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Abnormal synaptic pruning during adolescence underlying the development of psychotic disorders

Monique Germann, Sanne G. Brederoo, and Iris E.C. Sommer

Purpose of review
Excessive synaptic pruning has first been suggested by Irwin Feinberg (1982) as an important pillar in the pathophysiology in schizophrenia (SCZ). This article reviews recent developments highlighting factors implicated in aberrant synaptic pruning and its contribution to disease onset and emergence of cognitive symptoms in SCZ. Unraveling these factors provides new insights for potential prevention and treatment strategies for psychotic disorders.

Recent findings
Increased pruning in SCZ was recently confirmed by a positron emission tomography-study employing the novel tracer [11C]UCB-J, demonstrating the consequential loss of synaptic density. Recent evidence supports the contributing role of astrocytes and increased complement-mediated microglial pruning in disease onset and cognitive symptoms in SCZ. Increased microglial pruning is mediated specifically by C4. Furthermore, environmental factors (e.g., infections and stress) can lead to dysbiosis which was recently linked to microglial activation and pruning in SCZ.

Summary
Recent findings render the pruning machinery a potential target for early treatment and prevention in individuals at high risk for SCZ. Minocycline can improve cognition in SCZ, probably by reducing excessive pruning. Probiotics might also have beneficial effects on cognition, although recent findings are not encouraging. N-acetyl-cysteine recovers functional connectivity in SCZ both in vitro and in vivo, making it an interesting candidate.

Keywords
astrocytes, cognitive symptoms, microglia, schizophrenia, synaptic pruning

INTRODUCTION
Schizophrenia (SCZ) and related psychotic disorders are heterogeneous psychiatric disorders that are estimated to affect ~20 million people worldwide, placing a significant burden on individual well-being and global health [1]. In addition to positive and negative symptoms, disabling cognitive symptoms are present in a subpopulation of SCZ patients. Cognitive clustering shows three groups: one that is cognitively intact, one with moderate cognitive deficits in only one or a few domains, and one with severe cognitive dysfunction [2]. The latter group shows global deficits, affecting numerous cognitive domains ranging from executive functioning to working memory, planning, verbal fluency, and problem solving [3]. Whereas positive symptoms can be effectively mitigated by antipsychotic medication in the majority of patients, treatments alleviating cognitive symptoms are scarce [4,5]. Given the, as of yet, largely elusive pathophysiology underlying cognitive impairments in SCZ, a better understanding of the mechanisms behind this symptom is essential to aid the development of effective treatments and preventive strategies.

One of the most prominent hypotheses unraveling the pathophysiology of SCZ stems from the eighties. In 1982, Feinberg postulated that a crucial
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KEY POINTS

- Feinberg’s hypothesis (1982) has recently been supported in a PET-study demonstrating a decrease in synaptic density in SCZ.
- As extensive synaptic pruning peaks during late adolescence and early adulthood, this critical developmental period presents itself as most suitable for preventive interventions to protect cerebral connectivity and cognition in young individuals at high risk for SCZ.
- As the complement component system, microglia, astrocytes, and the microbiome are key players implicated in synaptic pruning in SCZ, they form primary targets for preventive interventions.

step in the development of SCZ is a faulty programmed synaptic pruning process [6]. Synaptic pruning is a healthy neurodevelopmental process important for the proper establishment and maturation of functional neural networks by eliminating infrequently used synapses whereas maintaining frequently used connections. Whereas pruning occurs throughout life, certain critical developmental periods are characterized by a peak in pruning. Particularly during adolescence, brain regions involved in higher cognitive functions have been found to be subject to extensive pruning [7*]. According to Feinberg, SCZ may arise due to a faulty programmed pruning process, resulting in excessive synaptic pruning during adolescence [6].

Recently, Feinberg’s hypothesis was supported in a PET-study employing the novel tracer [11C]UCB-J to image synaptic vesicle glycoprotein 2A (SV2A). This study demonstrated a significant and large decrease (Cohen’s $d = 0.8$) in synaptic density in vivo in 18 SCZ patients compared to 18 healthy controls (HC) [8*]. Such a reduction in synaptic density has a profound impact on cortical connectivity [7*]. In line with this, a recent meta-analysis across 14 studies (743 individuals with ultra-high risk [UHR] for psychosis, 588 HC) found abnormalities in gray matter volumes (an indirect measure of synaptic density) in UHR individuals. Compared to HC, UHR individuals showed increased gray matter volumes in the median cingulate, right fusiform gyrus, left superior temporal gyrus, and right thalamus ($P < 0.001$ in all studies) and decreased volumes in the superior frontal gyrus bilaterally ($P < 0.001$ in all studies) [10]. The loss of synaptic density may impair cognitive functioning by affecting certain neuronal circuitry, which would be reflected in an abnormal functional connectivity (FC). Indeed, Xian-Bin et al. (2019) recently assessed resting-state FC in 24 individuals with clinical high risk for psychosis (CHR), 19 patients with first-episode psychosis (FEP), and 47 HC. Results of this investigation revealed hypoconnectivities in CHR and FEP patients compared to HC between posterior insula and somatosensory areas (Effect sizes HC vs. FEP: 0.96–1.26; Effect sizes HC vs. CHR: 0.82–1.06) and between dorsal anterior insula and putamen (Effect sizes HC vs. FEP: 1.07–1.25; Effect sizes HC vs. CHR: 0.87–0.95) [11].

