Cell-type-specific epigenetic effects of early life stress on the brain

Mouly F. Rahman1,2 and Patrick O. McGowan3,4✉

© The Author(s) 2022

Early life stress (ELS) induces long-term phenotypic adaptations that contribute to increased vulnerability to a host of neuropsychiatric disorders. Epigenetic mechanisms, including DNA methylation, histone modifications and non-coding RNA, are a proposed link between environmental stressors, alterations in gene expression, and phenotypes. Epigenetic modifications play a primary role in shaping functional differences between cell types and can be modified by environmental perturbations, especially in early development. Together with contributions from genetic variation, epigenetic mechanisms orchestrate patterns of gene expression within specific cell types that contribute to phenotypic variation between individuals. To date, many studies have provided insights into epigenetic changes resulting from ELS. However, most of these studies have examined heterogeneous brain tissue, despite evidence of cell-type-specific epigenetic modifications in phenotypes associated with ELS. In this review, we focus on rodent and human studies that have examined epigenetic modifications induced by ELS in select cell types isolated from the brain or associated with genes that have cell-type-restricted expression in neurons, microglia, astrocytes, and oligodendrocytes. Although significant challenges remain, future studies using these approaches can enable important mechanistic insight into the role of epigenetic variation in the effects of ELS on brain function.

Translational Psychiatry (2022) 12:326; https://doi.org/10.1038/s41398-022-02076-9

EPIGENETICS AND CELLULAR PROGRAMMING

At the beginning of life, the unicellular fertilized egg, or zygote, gives rise to all the cells of an organism, possessing the property of “totipotency”. As the zygote becomes an embryo, and the embryo a fetus, there is a gradual decline in totipotency of the newly divided cells, narrowing the range of cell types they can become, rendering them “pluripotent”, a function of the plasticity of epigenetic marks. As the embryo develops and cells differentiate, pluripotency declines and epigenetic marks become more stable, determining and maintaining gene expression programs that underlie cell fate. In this way, cellular programming can be defined as the epigenetic process that contributes to stem cell differentiation into mature cell types [1, 2]. As the genetic sequence is virtual identical in all cells within each individual, the epigenome of each cell orchestrates the pattern of gene expression to confer cellular identity through histone modifications, DNA methylation (DNAm), and non-coding RNA (ncRNA).

Octamers of histone proteins coil DNA to form the nucleosomes, which are themselves wound to form chromatin. Modifications of histone N-terminal tails by acetylation, methylation, phosphorylation and ubiquitination play a role in determining DNA accessibility by modifying the positively charged N-terminal tails that tightly interact with the negatively charged DNA. For example, histone acetylation and deacetylation are understood to render chromatin more or less accessible to transcription factors (TFs), leading to enhanced or reduced transcriptional activity, respectively [3]. Histone methylation has also been associated with gene silencing or activation, depending on the amino acid modified [4, 5].

Methylation of DNA is a class of DNA modification that largely occurs at cytosine bases that are followed by guanine bases (CpG sites) in the mammalian genome. Non-CpG methylation, although less frequent, has been found in embryonic stem cells [6], neurons [7] and mature oocytes [8]. At gene promoters, first exons, and first introns, DNA methylation (DNAm) can suppress gene expression by inhibiting TF binding to regulatory elements [9]. DNAm within gene bodies and internal exons involves complex interactions with TF binding sites and conformational chromatin structure [10–12], the regulation of alternative splicing [13, 14], and is positively associated with transcription, especially for ubiquitously expressed genes [12, 15]. DNAm can also suppress gene expression through other mechanisms including histone deacetylase complex recruitment, which introduces histone modifications that result in chromatin silencing [16]. Conversely, TFs themselves can regulate DNAm by binding to specific DNA sequences to protect de novo methylation or recruit DNA methyltransferases to maintain, suppress, or initiate de novo DNAm [17].

Several types of ncRNAs, including micro-RNAs (miRs) and long non-coding RNA (lncRNAs) are sometimes considered an epigenetic mechanism due to their prominent roles in epigenetic regulation [18]. miRs are short sequences of nucleotides (~22) that...
largely repress gene expression post-transcriptionally through complementary binding to their target mRNAs, of which there can be hundreds, prompting the degradation of the corresponding mRNA and ultimately reducing its protein level [19]. The regulation of other genes at the RNA level by miRs is a property shared with other common post-transcriptional regulators of gene expression that are not typically considered part of epigenetics. Other ncRNAs are involved in epigenetic regulation at the nucleus, as is the case for lncRNA [20]. lncRNAs are >100 nucleotides in length, involved in processes including chromatin remodeling, modulation of histone and DNA methylation and acetylation, pre and post-transcription and translation, and have been functionally characterized mainly in the context of cancer [21, 22]. Generally, lncRNAs are processed and operate in the cytoplasm (see review [20]), however, lncRNAs involved in X-chromosome inactivation play a prominent role in the nucleus [22, 23].

The epigenetic program arises in response to fetal environmental signals that include extrinsic and intrinsic signaling molecules and growth factors (see review [24]), genomic imprinting through DNAm and histone modifications, and the DNA sequence itself. Genotypic differences can introduce DNAm sites (e.g. the presence of cytosines) and affect the binding of TFs, which in turn influences epigenetic modifications [17]. Cycles of epigenetic processes involved in genetic imprinting and sex-chromosome dosage compensation also occur in the zygote as discussed [25] and reviewed elsewhere [2, 26]. For example, soon after fertilization, in the pre-implanted embryo, or blastocyst, there is genome-wide demethylation in somatic cells, followed by a global wave of remethylation at implantation with locus-specific changes to methylation that continue into adulthood. Overall, during these periods of cellular production and maturation, ELS factors (e.g. stress hormones, immune stimulants) can alter cell type proportions as well as induce epigenetic reprogramming, leading to long-term changes in the brain. E = embryonic day, P = post-natal day.

Fig. 1 Timeline of cellular programming in the brain and windows of susceptibility for epigenetic effects by early life stress (ELS). Early to mid-gestation, neural stem cells produce neuronal progenitors, precursors for neurons, which migrate and form synapses from mid-gestation to toddler/weanling age. Synaptogenesis starts mid-gestation, peaks in toddler/weanling age, and continues throughout life. In late gestation to toddler/weanling age, neural stem cells give rise to glial progenitors, precursors for astrocytes and oligodendrocytes, which mature from birth to toddler/weanling age. In early and mid-gestation, myeloid-derived macrophages invade the embryonic nervous system to become microglia, which mature into the neonatal period. Microglia and glia continue to have local self-renewal and proliferative capacity into adulthood. Overall, during these periods of cellular production and maturation, ELS factors (e.g. stress hormones, immune stimulants) can alter cell type proportions as well as induce epigenetic reprogramming, leading to long-term changes in the brain.
Neurons are largely post-mitotic cells; most production and migration takes place prenatally and continues to a limited degree in the postnatal period [42], with a few neurogenic zones remaining active in adulthood [43]. The formation of synapses starts mid-gestation, peaking at toddler age in humans [44], and weanling age in rodents [42], and continues throughout life. Microglia continue to have local self-renewal and proliferative capacity [35, 36], and glial progenitors continue to proliferate, migrate and mature into adulthood [45]. Neurodevelopment also involves substantial apoptosis and synaptic pruning. Most neuronal cell death occurs early to late gestation, while glial cell death, synaptic pruning and experience-dependent modification occurs mostly at the post-natal stage [29].

