala Phenotypes and Subsequent Behavior at 2 Years of Age. Biol. Psychiatry 83, 109–119. doi:10.1016/j.biopsych.2017.05.027 [PubMed: 28754515]
- Gray TR, Choo RE, Concheiro M, Williams E, Elko A, Jansson LM, Jones HE, Huestis MA, 2010. Prenatal methadone exposure, meconium biomarker concentrations and neonatal abstinence syndrome. Addiction 105, 2151–2159. doi:10.1111/j.1360-0443.2010.03097.x [PubMed: 20854338]
- Griffiths SK, Campbell JP, 2015. Placental structure, function and drug transfer. Contin. Educ. Anaesth. Crit. Care Pain 15, 84–89. doi:10.1093/bjaceaccp/mku013
- Gumpp AM, Boeck C, Behnke A, Bach AM, Ramo-Fernández L, Welz T, Gündel H, Kolassa I-T, Karabatsiakis A, 2020. Childhood maltreatment is associated with changes in mitochondrial bioenergetics in maternal, but not in neonatal immune cells. Proc. Natl. Acad. Sci 117, 24778– 24784. doi:10.1073/pnas.2005885117 [PubMed: 33004627]
- Gustafsson HC, Sullivan EL, Battison EAJ, Holton KF, Graham AM, Karalunas SL, Fair DA, Loftis JM, Nigg JT, 2020. Evaluation of maternal inflammation as a marker of future offspring ADHD symptoms: A prospective investigation. Brain. Behav. Immun 89, 350–356. doi:10.1016/ j.bbi.2020.07.019 [PubMed: 32707260]
- Gustafsson HC, Sullivan EL, Nousen EK, Sullivan CA, Huang E, Rincon M, Nigg JT, Loftis JM, 2018. Maternal prenatal depression predicts infant negative affect via maternal inflammatory cytokine levels. Brain. Behav. Immun 73, 470–481. doi:10.1016/j.bbi.2018.06.011 [PubMed: 29920327]
- Guzmán DC, Vázquez IE, Brizuela NO, Alvarez RG, Mejía GB, García EH, Santamaría D, de Apreza M. la R., Olguín HJ, 2006. Assessment of Oxidative Damage Induced by Acute Doses of Morphine Sulfate in Postnatal and Adult Rat Brain. Neurochem. Res 31, 549–554. doi:10.1007/ s11064-006-9053-7 [PubMed: 16758364]
- Gyllenhammer LE, Entringer S, Buss C, Wadhwa PD, 2020. Developmental programming of mitochondrial biology: a conceptual framework and review. Proc. Biol. Sci 287, 20192713. doi:10.1098/rspb.2019.2713 [PubMed: 32345161]
- Hahn JW, Jagwani S, Kim E, Rendell VR, He J, Ezerskiy LA, Wesselschmidt R, Coscia CJ, Belcheva MM, 2010. Mu and kappa opioids modulate mouse embryonic stem cell-derived neural progenitor differentiation via MAP kinases. J. Neurochem 112, 1431–1441. doi:10.1111/ j.1471-4159.2009.06479.x [PubMed: 19895666]
- Hale KD, Weigent DA, Gauthier DK, Hiramoto RN, Ghanta VK, 2003. Cytokine and hormone profiles in mice subjected to handling combined with rectal temperature measurement stress and handling only stress. Life Sci. 72, 1495–1508. doi:10.1016/s0024-3205(02)02415-3 [PubMed: 12535717]
- Hans SL, Jeremy RJ, 2001. Postneonatal mental and motor development of infants exposed in utero to opioid drugs. Infant Ment. Health J 22, 300–315. doi:10.1002/imhj.1003
- Hansen H, Parker C, Netherland J, 2020. Race as a Ghost Variable in (White) Opioid Research. Sci. Technol. Hum. Values 45, 848–876. doi:10.1177/0162243920912812
- Hantsoo L, Jašarevi E, Criniti S, McGeehan B, Tanes C, Sammel MD, Elovitz MA, Compher C, Wu G, Epperson CN, 2019a. Childhood adversity impact on gut microbiota and inflammatory response to stress during pregnancy. Brain. Behav. Immun 75, 240–250. doi:10.1016/j.bbi.2018.11.005 [PubMed: 30399404]
- Hantsoo L, Kornfield S, Anguera MC, Epperson CN, 2019b. Inflammation: A Proposed Intermediary Between Maternal Stress and Offspring Neuropsychiatric Risk. Biol. Psychiatry, Prenatal Programming of Neuropsychiatric Disorders Across the Lifespan 85, 97–106. doi:10.1016/ j.biopsych.2018.08.018
- Hardeman RR, Murphy KA, Karbeah J, Kozhimannil KB, 2018. Naming Institutionalized Racism in the Public Health Literature: A Systematic Literature Review. Public Health Rep. 133, 240–249. doi:10.1177/0033354918760574 [PubMed: 29614234]

- Hartman S, Belsky J, 2018. Prenatal programming of postnatal plasticity revisited—And extended. Dev. Psychopathol 30, 825–842. doi:10.1017/S0954579418000548 [PubMed: 30068425]
- Hauser KF, Knapp PE, 2018. Opiate Drugs with Abuse Liability Hijack the Endogenous Opioid System to Disrupt Neuronal and Glial Maturation in the Central Nervous System. Front. Pediatr 5. doi:10.3389/fped.2017.00294
- Heim CM, Entringer S, Buss C, 2019. Translating basic research knowledge on the biological embedding of early-life stress into novel approaches for the developmental programming of lifelong health. Psychoneuroendocrinology 105, 123–137. doi:10.1016/j.psyneuen.2018.12.011 [PubMed: 30578047]
- Heinonen K, Räikkönen K, Pesonen A-K, Kajantie E, Andersson S, Eriksson JG, Niemelä A, Vartia T, Peltola J, Lano A, 2008. Prenatal and Postnatal Growth and Cognitive Abilities at 56 Months of Age: A Longitudinal Study of Infants Born at Term. Pediatrics 121, e1325–e1333. doi:10.1542/ peds.2007-1172 [PubMed: 18450875]
- Hendrix CL, Dilks DD, McKenna BG, Dunlop AL, Corwin EJ, Brennan PA, 2020. Maternal childhood adversity associates with frontoamygdala connectivity in neonates. Biol. Psychiatry Cogn. Neurosci. Neuroimaging 0. doi:10.1016/j.bpsc.2020.11.003
- Hodges LM, Markova SM, Chinn LW, Gow JM, Kroetz DL, Klein TE, Altman RB, 2011. Very important pharmacogene summary: ABCB1 (MDR1, P-glycoprotein). Pharmacogenet. Genomics 21, 152–161. doi:10.1097/FPC.0b013e3283385a1c [PubMed: 20216335]
- Hoffmann A, Spengler D, 2018. The Mitochondrion as Potential Interface in Early-Life Stress Brain Programming. Front. Behav. Neurosci 12. doi:10.3389/fnbeh.2018.00306
- Hol T, Niesink M, van Ree JM, Spruijt BM, 1996. Prenatal exposure to morphine affects juvenile play behavior and adult social behavior in rats. Pharmacol. Biochem. Behav., Neurobehavioral Teratology 55, 615–618. doi:10.1016/S0091-3057(96)00274-2
- Holdsworth EA, Appleton AA, 2020. Adverse childhood experiences and reproductive strategies in a contemporary U.S. population. Am. J. Phys. Anthropol 171, 37–49. doi:10.1002/ajpa.23967 [PubMed: 31710705]
- Hormozi A, Zarifkar A, Tatar M, Barazesh M, Rostami B, 2018. Effects of Post-Weaning Chronic Stress on Nociception, Spinal Cord μ-Opioid, and α2-Adrenergic Receptors Expression in Rats and Their Offspring. J. Mol. Neurosci 64, 567–573. doi:10.1007/s12031-018-1068-4 [PubMed: 29700767]
- Hung C-J, Wu C-C, Chen W-Y, Chang C-Y, Kuan Y-H, Pan H-C, Liao S-L, Chen C-J, 2013. Depression-like effect of prenatal buprenorphine exposure in rats. PloS One 8, e82262. doi:10.1371/journal.pone.0082262 [PubMed: 24367510]
- Hutchinson MR, Bland ST, Johnson KW, Rice KC, Maier SF, Watkins LR, 2007. Opioid-induced glial activation: mechanisms of activation and implications for opioid analgesia, dependence, and reward. Sci. World J 7, 98–111.
- Hutchinson MR, Northcutt AI, Hiranita T, Wang X, Lewis SS, Thomas J, Steeg K, Kopajtic TA, Loram LC, Stregola C, Galer E, Miles NE, Bland ST, Amat J, Rozeske RR, Maslanik T, Chapman TR, Strand KA, 2012. Opioid activation of toll-like receptor 4 contributes to drug reinforcement. J Neurosci 32, 11187–200. [PubMed: 22895704]
- Hutchinson MR, Shavit Y, Grace PM, Rice KC, Maier SF, Watking LR, 2011. Exploring the neuroimmunopharmacology of opioids: an integrative review of mechanisms of central immune signaling and their implications for opioid analgesia. Pharmacol Rev 63, 772–810. [PubMed: 21752874]
- Jantzie LL, Maxwell JR, Newville JC, Yellowhair TR, Kitase Y, Madurai N, Ramachandra S, Bakhireva LN, Northington FJ, Gerner G, Tekes A, Milio LA, Brigman JL, Robinson S, Allan A, 2020. Prenatal opioid exposure: The next neonatal neuroinflammatory disease. Brain. Behav. Immun 84, 45–58. doi:10.1016/j.bbi.2019.11.007 [PubMed: 31765790]
- Jaschke N, Pählig S, Pan Y-X, Hofbauer LC, Göbel A, Rachner TD, 2021. From Pharmacology to Physiology: Endocrine Functions of μ-Opioid Receptor Networks. Trends Endocrinol. Metab. TEM 32, 306–319. doi:10.1016/j.tem.2021.02.004 [PubMed: 33676828]
- Jirtle RL, Skinner MK, 2007. Environmental epigenomics and disease susceptibility. Nat. Rev. Genet 8, 253–262. doi:10.1038/nrg2045 [PubMed: 17363974]