Although these recent studies support Feinberg’s theory in showing decreased synaptic density and aberrant structural and FC in SCZ, the question remains which factors contribute to the emergence of aberrant synaptic pruning in SCZ. To advance our understanding of the pathophysiological machinery, the current review focuses on the latest scientific developments with respect to factors contributing to excessive synaptic pruning in SCZ. An overarching concept providing a well-established connection between genetic and environmental factors implicated in pruning is given by the notion of an immune system dysregulation in SCZ, with a crucial role assigned to abnormal microglial activation linked to increased complement system activation.

MICROGLIAL SYNAPTIC PRUNING

Microglia are resident macrophages of the CNS and play a crucial role in its immunity. Main functions of microglia entail phagocytosis of pathogens and disrupted or apoptotic neuronal parts such as infrequently used synapses [12]. The traditionally demarcated term ‘microglial activation’ nowadays comprises the highly diverse functional and morphological microglial reactions to triggers such as stress, trauma, neuroinflammation, or neurodegeneration [13]. Persistent overactivation of microglia can be induced by increased abundance of activated complement system proteins like C3 and C4 [14] and is associated with increased pro-inflammatory cytokine release (e.g., TNF-a, IL-1b, and IL-33) [15,16]. Moreover, prolonged microglial overactivation mediated by increased complement activation may cause excessive synaptic pruning [14,15]. Support for this comes from a recent meta-analysis reporting significantly higher microglial density and increased levels of pro-inflammatory cytokine proteins in postmortem brain samples of SCZ patients compared to HC [17].

In several lines of research, Stevens et al. impressively demonstrated the crucial role of microglial and the complement system in synaptic pruning during neurodevelopment. In one of their recent studies, the group revealed microglial pruning in hippocampal neurons to be phosphatidylserine (PS-) and TREM2-dependent in mice. PS has been found
to be a potential molecular component that tags certain neuronal elements, thereby signaling subsequent microglial engulfment. Interestingly, the involvement of PS in microglia-mediated pruning has been demonstrated to be highest during critical developmental periods characterized by a peak in microglial pruning. Finally, with their finding that C1q peaks during developmental pruning periods and the observation of decreased microglial engulfment of PS-marked elements in C1q knock-out mice, the group further highlights the role of complement during developmental microglial pruning [18]. Recent findings implicate dysfunctional complement system as an important driver of abnormal microglial activation [19**].

### The complement system as mediator of microglial synaptic pruning

The classical complement cascade is an integral part of the innate immune system. By opsonizing pathogens or abnormal cell parts and releasing inflammatory mediators, complement proteins direct microglial or abnormal cell parts and releasing inflammatory mediators. The complement system as mediator of microglial activation [19**] is fundamental for proper growth factor b (TGF-b) which enhances the expression of the complement cascade initiating C1q protein, ultimately resulting in the release of C3. Subsequent microglial activation leads to the release of C1q, IL-1a, and TNF-a, which results in reactive astrocytes with a diminished ability to promote synapse formation [20*]. Astrocytes may also use another phagocytic pathway, via receptors Multiple EGF-like domains 10 (MEG10) and MER Tyrosine Kinase (MERTK), which is independent of C1q. Both pathways mediate synaptic pruning during development [26]. Support comes from findings of excess synapses in MEGF10 and MERTK deficient mice, reflecting missing circuit refinement [26].

