Adult outcome of preterm birth: Implications for neurodevelopmental theories of psychosis

Lucy D. Vanes a,b,* , Robin M. Murray c , Chiara Nosarti a,b

a Centre for the Developing Brain, Department of Perinatal Imaging and Health, King’s College London, UK
b Department of Child and Adolescent Psychiatry, Institute of Psychiatry, Psychology and Neuroscience, King’s College London, UK
c Centre for the Developing Brain, Department of Perinatal Imaging and Health, King’s College London, UK

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A B S T R A C T

Preterm birth is associated with an elevated risk of developmental and adult psychiatric disorders, including psychosis. In this review, we evaluate the implications of neurodevelopmental, cognitive, motor, and social sequelae of preterm birth for developing psychosis, with an emphasis on outcomes observed in adulthood. Abnormal brain development precipitated by early exposure to the extra-uterine environment, and exacerbated by neuroinflammation, neonatal brain injury, and genetic vulnerability, can result in alterations of brain structure and function persisting into adulthood. These alterations, including abnormal regional brain volumes and white matter macro- and micro-structure, can critically impair functional (e.g. frontoparietal and thalamocortical) network connectivity in a manner characteristic of psychotic illness. The resulting executive, social, and motor dysfunctions may constitute the basis for behavioural vulnerability ultimately giving rise to psychotic symptomatology. There are many pathways to psychosis, but elucidating more precisely the mechanisms whereby preterm birth increases risk may shed light on that route consequent upon early neurodevelopmental insult.

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1. Introduction

Prenatal and perinatal complications have been implicated in the aetiology of psychosis since the 1930s (Rosanoff et al., 1934) but assumed a greater importance with the genesis of the neurodevelopmental hypothesis of schizophrenia (Lewis and Murray, 1987; Murray and Lewis, 1987; Weinberger, 1987). While the hypothesis has evolved over the years to include the complex interactions of a diverse set of genetic, environmental, and social risk factors occurring throughout development (Murray et al., 2017), a recent meta-analysis made it clear that associations between perinatal events and elevated psychosis risk remain important (Davies et al., 2020).

Being born preterm (i.e., before 37 completed weeks of gestation), very preterm (<33 weeks), or extremely preterm (<28 weeks) is associated with a heightened risk of adverse neurological and psychological outcomes from infancy to adulthood. This risk can be at least partly attributed to the sudden and premature exposure of the rapidly developing brain to the extrauterine environment during a critical period for synaptic formation, dendritic differentiation, layering of cortical neurons and glial proliferation (Kostović and Jovanov-Milošević, 2006; Kostović and Judaš, 2010). Preterm birth has been associated with severe brain injury (Volpe, 2009), but also with more subtle brain alterations involving white matter microstructure (Kelly et al., 2016), global structural topology (Batalle et al., 2017), functional connectivity (Ball et al., 2016) and cortical morphology (Ball et al., 2020). Mirroring this, outcomes of preterm birth range from subclinical psychological characteristics, such as mild inattention, to lifelong neurological disorders, such as cerebral palsy (Fawke, 2007; Johnson and Marlow, 2011).

Preterm birth is also a significant risk factor for psychiatric disorders (Johnson and Wolke, 2013; Nosarti et al., 2012; Walshe et al., 2008), though most research in this regard has focused on developmental childhood disorders such as attention-deficit hyperactivity disorder (ADHD) and autism-spectrum disorders (ASD) (Johnson and Marlow, 2011). Prospective research on adult psychiatric outcomes of preterm birth is more recent (Robinson et al., 2020; Taylor, 2017), as the generation of preterm infants who survived thanks to advances in neonatal intensive care practices (e.g. antenatal corticosteroids, surfactant therapy, and high-frequency ventilation (Manley et al., 2015)) has now reached adulthood. Hence, the need to better understand the long-term sequelae of prematurity is ever increasing.

Epidemiological studies have demonstrated an association between preterm birth and increased psychosis risk (Mathiason et al., 2011; Nosarti et al., 2012). However, the mechanism and specificity of this link are less clear. Psychosis typically first occurs in early adulthood, but is often preceded by prodromal signs of functional decline and
 attenuated psychotic symptoms (Poletti and Raballo, 2020), and is characterised by neural alterations that can be traced back to earlier stages of development (Walker and Bolliii, 2002), such as accelerated fronto-temporo grey matter loss (McIntosh et al., 2011) and reduced connectivity of large scale brain networks (Nath et al., 2020). There is a partial but notable overlap between the neural, cognitive, and behavioural profiles of adults with psychosis and those born preterm. In this review, we critically evaluate findings relating to adult outcomes of preterm birth which may shed light on the pathways to psychosis in this population. To this end, we draw in particular (but not exclusively) on findings from an unprecedented longitudinal study of individuals born very preterm and admitted to the neonatal unit of University College London Hospital (UCLH) between 1979 and 1985; this was the first cohort in the world to undergo intensive neonatal brain ultrasonography. The infants were then followed up at several time points into adulthood using neurocognitive and behavioural assessments as well as magnetic resonance imaging (MRI). Studies reporting on outcomes of the UCLH cohort in adulthood are listed in Table 1.

2. General overview: preterm birth and psychopathology

There has been substantial research on the psychological consequences of preterm birth in childhood and adolescence. Accumulating evidence in preterm-born children has converged on the identification of a so-called “preterm behavioural phenotype”, characterised by increases in anxiety, inattention, and social impairments, coupled with executive function deficits (Johnson and Marlow, 2011). This profile appears to extend into adolescence, manifesting in an increased prevalence of diagnosed ADHD, ASD, and affective disorders (Johnson and Wolke, 2013). Preterm adolescents are at a 3- to 4-fold increased risk of being diagnosed with any psychiatric disorder compared to their term-born peers, representing a prevalence of approximately 25% (Burnett et al., 2011; Johnson and Wolke, 2013). Sub-clinically, dimensional measures capture increased liability for experiencing a wide range of symptoms including social (Healy et al., 2013) and emotional difficulties (Indredavik et al., 2005) in adolescence, which may impact on daily function even in the absence of a formal diagnosis.

In adulthood, population linkage and meta-analytic studies have found that the risk of psychiatric hospitalisation is significantly related to the degree of prematurity (Lindström et al., 2009; Nosarti et al., 2012), and preterm-born young adults are substantially more likely to be diagnosed with a psychiatric disorder (Burnett et al., 2011; Mathiasen et al., 2011; Nosarti et al., 2012) or be prescribed psychotropic medication (Robinson et al., 2020) than their term-born counterparts. Comparatively few studies have assessed psychopathology dimensionally in preterm adults; however, a meta-analysis of 6 studies investigating self-reported mental health problems in adults born preterm at very low birth weight found significant increases in internalising problems and socially avoidant behaviour compared to term-born adults (Pyhältö et al., 2017). In a recent study assessing psychiatric symptoms dimensionally in the UCLH cohort, we found preterm-born adults aged 30 to have elevated total psychopathology as well as increased positive, negative, and cognitive symptoms compared to term-born controls (Kroll et al., 2018).