- Jones HE, Dengler E, Garrison A, O'Grady KE, Seashore C, Horton E, Andringa K, Jansson LM, Thorp J, 2014. Neonatal outcomes and their relationship to maternal buprenorphine dose during pregnancy. Drug Alcohol Depend. 134, 414–417. doi:10.1016/j.drugalcdep.2013.11.006 [PubMed: 24290979]
- Jones HE, Kaltenbach K, Heil SH, Stine SM, Coyle MG, Arria AM, O'Grady KE, Selby P, Martin PR, Fischer G, 2010. Neonatal Abstinence Syndrome after Methadone or Buprenorphine Exposure. N. Engl. J. Med 363, 2320–2331. doi:10.1056/NEJMoa1005359 [PubMed: 21142534]
- Jones MR, Viswanath O, Peck J, Kaye AD, Gill JS, Simopoulos TT, 2018. A Brief History of the Opioid Epidemic and Strategies for Pain Medicine. Pain Ther. 7, 13–21. doi:10.1007/s40122-018-0097-6 [PubMed: 29691801]
- Jovanovic T, Smith A, Kamkwalala A, Poole J, Samples T, Norrholm SD, Ressler KJ, Bradley B, 2011. Physiological markers of anxiety are increased in children of abused mothers. J. Child Psychol. Psychiatry 52, 844–852. doi:10.1111/j.1469-7610.2011.02410.x [PubMed: 21501167]
- Kammerer M, Adams D, Castelberg B. von, Glover V, 2002. Pregnant women become insensitive to cold stress. BMC Pregnancy Childbirth 2, 8. doi:10.1186/1471-2393-2-8 [PubMed: 12437774]
- Karkhanis AN, Rose JH, Weiner JL, Jones SR, 2016. Early-Life Social Isolation Stress Increases Kappa Opioid Receptor Responsiveness and Downregulates the Dopamine System. Neuropsychopharmacology 41, 2263–2274. doi:10.1038/npp.2016.21 [PubMed: 26860203]
- Kazemi M, Sahraei H, Azamia M, Dehghani L, Bahadoran H, Tekieh E, 2011. The effect of morphine consumption on plasma corticosteron concentration and placenta development in pregnant rats. Iran. J. Reprod. Med 9, 71–76. [PubMed: 25587250]
- Keevil BG, Clifton S, Tanton C, Macdowall W, Copas AJ, Lee D, Field N, Mitchell KR, Sonnenberg P, Bancroft J, Mercer CH, Johnson AM, Wellings K, Wu FCW, 2017. Distribution of Salivary Testosterone in Men and Women in a British General Population-Based Sample: The Third National Survey of Sexual Attitudes and Lifestyles (Natsal-3). J. Endocr. Soc 1, 14–25. doi:10.1210/js.2016-1029 [PubMed: 29264442]
- Kelty E, Preen DB, 2019. Risk Factors Associated with the Occurrence of Neonatal Opioid Withdrawal Syndrome: A Review. CNS Drugs 33, 1113–1120. doi:10.1007/s40263-019-00681-9 [PubMed: 31686408]
- Kharabian Masouleh S, Eickhoff SB, Hoffstaedter F, Genon S, Alzheimer's Disease Neuroimaging Initiative, 2019. Empirical examination of the replicability of associations between brain structure and psychological variables. eLife 8. doi:10.7554/eLife.43464
- Kiecolt-Glaser JK, Gouin J-P, Weng N, Malarkey WB, Beversdorf DQ, Glaser R, 2011. Childhood Adversity Heightens the Impact of Later-Life Caregiving Stress on Telomere Length and Inflammation. Psychosom. Med 73, 16–22. doi:10.1097/PSY.0b013e31820573b6 [PubMed: 21148804]
- Kinney DK, Munir KM, Crowley DJ, Miller AM, 2008. PRENATAL STRESS AND RISK FOR AUTISM. Neurosci. Biobehav. Rev 32, 1519–1532. doi:10.1016/j.neubiorev.2008.06.004 [PubMed: 18598714]
- Kirby ML, 1983. Recovery of spinal cord volume in postnatal rats following prenatal exposure to morphine. Dev. Brain Res 6, 211–217. doi:10.1016/0165-3806(83)90060-3
- Kirkegaard H, Möller S, Wu C, Häggström J, Olsen SF, Olsen J, Nohr EA, 2020. Associations of birth size, infancy, and childhood growth with intelligence quotient at 5 years of age: a Danish cohort study. Am. J. Clin. Nutr 112, 96–105. doi:10.1093/ajcn/nqaa051 [PubMed: 32232408]
- Knapp PE, Adjan VV, Hauser KF, 2009. Cell-specific loss of kappa-opioid receptors in oligodendrocytes of the dysmyelinating jimpy mouse. Neurosci. Lett 451, 114–118. doi:10.1016/ j.neulet.2008.12.022 [PubMed: 19110031]
- Knapp PE, Hauser KF, 1996. mu-Opioid receptor activation enhances DNA synthesis in immature oligodendrocytes. Brain Res. 743, 341–345. doi:10.1016/s0006-8993(96)01097-9 [PubMed: 9017266]
- Knapp PE, Itkis OS, Zhang L, Spruce BA, Bakalkin G, Hauser KF, 2001. Endogenous opioids and oligodendroglial function: possible autocrine/paracrine effects on cell survival and development. Glia 35, 156–165. doi:10.1002/glia.1080 [PubMed: 11460271]

- Knapp PE, Maderspach K, Hauser KF, 1998. Endogenous opioid system in developing normal and jimpy oligodendrocytes: mu and kappa opioid receptors mediate differential mitogenic and growth responses. Glia 22, 189–201. doi:10.1002/(sici)1098-1136(199802)22:2<189::aidglia10>3.0.co;2-u [PubMed: 9537839]
- Kocherlakota P, 2014. Neonatal Abstinence Syndrome. Pediatrics 134, e547–e561. doi:10.1542/ peds.2013-3524 [PubMed: 25070299]
- Koss KJ, Gunnar MR, 2018. Annual Research Review: Early adversity, the HPA axis, and child psychopathology. J. Child Psychol. Psychiatry 59, 327–346. doi:10.1111/jcpp.12784 [PubMed: 28714126]
- Krans EE, Patrick SW, 2016. Opioid Use Disorder in Pregnancy: Health Policy and Practice in the Midst of an Epidemic. Obstet. Gynecol 128, 4–10. doi:10.1097/AOG.00000000001446 [PubMed: 27275812]
- Ks B, Jp I, C M, Ba N, J F, Es R, Mr M, 2013. Power failure: why small sample size undermines the reliability of neuroscience. Nat. Rev. Neurosci 14, 365–376. doi:10.1038/nrn3475 [PubMed: 23571845]
- Kudryavtseva NN, Gerrits MAFM, Avgustinovich DF, Tenditnik MV, Van Ree JM, 2004. Modulation of anxiety-related behaviors by mu- and kappa-opioid receptor agonists depends on the social status of mice. Peptides 25, 1355–1363. doi:10.1016/j.peptides.2004.05.005 [PubMed: 15350704]
- Lacagnina MJ, Rivera PD, Bilbo SD, 2017. Glial and neuroimmune mechanisms as critical modulators of drug use and abuse. Neuropsychopharmacology 42, 156–77. [PubMed: 27402494]
- Lambertini L, Chen J, Nomura Y, 2015. Mitochondrial Gene Expression Profiles Are Associated with Maternal Psychosocial Stress in Pregnancy and Infant Temperament. PLOS ONE 10, e0138929. doi:10.1371/journal.pone.0138929 [PubMed: 26418562]
- Land BB, Bruchas MR, Lemos JC, Xu M, Melief EJ, Chavkin C, 2008. The Dysphoric Component of Stress Is Encoded by Activation of the Dynorphin κ-Opioid System. J. Neurosci 28, 407–414. doi:10.1523/JNEUROSCI.4458-07.2008 [PubMed: 18184783]
- Land BB, Bruchas MR, Schattauer S, Giardino WJ, Aita M, Messinger D, Hnasko TS, Palmiter RD, Chavkin C, 2009. Activation of the kappa opioid receptor in the dorsal raphe nucleus mediates the aversive effects of stress and reinstates drug seeking. Proc. Natl. Acad. Sci. U. S. A 106, 19168–19173. doi:10.1073/pnas.0910705106 [PubMed: 19864633]
- Lane N, Dean W, Erhardt S, Hajkova P, Surani A, Walter J, Reik W, 2003. Resistance of IAPs to methylation reprogramming may provide a mechanism for epigenetic inheritance in the mouse. Genes. N. Y. N 2000 35, 88–93. doi:10.1002/gene.10168
- Langridge AT, Glasson EJ, Nassar N, Jacoby P, Pennell C, Hagan R, Bourke J, Leonard H, Stanley FJ, 2013. Maternal Conditions and Perinatal Characteristics Associated with Autism Spectrum Disorder and Intellectual Disability. PLOS ONE 8, e50963. doi:10.1371/journal.pone.0050963 [PubMed: 23308096]
- Lehmer A, Yehuda R, 2018. Trauma across generations and paths to adaptation and resilience. Psychol. Trauma Theory Res. Pract. Policy 10, 22–29. doi:10.1037/tra0000302
- Lei M-K, Beach SRH, Simons RL, 2018. CHILDHOOD TRAUMA, PUBERTAL TIMING, AND CARDIOVASCULAR RISK IN ADULTHOOD. Health Psychol. Off. J. Div. Health Psychol. Am. Psychol. Assoc 37, 613–617. doi:10.1037/hea0000609
- Lester BM, Padbury JF, 2009. Third pathophysiology of prenatal cocaine exposure. Dev. Neurosci 31, 23–35. doi:10.1159/000207491 [PubMed: 19372684]
- Levine TA, Davie-Gray A, Kim HM, Lee SJ, Woodward LJ, 2021. Prenatal methadone exposure and child developmental outcomes in 2-year-old children. Dev. Med. Child Neurol, n/a. doi:10.1111/ dmcn.14808
- Levine TA, Woodward LJ, 2018. Early inhibitory control and working memory abilities of children prenatally exposed to methadone. Early Hum. Dev 116, 68–75. doi:10.1016/j.earlhumdev.2017.11.010 [PubMed: 29195088]
- Li Z, He Y, Wang D, Tang J, Chen X, 2017. Association between childhood trauma and accelerated telomere erosion in adulthood: A meta-analytic study. J. Psychiatr. Res 93, 64–71. doi:10.1016/ j.jpsychires.2017.06.002 [PubMed: 28601667]

