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Permalink

<https://escholarship.org/uc/item/4th7t646>

Journal

Schizophrenia bulletin, 46(Suppl 1)

ISSN

1787-9965

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Publication Date

2020-05-01

Peer reviewed

O5.6. ADVANCED DIFFUSION IMAGING IN PSYCHOSIS RISK: A CROSS-SECTIONAL AND LONGITUDINAL STUDY OF WHITE MATTER DEVELOPMENT

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Background: Studies in individuals at clinical high risk (CHR) for psychosis provide a powerful means to predict outcomes and inform putative mechanisms underlying conversion to psychosis. In previous work, we applied advanced diffusion imaging methods to reveal that white matter pathology in a CHR population is characterized by cellular-specific changes in white matter, suggesting a preexisting neurodevelopmental anomaly. However, it remains unknown whether these deficits relate to clinical symptoms and/or conversion to frank psychosis. To address this gap, we examined cross-sectional and longitudinal white matter maturation in the largest imaging population of CHR individuals to date, obtained from the North American Prodrome Longitudinal Study (NAPLS-3).

Methods: Multi-shell diffusion magnetic resonance imaging (MRI) data were collected across multiple timepoints (1–6 at ~2 month intervals) in 286 subjects (age range=12–32 years). These were 230 unmedicated CHR subjects, including 11% (n=25) who transitioned to psychosis (CHR-converters), as well as 56 age and sex-matched healthy controls. Raw diffusion signals were harmonized to remove scanner/site-induced effects, yielding a unified imaging dataset. Fractional anisotropy of cellular tissue (FA_t) and the volume fraction of extracellular free-water (FW) were assessed in 12 major tracts from the IIT Human Brain Atlas (v.5.0). Linear mixed effects (LME) models were fitted to infer developmental trajectories of FA_t and FW across age for CHR-converters, CHR-nonconverters and control groups, while accounting for the repeated measurements on each individual.

Results: The rate at which FA_t changed with age significantly differed between the three groups across commissural and association tracts (5 in total; p<0.05). In these tracts, FA_t increased with age in controls (0.002% change per year) and in CHR-nonconverters, albeit at a slower rate (0.00074% per year). In contrast, FA_t declined with age in CHR-converters at a rate that was significantly faster (-3.944% per year) than the rate of increase in the other two groups. By 25 years of age, FA_t was significantly lower in both CHR groups compared to controls (p<0.05). With regard to FW, the rate of change significantly differed between CHR-converters and controls across the forceps major and the left inferior longitudinal and fronto-occipital fasciculi (IFOF; 3 tracts in total; p<0.05). This was due to increased FW with age in the CHR-converters (0.0024% change per year) relative to controls (-0.0002% per year). Consequently, FW was significantly higher in CHR-converters compared to controls by 20 years of age (p<0.05). With regard to symptoms, there was a significant impact of IFOF FW on positive symptom severity across CHR subjects, regardless of conversion status (t=2.37, p<0.05).

Discussion: Our results revealed that clinical high-risk for psychosis is associated with cellular-specific alterations in white matter, regardless of conversion status. Only converters showed excess extracellular free-water,

which involved tracts connecting occipital, posterior temporal, and orbito-frontal areas. We also demonstrate a direct impact of free-water on positive symptomatology, collectively, suggesting that excess free-water may signal acute psychosis and its onset. This marker may be useful for patient selection for clinical trials and assessment of individuals with prodromal psychosis.

O6. Oral Sessions: Cognitive/ Other

O6.1. COGNITIVE CONTROL DIFFERENCES BETWEEN TREATMENT RESPONSIVE AND TREATMENT RESISTANT FIRST-EPISODE PSYCHOSIS PATIENTS

Abstract not included.

O6.2. A NETWORK FOR COGNITIVE, MOTOR, AND PSYCHOPATHOLOGICAL ALTERATIONS IN SCHIZOPHRENIA SPECTRUM DISORDERS

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Background: Schizophrenia spectrum disorders are complex syndromes involving multiple clinical manifestations. Besides psychopathological symptoms, cognitive and motor alterations are also highly relevant in the context of the comprehension, assessment, and treatment of these disorders. Moreover, these three domains of clinical manifestations display complex reciprocal interactions that require further characterization. This work aims to use network analysis to investigate the associations between cognitive, motor, and psychopathological alterations in schizophrenia spectrum disorders. This approach might prove to be advantageous in identifying key variables for the assessment and treatment of these disorders.

Methods: A sample of 732 patients with schizophrenia spectrum disorders from a multi-site cohort study was included in the analysis. We estimated a network using a regularized Gaussian Graphical Model and conducted network stability analyses. Twenty-six nodes were included, encompassing items from the Positive and Negative Syndrome Scale, multiple neuropsychological tests, and clinician-assessed extrapyramidal symptoms' scores. The results were further explored with centrality analyses and network comparisons between subgroups defined according to illness duration and remission status.

Results: We found that the estimated network was densely interconnected. Furthermore, nodes representing symptoms of disorganization were very central and, therefore, pivotal in connecting other psychopathological symptoms to cognitive and motor alterations. The estimated network for the subgroup of patients in remission showed a more sparse density and a different structure from the network of non-remitted patients.

Discussion: In conclusion, in the context of a broader representation of schizophrenia spectrum disorders' manifestations, our results of a network analysis confirm a close association between different symptom domains and unveil a highly influential role of disorganization symptoms. Moreover, structural differences in networks occur according to remission status. These results are relevant for research in nosology, clinical assessment, and treatment approaches.