Despite the many studies retrospectively examining a history of pre- and perinatal events in schizophrenia, there is very little research linking preterm birth specifically to psychotic disorders and symptoms. While preterm birth is indeed associated with an increased risk of being diagnosed with non-affective psychosis in adult life (Nosarti et al., 2012), this risk does not appear to be specific to psychotic disorder at the population level. Thus, it is possible that preterm birth confers a transdiagnostic biological vulnerability to psychopathology in adulthood, preceded by a more specific behavioural profile in childhood and adolescence. Different aspects of this profile in conjunction with intervening environmental factors may precipitate different trajectories in the expression of psychopathology during the transition to adulthood, leading to increased diagnostic heterogeneity and comorbidities amongst preterm-born adults. Potential risk factors for and precursors of psychosis arising as a result of preterm birth in adulthood will now be discussed.

3. Adult sequelae of preterm birth and psychosis risk

3.1. Neurodevelopment

Preterm birth is a leading cause of brain injury, mostly due to perinatal hypoxia-ischemia, which can result in damage in particular to the periventricular white matter and basal ganglia (Back, 2015; Logothetarajah et al., 2009; Volpe, 2009). However, even in the absence of severe or obvious brain injury, preterm birth can result in subtle changes in neural architecture that are evident throughout development and into adulthood (Nagy et al., 2009; Nosarti et al., 2002; Nosarti et al., 2014), and which have been related to a range of neurocognitive difficulties (Hadaya and Nosarti, 2020; Kanel et al., 2020).

There are several overlapping brain alterations associated with preterm birth and with psychotic disorder. One of the most commonly observed abnormalities following preterm birth is ventricular enlargement (often following peri/intraventricular haemorrhage, P/IVH), with a large proportion of preterm infants showing increased ventricle size at term-equivalent age (i.e., the age at which they would have been born had they not been premature) (Hart et al., 2008). Ventricular enlargement has also been observed in preterm adolescents and adults (Allin et al., 2011; Cooke and Abernethy, 1999; Hedderich et al., 2020; Nosarti et al., 2002; Stewart et al., 1999). Neurodevelopment of regions adjacent to the lateral ventricles are preferentially disrupted following P/IVH; consequently post-haemorrhagic ventricular enlargement is associated with reduced deep grey matter volumes (Brouwer et al., 2016) and periventricular white matter damage (Larroque et al., 2003) at term-equivalent age. Evidence from the UCLH study suggests that these alterations are likely to have long-lasting effects on brain structure and function. Thus, very preterm adults exhibit an association between ventricle size and impaired microstructural integrity of widespread white matter tracts (Allin et al., 2011), and ventricular enlargement resulting from perinatal brain injury is related to abnormal frontal (Kalpakidou et al., 2014) and frontoparietal (Froudust-Walsh et al., 2015) neural activation during working memory processing.

Increased ventricle size is also one of the most replicated neuroimaging findings in chronic schizophrenia (Olabi et al., 2011; van Erp et al., 2016), with additional evidence for ventricular enlargement at the stage of the prodrome (Chung et al., 2017) and first episode of psychosis (Steen et al., 2006; Vita et al., 2006), and even in male neonates at high genetic risk for schizophrenia (Gilmore et al., 2010). It appears to worsen over the course of the illness (Kempton et al., 2010) but at least some of this is related to the effects of antipsychotics (Murray et al., 2016). Numerous studies have found associations between ventricular abnormalities and a history of obstetric complications in patients with schizophrenia (Costas-Carrera et al., 2020), with a strong likelihood that complications during delivery interact with genetic risk for psychosis in contributing to these abnormalities (Cannon et al., 1989; Falkai et al., 2003). Ventricular enlargement following preterm birth may therefore represent a marker of increased underlying vulnerability to psychosis.

Similarly, white matter abnormalities are common in both preterm and psychosis populations. The most frequent form of brain injury following preterm birth is periventricular leukomalacia (PVL), defined as either focal (necrotic) or diffuse damage to the periventricular white matter. Diffuse PVL is thought to impact on premyelinating oligodendrocytes, which can subsequently result in an impairment of axonal myelination (Iida et al., 1995; Volpe, 2009). Myelination of the brains’ connective white matter tracts is known to extend well into adulthood (Miller et al., 2012) and underpins the efficiency of neuronal signal conduction, providing a crucial basis for brain connectivity. Healthy cognitive functioning is dependent on effective communication between
| First author (year) | N | Age | Type | Tasks(s)                                                                 | Findings                                                                                                                                                                                                 |
|---------------------|---|-----|------|--------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Allin et al. (2006b) |   |     |      | Neurological assessment                                                  | VPT > FT neurological signs; associated with reduced neuropsychological function                                                                                                                         |
| Allin et al. (2006a) | 18|     |      | Personality assessment                                                  | • VPT < FT extraversion scores                                                                                                             |
| Nosarti et al. (2007) |   |     |      | Cognitive assessment                                                    | • VPT < FT neuroticism and lie scores                                                                                                           |
| Nosarti et al. (2014) | 19|     |      | Cognitive assessments                                                   | • VPT < FT executive function involving response inhibition and mental flexibility                                                            |
| Kontis et al. (2009)  |   |     |      | Psychopathology assessment                                              | • VPT > FT risk for psychiatric disorder                                                                                                        |
| Kontis et al. (2011)  |   |     |      | Neurological assessment                                                  | • VPT < FT psychological disorder were particularly vulnerable                                                                              |
| Nosarti et al. (2009) |   |     |      | Neurological assessment                                                  | • VPT > FT MD in genu of the corpus callosum; associated with lower IQ                                                                            |
| Narberhaus et al. (2009) | 20|     |      | Task-based fMRI; structural T1                                          | • Differential correlation between verbal learning and callosal FA and MD between groups                                                   |
| Lawrence et al. (2009) |   |     |      | Structural T1                                                           | No group difference in task performance                                                                                                             |
| Lawrence et al. (2010) | 20|     |      | Task-based fMRI; structural T1                                          | • VPT > FT BOLD signal in anterior cingulate, right caudate, and left inferior frontal gyrus during easy trials; in left middle frontal gyrus during hard trials |
| Lawrence et al. (2011) | 19|     |      | Diffusion MRI, cognitive assessments                                    | • VPT > FT BOLD signal in right cerebellum and anterior cingulate during recognition                                                            |
| Kalpakidou et al. (2014) | 20|     |      | Task-based fMRI                                                         | Reduced grey matter in bilateral hippocampus in VPT compared to controls                                                                  |
| Lawrence et al. (2014) | 20|     |      | Task-based fMRI; structural T1                                          | No group difference in task performance                                                                                                             |
| Nosarti et al. (2014) |   |     |      | Structural T1, cognitive assessments                                    | VPT > FT grey matter volume in left parahippocampal and precentral gyri during encoding                                                       |
| Nosarti et al. (2014) |   |     |      | Hayling Sentence Completion Test, Controlled Oral Word Association Test, Animal and Object test, Trail-Making Test, Test of Attentional Performance | • VPT > FT grey matter volume in right cerebellum, lingual, parahippocampal, and middle temporal gyri                                         |
disparate brain regions and, accordingly, structural dysconnectivity (e.g., detected with diffusion imaging techniques) is associated with impaired cognitive functioning in various populations, including preterm-born individuals (reviewed in Hadaya and Nosarti, 2020) and those with psychosis (Kelly et al., 2019). Even in the absence of severe PVL observed with conventional MRI, premature infants typically exhibit altered patterns of white matter maturation (Anjari et al., 2007; Balle et al., 2017; Kelly et al., 2016).