- Lindley AA, Benson JE, Grimes C, Cole TM, Herman AA, 1999. The relationship in neonates between clinically measured head circumference and brain volume estimated from head CT-scans. Early Hum. Dev 56, 17–29. doi:10.1016/S0378-3782(99)00033-X [PubMed: 10530903]
- Lindqvist D, Epel ES, Mellon SH, Penninx BW, Révész D, Verhoeven JE, Reus VI, Lin J, Mahan L, Hough CM, Rosser R, Bersani FS, Blackburn EH, Wolkowitz OM, 2015. Psychiatric disorders and leukocyte telomere length: Underlying mechanisms linking mental illness with cellular aging. Neurosci. Biobehav. Rev 55, 333–364. doi:10.1016/j.neubiorev.2015.05.007 [PubMed: 25999120]
- Lovallo WR, Acheson A, Vincent AS, Sorocco KH, Cohoon AJ, 2018. Early life adversity diminishes the cortisol response to opioid blockade in women: Studies from the Family Health Patterns project. PLOS ONE 13, e0205723. doi:10.1371/journal.pone.0205723 [PubMed: 30312327]
- Lu X, Jin C, Yang J, Liu Q, Wu S, Li D, Guan Y, Cai Y, 2013. Prenatal and Lactational Lead Exposure Enhanced Oxidative Stress and Altered Apoptosis Status in Offspring Rats' Hippocampus. Biol. Trace Elem. Res 151, 75–84. doi:10.1007/s12011-012-9531-5 [PubMed: 23086308]
- Lutz P-E, Gross JA, Dhir SK, Maussion G, Yang J, Bramoulle A, Meaney MJ, Turecki G, 2018. Epigenetic Regulation of the Kappa Opioid Receptor by Child Abuse. Biol. Psychiatry, Opiates, Pain, and Addiction 84, 751–761. doi:10.1016/j.biopsych.2017.07.012
- Maeda A, Bateman BT, Clancy CR, Creanga AA, Leffert LR, 2014. Opioid Abuse and Dependence during Pregnancy: Temporal Trends and Obstetrical Outcomes. Anesthesiology 121, 1158–1165. doi:10.1097/ALN.000000000000472 [PubMed: 25405293]
- Magnan J, Tiberi M, 1989. Evidence for the presence of μ- and κ- but not of δ-opioid sites in the human fetal brain. Dev. Brain Res 45, 275–281. doi:10.1016/0165-3806(89)90045-X [PubMed: 2540923]
- Marek S, Tervo-Clemmens B, Calabro FJ, Montez DF, Kay BP, Hatoum AS, Donohue MR, Foran W, Miller RL, Feczko E, Miranda-Dominguez O, Graham AM, Earl EA, Perrone AJ, Cordova M, Doyle O, Moore LA, Conan G, Uriarte J, Snider K, Tam A, Chen J, Newbold DJ, Zheng A, Seider NA, Van AN, Laumann TO, Thompson WK, Greene DJ, Petersen SE, Nichols TE, Yeo BTT, Barch DM, Garavan H, Luna B, Fair DA, Dosenbach, N.U.F., 2020. Towards Reproducible Brain-Wide Association Studies (preprint). Neuroscience. doi:10.1101/2020.08.21.257758
- Marinelli PW, Quirion R, Gianoulakis C, 2004. An in vivo profile of β-endorphin release in the arcuate nucleus and nucleus accumbens following exposure to stress or alcohol. Neuroscience 127, 777– 784. doi:10.1016/j.neuroscience.2004.05.047 [PubMed: 15283974]
- Matthews SG, 2000. Antenatal glucocorticoids and programming of the developing CNS. Pediatr. Res 47, 291–300. doi:10.1203/00006450-200003000-00003 [PubMed: 10709726]
- McCormack C, Lauriola V, Feng T, Lee S, Spann M, Mitchell A, Champagne F, Monk C, 2020. Maternal childhood adversity and inflammation during pregnancy: Interactions with diet quality and depressive symptoms. Brain. Behav. Immun doi:10.1016/j.bbi.2020.09.023
- McGlone L, Mactier H, 2015. Infants of opioid-dependent mothers: neurodevelopment at six months. Early Hum. Dev 91, 19–21. doi:10.1016/j.earlhumdev.2014.10.006 [PubMed: 25460252]
- McLaughlin KA, Sheridan MA, Lambert HK, 2014. Childhood Adversity and Neural Development: Deprivation and Threat as Distinct Dimensions of Early Experience. Neurosci. Biobehav. Rev 47, 578–591. doi:10.1016/j.neubiorev.2014.10.012 [PubMed: 25454359]
- McLaughlin P, Mactier H, Gillis C, Hickish T, Parker A, Liang W-J, Osselton MD, 2017. Increased DNA Methylation of ABCB1, CYP2D6, and OPRM1 Genes in Newborn Infants of Methadone-Maintained Opioid-Dependent Mothers. J. Pediatr 190, 180–184.e1. doi:10.1016/ j.jpeds.2017.07.026 [PubMed: 28867064]
- Mehdizadeh H, Pourahmad J, Taghizadeh G, Vousooghi N, Yoonessi A, Naserzadeh P, Behzadfar L, Rouini MR, Sharifzadeh M, 2017. Mitochondrial impairments contribute to spatial learning and memory dysfunction induced by chronic tramadol administration in rat: Protective effect of physical exercise. Prog. Neuropsychopharmacol. Biol. Psychiatry 79, 426–433. doi:10.1016/j.pnpbp.2017.07.022 [PubMed: 28757160]
- Mei F, Mayoral SR, Nobuta H, Wang F, Desponts C, Lorrain DS, Xiao L, Green AJ, Rowitch D, Whistler J, Chan JR, 2016. Identification of the Kappa-Opioid Receptor as a Therapeutic Target for Oligodendrocyte Remyelination. J. Neurosci. Off. J. Soc. Neurosci 36, 7925–7935. doi:10.1523/JNEUROSCI.1493-16.2016

- Merhar SL, Parikh NA, Braimah A, Poindexter BB, Tkach J, Kline-Fath B, 2019. White Matter Injury and Structural Anomalies in Infants with Prenatal Opioid Exposue. Am. J. Neuroradiol ajnr;ajnr.A6282v1. doi:10.3174/ajnr.A6282
- Merrick MT, Ford DC, Haegerich TM, Simon T, 2020. Adverse Childhood Experiences Increase Risk for Prescription Opioid Misuse. J. Prim. Prev 41, 139–152. doi:10.1007/s10935-020-00578-0 [PubMed: 31989435]
- Messinger DS, Bauer CR, Das A, Seifer R, Lester BM, Lagasse LL, Wright LL, Shankaran S, Bada HS, Smeriglio VL, Langer JC, Beeghly M, Poole WK, 2004. The Maternal Lifestyle Study: Cognitive, Motor, and Behavioral Outcomes of Cocaine-Exposed and Opiate-Exposed Infants Through Three Years of Age. Pediatrics 113, 1677–1685. doi:10.1542/peds.113.6.1677 [PubMed: 15173491]
- Miranda JK, de la Osa N, Granero R, Ezpeleta L, 2013. Maternal Childhood Abuse, Intimate Partner Violence, and Child Psychopathology: The Mediator Role of Mothers' Mental Health. Violence Women 19, 50–68. doi:10.1177/1077801212475337
- Mitchell AM, Porter K, Christian LM, 2018. Examination of the role of obesity in the association between childhood trauma and inflammation during pregnancy. Health Psychol. Off. J. Div. Health Psychol. Am. Psychol. Assoc 37, 114–124. doi:10.1037/hea0000559
- Moe V, 2002. Foster-Placed and Adopted Children Exposed In Utero to Opiates and Other Substances: Prediction and Outcome at Four and a Half Years. J. Dev. Behav. Pediatr 23, 330–339. [PubMed: 12394521]
- Mohamed TM, Ghaffar HMA, El Husseiny RMR, 2015. Effects of tramadol, clonazepam, and their combination on brain mitochondrial complexes. Toxicol. Ind. Health 31, 1325–1333. doi:10.1177/0748233713491814 [PubMed: 23843224]
- Monnelly VJ, Anblagan D, Quigley A, Cabez MB, Cooper ES, Mactier H, Semple SI, Bastin ME, Boardman JP, 2018. Prenatal methadone exposure is associated with altered neonatal brain development. NeuroImage Clin. 18, 9–14. doi:10.1016/j.nicl.2017.12.033 [PubMed: 29326869]
- Montañez A, 2017. Beyond XX and XY: The Extraordinary Complexity of Sex Determination. Sci. Am 317, 50–51. doi:10.1038/scientificamerican0917-50 [PubMed: 28813372]
- Montoya-Barthelemy AG, Lee CD, Cundiff DR, Smith EB, 2020. COVID-19 and the Correctional Environment: The American Prison as a Focal Point for Public Health. Am. J. Prev. Med 58, 888–891. doi:10.1016/j.amepre.2020.04.001 [PubMed: 32387174]
- Moog NK, Buss C, Entringer S, Shahbaba B, Gillen DL, Hobel CJ, Wadhwa PD, 2016a. Maternal exposure to childhood trauma is associated during pregnancy with placental-fetal stress physiology. Biol. Psychiatry 79, 831–839. doi:10.1016/j.biopsych.2015.08.032 [PubMed: 26444076]
- Moog NK, Buss C, Entringer S, Shahbaba B, Gillen DL, Hobel CJ, Wadhwa PD, 2016b. Maternal Exposure to Childhood Trauma Is Associated During Pregnancy With Placental-Fetal Stress Physiology. Biol. Psychiatry 79, 831–839. doi:10.1016/j.biopsych.2015.08.032 [PubMed: 26444076]
- Moog NK, Entringer S, Rasmussen JM, Styner M, Gilmore JH, Kathmann N, Heim CM, Wadhwa PD, Buss C, 2018. Intergenerational Effect of Maternal Exposure to Childhood Maltreatment on Newborn Brain Anatomy. Biol. Psychiatry 83, 120–127. doi:10.1016/j.biopsych.2017.07.009 [PubMed: 28842114]
- Morgan HD, Sutherland HGE, Martin DIK, Whitelaw E, 1999. Epigenetic inheritance at the agouti locus in the mouse. Nat. Genet 23, 314–318. doi:10.1038/15490 [PubMed: 10545949]
- Morrison KE, Cole AB, Kane PJ, Meadows VE, Thompson SM, Bale TL, 2020. Pubertal adversity alters chromatin dynamics and stress circuitry in the pregnant brain. Neuropsychopharmacology 45, 1263–1271. doi:10.1038/s41386-020-0634-y [PubMed: 32045935]
- Morrison KE, Epperson CN, Sammel MD, Ewing G, Podcasy JS, Hantsoo L, Kim DR, Bale TL, 2017. Preadolescent Adversity Programs a Disrupted Maternal Stress Reactivity in Humans and Mice. Biol. Psychiatry, Stress and Neuroplasticity 81, 693–701. doi:10.1016/j.biopsych.2016.08.027
- Murrell A, Rakyan VK, Beck S, 2005. From genome to epigenome. Hum. Mol. Genet 14 Spec No 1, R3–R10. doi:10.1093/hmg/ddi110 [PubMed: 15809270]