Epigenetic reprogramming by early life stress
Developmental programming is the process by which cellular programming is fine-tuned by environmental factors in fetal and early postnatal stages, referred to as “early life”. This fine-tuning of cellular programs involves epigenetic reprogramming, whereby the epigenetic plan is altered in a persistent manner within a given cell type, can occur in multiple cell types, does not change cell type, and is typically stable across mitotic cell divisions [46, 47]. Epigenetic reprogramming can alter transcript abundance through long-term modifications that persist even in the absence of the initial environmental trigger. Although epigenetic reprogramming can occur later in life, the early life period is more sensitive to stress, or disruptions to homeostasis, since cell fates are established during this time.

Early life stress (ELS) is an acute or chronic factor that takes place at the prenatal, perinatal and/or pre-pubertal postnatal stages and elicits stress responses. The most studied forms of ELS are inflammation-based (e.g. infection, high-fat diets, xenobiotics) and psychosocial (e.g. emotional and physical abuse or neglect, and sexual abuse) [48]. ELS causes elevations in inflammatory cytokines and/or stress hormones that either impact offspring directly or indirectly via the placenta or breastmilk as maternal factors, inducing epigenetic modifications that impact neurodevelopmental trajectories [49]. In humans, inflammation-based and psychosocial ELS factors are most closely associated with neurodevelopmental conditions including autism spectrum disorder and psychiatric disorders related to anxiety, mood and psychosis (schizophrenia) [50–58]. In parallel, evidence from animal studies show that exposure to maternal high-fat diets, immunostimulants such as the bacterial mimic lipopolysaccharide (LPS) and psychosocial stress in early life lead to long-term changes in corresponding behavioral features of these disorders [59–65]. These findings are accompanied by evidence of long-lasting changes in microglial densities, cytokine expression and phagoctytic actions involved in synaptic pruning [59, 66, 67], neurotransmission, neural connectivity, and neuron-glia interactions [68–70]. In this context, accumulating evidence indicates that lasting phenotypic changes induced by ELS involve modifications to cellular programs in the brain [57, 71].

Animal models of ELS permit investigation of in and ex vivo changes taking place in the brain that are not easily feasible in humans. However, the correspondence to human conditions of ELS may only be approximate. For example, laboratory rodents are immunologically naive, unlike humans. Additional complications concern the assessment of different forms of psychosocial ELS (e.g. verbal versus physical abuse) and factors known to influence resiliency in humans (e.g. family income or educational attainment) [72, 73] that are difficult to model in animals. Nevertheless, given the complexities of studying the human brain in the context of ELS, rodent models appear necessary to discover causal pathways and enable the development of molecular targets for therapeutics.

Epigenetic reprogramming by ELS: the current landscape
The conventional approach to studying epigenetic reprogramming by ELS involves quantifying epigenetic modifications to select genes, or whole genomes, in heterogenous brain tissue as reviewed elsewhere [48, 49, 74–77]. One classic finding is changes of levels of hippocampal DNA methylation in the promoter region of the glucocorticoid receptor (Nr3c1; GR), a gene linked to impaired negative feedback inhibition of the hypothalamic-pituitary-adrenal (HPA) axis [78], in response to ELS (e.g. childhood abuse, neglect). However, studies of heterogenous brain tissue pose limitations on defining the mechanistic roles of such epigenetic modifications (Fig. 2). First, differences in the epigenetic mark (and expression levels) of a particular gene that is expressed in multiple cell types may be difficult to interpret if the function of a given gene varies between cell types. Second, differences in the epigenetic modifications of a gene expressed in multiple cell types can be masked by ‘background noise’ resulting from constitutive epigenetic modifications related to cell type-specific functions. Returning to the previous example, GR is expressed in all cell types in the brain [79], thus levels of DNA methylation of the Nr3c1 promoter in heterogenous brain tissue are not indicative of which cell types are driving such changes. GR can exert distinct genomic and rapid non-genomic functions in different cell types, regulating inflammation in glia [80, 81] and excitability in neurons [82], for example. Furthermore, perturbations in early development can influence cell fate determination, leading to differences in cell type proportions within tissues between experimental groups [83]. For example, neonatal maternal separation has been found to deplete oligodendrocyte progenitors in adult male mice [84]. Epigenetic modifications (e.g., DNAm) measured in particular genes in heterogenous tissue could then be reflective of cell type proportions rather than epigenetic reprogramming of a particular cell type(s). To mitigate these potential confounds, some recent studies have examined cell-type-specific epigenetic modifications.

Here we review studies that have examined the effects of ELS on epigenetic modifications in cell-type-enriched isolates, or in genes from heterogenous brain tissue where expression is known to be restricted to a single cell type in the brain (Table 1). We considered ELS studies that are inflammation-based or psychosocial that took place at the prenatal (maternal gestational), perinatal, or neonatal stages of development in female and male offspring aged post-natal day (PND)1 to adulthood. We considered both human and rodent studies, although only one human study was found within our scope. As our focus is on ELS, we did not review effects on offspring due to pre-conceptional parental or transgenerational stress as reviewed elsewhere [48, 74–77]. In the brain, the primary categorization of cell types includes neurons, glia (microglia, astrocytes, oligodendrocytes), pericytes and brain epithelial cells [89, 90], however cells in these broad categories can be subdivided based on cellular functions. For example, neurons can be classified as being either glutamatergic or GABA-ergic, excitatory or inhibitory, as well as by cortical layer of origin [91–93]. For the purposes of this review, and consistent with available data in the literature, the primary categorization for cell type was used.

Epigenetic modifications induced by ELS in neurons
In the adult human brain, there is approximately a 1:1 ratio of neurons to glia, although this varies across brain regions [94, 95], similar to rodents [96]. Epigenetic alterations to particular neurons as a consequence of ELS can alter phenotypes associated with neural circuits involved in the response to stress, fear memory, and cognition [97–99]. Dysregulation or impairments of related behaviors are associated with epigenomic alterations in the brain in response to ELS. For example, there are differences in DNA methylation of genes regulating the HPA axis, monoamines, and neuropeptides in humans with childhood trauma and in animals exposed to ELS [77, 97, 99]. However, it remains unclear whether neurons are ultimately the cell types driving the observed epigenetic changes.
To date, only a few studies have isolated neurons from rodent models of ELS to assess epigenetic modifications. One study found that an adverse caregiving environment from PND1–7 led to increased brain-derived neurotrophic factor (Bdnf) exon IV DNA methylation across 12 CpG sites in neurons isolated from the medial prefrontal-cortex (PFC) of adult female but not in male rats [100], mirroring findings found in heterogenous PFC tissue [101]. BDNF is involved in learning and memory and is a vital factor in neurodevelopment. Retinoic acid receptor, also critical to neurodevelopment, specifically neural differentiation, was found in another study to have increased promoter CpG island methylation in adult neural precursor cells isolated from the dentate gyrus of adult male rats (females unexamined) that underwent neonatal maternal separation (MS) [102]. While additional findings are needed to characterize the impacts of ELS on neurons, these studies indicate that neonatal psychosocial stressors may lead to aberrant gene expression programs essential for neural development.