| First author (year) | N | Age | Type | Task(s) | Findings |
|---------------------|---|-----|------|---------|----------|
| White et al. (2014) | VPT: 29 | 28 | Resting-state fMRI, cognitive assessments | WASI, Hayling Sentence Completion Test, | • VPT < FT white matter volume in frontal, temporal, and parietal, and callosal regions
| | FT: 23 | | | | • VPT > FT white matter volume in parahippocampal and occipital cortices and cerebellum
| | | | | | • White matter volume beneath inferior frontal gyrus associated with IQ; callosal white matter and temporal grey matter volume associated with executive function
| Cole et al. (2015) | VPT: 65 | 15 | Structural T1, Psychopathology assessment | Peters’ Delusional Inventory | • No group difference in hippocampal volumes
| | FT: 36 | 19 | | | • Hippocampal subregions deformations reflecting atrophy in VPT
| | | | | | • Baseline hippocampal volume associated with delusional ideation follow-up
| Nam et al. (2015) | VPT: 160 | 15 | Structural T1, cognitive assessments | WASI; Wechsler Memory Scale Revised; California Verbal Learning Test; Controlled Oral Word Association Test; Hayling Sentence Completion Test | • VPT < FT on all tasks except verbal memory
| | FT: 88 | 20 | | | • VPT > FT cortical thickness in frontal, occipital, temporal, and insular cortices
| | | | | | • VPT < FT cortical thickness in bilateral medial temporal and left insular cortices
| | | | | | • More extensive cortical thinning in VPT from 15 to 20
| | | | | | • Cortical thinning in left temporal pole, right occipitotemporal gyrus, and superior parietal lobe associated with executive functioning
| | | | | | • VPT-PBI < FT rich-club index
| | | | | | • Simulated lesion approach suggests altered role for basal ganglia and motor connections in VPT, associated with information flow, rule learning and verbal IQ.
| | | | | | • VPT-PBI: smaller dorsal cingulum volume associated with increased BOLD signal in periventricular cortex
| | | | | | • Increased BOLD signal in periventricular cortex associated with better task performance
| | | | | | • VPT > FT rich-club index
| | | | | | • Executive function score more strongly correlated with real-life achievement measures in VPT
| Kroll et al. (2017) | VPT: 122 | Cognitive assessments | WASI; Hayling Sentence Completion Test; Controlled Oral Word Association Test; CANTAB SOC; CANTAB IED; TMT-B; Continuous Performance Test; Global Role Functioning Index | | • VPT < FT executive function and IQ
| | FT: 89 | | | | • No group difference in task performance
| Froudist-Walsh et al. (2015) | VPT: 41 | 30 | Task-based fMRI, diffusion MRI | n-back task | • VPT-PBI < FT BOLD signal in fronto-parietal regions and increased BOLD signal in periventricular cortex
| | FT: 21 | | | | • VPT-PBI: smaller dorsal cingulum volume associated with increased BOLD signal in periventricular cortex
| | | | | | • Increased BOLD signal in periventricular cortex associated with better task performance
| Karolis et al. (2016) | VPT: 51 | 30 | Diffusion MRI, cognitive assessments | WASI; Trail Making Test; Controlled Oral Word Association Test; Animal Naming test and the Object Naming test; Hayling Sentence Completion Test; Visual Paired Associate Learning task; Intra-Extra-Dimensional Shift task | • VPT > FT rich-club index
| | FT: 60 | | | | • Simulated lesion approach suggests altered role for basal ganglia and motor connections in VPT, associated with information flow, rule learning and verbal IQ.
| | | | | | • VPT < FT executive function and IQ
| | | | | | • Executive function score more strongly correlated with real-life achievement measures in VPT
| Caldinelli et al. (2017) | VPT: 84 | 19 | Diffusion MRI, cognitive assessments | California Verbal Learning Test; Visual Reproduction Test of Wechsler Memory Scale-Revised; WASI | • VPT-PBI < FT BOLD signal in fronto-parietal regions and increased BOLD signal in periventricular cortex
| | FT: 48 | | | | • VPT-PBI: smaller dorsal cingulum volume associated with increased BOLD signal in periventricular cortex
| Kroll et al. (2018) | VPT: 152 | Cognitive and psychopathology assessment | WASI, Comprehensive Assessment of At-Risk Mental States | | • VPT < FT BOLD signal in right middle frontal gyrus and posterior cingulate cortex
| | FT: 96 | | | | • VPT > FT rich-club index
| Tseng et al. (2019) | VPT: 64 | 30 | Task-based fMRI, diffusion MRI, structural T1 | Recognition memory task | • VPT < FT BOLD signal in left inferior frontal gyrus and bilateral occipital cortex
| | FT: 36 | | | | • VPT > FT rich-club index
| | | | | | • VPT > FT rich-club index
| Kroll et al. (2019) | VPT: 258 | 8-31 | Cognitive assessment | WASI | • VPT < FT rich-club index
| | FT: 36 | | | | • VPT > FT rich-club index
| Papini, et al., 2020 | VPT: 79 | 30 | Structural T1, cognitive and psychopathology assessment | WASI; Comprehensive Assessment of At-Risk Mental States | • VPT < FT cortical gyriﬁcation in frontal, anterior temporal, and occipitoparietal lobes; was associated with lower IQ and higher psychopathology.