- Myhre MC, Dyb GA, Wentzel-Larsen T, Grøgaard JB, Thoresen S, 2014. Maternal childhood abuse predicts externalizing behaviour in toddlers: a prospective cohort study. Scand. J. Public Health 42, 263–269. doi:10.1177/1403494813510983 [PubMed: 24265163]
- Nakamoto K, Taniguchi A, Tokuyama S, 2020. Changes in opioid receptors, opioid peptides and morphine antinociception in mice subjected to early life stress. Eur. J. Pharmacol 881, 173173. doi:10.1016/j.ejphar.2020.173173 [PubMed: 32511976]
- Narita M, Miyatake M, Narita M, Shibasaki M, Shindo K, Nakamura A, Kuzumaki N, Nagumo Y, Suzuki T, 2006. Direct evidence of astrocytic modulation in the development of rewarding effects induced by drugs of abuse. Neuropsychopharmacol 31, 2476–88.
- Nelson LF, Yocum VK, Patel KD, Qeadan F, Hsi A, Weitzen S, 2020. Cognitive Outcomes of Young Children After Prenatal Exposure to Medications for Opioid Use Disorder: A Systematic Review and Meta-analysis. JAMA Netw. Open 3, e201195–e201195. doi:10.1001/ jamanetworkopen.2020.1195 [PubMed: 32186745]
- Niesink RJ, Vanderschuren LJ, van Ree JM, 1996. Social play in juvenile rats after in utero exposure to morphine. Neurotoxicology 17, 905–912. [PubMed: 9198792]
- Niu L, Cao B, Zhu H, Mei B, Wang M, Yang Y, Zhou Y, 2009. Impaired in vivo synaptic plasticity in dentate gyrus and spatial memory in juvenile rats induced by prenatal morphine exposure. Hippocampus 19, 649–657. doi:10.1002/hipo.20540 [PubMed: 19115391]
- Noll JG, Trickett PK, Long JD, Negriff S, Susman EJ, Shalev I, Li JC, Putnam FW, 2017. Childhood Sexual Abuse and Early Timing of Puberty. J. Adolesc. Health 60, 65–71. doi:10.1016/ j.jadohealth.2016.09.008 [PubMed: 27836531]
- Nygaard E, Moe V, Slinning K, Walhovd KB, 2015. Longitudinal cognitive development of children born to mothers with opioid and polysubstance use. Pediatr. Res 78, 330–335. doi:10.1038/ pr.2015.95 [PubMed: 25978800]
- Nygaard E, Slinning K, Moe V, Walhovd KB, 2016. Behavior and Attention Problems in Eight-Year-Old Children with Prenatal Opiate and Poly-Substance Exposure: A Longitudinal Study. PLOS ONE 11, e0158054. doi:10.1371/journal.pone.0158054 [PubMed: 27336798]
- O'Connor A, O'Brien L, Watson L, 2019. Implications of perinatal buprenorphine exposure on infant head circumference at birth. J. Matern. Fetal Neonatal Med 0, 1–5. doi:10.1080/14767058.2019.1599352
- O'Donnell KJ, Meaney MJ, 2016. Fetal Origins of Mental Health: The Developmental Origins of Health and Disease Hypothesis. Am. J. Psychiatry 174, 319–328. doi:10.1176/ appi.ajp.2016.16020138 [PubMed: 27838934]
- Ohmura Y, Kuniyoshi Y, 2017. A translational model to determine rodent's age from human foetal age. Sci. Rep 7, 17248. doi:10.1038/s41598-017-17571-z [PubMed: 29222462]
- Olsson M, Pauliny A, Wapstra E, Uller T, Schwartz T, Blomqvist D, 2011. Sex Differences in Sand Lizard Telomere Inheritance: Paternal Epigenetic Effects Increases Telomere Heritability and Offspring Survival. PLOS ONE 6, e17473. doi:10.1371/journal.pone.0017473 [PubMed: 21526170]
- Om A, 2018. The opioid crisis in black and white: the role of race in our nation's recent drug epidemic. J. Public Health 40, e614–e615. doi:10.1093/pubmed/fdy103
- Ornoy A, Segal J, Bar-Hamburger R, Greenbaum C, 2001. Developmental outcome of school-age children born to mothers with heroin dependency: importance of environmental factors. Dev. Med. Child Neurol 43, 668–675. doi:10.1017/s0012162201001219 [PubMed: 11665823]
- Pastor V, Antonelli MC, Pallarés ME, 2017. Unravelling the Link Between Prenatal Stress, Dopamine and Substance Use Disorder. Neurotox. Res 31, 169–186. doi:10.1007/s12640-016-9674-9 [PubMed: 27778246]
- Patrick SW, Schiff DM, Prevention, C. on S.U.A., 2017. A Public Health Response to Opioid Use in Pregnancy. Pediatrics, doi:10.1542/peds.2016-4070
- Pertschuk LP, Ford DH, Rainford EA, 1977. Localization of methadone in fetal rat eye by the immunofluorescence technic. Exp. Eye Res 24, 547–552. doi:10.1016/0014-4835(77)90111-7 [PubMed: 326569]
- Peterson BS, Rosen T, Dingman S, Toth ZR, Sawardekar S, Hao X, Liu F, Xu D, Dong Z, Peterson JB, Ryoo JH, Serino D, Branch CA, Bansal R, 2020. Associations of Maternal Prenatal Drug Abuse

With Measures of Newborn Brain Structure, Tissue Organization, and Metabolite Concentrations. JAMA Pediatr. doi:10.1001/jamapediatrics.2020.1622