Some studies that have used whole brain tissue to measure ELS-induced epigenetic changes have measured levels of miRs or DNA methylation (DNAm) of genes that are only expressed in a single cell type. In adult male rats born to dams with gestational psychosocial stress, miR-133b was found to have increased levels in the hippocampus (HPC) and decreased levels in the PFC [103]. miR-133b shows neuron-enriched expression in the brain [104, 105] and is involved in promoting neurite outgrowth [106]. The same ELS in another study in adult male rats was found to increase whole brain levels of the neuron-specific miR-323 [107], which is involved in host-pathogen interactions with viruses [108]. Together, these findings provide evidence that gestational ELS induces neuron-specific epigenetic reprogramming associated with neuronal growth and immune regulation.

Effects of postnatal stress on the neuronal epigenome have mostly been examined through MS paradigms, whereby neonatal rodents are typically separated from their dam for 3–4 hours/day from PND1–10 or 16. In one study, at PND16 immediately after MS, male mice were found to have increased hippocampal histone H4 acetylation at the promoter of activity-regulated cytoskeleton-associated protein (Arc) [109], which is only expressed in neurons and plays a critical role in learning and experience-induced synaptic plasticity [110, 111]. In adulthood, in the amygdala of female but not male mice, MS led to increased DNA methylation at the promoter of the neuron-specific serotonin-1A receptor (Htr1a) [112], which modulates emotional behavior [113]. MiR-326, which has neuron-enriched expression in the neocortex [105, 114, 115] and targets the dopamine D2 receptor [116] was found to be increased in the nucleus accumbens (NAc) and reduced in the striatum of adult male rats exposed to MS alone, and MS combined with maternal chronic unpredictable stress [117]. Also in adult rats (sex undeclared) with neonatal MS, there was increased hippocampal CA1 histone H3 acetylation and decreased methylation of CpGs in the promoter of the neuron-specific [118] corticotropin releasing hormone (Crh) [64]. Similar to the latter study, which also found evidence indicative of increased CRH expression, another study in adult rats of both sexes exposed to MS showed reduced Crh promoter CpG1 and CpG2 methylation, but in the paraventricular nucleus (PVN) of the hypothalamus [119]. PVN secretion of CRH kicksstart the HPA axis stress response, while CRH binding in the HPC is primarily responsible for
Table 1. Summary of reviewed articles whereby epigenetic modifications of genes from cell-type-enriched isolates, or genes with cell-type-enriched expression from heterogenous brain tissue, were measured in offspring exposed to early life stress.

| Cell type          | Analysis                                      | Stress timing   | Animal            | Early life stress type | Age point(s) | Sex | Brain region | Findings                                                                 | Reference |
|--------------------|-----------------------------------------------|-----------------|-------------------|------------------------|--------------|-----|-------------|--------------------------------------------------------------------------|-----------|
| Neuron             | Isolates                                      | Postnatal       | Rat               | Maternal separation    | PND56        | Male| DG          | ↑ Retinoic acid receptor promoter DNAm of CpG island                     | [102]     |
|                    |                                               |                 |                   |                        |              |     |             | ↑ Bdnf IV DNAm across 12 CpG sites (females only)                        | [100]     |
|                    | Enriched expression                            | Prenatal        | Rat               | Maternal stress        | PND0         | Male| Whole brain | ↑ miR-323                                                               | [108]     |
|                    |                                               |                 |                   | Maternal stress        | PND60        | Male| HPC & PFC   | ↑ miR-133b in HPC, ↓ miR-133b in PFC                                    | [103]     |
|                    | Postnatal                                      | Mouse           | Maternal separation | PND42-56               | Both         | AMY |             | ↑ Htr1a promoter DNAm (females)                                         | [113]     |
|                    |                                               |                 |                   | Maternal separation    | PND16        | Male| HPC         | ↑ Arc promoter histone H4ac                                              | [109]     |
|                    |                                               |                 |                   | Maternal separation    | PND60        | Both| PVN & CeA   | ↓ Crh promoter CpG1 and CpG2 methylation in PVN                         | [119]     |
|                    |                                               |                 |                   | Maternal separation and/or CUS | Adult        | Male| NAC & striatum | ↑ miR-326 in NAC, ↓ miR-326 in striatum (for CUS & MS+CUS only)         | [117]     |
|                    |                                               |                 |                   | Maternal separation    | Adult        | Undeclared | Ca1         | ↑ Crh promoter histone H3ac, ↓ Crh promoter CpG methylation            | [64]      |
| Microglia          | Isolates                                      | Prenatal        | Mouse             | Maternal allergic asthma | PND35        | Female| Whole brain | WGBS, DMR, functional annotation, pathway analysis                     | [129]     |
|                    |                                               |                 |                   | Maternal separation+ handling | ~PND60       | Male| NAC         | ↓ Il10 gene CpG island methylation                                      | [131]     |
|                    |                                               |                 |                   | Neonatal alcohol       | PND6 & 90    | Combined| Mediobasal hypoT | ↑ Tnfα promoter histone H3K9ac, ↑ Il6 promoter histone H3K9ac          | [130]     |
| Oligodendrocyte    | Isolates                                      | Prenatal        | Human             | Childhood abuse        | Childhood abuse: 19–85 years Controls: 15–81 years | Combined | ACC          | ↓ LINGO3 gene CpG methylation, ↓ Pou3f1 gene CpG methylation             | [139]     |
|                    |                                               |                 |                   | Maternal stress        | PND0         | Male| Whole brain | ↑ miR-219                                                              | [108]     |
| Astrocyte          | Enriched expression                            | Prenatal        | Rat               | Maternal separation    | PND35        | Male| Frontal cortex | ↓ Gfap promoter CpG methylation                                        | [138]     |

ac acetylation, ACC anterior cingulate cortex, AMY amygdala, Arc activity-regulated cytoskeleton-associated protein, Bdnf brain derived neurotrophic factor, CeA central amygdala, Crh corticotropin releasing hormone, CUS chronic unpredictable stress, DG dentate gyrus, DNAm DNA methylation, DMR differentially methylated regions, Gfap glial fibrillary acidic protein, HPC hippocampus, Htr1a serotonin receptor 1A, HypoT hypothalamus, Il10 interleukin 10, LINGO3 leucine-rich repeat and immunoglobulin-like domain-containing nogo receptor-interacting protein 3, LPS lipopolysaccharide, miR micro-RNA, mPFC medial prefrontal cortex, MS maternal separation, NAc nucleus accumbens, PND postnatal day, Pou3f1 Pou class 3 homeobox 1, PVN paraventricular nucleus of the hypothalamus, WGBS whole genome bisulfite sequencing.
regulating glutamatergic transmission and memory function [120]. Altogether, these findings suggest that MS leads to long-term alterations to emotion and learning processes via epigenetic reprogramming of neurotransmitter receptors and neurohormones.