Abbreviations: VPT = very preterm birth; FT = full term; WASI = Wechsler Abbreviated Scale of Intelligence; fMRI = functional magnetic resonance imaging; MD = mean diffusivity; FA = fractional anisotropy; BOLD = blood oxygen level dependent; PVH = periventricular haemorrhage; PBI = perinatal brain injury.
In very preterm adults enrolled in the UCLH study, we have observed abnormalities both in the microstructure (Allin et al., 2011; Froudist-Walsh et al., 2015; Tseng et al., 2019; Tseng et al., 2017) and volumes (Caldinelli et al., 2017; Nosarti et al., 2014; Tseng et al., 2017) of white matter structures compared to term-born controls. Overlapping observations have also been made in other adult cohorts, showing microstructural changes particularly in the corpus callosum and cingulum (Eikenes et al., 2011; Meng et al., 2016; Pascoe et al., 2019) but also extending to other core white matter tracts, which is further reflected in reduced global white matter volume in preterm individuals (Soria-Pastor et al., 2008). Several studies in preterm-born adults have shown impaired cognitive performance to be associated with reduced fractional anisotropy (FA) (Allin et al., 2011; Eikenes et al., 2011) or volume (Caldinelli et al., 2017) of specific white matter tracts, mirroring findings in younger cohorts (Skranes et al., 2007; Vollmer et al., 2017). Interestingly, however, several studies have shown that impaired structural white matter integrity may result in neural compensation at the functional level, allowing for comparable task performance to term-born controls (Froudist-Walsh et al., 2015; Salvan et al., 2017). This mechanism speaks in favour of adaptive plastic processes which are achieved in the preterm brain in order to compensate for the early insult. Using network analysis, we have found that despite a relative paucity of white matter resources, preterm adults display a stronger “rich-club” architecture compared to term-born controls, suggesting that in the face of anatomical constraints, global connectivity is prioritised over other, more peripheral, connections (Karolis et al., 2016). A simulated “lesion” approach further suggested that the basal ganglia in particular played an altered role in supporting global connectivity compared to term-born adults (Karolis et al., 2016). In addition, preterm adults from the same cohort showed compromised functional connectivity between striatal aspects of the salience network and the default mode network (White et al., 2014), suggesting that between-network connectivity is critically impaired in these individuals. Further evidence for an altered role played by the basal ganglia comes from a study showing that structural volumetric abnormalities in the striatum are associated with abnormal functional connectivity in a basal ganglia network in preterm-born adults (Bauml et al., 2015).

These observations raise interesting possibilities with respect to psychosis risk. Schizophrenia has frequently been conceptualised as a “disconnection syndrome” (Friston and Frith, 1995), whereby aberrant structural brain connectivity leads to a disintegration of effective mental functioning. It therefore seems plausible that an impairment in white matter integrity rooted in neodevelopmental brain injury may constitute a significant risk factor for psychosis (Bullmore et al., 1997; Kochunov and Hong, 2014). In fact, altered structural connectivity has been detected in infant offspring of women with schizophrenia, suggesting that genetic high risk impacts brain connectivity early on in development (Ahn et al., 2019; Shi et al., 2012). Fronto-striatal connectivity in particular has been implicated in psychosis (Dandash et al., 2017; Robbins, 1990), in line with the notion that psychotic symptoms arise as a result of aberrant integration of top-down cortical and bottom-up subcortical signals, precipitated by aberrant striatal dopamine function (Howes and Kapur, 2009). If, in the preterm brain, overall connectivity indeed relies more heavily on basal ganglia connections than normally expected (Bauml et al., 2015; Karolis et al., 2016), one might hypothesise that any disturbance of the basal ganglia occurring in early adulthood could disproportionately affect global connectivity, potentially mimicking striatal dysregulation typically observed in psychosis (Bullmore et al., 1997). Abnormal striatal dopamine functioning appears to be a key mechanism underlying positive psychotic symptoms (Howes and Murray, 2014). Preterm adults without brain injury were shown to have normal striatal dopamine synthesis capacity, and preterm adults with perinatal brain injury showed reduced dopamine synthesis capacity (Froudist-Walsh et al., 2017). This latter contrasts with the increased dopamine synthesis capacity typically observed in psychosis patients (Howes and Murray, 2014); however, another risk factor associated with schizophrenia, notably heavy cannabis use, has also been associated with decreased striatal dopamine (Bloomfield et al., 2014). One possibility is that this may be associated with supersensitivity of the dopamine D2 receptor (Murray et al., 2014), and that disruption of normal dopamine signalling either presynaptically or postsynaptically may lead to psychosis (Seeman and Seeman, 2014).

It has been suggested that gial cell abnormalities are critically involved in white matter pathology and subsequent symptomatology observed in psychosis and schizophrenia (Dietz et al., 2020). There is evidence for impaired gial progenitor cell differentiation in schizophrenia, resulting in delayed maturation of oligodendrocytes and therefore disrupted myelination in early development (Kerns et al., 2010; Windrem et al., 2017). Impaired progenitor cell differentiation may be precipitated by untimely microglial activation during late fetal development, e.g. as a result of maternal infection, and indeed increased risk of schizophrenia is associated with maternal infection in mid- to late pregnancy (Brown et al., 2004), when oligodendrocyte maturation is most sensitive to microglial activation (Chew et al., 2013). In terms of timing, preterm birth is of particular relevance here. Maternal-fetal inflammatory response and concomitant gial activation is indeed strongly implicated in prematurity (Mallard et al., 2019; Supramaniam et al., 2013) and is known to be associated with subsequent neurodevelopmental disorders such as ASD (Bokobza et al., 2019). Recent multimodal research has leveraged advances in imaging genomics to shed further light on gial cell involvement in altered brain development following preterm birth. Work here has demonstrated that the neurodevelopmental effects of prematurity are temporally and spatially coincident with developmental processes involving cortical gial cell populations (Ball et al., 2020), and that the microglial inflammatory response is implicated in structural white matter changes resulting from brain injury following preterm birth (Krishnan et al., 2017a). Thus, it is possible that neuroinflammation related to preterm birth and subsequent gial pathology could contribute to brain alterations characteristic of later psychotic illness.

On a macrostructural level, there is also evidence for volumetric grey matter abnormalities in preterm adults, with potential relevance for psychosis risk. Alterations of the grey matter are being increasingly recognised as important contributing factors to neurodevelopmental disorder and other psychopathologies following preterm birth (Fleiss et al., 2020). Young adults of the UCLH study showed reduced grey matter volume (GMV) in widespread regions including frontal, temporal, insular, subcortical, and occipital areas (Nosarti et al., 2014). Less extensive increases in GMV were found in medial and anterior frontal regions. Similar findings were also observed when the preterm individuals were adolescents (Nosarti et al., 2008), suggesting that volumetric differences are not a mere result of developmental delay, with individuals “catching up” by adulthood, but rather constitute permanent structural alterations. Studies in other cohorts confirm this, with reduced regional and global grey matter volumes being observed in both adult (Bauml et al., 2015; Meng et al., 2016; Pascoe et al., 2019; Shang et al., 2019) and younger (de Kievi et al., 2012) preterm samples. Perinatal brain injury was furthermore shown to exacerbate the observed structural changes (Nosarti et al., 2014). Abnormal GMV also partially accounted for altered brain activation during a verbal executive function task, once more suggesting processes of functional plasticity compensating for structural deficits (Nosarti et al., 2009).