### The emerging role of astrocytes in synaptic pruning

In addition to complement-mediated microglial pruning, astrocytes have recently been implicated in pruning [20*,25]. Astrocytes can produce cytokine interleukin (IL)-33, which directly enhances the phagocytic action of microglial cells. Interestingly, synaptic maturation has been reported to be accompanied by an increase in IL-33. This may reflect a homeostatic loop in which increased numbers of synapses constitute a signal for astrocytes to produce IL-33 for the induction of microglial pruning [25]. Astrocytes can also release transforming growth factor b (TGF-b) which enhances the expression of the complement cascade initiating C1q protein, ultimately resulting in the release of C3. Subsequent microglial activation leads to the release of C1q, IL-1a, and TNF-a, which results in reactive astrocytes with a diminished ability to promote synapse formation [20*]. Astrocytes may also use another phagocytic pathway, via receptors Multiple EGF-like domains 10 (MEG10) and MER Tyrosine Kinase (MERTK), which is independent of C1q. Both pathways mediate synaptic pruning during development [26]. Support comes from findings of excess synapses in MEGF10 and MERTK deficient mice, reflecting missing circuit refinement [26].

### Environmental factors implicated in synaptic pruning

Although the above-mentioned description focuses on the role of astrocytes and complement-mediated microglial pruning, certain environmental factors are known to affect synaptic pruning as well. These factors can be interpreted in light of the two-hit model of SCZ. This theory posits that an initial...
perinatal-immune activation in response to genetic and/or environmental alterations induces a first hit, and subsequent stressors or psychological trauma in childhood or adolescence poses a second hit [14,27]. Perinatal-immune activation can be influenced by a genetic liability that predisposes the individual for an overexpression of complement component proteins and pro-inflammatory cytokines [15,21,28,29]. Together, these factors can prime microglia which, in response to later stressful events, become overactivated, inducing excessive opsonization of synapses, leading to pathological pruning.

Maternal immune activation (MIA), perinatal insults, and early-life stress form environmental factors that can prime microglia [15,30]. MIA produces an increase in pro-inflammatory cytokines and MMP-9 release in the fetal body, both of which are inducers of microglial activity [28]. Prenatal infections and MIA are furthermore associated with microglial overactivation in children [31]. Interestingly, prenatal infections have recently been reported to cause astrocytes to react in a hypersensitive manner to stimuli in the future [32]. Furthermore, MIA and perinatal stress can lead to elevated microglial density and activity [15]. Although these perinatal immune activation factors prime microglia, a shift toward prolonged microglial overactivation phenotype takes place only after exposure to stress or traumatic stimuli during adolescence, a time which coincides with the onset of first clinical symptoms in SCZ [15]. Indeed, evidence points toward subtle, yet significant increases in pro-inflammatory cytokines in adults with childhood trauma, which itself has been associated with increased risk to develop SCZ [14]. Finally, cannabis usage during early adolescence may disturb neuro-developmental processes as it affects microglial function by causing an upregulation of CB2 receptors, thereby modifying synaptic pruning [33]. The effect of cannabis usage on brain connectivity has recently been demonstrated in a sample of 54 psychotic patients (29 patients who use cannabis, 25 patients who did not use cannabis) and 38 HC (16 HC who use cannabis, 22 HC who did not use cannabis). Results revealed increased connectivity in the dorsal attention network ($P = 0.019$) and visual dorsal attention internetwork ($P = 0.036$) in cannabis consuming patients compared to nonusing controls. Interestingly, however, there was no significant difference between patients with cannabis usage and patients without cannabis usage in terms of connectivity [34]. Given that the current state of literature on the effect of cannabis consumption on connectivity in SCZ patients points toward mixed results, further research is warranted.

The gut–brain axis allows for bidirectional communication between the gut and brain via pathways involving the immune system, autonomic nervous system, endocrine, and enteric nervous system [35]. The brain can influence the microbiota of the gut via these pathways. Conversely, the intestinal microbiota can affect cognition by influencing levels of short-chain fatty acids (SCFA), serotonin, GABA and other neurotransmitters [36]. Interestingly, the critical developmental period for cortical maturation characterized by a peak in synaptic pruning coincides with the time of maturation of the intestinal microbiota [37*], pointing toward a role of the microbiome in microglial activation [38]. Early life environmental factors such as cesarian section, antibiotic treatment, stress or infections can lead to dysbiosis, which can affect early brain maturation [35,39]. In germ-free mice, absence of gut microbiota negatively affects cognition, stress tolerance and social behavior by altering the regulation [39] and expression of genes implicated in synapse organization in microglia [40]. In humans, there is also evidence that microglial function and activation can be modified by the intestinal microbiota, especially during the critical developmental time [37*].