**Epigenetic modifications induced by ELS in microglia**

Microglia comprise about 5–12% of the total glia population in the CNS in adult humans [121] and rodents [122] depending on the brain region. In resting states, microglia are involved in synaptic remodeling, maintenance and monitoring of the CNS environment with cell surface receptors that bind to antigens, antibodies, cytokines, and hormones. When potential insults are recognized, such as infection, inflammation, and neurodegeneration, microglia become more ‘activated’, altering their morphology to become phagocytic and releasing inflammatory cytokines that alert neighboring cells and influence their functioning [123]. Towards the end of the insult, microglia release anti-inflammatory cytokines and phagocytes cellular debris. When the insult diminishes, microglia return to their ‘resting’ state. Potentiated and dysregulated states of microglial inflammation are harmful to the tissue environment and can kill healthy neurons. Ablent microglial activation has been associated with epigenetic dysregulation in anxiety, mood and autism spectrum disorders [124, 125] as well as neurodegenerative diseases [126, 127]. Notably, a similar phenotype indicative of exaggerated inflammation has been found in studies examining epigenetic effects of ELS in animal models [49, 108]. However, these studies utilized heterogenous brain tissue to form their epigenetic links. In addition to microglia, inflammation-related genes including cytokines are expressed by neurons, astrocytes and pericytes [128], making it difficult to determine the extent to which microglia drive neuroinflammatory changes in response to ELS.

Few studies have documented microglia-specific epigenetic changes in response to ELS. In microglia isolated from the whole brain of adolescent female mice subject to maternal allergic asthma exposure, whole-genome bisulfite sequencing of differentially-methylated regions showed enrichment of gene sets associated with cytokine signaling pathways [129]. In microglia isolated from the medial hypothalamus of PND6 and adult rats (sexes combined), neonatal alcohol exposure was found to increase acetylation of histone H3K9 at the promoter regions of the pro-inflammatory cytokines tumor necrosis factor alpha (TNFα) and interleukin (IL6) in baseline conditions and 2 h post-LPS challenge [130]. Similar to these findings that suggest there is an increased pro-inflammatory phenotype in offspring exposed to ELS, the anti-inflammatory cytokine IL10 showed reduced levels of CpG island methylation in microglia isolated from the NAc of adult male rats that underwent MS as well as handling, a change that was absent in whole tissue from the NAc [131]. In the whole brain of PND1 rats (sex undefined) subject to maternal LPS exposure 48 h earlier, there was reduced miR-126 and miR-146 [132], which regulate microglial inflammatory processes [107]. Since changes to both pro- and anti-inflammatory cytokines have been observed in these cell-type-specific studies, future studies can delineate how inflammatory pathways in microglia are epigenetically reprogrammed in response to ELS.

**Epigenetic modifications induced by ELS in astrocytes and oligodendrocytes**

In the human cerebral cortex, the glial population includes approximately 20% astrocytes and 75% oligodendrocytes [133], densities of which have also been examined in various brain regions in mice [134]. Astrocytes play critical roles in maintenance of homeostasis through ion buffering, immune signaling, blood-brain-barrier maintenance, regulation of neuronal synaptogenesis and removal [135]. Epigenetic dysregulation of astrocytes and reduced astrocyte cell proportions have been linked to psychiatric disorders associated with ELS, including major depressive disorder [136]. One study found that maternal separation led to reduced promoter CpG methylation of glial fibrillary acidic protein (Gfap), an intermediate filament protein expressed only in astrocytes [137], in the frontal cortex of adolescent male rats [138]. There is also some evidence of ELS-induced epigenetic reprogramming of oligodendrocytes, the myelinating cells of the CNS that enable salutary nerve conduction and axon integrity. Isolated oligodendrocytes from the anterior cingulate cortex of human suicide completers with a history of childhood abuse exhibited decreased CpG methylation of myelination-regulating genes, specifically, leucine-rich repeat and immunoglobulin-like domain-containing no go receptor-interacting protein 3 (LINGO3) and POU class 3 homeobox 1 (POU3F1), compared to controls who died of non-suicide causes [139]. An opposite effect was observed in male rats exposed to maternal gestational psychosocial stress, where there were increased levels of whole-brain miR-219 [108], which shows oligodendrocyte-enriched expression [107] and is necessary for enabling and promoting the maturation of oligodendrocyte precursor cells into myelinating oligodendrocytes [140, 141].

**Challenges and outlook**

Converging evidence supports the role of epigenetic modifications as mediators of the impact of ELS on long-term neurobiological alterations. Unlike genetic factors, the epigenome is potentially dynamic throughout life. This malleable quality of the epigenome may enable the identification of cell-specific epigenetic biomarkers of disease and aid in the development of targeted therapeutic approaches for neuropsychiatric or neurodevelopmental disorders, as is the case for certain cancers [142]. Examples of these developments include epigenetic reprogramming of rodent neuronal stem cells to ameliorate neurodegeneration [143] and de-methylation of the fragile X mental retardation (FMR) gene in human neurons derived from Fragile X syndrome patients to normalize neuronal activity [144]. Possible epigenetic treatments for neuropsychiatric conditions have also been reviewed elsewhere [145–147].

The study of cell-type-specific epigenetic modifications requires sophisticated technical approaches. Isolating a purified cell population with intact DNA/RNA/protein from the brain can necessitate large sample sizes and volumes of fresh tissue, requiring tools such as primary cell culture, density-based centrifugation, magnetic bead separation, flow cytometry, and single-cell sequencing. These methods require advanced expertise and may be difficult to access. While computational deconvolution tools exists for obtaining cell-type-specific information from bulk sequence reads from heterogenous tissue, they nonetheless provide a less accurate estimation [148].

Of course, epigenetic changes found in a certain cell type do not necessarily mean that similar changes are absent in other cell types. While measuring epigenetic changes in one cell type may provide information regarding mechanisms associated with its biological function, measuring them in multiple cell types would be needed to assess the specificity of such modifications. Absent the heroic effort of quantifying all major cell types within a brain region of interest, comparing epigenetic changes in a single cell type to those of heterogenous tissue is another potential strategy. In such a context, two types of information would be needed: 1) information pertaining to the epigenetic modification of the particular cell type (e.g., microglia) compared to whole tissue from the brain region of interest, and 2) quantification of the proportion of the cell type of interest relative to other cell types in the region of interest. This information would indicate both relative levels of the epigenetic modification in the cell type in the brain region of interest and cell proportion differences that may also contribute to the overall levels of the epigenetic modification (see Fig. 2). This could be particularly important in interpreting the observed epigenetic profiles, which may result from changes in the relative...
levels of the epigenetic modification in the cell type of interest (potentially indicating cellular reprogramming) and/or in the proportion of the cell type of interest within the region of interest. Notably, potential cell-proportion differences alone may constitute an epigenetically-determined phenotype [83, 84].