Meta-analyses of structural brain changes in individuals at high clinical risk of psychosis as well as those experiencing a first episode of psychosis (FEP) provide evidence for reduced GMV in frontal, temporal and insular cortices (Fusar-Poli et al., 2012b; Radua et al., 2012). Subcortical and insular GMV reductions are also seen in those at high familial risk of psychosis (Cooper et al., 2014). The reoccurrence of abnormalities of the insula may be of particular interest here, as this is a major hub of localisation of the transient bursting of spontaneous neuronal events that are critical for brain maturation in preterm infants (aged between 32 and 36 postmenstrual weeks) (Arichi et al., 2017) and as development of insular connections is preferably affected following preterm...
birth, with alterations modulated by genetic factors (Krishnan et al., 2017b). Furthermore, impaired integrity of the anterior insula appears to represent a transdagnostically shared neural substrate of mental illness across psychiatric disorders (Goodkind et al., 2015; McTeague et al., 2017). Thus, evidence again points towards an interaction of genetic liability for psychiatric disorder and perinatal factors including preterm birth in contributing to neurodevelopmental alterations that further increase the risk of experiencing psychopathology such as psychosis.

Reduced volumes of the hippocampus and parahippocampal gyrus have also been associated with clinical (Fusar-Poli et al., 2012b) and genetic (Boos et al., 2007; Fusar-Poli et al., 2014) high risk for psychosis. Schizophrenic patients who have suffered obstetric complications including prematurity were found to be especially likely to show decreased volume of the hippocampus (Stefanis et al., 1999). Reduced hippocampal volume is apparent in preterm-born infants by term-equivalent age (Ball et al., 2013), though findings at later ages are more mixed (Aanes et al., 2015; Fraello et al., 2011; Nosarti and Froudist-Walsh, 2016; Omizzolo et al., 2013). Of note, the hippocampus is known to follow a dynamic and highly heterogeneous maturational trajectory, with both gain and loss of regional volume observed throughout development (Gogtay et al., 2006). In adulthood, very preterm born UCLH study participants showed hippocampal shape changes consistent with atrophy, despite no overall volume difference compared to term-born adults (Cole et al., 2015). Intriguingly, this longitudinal study found that larger hippocampal volume in adolescence was associated with increased delusional ideation in early adulthood. Though initially counterintuitive, this finding is consistent with observations that individuals with an At Risk Mental State (ARMS) for psychosis show increased hippocampal volume before transitioning to psychosis, contrasting with reduced hippocampal volume in first episode psychosis patients (Buehlmann et al., 2010).

Finally, cortical gyriﬁcation (i.e., the folding of the cortical surface), which predominantly occurs in fetal life, is delayed in preterm infants (Dubois et al., 2019; Engelhardt et al., 2015). Recent work demonstrated that altered gyriﬁcation is also evident in preterm adults, and crucially mediates the effect of prematurity on general IQ reduction (Hedderich et al., 2019). These ﬁndings were extended on by work from the UCLH study showing that abnormal gyriﬁcation in preterm adults was related not only to lower IQ but also increased psychopathology (Papini et al., 2020). Strikingly, the pattern of abnormal cortical folding overlapped considerably with that observed in adolescents with a diagnosis of schizophrenia (Palaniyappan and Liddle, 2012) with alterations involving the inferior frontal, insular, and superior temporal cortices.

Taken together, neurobiological abnormalities following preterm birth are widespread and heterogeneous, and are likely to confer a general vulnerability to psychiatric disorder. However, within this heterogeneity, it is possible that certain neurodevelopmental trajectories are indicative of clinical risk for psychosis more speciﬁcally. Moreover, recent evidence supports the notion that genetic vulnerability to psychiatric illness (as measured by polygenic risk scores) interacts with the environmental stress caused by preterm birth to promote neuroanatomical abnormalities of the lentiform nucleus (Cullen et al., 2019) (which together with the caudate forms the striatum); this in turn may further compound the risk of experiencing psychopathology. Therefore, where prematurity coincides with genetic risk for psychosis and results in brain injury, the pre-existing vulnerability is likely to be exacerbated. Ventricular enlargement resulting from perinatal brain injury may be a particular marker of vulnerability to psychosis. In addition, brain tissue abnormalities, especially where they affect fronto-insular-temporal or hippocampal cortex, as well as fronto-striatal connectivity, likely underlie an increased risk for psychosis in preterm individuals. Importantly, many of the brain alterations observed in preterm or psychosis samples also mediate a range of cognitive deﬁcits, which will be discussed in more detail in the following section.

3.2. Cognitive function

Cognitive impairment in psychosis spans multiple domains including executive functioning (Reichenberg and Harvey, 2007), language-related abilities (Condray, 2005), and social cognition (covered in more detail in Section 3.4) (Sheffield et al., 2018). Executive dysfunction in particular is a hallmark of schizophrenia, with deﬁcits thought to be underpinned by neural abnormalities notably of the prefrontal and anterior cingulate cortices (Minzenberg et al., 2009). Importantly, cognitive deﬁcits are already evident in childhood in individuals who go on to develop schizophrenic psychosis (Fusar-Poli et al., 2012a; Jones et al., 1994). It has consequently been suggested that cognitive impairment lies, at least in part, on the causal pathway linking genetic or developmental risk factors with the development of psychosis (Reichenberg, 2005), and is therefore an early indicator of psychosis risk.

Cognitive development in children and adolescents born preterm has also been studied extensively (Johnson, 2007). A recent meta-analysis of cognitive outcomes concluded that very preterm born children exhibit medium to large deﬁcits in general intelligence, executive functioning, and processing speed (Brydges et al., 2018). Many of these deﬁcits are associated with early regionally speciﬁc neuroanatomical changes (Batalle et al., 2018). For example, thalamocortical (Ball et al., 2015) and callosal (Pannek et al., 2020) connectivity assessed at term-equivalent age are predictive of cognitive abilities in toddlers, whereas neonatal microstructural integrity of the arcuate fasciculus predicted language ability at the age of 2 (Salvan et al., 2017). Neonatal volumes of insula and putamen are associated with maths skills aged 5 and 7 (Ullman et al., 2015), and 8-year olds show smaller cortical volumes that are associated with general IQ (Peterson et al., 2000). Several studies have investigated which cognitive deﬁcits persist into adulthood (Breen et al., 2015; De Jong et al., 2012; Eryigit Madzwamuse et al., 2015; Lehagen et al., 2010); the UCLH study in particular has shed light on the neural correlates of cognition in preterm adults (Allin et al., 2011; Kalpakidou et al., 2014; Kontis et al., 2009; Lawrence et al., 2010; Lawrence et al., 2009; Narberhaus et al., 2009; Nosarti et al., 2007; Nosarti et al., 2006; Nosarti et al., 2009).