- Picard M, Prather AA, Puterman E, Cuillerier A, Coccia M, Aschbacher K, Burelle Y, Epel ES, 2018. A Mitochondrial Health Index Sensitive to Mood and Caregiving Stress. Biol. Psychiatry 84, 9–17. doi:10.1016/j.biopsych.2018.01.012 [PubMed: 29525040]
- Plant DT, Jones FW, Pariante CM, Pawlby S, 2017. Association between maternal childhood trauma and offspring childhood psychopathology: Mediation analysis from the ALSPAC cohort. Br. J. Psychiatry 211, 144–150. doi:10.1192/bjp.bp.117.198721 [PubMed: 28729357]
- Pritham UA, Paul JA, Hayes MJ, 2012. Opioid Dependency in Pregnancy and Length of Stay for Neonatal Abstinence Syndrome. J. Obstet. Gynecol. Neonatal Nurs 41, 180–190. doi:10.1111/ j.1552-6909.2011.01330.x
- Quinn K, Frueh BC, Scheidell J, Schatz D, Scanlon F, Khan MR, 2019. Internalizing and externalizing factors on the pathway from adverse experiences in childhood to non-medical prescription opioid use in adulthood. Drug Alcohol Depend. 197, 212–219. doi:10.1016/j.drugalcdep.2018.12.029 [PubMed: 30849646]
- Racine N, McDonald S, Chaput K, Tough S, Madigan S, 2020. Maternal substance use in pregnancy: Differential prediction by childhood adversity subtypes. Prev. Med 141, 106303. doi:10.1016/ j.ypmed.2020.106303 [PubMed: 33121963]
- Radford EJ, 2018. Exploring the extent and scope of epigenetic inheritance. Nat. Rev. Endocrinol 14, 345–355. doi:10.1038/s41574-018-0005-5 [PubMed: 29666451]
- Radhakrishnan R, Brown BP, Haas DM, Zang Y, Sparks C, Sadhasivam S, 2021a. Pilot study of fetal brain development and morphometry in prenatal opioid exposure and smoking on fetal MRI. J. Neuroradiol doi:10.1016/j.neurad.2020.12.004
- Radhakrishnan R, Elsaid NMH, Sadhasivam S, Reher TA, Hines AC, Yoder KK, Saykin AJ, Wu Y-C, 2021b. Resting state functional MRI in infants with prenatal opioid exposure—a pilot study. Neuroradiology 63, 585–591. doi:10.1007/s00234-020-02552-3 [PubMed: 32978671]
- Rakers F, Rupprecht S, Dreiling M, Bergmeier C, Witte OW, Schwab M, 2017. Transfer of maternal psychosocial stress to the fetus. Neurosci. Biobehav. Rev doi:10.1016/j.neubiorev.2017.02.019
- Rakyan VK, Chong S, Champ ME, Cuthbert PC, Morgan HD, Luu KVK, Whitelaw E, 2003. Transgenerational inheritance of epigenetic states at the murine Axin(Fu) allele occurs after maternal and paternal transmission. Proc. Natl. Acad. Sci. U. S. A 100, 2538–2543. doi:10.1073/ pnas.0436776100 [PubMed: 12601169]
- Rasmussen JM, Graham AM, Entringer S, Gilmore JH, Styner M, Fair DA, Wadhwa PD, Buss C, 2019. Maternal Interleukin-6 concentration during pregnancy is associated with variation in frontolimbic white matter and cognitive development in early life. NeuroImage 185, 825–835. doi:10.1016/j.neuroimage.2018.04.020 [PubMed: 29654875]
- Rehman KS, Sirianni R, Parker CR, Rainey WE, Carr BR, 2007. The regulation of adrenocorticotrophic hormone receptor by corticotropin-releasing hormone in human fetal adrenal definitive/transitional zone cells. Reprod. Sci. Thousand Oaks Calif 14, 578–587. doi:10.1177/1933719107307908
- Ridout KK, Levandowski M, Ridout SJ, Gantz L, Goonan K, Palermo D, Price LH, Tyrka AR, 2018. Early life adversity and telomere length: a meta-analysis. Mol. Psychiatry 23, 858–871. doi:10.1038/mp.2017.26 [PubMed: 28322278]
- Rijlaarsdam J, Stevens GWJM, Jansen PW, Ringoot AP, Jaddoe VWV, Hofman A, Ayer L, Verhulst FC, Hudziak JJ, Tiemeier H, 2014. Maternal Childhood Maltreatment and Offspring Emotional and Behavioral Problems: Maternal and Paternal Mechanisms of Risk Transmission. Child Maltreat. 19, 67–78. doi:10.1177/1077559514527639 [PubMed: 24642695]
- Rimanóczy Á, Šlamberová R, Vathy I, 2001. Prenatal morphine exposure alters estrogen regulation of κ receptors in the cortex and POA of adult female rats but has no effects on these receptors in adult male rats. Brain Res. 894, 154–156. doi:10.1016/S0006-8993(00)03326-6 [PubMed: 11245827]
- Roberts AL, Galea S, Austin SB, Corliss HL, Williams MA, Koenen KC, 2014. Women's experience of abuse in childhood and their children's smoking and overweight. Am. J. Prev. Med 46, 249– 258. doi:10.1016/j.amepre.2013.11.012 [PubMed: 24512863]

- Roberts AL, Gladish N, Gatev E, Jones MJ, Chen Y, MacIsaac JL, Tworoger SS, Austin SB, Tanrikut C, Chavarro JE, Baccarelli AA, Kobor MS, 2018. Exposure to childhood abuse is associated with human sperm DNA methylation. Transl. Psychiatry 8, 1–11. doi:10.1038/s41398-018-0252-1 [PubMed: 29317594]
- Roberts AL, Lyall K, Rich-Edwards JW, Ascherio A, Weisskopf MG, 2013. Association of maternal exposure to childhood abuse with elevated risk for autism in offspring. JAMA Psychiatry 70, 508–515. doi:10.1001/jamapsychiatry.2013.447 [PubMed: 23553149]
- Rodgers AB, Morgan CP, Bronson SL, Revello S, Bale TL, 2013. Paternal Stress Exposure Alters Sperm MicroRNA Content and Reprograms Offspring HPA Stress Axis Regulation. J. Neurosci 33, 9003–9012. doi:10.1523/JNEUROSCI.0914-13.2013 [PubMed: 23699511]
- Rodgers AB, Morgan CP, Leu NA, Bale TL, 2015. Transgenerational epigenetic programming via sperm microRNA recapitulates effects of paternal stress. Proc. Natl. Acad. Sci 112, 13699– 13704. doi:10.1073/pnas.1508347112 [PubMed: 26483456]
- Rudolph MD, Graham AM, Feczko E, Miranda-Dominguez O, Rasmussen JM, Nardos R, Entringer S, Wadhwa PD, Buss C, Fair DA, 2018. Maternal IL-6 during pregnancy can be estimated from newborn brain connectivity and predicts future working memory in offspring. Nat. Neurosci 21, 765–772. doi:10.1038/s41593-018-0128-y [PubMed: 29632361]
- Russell JA, Douglas AJ, Brunton PJ, 2008. Reduced Hypothalamo-pituitary-adrenal Axis Stress Responses in Late Pregnancy. Ann. N. Y. Acad. Sci 1148, 428–438. doi:10.1196/annals.1410.032 [PubMed: 19120138]
- Salzwedel A, Chen G, Chen Y, Grewen K, Gao W, 2020. Functional dissection of prenatal drug effects on baby brain and behavioral development. Hum. Brain Mapp n/a. doi:10.1002/hbm.25158
- SAMHSA, 2016. A Collaborative Approach to the Treatment of Pregnant Women with Opioid Use Disorders | Publications and Digital Products [WWW Document]. URL https://store.samhsa.gov/product/A-Collaborative-Approach-to-the-Treatment-of-Pregnant-Women-with-Opioid-Use-Disorders/SMA16-4978 (accessed 9.18.20).
- Sanchez-Delgado M, Court F, Vidal E, Medrano J, Monteagudo-Sánchez A, Martin-Trujillo A, Tayama C, Iglesias-Platas I, Kondova I, Bontrop R, Poo-Llanillo ME, Marques-Bonet T, Nakabayashi K, Simón C, Monk D, 2016. Human Oocyte-Derived Methylation Differences Persist in the Placenta Revealing Widespread Transient Imprinting. PLoS Genet. 12, e1006427. doi:10.1371/journal.pgen.1006427 [PubMed: 27835649]
- Sandman CA, Davis EP, Glynn LM, 2012. Psychobiological Stress and Preterm Birth, in: Morrison J (Ed.), Preterm Birth: Mother and Child. InTech, pp. 95–124.
- Sandman CA, Glynn L, Schetter CD, Wadhwa P, Garite T, Chicz-DeMet A, Hobel C, 2006. Elevated maternal cortisol early in pregnancy predicts third trimester levels of placental corticotropin releasing hormone (CRH): priming the placental clock. Peptides 27, 1457–1463. doi:10.1016/ j.peptides.2005.10.002 [PubMed: 16309788]
- Santoro TN, Santoro JD, 2018. Racial Bias in the US Opioid Epidemic: A Review of the History of Systemic Bias and Implications for Care. Cureus 10. doi:10.7759/cureus.3733
- Sarkaki A, Assaei R, Motamedi F, Badavi M, Pajouhi N, 2008. Effect of parental morphine addiction on hippocampal long-term potentiation in rats offspring. Behav. Brain Res 186, 72–77. doi:10.1016/j.bbr.2007.07.041 [PubMed: 17868930]
- Savulich G, Riccelli R, Passamonti L, Correia M, Deakin JFW, Elliott R, Flechais RSA, Lingford-Hughes AR, McGonigle J, Murphy A, Nutt DJ, Orban C, Paterson LM, Reed LJ, Smith DG, Suckling J, Tait R, Taylor EM, Sahakian BJ, Robbins TW, Ersche KD, 2017. Effects of naltrexone are influenced by childhood adversity during negative emotional processing in addiction recovery. Transl. Psychiatry 7, e1054. doi:10.1038/tp.2017.34 [PubMed: 28267152]
- Schneider JFL, Il'yasov KA, Hennig J, Martin E, 2004. Fast quantitative diffusion-tensor imaging of cerebral white matter from the neonatal period to adolescence. Neuroradiology 46, 258–266. doi:10.1007/s00234-003-1154-2 [PubMed: 14999435]
- Schreier HMC, Enlow MB, Ritz T, Gennings C, Wright RJ, 2015a. Childhood abuse is associated with increased hair cortisol levels among urban pregnant women. J. Epidemiol. Community Health 69, 1169–1174. doi:10.1136/jech-2015-205541 [PubMed: 26219886]