Based on the available literature, it is evident that cell-specific epigenetic changes in glia are presently very limited in the context of ELS studies, making this fertile ground for future discovery. As well, measurement of ncRNA has been limited to miRNA in the context of the analysis of cell-specific epigenetic changes due to ELS. Given the prominent role of IncRNA in cell type differentiation in the brain [149], and its association with ELS [150] and stress-related disease [22], cell-type-specific changes in IncRNA levels in the context of ELS is another important area for future inquiry.

Studying epigenetic modifications at a primary cell-specific resolution, while important, does not evade interpretational issues. As noted above, beyond the broad categorization of cell types in the brain, definitions of specific cell types are still contested [151]. Neurons can be further subdivided into categories that consider molecular, morphological, connectional, and functional properties, leading to conceptual difficulty in defining a cell type and a lack of a consensus on their taxonomy [152, 153]. Microglia are known to exist in a resting or an activation state, which can be divided into M1 and M2 states of activation that define the bounds of a spectrum of intermediate phenotypes that can vary according to where in the brain the microglia reside [123]. Thus, it is not guaranteed that similar epigenetic modifications will exist within defined cell types, even within a brain region. While single-cell sequencing can alleviate this problem by examining cells at an individual level, this does not seem to solve the problem entirely. To draw an example from cancer biology, histone methylation differences can exist among the same cell types at the center and periphery of the tumor [154]. As such, isolating particular cells involved in a ‘neural circuit’ or biological pathway may be more informative in delineating epigenetic changes contributing to particular phenotypes.

Analysis of the temporal dynamics of epigenetic modifications may also provide important information. Epigenetic modifications detected in temporal proximity to environmental factors are not always the same as the ones detected later. For example, one study found that in the PFC of female rats exposed to neonatal caregiver maltreatment, there was reduced Bdnf exon IV methylation at adolescence compared to normal care controls, but an increase at adulthood, and no differences at infancy [101]. Therefore, the temporal dynamics of epigenetic responses to environmental factors that lead to persistent effects on phenotypes is another important avenue for future research.

Another challenge concerns the interpretation of observed epigenetic modifications in relation to their role in gene expression and behavioral phenotypes. For example, such a relationship may not manifest as steady-state increases/reductions in transcript and may only be detected in specific conditions, such as after stress exposure. A study described earlier demonstrates this possibility; neonatal alcohol exposure was found to increase histone acetylation at Tnfα and il6 promoters in hypothalamic microglia in adult rats at baseline and post-LPS challenge, however increased transcript abundance of microglial mRNA of these genes only occurred in the post-LPS condition and not at baseline [130]. Interpretational challenges also extend to determining whether epigenetic modifications at select loci are ‘causal’ mechanisms of behavior, which may only be possible by altering target epigenetic modifications in vivo [155–157]. Tools used for cell-type-specific epigenetic editing include zinc-finger proteins, which can be fused with histone modifiers and target specific DNA sequences, and transcriptional activator-like effectors (TALEs), DNA binding proteins from bacteria that can be targeted to regulate gene transcription (see review [156]). More recent tools to alter epigenetic modifications include the CRISPR (clustered regularly interspersed short palindromic repeat)/Cas9 system, whereby the prokaryotic RNA-guided endonuclease can be targeted to a specific genomic locus using designed single guide RNA, and a catalytically dead (dCas9) is used to avoid genetic double-strand breaks [155]. The cell-type-specific expression of epigenetic modifiers (e.g., histone acetylation or methylation proteins) fused to dCas9 are then inducible by Cre recombinase, which can be transgenically or virally co-expressed [155, 156]. Epigenetic editing with CRISPR/dCas9 is not without its limitations, as it can involve off-target effects [155, 159]. However, the use of CRISPR, fusion and DNA binding proteins in combination with cell-specific analysis may help delineate the epigenetic mechanisms through which ELS leads to long-term perturbations on behavior, and improve therapeutic approaches for a variety of neuropsychiatric and neurodevelopmental disorders.