Preterm born adults tend to score lower on average than term-born controls on IQ tests (Breen et al., 2015; Kroll et al., 2019; Lehagen et al., 2010). However, even accounting for these differences in IQ, preterm adults exhibit further cognitive deﬁcits, particularly in executive functioning. In the ﬁrst broad assessment of executive function in very preterm young adults, our group reported impairments in tasks involving response inhibition and mental ﬂexibility (Nosarti et al., 2007). Impaired executive functioning was furthermore associated with poorer real-life achievements in these adults (Kroll et al., 2017). When investigating the neural correlates of executive functioning in the UCLH cohort, we also found that in spite of good performance on a response inhibition task, preterm adolescents and adults compared to controls displayed altered task-related haemodynamic responses, which may indicate alternative processing strategies in these individuals (Lawrence et al., 2009; Nosarti et al., 2006). This is underlined by ﬁndings from an independent cohort that preterm adults show a haemodynamic response characteristic of predominantly reactive- rather than proactive cognitive control, suggesting alternative neural strategies despite similar task performance (Olsen et al., 2018). Similar observations were made with respect to verbal ﬂuency (Kalpakidou et al., 2014; Nosarti et al., 2009) and verbal and visual associative learning (Lawrence et al., 2010; Narberhaus et al., 2009) tasks, during which preterm adults showed altered brain activation in the absence of performance differences. These ﬁndings in adults are in line with a considerable body of work in preterm children demonstrating engagement of alternative neural circuits, especially in the context of language processing (Barde et al., 2012; Barnes-Davis et al., 2018; Lubsen et al., 2011; Ment et al., 2006). Overall, while cognitive performance in preterm-born adults compared to term-born controls depends on the specific task domain (and difﬁculty) under study, performance does appear to
be associated with neuroanatomical alterations associated with preterm birth. For example, deficits in global executive functioning in preterm adults were found to be mediated by reduced temporal grey matter and callosal white matter volumes (Nosarti et al., 2014). Microstructural changes of the corpus callosum in preterm adults are also associated with lower IQ, verbal learning, and memory performance (Allin et al., 2011; Kontiokari et al., 2009).

Taken together, these findings suggest that neurodevelopmental abnormalities associated with preterm birth necessitate a functional reorganisation of executive processes, but nevertheless can result in persisting cognitive deficits. Some of these deficits and their associated neural changes overlap with those seen in psychosis: for example, impaired inhibitory control is associated with similar patterns of increased midline and attenuated fronto-parieto-cerebellar activation in psychosis (Minzenberg et al., 2009; Vercammen et al., 2012), and abnormal microstructural integrity of the corpus callosum is related to poor executive function in patients with schizophrenia (Oghal et al., 2019).

It is possible that executive function deficits following preterm birth are mediated by attentional difficulties, with inattention constituting the core behavioural difficulty in preterm individuals (Anderson et al., 2011; Elgen et al., 2002). This notion provides a compelling basis for a potential link between the cognitive profile of prematurely born individuals and elevated psychosis risk. Attentional impairments are not only pervasive in patients with psychotic disorder (Hoonakker et al., 2017; Luck and Gold, 2008), but may also constitute a neurodevelopmental marker of vulnerability to psychosis prior to illness onset (Seidman et al., 2016). Children with ADHD are at higher risk of developing schizophrenia in adulthood (Dalsgaard et al., 2014) and the severity of childhood ADHD symptoms retrospectively assessed in first-episode schizophrenia patients is associated with obstetric complications, delay of milestone attainment, and earlier onset of psychiatric symptoms (Peralta et al., 2011). Intriguingly, much of the altered neural activation observed in preterm adults during executive task performance occurs within a fronto-parietal network thought to underpin attention allocation (Lawrence et al., 2009; Nosarti et al., 2006). Corresponding with this, patients with psychosis exhibit altered activation and connectivity of the attentional fronto-parietal network not just during executive task processing (Godwin et al., 2017; Roiser et al., 2013), but even at rest (Chang et al., 2014; Tu et al., 2013). Attentional impairment is suggested to serve as a particularly useful endophenotypic marker of familial risk for psychosis (Cornblatt and Malhotra, 2001), predicting over half of individuals at high-risk for psychosis who develop schizophrenia (Erlenmeyer-Kimling et al., 2000). However, even in the absence of genetic risk for psychosis, it is possible that neurodevelopmental disruptions to networks underpinning attentional processes could have wide-reaching cognitive consequences increasing vulnerability to psychosis. More generally, it has been suggested that early neurodevelopmental insults might deplete cognitive reserves, resulting in a cascade of cognitive impairments that manifest increasingly throughout development as environmental demands grow, and culminate in psychosis (Mollon and Reichenberg, 2018).

In summary, there is overlap both in the cognitive domains and associated neural correlates implicated in psychosis and those affected by prematurity, notably executive functioning and attention. While not all studies show differences in cognitive performance between preterm and term-born adults, there is substantial evidence that cognitive abilities are associated with neural abnormalities in those born preterm, both during development (Ball et al., 2015; Pannek et al., 2020; Peterson et al., 2000; Salvan et al., 2017; Ullman et al., 2015) and in adulthood (Allin et al., 2011; Lawrence et al., 2014; Nam et al., 2015; Nosarti et al., 2014; Olsen et al., 2018). Given these findings, which suggest that alterations in neural systems underpinning cognitive functioning persist well into adulthood of preterm individuals, it is likely that these disruptions constitute significant vulnerability factors that could further interact with genetic or environmental risk for psychotic disorder.

### 3.3. Motor functioning

Even in the absence of obvious disability, neurological impairments are known to be more common in preterm-born individuals than in their term-born peers (Fawke, 2007). Delays in motor development are a key characteristic of preterm infants, with evidence that motor impairments can persist into adulthood (Allin et al., 2006b; de Kieviet et al., 2009).

Young adults of the UCLH study underwent comprehensive neurological assessment aged 18 (Allin et al., 2006b), with preterm-born individuals exhibiting increased motor confusion, alongside impaired sensory integration, compared to term-born controls. These integrative neurological abnormalities were furthermore associated with reduced general intelligence. In a further study, adults from the same cohort performed a movement generation task while undergoing functional imaging (Lawrence et al., 2014). Although many preterm and term-born participants performed near ceiling at this simple task and behavioural differences between groups were not detected, preterm individuals showed increased activation in a cerebellar-cortical network relative to controls. Similarly to several cognitive studies in this cohort, these findings imply recruitment of additional neural resources in order to perform the motor task at comparable levels to term-born subjects. This hyperactivation was correlated with structural grey matter deficits in right premotor cortex (Lawrence et al., 2014), lending further support to the notion of a neural compensatory strategy.

Research in other preterm cohorts also confirm that motor impairments following prematurity are not outgrown by adulthood. Extremely low birth weight survivors showed impaired motor coordination compared to controls from the age of 8 to 36, suggesting that the impairment is stable throughout development (Poole et al., 2015). Both fine and gross motor skills remained impaired from age 14 to 23 in preterm individuals born at very low birth weight (VLBW) (Husby et al., 2013), and these impairments were associated with microstructural abnormalities of interhemispheric (corpus callosum) and motor (corticospinal tract) pathways (Hollund et al., 2018). Motor coordination problems in adults born preterm have also been shown to be associated with reduced cortical surface area (Sripada et al., 2015). Crucially, poorer motor skills are associated with increased levels of psychiatric symptoms and lower quality of life ratings in preterm VLBW adults (Husby et al., 2016), leading to suggestions that manifestations of motor impairments may become more evident as the challenges related to the transition into adulthood increase.