- Schreier HMC, Enlow MB, Ritz T, Gennings C, Wright RJ, 2015b. Childhood abuse is associated with increased hair cortisol levels among urban pregnant women. J. Epidemiol. Community Health 69, 1169–1174. doi:10.1136/jech-2015-205541 [PubMed: 26219886]
- Schulson M, Liu A, Björkman T, Quinton A, Mann KP, Benzie R, Peek M, Nanan R, 2014. Mid-Gestational Enlargement of Fetal Thalami in Women Exposed to Methadone during Pregnancy. Front. Surg 1. doi:10.3389/fsurg.2014.00028
- Seisenberger S, Andrews S, Krueger F, Arand J, Walter J, Santos F, Popp C, Thienpont B, Dean W, Reik W, 2012. The dynamics of genome-wide DNA methylation reprogramming in mouse primordial germ cells. Mol. Cell 48, 849–862. doi:10.1016/j.molcel.2012.11.001 [PubMed: 23219530]
- Shea AK, Streiner DL, Fleming A, Kamath MV, Broad K, Steiner M, 2007. The effect of depression, anxiety and early life trauma on the cortisol awakening response during pregnancy: Preliminary results. Psychoneuroendocrinology 32, 1013–1020. doi:10.1016/ j.psyneuen.2007.07.006 [PubMed: 17855000]
- Shenoy SS, Lui F, 2020. Biochemistry, Endogenous Opioids, in: StatPearls. StatPearls Publishing, Treasure Island (FL).
- Sies H, Berndt C, Jones DP, 2017. Oxidative Stress. Annu. Rev. Biochem 86, 715–748. doi:10.1146/ annurev-biochem-061516-045037 [PubMed: 28441057]
- Simmons SC, Shepard RD, Gouty S, Langlois LD, Flerlage WJ, Cox BM, Nugent FS, 2020. Early life stress dysregulates kappa opioid receptor signaling within the lateral habenula. Neurobiol. Stress 13, 100267. doi:10.1016/j.ynstr.2020.100267 [PubMed: 33344720]
- Skumlien M, Ibsen IO, Kesmodel US, Nygaard E, 2020. Sex Differences in Early Cognitive Development After Prenatal Exposure to Opioids. J. Pediatr. Psychol 45, 475–485. doi:10.1093/ jpepsy/jsaa008 [PubMed: 32324876]
- Šlamberová R, Rimanóczy Á, Bar N, Schindler CJ, Vathy I, 2003. Density of μ-opioid receptors in the hippocampus of adult male and female rats is altered by prenatal morphine exposure and gonadal hormone treatment. Hippocampus 13, 461–471. doi:10.1002/hipo.10076 [PubMed: 12836915]
- Šlamberová R, Rimanóczy Á, Cao D, Schindler CJ, Vathy I, 2005. Alterations of prenatal morphine exposure in μ-opioid receptor density in hypothalamic nuclei associated with sexual behavior. Brain Res. Bull 65, 479–485. doi:10.1016/j.brainresbull.2005.02.030 [PubMed: 15862919]
- Slotkin RK, Martienssen R, 2007. Transposable elements and the epigenetic regulation of the genome. Nat. Rev. Genet 8, 272–285. doi:10.1038/nrg2072 [PubMed: 17363976]
- Smallwood SA, Tomizawa S, Krueger F, Ruf N, Carli N, Segonds-Pichon A, Sato S, Hata K, Andrews SR, Kelsey G, 2011. Dynamic CpG island methylation landscape in oocytes and preimplantation embryos. Nat. Genet 43, 811–814. doi:10.1038/ng.864 [PubMed: 21706000]
- Smith ZD, Chan MM, Mikkelsen TS, Gu H, Gnirke A, Regev A, Meissner A, 2012. A unique regulatory phase of DNA methylation in the early mammalian embryo. Nature 484, 339–344. doi:10.1038/nature10960 [PubMed: 22456710]
- Smyser CD, Inder TE, Shimony JS, Hill JE, Degnan AJ, Snyder AZ, Neil JJ, 2010. Longitudinal Analysis of Neural Network Development in Preterm Infants. Cereb. Cortex 20, 2852–2862. doi:10.1093/cercor/bhq035 [PubMed: 20237243]
- Song P, Zhao ZQ, 2001. The involvement of glial cells in the development of morphine tolerance. Neurosci Res 39, 281–6. [PubMed: 11248367]
- Stein MD, Conti MT, Kenney S, Anderson BJ, Flori JN, Risi MM, Bailey GL, 2017. Adverse childhood experience effects on opioid use initiation, injection drug use, and overdose among persons with opioid use disorder. Drug Alcohol Depend. 179, 325–329. doi:10.1016/ j.drugalcdep.2017.07.007 [PubMed: 28841495]
- Steine IM, LeWinn KZ, Lisha N, Tylavsky F, Smith R, Bowman M, Sathyanarayana S, Karr CJ, Smith AK, Kobor M, Bush NR, 2020. Maternal exposure to childhood traumatic events, but not multi-domain psychosocial stressors, predict placental corticotrophin releasing hormone across pregnancy. Soc. Sci. Med 266, 113461. doi:10.1016/j.socscimed.2020.113461 [PubMed: 33126094]
- Substance Abuse and Mental Health Services Administration (US), Office of the Surgeon General (US), 2018. Facing Addiction in America: The Surgeon General's Spotlight on Opioids,

Publications and Reports of the Surgeon General. US Department of Health and Human Services, Washington (DC).

- Swales DA, Stout-Oswald SA, Glynn LM, Sandman C, Wing DA, Davis EP, 2018. Exposure to traumatic events in childhood predicts cortisol production among high risk pregnant women. Biol. Psychol 139, 186–192. doi:10.1016/j.biopsycho.2018.10.006 [PubMed: 30359722]
- Terplan M, Smith EJ, Glavin SH, 2010. Trends in Injection Drug Use Among Pregnant Women Admitted into Drug Treatment: 1994–2006. J. Womens Health 19, 499–505. doi:10.1089/ jwh.2009.1562
- Tester G, 2017. "And then AIDS came along": A life course turning point and sub-cohorts of older gay men. J. Gay Lesbian Soc. Serv 30, 1–16. doi:10.1080/10538720.2017.1408516 [PubMed: 30880881]
- Thomas JC, Magel C, Tomfohr-Madsen L, Madigan S, Letourneau N, Campbell TS, Giesbrecht GF, 2018. Adverse childhood experiences and HPA axis function in pregnant women. Horm. Behav 102, 10–22. doi:10.1016/j.yhbeh.2018.04.004 [PubMed: 29630895]
- Thomas-Argyriou JC, Letourneau N, Dewey D, Campbell TS, Giesbrecht GF, the APrON Study Team, 2020. The role of HPA-axis function during pregnancy in the intergenerational transmission of maternal adverse childhood experiences to child behavior problems. Dev. Psychopathol 1–17. doi:10.1017/S0954579419001767
- Towers CV, Hyatt BW, Visconti KC, Chernicky L, Chattin K, Fortner KB, 2019. Neonatal Head Circumference in Newborns With Neonatal Abstinence Syndrome. Pediatrics 143. doi:10.1542/ peds.2018-0541
- Tripathi A, Khurshid N, Kumar P, Iyengar S, 2008. Expression of δ- and μ-opioid receptors in the ventricular and subventricular zones of the developing human neocortex. Neurosci. Res 61, 257–270. doi:10.1016/j.neures.2008.03.002 [PubMed: 18455254]
- Tyrka AR, Parade SH, Price LH, Kao H-T, Porton B, Philip NS, Welch ES, Carpenter LL, 2016. Alterations of Mitochondrial DNA Copy Number and Telomere Length With Early Adversity and Psychopathology. Biol. Psychiatry, Borderline Personality Disorder: Mechanisms of Emotion Dysregulation 79, 78–86. doi:10.1016/j.biopsych.2014.12.025
- Urakubo A, Jarskog LF, Lieberman JA, Gilmore JH, 2001. Prenatal exposure to maternal infection alters cytokine expression in the placenta, amniotic fluid, and fetal brain. Schizophr. Res 47, 27–36. doi:10.1016/s0920-9964(00)00032-3 [PubMed: 11163542]
- Valentino RJ, Volkow ND, 2018. Untangling the complexity of opioid receptor function. Neuropsychopharmacology 43, 2514–2520. doi:10.1038/s41386-018-0225-3 [PubMed: 30250308]
- van den Heuvel MP, Kersbergen KJ, de Reus MA, Keunen K, Kahn RS, Groenendaal F, de Vries LS, Benders MJNL, 2015. The Neonatal Connectome During Preterm Brain Development. Cereb. Cortex 25, 3000–3013. doi:10.1093/cercor/bhu095 [PubMed: 24833018]
- Vanderschuren LJMJ, Niesink RJM, Spruijt BM, Van Ree JM, 1995. μ- and κ-opioid receptormeiated opioid effects on social play in juvenile rats. Eur. J. Pharmacol 276, 257–266. doi:10.1016/0014-2999(95)00040-R [PubMed: 7601211]
- Vassoler FM, Oranges ML, Toorie AM, Byrnes EM, 2018. Oxycodone self-administration during pregnancy disrupts the maternal-infant dyad and decreases midbrain OPRM1 expression during early postnatal development in rats. Pharmacol. Biochem. Behav 173, 74–83. doi:10.1016/ j.pbb.2018.07.009 [PubMed: 30055180]
- Vassoler FM, Wright SJ, Byrnes EM, 2016. Exposure to opiates in female adolescents alters mu opiate receptor expression and increases the rewarding effects of morphine in future offspring. Neuropharmacology 103, 112–121. doi:10.1016/j.neuropharm.2015.11.026 [PubMed: 26700246]
- Vathy I, Katay L, 1992. Effects of prenatal morphine on adult sexual behavior and brain catecholamines in rats. Brain Res. Dev. Brain Res 68, 125–131. doi:10.1016/0165-3806(92)90254-t [PubMed: 1521318]
- Vathy I, Slamberová R, Rimanóczy A, Riley MA, Bar N, 2003. Autoradiographic evidence that prenatal morphine exposure sex-dependently alters mu-opioid receptor densities in brain regions that are involved in the control of drug abuse and other motivated behaviors. Prog.