REFERENCES

1. Olynik BM, Rastegar M. The genetic and epigenetic journey of embryonic stem cells into mature neural cells. Front Genet. 2012; 3. https://doi.org/10.3389/igne.2012.00081.
2. Reik W, Surani MA. Germline and pluripotent stem cells. Cold Spring Harbor Perspect Biol. 2015;7. https://doi.org/10.1101/cshperspect.a019422.
3. Perry M, Chalkley R. Histone acetylation increases the solubility of chromatin and occurs sequentially over most of the chromatin. A novel model for the biological role of histone acetylation. J Biol Chem. 1982;257:7336–47.
4. Shi Y, Lan F, Matson C, Mulligan P, Whetstone JR, Cole PA, et al. Histone demethylation mediated by the nuclear amine oxidase homolog LSD1. Cell. 2004;119:941–53.
5. Tsukada YI, Fang J, Erdjument-Bromage H, Warren ME, Leproust EM, Park IH, et al. Histone demethylation by a family of JmjC domain-containing proteins. Nature. 2005;439:811–6.
6. Lister R, Pelizzola M, Dowen RH, Hawkins RD, Hon G, Tonti-Filippini J, et al. Human DNA methylomes at base resolution show widespread epigenomic differences. Nature. 2009;462:315.
7. Kozlenkov A, Roussos P, Timashopsky A, Barbui M, Rudchenko S, Bibikova M, et al. Differences in DNA methylation between human neuronal and glial cells are concentrated in enhancers and non-CpG sites. Nucleic Acids Res. 2014;42:109.
8. Yu B, Dong X, Gravina S, Kartal O, Schimmel T, Cohen J, et al. Genome-wide, single-cell DNA methylation reveals increased non-CpG methylation during human oocyte maturation. Stem Cell Rep. 2017;9:397.
9. Anastasiadi D, Esteve-Codina A, Pifferer F. Consistent inverse correlation between DNA methylation of the first intron and gene expression across tissues and species. Epigenetics Chromatin. 2018;11:37.
10. Chuang T-I, Chen F-C, Chen Y-Z. Position-dependent correlations between DNA methylation and the evolutionary rates of mammalian coding exons. Proc Natl Acad Sci USA. 2012;109:15841–6.
11. Li S, Zhang J, Huang S, He X. Genome-wide analysis reveals that exon methylation facilitates its selective usage in the human transcriptome. Brief Bioinform. 2018;19:754–64.
12. Jingoo D, Conley A, Yi S, Lunyak V, Jordan I. On the presence and role of human gene-body DNA methylation. Oncotarget. 2012;3:642–74.
13. Maunakea AK, Chepelev I, Cui K, Zhao K. Intragenic DNA methylation modulates alternative splicing by recruiting MeCP2 to promote exon recognition. Cell Res. 2013;23:1256–69.
14. Shavievitch R, Askaday I, Keydar I, Ast G. The importance of DNA methylation of exons on alternative splicing. RNA. 2018;24:1351–62.
15. Intarte L, Li J, Gao Y, Lee JH, Leproust EM, Park H, et al. Targeted and genome-scale strategies reveal gene-body methylation signatures in human cells. Nat Biotechnol. 2009;27:361–8.
16. Nan X, Ng HH, Johnson CA, Laherty CD, Turner BM, Eisenman RN, et al. Transcriptional repression by the methyl-CpG-binding protein MeCP2 involves a histone deacetylase complex. Nature. 1998;393:386–9.
17. Moore LD, Le T, Fan G. DNA methylation and its basic function. Neuropsychopharmacol. 2013;38:23–38.
18. Aristizabal MJ, Andrei I, Hallidorsdottir T, Odgers CL, McDade TW, Goldberg A, et al. Biological embedding of experience: a primer on epigenetics. Proc Natl Acad Sci USA. 2020;117:23261–9.
19. Friedman RC, Farh KKH, Burge CB, Bartel DP. Most mammalian mRNAs are conserved targets of microRNAs. Genome Res. 2009;19:2–105.
20. Statello L, Guo CJ, Chen LL, Huarte M. Gene regulation by long non-coding RNAs and its biological functions. Nat Rev Mol Cell Biol. 2020;22:96–118.
21. Sahu A, Singhal U, Chinnaiyan AM. Long non-coding RNAs in cancer: from function to translation. Trends Cancer. 2015;1:93–109.
22. Daskalakis NP, Provenzale AC, Rohebeck AC, Giulian G. Noncoding RNAs: stress, glucocorticoids, and posttraumatic stress disorder. Biol Psychiatry. 2018;83:84–65.
23. Guttmann M, Amit I, Garber M, French C, Lin MF, Feldner D, et al. Chromatin signature reveals over a thousand highly conserved large non-coding RNAs in mammals. Nature. 2009;458:223–7.
24. Leckman JF. Commentary: What does immunology have to do with brain development and the onset of myelination in the human fetal brain. Front Neuroanat. 2009;3:5.
25. Rens W, Walduck MS, Lovell FL, Ferguson-Smith MA, Ferguson-Smith AC. Epigenetic modifications on X chromosomes in marsupial and monotreme mammals and implications for dosage compensation. Proc Natl Acad Sci USA. 2010;107:7657–62.
26. Fang H, Distelche GM, Berleth JB. X Inactivation and escape: epigenetic and structural features. Front Cell Dev Biol. 2019;7:219.
27. Mayer W, Niveleau A, Walter J, Fundele R, Haaf T. Demethylation of the zygotic genome and its biological functions. Nat Rev Mol Cell Biol. 2020;21:1–12.
28. Itskovitz-Eldor J, Schuldiner M, Karsenti D, Eden A, Yanuka O, Amit I, et al. Differentiation of human embryonic stem cells into embryoid bodies compromising the three embryonic germ layers. Mol Med. 2000;6:88.
29. Stiles J, Jernigan TL. The basics of brain development. Neuroscientist. 2010;20:327.
30. Andjelkovic AV, Nikolic B, Pachter JS, Zecevic N. Macrophages/microglial cells in human central nervous system development during an immunohistochemical study. Brain Res. 1998;814:15–25.
31. Monier A, Adle-Biassette H, Delezoide AL, Enard P, Gressens P, Vermey C. Entry and distribution of microglial cells in human embryonic and fetal cerebral cortex. J Neuropathol Exp Neurol. 2007;66:372–82.
32. Menassa DA, Gomez-Nicolta D. Microglial dynamics during human brain development. Front Immunol. 2018;9. doi:10.3389/fimmu.2018.01014.
33. Ginhoux F, Greter M, Leboeuf M, Nandi S, See P, Gokhan S, et al. Fate mapping analysis reveals that adult microglia derive from primitive macrophages. Science. 2010;330:841–5.
34. Ross C, Boroviakei T. Origin and function of the yolk sac in primate embryogenesis. Nat Commun. 2020;11:1–14.
35. Utz SG, See P, Mildenberger W, Thion MS, Silvin A, Lutz M, et al. Early fate regulators, different roles. Front Cell Neurosci. 2014;8:396.
36. Khashan AS, Abel KM, McMarnane R, Pedersen MG, Webb RT, Baker PN, et al. Higher risk of offspring schizophrenia following antenatal maternal exposure to severe adverse life events. Arch Gen Psychiatry. 2008;65:146–52.
37. Bilbo SD, Tsang V. Enduring consequences of maternal obesity for brain development and psychopathology? - A commentary on O’Connor et al. (2014). J Child Psychol Psychiatry Allied Discip. 2014;55:632–4.
38. Contu L, Hawkes C. A review of the impact of maternal obesity on the cognitive function and mental health of the offspring. Int J Mol Sci. 2017;18:1093.
39. Peter RH. Synaptic density in human frontal cortex - developmental changes and its biological functions. Nat Rev Mol Cell Biol. 2020;21:1–19.
40. Bilbo SD, Tsang V. Enduring consequences of maternal obesity for brain development and psychopathology? - A commentary on O’Connor et al. (2014). J Child Psychol Psychiatry Allied Discip. 2014;55:632–4.
41. Catale C, Gironda S, Iacono LL, Carola V. Microglial function in the effects of early-life stress on brain development: potential mechanisms for behavior, synaptic plasticity, and adrenocortical stress response. Behav Brain Res. 2013;240:1–12.
42. Smith KE, Pollak SD. Early life stress and development: potential mechanisms for behavior, synaptic plasticity, and adrenocortical stress response. Behav Brain Res. 2016;304:1–10.
43. Sominsky L, Walker FR, Hodgson DM. Neonatal endotoxin exposure modifies the acoustic startle response and circulating levels of corticosterone in the adult rat but only following acute stress. J Psychiatr Res. 2008;42:1094–103.
44. Wang A, Nie W, Li H, Hou Y, Yu Z, Fan Q, et al. Epigenetic upregulation of corticotrophin-releasing hormone mediates postnatal maternal separation-induced memory deficiency. PLoS ONE. 2014;9. doi:10.1371/journal.pone.0094394.
45. Weinstock M. Prenatal stressors in rodents: effects on behavior. Neurobiol Stress. 2017;6:3–13.
46. Catala C, Gironda S, Iacono LL, Carola V. Microglial function in the effects of early-life stress on brain and behavioral development. J Clin Med. 2020;9. doi:10.3390/JC9020468.
47. Sominsky L, Walker FR, Ong UK, Tynan RJ, Walker FR, Hodgson DM. Increased microglial activation in the rat brain following neonatal exposure to a bacterial mimetic. Behavioural Brain Res. 2012;226:351–6.
48. Bale TL, Baram TZ, Brown AS, Goldstein JM, Insel TR, McCarthy MM, et al. Early life programming and neurodevelopmental disorders. Biol Psychiatry. 2010;68:314–9.
101. Blaze J, Scheuing L, Roth TL. Differential methylation of genes in the medial prefrontal cortex of developing and adult rats following exposure to maltreatment or nurturing care during infancy. Developmental Neurosci. 2013;35:306–16.