Psychosis is known to be associated with an excess of neurological soft signs including impairments in motor coordination, sequencing, and sensory integration (Dazzan and Murray, 2002). Given the preponderance of motor deficits in first-degree relatives of individuals with schizophrenia, developmental motor symptoms – especially impaired coordination – are thought to represent an endophenotype of schizophrenia indicating disease risk (Burton et al., 2016). Motor deficits in schizophrenia are typically associated with negative and cognitive symptoms (Bomkin et al., 2005), but in individuals at high risk for psychosis, motor dysfunction was also highly related to premorbid positive symptomatology and was furthermore indicative of transition to psychosis (Masucci et al., 2018). Indeed, gross motor skills have been shown to have unusually high sensitivity (75%) in predicting schizophrenia-related psychoses in offspring of patients with schizophrenia (Erlenmeyer-Kimling et al., 2000).

Motor neurological soft signs in psychosis are associated with structural alterations in a cerebello-thalamo-prefrontal network (Mouchet-Mages et al., 2011) which is also implicated in “cognitive dysmetria” (i.e., disruption in the fluid coordination of mental activity) in schizophrenia (Andreasen et al., 1999; Andreasen et al., 1996). Cerebellar and thalamocortical connectivity are known to be affected by prematurity (Ball et al., 2013; Herzenmann et al., 2019), likely playing a role in the emergence of motor dysfunction in preterm infants (Hoon Jr et al., 2009; Messerschmidt et al., 2008). Furthermore structural alterations of the basal ganglia, which are at particular risk of neonatal brain injury
following preterm birth (Logitharajah et al., 2009) are associated with motor abnormalities in first episode psychosis patients (Cuesta et al., 2020). These findings suggest that preterm birth may cause disruption to the neural circuitry crucially involved in early neurological signs associated with psychosis. Early motor difficulties could lead to impaired self-other distinction and development of the embodied self, which are implicated in the genesis of psychotic symptoms by virtue of a failure to distinguish between internally and externally generated sensation (Poletti et al., 2019). Recent evidence of impaired body representation and deficits in sensorimotor representation of self and other generated action in preterm children (Butti et al., 2020; Montiroso et al., 2019) provides a possible mechanistic link between abnormal motor development following preterm birth and vulnerability to psychosis. This is further underlined by evidence that neuromotor abnormalities are significantly associated with obstetric complications in high risk (Marcus et al., 1993) and psychotic (Peralta and Cuesta, 2011) individuals.

3.4. Social functioning

Social difficulties and deficits in social cognition (referring to the mental operations underlying social behaviour) are a key characteristic of the preterm behavioural phenotype (Johnson and Marlow, 2011). Preterm-born children and adolescents are at increased risk of developing autism-spectrum disorders (ASD), whereby the symptomatic presentation is more strongly characterised by social communication problems than repetitive or stereotyped behaviour (Indredavik et al., 2005). Both alterations in brain development (Fischi-Gómez et al., 2015; Healy et al., 2013) and exposure to socio-environmental risk factors (Montagna and Nosarti, 2016) are implicated in adverse childhood social outcomes following preterm birth, and general neurocognitive impairments in functions such as attention and memory are likely to contribute to social cognitive deficits in these individuals (Dean et al., 2021). Socio-emotional problems following preterm birth can already be observed in early childhood, and these problems have been associated with structural and functional brain alterations. For example, disrupted orbitofrontal white matter integrity at term-equivalent age is predictive of socio-emotional difficulties at age 5 (Rogers et al., 2012), and in school-aged children born preterm poorer performance on social tasks is associated with hypoactivation of relevant fronto-parietal circuits (Mossad et al., 2017; Urbain et al., 2019).

In contrast to childhood and adolescence, comparatively little research has focused on social cognition in adults born preterm, but evidence suggests that social difficulties persist into adulthood, with those born preterm reporting poorer social life and fewer social interactions (Kajantie et al., 2008; Lund et al., 2012; Saigal, 2014). Personality assessments in preterm adults also showed that they are characterised by a distinct socially withdrawn personality type (Eryigit-Madzwamuse et al., 2015), lower extraversion and higher neuroticism (Allin et al., 2006a).

Psychotic disorders, too, are characterised by profound problems with social interactions and social cognitive deficits. These difficulties are likely rooted in early development, with children who later develop schizophrenia showing greater social maladjustment than healthy controls (Done et al., 1994). In fact, the magnitude of social cognitive deficits individuals at high clinical risk for psychosis substantially exceeds deficits in other cognitive domains (Fusar-Poli et al., 2012a). Similarly, to other cognitive deficits, attention is suggested to play an important role in the development of social difficulties in psychosis (Cornblatt and Malhotra, 2001), whereby attentional problems may cause an inefficiency to process information from the (social) environment, resulting in a disruption of social competence. This is underscored by observations that attention deficits tend to be more highly correlated with social compared to psychotic symptoms (Green, 1996).

Good social functioning likely acts as a protective factor against psychopathology (McLaughlin et al., 2020; Selten and Cantor-Graae, 2005), thus social difficulties arising as a result of preterm birth may transdiagnostically increase the risk for developing psychiatric disorder in later life. With particular relevance for psychosis, social stress, disadvantage, or isolation (encapsulated in the concept of “social defeat”) has furthermore been suggested to cause sensitisation and/or overactivity of the mesolimbic dopamine system, thus precipitating the emergence of psychotic symptoms more specifically (Gevonden et al., 2014; Selten and Cantor-Graae, 2005). Conversely, the dysfunction of striatal dopamine which we found in adults with perinatal brain injury from the UCHL cohort (Froudast-Walsh et al., 2017) may underpin vulnerability to psychosocial stress (Pyhälä et al., 2009).

In summary, psychosocial stress associated with poor interpersonal skills following preterm birth is highly likely to increase general vulnerability to mental health problems, although the specificity of this for psychosis is less clear. However, social impairments following preterm birth may also be a marker of disrupted development of neural systems not only underpinning social but also general cognitive functioning (such as attentional processes) (Montagna and Nosarti, 2016). Key regions of the social brain are also relevant for cognitive processes implicated in psychosis development, such as temporoparietal junction (Gromann et al., 2013) and medial prefrontal cortex (Ilzarbe et al., 2019). Thus, abnormalities in these regions associated with prematurity may manifest socially during earlier development, but contribute to elevated psychosis risk in the transition to adulthood.