Neuropsychopharmacol. Biol. Psychiatry 27, 381–393. doi:10.1016/S0278-5846(02)00355-X [PubMed: 12691773]

- Vazquez V, Penit-Soria J, Durand C, Besson MJ, Giros B, Daugé V, 2005. Maternal deprivation increases vulnerability to morphine dependence and disturbs the enkephalinergic system in adulthood. J. Neurosci. Off. J. Soc. Neurosci 25, 4453–4462. doi:10.1523/ JNEUROSCI.4807-04.2005
- Visconti KC, Hennessy KC, Towers CV, Howard BC, 2015. Chronic opiate use in pregnancy and newborn head circumference. Am. J. Perinatol 32, 27–32. doi:10.1055/s-0034-1374817 [PubMed: 24792769]
- Volkow ND, Blanco C, 2021. The changing opioid crisis: development, challenges and opportunities. Mol. Psychiatry 26, 218–233. doi:10.1038/s41380-020-0661-4 [PubMed: 32020048]
- Wachman EM, Hayes MJ, Lester BM, Terrin N, Brown MS, Nielsen DA, Davis JM, 2014. Epigenetic variation in the mu-opioid receptor gene in infants with neonatal abstinence syndrome. J Pediatr 165, 472–8. [PubMed: 24996986]
- Wachman EM, Hunter RG, Shrestha H, Lapp HE, Meyer J, Alvarez CD, Tronick E, 2020. Maternal hair cortisol levels as a novel predictor of neonatal abstinence syndrome severity: A pilot feasibility study. Dev. Psychobiol 62, 116–122. doi:10.1002/dev.21896 [PubMed: 31342518]
- Wadhwa PD, Garite TJ, Porto M, Glynn L, Chicz-DeMet A, Dunkel-Schetter C, Sandman CA, 2004. Placental corticotropin-releasing hormone (CRH), spontaneous preterm birth, and fetal growth restriction: a prospective investigation. Am. J. Obstet. Gynecol 191, 1063–1069. doi:10.1016/ j.ajog.2004.06.070 [PubMed: 15507922]
- Walhovd KB, Watts R, Amlien I, Woodward LJ, 2012. Neural Tract Development of Infants Born to Methadone-Maintained Mothers. Pediatr. Neurol 47, 1–6. doi:10.1016/ j.pediatrneurol.2012.04.008 [PubMed: 22704008]
- Walsh K, Basu A, Werner E, Lee S, Feng T, Osborne LM, Rainford A, Gilchrist M, Monk C, 2016. Associations Among Child Abuse, Depression, and Interleukin 6 in Pregnant Adolescents. Psychosom. Med 78, 920–930. doi:10.1097/PSY.0000000000000344 [PubMed: 27187846]
- Wang X, Dow-Edwards D, Anderson V, Minkoff H, Hurd YL, 2006. Discrete opioid gene expression impairment in the human fetal brain associated with maternal marijuana use. Pharmacogenomics J. 6, 255–264. doi:10.1038/sj.tpj.6500375 [PubMed: 16477274]
- Wang X, Loran LC, Ramos K, Jess AJ, Thomas J, Cheng K, Reddy A, Somogyi AA, Hutchinson MR, Watkins LR, Yin H, 2012. Morphine activates neuroinflammation in a manner parallel to endotoxin. PNAS 109, 6325–30. [PubMed: 22474354]
- Wang X, Luo H, Chen C, Chen K, Wang J, Cai Y, Zheng S, Yang X, Zhou L, Jose PA, Zeng C, 2014. Prenatal lipopolysaccharide exposure results in dysfunction of the renal dopamine D1 receptor in offspring. Free Radic. Biol. Med 76, 242–250. doi:10.1016/j.freeradbiomed.2014.08.010 [PubMed: 25236748]
- Wang Y, Han T-Z, 2009. Prenatal exposure to heroin in mice elicits memory deficits that can be attributed to neuronal apoptosis. Neuroscience 160, 330–338. doi:10.1016/ j.neuroscience.2009.02.058 [PubMed: 19272431]
- Wang Y, Yao Y, Nie H, He X, 2017. Implication of protein kinase C of the left intermediate medial mesopallium in memory impairments induced by early prenatal morphine exposure in one-day old chicks. Eur. J. Pharmacol 795, 94–100. doi:10.1016/j.ejphar.2016.12.011 [PubMed: 27940175]
- Winklbaur B, Baewert A, Jagsch R, Rohrmeister K, Metz V, Aeschbach Jachmann C, Thau K, Fischer G, 2009. Association between Prenatal Tobacco Exposure and Outcome of Neonates Born to Opioid-Maintained Mothers. Eur. Addict. Res 15, 150–156. doi:10.1159/000216466 [PubMed: 19420947]
- Wolffe AP, Matzke MA, 1999. Epigenetics: regulation through repression. Science 286, 481–486. doi:10.1126/science.286.5439.481 [PubMed: 10521337]
- World Health Organization, 2014. Guidelines for the identification and management of substance use and substance use disorders in pregnancy. World Health Organization, Geneva.
- Wright CM, Emond A, 2015. Head Growth and Neurocognitive Outcomes. Pediatrics 135, e1393– e1398. doi:10.1542/peds.2014-3172 [PubMed: 25986019]

- Wu C-C, Hung C-J, Lin S-Y, Wang Y-Y, Chang C-Y, Chen W-Y, Liao S-L, Raung S-L, Yang C-P, Chen C-J, 2017. Treadmill exercise alleviated prenatal buprenorphine exposure-induced depression in rats. Neurochem. Int 110, 91–100. doi:10.1016/j.neuint.2017.09.012 [PubMed: 28962922]
- Wu Y, Zhang H, Wang C, Broekman BFP, Chong Y-S, Shek LP, Gluckman PD, Meaney MJ, Fortier MV, Qiu A, 2020. Inflammatory modulation of the associations between prenatal maternal depression and neonatal brain. Neuropsychopharmacology 1–8. doi:10.1038/s41386-020-0774-0
- Yamamoto M, Komori T, Matsumoto T, Zhang K, Miyahara S, Shizuya K, Okazaki Y, 2003. Effects of single and repeated prolonged stress on mu-opioid receptor mRNA expression in rat gross hypothalamic and midbrain homogenates. Brain Res. 980, 191–196. doi:10.1016/ s0006-8993(03)02969-x [PubMed: 12867258]
- Yang Z, Ye J, Li C, Zhou D, Shen Q, Wu J, Cao L, Wang T, Cui D, He S, Qi G, He L, Liu Y, 2013. Drug addiction is associated with leukocyte telomere length. Sci. Rep 3, 1542. doi:10.1038/ srep01542 [PubMed: 23528991]
- Yehuda R, Daskalakis NP, Lehrner A, Desarnaud F, Bader HN, Makotkine I, Flory JD, Bierer LM, Meaney MJ, 2014. Influences of maternal and paternal PTSD on epigenetic regulation of the glucocorticoid receptor gene in Holocaust survivor offspring. Am. J. Psychiatry 171, 872–880. doi:10.1176/appi.ajp.2014.13121571 [PubMed: 24832930]
- Yeoh SL, Eastwood J, Wright IM, Morton R, Melhuish E, Ward M, Oei JL, 2019. Cognitive and Motor Outcomes of Children With Prenatal Opioid Exposure: A Systematic Review and Meta-analysis. JAMA Netw. Open 2, e197025–e197025. doi:10.1001/jamanetworkopen.2019.7025 [PubMed: 31298718]
- Yoon BH, Jun JK, Romero R, Park KH, Gomez R, Choi J-H, Kim I-O, 1997. Amniotic fluid inflammatory cytokines (interleukin-6, interleukin-1β, and tumor necrosis factor-α), neonatal brain white matter lesions, and cerebral palsy. Am. J. Obstet. Gynecol 177, 19–26. doi:10.1016/ S0002-9378(97)70432-0 [PubMed: 9240577]
- Yuan Q, Rubic M, Seah J, Rae C, Wright IMR, Kaltenbach K, Feller JM, Abdel-Latif ME, Chu C, Oei JL, 2014. Do maternal opioids reduce neonatal regional brain volumes? A pilot study. J. Perinatol 34, 909–913. doi:10.1038/jp.2014.111 [PubMed: 24945162]
- Zaidan H, Leshem M, Gaisler-Salomon I, 2013. Prereproductive stress to female rats alters corticotropin releasing factor type 1 expression in ova and behavior and brain corticotropin releasing factor type 1 expression in offspring. Biol. Psychiatry 74, 680–687. doi:10.1016/ j.biopsych.2013.04.014 [PubMed: 23726318]
- Zedler BK, Mann AL, Kim MM, Amick HR, Joyce AR, Murrelle EL, Jones HE, 2016. Buprenorphine compared with methadone to treat pregnant women with opioid use disorder: a systematic review and meta-analysis of safety in the mother, fetus and child. Addiction 111, 2115–2128. doi:10.1111/add.13462 [PubMed: 27223595]
- Zhang H, Largent-Milnes TM, Vanderah TW, 2020. Brain Res Bull, 155.
- Zhang H, Wong T-Y, Broekman BFP, Chong Y-S, Shek LP, Gluckman PD, Tan KH, Meaney MJ, Fortier MV, Qiu A, 2021. Maternal Adverse Childhood Experience and Depression in Relation with Brain Network Development and Behaviors in Children: A Longitudinal Study. Cereb. Cortex doi:10.1093/cercor/bhab081
- Zhang X-Q, Cui Y, Cui Y, Chen Y, Na X-D, Chen F-Y, 2012. Activation of p38 signaling in the microglia in the nucleus accumbens contributes to the acquisition and maintenance of morphine-induced conditioned place preference. Brain Behav Immun 26, 318–25. [PubMed: 22004988]
- Zhou Y, Leri F, 2016. Chapter 12 Neuroscience of opiates for addiction medicine: From stressresponsive systems to behavior, in: Ekhtiari H, Paulus M (Eds.), Progress in Brain Research, Neuroscience for Addiction Medicine: From Prevention to Rehabilitation - Constructs and Drugs. Elsevier, pp. 237–251. doi:10.1016/bs.pbr.2015.09.001
- Zhou Y, Proudnikov D, Yuferov V, Kreek MJ, 2010. Drug-induced and genetic alterations in stressresponsive systems: Implications for specific addictive diseases. Brain Res., Neuropeptides in Stress and Addiction 1314, 235–252. doi:10.1016/j.brainres.2009.11.015
- Zhu H, Belcher M, van der Harst P, 2011. Healthy aging and disease: role for telomere biology? Clin. Sci 120, 427–440. doi:10.1042/CS20100385

Zhuo H-Q, Huang L, Huang H-Q, Cai Z, 2012. Effects of chronic tramadol exposure the zebrafish brain: A proteomic study. J. Proteomics 75, 3351–3364. doi:10.1016/j.jprot.2012.03.038 [PubMed: 22507199]

Highlights

- Opioid-using pregnant individuals are often affected by many co-occurring risk factors
- Disentangling effects of prenatal opioid use from co-exposures is challenging
- Prenatal opioid use and maternal CA affect fetal brains through similar pathways

Table 1.