102. Boku S, Toda H, Nakagawa S, Kato A, Inoue T, Koyama T, et al. Neonatal maternal separation alters the capacity of adult neural precursor cells to differentiate into neurons via methylation of retinoic acid receptor gene promoter. Biol Psychiatry. 2015;77:335–44.

103. Monteleone MC, Adrover E, Pallarés ME, Antonelli MC, Frasch AC, Brocco MA. Prenatal stress changes the glycoprotein gmp6a gene expression and induces epigenetic changes in rat offspring brain. Epigenetics. 2014;9:152–60.

104. He M, Liu Y, Wang X, Zhang MQ, Hannon GJ, Huang ZJ. Cell-type-based analysis of MicroRNA profiles in the mouse brain. Neuron. 2012;73:35–48.

105. Pomper N, Liu Y, Hoyle ML, Dougherty JD, Miller TM. CNS microRNA profiles: a novel high-throughput approach for cell type identification. Cell. 2002;115:351–61.

106. Lu XC, Zhang JY, Tang LJ, Huang BS, Li K, Tao Y, et al. MiR-133b promotes neurite outgrowth by targeting RhoA expression. Cell Physiol Biochem. 2015;35:246–58.

107. Jovičić A, Roshan R, Moisoi N, Pradravad S, Moser R, Pillai B, et al. Comprehensively explore expression arrays of neural cell-type-specific miRNAs to identify new determinants of the specification and maintenance of neuronal phenotypes. Ann Intern Med. 2013;158:5127–37.

108. Zucchi FCR, Yao Y, Ward ID, Ilintsky S, Olson DM, Benzkes K, et al. Maternal stress induces epigenetic signatures of psychiatric and neurological diseases in the offspring. PLoS ONE. 2013;8:e56967.

109. Xie L, Korkmaz KS, Braun K, Bock J. Early life stress-induced histone acetylations correlate with activation of the synaptic plasticity genes Arc and Egr1 in the mouse hippocampus. J Neurochemistry. 2013;125:457–64.

110. Pastuzyn ED, Day CE, Kearns RB, Kyker-Smith T, Maib AI, McCormick J, et al. The neuronal gene Arc encodes a repurposed retrotransposon Gag protein that mediates intercellular RNA transfer. Cell. 2018;172:275–288.e18.

111. Steward O, Wallace CS, Lyford GL, Worley PF. Synaptic activation causes the maturation of the leg Arc locus to a transcriptionally active near activated postsynaptic sites on dendrites. Neurosci. 1998;281:741–51.

112. Albert PR. Transcriptional regulation of the S-HT1A receptor: implications for mental illness. Philos Trans R Soc B: Biol Sci. 2012;367:2402–15.

113. Mattern F, Post A, Solger F, Oacciones A, Franke A, et al. Prenatal and postnatal experiences associated with epigenetic changes in the adult mouse brain. Behavioural Brain Res. 2013;259:143–8.

114. Kefas B, Comeau L, Floyd DH, Seleverstov O, Godlewski J, Schmittgen T, et al. The neuronal microRNA miR-326 acts in a feedback loop with Notch and has therapeutic potential against brain tumors. J Neurosci. 2009;29:15161–8.

115. Kim J, Krichenovsky A, Grad Y, Hayes GD, Kosik KS, Church GM, et al. Identification of many microRNAs that copiously polypurinobos in mammalian neurons. Proc Natl Acad Sci USA. 2004;101:360–5.

116. Allen L, Dwivedi Y, MicroRNA mediators of early life stress vulnerability to depression and suicidal behavior. Mol Psychiatry. 2020;25:308–20.

117. Zhang Y, Wang Y, Wang L, Bai M, Zhang X, Zhu X. Dopamine receptor D2 and associated microRNAs are involved in stress susceptibility and resistance to esclatopram treatment. Int J Neuropsychopharmacol. 2015;18:1–10.

118. Zhou JN, Fang H. Transcriptional regulation of corticotropin-releasing hormone gene in stress response. iBRO Rep. 2018;5:137–46.

119. Chen J, Evans AN, Liu Y, Honda M, Saavedra JM, Aguilera G. Maternal deprivation in rats is associated with corticotrophin-releasing hormone (CRH) promoter hypomethylation and enhances CRH transcriptional responses to stress in adulthood. J Neuroendocrinol. 2012;24:1055–64.

120. Refojo D, Schweizer M, Kuehne C, Ehrenberg S, Thoeringer C, Vogl AM, et al. Glutamate and dopamine neurons mediate anxiogenic and anxiolytic effects of CRHR1. Science. 2011;333:1903–7.

121. Kim YS, Joh TH. Microglia, major player in the brain inflammation: their roles in the pathogenesis of Parkinson’s disease. Exp Mol Med. 2006;38:333–47.

122. Kaur C, Rathnasamy G, Ling EA. Biology of microglia in the developing brain. J Neurodevelopmental Disorders. 2013;5:48.

123. Carreira AM, Paleczna M, Kuter Z. Overview of general and discriminating markers of afferential microglial phenotypes. Front Cell Neurosci. 2020;14:198.

124. Martins-Ferreira R, Leal B, Costa PP, Ballestar E. Microglial innate memory and epigenetic reprogramming in neurological disorders. Prog Neurobiol. 2021;200:101971.

125. Frank MG, Weber MD, Watkins LR, Maier SF. Stress-induced neuroinflammatory priming: a liability factor in the etiology of psychiatric disorders. Neurobiol Stress. 2016;4:62.

126. Harbets E, Datta D, Chen S, Wohonier J, Oh U, Jacobson S. Translocator protein 18 KDa (TSPO) expression in multiple sclerosis patients. J Neuroimmunol. 2013;6:51.

127. Liu Y, Wang M, Marcora EM, Zhang B, Goate AM. Promoter DNA hypermethylation – implications for Alzheimer’s disease. Neurosci Lett. 2019;711:134403.

128. Galic MA, Riazi K, Pittman QJ. Cytokines and brain excitability. Front Neuroendocrinol. 2012;33:116–25.

129. Vogel Cernia A, Careaga M, LaSalle JM, Ashwood P. Microglia from offspring of dams with allergic asthma exhibit epigenetic alterations in genes dysregulated in asthma. GLIA. 2018;66:505–21.
130. Chastain LG, Franklin T, Gangisetty O, Cabrera MA, Mukherjee S, Shrivastava P, et al. Early life alcohol exposure primes hypothalamic microglia to later-life hypersensitivity to immune stress: possible epigenetic mechanism. Neuropsychopharmacology. 2019;44:1579–88.