4. Summary and outlook

We have considered the implications of the neurodevelopmental, cognitive, motor, and social sequelae of preterm birth for developing vulnerability to psychosis, with an emphasis on outcomes observed in adulthood. There is remarkably little research explicitly investigating the occurrence of psychotic symptoms following preterm birth. This may be a consequence of the difficulties caused by the long interval between preterm birth and onset of psychosis, but it is also a reflection of a lack of specificity in the link between the two. Indeed research indicates that preterm birth results in a transdiagnostically increased risk for psychiatric disorder in adulthood at the population level (Nosarti et al., 2012; Walshe et al., 2008). However, at the individual level, it is nevertheless important to consider the mechanisms by which adverse outcomes of preterm birth may increase vulnerability specifically to psychosis. Outcomes of preterm birth are highly heterogeneous, placing those exposed onto one of a wide array of possible trajectories. Similarly, psychotic disorders are highly heterogeneous both in presentation and in origin. The aim of this review was to tease out the pathways that could link prematurity to psychosis risk within this diverseness. Key potential mechanisms and markers identified here are depicted in Fig. 1.

Certain aspects of the developmental preterm behavioural phenotype are particularly relevant for potential psychosis risk. Specifically, deficits in attention, social difficulties, and motor impairments, all of which are increased following preterm birth, are known to be predictive of psychosis. Each of these factors has separately been identified as endophenotypes signalling familial risk for psychosis (Burton et al., 2016; Cornblatt and Malhotra, 2001; Tikka et al., 2020). As such, they could be considered simple epiphenomena of underlying genetic vulnerability; however, it is more likely that they lie on the causal pathway towards developing psychotic symptoms and therefore pose risk even in people with relatively low genetic vulnerability. Attentional deficits likely induce greater social and cognitive impairment engendering abnormal information processing; social difficulties increase the likelihood of experiencing psychosocial stress and isolation resulting in social defeat; and motor difficulties could lead to impaired self-other distinction and development of the embodied self, all of which represent outcomes known to be strongly associated with psychotic experiences. Each of these factors warrant more detailed investigations in adult preterm cohorts in conjunction with assessments of psychosocial proneness.
The behavioural vulnerability factors described here are underpinned by neural abnormalities that show overlapping structure in preterm and psychosis patient samples. Neural changes observed in psychosis have long been considered to be at least partly neurodevelopmental in nature (Murray and Lewis, 1987), and many of these alterations could be caused by insults to the brain and/or subsequent altered neurodevelopment as a result of preterm birth, including those resulting from neuroinflammatory glial response. Of course, individuals subject to preterm birth suffer very different impacts on the neonatal brain. Those candidate impacts most likely to increase risk of psychosis include white matter injury resulting in impaired microstructural integrity of fronto-striatal and interhemispheric tracts (Allin et al., 2011; Dandash et al., 2017; Karolis et al., 2016; Ohoshi et al., 2019), reduced grey matter volume in basal-ganglia, fronto-insular, and temporal regions (Cuesta et al., 2020; Fusar-Poli et al., 2012b; Logitharajah et al., 2009; Nosarti et al., 2014), and functional dysconnectivity in fronto-parietal and cerebellono-cortical circuits (Ball et al., 2013; Mouchet-Mages et al., 2011; Narberhaus et al., 2009; Roiser et al., 2013).

Brain alterations resulting from preterm birth may manifest in symptoms such as inattention or autism early on, but then change in the transition to adulthood as the demands of the surrounding world increase, ultimately leading to psychosis. Equally, individuals could appear behaviourally unaffected for the most part (as seen in the frequently normal cognitive task performance in preterm and term-born adults), but the neural compensatory changes necessary for this intact behaviour could signal vulnerability that renders the system less resilient to later stressors and risk factors. Importantly, substantial evidence suggests that there is at least an additive effect between genetic risk for psychosis and obstetric complications (Cannon et al., 1989; Cullen et al., 2019; Falkai et al., 2003), whereby neonatal brain injury as a result of preterm birth likely increases an already existing risk. In addition, social factors of the caregiving environment such as parental distress (Wadhwa et al., 2001) likely poses a greater risk for subsequent psychosis-proneness (due to the established link between inflammation and psychosis, as discussed above) compared to indicated preterm birth due to pre-eclampsia (Davies et al., 2020). Although neonatal morbidity (Bastek et al., 2010) and mortality (Delorme et al., 2016) are known to depend on the subtype of preterm birth, the impact of different causes of preterm birth on long-term outcomes is an area which remains largely unexplored (Crump, 2020). However, the possibility of a genetic confounding of the association between prematurity and psychopathology must be considered, given that women with mental illness are at increased risk of preterm birth (Baer et al., 2016), especially spontaneous preterm birth (Sanchez et al., 2013; Venkatesh et al., 2019). That said, a recent study found little evidence for an association between maternal polygenic risk for schizophrenia and preterm delivery (Leppert et al., 2019), although psychosocial and lifestyle factors associated with psychiatric disorder could undermine the increased risk of giving birth prematurely (Behrman and Butler, 2007). Taken together, it must be acknowledged that an association between preterm birth and psychosis risk could in some cases be at least partly attributed to common underlying risk factors for both, including genetic susceptibility.

Moreover, prematurely born infants are subject to a wide range of different postnatal experiences associated with varying degrees of neonatal illness (and treatment thereof) in neonatal intensive care, such as exposure to pain, need for mechanical ventilation, surgery, or anaesthesia. While it is difficult to separate out potential effects of intensive care strategies from effects of morbidity on later outcomes (Marlow, 2014), many of the factors characterising early hospital care are important risk factors for poor neurodevelopmental outcomes (Kocak et al., 2016; Smith et al., 2011; Xiong et al., 2012). Gaining a better understanding the long-term consequences of neonatal intensive care remains challenging and will be important in order to elucidate potential links between prematurity and psychiatric outcomes, including psychosis.
It is important to remember that a range of factors can increase risk of psychosis and that the pathways between these different factors and psychotic symptoms may be quite different. For example, already it is clear that those individuals who develop psychosis following heavy cannabis use show relatively little in the way of neurodevelopmental or social difficulties in childhood. Future research can further improve our understanding of psychosis by taking into account this heterogeneity. Risk related to preterm birth should be studied firstly by more explicitly assessing the prevalence of psychotic symptoms dimensionally in large developmental and adult preterm cohorts. Here, a better understanding of the long-term outcomes following varying subtypes of preterm birth with different underlying causes will substantially improve further risk stratification. In addition, testing whether symptoms of psychosis are more strongly related to brain structure and function in preterm compared to term-born individuals will shed light on specific pathways to psychosis related to neurodevelopmental disruption. Finally, it will be crucial to investigate the predictive validity of developmental symptoms in conjunction with neural abnormalities for transition to psychosis specifically in preterm cohorts. Overall, this will improve clinicians’ ability to anticipate the type of psychosisopathology preterm-born individuals are most likely to suffer from long-term, and thereby provide more tailored care on an individual level.

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