Common mechanistic pathways of influence for maternal opioid use during pregnancy and maternal CA on offspring brain development.

	Prenatal opioid exposure		Maternal CA	
Pathways	Main Findings	Population studied	Main findings	Population studied
Endogenous opioids	↓ MOR expression in midbrain	Rats on PND1 prenatally exposed to oxycodone (Vassoler et al., 2018)	↓ MOR expression in spinal cord and hyperalgesia	Non-stressed male offspring of female rats exposed to chronic stress post-weaning through adulthood (Hormozi et al.,2018)
	↓ MOR binding in spinal cord	Rats in early postnatal period prenatally exposed to opioids (Chiou et al., 2003; Kirby, 1983)	↓ MOR expression in spinal cord	Adult male rats exposed to chronic stress post-weaning through adulthood (Hormozi et al., 2018
			\downarrow MOR mRNA in NAc	Neonatal rats exposed to predator odor (Chang et al., 2019)
	↓ MOR density in striatum, thalamus, amygdala		\downarrow MOR mRNA in PAG	Adult mice exposed to CA (Nakamoto et al., 2020)
	↓ MOR binding and expression in whole brain		↑ MOR mRNA in NAc	Juvenile rats exposed to neonatal predator odor (Chang et al., 2019)
	↓ MOR binding in MPOA	Adult rats prenatally exposed to morphine (Vathy et al., 2003)	↓ expression of KOR in the PAG and lateral habenula	Adult rodents exposed to CA (Nakamoto et al., 2020; Simmons et al., 2020)
	↓ MOR binding in BLA	Adult male rats prenatally exposed to morphine (Šlamberová et al., 2005)		
	↑ MOR binding in spinal cord	Adult rats prenatally exposed to morphine (Bhat et al., 2006; Kirby, 1983; Vathy et al., 2003)	↓ expression of KOR in NAc	Neonatal rats exposed to predator odor (Chang et al., 2019)
	↑ MOR binding in CeA, PMCoA, and NAc		↑ expression of KOR in amygdala	Adult mice exposed to CA (Nakamoto et al., 2020)
	↑ MOR binding and expression in whole brain		↑ dynorphin-dependent KOR activation in the basolateral amygdala, NAc, dorsal raphe, and hippocampus	Adult mice exposed to chronic stress (Land et al., 2008)
	↑ MOR density in hippocampus	Female adult rats prenatally exposed to morphine (Šlamberová et al., 2003)	↑ inhibition of DA release (suggesting ↑ KOR sensitivity)	Adult rats exposed to CA (Karkhanis et al., 2016)
	↑ KOR binding in POA	Adult ovariectomized female rats prenatally exposed to morphine (Rimanóczy et al., 2001)	\downarrow DOR mRNA in PAG	Adult mice exposed to CA (Nakamoto et al., 2020)
	↑ endogenous opioid release (substantia nigra, piriform cortex, septum)	Adult rats exposed to morphine during gestation (Buisman-Pijlman et al., 2009b)	↑ DOR mRNA in NAc	Juvenile rats exposed to neonatal predator odor (Chang et al., 2019)
			↑ dynorphin levels in lateral habenula	Rats across development that were exposed to CA (Simmons et al., 2020)
			↓ cortisol increase in response to naltrexone (suggesting ↓ endogenous opioid signaling)	Adult individuals AFAB exposed to CA (Lovallo et al., 2018)

	Prenatal opioid exposure		Maternal CA	
Pathways	Main Findings	Population studied	Main findings	Population studied
			↓ morphine antinociception ↓ HPA axis response to naltrexone	Adult mice exposed to CA (Nakamoto et al., 2020) Humans AFAB with preconception CA compared to humans AFAB unexposed to CA (Lovallo et al., 2018)
HPA axis	↑ glucocorticoid levels	Pregnant rats administered daily morphine compared to opioid- naïve pregnant rats	↓ baseline cortisol immediately after waking	Pregnant humans with CA history (Bublitz et al., 2014;Bublitz and Stroud, 2012b; Shea et al., 2007; Thomas et al., 2018; Thomas- Argyriou et al., 2020)
			↑ cortisol awakening response	
			Flattened diurnal cortisol slope	
			↑ hair cortisol concentrations	Individuals in mid- to late- pregnancy (Schreier et al., 2015b; Swales et al., 2018)
	↑ maternal cortisol correlated with ↓ offspring NOWS	Pregnant humans using opioids (Wachman et al., 2020)	Steeper increase of pCRH during third trimester of pregnancy	Pregnant humans in third trimester (Moog et al., 2016a; Steine et al., 2020)
	Mixed effects of prenatal opioid exposure on offspring HPA axis activity	Animals prenatally exposed to opioids (Byrnes and Vassoler, 2018)		
Immune	[↑] systemic TNF-α, IFN-γ, IL1-β, IL-6, IL-10, CCL4, CCL16 via opioid interaction with TLR4-MD2-LPS complex	Animals administered opioids (Buchanan et al., 2010; Hutchinson et al., 2007; Wang et al., 2012; Zhang et al., 2020)	↑ serum CRP	Pregnant individuals with CA (Finy and Christian, 2018; Mitchell et al., 2018)
	↑ opioid-seeking behavior after opioid activation of glial cells	Adult rodents administered opioids (Arezoomandan et al., 2016; Bachtell et al., 2015; Hutchinson et al., 2012; Narita et al., 2006; Song and Zhao, 2001; Zhang et al., 2012)	↑ serum IL-6	
	↑ systemic TNF-α, IL-1β, IL-6, CXCL1	10-day-old rats prenatally exposed to morphine (Jantzie et al., 2020)	↑ serum IL-15	Pregnant individuals with gestational diabetes and a history of CA (Bublitz et al., 2017)
	↑ systemic IL-1β, no difference in TNF-α, IL-6, CXCL1	21-day-old rats prenatally exposed to morphine (Jantzie et al., 2020)		
	↑ brain TNF-α, IL-6, TLR4, and Myd88			
	\downarrow glial cell branching			
Oxidative stress	↑ oxidative stress	Offspring prenatally exposed to opioids across development (Aboulhoda and Hassan, 2018; Guzmán et al., 2006; Hung et al., 2013)	↑ ROS production associated with ↑ oxidative stress and ↑ mitochondrial activity	Non-pregnant individuals with CA exposure compared to unexposed controls (Boeck et al., 2016)
	↑ oxidative stress and mitochondrial damage	Animals administered opioids (Cunha-Oliveira et al., 2008; Faria et al., 2016; Mehdizadeh et al., 2017; Mohamed et al., 2015; Zhuo et al., 2012)	↑ mitochondrial activity and density	Individuals with CA exposure shortly after parturition compared to unexposed controls (Gumpp et al., 2020)
			↓ reduced placental mtDNA content associated with ↑ stress	Pregnant individuals (Brunst et al., 2017)

	Prenatal opioid exposure		Maternal CA	
Pathways	Main Findings	Population studied	Main findings	Population studied
			during pregnancy and ↑ lifetime stress	
			No changes found in mitochondrial respiration or density	Neonates with maternal CA exposure compared to unexposed controls (Gumpp et al., 2020)
Epigenetics	↓ hippocampal synaptic plasticity	Rat offspring during puberty exposed to maternal preconception, prenatal, and lactation morphine (Sarkaki et al., 2008)	Epigenetic alterations to paternal germ lines (methylation and sncRNA)	Paternal germ lines of male mice exposed to chronic stress/ odor-paired fear conditioning (Dias and Ressler, 2014; Rodgers et al., 2013), and humans exposed to childhood abuse (Roberts et al., 2018)
	↑ methvlation of	Human neonates prenatally	Lower glucocorticoid receptor sensitivity	Adult offspring of AMAB Holocaust survivors (Yehuda et al., 2014)
	ABCB1, CYP2D6, MOR mRNA	exposed to opioids (McLaughlin et al., 2017)	↑ methylation of the NR3C1 promotor region	
	↓ telomere length	Human adults with chronic heroin exposure (Yang et al., 2013)	↓ telomere length	4-, 12-, and 18-month-old infants with maternal CA history (Esteves et al., 2019)
			↑ CRF1 mRNA in the frontal cortex	Offspring of female rats that underwent chronic unpredictable stress in adulthood (Zaidan et al., 2013)
			↑ CRF1 expression	Female rats exposed to chronic unpredictable stress and their offspring (Zaidan et al., 2013)

Abbreviations: adenosine triphosphate (ATP)-binding cassette sub-family B member 1 (ABCB1), assigned male at birth (AMAB), assigned female at birth (AFAB), bilateral amygdala (BLA), childhood adversity (CA), chemokine ligand 4 (CCL4), chemokine ligand 16 (CCL16), central amygdaloid nuclei (CeA), corticotropin releasing factor type 1 (CRF1), corticotropin releasing hormone (CRH), chemokine CXC ligand 1 (CXCL1), cytochrome P450 family 2 subfamily D member 6 (CYP2D6), Dopamine (DA), δ -opioid receptor (DOR), early life stress (ELS), hypothalamic-pituitary-adrenal (HPA), interferon gamma (IFN- γ), interleukin 1 beta (IL1- β), interleukin 6 (IL-6), interleukin 10 (IL-10), interleukin 15 (IL-15), κ -opioid receptor (KOR), lipopolysaccharide (LPS), medial preoptic area (MPOA), mitochondrial DNA (mtDNA), μ -opioid receptor (MOR), myeloid differentiation protein 2 (MD2), myeloid differentiation primary response protein (Myd88), nucleus accumbens (NAc), nuclear-factor kappa-B (NF- κ B), neonatal opioid withdrawal syndrome (NOWS), nuclear receptor subfamily 3 group C member 1 (NR3C1), periaqueductal gray (PAG), placental corticotropin releasing hormone (pCRH), posteromedial cortical amygdaloid nuclei (PMCoA), postnatal day (PND), preoptic area (POA), reactive oxygen species (ROS), tumor necrosis factor alpha (TNF- α), toll-like receptor 4 (TLR4).