131. Schwartz JM, Hutchinson MR, Bilbo SD. Early-life experience decreases drug-induced reinstatement of morphine CPP in adulthood via microglial-specific epigenetic programming of anti-inflammatory IL-10 expression. J Neurosci. 2011;31:17835–47.

132. Carloni S, Favrais G, Saliba E, Albertini MC, Chalon S, Longini M, et al. Melatonin modulates neonatal brain inflammation through endoplasmic reticulum stress, autophagy, and mir-34a/silent information regulator 1 pathway. J Pineal Res. 2016;61:370–80.

133. Pelvig DP, Pakkenberg H, Stark AK, Pakkenberg B. Neocortical glial cell numbers. Front Neuroanat. 2018;12. https://doi.org/10.3389/fnana.2018.00083.

134. Keller D, Ero C, Markram H. Cell densities in the mouse brain: a systematic review. Front Neuroanat. 2018;12:18. doi:10.3389/fnana.2018.00083.

135. Miller SJ. Astrocyte heterogeneity in the adult central nervous system. Front Cell Neurosci. 2018;12:15.

136. Ahmed OG, Shehata GA, Ali RM, Makboul R, Abd Allah ESH, Abd El-Rady NM. Functional LncRNA in OPC fate determination. PLoS Genet. 2015;11:1005669.

137. Brenner M, Kisseberth WC, Su Y, Besnard F, Messing A. GFAP promoter directs astrocyte-specific expression in transgenic mice. J Neurosci. 1994;14:1030–7.

138. Ahmed OG, Shehata GA, Ali RM, Makboul R, Abd Allah ESH, Abd El-Rady NM. Folic acid ameliorates neonatal isolation-induced autistic like behaviors in rats: Epigenetic modifications of bdnf and gfap promoters. Appl Physiol Nutr Metab. 2021;46:964–75.

139. Lutz PE, Tanti A, Gasecka A, Barnett-Burns S, Kim JJ, Zhou Y, et al. Association of early life alcohol exposure primes hypothalamic microglia to later-life defeat stress. Neuropsychopharmacology. 2018;43:272–84.

140. Fan HB, Chen LX, Ou XB, Ren CL, Wu XJ, Dong FX, et al. Transplanted mir-219-oversexpressing oligodendrocyte precursor cells promoted remyelination and improved functional recovery in a chronic demyelinated model. Sci Rep. 2017;1:1–18.

141. Bruinsma IB, van Dijk M, Bridel C, van de Lisdonk T, Haverkort SQ, Runia TF, et al. Expression of long non-coding RNAs in purifying brain cell types from the brain reveals functional LncRNA in OPC fate determination. Plas Genet. 2015;11:1005669.

142. Cheng Y, He C, Wang M, Ma X, Mo F, Yang S, et al. Targeting epigenetic regulations of bdnf and gfap promotors. Appl Physiol Nutr Metab. 2014;39:637–47.

143. Liu X, Wu H, Krzisz M, Wu X, Graef J, Muffat J, et al. Rescue of fragile X syndrome neurons by DNA methylation editing of the FMR1 gene. Cell. 2018;172:979–992.e6.

144. Millan M. An epigenetic framework for neurodevelopmental disorders: from pathogenesis to potential therapy. Neuropsychopharmacology. 2013;38:62–82.

145. Schiele MA, Gottschalk MG, Domshke K. The applied implications of epigenetics in anxiety, affective and stress-related disorders - A review and synthesis on psychosocial stress, psychotherapy and prevention. Clin Psychol Rev. 2020;77:101830.

146. Moody L, Chen H, Fan Y-X. Early-life nutritional programming of cognition—the fundamental role of epigenetic mechanisms in mediating the relation between early-life environment and learning and memory. Process Adv Nutr Rev J. 2017;8:337–50.

147. Avila Cobos F, Alquicira-Hernandez J, Powell JE, Mestdagh P, De Preter K. Benchmarking of cell type deconvolution pipelines for transcriptomics data. Nat Commun. 2020;11:1–14.

148. Dong X, Chen K, Cuevas-Diaz Duran R, You Y, Sloan SA, Zhang Y, et al. Comprehensive identification of long non-coding RNAs in purified cell types from the brain reveals functional LncRNA in OPC fate determination. PLoS Genet. 2015;11:1005669.

149. Azrza T, Jiang C, Yan Y, Bai X. The importance of non-coding RNAs in environmental stress-related developmental brain disorders: A systematic review of evidence associated with exposure to alcohol, anesthetic drugs, nicotine, and viral infections. Neurosci Biobehav Rev. 2021;128:633–47.

150. Clevens H. Cell Systems Voices What Is Your Conceptual Definition of “Cell Type” in the Context of a Mature Organism? What Is an Adult Cell Type, Really? 2017 https://doi.org/10.1016/j.cell.2017.03.006.

151. Zeng H, Sanes JR. Neuronal cell-type classification: challenges, opportunities and the path forward. Nat Rev Neurosci. 2017;18:530–46.

152. Mukamel EA, Ngai J. Perspectives on defining cell types in the brain. Curr Opin Neurobiol. 2019;56:61–68.

153. Assenov Y, Brooks D, Gerhäuser C. Intratumor heterogeneity in epigenetic patterns. Semin Cancer Biol. 2018;51:12–21.

154. Assenov Y, Brooks D, Gerhäuser C. Intratumor heterogeneity in epigenetic patterns. Semin Cancer Biol. 2018;51:12–21.

155. Liu XS, Jaenisch R. Editing the epigenome to tackle brain disorders. Trends Neurosci. 2019;42:861–70.

156. Hamilton P, Burek D, Lombrosso S, Neve R, Robison A, Nestler E, et al. Cell-type-specific epigenetic editing at the Foxb gene controls susceptibility to social defeat stress. Neuropsychopharmacology. 2018;43:272–84.

157. Zhang KH, Tee LY, Wang XG, Huang QS, Yang SH. Off-target effects in CRISPR/Cas9-mediated genome engineering. Mol Ther - Nucleic Acids. 2015;4:e264.

ACKNOWLEDGEMENTS
This work was supported by a Discovery Grant from the Natural Sciences and Engineering Research Council of Canada (NSERC) to POM and an Ontario Graduate Scholarship to MFR.

AUTHOR CONTRIBUTIONS
MFR and POM contributed to the conception, design, drafting, revisions, and final approval of the article.

COMPETING INTERESTS
The authors declare no competing interests.

ADDITIONAL INFORMATION
Correspondence and requests for materials should be addressed to Patrick O. McGowan.

Reprints and permission information is available at http://www.nature.com/reprints

Publisher’s note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit http://creativecommons.org/licenses/by/4.0/.

© The Author(s) 2022