**CLINICAL IMPLICATIONS**

Converging lines of evidence point toward an overactive pruning machinery in SCZ as a potent target for preventive and therapeutic actions. Minocycline is a tetracyclic antibiotic with anti-inflammatory properties and has been proven to be safe, well-tolerated and effective in SCZ patients [41*,42], as a recent randomized double-blind, placebo-controlled study demonstrated that the administration of minocycline to 75 SCZ patients led to significantly reduced cognitive deficits and levels of IL-6 and IL-1b compared to placebo [43]. Further support comes from a meta-analysis on randomized placebo-controlled trials in SCZ patients investigating the effect of adjuvant anti-inflammatory agents on distinct clinical outcome variables. Based on two trials (133 SCZ patients), they concluded that minocycline significantly improved cognitive functioning in SCZ patients compared to placebo, specifically with regard to executive functioning (Hedges’ g: first trial = 1.35; second trial = 0.17) and visual learning/memory (effect size 0.94). Furthermore, no significant differences in side effects between minocycline and placebo have been found [44]. A potential explanation for this positive effect of minocycline is that it reduces extensive pruning, as shown in cell lines by Sellgren [19*]. Alternatively, minocycline may primarily affect the microbiome, which could lead
to alterations in SCFA and GABA production, also known to affect cognition [45].

Manipulation of the microbial environment by the administration of probiotics (beneficiary microbiota bacteria) or prebiotics (nutritional fibers that serve as food for these bacteria) may affect brain function and improve cognition [35,36]. A recent meta-analysis investigated the effect of probiotics (11 studies, 724 participants), prebiotics (5 studies, 355 participants), and fermented food (6 studies, 472 participants) in a mixed group of HC and patients with a wide range of brain and somatic disorders on cognition. Pooled results did not demonstrate beneficial effects of probiotics, prebiotics or fermented food on cognitive outcome measures. Authors of this meta-analysis suggested that the absence of significant findings for pooled estimates may be due to high clinical heterogeneity, which ranged from fibromyalgia, to cirrhosis, to Alzheimer, depression and more [46]. Findings of another meta-analysis assessing 7 controlled clinical trials in a mixed sample of patients with Alzheimer, fibromyalgia, depression, minimal hepatic encephalopathy, and HC, and 11 animal studies did support the beneficial effect of probiotics on cognitive functioning in both human and animals. Interestingly, the beneficial effect of probiotics was more pronounced in cognitively impaired individuals compared to HC [47]. Overall, mixed results of the effects of probiotics and the absence of investigations of the effect of pre or probiotics on cognition in SCZ patients warrant further research.

Finally, to prevent the loss of brain connectivity by increased pruning, the administration of N-acetyl-cysteine (NAC) supplementation may be beneficial. Mullier et al. [48] compared the effect of 6 months NAC supplementation to placebo on FC between cingulate cortex areas in 20 early psychotic patients and 74 HC. This pilot study demonstrated that, compared to placebo, supplementation of NAC in early psychotic individuals increased the FC between designated areas implicated in positive symptoms and processing speed [48]. However, replication studies are warranted.

Based on the peak of excessive pruning in late adolescence and early adulthood, it may prove most beneficial to administer minocycline, NAC, or possibly pro/prebiotics to prevent over-pruning, early on in adolescent individuals who are at high risk to develop SCZ. Even though the peak in synaptic pruning occurs during adolescence, it is a process part of brain homeostasis taking place throughout life. Therefore, interventions with minocycline may still be useful after adolescence. It is encouraging that these three interventions are all well tolerated with few if any side-effects [44]. People who could potentially benefit from such interventions are those with first degree family with SCZ and polygenic risk scores involving the complement system, as has been elegantly demonstrated by Sellgren et al. in vitro [19**].

CONCLUSION

The present review outlines recent developments in the elucidation of the pathophysiological mechanism underlying aberrant synaptic pruning in SCZ, and alterations succeeding it. Genetically overexpressed complement proteins C3 and C4 contribute to a generally enhanced pro-inflammatory state of both periphery and CNS. Together, this results in microglial overactivation for a prolonged period of time, thereby inducing excessive synaptic pruning. The resulting loss of synaptic density has recently been confirmed by a PET-study employing a novel tracer [11C]UCB-J. Astrocytes contribute to excessive pruning through their ability to directly enhance microglial phagocytic actions. The ability of the intestinal microbiota to modify microglial function and activation implicate the human microbiome as a modifiable factor in pruning as well. Minocycline has been demonstrated to be safe and beneficial in improving cognitive functioning in SCZ. Although individual studies reported minocycline and NAC to improve cognitive functioning, further research is warranted to replicate their unambiguous beneficial effects on cognition.

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Conflicts of interest

I.S. is a consultant to Gabather, received research support from Janssen Pharmaceuticals Inc. and Sunovion Pharmaceuticals Inc. The rest of the authors do not have any conflicts of interest.

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Papers of particular interest, published within the annual period of review, have been highlighted as:

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